

Tina's Random Number Generator Library

version 4.3

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Contents

1	TRNG in a nutshell	4
1.1	Introduction	4
1.2	History	5
2	Pseudo-random numbers for parallel Monte Carlo simulations	6
2.1	Pseudo-random numbers	6
2.2	General parallelization techniques for PRNGs	6
2.3	Playing fair	8
2.4	Linear feedback shift register sequences	9
2.4.1	Parallelization of LFSR sequences	10
2.4.2	Choice of modulus	12
2.5	YARN sequences	13
3	Basic concepts	15
3.1	Random number engines	15
3.2	Random number distributions	18
4	TRNG classes	21
4.1	Random number engines	21
4.1.1	Linear congruential generator	22
4.1.2	Multiple recursive generators	24
4.1.3	YARN generators	30
4.1.4	Lagged Fibonacci generators	37
4.2	Random number distributions	40
4.2.1	Uniform distributions	41
4.2.2	Exponential distribution	45
4.2.3	Normal distribution	46
4.2.4	Cauchy distribution	49
4.2.5	Logistic distribution	50
4.2.6	Lognormal distribution	52
4.2.7	Pareto distribution	53
4.2.8	Power-law distribution	54
4.2.9	Tent distribution	55
4.2.10	Weibull distribution	57
4.2.11	Extreme value distribution	58
4.2.12	Γ -distribution	59
4.2.13	χ^2 -distribution	60
4.2.14	Rayleigh distribution	62
4.2.15	Bernoulli distribution	63

Contents

4.2.16	Binomial distribution	64
4.2.17	Geometric distribution	66
4.2.18	Poisson distribution	67
4.2.19	Discrete distribution	68
4.3	Function template generate_canonical	69
5	Installation	70
6	Examples	72
6.1	Hello world!	72
6.2	Hello parallel world!	74
6.2.1	Block splitting	74
6.2.2	Leapfrog	76
6.2.3	Block splitting or leapfrog?	78
6.3	Using TRNG with STL and Boost	82
7	Implementation details and efficiency	86
7.1	Efficient modular reduction	86
7.2	Fast delinearization	88
7.3	Performance	88
8	Quality	90
9	Frequently asked questions	99
	Bibliography	101

1 TRNG in a nutshell

1.1 Introduction

The Monte Carlo method is a widely used and commonly accepted simulation technique in physics, operations research, artificial intelligence, and other fields, and pseudo-random numbers (PRNs) are its key resource. All Monte Carlo simulations include some sort of averaging independent samples, a calculation that is embarrassingly parallel. Hence it is no surprise that more and more large scale simulations are run on parallel systems like networked workstations or clusters. For each Monte Carlo simulation the quality of the PRN generator (PRNG) is a crucial factor. In a parallel environment the quality of a PRNG is even more important than in a non parallel environment to some extent, because feasible sample sizes are easily $10 \dots 10^4$ times as large as on a sequential machine. The main problem is the parallelization of the PRNG itself.

Application programmers and scientists need not to grapple with all the technical details of pseudo-random number generation if a PRNG library is used. The following requirements are frequently demanded from a library for (parallel) pseudo-random number generation:

- The library should provide a set of different interchangeable algorithms for pseudo-random number generation.
- For each algorithm different well tested parameter sets should be provided that guarantee a long period and good statistical properties.
- The internal state of a PRNG can be saved for later use and restored. This makes it possible to stop a simulation and to carry on later.
- PRNGs have to support block splitting and leapfrog, see section 2.1.
- The library should provide methods for generating random variables with various distributions, uniform and non-uniform.
- The library should be implemented in a portable, speed-optimized fashion.

If these are also your requirements for a PRNG library, you should go with Tina's Random Number Generator Library.

Tina's Random Number Generator Library (TRNG) is a state of the art C++ pseudo-random number generator library for sequential and parallel Monte Carlo simulations. Its design principles are based on a proposal [5] for an extensible random number generator facility, that will be part of the forthcoming revision of the ISO C++ standard. The TRNG library features an object oriented design, is easy to use and has been speed optimized. Its implementation does not depend on any communication library or hardware architecture. TRNG is suited for shared memory as well as for distributed memory computers and may be used in any parallel programming environment, e. g. Message Passing Standard or OpenMP. All generators, that

are implemented by TRNG, have been subjected to thorough statistical tests in sequential and parallel setups, see also section 8.

This reference is organized as follows. In chapter 2 we present some basic techniques for parallel random number generation, chapter 3 introduces the basic concepts of TRNG, whereas chapter 4 describes all classes of TRNG in detail. In chapter 5 we give installation instructions, and chapter 6 presents some example programs, that demonstrate the usage of TRNG in sequential as well as in parallel Monte Carlo applications. Chapter 7 deals with some implementation details and performance issues. We complete the TRNG reference with the presentation of some statistical tests of the PRNGs of TRNG in chapter 8 and answer some FAQs in chapter 9.

This manual can be read in several ways. You might read this manual chapter by chapter from the beginning to its end. Impatient readers should read at least chapter 2 to familiarize themselves with some basic terms, that are used in this text, before they jump to chapter 5 and chapter 6. These chapters deal with the installation and give some example code. Chapters 3 and 4 are mainly for reference and the reader will come back to them again and again.

The TRNG manual is not written as an introduction to the Monte Carlo method. It is assumed that the reader already knows the basic concepts of Monte Carlo. Novices in the Monte Carlo business find further information in various textbooks on this topic [12, 49, 42, 22, 21, 38].

1.2 History

TRNG started in 2000 as a student research project. Its implementation as well as its technical design has changed several times. Starting with version 4.0 we adopted the interface proposed by [5].

Version 4.0 Initial release of TRNG that implements the interface proposed by [5].

Version 4.1 Additive and exclusive-or lagged Fibonacci generators with two and four feedback taps have been added to the set of PRNGs. Lagged Fibonacci generators do not provide any splitting facilities. TRNG implements the template function `generate_cononical` introduced by [5].

Version 4.2 Documentation has been revised. Minor bug-fixes to lagged Fibonacci generators.

Version 4.3 Rayleigh distribution and class for correlated normal distributed random numbers added. Changed default parameter sets for generators `mrg3s`, `mrg5s`, `yarn3s`, and `yarn5s`. The new parameter sets perform better in the spectral test.

2 Pseudo-random numbers for parallel Monte Carlo simulations

2.1 Pseudo-random numbers

Monte Carlo methods are a class of computational algorithms for simulating the behavior of various physical and mathematical systems by a stochastic process. While simulating such a stochastic process on a computer, large amounts of random numbers are consumed. Actually, a computer as a deterministic machine is not able to generate random digits. John von Neumann, pioneer in Monte Carlo simulation, summarized this problem in his famous quote:

“Anyone who considers arithmetical methods of producing random digits is, of course, in a state of sin.”

For computer simulations we have to content ourselves with something weaker than random numbers, namely *pseudo-random* numbers. We define a stream of PRNs r_i in the following in an informal manner:

- PRNs are generated by a deterministic rule.
- A stream of PRNs r_i cannot be distinguished from a true random sequence by means of practicable methods applying a *finite* set of statistical tests on *finite* samples.

Almost all PRNGs produce a sequence r_0, r_1, r_2, \dots of PRNs by a recurrence

$$r_i = f(r_{i-1}, r_{i-2}, \dots, r_{i-k}), \quad (2.1)$$

and the art of random number generation lies in the design of the function $f(\cdot)$.

The objective in PRNG design is to find a transition algorithm $f(\cdot)$ that yields a PRNG with a long period and good statistical properties within the stream of PRNs. Statistical properties of a PRNG may be investigated by theoretical or empirical means, see [20]. But experience shows, there is nothing like an ideal PRNG. A PRNG may behave like a perfect source of randomness in one kind of Monte Carlo simulation, whereas it may suffer from significant statistical correlations if it is used in another context, which makes the Monte Carlo simulation unreliable.

2.2 General parallelization techniques for PRNGs

In parallel applications we need to generate streams $t_{j,i}$ of random numbers. Streams are numbered by $j = 0, 1, \dots, p - 1$, where p is the number of processes. We require statistical independency of the $t_{j,i}$ within each stream and between streams as well. Four different parallelization techniques are used in practice:

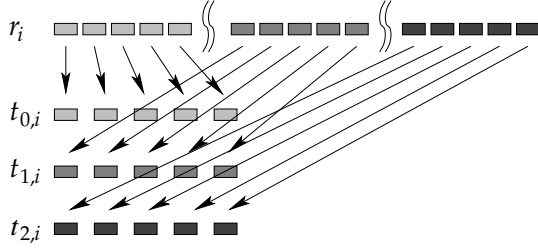


Figure 2.1: Parallelization by block splitting.

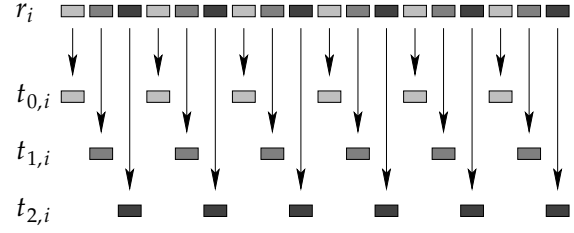


Figure 2.2: Parallelization by leapfrogging.

Random seeding: All processes use the same PRNG but a different “random” seed. The hope is that they will generate non-overlapping and uncorrelated subsequences of the original PRNG. This hope, however, has no theoretical foundation. Random seeding is a violation of Donald Knuth’s advice “Random number generators should not be chosen at random” [20].

Parameterization: All processes use the same type of generator but with different parameters for each processor. Example: linear congruential generators with additive constant b_j for the j th stream [46]:

$$t_{j,i} = a \cdot t_{j,i-1} + b_j \bmod 2^e, \quad (2.2)$$

where b_j is the $(j + 2)$ th prime number. Another variant uses different multipliers a for different streams [33]. The theoretical foundation of these methods is weak, and empirical tests have revealed serious correlations between streams [36]. On massive parallel system you may need thousands of parallel streams, and it is not trivial to find a type of PRNG with thousands of “well tested” parameter sets.

Block splitting: Let M be the maximum number of calls to a PRNG by each processor, and let p be the number of processes. Then we can split the sequence r_i of a sequential PRNG into consecutive blocks of length M such that

$$\begin{aligned} t_{0,i} &= r_i \\ t_{1,i} &= r_{i+M} \\ &\dots \\ t_{p-1,i} &= r_{i+M(p-1)}. \end{aligned} \quad (2.3)$$

This method works only if we know M in advance or can at least safely estimate an upper bound for M . To apply block splitting it is necessary to jump from the i th random number to the $(i + M)$ th number without calculating all the numbers in between, which cannot be done efficiently for many PRNGs. A potential disadvantage of this method is that long range correlations, usually not observed in sequential simulations, may become short range correlations between sub-streams [37, 9]. Block splitting is illustrated in Figure 2.1.

Leapfrog: The leapfrog method distributes a sequence r_i of random numbers over p processes

by decimating this base sequence such that

$$\begin{aligned} t_{0,i} &= r_{pi} \\ t_{1,i} &= r_{pi+1} \\ &\dots \\ t_{p-1,i} &= r_{pi+(p-1)} \cdot \end{aligned} \tag{2.4}$$

Leapfrogging is illustrated in Figure 2.2. It is the most versatile and robust method for parallelization and it does not require an a priori estimate of how many random numbers will be consumed by each processor. An efficient implementation requires a PRNG that can be modified to generate directly only every p th element of the original sequence. Again this excludes many popular PRNGs.

At first glance block splitting and leapfrog seem to be quite different approaches. But in fact, these are closely related to each other. Because if leapfrog is applied to any *finite* base sequence the leapfrog sequences are cyclic shifts of each other. Consider an arbitrary sequence r_i with period T . If $\gcd(T, p) = 1$, all leapfrog sequences $t_{1,i}, t_{2,i}, \dots, t_{p,i}$ are cyclic shifts of each other, i. e., for every pair of leapfrog sequences $t_{j_1,i}$ and $t_{j_2,i}$ of a common base sequence r_i with period T there is a constant s , such that $t_{j_1,i} = t_{j_2,i+s}$ for all i , and s is at least $\lfloor T/p \rfloor$. Furthermore, if $\gcd(T, p) = d > 1$, the period of each leapfrog sequence equals T/d and there are d classes of leapfrog sequences. Within a class of leapfrog sequences there are p/d sequences, each sequence is just a cyclic shift of another and the size of the shift is at least $\lfloor T/p \rfloor$.

The first two methods, random seeding and parameterization, have little or no theoretical backup, but their weakest point is yet another. The results of a simulation should not depend on the number of processors it runs on. Leapfrog and block splitting do allow to organize simulations such that the same random numbers are used independently of the number of processors. With parameterization or random seeding the results will always depend on the parallelization, see section 6.2 for details. PRNGs that do not support leapfrog and block splitting should not be used in parallel simulations.

2.3 Playing fair

We say that a parallel Monte Carlo simulation *plays fair*, if its outcome is strictly independent of the underlying hardware. Fair play implies the use of the same PRNs in the same context, independently of the number of parallel processes. It is mandatory for debugging, especially in parallel environments where the number of parallel processes varies from run to run, but another benefit of playing fair is even more important: Fair play guarantees that the quality of a PRNG with respect to an application does not depend on the degree of parallelization.

Obviously the use of parameterization or random seeding prevent a simulation from playing fair. Leapfrog and block splitting, on the other hand, do allow to use the same PRNs within the same context independently of the number of parallel streams.

Consider the site percolation problem. A site in a lattice of size N is occupied with some probability, and the occupancy is determined by a PRN. M random configurations are generated. A naive parallel simulation on p processes could split a base sequence into p leapfrog streams and having each process generate $\approx M/p$ lattice configurations, independently of the other processes. Obviously this parallel simulation is not equivalent to its sequential version,

that consumes PRNs from the base sequence to generate one lattice configuration after another. The effective shape of the resulting lattice configurations depends on the number of processes. This parallel algorithm does not play fair.

We can turn the site percolation simulation into a fair playing algorithm by leapfrogging on the level of lattice configurations. Here each process consumes distinct contiguous blocks of PRNs from the sequence r_i , and the workload is spread over p processors in such a way, that each process analyzes each p th lattice. If we number the processes by their rank i from 0 to $p - 1$ and the lattices from 0 to $M - 1$, each process starts with a lattice whose number equals its own rank. That means process i has to skip $i \cdot N$ PRNs from the sequence r_i before the first lattice configuration is generated. Thereafter each process can skip $p - 1$ lattices, i. e., $(p - 1) \cdot N$ PRNs and continue with the next lattice. In section 6.2 we investigate this approach in more detail and will give further examples of fair playing Monte Carlo algorithms and their implementation.

Organizing simulation algorithms such that they play fair is not always as easy as in the above example, but with a little effort one can achieve fair play in more complicated situations, too. This may require the combination of block splitting and the leapfrog method, or iterated leapfrogging. Sometimes it is also necessary to use more than one stream of PRNs per process, e. g. in the Swendsen Wang cluster algorithm [53, 42] one may use one PRNG to construct the bond percolation clusters and another PRNG to decide if a cluster has to be flipped.

2.4 Linear feedback shift register sequences

The majority of the PRNG algorithms that are implemented by TRNG are based on so-called linear feedback shift register sequences. Therefore we review some of its mathematical properties in this section. Readers how do not want to bother with mathematical details might skip this and the next section on YARN generators and may come back later.

