

L'Ecuyer, Pierre

Working Paper

Random number generation

Papers, No. 2004,21

Provided in Cooperation with:

CASE - Center for Applied Statistics and Economics, Humboldt University Berlin

Suggested Citation: L'Ecuyer, Pierre (2004) : Random number generation, Papers, No. 2004,21, Humboldt-Universität zu Berlin, Center for Applied Statistics and Economics (CASE), Berlin

This Version is available at:

<https://hdl.handle.net/10419/22195>

Standard-Nutzungsbedingungen:

Die Dokumente auf EconStor dürfen zu eigenen wissenschaftlichen Zwecken und zum Privatgebrauch gespeichert und kopiert werden.

Sie dürfen die Dokumente nicht für öffentliche oder kommerzielle Zwecke vervielfältigen, öffentlich ausstellen, öffentlich zugänglich machen, vertreiben oder anderweitig nutzen.

Sofern die Verfasser die Dokumente unter Open-Content-Lizenzen (insbesondere CC-Lizenzen) zur Verfügung gestellt haben sollten, gelten abweichend von diesen Nutzungsbedingungen die in der dort genannten Lizenz gewährten Nutzungsrechte.

Terms of use:

Documents in EconStor may be saved and copied for your personal and scholarly purposes.

You are not to copy documents for public or commercial purposes, to exhibit the documents publicly, to make them publicly available on the internet, or to distribute or otherwise use the documents in public.

If the documents have been made available under an Open Content Licence (especially Creative Commons Licences), you may exercise further usage rights as specified in the indicated licence.

Random Number Generation

Pierre L'Ecuyer¹

Département d'Informatique et de Recherche Opérationnelle, Université de
Montréal, C.P. 6128, Succ. Centre-Ville, Montréal (Québec), H9S 5B8, Canada.
<http://www.iro.umontreal.ca/~lecuyer>

1 Introduction

The fields of probability and statistics are built over the abstract concepts of probability space and random variable. This has given rise to elegant and powerful mathematical theory, but exact implementation of these concepts on conventional computers seems impossible. In practice, random variables and other random objects are *simulated* by *deterministic algorithms*. The purpose of these algorithms is to produce sequences of numbers or objects whose behavior is very hard to distinguish from that of their “truly random” counterparts, at least for the application of interest. Key requirements may differ depending on the context. For Monte Carlo methods, the main goal is to reproduce the statistical properties on which these methods are based, so that the Monte Carlo estimators behave as expected, whereas for gambling machines and cryptology, observing the sequence of output values for some time should provide no practical advantage for predicting the forthcoming numbers better than by just guessing at random.

In computational statistics, random variate generation is usually made in two steps: (1) generating imitations of independent and identically distributed (i.i.d.) random variables having the uniform distribution over the interval $(0, 1)$ and (2) applying transformations to these i.i.d. $U(0, 1)$ random variates in order to generate (or imitate) random variates and random vectors from arbitrary distributions. These two steps are essentially independent and the world's best experts on them are two different groups of scientists, with little overlap. The expression *(pseudo)random number generator* (RNG) usually refers to an algorithm used for step (1).

In principle, the simplest way of generating a random variate X with distribution function F from a $U(0, 1)$ random variate U is to apply the inverse of F to U :

$$X = F^{-1}(U) \stackrel{\text{def}}{=} \min\{x \mid F(x) \geq U\}. \quad (1)$$

This is the *inversion* method. It is easily seen that X has the desired distribution: $P[X \leq x] = P[F^{-1}(U) \leq x] = P[U \leq F(x)] = F(x)$. Other methods are sometimes preferable when F^{-1} is too difficult or expensive to compute, as will be seen later.

The remainder of this chapter is organized as follows. In the next section, we give a definition and the main requirements of a uniform RNG. Generators based on linear recurrences modulo a large integer m , their lattice structure and quality criteria, and their implementation, are covered in Sect. 3. In Sect. 4, we have a similar discussion for RNGs based on linear recurrences modulo 2. Nonlinear RNGs are briefly presented in Sect. 5. In Sect. 6, we discuss empirical statistical testing of RNGs and give some examples. Sect. 7 contains a few pointers to recommended RNGs and software. In Sect. 8, we cover non-uniform random variate generators. We first discuss inversion and its implementation in various settings. We then explain the alias, rejection, ratio-of-uniform, composition, and convolution methods, and provide pointers to the several other methods that apply in special cases.

Important basic references that we recommend are Knuth (1998); L'Ecuyer (1994, 1998); Niederreiter (1992), and Tezuka (1995) for uniform RNGs, and Devroye (1986); Gentle (2003), and Hörmann et al. (2004) for non-uniform RNGs.

2 Uniform Random Number Generators

2.1 Physical Devices

Random numbers can be generated via physical mechanisms such as the timing between successive events in atomic decay, thermal noise in semiconductors, and the like. A key issue when constructing a RNG based on a physical device is that a “random” or “chaotic” output does not suffice; the numbers produced must be, at least to a good approximation, realizations of *independent and uniformly distributed* random variables. If the device generates a stream of bits, which is typical, then each bit should be 0 or 1 with equal probability, and be independent of all the other bits. In general, this cannot be proved, so one must rely on the results of empirical statistical testing to get convinced that the output values have the desired statistical behavior. Not all these devices are reliable, but some apparently are. I did test two of them recently and they passed all statistical tests that I tried.

For computational statistics, physical devices have several disadvantages compared to a good algorithmic RNG that stands in a few lines of code. For example, (a) they are much more cumbersome to install and run; (b) they are more costly; (c) they are slower; (d) they cannot reproduce exactly the same sequence twice. Item (d) is important in several contexts, including program verification and debugging as well as comparison of similar systems

by simulation with common random numbers to reduce the variance (Bratley et al., 1987; Fishman, 1996; Law and Kelton, 2000). Nevertheless, these physical RNGs can be useful for selecting the seed of an algorithmic RNG, more particularly for applications in cryptology and for gaming machines, where frequent reseeding of the RNG with an external source of entropy (or randomness) is important. A good algorithmic RNG whose seed is selected at random can be viewed as an extensor of randomness, stretching a short random seed into a long sequence of *pseudorandom* numbers.

2.2 Generators Based on a Deterministic Recurrence

RNGs used for simulation and other statistical applications are almost always based on deterministic algorithms that fit the following framework, taken from L'Ecuyer (1994): a RNG is a structure $(\mathcal{S}, \mu, f, \mathcal{U}, g)$ where $\mathcal{S}$ is a finite set of *states* (the *state space*), μ is a probability distribution on $\mathcal{S}$ used to select the *initial state* (or *seed*) s_0 , $f : \mathcal{S} \rightarrow \mathcal{S}$ is the *transition function*, $\mathcal{U}$ is the *output space*, and $g : \mathcal{S} \rightarrow \mathcal{U}$ is the *output function*. Usually, $\mathcal{U} = (0, 1)$, and we shall assume henceforth that this is the case. The state of the RNG evolves according to the recurrence $s_i = f(s_{i-1})$, for $i \geq 1$, and the *output* at step i is $u_i = g(s_i) \in \mathcal{U}$. The output values $u_0, u_1, u_2, \dots$ are the so-called *random numbers* produced by the RNG.

Because $\mathcal{S}$ is finite, there must be some finite $l \geq 0$ and $j > 0$ such that $s_{l+j} = s_l$. Then, for all $i \geq l$, one has $s_{i+j} = s_i$ and $u_{i+j} = u_i$, because both f and g are deterministic. That is, the state and output sequences are eventually periodic. The smallest positive j for which this happens is called the *period length* of the RNG, and is denoted by ρ . When $l = 0$, the sequence is said to be *purely periodic*. Obviously, $\rho \leq |\mathcal{S}|$, the cardinality of $\mathcal{S}$. If the state has a k -bit representation on the computer, then $\rho \leq 2^k$. Good RNGs are designed so that their period length ρ is not far from that upper bound. In general, the value of ρ may depend on the seed s_0 , but good RNGs are normally designed so that the period length is the same for all admissible seeds.

In practical implementations, it is important that the output be *strictly* between 0 and 1, because $F^{-1}(U)$ is often infinite when U is 0 or 1. All good implementations take care of that. However, for the mathematical analysis of RNGs, we often assume that the output space is $[0, 1)$ (i.e., 0 is admissible), because this simplifies the analysis considerably without making much difference in the mathematical structure of the generator.

2.3 Quality Criteria

What important quality criteria should we consider when designing RNGs? An extremely *long period* is obviously essential, to make sure that no wrap-around over the cycle can occur in practice. The length of the period must be guaranteed by a mathematical proof. The RNG must also be *efficient* (run fast and use only a small amount of memory), *repeatable* (able to reproduce

exactly the same sequence as many times as we want), and *portable* (work the same way in different software/hardware environments). The availability of efficient *jump-ahead* methods that can quickly compute $s_{i+\nu}$ given s_i , for any large ν and any i , is also very useful, because it permits one to partition the RNG sequence into long disjoint *streams* and *substreams* of random numbers, in order to create an arbitrary number of *virtual generators* from a single RNG (Law and Kelton, 2000; L'Ecuyer et al., 2002a). These virtual generators can be used on parallel processors or to support different sources of randomness in a large simulation model, for example.

Consider a RNG with state space $\mathcal{S} = \{1, \dots, 2^{1000} - 1\}$, transition function $s_{i+1} = f(s_i) = (s_i + 1) \bmod 2^{1000}$, and $u_i = g(s_i) = s_i / 2^{1000}$. This RNG has period length 2^{1000} and enjoys all the nice properties described in the preceding paragraph, but is far from imitating “randomness.” In other words, these properties are *not sufficient*.

A sequence of real-valued random variables $u_0, u_1, u_2, \dots$ are i.i.d. $U(0, 1)$ if and only if for every integers $i \geq 0$ and $t > 0$, the vector $\mathbf{u}_{i,t} = (u_i, \dots, u_{i+t-1})$ is uniformly distributed over the t -dimensional unit hypercube $(0, 1)^t$. Of course, this cannot hold for algorithmic RNGs because any vector of t successive values produced by the generator must belong to the *finite set*

$$\Psi_t = \{(u_0, \dots, u_{t-1}) : s_0 \in \mathcal{S}\},$$

which is the set of all vectors of t successive output values, from all possible initial states. Here we interpret Ψ_t as a *multiset*, which means that the vectors are counted as many times as they appear, and the cardinality of Ψ_t is exactly equal to that of $\mathcal{S}$.

Suppose we select the seed s_0 at random, uniformly over $\mathcal{S}$. This can be approximated by using some physical device, for example. Then, the vector $\mathbf{u}_{0,t}$ has the uniform distribution over the finite set Ψ_t . And if the sequence is purely periodic for all s_0 , $\mathbf{u}_{i,t} = (u_i, \dots, u_{i+t-1})$ is also uniformly distributed over Ψ_t for all $i \geq 0$. Since the goal is to approximate the uniform distribution over $(0, 1)^t$, it immediately becomes apparent that Ψ_t should be evenly spread over this unit hypercube. In other words, Ψ_t *approximates* $(0, 1)^t$ as the *sample space* from which the vectors of successive output values are drawn randomly, so it must be a good approximation of $(0, 1)^t$ in some sense. The design of good-quality RNGs must therefore involve practical ways of measuring the uniformity of the corresponding sets Ψ_t even when they have huge cardinalities. In fact, a large state space $\mathcal{S}$ is necessary to obtain a long period, but an even more important reason for having a huge number of states is to make sure that Ψ_t can be large enough to provide a good uniform coverage of the unit hypercube, at least for moderate values of t .

More generally, we may also want to measure the uniformity of sets of the form

$$\Psi_I = \{(u_{i_1}, \dots, u_{i_t}) \mid s_0 \in \mathcal{S}\},$$

where $I = \{i_1, \dots, i_t\}$ is a fixed set of non-negative integers such that $0 \leq i_1 < \dots < i_t$. As a special case, we recover $\Psi_t = \Psi_I$ when $I = \{0, \dots, t-1\}$.

The uniformity of a set Ψ_I is typically assessed by measuring the *discrepancy* between the empirical distribution of its points and the uniform distribution over $(0, 1)^t$ (Niederreiter, 1992; Hellekalek and Larcher, 1998; L'Ecuyer and Lemieux, 2002). Discrepancy measures are equivalent to goodness-of-fit test statistics for the multivariate uniform distribution. They can be defined in many different ways. In fact, the choice of a specific definition typically depends on the mathematical structure of the RNG to be studied and the reason for this is very pragmatic: we must be able to compute these measures quickly even when $\mathcal{S}$ has very large cardinality. This obviously excludes any method that requires explicit generation of the sequence over its entire period. The selected discrepancy measure is usually computed for each set I in some predefined class $\mathcal{J}$, these values are weighted or normalized by factors that depend on I , and the worst-case (or average) over $\mathcal{J}$ is adopted as a *figure of merit* used to rank RNGs. The choice of $\mathcal{J}$ and of the weights are arbitrary. Typically, $\mathcal{J}$ would contain sets I such that t and $i_t - i_1$ are rather small. Examples of such figures of merit will be given when we discuss specific classes of RNGs.

2.4 Statistical Testing

Good RNGs are designed based on mathematical analysis of their properties, then implemented and submitted to batteries of *empirical statistical tests*. These tests try to detect empirical evidence against the null hypothesis $\mathcal{H}_0$: “the u_i are realizations of i.i.d. $U(0, 1)$ random variables.” A test can be defined by any function T that maps a sequence $u_0, u_1, \dots$ in $(0, 1)$ to a real number X , and for which a good approximation is available for the distribution of the random variable X under $\mathcal{H}_0$. For the test to be implementable, X must depend on only a finite (but perhaps random) number of u_i 's. Passing many tests may improve one's confidence in the RNG, but never guarantees that the RNG is foolproof for all kinds of simulations.

Building a RNG that *passes all statistical tests* is an impossible dream. Consider, for example, the class of all tests that examine the first (most significant) b bits of n successive output values, $u_0, \dots, u_{n-1}$, and return a binary value $X \in \{0, 1\}$. Select $\alpha \in (0, 1)$ so that αb^n is an integer and let $\mathcal{T}_{n,b,\alpha}$ be the tests in this class that return $X = 1$ for *exactly* αb^n of the b^n possible output sequences. We may say that the sequence *fails* the test when $X = 1$. The number of tests in $\mathcal{T}_{n,b,\alpha}$ is equal to the number of ways of choosing αb^n distinct objects among b^n . The chosen objects are the sequences that fail the test. Now, for any given output sequence, the number of such tests that return 1 for this particular sequence is equal to the number of ways of choosing the other $\alpha b^n - 1$ sequences that also fail the test. This is the number of ways of choosing $\alpha b^n - 1$ distinct objects among $b^n - 1$. In other words, as pointed out by Leeb (1995), every output sequence fails exactly the same number of tests! This result should not be surprising. Viewed from a different angle, it is essentially a restatement of the well-known fact that under $\mathcal{H}_0$, each of the b^n

possible sequences has the same probability of occurring, so one could argue that none should be considered more random than any other (Knuth, 1998).

This viewpoint seems to lead into a dead end. For statistical testing to be meaningful, all tests should not be considered on equal footing. So which ones are more important? Any answer is certainly tainted with its share of arbitrariness. However, for large values of n , the number of tests is huge and all but a tiny fraction are too complicated even to be implemented. So we may say that *bad* RNGs are those that fail simple tests, whereas *good* RNGs fail only complicated tests that are hard to find and run. This common-sense compromise has been generally adopted in one way or another.

Experience shows that RNGs with very long periods, good structure of their set Ψ_t , and based on recurrences that are not too simplistic, pass most reasonable tests, whereas RNGs with short periods or bad structures are usually easy to crack by standard statistical tests. For sensitive applications, it is a good idea, when this is possible, to apply additional statistical tests designed in close relation with the random variable of interest (e.g., based on a *simplification* of the stochastic model being simulated, and for which the theoretical distribution can be computed).

Our discussion of statistical tests continues in Sect. 6.

2.5 Cryptographically Strong Generators

One way of defining an ideal RNG would be that no statistical test can distinguish its output sequence from an i.i.d. $U(0, 1)$ sequence. If an unlimited computing time is available, no finite-state RNG can satisfy this requirement, because by running it long enough one can eventually figure out its periodicity. But what if we impose a limit on the computing time? This can be analyzed formally in the framework of asymptotic *computational complexity* theory, under the familiar “rough-cut” assumption that polynomial-time algorithms are practical and others are not.

Consider a family of RNGs $\{\mathcal{G}_k = (\mathcal{S}_k, \mu_k, f_k, \mathcal{U}_k, g_k), k = 1, 2, \dots\}$ where $\mathcal{S}_k$ of cardinality 2^k (i.e., $\mathcal{G}_k$ has a k -bit state). Suppose that the transition and output functions f and g can be computed in time bounded by a polynomial in k . Let $\mathcal{T}$ be the class of statistical tests that run in time bounded by a polynomial in k and try to differentiate between the output sequence of the RNG and an i.i.d. $U(0, 1)$ sequence. The RNG family is called *polynomial-time perfect* if there is a constant $\epsilon > 0$ such that for all k , no test in $\mathcal{T}$ can differentiate correctly with probability larger than $1/2 + e^{-k\epsilon}$. This is equivalent to asking that no polynomial-time algorithm can predict any given bit of u_i with probability of success larger than $1/2 + e^{-k\epsilon}$, after observing $u_0, \dots, u_{i-1}$. This links unpredictability with statistical uniformity and independence. For the proofs and additional details, see, e.g. Blum et al. (1986); L'Ecuyer and Proulx (1989); Lagarias (1993), and Luby (1996). This theoretical framework has been used to define a notion of reliable RNG in the context of cryptography. But the guarantee is only asymptotic; it does not necessarily tell what

value of k is large enough for the RNG to be secure in practice. Moreover, specific RNG families have been proved to be polynomial-time perfect only under yet unproven conjectures. So far, no one has been able to prove even their existence. Most RNGs discussed in the remainder of this chapter are known *not* to be polynomial-time perfect. However, they are fast, convenient, and have good enough statistical properties when their parameters are chosen carefully.

3 Linear Recurrences Modulo m

3.1 The Multiple Recursive Generator

The most widely used RNGs are based on the linear recurrence

$$x_i = (a_1x_{i-1} + \cdots + a_kx_{i-k}) \bmod m, \quad (2)$$

where m and k are positive integers called the *modulus* and the *order*, and the *coefficients* $a_1, \dots, a_k$ are in $\mathbb{Z}_m$, interpreted as the set $\{0, \dots, m-1\}$ on which all operations are performed with reduction modulo m . The *state* at step i is $s_i = \mathbf{x}_i = (x_{i-k+1}, \dots, x_i)^\top$. When m is a prime number, the finite ring $\mathbb{Z}_m$ is a finite field and it is possible to choose the coefficients a_j so that the period length reaches $\rho = m^k - 1$ (the largest possible value) (Knuth, 1998). This maximal period length is achieved if and only if the characteristic polynomial of the recurrence, $P(z) = z^k - a_1z^{k-1} - \cdots - a_k$, is a primitive polynomial over $\mathbb{Z}_m$, i.e., if and only if the smallest positive integer ν such that $(z^\nu \bmod P(z)) \bmod m = 1$ is $\nu = m^k - 1$. Knuth (1998) explains how to verify this for a given $P(z)$. For $k > 1$, for $P(z)$ to be a primitive polynomial, it is necessary that a_k and at least another coefficient a_j be nonzero. Finding primitive polynomials of this form is generally easy and they yield the simplified recurrence:

$$x_n = (a_r x_{n-r} + a_k x_{n-k}) \bmod m. \quad (3)$$

A *multiple recursive generator* (MRG) uses (2) with a large value of m and defines the output as $u_i = x_i/m$. For $k = 1$, this is the classical *linear congruential generator* (LCG). In practice, the output function is modified slightly to make sure that u_i never takes the value 0 or 1 (e.g., one may define $u_i = (x_i + 1)/(m + 1)$, or $u_i = x_i/(m + 1)$ if $x_i > 0$ and $u_i = m/(m + 1)$ otherwise) but to simplify the theoretical analysis, we will follow the common convention of assuming that $u_i = x_i/m$ (in which case u_i *does* take the value 0 occasionally).

3.2 The Lattice Structure

Let $\mathbf{e}_i$ denote the i th unit vector in k dimensions, with a 1 in position i and 0's elsewhere. Denote by $x_{i,0}, x_{i,1}, x_{i,2}, \dots$ the values of $x_0, x_1, x_2, \dots$ produced by

the recurrence (2) when the initial state $\mathbf{x}_0$ is $\mathbf{e}_i$. An arbitrary initial state $\mathbf{x}_0 = (z_1, \dots, z_k)^\top$ can be written as the linear combination $z_1 \mathbf{e}_1 + \dots + z_k \mathbf{e}_k$ and the corresponding sequence is a linear combination of the sequences $(x_{i,0}, x_{i,1}, \dots)$, with reduction of the coordinates modulo m . Reciprocally, any such linear combination reduced modulo m is a sequence that can be obtained from some initial state $\mathbf{x}_0 \in \mathcal{S} = \mathbb{Z}_m^k$. If we divide everything by m we find that for the MRG, for each $t \geq 1$, $\Psi_t = L_t \cap [0, 1)^t$ where

$$L_t = \left\{ \mathbf{v} = \sum_{i=1}^t z_i \mathbf{v}_i \mid z_i \in \mathbb{Z} \right\},$$

is a t -dimensional *lattice* in $\mathbb{R}^t$, with basis

$$\begin{aligned} \mathbf{v}_1 &= (1, 0, \dots, 0, x_{1,k}, \dots, x_{1,t-1})^\top / m \\ &\vdots \\ \mathbf{v}_k &= (0, 0, \dots, 1, x_{k,k}, \dots, x_{k,t-1})^\top / m \\ \mathbf{v}_{k+1} &= (0, 0, \dots, 0, 1, \dots, 0)^\top \\ &\vdots \\ \mathbf{v}_t &= (0, 0, \dots, 0, 0, \dots, 1)^\top. \end{aligned}$$

For $t \leq k$, L_t contains all vectors whose coordinates are multiples of $1/m$. For $t > k$, it contains a fraction m^{k-t} of those vectors.

This lattice structure implies that the points of Ψ_t are distributed according to a very regular pattern, in equidistant parallel hyperplanes. Graphical illustrations of this, usually for LCGs, can be found in a myriad of papers and books; e.g., Gentle (2003); Knuth (1998); Law and Kelton (2000), and L'Ecuyer (1998). Define the *dual lattice* to L_t as

$$L_t^* = \{ \mathbf{h} \in \mathbb{R}^t : \mathbf{h}^\top \mathbf{v} \in \mathbb{Z} \text{ for all } \mathbf{v} \in L_t \}.$$

Each $\mathbf{h} \in L_t^*$ is a normal vector that defines a family of equidistant parallel hyperplanes, at distance $1/\|\mathbf{h}\|_2$ apart, and these hyperplanes cover all the points of L_t unless $\mathbf{h}$ is an integer multiple of some other vector $\mathbf{h}' \in L_t^*$. Therefore, if ℓ_t is the euclidean length of a shortest non-zero vector $\mathbf{h}$ in L_t^* , then there is a family of hyperplanes at distance $1/\ell_t$ apart that cover all the points of L_t . A small ℓ_t means thick slices of empty space between the hyperplanes and we want to avoid that. A large ℓ_t means a better (more uniform) coverage of the unit hypercube by the point set Ψ_t . Computing the value of $1/\ell_t$ is often called the *spectral test* (Knuth, 1998; Fishman, 1996).

The lattice property holds as well for the point sets Ψ_I formed by values at arbitrary lags defined by a fixed set of indices $I = \{i_1, \dots, i_t\}$. One has $\Psi_I = L_I \cap [0, 1)^t$ for some lattice L_I , and the largest distance between successive hyperplanes for a family of hyperplanes that cover all the points of L_I is $1/\ell_I$,

where ℓ_I is the euclidean length of a shortest nonzero vector in L_I^* , the dual lattice to L_I .

The lattice L_I and its dual can be constructed as explained in Couture and L'Ecuyer (1996) and L'Ecuyer and Couture (1997). Finding the shortest nonzero vector in a lattice with basis $\mathbf{v}_1, \dots, \mathbf{v}_t$ can be formulated as an integer programming problem with a quadratic objective function:

$$\text{Minimize } \|\mathbf{v}\|^2 = \sum_{i=1}^t \sum_{j=1}^t z_i \mathbf{v}_i^\top \mathbf{v}_j z_j$$

subject to $z_1, \dots, z_t$ integers and not all zero. This problem can be solved by a branch-and-bound algorithm (Fincke and Pohst, 1985; L'Ecuyer and Couture, 1997; Tezuka, 1995).

For any given dimension t and m^k points per unit of volume, there is an absolute upper bound on the best possible value of ℓ_I (Conway and Sloane, 1999; Knuth, 1998; L'Ecuyer, 1999b). Let $\ell_t^*(m^k)$ denote such an upper bound. To define a figure of merit that takes into account several sets I , in different numbers of dimensions, it is common practice to divide ℓ_I by an upper bound, in order to obtain a standardized value between 0 and 1, and then take the worst case over a given class $\mathcal{J}$ of sets I . This gives a figure of merit of the form

$$M_{\mathcal{J}} = \min_{I \in \mathcal{J}} \ell_I / \ell_{|I|}^*(m^k).$$

A value of $M_{\mathcal{J}}$ too close to zero means that L_I has a bad lattice structure for at least one of the selected sets I . We want a value as close to 1 as possible. Computer searches for good MRGs with respect to this criterion have been reported by L'Ecuyer et al. (1993); L'Ecuyer and Andres (1997); L'Ecuyer (1999a), for example. In most cases, $\mathcal{J}$ was simply the sets of the form $I = \{1, \dots, t\}$ for $t \leq t_1$, where t_1 was an arbitrary integer ranging from 8 to 45. L'Ecuyer and Lemieux (2000) also consider the small dimensional sets I with indices not too far apart. They suggest taking $\mathcal{J} = \{\{0, 1, \dots, i\} : i < t_1\} \cup \{\{i_1, i_2\} : 0 = i_1 < i_2 < t_2\} \cup \dots \cup \{\{i_1, \dots, i_d\} : 0 = i_1 < \dots < i_d < t_d\}$ for some positive integers $d, t_1, \dots, t_d$. We could also take a weighted average instead of the minimum in the definition of $M_{\mathcal{J}}$.

An important observation is that for $t > k$, the t -dimensional vector $\mathbf{h} = (-1, a_1, \dots, a_k, 0, \dots, 0)^\top$ always belong to L_t^* , because for any vector $\mathbf{v} \in L_t$, the first $k+1$ coordinates of $m\mathbf{v}$ must satisfy the recurrence (2), which implies that $(-1, a_1, \dots, a_k, 0, \dots, 0)\mathbf{v}$ must be an integer. Therefore, one always has $\ell_t^2 \leq 1 + a_1^2 + \dots + a_k^2$. Likewise, if I contains 0 and all indices j such that $a_{k-j} \neq 0$, then $\ell_I^2 \leq 1 + a_1^2 + \dots + a_k^2$ (L'Ecuyer, 1997). This means that the sum of squares of the coefficients a_j *must be large* if we want to have any chance that the lattice structure be good.