Numerous recipes for $f(\cdot)$ in (2.1) have been discussed in the literature, see [20, 27] and references therein. One of the oldest and most popular PRNGs is the linear congruential generator (LCG)

$$r_i = a \cdot r_{i-1} + b \bmod m \quad (2.5)$$

introduced by Lehmer [29]. The additive constant b may be zero. Knuth [19] proposed a generalization of Lehmer's method known as multiple recurrence generator (MRG)

$$r_i = a_1 r_{i-1} + a_2 r_{i-2} + \dots + a_n r_{i-n} \bmod m. \quad (2.6)$$

In the theory of finite fields a sequence of type (2.6) is called *linear feedback shift register sequence*, or LFSR sequence for short. Note that a LFSR sequence is fully determined by specifying n coefficients (a_1, a_2, \dots, a_n) plus n initial values (r_1, r_2, \dots, r_n) . There is a wealth of rigorous results on LFSR sequences that can (and should) be used to construct a good PRNG. Here we only discuss a few but important facts without proofs. A detailed discussion of LFSR sequences including proofs can be found in [14, 17, 30, 31, 11, 55].

Since the all zero tuple $(0, 0, \dots, 0)$ is a fixed-point of (2.6), the maximum period of a LFSR sequence cannot exceed $m^n - 1$. The following theorem tells us precisely how to choose the coefficients (a_1, a_2, \dots, a_n) to achieve this period [20]:

Theorem 1 The LFSR sequence (2.6) over \mathbb{F}_m has period $m^n - 1$, if and only if the characteristic polynomial

$$f(x) = x^n - a_1x^{n-1} - a_2x^{n-2} - \dots - a_n \quad (2.7)$$

is *primitive* modulo m .

A monic polynomial $f(x)$ of degree n over \mathbb{F}_m is primitive modulo m , if and only if it is irreducible (i. e., cannot be factorized over \mathbb{F}_m), and if it has a primitive element of the extension field \mathbb{F}_{m^n} as one of its roots. The number of primitive polynomials of degree n modulo m is equal to $\phi(m^n - 1)/n = \mathcal{O}(m^n / (n \ln(n \ln m)))$ [54], where $\phi(x)$ denotes Euler's totient function. As a consequence a random polynomial of degree n is primitive modulo m with probability $\simeq 1/(n \ln(n \ln m))$, and finding primitive polynomials reduces to testing whether a given polynomial is primitive. The latter can be done efficiently, if the factorization of $m^n - 1$ is known [17], and most computer algebra systems offer a procedure for this test.

Theorem 2 Let r_i be an LFSR sequence (2.6) with a primitive characteristic polynomial. Then each k -tuple $(r_{i+1}, \dots, r_{i+k})$ occurs m^{n-k} times per period for $k \leq n$ (except the all zero tuple for $k = n$).

From this theorem it follows that, if a k -tuple of consecutive numbers with $k \leq n$ is chosen randomly from a LFSR sequence, the outcome is uniformly distributed over all possible k -tuples in \mathbb{F}_m . This is exactly what one would expect from a truly random sequence. In terms of Compagner's ensemble theory tuples of size less than or equal to n drawn from a LFSR sequence with primitive characteristic polynomial are indistinguishable from truly random tuples [6, 7].

Theorem 3 Let r_i be an LFSR sequence (2.6) with period $T = m^n - 1$ and let α be a complex m th root of unity and $\bar{\alpha}$ its complex conjugated. Then

$$C(h) := \sum_{i=1}^T \alpha^{r_i} \cdot \bar{\alpha}^{r_{i+h}} = \begin{cases} T & \text{if } h = 0 \bmod T \\ -1 & \text{if } h \neq 0 \bmod T \end{cases} \quad (2.8)$$

$C(h)$ can be interpreted as autocorrelation function of the sequence, and Theorem 3 tells us that LFSR sequences with maximum period have autocorrelations that are very similar to the autocorrelations of a random sequence with period T . Together with the nice equidistribution properties (Theorem 2) this qualifies LFSR sequences with maximum period as *pseudo-noise sequences*, a term originally coined by Golomb for binary sequences [14, 17].

2.4.1 Parallelization of LFSR sequences

As a matter of fact, LFSR sequences do support leapfrog and block splitting very well. Block splitting means basically jumping ahead in a PRN sequence. In the case of LFSR sequences this can be done quite efficiently. Note, that by introducing a companion matrix A the linear recurrence (2.6) can be written as a vector matrix product.

$$\begin{pmatrix} r_{i-(n-1)} \\ \vdots \\ r_{i-1} \\ r_i \end{pmatrix} = \underbrace{\begin{pmatrix} 0 & 1 & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & 1 \\ a_n & a_{n-1} & \dots & a_1 \end{pmatrix}}_A \begin{pmatrix} r_{i-n} \\ \vdots \\ r_{i-2} \\ r_{i-1} \end{pmatrix} \bmod m \quad (2.9)$$

From this formula it follows immediately that the M -fold successive iteration of (2.6) may be written as

$$\begin{pmatrix} r_{i-(n-1)} \\ \vdots \\ r_{i-1} \\ r_i \end{pmatrix} = A^M \begin{pmatrix} r_{i-M-(n-1)} \\ \vdots \\ r_{i-M-1} \\ r_{i-M} \end{pmatrix} \bmod m. \quad (2.10)$$

Matrix exponentiation can be accomplished in $\mathcal{O}(n^3 \ln M)$ steps via binary exponentiation (also known as exponentiation by squaring).

Implementing leapfrogging efficiently is less straightforward. Calculating $t_{j,i} = r_{pi+j}$ via

$$\begin{pmatrix} r_{pi+j-(n-1)} \\ \vdots \\ r_{pi+j-1} \\ r_{pi+j} \end{pmatrix} = A^p \begin{pmatrix} r_{p(i-1)+j-(n-1)} \\ \vdots \\ r_{p(i-1)+j-1} \\ r_{p(i-1)+j} \end{pmatrix} \bmod m \quad (2.11)$$

is no option, because A^p is usually a dense matrix, in which case calculating a new element from the leapfrog sequence requires $\mathcal{O}(n^2)$ operations instead of $\mathcal{O}(n)$ operations in the base sequence.

The following theorem assures that the leapfrog subsequences of LFSR sequences are again LFSR sequences [17]. This will provide us with a very efficient way to generate leapfrog sequences.

Theorem 4 Let r_i be a LFSR sequence based on a primitive polynomial of degree n with period $m^n - 1$ (pseudo-noise sequence) over \mathbb{F}_m , and let (t) be the decimated sequence with lag $p > 0$ and offset j , e. g.

$$t_{j,i} = r_{pi+j}. \quad (2.12)$$

Then $t_{j,i}$ is a LFSR sequence based on a primitive polynomial of degree n , too, if and only if p and $m^n - 1$ are coprime, e. g. $\gcd(m^n - 1, p) = 1$. In addition, r_i and $t_{j,i}$ are not just cyclic shifts of each other, except when

$$p = m^h \bmod (m^n - 1) \quad (2.13)$$

for some $0 \leq h < n$. If $\gcd(m^n - 1, p) > 1$ the sequence $t_{j,i}$ is still a LFSR sequence, but not a pseudo-noise sequence.

It is not hard to find prime numbers m such that $m^n - 1$ has very few (and large) prime factors. For such numbers, the leapfrog method yields pseudo-noise sequences for any reasonable number of parallel streams [3]. While Theorem 4 ensures that leapfrog sequences are not just cyclic shifts of the base sequence (unless (2.13) holds), the leapfrog sequences are cyclic shifts of each other, see section 2.2.

Theorem 4 tells us that all leapfrog sequences of a LFSR sequence of degree n can be generated by another LFSR of degree n or less. The following theorem gives us a recipe to calculate the coefficients (b_1, b_2, \dots, b_n) of the corresponding leapfrog feedback polynomial.

Theorem 5 Let t_i be a (periodic) LFSR sequence over the field \mathbb{F}_m and $f(x)$ its characteristic polynomial of degree n . Then the coefficients (b_1, b_2, \dots, b_n) of $f(x)$ can be computed from $2n$

successive elements of t_i by solving the linear system

$$\begin{pmatrix} t_{i+n} \\ t_{i+n+1} \\ \vdots \\ t_{i+2n-1} \end{pmatrix} = \begin{pmatrix} t_{i+n-1} & \cdots & t_{i+1} & t_i \\ t_{i+n} & \cdots & t_{i+2} & t_{i+1} \\ \vdots & \ddots & \vdots & \vdots \\ t_{i+2n-2} & \cdots & t_{i+n} & t_{i+n-1} \end{pmatrix} \begin{pmatrix} b_1 \\ b_2 \\ \vdots \\ b_n \end{pmatrix} \pmod{m} \quad (2.14)$$

over \mathbb{F}_m .

Starting from the base sequence we determine $2n$ values of the sequence t_i by applying the leapfrog rule. Then we solve (2.14) by Gaussian elimination to get the characteristic polynomial for a new LFSR generator that yields the elements of the leapfrog sequence directly with each call. If the matrix in (2.14) is singular, the linear system has more than one solution, and it is sufficient to pick one of them. In this case it is always possible to generate the leapfrog sequence by a LFSR of degree less than the degree of the original sequence.

2.4.2 Choice of modulus

LFSR sequences can be defined over any prime field. In particular LFSR sequences over \mathbb{F}_2 with sparse feedback polynomials are popular sources of PRNs [18, 56, 20] and generators of this type can be found in various software libraries. This is due to the fact that multiplication in \mathbb{F}_2 is trivial, addition reduces to exclusive-or and the modulo operation comes for free. As a result, generators that operate in \mathbb{F}_2 are extremely fast. Unfortunately, these generators suffer from serious statistical defects [10, 15, 51, 56] that can be blamed to the small size of the underlying field [1]. In parallel applications we have the additional drawback, that, if the leapfrog method is applied to a LFSR sequence with sparse characteristic polynomial, the new sequence will have a dense polynomial. The computational complexity of generating values of the LFSR sequence grows from $\mathcal{O}(1)$ to $\mathcal{O}(n)$. Remember that for generators in \mathbb{F}_2 , n is typically of order 1000 or even larger to get a long period $2^n - 1$ and reasonable statistical properties.

The theorems and parallelization techniques we have presented so far do apply to LFSR sequences over any finite field \mathbb{F}_m . Therefore we are free to choose the prime modulus m . In order to get maximum entropy on the macrostate level [39] m should be as large as possible. A good choice is to set m to a value that is of the order of the largest representable integer of the computer. If the computer deals with e -bit registers, we may write the modulus as $m = 2^e - k$, with k reasonably small. In fact if $k(k+2) \leq m$ modular reduction can be done reasonably fast by a few bit-shifts, additions and multiplications, see chapter 7. Furthermore a large modulus allows us to restrict the degree of the LFSR to rather small values, e. g. $n \approx 4$, while the PRNG has a large period and good statistical properties.

In accordance with Theorem 4 a leapfrog sequence of a pseudo-noise sequence is a pseudo-noise sequence, too, if and only if its period $m^n - 1$ and the lag p are coprime. For that reason $m^n - 1$ should have a small number of prime factors. It can be shown that $m^n - 1$ has at least three prime factors and if the number of prime factors does not exceed three, then m is necessarily a Sophie-Germain Prime and n a prime larger than two [3].

To sum up, the modulus m of a LFSR sequence should be a Sophie-Germain Prime, such that $m^n - 1$ has not more than three prime factors and such that $m = 2^e - k$ and $k(k+2) \leq m$ for some integers e and k .

2.5 YARN sequences

LFSR sequences over prime fields with a large prime modulus seem to be ideally suited as PRNGs. They have, however, a well known weakness. When used to sample coordinates in d -dimensional space, pseudo-noise sequences cover every point for $d < n$, and every point except $(0, 0, \dots, 0)$ for $d = n$. For $d > n$ the set of positions generated is obviously sparse, and the linearity of the production rule (2.6) leads to the concentration of the sampling points on n -dimensional hyper-planes [16, 24], see also Figure 2.3. This phenomenon, first noticed by Marsaglia in 1968 [32], constitutes one of the well known tests of randomness applied to PRNGs, the so-called spectral test [20]. The spectral test checks the behavior of a generator when its outputs are used to form d -tuples. LFSR sequences do fail the spectral test in dimensions larger than the register size n . Closely related to this mechanism are the observed correlations in other empirical tests like the birthday spacings test and the collision test [26, 28]. Non-linear generators do quite well in all these tests, but compared to LFSR sequences they have much less nice and *provable* properties and they are not suited for fair playing parallelization.

To get the best of both worlds we propose a delinearization that preserves all the nice properties of linear pseudo-noise sequences:

Theorem 6 Let q_i be a pseudo-noise sequence in \mathbb{F}_m , and let g be a generating element of the multiplicative group \mathbb{F}_m^* . Then the sequence r_i with

$$r_i = \begin{cases} g^{q_i} \bmod m & \text{if } q_i > 0 \\ 0 & \text{if } q_i = 0 \end{cases} \quad (2.15)$$

is a pseudo-noise sequence, too.

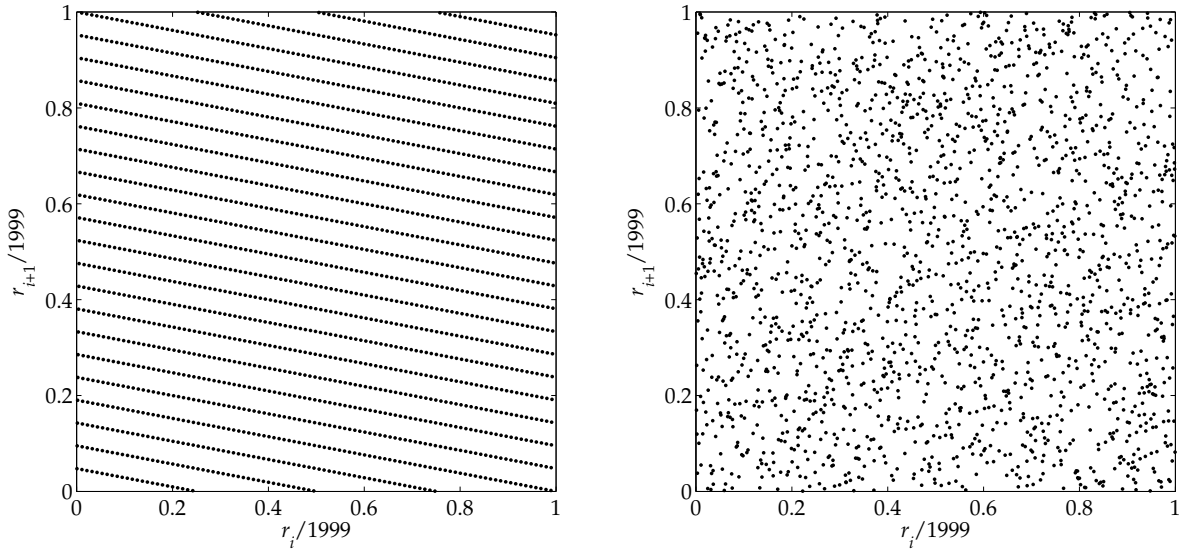


Figure 2.3: Exponentiation of a generating element in a prime field is an effective way to destroy the linear structures of LFSR sequences. Both pictures show the full period of the generator. Left: $r_i = 95 \cdot r_{i-1} \bmod 1999$. Right: $r_i = 1099^{q_i} \bmod 1999$ with $q_i = 95 \cdot q_{i-1} \bmod 1999$.