Constructing MRGs with only two nonzero coefficients and taking these coefficients small has been a very popular idea, because this makes the implementation easier and faster (Deng and Lin, 2000; Knuth, 1998). However,

MRGs thus obtained have a bad structure. As a worst-case illustration, consider the widely-available additive or subtractive *lagged-Fibonacci* generator, based on the recurrence (2) where the two coefficients a_r and a_k are both equal to ± 1 . In this case, whenever I contains $\{0, k-r, k\}$, one has $\ell_I^2 \leq 3$, so the distance between the hyperplanes is at least $1/\sqrt{3}$. In particular, for $I = \{0, k-r, k\}$, all the points of Ψ_I (aside from the zero vector) are contained in only two planes! This type of structure can have a dramatic effect on certain simulation problems and is a good reason for staying away from these lagged-Fibonacci generators, regardless of their parameters.

A similar problem occurs for the “fast MRG” proposed by Deng and Lin (2000), based on the recurrence

$$x_i = (-x_{i-1} + ax_{i-k}) \bmod m = ((m-1)x_{i-1} + ax_{i-k}) \bmod m,$$

with $a^2 < m$. If a is small, the bound $\ell_I^2 \leq 1+a^2$ implies a bad lattice structure for $I = \{0, k-1, k\}$. A more detailed analysis by L'Ecuyer and Touzin (2004) shows that this type of generator cannot have a good lattice structure even if the condition $a^2 < m$ is removed. Another special case proposed by Deng and Xu (2003) has the form

$$x_i = a(x_{i-j_2} + \cdots + x_{i-j_t}) \bmod m. \quad (4)$$

In this case, for $I = \{0, k-j_{t-1}, \dots, k-j_2, k\}$, the vectors $(1, a, \dots, a)$ and $(a^*, 1, \dots, 1)$ both belong to the dual lattice L_I^* , where a^* is the multiplicative inverse of a modulo m . So neither a nor a^* should be small.

To get around this structural problem when I contains certain sets of indices, Lüscher (1994) and Knuth (1998) recommend to skip some of the output values in order to break up the bad vectors. For the lagged-Fibonacci generator, for example, one can output k successive values produced by the recurrence, then skip the next d values, output the next k , skip the next d , and so on. A large value of d (e.g., $d = 5k$ or more) may get rid of the bad structure, but slows down the generator. See Wegenkittl and Matsumoto (1999) for further discussion.

3.3 MRG Implementation Techniques

The modulus m is often taken as a large prime number close to the largest integer directly representable on the computer (e.g., equal or near $2^{31} - 1$ for 32-bit computers). Since each x_{i-j} can be as large as $m-1$, one must be careful in computing the right side of (2) because the product $a_j x_{i-j}$ is typically not representable as an ordinary integer. Various techniques for computing this product modulo m are discussed and compared by Fishman (1996); L'Ecuyer and Tezuka (1991); L'Ecuyer (1999a), and L'Ecuyer and Simard (1999). Note that if $a_j = m - a'_j > 0$, using a_j is equivalent to using the negative coefficient $-a'_j$, which is sometimes more convenient from the implementation viewpoint. In what follows, we assume that a_j can be either positive or negative.

One approach is to perform the arithmetic modulo m in 64-bit (double precision) floating-point arithmetic (L'Ecuyer, 1999a). Under this representation, assuming that the usual IEEE floating-point standard is respected, all positive integers up to 2^{53} are represented exactly. Then, if each coefficient a_j is selected to satisfy $|a_j|(m-1) \leq 2^{53}$, the product $|a_j|x_{i-j}$ will always be represented exactly and $z_j = |a_j|x_{i-j} \bmod m$ can be computed by the instructions

$$y = |a_j|x_{i-j}; \quad z_j = y - m \lfloor y/m \rfloor.$$

Similarly, if $(|a_1| + \dots + |a_k|)(m-1) \leq 2^{53}$, $a_1x_{i-1} + \dots + a_kx_{i-k}$ will always be represented exactly.

A second technique, called *approximate factoring* (L'Ecuyer and Côté, 1991), uses only the integer representation and works under the condition that $|a_j| = i$ or $|a_j| = \lfloor m/i \rfloor$ for some integer $i < \sqrt{m}$. One precomputes $q_j = \lfloor m/|a_j| \rfloor$ and $r_j = m \bmod |a_j|$. Then, $z_j = |a_j|x_{i-j} \bmod m$ can be computed by

$$\begin{aligned} y &= \lfloor x_{i-j}/q_j \rfloor; & z &= |a_j|(x_{i-j} - yq_j) - yr_j; \\ \text{if } z < 0 & \text{ then } z_j = z + m \text{ else } z_j = z. \end{aligned}$$

All quantities involved in these computations are integers between $-m$ and m , so no overflow can occur if m can be represented as an ordinary integer (e.g., $m < 2^{31}$ on a 32-bit computer).

The *powers-of-two decomposition* approach selects coefficients a_j that can be written as a sum or difference of a small number of powers of 2 (Wu, 1997; L'Ecuyer and Simard, 1999; L'Ecuyer and Touzin, 2000). For example, one may take $a_j = \pm 2^q \pm 2^r$ and $m = 2^e - h$ for some positive integers q , r , e , and h . To compute $y = 2^q x \bmod m$, decompose $x = z_0 + 2^{e-q} z_1$ (where $z_0 = x \bmod 2^{e-q}$) and observe that

$$y = 2^q(z_0 + 2^{e-q} z_1) \bmod (2^e - h) = (2^q z_0 + h z_1) \bmod (2^e - h).$$

Suppose now that

$$h < 2^q \quad \text{and} \quad h(2^q - (h+1)2^{-e+q}) < m. \quad (5)$$

Then, $2^q z_0 < m$ and $h z_1 < m$, so y can be computed by shifts, masks, additions, subtractions, and a single multiplication by h . Intermediate results never exceed $2m - 1$. Things simplify further if $q = 0$ or $q = 1$ or $h = 1$. For $h = 1$, y is obtained simply by swapping the blocks of bits z_0 and z_1 (Wu, 1997). It has been pointed out by L'Ecuyer and Simard (1999) that LCGs with parameters of the form $m = 2^e - 1$ and $a = \pm 2^q \pm 2^r$ have bad statistical properties because the recurrence does not “mix the bits” well enough. However, good and fast MRGs can be obtained via the power-of-two decomposition method, as explained in L'Ecuyer and Touzin (2000).

Another interesting idea for improving efficiency is to take all nonzero coefficients a_j equal to the same constant a (Marsaglia, 1996; Deng and Xu,

2003). Then, computing the right side of (2) requires a single multiplication. Deng and Xu (2003) provide specific parameter sets and concrete implementations for MRGs of this type, for prime m near 2^{31} , and $k = 102, 120$, and 1511 .

One may be tempted to take m equal to a power of two, say $m = 2^e$, because then the “mod m ” operation is much easier: it suffices to keep the e least significant bits and mask-out all others. However, taking a power-of-two modulus is not recommended because it has several strong disadvantages in terms of the quality of the RNG (L'Ecuyer, 1990, 1998). In particular, the least significant bits have very short periodicity and the period length of the recurrence (2) cannot exceed $(2^k - 1)2^{e-1}$ if $k > 1$, and 2^{e-2} if $k = 1$ and $e \geq 4$. The maximal period length achievable with $k = 7$ and $m = 2^{31}$, for example, is more than 2^{180} times smaller than the maximal period length achievable with $k = 7$ and $m = 2^{31} - 1$ (a prime number).

3.4 Combined MRGs and LCGs

The conditions that make MRG implementations run faster (e.g., only two nonzero coefficients both close to zero) are generally in conflict with those required for having a good lattice structure and statistical robustness. *Combined MRGs* are one solution to this problem. Consider J distinct MRGs evolving in parallel, based on the recurrences

$$x_{j,i} = (a_{j,1}x_{j,i-1} + \cdots + a_{j,k}x_{j,i-k}) \bmod m_j \quad (6)$$

where $a_{j,k} \neq 0$, for $j = 1, \dots, J$. Let $\delta_1, \dots, \delta_J$ be arbitrary integers,

$$z_i = (\delta_1 x_{1,i} + \cdots + \delta_J x_{J,i}) \bmod m_1, \quad u_i = z_i / m_1, \quad (7)$$

and

$$w_i = (\delta_1 x_{1,i} / m_1 + \cdots + \delta_J x_{J,i} / m_J) \bmod 1. \quad (8)$$

This defines two RNGs, with output sequences $\{u_i, i \geq 0\}$ and $\{w_i, i \geq 0\}$.

Suppose that the m_j are pairwise relatively prime, that δ_j and m_j have no common factor for each j , and that each recurrence (6) is purely periodic with period length ρ_j . Let $m = m_1 \cdots m_J$ and let ρ be the least common multiple of $\rho_1, \dots, \rho_J$. Under these conditions, the following results have been proved by L'Ecuyer and Tezuka (1991) and L'Ecuyer (1996a): (a) the sequence (8) is exactly equivalent to the output sequence of a MRG with (composite) modulus m and coefficients a_j that can be computed explicitly as explained in L'Ecuyer (1996a); (b) the two sequences in (7) and (8) have period length ρ ; and (c) if both sequences have the same initial state, then $u_i = w_i + \epsilon_i$ where $\max_{i \geq 0} |\epsilon_i|$ can be bounded explicitly by a constant ϵ which is very small when the m_j are close to each other.

Thus, these combined MRGs can be viewed as practical ways of implementing an MRG with a large m and several large nonzero coefficients. The

idea is to cleverly select the components so that: (1) each one is easy to implement efficiently (e.g., has only two small nonzero coefficients) and (2) the MRG that corresponds to the combination has a good lattice structure. If each m_j is prime and if each component j has maximal period length $\rho_j = m_j^k - 1$, then each ρ_j is even and ρ cannot exceed $\rho_1 \cdots \rho_J / 2^{J-1}$. Tables of good parameters for combined MRGs of different sizes that reach this upper bound are given in L'Ecuyer (1999a) and L'Ecuyer and Touzin (2000), together with C implementations.

3.5 Jumping Ahead

The recurrence (2) can be written in matrix form as

$$\mathbf{x}_i = \mathbf{A}\mathbf{x}_{i-1} \bmod m = \begin{pmatrix} 0 & 1 & \cdots & 0 \\ \vdots & & \ddots & \vdots \\ 0 & 0 & \cdots & 1 \\ a_k & a_{k-1} & \cdots & a_1 \end{pmatrix} \mathbf{x}_{i-1} \bmod m.$$

To jump ahead directly from $\mathbf{x}_i$ to $\mathbf{x}_{i+\nu}$, for an arbitrary integer ν , it suffices to exploit the relationship

$$\mathbf{x}_{i+\nu} = \mathbf{A}^\nu \mathbf{x}_i \bmod m = (\mathbf{A}^\nu \bmod m) \mathbf{x}_i \bmod m.$$

If this is to be done several times for the same ν , the matrix $\mathbf{A}^\nu \bmod m$ can be precomputed once for all. For a large ν , this can be done in $O(\log_2 \nu)$ matrix multiplications via a standard divide-and-conquer algorithm (Knuth, 1998):

$$\mathbf{A}^\nu \bmod m = \begin{cases} (\mathbf{A}^{\nu/2} \bmod m)(\mathbf{A}^{\nu/2} \bmod m) \bmod m & \text{if } \nu \text{ is even;} \\ \mathbf{A}(\mathbf{A}^{\nu-1} \bmod m) \bmod m & \text{if } \nu \text{ is odd.} \end{cases}$$

3.6 Linear Recurrences With Carry

These types of recurrences were introduced by Marsaglia and Zaman (1991) to obtain a large period even when m is a power of two (in which case the implementation may be faster). They were studied and generalized by Tezuka et al. (1994); Couture and L'Ecuyer (1994, 1997), and Goresky and Klapper (2003). The basic idea is to add a *carry* to the linear recurrence (2). The general form of this RNG, called *multiply-with-carry* (MWC), can be written as

$$\begin{aligned} x_i &= (a_1 x_{i-1} + \cdots + a_k x_{i-k} + c_{i-1})d \bmod b, \\ c_i &= \lfloor (a_0 x_i + a_1 x_{i-1} + \cdots + a_k x_{i-k} + c_{i-1})/b \rfloor, \\ u_i &= \sum_{\ell=1}^{\infty} x_{i+\ell-1} b^{-\ell}, \end{aligned}$$

where b is a positive integer (e.g., a power of two), $a_0, \dots, a_k$ are arbitrary integers such that a_0 is relatively prime to b , and d is the multiplicative inverse of $-a_0$ modulo b . The state at step i is $s_i = (x_{i-k+1}, \dots, x_i, c_i)^\top$. In practice, the sum in (9) is truncated to a few terms (it could be a single term if b is large), but the theoretical analysis is much easier for the infinite sum.

Define $m = \sum_{\ell=0}^k a_\ell b^\ell$ and let a be the inverse of b in arithmetic modulo m , assuming for now that $m > 0$. A major result proved in Tezuka et al. (1994); Couture and L'Ecuyer (1997), and Goresky and Klapper (2003) is that if the initial states agree, the output sequence $\{u_i, i \geq 0\}$ is exactly the same as that produced by the LCG with modulus m and multiplier a . Therefore, the MWC can be seen as a clever way of implementing a LCG with very large modulus. It has been shown by Couture and L'Ecuyer (1997) that the value of ℓ_t for this LCG satisfies $\ell_t^2 \leq a_0^2 + \dots + a_k^2$ for $t \geq k$, which means that the lattice structure will be bad unless the sum of squares of coefficients a_j is large.

In the original proposals of Marsaglia and Zaman (1991), called *add-with-carry* and *subtract-with-borrow*, one has $-a_0 = \pm a_r = \pm a_k = 1$ for some $r < k$ and the other coefficients a_j are zero, so $\ell_t^2 \leq 3$ for $t \geq k$ and the generator has essentially the same structural defect as the additive lagged-Fibonacci generator. In the version studied by Couture and L'Ecuyer (1997), it was assumed that $-a_0 = d = 1$. Then, the period length cannot exceed $(m-1)/2$ if b is a power of two. A concrete implementation was given in that paper. Goresky and Klapper (2003) pointed out that the maximal period length of $\rho = m-1$ can be achieved by allowing a more general a_0 . They provided specific parameters that give a maximal period for b ranging from 2^{21} to 2^{35} and ρ up to approximately 2^{2521} .

4 Generators Based on Recurrences Modulo 2

4.1 A General Framework

It seems natural to exploit the fact that computers work in binary arithmetic and to design RNGs defined directly in terms of bit strings and sequences. We do this under the following framework, taken from L'Ecuyer and Panneton (2002). Let $\mathbb{F}_2$ denote the finite field with two elements, 0 and 1, in which the operations are equivalent to addition and multiplication modulo 2. Consider the RNG defined by a matrix linear recurrence over $\mathbb{F}_2$, as follows:

$$\mathbf{x}_i = \mathbf{A}\mathbf{x}_{i-1}, \quad (9)$$

$$\mathbf{y}_i = \mathbf{B}\mathbf{x}_i, \quad (10)$$

$$u_i = \sum_{\ell=1}^w y_{i,\ell-1} 2^{-\ell} = .y_{i,0} y_{i,1} y_{i,2} \dots, \quad (11)$$

where $\mathbf{x}_i = (x_{i,0}, \dots, x_{i,k-1})^\top \in \mathbb{F}_2^k$ is the k -bit *state vector* at step i , $\mathbf{y}_i = (y_{i,0}, \dots, y_{i,w-1})^\top \in \mathbb{F}_2^w$ is the w -bit *output vector* at step i , k and w are positive integers, $\mathbf{A}$ is a $k \times k$ *transition matrix* with elements in $\mathbb{F}_2$, $\mathbf{B}$ is a $w \times k$ *output transformation matrix* with elements in $\mathbb{F}_2$, and $u_i \in [0, 1)$ is the *output* at step i . All operations in (9) and (10) are performed in $\mathbb{F}_2$.

It is well-known (Niederreiter, 1992; L'Ecuyer, 1994) that when the $\mathbf{x}_i$'s obey (9), for each j , the sequence $\{x_{i,j}, i \geq 0\}$ follows the linear recurrence

$$x_{i,j} = (\alpha_1 x_{i-1,j} + \dots + \alpha_k x_{i-k,j}) \bmod 2, \quad (12)$$

whose *characteristic polynomial* $P(z)$ is the characteristic polynomial of $\mathbf{A}$, i.e.,

$$P(z) = \det(\mathbf{A} - z\mathbf{I}) = z^k - \alpha_1 z^{k-1} - \dots - \alpha_{k-1} z - \alpha_k,$$

where $\mathbf{I}$ is the identity matrix and each α_j is in $\mathbb{F}_2$. The sequences $\{y_{i,j}, i \geq 0\}$, for $0 \leq j < w$, also obey the same recurrence (although some of them may follow recurrences of shorter order as well in certain situations, depending on $\mathbf{B}$). We assume that $\alpha_k = 1$, so that the recurrence (12) has *order* k and is purely periodic. Its period length is $2^k - 1$ (i.e., maximal) if and only if $P(z)$ is a primitive polynomial over $\mathbb{F}_2$ (Niederreiter, 1992; Knuth, 1998).

To jump ahead directly from $\mathbf{x}_i$ to $\mathbf{x}_{i+\nu}$ with this type of generator, it suffices to precompute the matrix $\mathbf{A}^\nu$ (in $\mathbb{F}_2$) and then multiply $\mathbf{x}_i$ by this matrix.

Several popular classes of RNGs fit this framework as special cases, by appropriate choices of the matrices $\mathbf{A}$ and $\mathbf{B}$. This includes the Tausworthe or LFSR, polynomial LCG, GFSR, twisted GFSR, Mersenne twister, multiple recursive matrix generators, and combinations of these (L'Ecuyer and Panneton, 2002; Matsumoto and Nishimura, 1998; Niederreiter, 1995; Tezuka, 1995). We detail some of them after discussing measures of uniformity.

4.2 Measures of Uniformity

The uniformity of point sets Ψ_I produced by RNGs based on linear recurrences over $\mathbb{F}_2$ is usually assessed by measures of equidistribution defined as follows (L'Ecuyer, 1996b; L'Ecuyer and Panneton, 2002; L'Ecuyer, 2004; Tezuka, 1995). For an arbitrary vector $\mathbf{q} = (q_1, \dots, q_t)$ of non-negative integers, partition the unit hypercube $[0, 1)^t$ into 2^{q_j} intervals of the same length along axis j , for each j . This determines a partition of $[0, 1)^t$ into $2^{q_1 + \dots + q_t}$ rectangular boxes of the same size and shape. We call this partition the $\mathbf{q}$ -equidissection of the unit hypercube.

For some index set $I = \{i_1, \dots, i_t\}$, if Ψ_I has 2^k points, we say that Ψ_I is **$\mathbf{q}$ -equidistributed in base 2** if there are exactly 2^q points in each box of the $\mathbf{q}$ -equidissection, where $k - q = q_1 + \dots + q_t$. This means that among the 2^k points $(x_{j_1}, \dots, x_{j_t})$ of Ψ_I , if we consider the first q_1 bits of x_{j_1} , the first q_2 bits of x_{j_2} , ..., and the first q_t bits of x_{j_t} , each of the 2^{k-q} possibilities occurs exactly the same number of times. This is possible only if $q \leq k$.

The $\mathbf{q}$ -equidistribution of Ψ_I depends only on the first q_j bits of x_{i_j} for $1 \leq j \leq t$, for the points $(x_{i_1}, \dots, x_{i_t})$ that belong to Ψ_I . The vector of these $q_1 + \dots + q_t = k - q$ bits can always be expressed as a linear function of the k bits of the initial state $\mathbf{x}_0$, i.e., as $\mathbf{M}_{\mathbf{q}}\mathbf{x}_0$ for some $(k - q) \times k$ binary matrix $\mathbf{M}_{\mathbf{q}}$, and it is easily seen that Ψ_I is $\mathbf{q}$ -equidistributed if and only if $\mathbf{M}_{\mathbf{q}}$ has full rank $k - q$. This provides an easy way of checking equidistribution (Fushimi, 1983; L'Ecuyer, 1996b; Tezuka, 1995).

If Ψ_I is $(\ell, \dots, \ell)$ -equidistributed for some $\ell \geq 1$, it is called *t-distributed with ℓ bits of accuracy*, or *(t, ℓ)-equidistributed* (L'Ecuyer, 1996b). The largest value of ℓ for which this holds is called the *resolution* of the set Ψ_I and is denoted by ℓ_I . This value has the upper bound $\ell_t^* = \min(\lfloor k/t \rfloor, w)$. The *resolution gap* of Ψ_I is defined as $\delta_I = \ell_t^* - \ell_I$. In the same vein as for MRGs, a worst-case figure of merit can be defined here by

$$\Delta_{\mathcal{J}} = \max_{I \in \mathcal{J}} \delta_I,$$

where $\mathcal{J}$ is a preselected class of index sets I .

The point set Ψ_I is a *(q, k, t)-net in base 2* (often called a *(t, m, s)-net* in the context of quasi-Monte Carlo methods, where a different notation is used (Niederreiter, 1992)), if it is $(q_1, \dots, q_t)$ -equidistributed in base 2 for all non-negative integers $q_1, \dots, q_t$ summing to $k - q$. We call the smallest such q the *q-value* of Ψ_I . The smaller it is, the better. One candidate for a figure of merit could be the *q-value* of Ψ_t for some large t . A major drawback of this measure is that it is extremely difficult to compute for good long-period generators (for which $k - q$ is large), because there are too many vectors $\mathbf{q}$ for which equidistribution needs to be checked. In practice, one must settle for figures of merit that involve a smaller number of equidissections.

When $\delta_I = 0$ for all sets I of the form $I = \{0, \dots, t - 1\}$, for $1 \leq t \leq k$, the RNG is said to be *maximally equidistributed* or *asymptotically random* for the word size w (L'Ecuyer, 1996b; Tezuka, 1995; Tootill et al., 1973). This property ensures perfect equidistribution of all sets Ψ_t , for any partition of the unit hypercube into subcubes of equal sizes, as long as $\ell \leq w$ and the number of subcubes does not exceed the number of points in Ψ_t . Large-period maximally equidistributed generators, together with their implementations, can be found in L'Ecuyer (1999c); L'Ecuyer and Panneton (2002), and Panneton and L'Ecuyer (2004), for example.

4.3 Lattice Structure in Spaces of Polynomials and Formal Series

The RNGs defined via (9)–(11) do not have a lattice structure in the real space like MRGs, but they do have a lattice structure in a space of formal series, as explained in Couture and L'Ecuyer (2000); L'Ecuyer (2004); Lemieux and L'Ecuyer (2003), and Tezuka (1995). The real space $\mathbb{R}$ is replaced by the space $\mathbb{L}_2$ of formal power series with coefficients in $\mathbb{F}_2$, of the form $\sum_{\ell=\omega}^{\infty} x_{\ell} z^{-\ell}$ for some integer ω . In that setting, the lattices have the form

$$\mathcal{L}_t = \left\{ \mathbf{v}(z) = \sum_{j=1}^t h_j(z) \mathbf{v}_j(z) \text{ such that each } h_j(z) \in \mathbb{F}_2[z] \right\},$$

where $\mathbb{F}_2[z]$ is the ring of polynomials with coefficients in $\mathbb{F}_2$, and the basis vectors $\mathbf{v}_j$ are in $\mathbb{L}_2^t$. The elements of the dual lattice $\mathcal{L}_t^*$ are the vectors $\mathbf{h}(z)$ in $\mathbb{L}_2^t$ whose scalar product with any vector of $\mathcal{L}_t$ is a polynomial (in $\mathbb{F}_2[z]$). We define the mapping $\varphi : \mathbb{L}_2 \rightarrow \mathbb{R}$ by

$$\varphi \left(\sum_{\ell=\omega}^{\infty} x_{\ell} z^{-\ell} \right) = \sum_{\ell=\omega}^{\infty} x_{\ell} 2^{-\ell}.$$

Then, it turns out that the point set Ψ_t produced by the generator is equal to $\varphi(\mathcal{L}_t) \cap [0, 1)^t$. Moreover, the equidistribution properties examined in Sect. 4.2 can be expressed in terms of lengths of shortest vectors in the dual lattice, with appropriate definitions of the length (or norm). Much of the theory and algorithms developed for lattices in the real space can be adapted to these new types of lattices (Couture and L'Ecuyer, 2000; L'Ecuyer, 2004; Lemieux and L'Ecuyer, 2003; Tezuka, 1995).

4.4 The LFSR Generator

The *Tausworthe* or *linear feedback shift register* (LFSR) generator (Tausworthe, 1965; L'Ecuyer, 1996b; Tezuka, 1995) is a special case of (9)–(11) with $\mathbf{A} = \mathbf{A}_0^s$ (in $\mathbb{F}_2$) for some positive integer s , where

$$\mathbf{A}_0 = \begin{pmatrix} & & 1 & & \\ & & & \ddots & \\ & & & & 1 \\ a_k & a_{k-1} & \dots & a_1 & \end{pmatrix}, \quad (13)$$

$a_1, \dots, a_k$ are in $\mathbb{F}_2$, $a_k = 1$, and all blank entries in the matrix are zeros. We take $w \leq k$ and the matrix $\mathbf{B}$ contains the first w lines of the $k \times k$ identity matrix. The RNG thus obtained can be defined equivalently by

$$x_i = a_1 x_{i-1} + \dots + a_k x_{i-k} \pmod{2}, \quad (14)$$

$$u_i = \sum_{\ell=1}^w x_{is+\ell-1} 2^{-\ell}. \quad (15)$$

Here, $P(z)$ is the characteristic polynomial of the matrix $\mathbf{A}_0^s$, not the characteristic polynomial of the recurrence (14), and the choice of s is important for determining the quality of the generator. A frequently encountered case is when a single a_j is nonzero in addition to a_k ; then, $P(z)$ is a trinomial and we say that we have a *trinomial-based* LFSR generator. These generators

are known to have important statistical deficiencies (Matsumoto and Kurita, 1996; Tezuka, 1995) but they can be used as components of combined RNGs.

LFSR generators can be expressed as LCGs in a space of polynomials (Tezuka and L'Ecuyer, 1991; Tezuka, 1995; L'Ecuyer, 1994). With this representation, their lattice structure as discussed in Sect. 4.3 follows immediately.