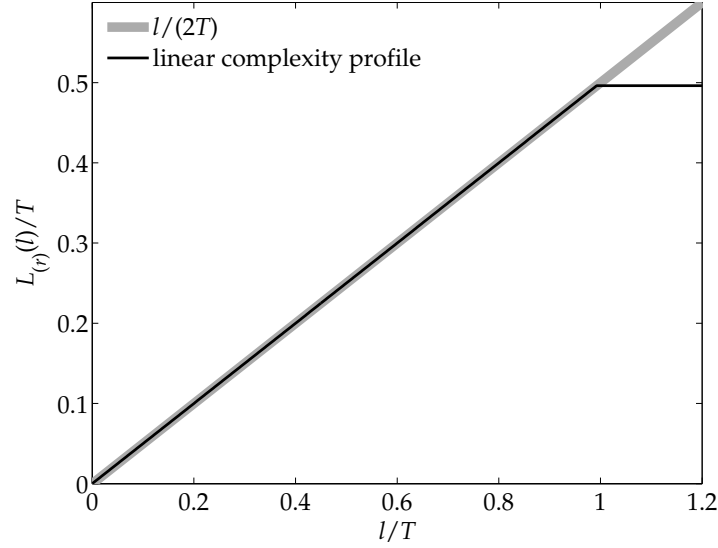


Figure 2.4: Linear complexity profile $L_{(r)}(l)$ of a YARN sequence, produced by the recurrence $q_i = 173 \cdot q_{i-1} + 219 \cdot q_{i-2} \bmod 317$ and $r_i = 151^{q_i} \bmod 317$. The period of this sequence equals $T = 317^2 - 1$.

The proof of this theorem is trivial: since g is a generator of \mathbb{F}_m^* , the map (2.15) is bijective. We call delinearized generators based on Theorem 6 YARN generators (yet another random number).

The linearity is completely destroyed by the map (2.15), see Figure 2.3. Let $L_{(r)}(l)$ denote the linear complexity of the subsequence (r_1, r_2, \dots, r_l) . This function is known as the linear complexity profile of r_i . For a truly random sequence it grows on average like $l/2$. Figure 2.4 shows the linear complexity profile $L_{(r)}(l)$ of a typical YARN sequence. It shows the same growth rate as a truly random sequence up to the point where more than 99% of the period have been considered. Sharing the linear complexity profile with a truly random sequence, we may say that the YARN generator is as nonlinear as it can get.

3 Basic concepts

The TRNG library consists of a loose bunch of classes. These classes can be divided into to kinds of classes, *random number engines* and *random number distributions*.

Random number engines are the workhorses of TRNG. Each random number engine implements some algorithm that is used to produce pseudo-random numbers. The notion of a random number engine as it is used by TRNG was introduced by [5] and it is a very general concept. For example the random number engine concept does not specify what kind of pseudo-random numbers (integers, floating point numbers or just bits) are generated. All random number engine classes of TRNG implement the concept of a random number engine as introduced in [5]. But in TRNG the notion of a random number engine is extended to a *parallel random number engine*. To fulfill the requirements of a parallel random number engine, a class has to fulfill all the requirements of a random number engine and in addition some further requirements that make them applicable for parallel Monte Carlo simulations. The random number engine concept and the parallel random number engine concept will be discussed in detail in section 3.1.

A random number engine is not very useful by itself. To write some real world Monte Carlo applications we need random number distribution classes, too. A random number distribution class converts the output of an arbitrary random number engine into a pseudo-random number with some specific distribution. The general concept of a random number distribution is discussed in section 3.2.

3.1 Random number engines

To be a random number engine, a class has to fulfill a set of requirements that we will summarize as follows, for details see [5]. A class X satisfies the requirements of a random number engine, if the expressions as shown in Table 3.1 are valid and have the indicated semantics. In that table and throughout this section,

- T is the type named by X 's associated `result_type`;
- t is a value of T ;
- u is a value of X , v is an lvalue of X , x and y are (possibly `const`) values of X ;
- s is a value of integral type;
- g is an lvalue, of a type other than X , that defines a zero-argument function object returning values of type `unsigned long`;
- os is an lvalue of the type of some class template specialization `std::basic_ostream<charT, traits>`; and

Table 3.1: Random number engine requirements.

expression	return type	pre/post-condition	complexity
<code>X::result_type</code>	<code>T</code>	<code>T</code> is an arithmetic type other than <code>bool</code> .	compile-time
<code>u()</code>	<code>T</code>	Sets the state to $u_{i+1} = \text{TA}(u_i)$ and returns $\text{GA}(u_i)$. If <code>X</code> is integral, returns a value in the closed interval $[\text{X}::\text{min}, \text{X}::\text{max}]$; otherwise, returns a value in the open interval $(0, 1)$.	amortized constant
<code>X::min</code>	<code>T</code> , if <code>X</code> is integral; otherwise <code>int</code> .	If <code>X</code> is integral, denotes the least value potentially returned by <code>operator()</code> ; otherwise denotes 0.	compile-time
<code>X::max</code>	<code>T</code> , if <code>X</code> is integral; otherwise <code>int</code> .	If <code>X</code> is integral, denotes the greatest value potentially returned by <code>operator()</code> ; otherwise denotes 1.	compile-time
<code>X()</code>		Creates an engine with the same initial state as all other default-constructed engines of type <code>X</code> .	\mathcal{O} (size of state)
<code>X(s)</code>		Creates an engine with initial state determined by <code>static_cast<unsigned long>(s)</code> .	\mathcal{O} (size of state)
<code>X(g)</code>		Creates an engine with initial state determined by the results of successive invocations of <code>g</code> . Throws what and when <code>g</code> throws.	\mathcal{O} (size of state)
<code>u.seed()</code>	<code>void</code>	post: <code>u==X()</code>	same as <code>X()</code>
<code>u.seed(s)</code>	<code>void</code>	post: <code>u==X(s)</code>	same as <code>X(s)</code>
<code>u.seed(g)</code>	<code>void</code>	post: If <code>g</code> does not throw, <code>u==v</code> , where the state of <code>v</code> is as if constructed by <code>X(g)</code> . Otherwise, the exception is re-thrown and the engine <code>s</code> state is deemed invalid. Thereafter, further use of <code>u</code> is undefined except for destruction or invoking a function that establishes a valid state.	same as <code>X(g)</code>
<code>x==y</code>	<code>bool</code>	With S_x and S_y as the infinite sequences of values that would be generated by repeated calls to <code>x()</code> and <code>y()</code> , respectively, returns <code>true</code> if $S_x = S_y$; returns <code>false</code> otherwise.	\mathcal{O} (size of state)
<code>x!=y</code>	<code>bool</code>	<code>!(x==y)</code>	\mathcal{O} (size of state)

- `is` is an lvalue of the type of some class template specialization `std::basic_istream<charT, traits>`.

A random number engine object `x` has at any given time a state x_i for some integer $i \geq 0$. Upon construction, a random number engine `x` has an initial state x_0 . The state of an engine may be established by invoking its constructor, seed member function, `operator=`, or a suitable `operator>>`.

The specification of each random number engine defines the size of its state in multiples of the size of its `result_type`, given as an integral constant expression. The specification of each random number engine also defines

- the *transition algorithm* `TA` by which the state x_i of an engine is advanced to its *successor*

Table 3.1: Random number engine requirements continued.

expression	return type	pre/post-condition	complexity
<code>os << x</code>	reference to the type of <code>os</code>	With <code>os.fmtflags</code> set to <code>std::ios_base::dec std::ios_base::fixed std::ios_base::left</code> and the fill character set to the space character, writes to <code>os</code> the textual representation of <code>x</code> 's current state. In the output, adjacent numbers are separated by one or more space characters. post: The <code>os.fmtflags</code> and fill character are unchanged.	\mathcal{O} (size of state)
<code>is >> v</code>	reference to the type of <code>is</code>	Sets <code>v</code> 's state as determined by reading its textual representation from <code>is</code> . If bad input is encountered, ensures that <code>v</code> 's state is unchanged by the operation and calls <code>is.setstate(std::ios::failbit)</code> (which may throw <code>std::ios::failure</code>). pre: The textual representation was previously written using an <code>os</code> whose imbued locale and whose type's template specialization arguments <code>charT</code> and <code>traits</code> were the same as those of <code>is</code> . post: The <code>is.fmtflags</code> are unchanged.	\mathcal{O} (size of state)

Table 3.2: Parallel random number engine requirements.

expression	return type	pre/post-condition	complexity
<code>split(s, p)</code>	void	pre: <code>s</code> and <code>p</code> are of type <code>unsigned int</code> with <code>s < p</code> . If <code>s ≥ p</code> an exception <code>std::invalid_argument</code> is thrown. post: Internal parameters of the random number engine are changed in such a way, that future calls to <code>operator()</code> will generate the <code>sth</code> sub-stream of <code>p</code> sub-streams. Sub-streams are numbered from 0 to <code>p - 1</code> . The complexity of <code>operator()</code> will not change.	polynomial in size of state, <code>p</code> and <code>s</code>
<code>jump2(s)</code>	void	pre: <code>s</code> is of type <code>unsigned int</code> . post: Internal state of the random number engine is changed in such a way, that the engine jumps 2^s steps ahead.	polynomial in size of state and <code>s</code>
<code>jump(s)</code>	void	pre: <code>s</code> is of type <code>unsigned long long</code> . post: Internal state of the random number engine is changed in such a way, that the engine jumps <code>s</code> steps ahead.	polynomial in size of state and the logarithm of <code>s</code>

state x_{i+1} , and

- the *generation algorithm* GA by which the state of an engine is mapped to a value of type `result_type`.

Furthermore, a random number engine shall fulfill the requirements of the concepts “Copy-Constructible” and of “Assignable”. That means roughly, random number engines support copy and assignment operations with the same semantic like build-in types as `int` or `double`. Copy construction and assignment shall each be of complexity \mathcal{O} (size of state).

Random number engine requirements had been adopted from [5]. For parallel Monte Carlo applications we extend the concept of a random number engine to a parallel random number engine. Such an engine has to meet all the requirements of a parallel random number engine and additionally the requirements shown in Table 3.2.

A parallel random number engine provides block splitting and leapfrog. Note that it is demanded that leapfrog is implemented in such a way, that the complexity of `operator()` will not depend on, in how many sub-streams a stream has been split. That means, a valid implementation of leapfrog will not just calculate all random numbers of a stream and then throw away bunches of numbers to derive the random numbers of a leapfrog sub-stream. This rather strong requirement restricts the number of pseudo-random number generator algorithms that are proper for parallel random number engines. But LFSR sequences and YARN generators, which had been discussed in sections 2.4 and 6, meet these conditions easily.

3.2 Random number distributions

To model the concept of a random number distribution a class has to fulfill a set of requirements that we will summarize as follows, refer to [5] for details. A class `X` satisfies the requirements of a random number distribution if the expressions shown in Table 3.3 are valid and have the indicated semantics, and if `X` and its associated types also satisfies all other requirements of this section. In that table and throughout this section,

- `T` is the type named by `X`’s associated `result_type`;
- `P` is the type named by `X`’s associated `param_type`;
- `u` is a value of `X` and `x` is a (possibly `const`) value of `X`;
- `glb` and `lub` are values of `T` respectively corresponding to the greatest lower bound and the least upper bound on the values potentially returned by `u`’s `operator()`, as determined by the current values of `u`’s parameters;
- `p` is a value of `P`;
- `e` is an lvalue of an arbitrary type that satisfies the requirements of a uniform random number generator;
- `os` is an lvalue of the type of some class template specialization `basic_ostream<charT, traits>`; and
- `is` is an lvalue of the type of some class template specialization `basic_istream<charT, traits>`.

3 Basic concepts

The specification of each random number distribution identifies an associated mathematical *probability density function* $p(z)$ or an associated discrete *probability function* $P(z_i)$. Such functions are typically expressed using certain externally supplied quantities known as the *parameters of the distribution*. Such distribution parameters are identified in this context by writing, for example, $p(z|a, b)$ or $P(z_i|a, b)$, to name specific parameters, or by writing, for example, $p(z|\{p\})$ or $P(z_i|\{p\})$, to denote the parameters p of a distribution taken as a whole.

Furthermore a random number distribution shall fulfill the requirements of the concepts “CopyConstructible” and of “Assignable”. That means roughly, random number distributions support copy and assignment operations with the same semantic like build-in types like `int` or `double`. Copy construction and assignment shall each be of complexity \mathcal{O} (size of state).

For each of the constructors of `X` taking arguments corresponding to parameters of the distribution, `P` shall have a corresponding constructor subject to the same requirements and taking arguments identical in number, type, and default values. Moreover, for each of the member functions of `X` that return values corresponding to parameters of the distribution, `P` shall have a corresponding member function with the identical name, type, and semantics.

Table 3.3: Random number distribution requirements.

expression	return type	pre/post-condition	complexity
<code>X::result_type</code>	T	T is an arithmetic type.	compile-time
<code>X::param_type</code>	P		compile-time
<code>X(p)</code>		Creates a distribution whose behavior is indistinguishable from that of a distribution newly constructed directly from the values used to construct p.	same as p's construction
<code>u.reset()</code>	void	Subsequent uses of u do not depend on values produced by e prior to invoking reset.	constant
<code>x.param()</code>	P	Returns a value p such that <code>X(p).param()==p</code> .	no worse than the complexity of X(p)
<code>u.param(p)</code>	void	post: <code>u.param() == p</code> .	no worse than the complexity of X(p)
<code>u(e)</code>	T	With <code>p=u.param()</code> , the sequence of numbers returned by successive invocations with the same object e is randomly distributed according to the associated $p(z \{p\})$ or $P(z_i \{p\})$ function.	amortized constant number of invocations of e
<code>u(e, p)</code>	T	The sequence of numbers returned by successive invocations with the same objects e and p is randomly distributed according to the associated $p(z \{p\})$ or $P(z_i \{p\})$ function	
<code>x.min()</code>	T	Returns glb.	constant
<code>x.max()</code>	T	Returns lub.	constant
<code>os << x</code>	reference to the type of os	Writes to os a textual representation for the parameters and the additional internal data of x. post: The <code>os.fmtflags</code> and fill character are unchanged.	
<code>is >> u</code>	reference to the type of is	Restores from is the parameters and additional internal data of u. If bad input is encountered, ensures that u's state is unchanged by the operation and calls <code>is.setstate(ios::failbit)</code> (which may throw <code>std::ios::failure</code>). pre: is provides a textual representation that was previously written using an os whose imbued locale and whose type's template specialization arguments <code>charT</code> and <code>traits</code> were the same as those of is. post: The <code>is.fmtflags</code> are unchanged.	

4 TRNG classes

In chapter 3 the abstract concepts of (parallel) random number engines and random number distributions had been introduced. Now we look at some actual realizations of these concepts. TRNG provides several (parallel) random number engines and random number distributions. Each engine and each distribution is implemented by its own class that resides in the name space `trng`.