4.5 The GFSR and Twisted GFSR

Here we take $\mathbf{A}$ as the $pq \times pq$ matrix

$$\mathbf{A} = \begin{pmatrix} & \mathbf{I}_p & \mathbf{S} \\ \mathbf{I}_p & & \\ & \mathbf{I}_p & \\ & & \ddots \\ & & & \mathbf{I}_p \end{pmatrix}$$

for some positive integers p and q , where $\mathbf{I}_p$ is the $p \times p$ identity matrix, $\mathbf{S}$ is a $p \times p$ matrix, and the matrix $\mathbf{I}_p$ on the first line is in columns $(r-1)p+1$ to rp for some positive integer r . Often, $w = p$ and $\mathbf{B}$ contains the first w lines of the $pq \times pq$ identity matrix. If $\mathbf{S}$ is also the identity matrix, the generator thus obtained is the trinomial-based *generalized feedback shift register* (GFSR), for which $\mathbf{x}_i$ is obtained by a bitwise exclusive-or of $\mathbf{x}_{i-r}$ and $\mathbf{x}_{i-q}$. This gives a very fast RNG, but its period length cannot exceed $2^q - 1$, because each bit of $\mathbf{x}_i$ follows the same binary recurrence of order $k = q$, with characteristic polynomial $P(z) = z^q - z^{q-r} - 1$.

More generally, we can define $\mathbf{x}_i$ as the bitwise exclusive-or of $\mathbf{x}_{i-r_1}, \mathbf{x}_{i-r_2}, \dots, \mathbf{x}_{i-r_d}$ where $r_d = q$, so that each bit of $\mathbf{x}_i$ follows a recurrence in $\mathbb{F}_2$ whose characteristic polynomial $P(z)$ has $d+1$ nonzero terms. However, the period length is still bounded by $2^q - 1$, whereas considering the pq -bit state, we should rather expect a period length close to 2^{pq} . This was the main motivation for the *twisted GFSR* (TGFSR) generator. In the original version introduced by Matsumoto and Kurita (1992), $w = p$ and the matrix $\mathbf{S}$ is defined as the transpose of $\mathbf{A}_0$ in (13), with k replaced by p . The characteristic polynomial of $\mathbf{A}$ is then $P(z) = P_S(z^q + z^m)$, where $P_S(z) = z^p - a_p z^{p-1} - \dots - a_1$ is the characteristic polynomial of S , and its degree is $k = pq$. If the parameters are selected so that $P(z)$ is primitive over $\mathbb{F}_2$, then the TGFSR has period length $2^k - 1$. Matsumoto and Kurita (1994) pointed out important weaknesses of the original TGFSR and proposed an improved version that uses a well-chosen matrix $\mathbf{B}$ whose lines differ from those of the identity. The operations implemented by this matrix are called *tempering* and their purpose is to improve the uniformity of the points produced by the RNG. The *Mersenne twister* (Matsumoto and Nishimura, 1998; Nishimura, 2000) is a variant of the TGFSR where k is slightly less than pq and can be a prime number. A specific instance proposed by Matsumoto and Nishimura (1998) is

fast, robust, has the huge period length of $2^{19937} - 1$, and has become quite popular.

In the *multiple recursive matrix method* of Niederreiter (1995), the first row of $p \times p$ matrices in $\mathbf{A}$ contains arbitrary matrices. However, a fast implementation is possible only when these matrices are sparse and have a special structure.

4.6 Combined Linear Generators Over $\mathbb{F}_2$

Many of the best generators based on linear recurrences over $\mathbb{F}_2$ are constructed by combining the outputs of two or more RNGs having a simple structure. The idea is the same as for MRGs: select simple components that can run fast but such that their combination has a more complicated structure and highly-uniform sets Ψ_I for the sets I considered important.

Consider J distinct recurrences of the form (9)–(10), where the j th recurrence has parameters $(k, w, \mathbf{A}, \mathbf{B}) = (k_j, w, \mathbf{A}_j, \mathbf{B}_j)$ and state $\mathbf{x}_{j,i}$ at step i , for $j = 1, \dots, J$. The output of the combined generator at step i is defined by

$$\begin{aligned} \mathbf{y}_i &= \mathbf{B}_1 \mathbf{x}_{1,i} \oplus \dots \oplus \mathbf{B}_J \mathbf{x}_{J,i}, \\ u_i &= \sum_{\ell=1}^w y_{i,\ell-1} 2^{-\ell}, \end{aligned}$$

where $\oplus$ denotes the bitwise exclusive-or operation. One can show (Tezuka, 1995) that the period length ρ of this combined generator is the least common multiple of the period lengths ρ_j of its components. Moreover, this combined generator is equivalent to the generator (9)–(11) with $k = k_1 + \dots + k_J$, $\mathbf{A} = \text{diag}(\mathbf{A}_1, \dots, \mathbf{A}_J)$, and $\mathbf{B} = (\mathbf{B}_1, \dots, \mathbf{B}_J)$.

With this method, by selecting the parameters carefully, the combination of LFSR generators with characteristic polynomials $P_1(z), \dots, P_J(z)$ gives yet another LFSR with characteristic polynomial $P(z) = P_1(z) \dots P_J(z)$ and period length equal to the product of the period lengths of the components (Tezuka and L'Ecuyer, 1991; Wang and Compagner, 1993; L'Ecuyer, 1996b; Tezuka, 1995). Tables and fast implementations of maximally equidistributed combined LFSR generators are given in L'Ecuyer (1996b).

The TGFSR and Mersenne twister generators proposed in Matsumoto and Kurita (1994); Matsumoto and Nishimura (1998) and Nishimura (2000) cannot be maximally equidistributed. However, concrete examples of maximally equidistributed combined TGFSR generators with period lengths near 2^{466} and 2^{1250} are given in L'Ecuyer and Panneton (2002). These generators have the additional property that the resolution gaps δ_I are zero for a class of small sets I with indices not too far apart.

5 Nonlinear RNGs

All RNGs discussed so far are based on linear recurrences and their structure may be deemed too regular. There are at least two ways of getting rid of this regular linear structure: (1) use a nonlinear transition function f or (2) keep the transition function linear but use a nonlinear output function g . Several types of nonlinear RNGs have been proposed over the years; see, e.g., Blum et al. (1986); Eichenauer-Herrmann (1995); Eichenauer-Herrmann et al. (1997); Hellekalek and Wegenkittl (2003); Knuth (1998); L'Ecuyer (1994); Niederreiter and Shparlinski (2002), and Tezuka (1995). Their nonlinear mappings are defined in various ways by multiplicative inversion in a finite field, quadratic and cubic functions in the finite ring of integers modulo m , and other more complicated transformations. Many of them have output sequences that tend to behave much like i.i.d. $U(0, 1)$ sequences even over their entire period length, in contrast with “good” linear RNGs, whose point sets Ψ_t are much more regular than typical random points (Eichenauer-Herrmann et al., 1997; L'Ecuyer and Hellekalek, 1998; L'Ecuyer and Granger-Piché, 2003; Niederreiter and Shparlinski, 2002). On the other hand, their statistical properties have been analyzed only empirically or via asymptotic theoretical results. For specific nonlinear RNGs, the uniformity of the point sets Ψ_t is very difficult to measure theoretically. Moreover, the nonlinear RNGs are generally significantly slower than the linear ones. The RNGs recommended for cryptology are all nonlinear.

An interesting idea for adding nonlinearity without incurring an excessive speed penalty is to combine a small nonlinear generator with a fast long-period linear one (Aiello et al., 1998; L'Ecuyer and Granger-Piché, 2003). L'Ecuyer and Granger-Piché (2003) show how to do this while ensuring theoretically the good uniformity properties of Ψ_t for the combined generator. A very fast implementation can be achieved by using precomputed tables for the nonlinear component. Empirical studies suggest that mixed linear-nonlinear combined generators are more robust than the linear ones with respect to statistical tests, because of their less regular structure.

Several authors have proposed various ways of combining RNGs to produce streams of random numbers with less regularity and better “randomness” properties; see, e.g., Collings (1987); Knuth (1998); Gentle (2003); Law and Kelton (2000); L'Ecuyer (1994); Fishman (1996); Marsaglia (1985), and other references given there. This includes *shuffling* the output sequence of one generator using another one (or the same one), alternating between several streams, or just adding them in different ways. Most of these techniques are heuristics. They usually improve the uniformity (empirically), but they can also make it worse. For *random variables* in the mathematical sense, certain types of combinations (e.g., addition modulo 1) can *provably* improve the uniformity, and some authors have used this fact to argue that combined RNGs are provably better than their components alone (Brown and Solomon, 1979; Deng and George, 1990; Marsaglia, 1985; Gentle, 2003), but this argument is

faulty because the output sequences of RNGs are deterministic, not sequences of independent random variables. To assess the quality of a combined generator, one must analyze the mathematical structure of the combined generator itself rather than the structure of its components (L'Ecuyer, 1996b,a, 1998; L'Ecuyer and Granger-Piché, 2003; Tezuka, 1995).

6 Examples of Statistical Tests

As mentioned earlier, a statistical test for RNGs is defined by a random variable X whose distribution under $\mathcal{H}_0$ can be well approximated. When X takes the value x , we define the right and left p -values of the test by

$$p_R = P[X \geq x \mid \mathcal{H}_0] \quad \text{and} \quad p_L = P[X \leq x \mid \mathcal{H}_0].$$

When testing RNGs, there is no need to prespecify the level of the test. If any of the right or left p -value is extremely close to zero, e.g., less than 10^{-15} , then it is clear that $\mathcal{H}_0$ (and the RNG) must be rejected. When a *suspicious* p -value is obtained, e.g., near 10^{-2} or 10^{-3} , one can just repeat this particular test a few more times, perhaps with a larger sample size. Almost always, things will then clarify.

Most tests are defined by partitioning the possible realizations of $(u_0, \dots, u_{\tau-1})$ into a finite number of subsets (where the integer τ can be random or deterministic), computing the probability p_j of each subset j under $\mathcal{H}_0$, and measuring the discrepancy between these probabilities and empirical frequencies from realizations simulated by the RNG.

A special case that immediately comes to mind is to take $\tau = t$ (a constant) and cut the interval $[0, 1)$ into d equal segments for some positive integer d , in order to partition the hypercube $[0, 1)^t$ into $k = d^t$ subcubes of volume $1/k$. We then generate n points $\mathbf{u}_i = (u_{ti}, \dots, u_{ti+t-1}) \in [0, 1)^t$, for $i = 0, \dots, n-1$, and count the number N_j of points falling in subcube j , for $j = 0, \dots, k-1$. Any measure of distance (or divergence) between the numbers N_j and their expectations n/k can define a test statistic X . The tests thus defined are generally called *serial tests* of uniformity (Knuth, 1998; L'Ecuyer et al., 2002b). They can be *sparse* (if $n/k \ll 1$), or *dense* (if $n/k \gg 1$), or somewhere in between. There are also *overlapping* versions, where the points are defined by $\mathbf{u}_i = (u_i, \dots, u_{i+t-1})$ for $i = 0, \dots, n-1$ (they have overlapping coordinates).

Special instances for which the distribution under $\mathcal{H}_0$ is well-known are the chi-square, the (negative) empirical entropy, and the number of collisions (L'Ecuyer and Hellekalek, 1998; L'Ecuyer et al., 2002b; Read and Cressie, 1988). For the latter, the test statistic X is the number of times a point falls in a subcube that already had a point in it. Its distribution under $\mathcal{H}_0$ is approximately Poisson with mean $\lambda_1 = n^2/(2k)$, if n is large and λ_1 not too large.

A variant is the *birthday spacings* test, defined as follows (Marsaglia, 1985; Knuth, 1998; L'Ecuyer and Simard, 2001). Let $I_{(1)} \leq \dots \leq I_{(n)}$ be the numbers of the subcubes that contain the points, sorted by increasing order. Define the *spacings* $S_j = I_{(j+1)} - I_{(j)}$, for $j = 1, \dots, n-1$, and let X be the number of collisions between these spacings. Under $\mathcal{H}_0$, X is approximately Poisson with mean $\lambda_2 = n^3/(4k)$, if n is large and λ_2 not too large.

Consider now a MRG, for which Ψ_t has a regular lattice structure. Because of this regularity the points of Ψ_t will tend to be more evenly distributed among the subcubes than random points. For a well-chosen k and large enough n , we expect the collision test to detect this: it is likely that there will be too few collisions. In fact, the same applies to any RNG whose set Ψ_t is very evenly distributed. When a birthday spacings test with a very large k is applied to a MRG, the numbers of the subcubes that contain one point of Ψ_t tend to be too evenly spaced and the test detects this by finding too many collisions.

These specific interactions between the test and the structure of the RNG lead to systematic patterns in the p -values of the tests. To illustrate this, suppose that we take k slightly larger than the cardinality of Ψ_t (so $k \approx \rho$) and that due to the excessive regularity, no collision is observed in the collision test. The left p -value will then be $p_L \approx P[X \leq 0 \mid X \sim \text{Poisson}(\lambda_1)] = \exp[-n^2/(2k)]$. For this p -value to be smaller than a given ϵ , we need a sample size n proportional to the square root of the period length ρ . And after that, p_L decreases exponentially fast in n^2 .

Extensive experiments with LCGs, MRGs, and LFSR generators confirms that this is actually what happens with these RNGs (L'Ecuyer and Hellekalek, 1998; L'Ecuyer, 2001; L'Ecuyer et al., 2002b). For example, if we take $\epsilon = 10^{-15}$ and define n_0 as the minimal sample size n for which $p_L < \epsilon$, we find that $n_0 \approx 16\rho^{1/2}$ (plus some noise) for LCGs that behave well in the spectral test as well as for LFSR generators. For the birthday spacings test, the rule for LCGs is $n_0 \approx 16\rho^{1/3}$ instead (L'Ecuyer and Simard, 2001). So to be safe with respect to these tests, the period length ρ must be so large that generating more than $\rho^{1/3}$ numbers is practically unfeasible. This certainly disqualifies all LCGs with modulus smaller than 2^{100} or so.