4.1 Random number engines

In this section we give a detailed documentation of all random number engines. Each subsection describes the public interface of one random number engine and focuses on aspects that are specific for a particular random number engine. This includes extensions to the random number engine interface as well as algorithmic details. That part of the public interface, that is mandatory for each (parallel) random number engine, will not be discussed in detail. Read section 3.1 instead. Table 4.1 gives an overview over all random number engines of TRNG.

All classes that will be describe in this section model either a random number engine or a

Table 4.1: Random number engines of TRNG.

random number engine	description	concept
<code>trng::lcg64</code>	linear congruential generator with modulus 2^{64}	parallel random number engine
<code>trng::mrngn</code>	multiple recurrence generator based on a linear feedback shift register sequence over $\mathbb{F}_{2^{31}-1}$ of depth n	parallel random number engine
<code>trng::mrngns</code>	multiple recurrence generator based on a linear feedback shift register sequence over \mathbb{F}_m of depth n , with m being a Sophie-Germain Prime	parallel random number engine
<code>trng::yarnn</code>	YARN sequence based on a linear feedback shift register sequence over $\mathbb{F}_{2^{31}-1}$ of depth n	parallel random number engine
<code>trng::yarnns</code>	YARN sequence based on a linear feedback shift register sequence over \mathbb{F}_m of depth n , with m being a Sophie-Germain Prime	parallel random number engine
<code>trng::lagfibn xor</code>	lagged Fibonacci generator with n feedback taps and exclusive-or operation	random number engine
<code>trng::lagfibn plus</code>	lagged Fibonacci generator with n feedback taps and addition	random number engine

parallel random number engine and therefore fulfill the requirements introduced in section 3.1. But for convenience their interface provides even more. For example all random number engines model a *random number generator* as well. The notion of a random number generator had been introduced by the C++ Standard Template Library. A random number generator is a class that provides an `operator() (long)` that returns a uniformly distributed random integer larger than or equal to zero but less than its argument. That makes TRNG (parallel) random number engines applicable to the STL algorithm `std::random_shuffle`. Additionally TRNG (parallel) random number engines provide a function `name()` that returns a string with the name of the random number engine.

4.1.1 Linear congruential generator

The class `trng::lcg64` implements a linear congruential generator, for which the transition algorithm reads [29, 20]

$$r_{i+1} = a \cdot r_i + b \bmod 2^{64}.$$

The state of this generator at time i is given by r_i . Its period equals 2^{64} if and only if b is odd and $a \bmod 4 = 1$ [20]. The statistical properties of linear congruential generators depend crucial on the choice of the multiplier a , which has to be chosen carefully.

This linear congruential generator is the quick and dirty generator of TRNG. It's dammed fast, see section 7, but even for proper chosen parameters a and b the lower bits of r_i are less random than the higher order bits. The class `trng::lcg64` should be avoided whenever the randomness of lower bits have a significant impact to the simulation. In [23] L'Ecuyer warns about multiplicative linear congruential generators (MLCG) with $r_{i+1} = a \cdot r_i \bmod m$:

"If $m = 2^e$ where e is the number of bits on the computer word, and if one can use unsigned integers without overflow checking, the products modulo m are easy to compute: just discard the overflow. This is quick and simple. For that reason, MLCGs with moduli of this form are used abundantly in practice, despite their serious drawbacks. Some nuclear physicists, for instance, perform simulations that use billions of random numbers on supercomputers and are quite reluctant to give up using them [...]. Usually, they also generate many substreams in parallel. In view of the above remarks, all this appears dangerous. Perhaps some people like playing with fire."

The same warning applies if $b \neq 0$. In spite of its weakness this generator is well suited for a large classes of generic Monte Carlo schemes, e. g. simulating a (biased) coin or cluster Monte Carlo [10].

The class `trng::lcg64` is declared in the header file `trng/lcg64.hpp` and its public interface is given as follows:

```
namespace trng {
    class lcg64 {
    public:
```

First the necessary type, static class constants, and the call operator are declared.

```
typedef unsigned long long result_type;
result_type operator()() const;
static const result_type min;
static const result_type max;
```

4 TRNG classes

We also define some parameter and status classes that will be used internally and by the constructor.

```
class parameter_type;  
class status_type;
```

TRNG provides four parameter sets for a and b , which are chosen to give good statistical properties. Three of these are taken from [25], the default parameter set had been found by the author of TRNG.

$$a = 18\,145\,460\,002\,477\,866\,997, \quad b = 1$$

```
static const parameter_type Default;
```

$$a = 2\,862\,933\,555\,777\,941\,757, \quad b = 1$$

```
static const parameter_type LEcuyer1;
```

$$a = 3\,202\,034\,522\,624\,059\,733, \quad b = 1$$

```
static const parameter_type LEcuyer2;
```

$$a = 3\,935\,559\,000\,370\,003\,845, \quad b = 1$$

```
static const parameter_type LEcuyer3;
```

An instance of class `trng::lcg64` can be instantiated by various constructors as specified for a random number engine. Additionally a non-default parameter set may be given.

```
explicit lc64(parameter_type=Default);  
explicit lc64(unsigned long, parameter_type=Default);  
template<typename gen>  
explicit lc64(gen &, parameter_type P=Default);
```

Class `trng::lcg64` provides all necessary seeding functions (see Table 3.1) and an additional function that sets r_i .

```
void seed();  
void seed(unsigned long);  
template<typename gen>  
void seed(gen &);  
void seed(result_type);
```

The following three methods are necessary for a parallel random number engine.

```
void split(unsigned int, unsigned int);  
void jump2(unsigned int);  
void jump(unsigned long long);
```

Furthermore the class `trng::lcg64` provides a function that returns the string `lcg64` and an operator `operator()`.

```
static const char * name();  
long operator()(long) const;  
};
```

Random number engines are comparable and can be written to or read from a stream.

```

bool operator==(const lcg64 &, const lcg64 &);
bool operator!=(const lcg64 &, const lcg64 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const lcg64 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, lcg64 &);
}

```

4.1.2 Multiple recursive generators

TRNG offers several multiple recursive generators based on LFSR sequences over prime fields \mathbb{F}_m with different numbers of feedback taps. These are implemented by the classes `trng::mrg2`, `trng::mrg3`, `trng::mrg3s`, `trng::mrg4`, `trng::mrg5`, and `trng::mrg5s`. Table 4.2 summarizes the key features of these classes. The transition algorithm of a multiple recursive generator with n feedback taps reads

$$r_i = a_1 \cdot r_{i-1} + a_2 \cdot r_{i-2} + \dots + a_n \cdot r_{i-n} \bmod m.$$

The state of this generator at time i is given by $(r_{i-1}, r_{i-2}, \dots, r_{i-n})$. See section 2.4 for details on LFSR sequences.

The prime modulus m that characterizes the prime field \mathbb{F}_m was either chosen as the Mersenne Prime (classes `trng::mrgn`) or a Sophie-Germain Prime such that $m^n - 1$ has as few prime factors as possible (classes `trng::mrgns`). The former choice gives us some performance benefits, see section 7.1, whereas the second has some theoretical advantages, see section 2.4.2.

The classes `trng::mrgn` and `trng::mrgns` implement the interface described in section 3.1. Each class defines some parameter and status classes that will be used internally and by the constructor. Furthermore for each generator several parameter sets are given, see Table 4.3. Most of the parameter sets are taken from [24] and chosen to give generators with good statistical properties.

An instance of a class `trng::mrgn` or `trng::mrgns` can be instantiated by various constructors as specified for a random number engine. Additionally a non-default parameter set may be chosen. The classes `trng::mrgn` and `trng::mrgns` provide all necessary seeding functions (see Table 3.1) and additionally a function that sets the internal state $(r_{i-1}, r_{i-2}, \dots, r_{i-n})$. This function should never be called with all arguments set to zero. The classes `trng::mrgn` and `trng::mrgns` model the concept of a parallel random number engine and therefore the methods

```

void split(unsigned int, unsigned int);
void jump2(unsigned int);
void jump(unsigned long long);

```

are implemented. Furthermore the classes `trng::mrgn` or `trng::mrgns` provide a function that returns a string with its name and an operator `operator()`. Random number engines are comparable and can be written to or read from a stream.

The detailed interface of the classes `trng::mrgn` or `trng::mrgns` is given as follows:

```

namespace trng {
    class mrg2 {

```


Table 4.2: Key features of multiple recursive generator classes.

class	header file	feedback taps n	prime field \mathbb{F}_m	period	return value of name()
trng::mrg2	trng/mrg2.hpp	2	$\mathbb{F}_{2^{31}-1}$	$m^2 - 1 \approx 2^{62} \approx 4.61 \cdot 10^{18}$	mrg2
trng::mrg3	trng/mrg3.hpp	3	$\mathbb{F}_{2^{31}-1}$	$m^3 - 1 \approx 2^{93} \approx 9.90 \cdot 10^{27}$	mrg3
trng::mrg3s	trng/mrg3s.hpp	3	$\mathbb{F}_{2^{31}-21069}$	$m^3 - 1 \approx 2^{93} \approx 9.90 \cdot 10^{27}$	mrg4s
trng::mrg4	trng/mrg4.hpp	4	$\mathbb{F}_{2^{31}-1}$	$m^4 - 1 \approx 2^{124} \approx 2.13 \cdot 10^{37}$	mrg4
trng::mrg5	trng/mrg5.hpp	5	$\mathbb{F}_{2^{31}-1}$	$m^5 - 1 \approx 2^{155} \approx 4.57 \cdot 10^{46}$	mrg5
trng::mrg5s	trng/mrg5s.hpp	5	$\mathbb{F}_{2^{31}-22641}$	$m^5 - 1 \approx 2^{155} \approx 4.57 \cdot 10^{46}$	mrg5s

Table 4.3: Parameter sets for multiple recursive generators.

parameter set	a_1	a_2	a_3	a_4	a_5
trng::mrg2::LEcuyer1	1 498 809 829	1 160 990 996			
trng::mrg2::LEcuyer2	46 325	1 084 587			
trng::mrg3::LEcuyer1	2 021 422 057	1 826 992 351	1 977 753 457		
trng::mrg3::LEcuyer2	1 476 728 729	0	1 155 643 113		
trng::mrg3::LEcuyer3	65 338	0	64 636		
trng::mrg3s::trng0	2 025 213 985	1 112 953 677	2 038 969 601		
trng::mrg3s::trng1	1 287 767 370	1 045 931 779	58 150 106		
trng::mrg4::LEcuyer1	2 001 982 722	1 412 284 257	1 155 380 217	1 668 339 922	
trng::mrg4::LEcuyer2	64 886	0	0	64 322	
trng::mrg5::LEcuyer1	107 374 182	0	0	0 104 480	
trng::mrg5s::trng0	1 053 223 373	1 530 818 118	1 612 122 482	133 497 989	573 245 311
trng::mrg5s::trng1	2 068 619 238	2 138 332 912	671 754 166	1 442 240 992	1 526 958 817

```

public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;
    static const parameter_type LEcuyer2;

    explicit mrg2(parameter_type=LEcuyer1);
    explicit mrg2(unsigned long, parameter_type=LEcuyer1);
    template<typename gen>
    explicit mrg2(gen &, parameter_type P=LEcuyer1);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type);

    void split(unsigned int, unsigned int);
    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

bool operator==(const mrg2 &, const mrg2 &);
bool operator!=(const mrg2 &, const mrg2 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const mrg2 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, mrg2 &);
}

```

```

namespace trng {

class mrg3 {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;
    static const parameter_type LEcuyer2;
    static const parameter_type LEcuyer3;

```

```

explicit mrg3(parameter_type=LEcuyer1);
explicit mrg3(unsigned long, parameter_type=LEcuyer1);
template<typename gen>
explicit mrg3(gen &, parameter_type P=LEcuyer1);

void seed();
void seed(unsigned long);
template<typename gen>
void seed(gen &);
void seed(result_type, result_type, result_type);

void split(unsigned int, unsigned int);
void jump2(unsigned int);
void jump(unsigned long long);

static const char * name();
long operator()(long) const;
};

bool operator==(const mrg3 &, const mrg3 &);
bool operator!=(const mrg3 &, const mrg3 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const mrg3 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, mrg3 &);
}

```

```

namespace trng {

class mrg3s {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type trng0;
    static const parameter_type trng1;

    explicit mrg3s(parameter_type=trng0);
    explicit mrg3s(unsigned long, parameter_type=trng0);
    template<typename gen>
    explicit mrg3s(gen &, parameter_type P=trng0);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type, result_type);

    void split(unsigned int, unsigned int);

```

```

    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

bool operator==(const mrg3s &, const mrg3s &);
bool operator!=(const mrg3s &, const mrg3s &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const mrg3s &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, mrg3s &);
}

```

```

namespace trng {

class mrg4 {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;
    static const parameter_type LEcuyer2;

    explicit mrg4(parameter_type=LEcuyer1);
    explicit mrg4(unsigned long, parameter_type=LEcuyer1);
    template<typename gen>
    explicit mrg4(gen &, parameter_type P=LEcuyer1);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type, result_type, result_type);

    void split(unsigned int, unsigned int);
    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

bool operator==(const mrg4 &, const mrg4 &);
bool operator!=(const mrg4 &, const mrg4 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const mrg4 &);
template<typename char_t, typename traits_t>

```

4 TRNG classes

```
std::basic_istream<char_t, traits_t> &  
operator>>(std::basic_istream<char_t, traits_t> &, mrg4 &);  
}
```

```
namespace trng {  
  
class mrg5 {  
public:  
    typedef long result_type;  
    result_type operator()() const;  
    static const result_type min;  
    static const result_type max;  
  
    class parameter_type;  
    class status_type;  
  
    static const parameter_type LEcuyer1;  
  
    explicit mrg5(parameter_type=LEcuyer1);  
    explicit mrg5(unsigned long, parameter_type=LEcuyer1);  
    template<typename gen>  
    explicit mrg5(gen &, parameter_type P=LEcuyer1);  
  
    void seed();  
    void seed(unsigned long);  
    template<typename gen>  
    void seed(gen &);  
    void seed(result_type, result_type, result_type, result_type, result_type);  
  
    void split(unsigned int, unsigned int);  
    void jump2(unsigned int);  
    void jump(unsigned long long);  
  
    static const char * name();  
    long operator()(long) const;  
};  
  
bool operator==(const mrg5 &, const mrg5 &);  
bool operator!=(const mrg5 &, const mrg5 &);  
template<typename char_t, typename traits_t>  
std::basic_ostream<char_t, traits_t> &  
operator<<(std::basic_ostream<char_t, traits_t> &, const mrg5 &);  
template<typename char_t, typename traits_t>  
std::basic_istream<char_t, traits_t> &  
operator>>(std::basic_istream<char_t, traits_t> &, mrg5 &);  
}
```

```
namespace trng {  
  
class mrg5s {  
public:  
    typedef long result_type;  
    result_type operator()() const;  
    static const result_type min;  
    static const result_type max;  
  
    class parameter_type;
```

```

class status_type;

static const parameter_type trng0;
static const parameter_type trng1;

explicit mrg5s(parameter_type=trng0);
explicit mrg5s(unsigned long, parameter_type=trng0);
template<typename gen>
explicit mrg5s(gen &, parameter_type P=trng0);

void seed();
void seed(unsigned long);
template<typename gen>
void seed(gen &);
void seed(result_type, result_type, result_type, result_type, result_type);

void split(unsigned int, unsigned int);
void jump2(unsigned int);
void jump(unsigned long long);

static const char * name();
long operator()(long) const;
};

bool operator==(const mrg5s &, const mrg5s &);
bool operator!=(const mrg5s &, const mrg5s &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const mrg5s &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, mrg5s &);
}

```

4.1.3 YARN generators

The classes `trng::yarnn` and `trng::yarnns` implement so-called YARN generators (yet another random number generator). Table 4.4 summarizes the key features of these classes. Each of them is based on a multiple recursive generator with n feedback taps, for which the transition algorithm reads

$$r_i = a_1 \cdot r_{i-1} + a_2 \cdot r_{i-2} + \dots + a_n \cdot r_{i-n} \bmod m.$$

The state of this generator at time i is given by $(r_{i-1}, r_{i-2}, \dots, r_{i-n})$. See section 2.4 for details on LFSR sequences.