Other types of tests for RNGs include tests based on the closest pairs of points among n points generated in the hypercube, tests based on random walks on the real line or over the integers, tests based on the linear complexity of a binary sequence, tests based on the simulation of dices or poker hands, and many others (Gentle, 2003; Knuth, 1998; L'Ecuyer and Simard, 2002; Marsaglia, 1996; Rukhin et al., 2001; Vattulainen et al., 1995).

When testing RNGs, there is no specific alternative hypothesis to $\mathcal{H}_0$. Different tests are needed to detect different types of departures from $\mathcal{H}_0$. *Test suites* for RNGs include a selection of tests, with predetermined parameters and sample sizes. The best known are probably DIEHARD (Marsaglia, 1996) and the NIST test suite (Rukhin et al., 2001). The library *TestU01* (L'Ecuyer and Simard, 2002) implements a large selection of tests in the C language

and provides a variety of test suites, some designed for i.i.d. $U(0, 1)$ output sequences and others for strings of bits.

7 Available Software and Recommendations

When we apply test suites to RNGs currently found in commercial software (statistical and simulation software, spreadsheets, etc.), we find that many of them fail the tests spectacularly (L'Ecuyer, 1997, 2001). There is no reason to use these poor RNGs, because there are also several good ones that are fast, portable, and pass all these test suites with flying colors. Among them we recommend, for example, the combined MRGs, combined LFSRs, and Mersenne twisters proposed in L'Ecuyer (1999c,a); L'Ecuyer and Panneton (2002); Matsumoto and Nishimura (1998), and Nishimura (2000).

A convenient object-oriented software package with multiple streams and substreams of random numbers, is described in L'Ecuyer et al. (2002a) and is available in Java, C, and C++, at <http://www.iro.umontreal.ca/~lecuyer>.

8 Non-Uniform Random Variate Generation

Like for the uniform case, non-uniform variate generation often involves approximations and compromises. The first requirement is, of course, *correctness*. This does not mean that the generated random variate X must always have *exactly* the required distribution, because this would sometimes be much too costly or even impossible. But we must have a *good approximation* and, preferably, some understanding of the quality of that approximation. *Robustness* is also important: when the accuracy depends on the parameters of the distribution, it must be good *uniformly* over the entire range of parameter values that we are interested in.

The method must also be *efficient* both in terms of speed and memory usage. Often, it is possible to increase the speed by using more memory (e.g., for larger precomputed tables) or by relaxing the accuracy requirements. Some methods need a one-time setup to compute constants and construct tables. The setup time can be significant but may be well worth spending if it is amortized by a large number of subsequent calls to the generator. For example, it makes sense to invest in a more extensive setup if we plan to make a million calls to a given generator than if we expect to make only a few calls, assuming that this investment can improve the speed of the generator sufficiently.

In general, compromises must be made between simplicity of the algorithm, quality of the approximation, robustness with respect to the distribution parameters, and efficiency (generation speed, memory requirements, and setup time).

In many situations, compatibility with variance reduction techniques is another important issue (Bratley et al., 1987; Law and Kelton, 2000). We may be willing to sacrifice the speed of the generator to preserve inversion, because the gain in efficiency obtained via the variance reduction methods may more than compensate (sometimes by orders of magnitude) for the slightly slower generator.

8.1 Inversion

The inversion method, defined in the introduction, should be the method of choice for generating non-uniform random variates in a majority of situations. The fact that $X = F^{-1}(U)$ is a monotone (non-decreasing) function of U makes this method compatible with important variance reductions techniques such as common random numbers, antithetic variates, latin hypercube sampling, and randomized quasi-Monte Carlo methods (Bratley et al., 1987; Law and Kelton, 2000; L'Ecuyer and Lemieux, 2000).

For some distributions, an analytic expression can be obtained for the inverse distribution function F^{-1} and inversion can be easily implemented. As an example, consider the *Weibull* distribution function with parameters $\alpha > 0$ and $\beta > 0$, defined by $F(x) = 1 - \exp[-(x/\beta)^\alpha]$ for $x > 0$. It is easy to see that $F^{-1}(U) = \beta[-\ln(1 - U)]^{1/\alpha}$. For $\alpha = 1$, we have the special case of the exponential distribution with mean β .

For an example of a simple discrete distribution, suppose that $P[X = i] = p_i$ where $p_0 = 0.6$, $p_1 = 0.3$, $p_2 = 0.1$, and $p_i = 0$ elsewhere. The inversion method in this case will return 0 if $U < 0.6$, 1 if $0.6 \leq U < 0.9$, and 2 if $U \geq 0.9$. For the discrete uniform distribution over $\{0, \dots, k-1\}$, return $X = \lfloor kU \rfloor$. As another example, let X have the *geometric* distribution with parameter p , so $P[X = x] = p(1-p)^x$ for $x = 0, 1, 2, \dots$, where $0 < p < 1$. Then, $F(x) = 1 - (1-p)^{\lfloor x+1 \rfloor}$ for $x \geq 0$ and one can show that $X = F^{-1}(U) = \lceil \ln(1 - U) / \ln(1 - p) \rceil - 1$.

There are other distributions (e.g., the normal, Student, chi-square) for which there is no closed-form expression for F^{-1} but good numerical approximations are available (Bratley et al., 1987; Gentle, 2003; Marsaglia et al., 1994). When the distribution has only scale and location parameters, we need to approximate F^{-1} only for a standardized version of the distribution. For the normal distribution, for example, it suffices to have an efficient method for evaluating the inverse distribution function of a $N(0, 1)$ random variable Z , since a normal with mean μ and variance σ^2 can be generated by $X = \sigma Z + \mu$. When shape parameters are involved (e.g., the gamma and beta distributions), things are more complicated because F^{-1} then depends on the parameters in a more fundamental manner.

Hörmann and Leydold (2003) propose a general adaptive and automatic method that constructs a highly accurate Hermite interpolation method of F^{-1} . In a one-time setup, their method produces tables for the interpolation

points and coefficients. Random variate generation using these tables is then quite fast.

A less efficient but simpler way of implementing inversion when a method is available for computing F is via binary search (Cheng, 1998). If the density is also available and if it is unimodal with known mode, a Newton-Raphson iteration method can advantageously replace the binary search (Cheng, 1998; Devroye, 1986).

To implement inversion for general discrete distributions, sequential search and binary search with look-up tables are the standard methods (Bratley et al., 1987; Cheng, 1998). For a discrete distribution over the values $x_1 < \dots < x_k$, one first tabulates the pairs $(x_i, F(x_i))$, where $F(x_i) = P[X \leq x_i]$ for $i = 1, \dots, k$. To generate X , it then suffices to generate $U \sim U(0, 1)$, find $I = \min\{i \mid F(x_i) \geq U\}$, and return $X = x_I$. The following algorithms do that.

Sequential search (needs $O(k)$ iterations in the worst case);

```
generate  $U \sim U(0, 1)$ ;    let  $i = 1$ ;
while  $F(x_i) < U$  do  $i = i + 1$ ;
return  $x_i$ .
```

Binary search (needs $O(\log k)$ iterations in the worst case);

```
generate  $U \sim U(0, 1)$ ;    let  $L = 0$  and  $R = k$ ;
while  $L < R - 1$  do
   $m = \lfloor (L + R)/2 \rfloor$ ;
  if  $F(x_m) < U$  then  $L = m$  else  $R = m$ ;
/* Invariant: at this stage, the index  $I$  is in  $\{L + 1, \dots, R\}$ . */
return  $x_R$ .
```

These algorithms can be modified in many different ways. For example, if $k = \infty$, in the binary search, one can start with an arbitrary value of R , double it until $F(x_R) \geq U$, and start the algorithm with this R and $L = R/2$. Of course, only a finite portion of the table (a portion that contains most of the probability mass) would be precomputed in this case, the other values can be computed only when needed. This can also be done if k is finite but large.

Another class of techniques use indexing or buckets to speed up the search (Chen and Asau, 1974; Bratley et al., 1987; Devroye, 1986). For example, one can partition the interval $(0, 1)$ into c subintervals of equal sizes and use (pre-tabulated) initial values of (L, R) that depend on the subinterval in which U falls. For the subinterval $[j/c, (j+1)/c)$ the values of L and R would be $L_j = F^{-1}(j/c)$ and $R_j = F^{-1}((j+1)/c)$, for $j = 0, \dots, c-1$. The subinterval number that corresponds to a given U is simply $J = \lfloor cU \rfloor$. Once we know that subinterval, we can search it by linear or binary search. With a larger value of c the search is faster (on the average) but the setup is more costly and a larger amount of memory is needed. So a compromise must be made depending on the situation (e.g., the value of k , the number of variates we expect to generate, etc.). For $c = 1$, we recover the basic sequential and binary

search algorithms given above. A well-implemented indexed search with a large enough c is generally competitive with the alias method (described in the next paragraph). A combined indexed/binary search algorithm is given below. An easy adaptation gives the combined indexed/sequential search, which is generally preferable when k/c is small, because it has smaller overhead.

Indexed search (combined with binary search);
 generate $U \sim U(0, 1)$; let $J = \lfloor cU \rfloor$, $L = L_J$, and $R = R_J$;
 while $L < R - 1$ do
 $m = \lfloor (L + R)/2 \rfloor$;
 if $F(x_m) < U$ then $L = m$ else $R = m$;
 return x_R .

These search methods are also useful for piecewise-linear (or piecewise-polynomial) distribution functions. Essentially, it suffices to add an interpolation step at the end of the algorithm, after the appropriate linear (or polynomial) piece has been determined (Bratley et al., 1987).

Finally, the stochastic model itself can sometimes be selected in a way that makes inversion easier. For example, one can fit a parametric, highly-flexible, and easily computable inverse distribution function F^{-1} to the data, directly or indirectly (Nelson and Yamnitsky, 1998; Wagner and Wilson, 1996).

There are situations where *speed* is important and where non-inversion methods are appropriate. In forthcoming subsections, we outline the main non-inversion methods.

8.2 The Alias Method

Sequential and binary search require $O(k)$ and $O(\log k)$ time, respectively, in the worst case, to generate a random variate X by inversion over the finite set $\{x_1, \dots, x_k\}$. The *alias method* (Walker, 1974, 1977) can generate such a X in $O(1)$ time per variate, after a table setup that takes $O(k)$ time and space. On the other hand, it does not implement inversion, i.e., the transformation from U to X is not monotone.

To explain the idea, consider a bar diagram of the distribution, where each index i has a bar of height $p_i = P[X = x_i]$. The idea is to “equalize” the bars so that they all have height $1/k$, by cutting-off bar pieces and transferring them to other bars. This is done in a way that in the new diagram, each bar i contains one piece of size q_i (say) from the original bar i and one piece of size $1/k - q_i$ from another bar whose index j , denoted $A(i)$, is called the *alias* value of i . The setup procedure initializes two tables, A and R , where $A(i)$ is the alias value of i and $R(i) = (i - 1)/k + q_i$. See Devroye (1986) and Law and Kelton (2000) for the details. To generate X , we generate $U \sim U[0, 1]$, define $I = \lceil kU \rceil$, and return $X = x_I$ if $U < R(I)$ and $X = x_{A(I)}$ otherwise.

There is a version of the alias method for continuous distributions, called the *acceptance-complement* method (Kronmal and Peterson, 1984; Devroye,

1986; Gentle, 2003). The idea is to decompose the density f of the target distribution as the convex combination of two densities f_1 and f_2 , $f = wf_1 + (1-w)f_2$ for some real number $w \in (0, 1)$, in a way that $wf_1 \leq g$ for some other density g and so that it is easy to generate from g and f_2 . The algorithm works as follows: Generate X from density g and $U \sim U(0, 1)$; if $Ug(X) \leq wf_1(X)$ return X , otherwise generate a new X from density f_2 and return it.

8.3 Kernel Density Estimation and Generation

Instead of selecting a parametric distribution that is hard to invert and estimating the parameters, one can estimate the density via a *kernel density estimation* method for which random variate generation is very easy (Devroye, 1986; Hörmann and Leydold, 2000). In the case of a gaussian kernel, for example, one can generate variates simply by selecting one observation at random from the data and adding random noise generated from a normal distribution with mean zero. However, this method *is not* equivalent to inversion. Because of the added noise, selecting a larger observation does not necessarily guarantee a larger value for the generated variate.

8.4 The Rejection Method

Now suppose we want to generate X from a complicated density f . In fact f may be known only up to a multiplicative constant $\kappa > 0$, i.e., we know only κf . If we know f , we may just take $\kappa = 1$. We select another density r such that $\kappa f(x) \leq t(x) \stackrel{\text{def}}{=} ar(x)$ for all x for some constant a , and such that generating variates Y from the density r is easy. The function t is called a *hat function* or *majorizing function*. By integrating this inequality with respect to x on both sides, we find that $\kappa \leq a$. The following *rejection* method generates X with density f (von Neumann, 1951; Devroye, 1986; Evans and Swartz, 2000):

Rejection method;

```
repeat
  generate  $Y$  from the density  $r$  and  $U \sim U(0, 1)$ , independent;
until  $Ut(Y) \leq \kappa f(Y)$ ;
return  $X = Y$ .
```

The number R of turns into the “repeat” loop is one plus a geometric random variable with parameter κ/a , so $E[R] = a/\kappa$. Thus, we want $a/\kappa \geq 1$ to be as small as possible, i.e., we want to minimize the area between κf and t . There is generally a compromise between bringing a/κ close to 1 and keeping r simple.