The prime modulus m that characterizes the prime field \mathbb{F}_m was either chosen as the Mersenne Prime (classes `trng::mrgn`) or a Sophie-Germain Prime such that $m^n - 1$ has as few prime factors as possible (classes `trng::mrgns`). The former choice gives us some performance benefits, see section 7.1, whereas the second has some theoretical advantages, see section 2.4.2.

While pure multiple recursive generators return the r_i as pseudo-random numbers directly, a YARN generator “shuffles” the output of the underlying multiple recursive generator by a

bijjective mapping. In the case of a YARN generator with modulus m this mapping reads

$$q_i = \begin{cases} b^{r_i} \bmod m & \text{if } r_i > 0 \\ 0 & \text{if } r_i = 0 \end{cases} ,$$

where b is a generating element of the multiplicative group modulo m . This bijjective mapping destroys the linear structures of the linear feedback shift register sequence. But on the other hand the new sequence q_i inherits all the nice features of the linear feedback shift register sequence r_i , e. g. its period. Block splitting and leapfrog methods can be implemented as easily as for multiple recursive generators, see section 2.4 and 2.5 for details.

The classes `trng::yarnn` and `trng::yarnns` implement the interface described in section 3.1. Each class defines some parameter and status classes that will be used internally and by the constructor. Furthermore for each generator several parameter sets are given, see Table 4.3. Most of the parameter sets are taken from [24] and chosen to give generators with good statistical properties.

An instance of a class `trng::yarnn` or `trng::yarnns` can be instantiated by various constructors as specified for a random number engine. Additionally a non-default parameter set may be chosen. The classes `trng::yarnn` and `trng::yarnns` provide all necessary seeding functions (see Table 3.1) and additionally a function that sets the internal state $(r_{i-1}, r_{i-2}, \dots, r_{i-n})$. This function should never be called with all arguments set to zero. The classes `trng::yarnn` and `trng::yarnns` model the concept of a parallel random number engine and therefore the methods

```
void split(unsigned int, unsigned int);
void jump2(unsigned int);
void jump(unsigned long long);
```

are implemented. Furthermore the classes `trng::yarnn` or `trng::yarnns` provide a function that returns a string with its name and an operator `operator()`. Random number engines are comparable and can be written to or read from a stream.

The detailed interface of the classes `trng::mrgn` or `trng::mrgns` is given as follows:

```
namespace trng {

class yarn2 {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;
    static const parameter_type LEcuyer2;

    explicit yarn2(parameter_type=LEcuyer1);
    explicit yarn2(unsigned long, parameter_type=LEcuyer1);
    template<typename gen>
    explicit yarn2(gen &, parameter_type P=LEcuyer1);

    void seed();
```

Table 4.4: Key features of YARN generator classes.

class	header file	feedback taps n	prime field \mathbb{F}_m	period	return value of name()
trng::yarn2	trng/yarn2.hpp	2	$\mathbb{F}_{2^{31}-1}$	$m^2 - 1 \approx 2^{62} \approx 4.61 \cdot 10^{18}$	yarn2
trng::yarn3	trng/yarn3.hpp	3	$\mathbb{F}_{2^{31}-1}$	$m^3 - 1 \approx 2^{93} \approx 9.90 \cdot 10^{27}$	yarn3
trng::yarn3s	trng/yarn3s.hpp	3	$\mathbb{F}_{2^{31}-21069}$	$m^3 - 1 \approx 2^{93} \approx 9.90 \cdot 10^{27}$	yarn4s
trng::yarn4	trng/yarn4.hpp	4	$\mathbb{F}_{2^{31}-1}$	$m^4 - 1 \approx 2^{124} \approx 2.13 \cdot 10^{37}$	yarn4
trng::yarn5	trng/yarn5.hpp	5	$\mathbb{F}_{2^{31}-1}$	$m^5 - 1 \approx 2^{155} \approx 4.57 \cdot 10^{46}$	yarn5
trng::yarn5s	trng/yarn5s.hpp	5	$\mathbb{F}_{2^{31}-22641}$	$m^5 - 1 \approx 2^{155} \approx 4.57 \cdot 10^{46}$	yarn5s

Table 4.5: Parameter sets for YARN generators.

parameter set	a_1	a_2	a_3	a_4	a_5	b
trng::yarn2::LEcuyer1	1 498 809 829	1 160 990 996				123 567 893
trng::yarn2::LEcuyer2	46 325	1 084 587				123 567 893
trng::yarn3::LEcuyer1	2 021 422 057	1 826 992 351	1 977 753 457			123 567 893
trng::yarn3::LEcuyer2	1 476 728 729	0	1 155 643 113			123 567 893
trng::yarn3::LEcuyer3	65 338	0	64 636			123 567 893
trng::yarn3s::trng0	2 025 213 985	1 112 953 677	2 038 969 601			1616 076 847
trng::yarn3s::trng1	1 287 767 370	1 045 931 779	58 150 106			1616 076 847
trng::yarn4::LEcuyer1	2 001 982 722	1 412 284 257	1 155 380 217	1 668 339 922		123 567 893
trng::yarn4::LEcuyer2	64 886	0	0	64 322		123 567 893
trng::yarn5::LEcuyer1	107 374 182	0	0	0 104 480	123 567 893	
trng::yarn5s::trng0	1 053 223 373	1 530 818 118	1 612 122 482	133 497 989	573 245 311	889 744 251
trng::yarn5s::trng1	2 068 619 238	2 138 332 912	671 754 166	1 442 240 992	1 526 958 817	889 744 251


```

    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type);

    void split(unsigned int, unsigned int);
    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

bool operator==(const yarn2 &, const yarn2 &);
bool operator!=(const yarn2 &, const yarn2 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &t, const yarn2 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, yarn2 &);
}

```

```

namespace trng {

class yarn3 {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;
    static const parameter_type LEcuyer2;

    static const parameter_type LEcuyer3;

    explicit yarn3(parameter_type=LEcuyer1);
    explicit yarn3(unsigned long, parameter_type=LEcuyer1);
    template<typename gen>
    explicit yarn3(gen &, parameter_type P=LEcuyer1);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type, result_type);

    void split(unsigned int, unsigned int);
    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

```

```
};

bool operator==(const yarn3 &, const yarn3 &);
bool operator!=(const yarn3 &, const yarn3 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const yarn3 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, yarn3 &);
}
```

```
namespace trng {

class yarn3s {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type trng0;
    static const parameter_type trng1;

    explicit yarn3s(parameter_type=trng0);
    explicit yarn3s(unsigned long, parameter_type=trng0);
    template<typename gen>
    explicit yarn3s(gen &, parameter_type P=trng0);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type, result_type);

    void split(unsigned int, unsigned int);
    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

bool operator==(const yarn3s &, const yarn3s &);
bool operator!=(const yarn3s &, const yarn3s &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const yarn3s &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, yarn3s &);
}
```

```

namespace trng {

class yarn4 {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;
    static const parameter_type LEcuyer2;

    explicit yarn4(parameter_type=LEcuyer1);
    explicit yarn4(unsigned long, parameter_type=LEcuyer1);
    template<typename gen>
    explicit yarn4(gen &, parameter_type P=LEcuyer1);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type, result_type, result_type);

    void split(unsigned int, unsigned int);
    void jump2(unsigned int);
    void jump(unsigned long long);

    static const char * name();
    long operator()(long) const;
};

bool operator==(const yarn4 &, const yarn4 &);
bool operator!=(const yarn4 &, const yarn4 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const yarn4 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, yarn4 &);
}

```

```

namespace trng {

class yarn5 {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type LEcuyer1;

```

```

explicit yarn5(parameter_type=LEcuyer1);
explicit yarn5(unsigned long, parameter_type=LEcuyer1);
template<typename gen>
explicit yarn5(gen &, parameter_type P=LEcuyer1);

void seed();
void seed(unsigned long);
template<typename gen>
void seed(gen &);
void seed(result_type, result_type, result_type, result_type, result_type);

void split(unsigned int, unsigned int);
void jump2(unsigned int);
void jump(unsigned long long);

static const char * name();
long operator()(long) const;
};

bool operator==(const yarn5 &, const yarn5 &);
bool operator!=(const yarn5 &, const yarn5 &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const yarn5 &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, yarn5 &);
}

```

```

namespace trng {

class yarn5s {
public:
    typedef long result_type;
    result_type operator()() const;
    static const result_type min;
    static const result_type max;

    class parameter_type;
    class status_type;

    static const parameter_type trng0;
    static const parameter_type trng1;

    explicit yarn5s(parameter_type=trng0);
    explicit yarn5s(unsigned long, parameter_type=trng0);
    template<typename gen>
    explicit yarn5s(gen &, parameter_type P=trng0);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
    void seed(result_type, result_type, result_type, result_type, result_type);

    void split(unsigned int, unsigned int);

```

```

void jump2(unsigned int);
void jump(unsigned long long);

static const char * name();
long operator()(long) const;
};

bool operator==(const yarn5s &, const yarn5s &);
bool operator!=(const yarn5s &, const yarn5s &);
template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const yarn5s &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, yarn5s &);
}

```

4.1.4 Lagged Fibonacci generators

The template classes `trng::lagfib2xor`, `trng::lagfib4xor`, `trng::lagfib2plus`, `trng::lagfib4plus` model random number engines (no splitting facilities) and implement lagged Fibonacci generators with two or four feedback taps and exclusive-or or additive operation. The recursion relation of these types of generators read

$$\begin{aligned}
r_i &= r_{i-A} \oplus r_{i-B} \\
r_i &= r_{i-A} \oplus r_{i-B} \oplus r_{i-C} \oplus r_{i-D} \\
r_i &= r_{i-A} + r_{i-B} \bmod 2^l \\
r_i &= r_{i-A} + r_{i-B} + r_{i-C} + r_{i-D} \bmod 2^l.
\end{aligned}$$

These template classes are parameterized by an unsigned integer type, e.g. `unsigned int` or `unsigned long long`, and the position of the feedback taps with $A < B < C < D$. For properly chosen feedback taps the period of an exclusive-or generator is $2^B - 1$ or $2^D - 1$ respectively, and the period of an plus generator is $(2^B - 1)2^{l-1}$ or $(2^D - 1)2^{l-1}$ respectively, where l denotes the number of significant bits of the integer type given as a template argument. Template classes are declared in the header files `trng/lagfib2xor.hpp`, `trng/lagfib4xor.hpp`, `trng/lagfib2plus.hpp`, and `trng/lagfib4plus.hpp`. For convenience TRNG provides some typedefs for some realizations of lagged Fibonacci generators with two or four feedback taps.

The detailed interface of the classes `trng::lagfib2xor`, `trng::lagfib4xor`, `trng::lagfib2plus`, `trng::lagfib4plus` is given as follows:

```

namespace trng {

template<typename integer_type,
        unsigned int A, unsigned int B>
class lagfib2xor {
public:
    typedef integer_type result_type;
    result_type operator()();
    static const result_type min;
    static const result_type max;
};

```

```

class status_type;

lagfib2xor();
explicit lagfib2xor(unsigned long);
template<typename gen>
explicit lagfib2xor(gen &);

void seed();
void seed(unsigned long);
template<typename gen>
void seed(gen &);
};

typedef lagfib2xor<unsigned long,      103,   250> r250_ul;
typedef lagfib2xor<unsigned long long, 103,   250> r250_ull;
typedef lagfib2xor<unsigned long,      168,   521> lagfib2xor_521_ul;
typedef lagfib2xor<unsigned long long, 168,   521> lagfib2xor_521_ull;
typedef lagfib2xor<unsigned long,      273,   607> lagfib2xor_607_ul;
typedef lagfib2xor<unsigned long long, 273,   607> lagfib2xor_607_ull;
typedef lagfib2xor<unsigned long,      418,  1279> lagfib2xor_1279_ul;
typedef lagfib2xor<unsigned long long, 418,  1279> lagfib2xor_1279_ull;
typedef lagfib2xor<unsigned long,     1029,  2281> lagfib2xor_2281_ul;
typedef lagfib2xor<unsigned long long, 1029,  2281> lagfib2xor_2281_ull;
typedef lagfib2xor<unsigned long,      576,  3217> lagfib2xor_3217_ul;
typedef lagfib2xor<unsigned long long, 576,  3217> lagfib2xor_3217_ull;
typedef lagfib2xor<unsigned long,     2098,  4423> lagfib2xor_4423_ul;
typedef lagfib2xor<unsigned long long, 2098,  4423> lagfib2xor_4423_ull;
typedef lagfib2xor<unsigned long,     4187,  9689> lagfib2xor_9689_ul;
typedef lagfib2xor<unsigned long long, 4187,  9689> lagfib2xor_9689_ull;
typedef lagfib2xor<unsigned long,     9842, 19937> lagfib2xor_19937_ul;
typedef lagfib2xor<unsigned long long, 9842, 19937> lagfib2xor_19937_ull;
}

```

```

namespace trng {

template<typename integer_type,
        unsigned int A, unsigned int B, unsigned int C, unsigned int D>
class lagfib4xor {
public:
    typedef integer_type result_type;
    result_type operator()();
    static const result_type min;
    static const result_type max;

    class status_type;

    lagfib4xor();
    explicit lagfib4xor(unsigned long);
    template<typename gen>
    explicit lagfib4xor(gen &);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
};

```

```

};

typedef lagfib4xor<unsigned long,      471, 1586,  6988,  9689> Ziff_ul;
typedef lagfib4xor<unsigned long long, 471, 1586,  6988,  9689> Ziff_ull;
typedef lagfib4xor<unsigned long,      168,  205,   242,   521> lagfib4xor_521_ul;
typedef lagfib4xor<unsigned long long, 168,  205,   242,   521> lagfib4xor_521_ull;
typedef lagfib4xor<unsigned long,      147,  239,   515,   607> lagfib4xor_607_ul;
typedef lagfib4xor<unsigned long long, 147,  239,   515,   607> lagfib4xor_607_ull;
typedef lagfib4xor<unsigned long,      418,  705,   992,  1279> lagfib4xor_1279_ul;
typedef lagfib4xor<unsigned long long, 418,  705,   992,  1279> lagfib4xor_1279_ull;
typedef lagfib4xor<unsigned long,      305,  610,   915,  2281> lagfib4xor_2281_ul;
typedef lagfib4xor<unsigned long long, 305,  610,   915,  2281> lagfib4xor_2281_ull;
typedef lagfib4xor<unsigned long,      576,  871,  1461,  3217> lagfib4xor_3217_ul;
typedef lagfib4xor<unsigned long long, 576,  871,  1461,  3217> lagfib4xor_3217_ull;
typedef lagfib4xor<unsigned long,      1419, 1736,  2053,  4423> lagfib4xor_4423_ul;
typedef lagfib4xor<unsigned long long, 1419, 1736,  2053,  4423> lagfib4xor_4423_ull;
typedef lagfib4xor<unsigned long,      471, 2032,  4064,  9689> lagfib4xor_9689_ul;
typedef lagfib4xor<unsigned long long, 471, 2032,  4064,  9689> lagfib4xor_9689_ull;
typedef lagfib4xor<unsigned long,      3860, 7083, 11580, 19937> lagfib4xor_19937_ul;
typedef lagfib4xor<unsigned long long, 3860, 7083, 11580, 19937> lagfib4xor_19937_ull;

}

```

```

namespace trng {

template<typename integer_type,
        unsigned int A, unsigned int B>
class lagfib2plus {
public:
    typedef integer_type result_type;
    result_type operator()();
    static const result_type min;
    static const result_type max;

    class status_type;

    lagfib2plus();
    explicit lagfib2plus(unsigned long);
    template<typename gen>
    explicit lagfib2plus(gen &);

    void seed();
    void seed(unsigned long);
    template<typename gen>
    void seed(gen &);
};

typedef lagfib2plus<unsigned long,      168,   521> lagfib2plus_521_ul;
typedef lagfib2plus<unsigned long long, 168,   521> lagfib2plus_521_ull;
typedef lagfib2plus<unsigned long,      273,   607> lagfib2plus_607_ul;
typedef lagfib2plus<unsigned long long, 273,   607> lagfib2plus_607_ull;
typedef lagfib2plus<unsigned long,      418,  1279> lagfib2plus_1279_ul;
typedef lagfib2plus<unsigned long long, 418,  1279> lagfib2plus_1279_ull;
typedef lagfib2plus<unsigned long,      1029, 2281> lagfib2plus_2281_ul;
typedef lagfib2plus<unsigned long long, 1029, 2281> lagfib2plus_2281_ull;
typedef lagfib2plus<unsigned long,      576,  3217> lagfib2plus_3217_ul;
typedef lagfib2plus<unsigned long long, 576,  3217> lagfib2plus_3217_ull;

```

```

typedef lagfib2plus<unsigned long,      2098,  4423> lagfib2plus_4423_ul;
typedef lagfib2plus<unsigned long long, 2098,  4423> lagfib2plus_4423_ull;
typedef lagfib2plus<unsigned long,      4187,  9689> lagfib2plus_9689_ul;
typedef lagfib2plus<unsigned long long, 4187,  9689> lagfib2plus_9689_ull;
typedef lagfib2plus<unsigned long,      9842, 19937> lagfib2plus_19937_ul;
typedef lagfib2plus<unsigned long long, 9842, 19937> lagfib2plus_19937_ull;

}

namespace trng {

    template<typename integer_type,
             unsigned int A, unsigned int B, unsigned int C, unsigned int D>
    class lagfib4plus {
    public:
        typedef integer_type result_type;
        result_type operator()();
        static const result_type min;
        static const result_type max;

        class status_type;

        lagfib4plus();
        explicit lagfib2plus(unsigned long);
        template<typename gen>
        explicit lagfib4plus(gen &);

        void seed();
        void seed(unsigned long);
        template<typename gen>
        void seed(gen &);
    };

    typedef lagfib4plus<unsigned long,      168,  205,  242,  521> lagfib4plus_521_ul;
    typedef lagfib4plus<unsigned long long, 168,  205,  242,  521> lagfib4plus_521_ull;
    typedef lagfib4plus<unsigned long,      147,  239,  515,  607> lagfib4plus_607_ul;
    typedef lagfib4plus<unsigned long long, 147,  239,  515,  607> lagfib4plus_607_ull;
    typedef lagfib4plus<unsigned long,      418,  705,  992, 1279> lagfib4plus_1279_ul;
    typedef lagfib4plus<unsigned long long, 418,  705,  992, 1279> lagfib4plus_1279_ull;
    typedef lagfib4plus<unsigned long,      305,  610,  915, 2281> lagfib4plus_2281_ul;
    typedef lagfib4plus<unsigned long long, 305,  610,  915, 2281> lagfib4plus_2281_ull;
    typedef lagfib4plus<unsigned long,      576,  871, 1461, 3217> lagfib4plus_3217_ul;
    typedef lagfib4plus<unsigned long long, 576,  871, 1461, 3217> lagfib4plus_3217_ull;
    typedef lagfib4plus<unsigned long,      1419, 1736, 2053, 4423> lagfib4plus_4423_ul;
    typedef lagfib4plus<unsigned long long, 1419, 1736, 2053, 4423> lagfib4plus_4423_ull;
    typedef lagfib4plus<unsigned long,      471, 2032, 4064, 9689> lagfib4plus_9689_ul;
    typedef lagfib4plus<unsigned long long, 471, 2032, 4064, 9689> lagfib4plus_9689_ull;
    typedef lagfib4plus<unsigned long,      3860, 7083, 11580, 19937> lagfib4plus_19937_ul;
    typedef lagfib4plus<unsigned long long, 3860, 7083, 11580, 19937> lagfib4plus_19937_ull;
}

```

4.2 Random number distributions

This section gives a detailed description of all random number distributions, that have been implemented by TRNG. Each subsection presents the public interface of one random num-

ber distribution. The part of the public interface, that is mandatory for a random number distribution, will not be discussed in detail, read section 3.2 instead.

Additionally to the requirements in section 3.2 each random number distribution class provides a member function that calculates its probability distribution function, its cumulative distribution function and in the case of continuous distributions its inverse cumulative distribution function as well. These member functions have the signatures

```
double pdf(double x) const;
double cdf(double x) const;
double icdf(double x) const;
```

and for discrete random variables

```
double pdf(int x) const;
double cdf(int x) const;
```

The concept of a random number distribution requires two functions, that take a random number engine as its argument and generate a random variable with some specific distribution by calling `operator()` of the given random number engine. Note, the concept of a random number distribution does not specify how often `operator()` is called. This allows the implementor of a random number distribution to choose between various algorithms [20] that transform uniform random numbers into non-uniform distributed numbers. Some of these algorithms transform exactly one uniform random number into one non-uniform number, while some other algorithms have to call `operator()` more than once. How often `operator()` is called may even vary at runtime. If not otherwise stated, all random number distributions in TRNG are implemented in such a way, that `operator()` is called exactly once. Because of this special feature it is much more easy to write parallel Monte Carlo simulations that give the same result (and statistical error) independent of the number of parallel processes. We say such algorithms play fair, see section 2.3 and 6.

4.2.1 Uniform distributions

TRNG provides by the classes `uniform01_dist`, `uniform_dist` and `uniform_int_dist` three different kinds of uniform distributions.

Class `uniform01_dist` provides random numbers with distribution function

$$p(x) = \begin{cases} 1 & \text{if } 0 \leq x < 1 \\ 0 & \text{otherwise} \end{cases}.$$

The class `uniform01_dist` is declared in the header file `trng/uniform01_dist.hpp` and its public interface is given as follows:

```
namespace trng {

class uniform01_dist {
public:
    typedef double result_type;
    class param_type;
    explicit uniform01_dist();
    explicit uniform01_dist(const param_type &);
    void reset();
    template<typename R>
```

```

double operator()(R &);
template<typename R>
double operator()(R &, const param_type &);
double min() const;
double max() const;
param_type param() const;
void param(const param_type &);
double pdf(double x) const;
double cdf(double x) const;
double icdf(double x) const;
};

bool operator==(const uniform01_dist::param_type &, const uniform01_dist::param_type &);
bool operator!=(const uniform01_dist::param_type &, const uniform01_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const uniform01_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, uniform01_dist::param_type &);

bool operator==(const uniform01_dist &, const uniform01_dist &);
bool operator!=(const uniform01_dist &, const uniform01_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const uniform01_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, uniform01_dist &);
}

```

Class `uniform_dist` is a generalization of class `uniform01_dist` and it provides random numbers with distribution function

$$p(x|a,b) = \begin{cases} 1/(b-a) & \text{if } a \leq x < b \\ 0 & \text{otherwise} \end{cases}.$$

Valid parameters for this distribution are $a, b \in \mathbb{R}$ with $a < b$. If pseudo-random numbers uniformly distributed in $[0, 1)$ are desired, class `uniform01_dist` might be faster than `uniform_dist` with parameters $a = 0$ and $b = 1$.

The class `uniform_dist` is declared in the header file `trng/uniform_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class uniform_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double a() const;
        void a(double);
        double b() const;
        void b(double);
    };
};

```

```

    explicit param_type(double a, double b);
};
explicit uniform_dist(double a, double b);
explicit uniform_dist(const param_type &);
void reset();
template<typename R>
double operator()(R &);
template<typename R>
double operator()(R &, const param_type &)
double min() const;
double max() const;
param_type param() const;
void param(const param_type &);
double a() const;
void a(double);
double b();
void b(double);
double pdf(double x) const;
double cdf(double x) const;
double icdf(double x) const;
};

bool operator==(const uniform_dist::param_type &, const uniform_dist::param_type &);
bool operator!=(const uniform_dist::param_type &, const uniform_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const uniform_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, uniform_dist::param_type &);

bool operator==(const uniform_dist &, const uniform_dist &);
bool operator!=(const uniform_dist &, const uniform_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const uniform_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, uniform_dist &);
}

```

Class `uniform_int_dist` is a variant of `uniform_dist` for integer valued random variables. It provides random numbers with distribution function

$$p(x|a,b) = \begin{cases} 1/(b-a) & \text{if } a \leq x < b \\ 0 & \text{otherwise} \end{cases} \quad \text{for } x \in \mathbb{Z}.$$

Valid parameters for this distribution are $a, b \in \mathbb{Z}$ with $a < b$.

The class `uniform_int_dist` is declared in the header file `trng/uniform_int_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class uniform_int_dist {

```

```

public:
    typedef int result_type;
    class param_type {
    public:
        int a() const;
        void a(int);
        int b() const;
        void b(int);
        explicit param_type(int a, int b);
    };
    explicit uniform_int_dist(int a, int b);
    explicit uniform_int_dist(const param_type &)
    void reset();
    template<typename R>
    int operator()(R &);
    template<typename R>
    int operator()(R &, const param_type &);
    int min() const;
    int max() const;
    param_type param() const;
    void param(const param_type &);
    int a() const;
    void a(int);
    int b() const;
    void b(int);
    double pdf(int x) const;
    double cdf(int x) const;
};

bool operator==(const uniform_int_dist::param_type &, const uniform_int_dist::param_type &);
bool operator!=(const uniform_int_dist::param_type &, const uniform_int_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const uniform_int_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, uniform_int_dist::param_type &);

bool operator==(const uniform_int_dist &, const uniform_int_dist &);
bool operator!=(const uniform_int_dist &, const uniform_int_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const uniform_int_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, uniform_int_dist &);
}

```

4.2.2 Exponential distribution

Class `exponential_dist` provides random numbers with exponential distribution with mean μ . The probability distribution function reads

$$p(x|\mu) = \begin{cases} \frac{1}{\mu}e^{-x/\mu} & \text{if } x \geq 0 \\ 0 & \text{otherwise} \end{cases} \quad \text{for } x \in \mathbb{R}.$$

Valid parameter for this distribution is $\mu \in \mathbb{R}$ with $\mu > 0$.

The class `exponential_dist` is declared in the header file `trng/exponential_dist.hpp` and its public interface is given as follows:

```
namespace trng {

class exponential_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double mu() const;
        void mu(double);
        explicit param_type(double mu);
    };
    explicit exponential_dist(double mu);
    explicit exponential_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double mu() const;
    void mu(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const exponential_dist::param_type &, const exponential_dist::param_type &);
bool operator!=(const exponential_dist::param_type &, const exponential_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const exponential_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, exponential_dist::param_type &);

bool operator==(const exponential_dist &, const exponential_dist &);
bool operator!=(const exponential_dist &, const exponential_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
```

```

operator<<(std::basic_ostream<char_t, traits_t> &, const exponential_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, exponential_dist &);
}

```

4.2.3 Normal distribution

There are two classes for producing random numbers with normal distribution, `normal_dist` and `correlated_normal_dist`. Class `normal_dist` provides uncorrelated random numbers with normal distribution with mean μ and standard deviation σ . The probability distribution function reads

$$p(x|\mu, \sigma) = \frac{1}{\sqrt{2\pi}\sigma} e^{-(x-\mu)^2/(2\sigma^2)}.$$

Valid parameters for this distribution are $\mu, \sigma \in \mathbb{R}$ with $\sigma > 0$. The normal distribution is also known as Gaussian distribution.

The class `normal_dist` is declared in the header file `trng/normal_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class normal_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double mu() const;
        void mu(double);
        double sigma() const;
        void sigma(double);
        param_type(double mu, double sigma);
    };
    normal_dist(double mu, double sigma);
    explicit normal_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double mu() const;
    void mu(double);
    double sigma() const;
    void sigma(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const normal_dist::param_type &, const normal_dist::param_type &);
bool operator!=(const normal_dist::param_type &, const normal_dist::param_type &);

```

```

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const normal_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, normal_dist::param_type &);

bool operator==(const normal_dist &, const normal_dist &);
bool operator!=(const normal_dist &, const normal_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const normal_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, normal_dist &);
}

```

If $\mathbf{x} = (x_1, x_2, \dots, x_d)$ are d random variables, then the multivariate normal density function for \mathbf{x} is

$$p(\mathbf{x}|\mathbf{V}) = \frac{1}{\sqrt{(2\pi)^d \det \mathbf{V}}} \exp\left(-\frac{1}{2}\mathbf{x}^T \mathbf{V}^{-1} \mathbf{x}\right). \quad (4.1)$$

Each variable x_1, x_2, \dots, x_d has mean zero and the the covariance matrix of x_1, x_2, \dots, x_d is given by the symmetric positive definite $d \times d$ matrix \mathbf{V} . Class `correlated_normal_dist` provides correlated random numbers with normal distribution by the transformation of uncorrelated random numbers [8].

The class `normal_dist` is declared in the header file `trng/normal_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class correlated_normal_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        template<typename iter>
        param_type(iter first, iter last);
    };
    template<typename iter>
    correlated_normal_dist(iter first, iter last);
    explicit correlated_normal_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &p_new);
};

bool operator==(const correlated_normal_dist::param_type &,
                const correlated_normal_dist::param_type &);

```

```

bool operator!=(const correlated_normal_dist::param_type &,
               const correlated_normal_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const correlated_normal_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, correlated_normal_dist::param_type &);

bool operator==(const correlated_normal_dist &, const correlated_normal_dist &);
bool operator!=(const correlated_normal_dist &, const correlated_normal_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const correlated_normal_dist &);

template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, correlated_normal_dist &);
}

```

The covariance matrix \mathbf{V} has to be passed to the constructor of `correlated_normal_dist` by two iterators. It is not checked, if the matrix is positive definite. The call operator `operator()` returns a single random number and has complexity $\mathcal{O}(d)$. As a consequence, the generation of a tuple of d correlated random numbers takes $\mathcal{O}(d^2)$ operations.