When κf is expensive to compute, we can also use *squeeze functions* q_1 and q_2 that are faster to evaluate and such that $q_1(x) \leq \kappa f(x) \leq q_2(x) \leq t(x)$ for all x . To verify the condition $Ut(Y) \leq \kappa f(Y)$, we first check if $Ut(Y) \leq q_1(Y)$,

in which case we accept Y immediately, otherwise we check if $Ut(Y) \geq q_2(Y)$, in which case we reject Y immediately. The value of $\kappa f(Y)$ must be computed only when $Ut(Y)$ falls between the two squeezes. Sequences of imbedded squeezes can also be used, where the primary ones are the least expensive to compute, the secondary ones are a little more expensive but closer to κf , etc.

It is common practice to transform the density f by a smooth increasing function T (e.g., $T(x) = \log x$ or $T(x) = -x^{-1/2}$) selected so that it is easier to construct good hat and squeeze functions (often piecewise linear) for the transformed density $T(f(\cdot))$. By transforming back to the original scale, we get hat and squeeze functions for f . This is the *transformed density rejection* method, which has several variants and extensions (Devroye, 1986; Evans and Swartz, 2000; Hörmann et al., 2004).

The rejection method works for discrete distributions as well; it suffices to replace densities by probability mass functions.

8.5 Thinning for Point Processes with Time-Varying Rates

Thinning is a cousin of acceptance-rejection, often used for generating events from a non-homogeneous Poisson process. Suppose the process has rate $\lambda(t)$ at time t , with $\lambda(t) \leq \bar{\lambda}$ for all t , where $\bar{\lambda}$ is a finite constant. One can generate Poisson *pseudo-arrivals* at constant rate $\bar{\lambda}$ by generating interarrival times that are i.i.d. exponentials of mean $1/\bar{\lambda}$. Then, a pseudo-arrival at time t is accepted (becomes an arrival) with probability $\lambda(t)/\bar{\lambda}$ (i.e., if $U \leq \lambda(t)/\bar{\lambda}$, where U is an independent $U[0, 1]$), and rejected with probability $1 - \lambda(t)/\bar{\lambda}$. Non-homogeneous Poisson processes can also be generated by inversion (Bratley et al., 1987). The idea is to apply a nonlinear transformation to the time scale to make the process homogeneous with rate 1 in the new time scale. Arrival times are generated in this new time scale (which is easy), and then transformed back to the original time scale. The method can be adapted to other types of point processes with time-varying rates.

8.6 The Ratio-of-Uniforms Method

If f is a density over the real-line, κ an arbitrary positive constant, and the pair (U, V) has the uniform distribution over the set

$$\mathcal{C} = \left\{ (u, v) \in \mathbb{R}^2 \text{ such that } 0 \leq u \leq \sqrt{\kappa f(v/u)} \right\},$$

then V/U has density f (Kinderman and Monahan, 1977; Devroye, 1986; Gentle, 2003). This interesting property can be exploited to generate X with density f : generate (U, V) uniformly over $\mathcal{C}$ and return $X = V/U$. This is the *ratio-of-uniforms* method. The key issue is how do we generate a point uniformly over $\mathcal{C}$. In the cases where this can be done efficiently, we immediately have an efficient way of generating X .

The most frequent approach for generating (U, V) uniformly over $\mathcal{C}$ is the rejection method: Define a region $\mathcal{C}_2$ that contains $\mathcal{C}$ and in which it is easy to generate a point uniformly (for example, a rectangular box or a polygonal region). To generate X , repeat: generate (U, V) uniformly over $\mathcal{C}_2$, until it belongs to $\mathcal{C}$. Then return $X = V/U$. If there is another region $\mathcal{C}_1$ contained in $\mathcal{C}$ and for which it is very fast to check if a point (U, V) is in $\mathcal{C}_1$, this $\mathcal{C}_1$ can also be used as a squeeze to accelerate the verification that the point belongs to $\mathcal{C}$. Several special cases and refinements are described in Devroye (1986); Gentle (2003); Leydold (2000), and other references given there.

8.7 Composition and Convolution

Suppose F is a convex combination of several distributions, i.e., $F(x) = \sum_j p_j F_j(x)$, or more generally $F(x) = \int F_y(x) dH(y)$. To generate from F , one can generate $J = j$ with probability p_j (or Y from H), then generate X from F_J (or F_Y). This method, called the *composition algorithm*, is useful for generating from *compound* distributions such as the hyperexponential or from compound Poisson processes. It is also frequently used to design specialized algorithms for generating from complicated densities. The idea is to partition the area under the complicated density into pieces, where piece j has surface p_j . To generate X , first select a piece (choose piece j with probability p_j), then draw a random point uniformly over that piece and project it to the horizontal axis. If the partition is defined so that it is fast and easy to generate from the large pieces, then X will be returned very quickly most of the time. The rejection method with a squeeze is often used to generate from some of the pieces.

A dual method to composition is the *convolution method*, which can be used when $X = Y_1 + Y_2 + \dots + Y_n$, where the Y_i 's are independent with specified distributions. With this method, one just generates the Y_i 's and sum up. This requires at least n uniforms. Examples of random variables that can be expressed as sums like this include the hypoexponential, Erlang, and binomial distributions.

8.8 Other Special Techniques

Besides the general methods, several specialized and sometimes very elegant techniques have been designed for commonly used distributions such as the Poisson distribution with small mean, the normal (e.g., the Box-Muller method), for generating points uniformly on a k -dimensional sphere, for generating random permutations, and so on. Details can be found, e.g., in Bratley et al. (1987); Cheng (1998); Devroye (1986); Fishman (1996); Gentle (2003).

Recently, there has been an effort in developing *automatic* or *black box* algorithms for generating variates from an arbitrary (known) density, and reliable software that implements these methods (Hörmann and Leydold, 2000; Hörmann et al., 2004; Leydold and Hörmann, 2002; Leydold et al., 2002).

Acknowledgements

This work has been supported by the Natural Sciences and Engineering Research Council of Canada (NSERC) Grant No. ODGP0110050, NATEQ-Québec grant No. 02ER3218, and a Canada Research Chair to the author. Wolfgang Hörmann, Josef Leydold, François Panneton, and Richard Simard made helpful comments and corrections on an earlier draft. The author has been asked to write chapters on Random Number Generation for several handbooks and encyclopedia recently. Inevitably, there is a large amount of duplication between these chapters.

References

- Aiello, W., Rajagopalan, S., and Venkatesan, R. (1998). Design of practical and provably good random number generators. *Journal of Algorithms*, 29(2):358–389.
- Blum, L., Blum, M., and Schub, M. (1986). A simple unpredictable pseudorandom number generator. *SIAM Journal on Computing*, 15(2):364–383.
- Bratley, P., Fox, B. L., and Schrage, L. E. (1987). *A Guide to Simulation*. Springer-Verlag, New York, second edition.
- Brown, M. and Solomon, H. (1979). On combining pseudorandom number generators. *Annals of Statistics*, 1:691–695.
- Chen, H. C. and Asau, Y. (1974). On generating random variates from an empirical distribution. *AIEE Transactions*, 6:163–166.
- Cheng, R. C. H. (1998). Random variate generation. In Banks, J., editor, *Handbook of Simulation*, pages 139–172. Wiley. chapter 5.
- Collings, B. J. (1987). Compound random number generators. *Journal of the American Statistical Association*, 82(398):525–527.
- Conway, J. H. and Sloane, N. J. A. (1999). *Sphere Packings, Lattices and Groups*. Grundlehren der Mathematischen Wissenschaften 290. Springer-Verlag, New York, 3rd edition.
- Couture, R. and L'Ecuyer, P. (1994). On the lattice structure of certain linear congruential sequences related to AWC/SWB generators. *Mathematics of Computation*, 62(206):798–808.
- Couture, R. and L'Ecuyer, P. (1996). Orbits and lattices for linear random number generators with composite moduli. *Mathematics of Computation*, 65(213):189–201.
- Couture, R. and L'Ecuyer, P. (1997). Distribution properties of multiply-with-carry random number generators. *Mathematics of Computation*, 66(218):591–607.
- Couture, R. and L'Ecuyer, P. (2000). Lattice computations for random numbers. *Mathematics of Computation*, 69(230):757–765.

- Deng, L.-Y. and George, E. O. (1990). Generation of uniform variates from several nearly uniformly distributed variables. *Communications in Statistics*, B19(1):145–154.
- Deng, L.-Y. and Lin, D. K. J. (2000). Random number generation for the new century. *The American Statistician*, 54(2):145–150.
- Deng, L.-Y. and Xu, H. (2003). A system of high-dimensional, efficient, long-cycle and portable uniform random number generators. *ACM Transactions on Modeling and Computer Simulation*, 13(4):299–309.
- Devroye, L. (1986). *Non-Uniform Random Variate Generation*. Springer-Verlag, New York.
- Eichenauer-Herrmann, J. (1995). Pseudorandom number generation by non-linear methods. *International Statistical Reviews*, 63:247–255.
- Eichenauer-Herrmann, J., Herrmann, E., and Wegenkittl, S. (1997). A survey of quadratic and inversive congruential pseudorandom numbers. In Hellekalek, P., Larcher, G., Niederreiter, H., and Zinterhof, P., editors, *Monte Carlo and Quasi-Monte Carlo Methods in Scientific Computing*, volume 127 of *Lecture Notes in Statistics*, pages 66–97, New York. Springer.
- Evans, M. and Swartz, T. (2000). *Approximating Integrals via Monte Carlo and Deterministic Methods*. Oxford University Press, Oxford, UK.
- Fincke, U. and Pohst, M. (1985). Improved methods for calculating vectors of short length in a lattice, including a complexity analysis. *Mathematics of Computation*, 44:463–471.
- Fishman, G. S. (1996). *Monte Carlo: Concepts, Algorithms, and Applications*. Springer Series in Operations Research. Springer-Verlag, New York.
- Fushimi, M. (1983). Increasing the orders of equidistribution of the leading bits of the Tausworthe sequence. *Information Processing Letters*, 16:189–192.
- Gentle, J. E. (2003). *Random Number Generation and Monte Carlo Methods*. Springer, New York, second edition.
- Goresky, M. and Klapper, A. (2003). Efficient multiply-with-carry random number generators with maximal period. *ACM Transactions on Modeling and Computer Simulation*, 13(4):310–321.
- Hellekalek, P. and Larcher, G., editors (1998). *Random and Quasi-Random Point Sets*, volume 138 of *Lecture Notes in Statistics*. Springer, New York.
- Hellekalek, P. and Wegenkittl, S. (2003). Empirical evidence concerning AES. *ACM Transactions on Modeling and Computer Simulation*, 13(4):322–333.
- Hörmann, W. and Leydold, J. (2000). Automatic random variate generation for simulation input. In Joines, J. A., Barton, R. R., Kang, K., and Fishwick, P. A., editors, *Proceedings of the 2000 Winter Simulation Conference*, pages 675–682, Piscataway, NJ. IEEE Press.
- Hörmann, W. and Leydold, J. (2003). Continuous random variate generation by fast numerical inversion. *ACM Transactions on Modeling and Computer Simulation*, 13(4):347–362.
- Hörmann, W., Leydold, J., and Derflinger, G. (2004). *Automatic Nonuniform Random Variate Generation*. Springer-Verlag, Berlin.

- Kinderman, A. J. and Monahan, J. F. (1977). Computer generation of random variables using the ratio of uniform deviates. *ACM Transactions on Mathematical Software*, 3:257–260.
- Knuth, D. E. (1998). *The Art of Computer Programming, Volume 2: Seminumerical Algorithms*. Addison-Wesley, Reading, Mass., third edition.
- Kronmal, R. A. and Peterson, A. V. (1984). An acceptance-complement analogue of the mixture-plus-acceptance-rejection method for generating random variables. *ACM Transactions on Mathematical Software*, 10:271–281.
- Lagarias, J. C. (1993). Pseudorandom numbers. *Statistical Science*, 8(1):31–39.
- Law, A. M. and Kelton, W. D. (2000). *Simulation Modeling and Analysis*. McGraw-Hill, New York, third edition.
- L'Ecuyer, P. (1990). Random numbers for simulation. *Communications of the ACM*, 33(10):85–97.
- L'Ecuyer, P. (1994). Uniform random number generation. *Annals of Operations Research*, 53:77–120.
- L'Ecuyer, P. (1996a). Combined multiple recursive random number generators. *Operations Research*, 44(5):816–822.
- L'Ecuyer, P. (1996b). Maximally equidistributed combined Tausworthe generators. *Mathematics of Computation*, 65(213):203–213.
- L'Ecuyer, P. (1997). Bad lattice structures for vectors of non-successive values produced by some linear recurrences. *INFORMS Journal on Computing*, 9(1):57–60.
- L'Ecuyer, P. (1998). Random number generation. In Banks, J., editor, *Handbook of Simulation*, pages 93–137. Wiley. chapter 4.
- L'Ecuyer, P. (1999a). Good parameters and implementations for combined multiple recursive random number generators. *Operations Research*, 47(1):159–164.
- L'Ecuyer, P. (1999b). Tables of linear congruential generators of different sizes and good lattice structure. *Mathematics of Computation*, 68(225):249–260.
- L'Ecuyer, P. (1999c). Tables of maximally equidistributed combined LFSR generators. *Mathematics of Computation*, 68(225):261–269.
- L'Ecuyer, P. (2001). Software for uniform random number generation: Distinguishing the good and the bad. In *Proceedings of the 2001 Winter Simulation Conference*, pages 95–105, Piscataway, NJ. IEEE Press.
- L'Ecuyer, P. (2004). Polynomial integration lattices. In Niederreiter, H., editor, *Monte Carlo and Quasi-Monte Carlo Methods 2002*, pages 73–98, Berlin. Springer-Verlag.
- L'Ecuyer, P. and Andres, T. H. (1997). A random number generator based on the combination of four LCGs. *Mathematics and Computers in Simulation*, 44:99–107.
- L'Ecuyer, P., Blouin, F., and Couture, R. (1993). A search for good multiple recursive random number generators. *ACM Transactions on Modeling and Computer Simulation*, 3(2):87–98.