Successive calls return random numbers with variance $\mathbf{V}_{1,1}$, $\mathbf{V}_{2,2}$ and so on, until the `operator()` has been called d times, which returns a random number with variance $\mathbf{V}_{d,d}$. A sequence of further calls of `operator()` will return random numbers with the same sequences of variances. The method `reset` resets the internal state of the distribution such that, of further calls of `operator()` will return random numbers starting with a number with variance $\mathbf{V}_{1,1}$. Listing 4.1 illustrates the usage of class `correlated_normal_dist`.

Listing 4.1: Demonstration program illustrating the usage of `correlated_normal_dist`.

```

#include <cstdlib>
#include <iostream>
#include <iomanip>
#include <vector>
#include <trng/lcg64.hpp>
#include <trng/correlated_normal_dist.hpp>

double covariance(const std::vector<double> &v1, const std::vector<double> &v2) {
    std::vector<double>::size_type n=v1.size();
    double m1=0.0, m2=0.0, c=0.0;
    for (std::vector<double>::size_type i=0; i<n; ++i) {
        m1+=v1[i]/n; m2+=v2[i]/n;
    }
    for (std::vector<double>::size_type i=0; i<n; ++i)
        c+=(v1[i]-m1)*(v2[i]-m2)/n;
    return c;
}

int main() {
    const int d=4;

```



```

// covariance matrix
double sig[d][d] = { { 2.0, -0.5, 0.3, -0.3},
                     {-0.5, 3.0, -0.3, 0.3},
                     { 0.3, -0.3, 1.0, -0.3},
                     {-0.3, 0.3, -0.3, 1.0} };
trng::correlated_normal_dist D(&sig[0][0], &sig[d-1][d-1]+1);
trng::lcg64 R;

std::vector<double> x1, x2, x3, x4;
// generate 4-tuples of correlated normal variables
for (int i=0; i<1000000; ++i) {
    x1.push_back(D(R)); x2.push_back(D(R)); x3.push_back(D(R)); x4.push_back(D(R));
}
// print (empirical) covariance matrix
std::cout << std::setprecision(4)
    << covariance(x1, x1) << '\t' << covariance(x1, x2) << '\t'
    << covariance(x1, x3) << '\t' << covariance(x1, x4) << '\n'
    << covariance(x2, x1) << '\t' << covariance(x2, x2) << '\t'
    << covariance(x2, x3) << '\t' << covariance(x2, x4) << '\n'
    << covariance(x3, x1) << '\t' << covariance(x3, x2) << '\t'
    << covariance(x3, x3) << '\t' << covariance(x3, x4) << '\n'
    << covariance(x4, x1) << '\t' << covariance(x4, x2) << '\t'
    << covariance(x4, x3) << '\t' << covariance(x4, x4) << '\n';
return EXIT_SUCCESS;
}

```

4.2.4 Cauchy distribution

Class `cauchy_dist` provides random numbers with Cauchy distribution with parameters θ and η . The probability distribution function reads

$$p(x|\theta, \eta) = \frac{1}{\theta\pi \left(1 + \left(\frac{x-\eta}{\theta}\right)^2\right)}.$$

Valid parameters for this distribution are $\theta, \eta \in \mathbb{R}$ with $\theta > 0$. The Cauchy distribution is also known as Lorentz distribution or Breit-Wigner distribution.

The class `cauchy_dist` is declared in the header file `trng/cauchy_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class cauchy_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double theta() const;
        void theta(double);
        double eta() const;
        void eta(double);
        explicit param_type(double theta, double eta);
    };
    explicit cauchy_dist(double theta, double eta);
    explicit cauchy_dist(const param_type &);
};

```

```

void reset();
template<typename R>
double operator()(R &);
template<typename R>
double operator()(R &, const param_type &);
double min() const;
double max() const;
param_type param() const;
void param(const param_type &);
double theta() const;
void theta(double);
double eta() const;
void eta(double);
double pdf(double) const;
double cdf(double) const;
double icdf(double) const;
};

bool operator==(const cauchy_dist::param_type &, const cauchy_dist::param_type &);
bool operator!=(const cauchy_dist::param_type &, const cauchy_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const cauchy_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, cauchy_dist::param_type &);

bool operator==(const cauchy_dist &, const cauchy_dist &);
bool operator!=(const cauchy_dist &, const cauchy_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const cauchy_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, cauchy_dist &);
}

```

4.2.5 Logistic distribution

Class `logistic_dist` provides random numbers with Logistic distribution with parameters θ and η . The probability distribution function reads

$$p(x|\theta, \eta) = \frac{e^{-(x-\eta)/\theta}}{\theta (1 + e^{-(x-\eta)/\theta})^2}.$$

Valid parameters for this distribution are $\theta, \eta \in \mathbb{R}$ with $\theta > 0$.

The class `logistic_dist` is declared in the header file `trng/logistic_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class logistic_dist {
public:

```

```

typedef double result_type;
class param_type {
public:
    double theta() const;
    void theta(double);
    double eta() const;
    void eta(double);
    explicit param_type(double theta, double eta);
};
explicit logistic_dist(double theta, double eta);
explicit logistic_dist(const param_type &);
void reset();
template<typename R>
double operator()(R &);
template<typename R>
double operator()(R &, const param_type &);
double min() const;
double max() const;
param_type param() const;
void param(const param_type &);
double theta() const;
void theta(double);
double eta() const;
void eta(double);
double pdf(double) const;
double cdf(double) const;
double icdf(double) const;
};

bool operator==(const logistic_dist::param_type &, const logistic_dist::param_type &);
bool operator!=(const logistic_dist::param_type &, const logistic_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const logistic_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, logistic_dist::param_type &);

bool operator==(const logistic_dist &, const logistic_dist &);
bool operator!=(const logistic_dist &, const logistic_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const logistic_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, logistic_dist &);
}

```

4.2.6 Lognormal distribution

Class `lognormal_dist` provides random numbers with lognormal distribution with parameters μ and σ . The probability distribution function reads

$$p(x|\mu, \sigma) = \begin{cases} 0 & \text{for } x \leq 0 \\ \frac{1}{x\sqrt{2\pi\sigma^2}} e^{-(\ln x - \mu)^2 / (2\sigma^2)} & \text{for } x > 0 \end{cases}.$$

Valid parameters for this distribution are $\mu, \sigma \in \mathbb{R}$ with $\sigma > 0$.

The class `lognormal_dist` is declared in the header file `trng/lognormal_dist.hpp` and its public interface is given as follows:

```
namespace trng {

class lognormal_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double mu() const;
        void mu(double);
        double sigma() const;
        void sigma(double);
        explicit param_type(double mu, double sigma);
    };
    explicit lognormal_dist(double mu, double sigma);
    explicit lognormal_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double mu() const;
    void mu(double);
    double sigma() const;
    void sigma(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const lognormal_dist::param_type &, const lognormal_dist::param_type &);
bool operator!=(const lognormal_dist::param_type &, const lognormal_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const lognormal_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, lognormal_dist::param_type &);
```

```

bool operator==(const lognormal_dist &, const lognormal_dist &);
bool operator!=(const lognormal_dist &, const lognormal_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const lognormal_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, lognormal_dist &);
}

```

4.2.7 Pareto distribution

Class `pareto_dist` provides random numbers with Pareto distribution with parameters γ and θ . The probability distribution function reads

$$p(x|\gamma, \theta) = \begin{cases} 0 & \text{for } x < 0 \\ \frac{\gamma}{\theta} \left(1 + \frac{x}{\theta}\right)^{-\gamma-1} & \text{for } x \geq 0 \end{cases}.$$

Valid parameters for this distribution are $\gamma, \theta \in \mathbb{R}$ with $\gamma > 0$ and $\theta > 0$. Actually in mathematics literature one can find two different kinds of probability distributions, that are referred as Pareto distribution. Section 4.2.8 introduces another probability distribution that is also sometimes called Pareto distribution.

The class `pareto_dist` is declared in the header file `trng/pareto_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class pareto_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double gamma() const;
        void gamma(double);
        double theta() const;
        void theta(double);
        explicit param_type(double gamma, double theta);
    };
    explicit pareto_dist(double gamma, double theta);
    explicit pareto_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double gamma() const;
    void gamma(double);
    double theta() const;

```

```

    void theta(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const pareto_dist::param_type &, const pareto_dist::param_type &);
bool operator!=(const pareto_dist::param_type &, const pareto_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const pareto_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, pareto_dist::param_type &);

bool operator==(const pareto_dist &, const pareto_dist &);
bool operator!=(const pareto_dist &, const pareto_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const pareto_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, pareto_dist &);
}

```

4.2.8 Power-law distribution

Class `powerlaw_dist` provides random numbers with power-law distribution with parameters γ and θ . This distribution is related to the Pareto distribution and its probability distribution function reads

$$p(x|\gamma, \theta) = \begin{cases} 0 & \text{for } x < \theta \\ \frac{\gamma}{\theta} \left(\frac{x}{\theta}\right)^{-\gamma-1} & \text{for } x \geq \theta \end{cases}.$$

Valid parameters for this distribution are $\gamma, \theta \in \mathbb{R}$ with $\gamma > 0$ and $\theta > 0$.

The class `powerlaw_dist` is declared in the header file `trng/powerlaw_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class powerlaw_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double gamma() const;
        void gamma(double);
        double theta() const;
        void theta(double);
        explicit param_type(double gamma, double theta);
    };
    explicit powerlaw_dist(double gamma, double theta);
    explicit powerlaw_dist(const param_type &);
};

```

```

void reset();
template<typename R>
double operator()(R &);
template<typename R>
double operator()(R &, const param_type &);
double min() const;
double max() const;
param_type param() const;
void param(const param_type &);
double gamma() const;
void gamma(double);
double theta() const;
void theta(double);
double pdf(double) const;
double cdf(double) const;
double icdf(double) const;
};

bool operator==(const powerlaw_dist::param_type &, const powerlaw_dist::param_type &);
bool operator!=(const powerlaw_dist::param_type &, const powerlaw_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const powerlaw_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, powerlaw_dist::param_type &);

bool operator==(const powerlaw_dist &, const powerlaw_dist &);
bool operator!=(const powerlaw_dist &, const powerlaw_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const powerlaw_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, powerlaw_dist &);
}

```

4.2.9 Tent distribution

Class `tent_dist` provides random numbers with tent distribution with parameters m and d . This distribution is symmetric around m and its support is the interval $[m - d, m + d]$. The probability distribution function reads

$$p(x|m,d) = \begin{cases} 0 & \text{for } x < m - d \text{ or } x > m + d \\ \frac{1 + (x - m)/d}{d} & \text{for } m - d \leq x \leq m \\ \frac{1 - (x - m)/d}{d} & \text{for } m \leq x \leq m + d \end{cases}.$$

Valid parameters for this distribution are $m, d \in \mathbb{R}$ with $d > 0$.

The class `tent_dist` is declared in the header file `trng/tent_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class tent_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double m() const;
        void m(double);
        double d() const;
        void d(double);
        explicit param_type(double m, double d);
    };
    explicit tent_dist(double m, double d);
    explicit tent_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double m() const;
    void m(double);
    double d() const;
    void d(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const tent_dist::param_type &, const tent_dist::param_type &);
bool operator!=(const tent_dist::param_type &, const tent_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const tent_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, tent_dist::param_type &);

bool operator==(const tent_dist &, const tent_dist &);
bool operator!=(const tent_dist &, const tent_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const tent_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, tent_dist &);
}

```


4.2.10 Weibull distribution

Class `weibull_dist` provides random numbers with Weibull distribution with parameters β and θ . The probability distribution function reads

$$p(x|\beta, \theta) = \begin{cases} 0 & \text{for } x < \theta \\ \frac{\beta}{\theta} \left(\frac{x}{\theta}\right)^{\beta-1} e^{-(x/\theta)^\beta} & \text{for } x \geq \theta \end{cases}.$$

Valid parameters for this distribution are $\beta, \theta \in \mathbb{R}$ with $\beta > 0$ and $\theta > 0$. For $\beta = 1$ Weibull distribution degenerates to an exponential distribution and for $\beta = 2$ and $\theta = \sqrt{2} \cdot \sigma$ this distribution is also known as Rayleigh distribution with parameter σ .

The class `weibull_dist` is declared in the header file `trng/weibull_dist.hpp` and its public interface is given as follows:

```
namespace trng {

class weibull_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double beta() const;
        void beta(double);
        double theta() const;
        void theta(double);
        explicit param_type(double beta, double theta);
    };
    explicit weibull_dist(double beta, double theta);
    explicit weibull_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double beta() const;
    void beta(double);
    double theta() const;
    void theta(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const weibull_dist::param_type &, const weibull_dist::param_type &);
bool operator!=(const weibull_dist::param_type &, const weibull_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const weibull_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
```

```

operator>>(std::basic_istream<char_t, traits_t> &, weibull_dist::param_type &);

bool operator==(const weibull_dist &, const weibull_dist &);
bool operator!=(const weibull_dist &, const weibull_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const weibull_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, weibull_dist &);
}

```

4.2.11 Extreme value distribution

Class `extreme_value_dist` provides random numbers with extreme value distribution with parameters θ and η . The probability distribution function reads

$$p(x|\theta, \eta) = \frac{1}{\theta} \exp\left(\frac{x - \eta}{\theta} - \exp \frac{x - \eta}{\theta}\right).$$

Valid parameters for this distribution are $\theta, \eta \in \mathbb{R}$ with $\theta > 0$.

The class `extreme_value_dist` is declared in the header file `trng/extreme_value_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class extreme_value_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double theta() const;
        void theta(double);
        double eta() const;
        void eta(double);
        explicit param_type(double theta, double eta);
    };
    explicit extreme_value_dist(double theta, double eta);
    explicit extreme_value_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    double theta() const;
    void theta(double);
    double eta() const;
    void eta(double);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

```

```

bool operator==(const extreme_value_dist::param_type &,
                const extreme_value_dist::param_type &);
bool operator!=(const extreme_value_dist::param_type &,
                const extreme_value_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const extreme_value_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, extreme_value_dist::param_type &);

bool operator==(const extreme_value_dist &, const extreme_value_dist &);
bool operator!=(const extreme_value_dist &, const extreme_value_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const extreme_value_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, extreme_value_dist &);
}

```

4.2.12 Γ -distribution

Class `gamma_dist` provides random numbers with Γ -distribution with parameters θ and κ . The probability distribution function reads

$$p(x|\theta, \kappa) = \begin{cases} 0 & \text{if } x < 0 \\ \frac{1}{\theta \Gamma(\kappa)} \left(\frac{x}{\theta}\right)^{\kappa-1} e^{-x/\theta} & \text{if } x \geq 0 \end{cases}.$$

Valid parameters for this distribution are $\kappa, \theta \in \mathbb{R}$ with $\kappa \geq 1$ and $\theta > 0$. Note, Γ -distribution is defined for arbitrary $\kappa \geq 0$, but class `gamma_dist` can handle only Γ -distributions with $\kappa \geq 1$ correctly. For $\kappa = 1$ Γ -distribution degenerates to an exponential distribution.