- L'Ecuyer, P. and Côté, S. (1991). Implementing a random number package with splitting facilities. *ACM Transactions on Mathematical Software*, 17(1):98–111.
- L'Ecuyer, P. and Couture, R. (1997). An implementation of the lattice and spectral tests for multiple recursive linear random number generators. *INFORMS Journal on Computing*, 9(2):206–217.
- L'Ecuyer, P. and Granger-Piché, J. (2003). Combined generators with components from different families. *Mathematics and Computers in Simulation*, 62:395–404.
- L'Ecuyer, P. and Hellekalek, P. (1998). Random number generators: Selection criteria and testing. In Hellekalek, P. and Larcher, G., editors, *Random and Quasi-Random Point Sets*, volume 138 of *Lecture Notes in Statistics*, pages 223–265. Springer, New York.
- L'Ecuyer, P. and Lemieux, C. (2000). Variance reduction via lattice rules. *Management Science*, 46(9):1214–1235.
- L'Ecuyer, P. and Lemieux, C. (2002). Recent advances in randomized quasi-Monte Carlo methods. In Dror, M., L'Ecuyer, P., and Szidarovszki, F., editors, *Modeling Uncertainty: An Examination of Stochastic Theory, Methods, and Applications*, pages 419–474. Kluwer Academic Publishers, Boston.
- L'Ecuyer, P. and Panneton, F. (2002). Construction of equidistributed generators based on linear recurrences modulo 2. In Fang, K.-T., Hickernell, F. J., and Niederreiter, H., editors, *Monte Carlo and Quasi-Monte Carlo Methods 2000*, pages 318–330. Springer-Verlag, Berlin.
- L'Ecuyer, P. and Proulx, R. (1989). About polynomial-time “unpredictable” generators. In *Proceedings of the 1989 Winter Simulation Conference*, pages 467–476. IEEE Press.
- L'Ecuyer, P. and Simard, R. (1999). Beware of linear congruential generators with multipliers of the form $a = \pm 2^q \pm 2^r$. *ACM Transactions on Mathematical Software*, 25(3):367–374.
- L'Ecuyer, P. and Simard, R. (2001). On the performance of birthday spacings tests for certain families of random number generators. *Mathematics and Computers in Simulation*, 55(1–3):131–137.
- L'Ecuyer, P. and Simard, R. (2002). *TestU01: A Software Library in ANSI C for Empirical Testing of Random Number Generators*. Software user's guide.
- L'Ecuyer, P., Simard, R., Chen, E. J., and Kelton, W. D. (2002a). An object-oriented random-number package with many long streams and substreams. *Operations Research*, 50(6):1073–1075.
- L'Ecuyer, P., Simard, R., and Wegenkittl, S. (2002b). Sparse serial tests of uniformity for random number generators. *SIAM Journal on Scientific Computing*, 24(2):652–668.
- L'Ecuyer, P. and Tezuka, S. (1991). Structural properties for two classes of combined random number generators. *Mathematics of Computation*, 57(196):735–746.

- L'Ecuyer, P. and Touzin, R. (2000). Fast combined multiple recursive generators with multipliers of the form $a = \pm 2^q \pm 2^r$. In Joines, J. A., Barton, R. R., Kang, K., and Fishwick, P. A., editors, *Proceedings of the 2000 Winter Simulation Conference*, pages 683–689, Piscataway, NJ. IEEE Press.
- L'Ecuyer, P. and Touzin, R. (2004). On the Deng-Lin random number generators and related methods. *Statistics and Computing*, 14:5–9.
- Leeb, H. (1995). Random numbers for computer simulation. Master's thesis, University of Salzburg.
- Lemieux, C. and L'Ecuyer, P. (2003). Randomized polynomial lattice rules for multivariate integration and simulation. *SIAM Journal on Scientific Computing*, 24(5):1768–1789.
- Leydold, J. (2000). Automatic sampling with the ratio-of-uniform method. *ACM Transactions on Mathematical Software*, 26(1):78–98.
- Leydold, J. and Hörmann, W. (2002). *UNURAN—A Library for Universal Non-Uniform Random Number Generators*. Available at <http://statistik.wu-wien.ac.at/unuran>.
- Leydold, J., Janka, E., and Hörmann, W. (2002). Variants of transformed density rejection and correlation induction. In Fang, K.-T., Hickernell, F. J., and Niederreiter, H., editors, *Monte Carlo and Quasi-Monte Carlo Methods 2000*, pages 345–356, Berlin. Springer-Verlag.
- Luby, M. (1996). *Pseudorandomness and Cryptographic Applications*. Princeton University Press, Princeton.
- Lüscher, M. (1994). A portable high-quality random number generator for lattice field theory simulations. *Computer Physics Communications*, 79:100–110.
- Marsaglia, G. (1985). A current view of random number generators. In *Computer Science and Statistics, Sixteenth Symposium on the Interface*, pages 3–10, North-Holland, Amsterdam. Elsevier Science Publishers.
- Marsaglia, G. (1996). The Marsaglia random number CDROM including the DIEHARD battery of tests of randomness. See <http://stat.fsu.edu/pub/diehard>.
- Marsaglia, G. and Zaman, A. (1991). A new class of random number generators. *The Annals of Applied Probability*, 1:462–480.
- Marsaglia, G., Zaman, A., and Marsaglia, J. C. W. (1994). Rapid evaluation of the inverse normal distribution function. *Statistic and Probability Letters*, 19:259–266.
- Matsumoto, M. and Kurita, Y. (1992). Twisted GFSR generators. *ACM Transactions on Modeling and Computer Simulation*, 2(3):179–194.
- Matsumoto, M. and Kurita, Y. (1994). Twisted GFSR generators II. *ACM Transactions on Modeling and Computer Simulation*, 4(3):254–266.
- Matsumoto, M. and Kurita, Y. (1996). Strong deviations from randomness in m -sequences based on trinomials. *ACM Transactions on Modeling and Computer Simulation*, 6(2):99–106.

- Matsumoto, M. and Nishimura, T. (1998). Mersenne twister: A 623-dimensionally equidistributed uniform pseudo-random number generator. *ACM Transactions on Modeling and Computer Simulation*, 8(1):3–30.
- Nelson, B. L. and Yamnitsky, M. (1998). Input modeling tools for complex problems. In *Proceedings of the 1998 Winter Simulation Conference*, pages 105–112, Piscataway, NJ. IEEE Press.
- Niederreiter, H. (1992). *Random Number Generation and Quasi-Monte Carlo Methods*, volume 63 of *SIAM CBMS-NSF Regional Conference Series in Applied Mathematics*. SIAM, Philadelphia.
- Niederreiter, H. (1995). The multiple-recursive matrix method for pseudorandom number generation. *Finite Fields and their Applications*, 1:3–30.
- Niederreiter, H. and Shparlinski, I. E. (2002). Recent advances in the theory of nonlinear pseudorandom number generators. In Fang, K.-T., Hickernell, F. J., and Niederreiter, H., editors, *Monte Carlo and Quasi-Monte Carlo Methods 2000*, pages 86–102, Berlin. Springer-Verlag.
- Nishimura, T. (2000). Tables of 64-bit Mersenne twisters. *ACM Transactions on Modeling and Computer Simulation*, 10(4):348–357.
- Panneton, F. and L’Ecuyer, P. (2004). Random number generators based on linear recurrences in F_{2^w} . In Niederreiter, H., editor, *Monte Carlo and Quasi-Monte Carlo Methods 2002*, pages 367–378, Berlin. Springer-Verlag.
- Read, T. R. C. and Cressie, N. A. C. (1988). *Goodness-of-Fit Statistics for Discrete Multivariate Data*. Springer Series in Statistics. Springer-Verlag, New York.
- Rukhin, A., Soto, J., Nechvatal, J., Smid, M., Barker, E., Leigh, S., Levenson, M., Vangel, M., Banks, D., Heckert, A., Dray, J., and Vo, S. (2001). A statistical test suite for random and pseudorandom number generators for cryptographic applications. NIST special publication 800-22, National Institute of Standards and Technology (NIST), Gaithersburg, Maryland, USA. See <http://csrc.nist.gov/rng/>.
- Tausworthe, R. C. (1965). Random numbers generated by linear recurrence modulo two. *Mathematics of Computation*, 19:201–209.
- Tezuka, S. (1995). *Uniform Random Numbers: Theory and Practice*. Kluwer Academic Publishers, Norwell, Mass.
- Tezuka, S. and L’Ecuyer, P. (1991). Efficient and portable combined Tausworthe random number generators. *ACM Transactions on Modeling and Computer Simulation*, 1(2):99–112.
- Tezuka, S., L’Ecuyer, P., and Couture, R. (1994). On the add-with-carry and subtract-with-borrow random number generators. *ACM Transactions of Modeling and Computer Simulation*, 3(4):315–331.
- Tootill, J. P. R., Robinson, W. D., and Eagle, D. J. (1973). An asymptotically random Tausworthe sequence. *Journal of the ACM*, 20:469–481.
- Vattulainen, I., Ala-Nissila, T., and Kankaala, K. (1995). Physical models as tests of randomness. *Physical Review E*, 52(3):3205–3213.
- von Neumann, J. (1951). Various techniques used in connection with random digits. In Householder, A. S. et al., editors, *The Monte Carlo Method*,

- number 12 in National Bureau of Standards Applied Mathematics Series, pages 36–38.
- Wagner, M. A. F. and Wilson, J. R. (1996). Using univariate Bézier distributions to model simulation input processes. *IIE Transactions*, 28:699–711.
- Walker, A. J. (1974). New fast method for generating discrete random numbers with arbitrary frequency distributions. *Electronic Letters*, 10:127–128.
- Walker, A. J. (1977). An efficient method for generating discrete random variables with general distributions. *ACM Transactions on Mathematical Software*, 3:253–256.
- Wang, D. and Compagner, A. (1993). On the use of reducible polynomials as random number generators. *Mathematics of Computation*, 60:363–374.
- Wegenkittl, S. and Matsumoto, M. (1999). Getting rid of correlations among pseudorandom numbers: Discarding versus tempering. *ACM Transactions on Modeling and Computer Simulation*, 9(3):282–294.
- Wu, P.-C. (1997). Multiplicative, congruential random number generators with multiplier $\pm 2^{k_1} \pm 2^{k_2}$ and modulus $2^p - 1$. *ACM Transactions on Mathematical Software*, 23(2):255–265.

Index

- (t, m, s) -net, 16
- p -value, 21
- acceptance-complement method, 26
- add-with-carry, 14
- alias method, 26
- asymptotically random, 16
- automatic methods, 24, 29
- binary search, 25
- birthday spacings test, 22
- characteristic polynomial, 15, 17–19
- collision test, 21, 22
- combined generators, 12, 19
- composition method, 28
- convolution method, 28
- discrepancy, 5
- dual lattice, 8, 17
- entropy, 21
- equidissection, 15
- equidistribution, 15
- generalized feedback shift register (GFSR), 15, 18
- goodness-of-fit, 5
- hat function, 27
- inversion method, 2, 24
- kernel density estimation, 26
- lagged-Fibonacci generator, 10
- lattice, 7, 16, 22
- linear congruential generator (LCG), 7, 14, 22
- linear feedback shift register (LFSR), 15, 17, 19, 22
- linear recurrence, 7
- linear recurrence modulo 2, 14, 15
- linear recurrence with carry, 13
- matrix linear recurrence, 14
- maximally equidistributed, 16, 19
- Mersenne twister, 15, 18, 19, 23
- Monte Carlo methods, 1
- multiple recursive generator (MRG), 7
- multiple recursive matrix generator, 19
- multiply-with-carry generator, 13
- Poisson distribution, 21
- Poisson process, 27
- polynomial lattice, 17
- polynomial LCG, 18
- primitive polynomial, 7, 15
- pseudorandom number generator, 1
- random number generator, 1
 - approximate factoring, 11
 - combined generators, 12, 19, 20, 23
 - definition, 3
 - figure of merit, 9, 16
 - floating-point implementation, 11
 - implementation, 10, 20
 - jumping ahead, 4, 13, 15
 - non-uniform, 23

- nonlinear, 20
- period length, 3, 15, 18
- physical device, 2
- power-of-two modulus, 12
- powers-of-two-decomposition, 11
- purely periodic, 3
- quality criteria, 3, 23
- seed, 3
- state, 3
- statistical tests, 5, 21
- streams and substreams, 4, 23
- ratio-of-uniforms method, 28
- rejection method, 26, 28
- serial test, 21
- spectral test, 8
- squeeze function, 27
- subtract-with-borrow, 14
- Tausworthe generator, 15, 17
- tempering, 18
- thinning, 27
- transformed density rejection, 27
- twisted GFSR, 15, 18, 19
- uniform distribution, 1
- uniformity measure, 4, 15
- unpredictability, 6
- variance reduction, 24