The class `gamma_dist` is declared in the header file `trng/gamma_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class gamma_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double kappa() const;
        void kappa(double);
        double theta() const;
        void theta(double);
        explicit param_type(double kappa, double theta);
    };
    explicit gamma_dist(double kappa, double theta);
    explicit gamma_dist(const param_type &);

```

```

void reset();
template<typename R>
double operator()(R &);
template<typename R>
double operator()(R &, const param_type &);
double min() const;
double max() const;
param_type param() const;
void param(const param_type &);
double kappa() const;
void kappa(double);
double theta() const;
void theta(double);
double pdf(double) const;
double cdf(double) const;
double icdf(double) const;
};

bool operator==(const gamma_dist::param_type &, const gamma_dist::param_type &);
bool operator!=(const gamma_dist::param_type &, const gamma_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const gamma_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, gamma_dist::param_type &);

bool operator==(const gamma_dist &, const gamma_dist &);
bool operator!=(const gamma_dist &, const gamma_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const gamma_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, gamma_dist &);
}

```

4.2.13 χ^2 -distribution

Class `chi_square_dist` provides random numbers with χ^2 -distribution with ν degrees of freedom. The probability distribution function reads

$$p(x|\nu) = \begin{cases} 0 & \text{if } x < 0 \\ \frac{x^{\nu/2-1} e^{-x/2}}{2^{\nu/2} \Gamma(\nu/2)} & \text{if } x \geq 0 \end{cases}.$$

A valid parameter for this distribution is $\nu \in \mathbb{N}$ with $\nu \geq 1$. Note, χ^2 -distribution is a special case of Γ -distribution with $\kappa = \nu/2$ and $\theta = 2$.

The class `chi_square_dist` is declared in the header file `trng/chi_square_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class chi_square_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        int nu() const;
        void nu(int);
        explicit param_type(int nu);
    };
    explicit chi_square_dist(int nu);
    explicit chi_square_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const;
    void param(const param_type &);
    int nu() const;
    void nu(int);
    double pdf(double) const;
    double cdf(double) const;
    double icdf(double) const;
};

bool operator==(const chi_square_dist::param_type &, const chi_square_dist::param_type &);
bool operator!=(const chi_square_dist::param_type &, const chi_square_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const chi_square_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, chi_square_dist::param_type &);

bool operator==(const chi_square_dist &, const chi_square_dist &);
bool operator!=(const chi_square_dist &, const chi_square_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const chi_square_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, chi_square_dist &);
}

```

4.2.14 Rayleigh distribution

Class `rayleigh_dist` provides random numbers with Rayleigh distribution with parameter ν . The probability distribution function reads

$$p(x|\nu) = \begin{cases} 0 & \text{if } x \leq 0 \\ \frac{x}{\nu^2} e^{-x^2/(2\nu^2)} & \text{if } x > 0 \end{cases}.$$

A valid parameter for this distribution is $\nu > 0$.

The class `rayleigh_dist` is declared in the header file `trng/rayleigh_dist.hpp` and its public interface is given as follows:

```
namespace trng {

class rayleigh_dist {
public:
    typedef double result_type;
    class param_type {
    public:
        double nu() const;
        void nu(double nu_new);
        explicit param_type(double nu);
        friend class rayleigh_dist;
    };

    explicit rayleigh_dist(double nu);
    explicit rayleigh_dist(const param_type &);
    void reset();
    template<typename R>
    double operator()(R &);
    template<typename R>
    double operator()(R &, const param_type &);
    double min() const;
    double max() const;
    param_type param() const { return p; }
    void param(const param_type &);
    double nu() const;
    void nu(double);
    double pdf(double x) const;
    double cdf(double x) const;
    double icdf(double x) const;
};

bool operator==(const rayleigh_dist::param_type &, const rayleigh_dist::param_type &);
bool operator!=(const rayleigh_dist::param_type &, const rayleigh_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const rayleigh_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, rayleigh_dist::param_type &);

bool operator==(const rayleigh_dist &, const rayleigh_dist &);
bool operator!=(const rayleigh_dist &, const rayleigh_dist &);
```

```

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const rayleigh_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, rayleigh_dist &);
}

```

4.2.15 Bernoulli distribution

The template class `bernoulli_dist` provides random objects with Bernoulli distribution with parameter p . The probability distribution function reads

$$P(x|p) = \begin{cases} p & \text{if } x = \text{head} \\ 1 - p & \text{if } x = \text{tail} \\ 0 & \text{else} \end{cases}.$$

A valid parameter for this distribution is $p \in [0, 1]$.

The class `bernoulli_dist` is declared in the header file `trng/bernoulli_dist.hpp` and its public interface is given as follows:

```

namespace trng {

template<typename T>
class bernoulli_dist {
public:
    typedef T result_type;

    class param_type {
public:
        double p() const;
        void p(double);
        T head() const;
        void head(const T &);
        T tail() const;
        void tail(const T &);
        explicit param_type(double p, const T &head, const T &tail);
    };

    explicit bernoulli_dist(double p, const T &head, const T &tail);
    explicit bernoulli_dist(const param_type &);
    void reset();
    template<typename R>
    T operator()(R &);
    template<typename R>
    T operator()(R &, const param_type &);
};

```

Method `min` returns “head” and method `max` returns “tail”.

```

T min() const;
T max() const;
param_type param() const;
void param(const param_type &);
double p() const;
void p(double);

```

```

T head() const;
void head(const T &);
T tail() const;
void tail(const T &);

```

Method pdf will return p if its argument is “head”, $1 - p$ if its argument is “tail” and 0 otherwise.

```
double pdf(const T &) const;
```

Method cdf will return p if its argument is “head”, 1 if its argument is “tail” and 0 otherwise.

```

double cdf(const T &) const;
};

template<typename T>
bool operator==(const typename bernoulli_dist<T>::param_type &,
               const typename bernoulli_dist<T>::param_type &);
template<typename T>
bool operator!=(const typename bernoulli_dist<T>::param_type &,
               const typename bernoulli_dist<T>::param_type &);

template<typename char_t, typename traits_t, typename T>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &,
          const typename bernoulli_dist<T>::param_type &);
template<typename char_t, typename traits_t, typename T>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &,
          typename bernoulli_dist<T>::param_type &);

template<typename T>
bool operator==(const bernoulli_dist<T> &, const bernoulli_dist<T> &);
template<typename T>
bool operator!=(const bernoulli_dist<T> &, const bernoulli_dist<T> &);

template<typename char_t, typename traits_t, typename T>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const bernoulli_dist<T> &);
template<typename char_t, typename traits_t, typename T>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, bernoulli_dist<T> &);
}

```

4.2.16 Binomial distribution

Class `binomial_dist` provides random integers with binomial distribution with parameters p and n . The probability distribution function reads

$$P(x|p, n) = \begin{cases} \binom{n}{x} p^x (1-p)^{n-x} & \text{if } x \in \{0, 1, \dots, n\} \\ 0 & \text{else} \end{cases}.$$

Valid parameters for this distribution are $p \in [0, 1]$ and $n \in \mathbb{N}$.

The class `binomial_dist` is declared in the header file `trng/binomial_dist.hpp` and its public interface is given as follows:


```

namespace trng {

class binomial_dist {
public:
    typedef int result_type;

    class param_type {
public:
        double p() const;
        void p(double);
        int n() const;
        void n(int);
        explicit param_type(double p, int n);
    };

    explicit binomial_dist(double p, int n);
    explicit binomial_dist(const param_type &);
    void reset();
    template<typename R>
    int operator()(R &);
    template<typename R>
    int operator()(R &, const param_type &);
    int min() const;
    int max() const;
    param_type param() const;
    void param(const param_type &);
    double p() const;
    void p(double);
    int n() const;
    void n(int);
    double pdf(int) const;
    double cdf(int) const;
};

bool operator==(const binomial_dist::param_type &, const binomial_dist::param_type &);
bool operator!=(const binomial_dist::param_type &, const binomial_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const binomial_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, binomial_dist::param_type &);

bool operator==(const binomial_dist &, const binomial_dist &);
bool operator!=(const binomial_dist &, const binomial_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const binomial_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, binomial_dist &);
}

```

4.2.17 Geometric distribution

Class `geometric_dist` provides random integers with geometric distribution with parameter p . The probability distribution function reads

$$P(x|p) = p(1 - p)^x \quad \text{for } x \in \{0, 1, 2, \dots\}.$$

A valid parameter p is $p \in (0, 1)$.

The class `geometric_dist` is declared in the header file `trng/geometric_dist.hpp` and its public interface is given as follows:

```
namespace trng {

class geometric_dist {
public:
    typedef int result_type;

    class param_type {
    public:
        double p() const;
        void p(double);
        explicit param_type(double p);
    };

    explicit geometric_dist(double p);
    explicit geometric_dist(const param_type &);
    void reset();
    template<typename R>
    int operator()(R &);
    template<typename R>
    int operator()(R &, const param_type &);
    int min() const;
    int max() const;
    param_type param() const;
    void param(const param_type &);
    double p() const;
    void p(double);
    double pdf(int) const;
    double cdf(int) const;
};

bool operator==(const geometric_dist::param_type &, const geometric_dist::param_type &);
bool operator!=(const geometric_dist::param_type &, const geometric_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const geometric_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, geometric_dist::param_type &);

bool operator==(const geometric_dist &, const geometric_dist &);
bool operator!=(const geometric_dist &, const geometric_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const geometric_dist &);
```

```

template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, geometric_dist &);
}

```

4.2.18 Poisson distribution

Class `poisson_dist` provides random integers with poisson distribution with mean μ . The probability distribution function reads

$$P(x|\mu) = \frac{e^{-\mu}\mu^x}{x!} \quad \text{for } x \in \{0, 1, 2, \dots\}.$$

A valid parameter μ is $\mu \in [0, \infty)$.

The class `poisson_dist` is declared in the header file `trng/poisson_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class poisson_dist {
public:
    typedef int result_type;

    class param_type {
public:
        double mu() const;
        void mu(double);
        explicit param_type(double mu);
    };

    explicit poisson_dist(double mu);
    explicit poisson_dist(const param_type &);
    void reset();
    template<typename R>
    int operator()(R &);
    template<typename R>
    int operator()(R &, const param_type &);
    int min() const;
    int max() const;
    param_type param() const;
    void param(const param_type &);
    double mu() const;
    void mu(double);
    double pdf(int) const;
    double cdf(int) const;
};

bool operator==(const poisson_dist::param_type &, const poisson_dist::param_type &);
bool operator!=(const poisson_dist::param_type &, const poisson_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const poisson_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, poisson_dist::param_type &);

```

```

bool operator==(const poisson_dist &, const poisson_dist &);
bool operator!=(const poisson_dist &, const poisson_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const poisson_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, poisson_dist &);
}

```

4.2.19 Discrete distribution

Class `discrete_dist` provides random integers with arbitrary discrete distribution. The probability distribution function is given by a set of n positive weights p_i ($i = 0, 1, \dots, n - 1$) and reads

$$P(x|\{p_i\}) = \frac{p_x}{\sum_{i=0}^{n-1} p_i} \quad \text{for } x \in \{0, 1, \dots, n - 1\}.$$

The weights p_i have to be passed to the constructor of class `discrete_dist` by some iterator range. The class `discrete_dist` is declared in the header file `trng/discrete_dist.hpp` and its public interface is given as follows:

```

namespace trng {

class discrete_dist {
public:
    typedef int result_type;
    class param_type {
    public:
        template<typename iter>
        explicit param_type(iter first, iter last);
    };

    template<typename iter>
    explicit discrete_dist(iter first, iter last);
    explicit discrete_dist(const param_type &);
    void reset();
    template<typename R>
    int operator()(R &);
    template<typename R>
    int operator()(R &, const param_type &);
    int min() const;
    int max() const;
    param_type param() const;
    void param(const param_type &);
    double pdf(int) const;
    double cdf(int) const;
};

bool operator==(const discrete_dist::param_type &, const discrete_dist::param_type &);
bool operator!=(const discrete_dist::param_type &, const discrete_dist::param_type &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &

```

```

operator<<(std::basic_ostream<char_t, traits_t> &, const discrete_dist::param_type &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, discrete_dist::param_type &);

bool operator==(const discrete_dist &, const discrete_dist &);
bool operator!=(const discrete_dist &, const discrete_dist &);

template<typename char_t, typename traits_t>
std::basic_ostream<char_t, traits_t> &
operator<<(std::basic_ostream<char_t, traits_t> &, const discrete_dist &);
template<typename char_t, typename traits_t>
std::basic_istream<char_t, traits_t> &
operator>>(std::basic_istream<char_t, traits_t> &, discrete_dist &);
}

```

4.3 Function template `generate_canonical`

In this section we describe a function template introduced by [5]. Each function instantiated from the template `generate_canonical` maps the result of a single invocation of a supplied uniform random number generator to one member of the set \mathcal{L} (described below) such that, if the values produced by the generator are uniformly distributed, the results of the instantiation are distributed as uniformly as possible according to the uniformity requirements described below.

Let \mathcal{L} consist of all values t of type `result_type` such that:

- If `result_type` is a floating-point type, `result_type(0) < t < result_type(1)`.
- If `result_type` is a signed or unsigned integral type, `numeric_limits<result_type>::min() ≤ t ≤ numeric_limits<result_type>::max()`.

Obtaining a value in \mathcal{L} can be a useful step in the process of transforming a value generated by a uniform random number generator into a value that can be delivered by a random number distribution. The function template

```

template<class result_type, class UniformRandomNumberGenerator>
result_type generate_canonical(UniformRandomNumberGenerator & g);

```

returns a value from \mathcal{L} by exactly one invocation of `g`, see [5] for details.

5 Installation

To make the installation procedure portable and comfortable, TRNG utilizes the GNU build system. For a proper installation you will need

- GNU autotools (autoconf, automake, libtool) and
- a recent C++ compiler, that has a good template support and knows long long as a build-in data type (e. g. the GNU C++ compiler version 3.0 or newer).

TRNG comes with numerous sample programs, that illustrate the usage of the TRNG library. Some of these sample programs will use external libraries, namely:

- Boost C++ libraries, [4]
- An implementation of the Message Passing Interface (MPI) standard, various open source implementations can be found at [43, 40].

If you want to compile all sample programs, you will have to install these libraries as well. But TRNG does not depend on the libraries listed above by itself.

After the tar-ball had been extracted, you have to call the configure script. This script tries to find your C++ compiler and generates a set of Makefiles. On most Unix-like boxes, just calling

```
bauke@hal:~/trng-4.3$ ./configure
```

will work fine. The configure script can be controlled by various options and shell variables. If no options are provided to configure TRNG will be installed in the /usr/local hierarchy. Call

```
bauke@hal:~/trng-4.3$ ./configure --help
```

to get an overview about all options. Here a complex example: to compile TRNG with the Intel C++ compiler `icc` and to install the library and the header files in `/opt/trng` call

```
bauke@hal:~/trng-4.3$ export CXX=icc
bauke@hal:~/trng-4.3$ export LDFLAGS=-lstdc++
bauke@hal:~/trng-4.3$ ./configure --prefix=/opt/trng
```

After TRNG had been configured, the library will be compiled and installed by the make tool.

```
bauke@hal:~/trng-4.3$ make
bauke@hal:~/trng-4.3$ make install
```

The TRNG library can be uninstalled by calling make with the `uninstall` target.

```
bauke@hal:~/trng-4.3$ make uninstall
```

5 *Installation*

Depending on your system further steps might be necessary to make the TRNG shared library known to the dynamic linker. On a Linux system the system administrator has to call `ldconfig` or you might set the `LD_LIBRARY_PATH` environment variable, see also the `ld.so` man page for further information.

In the source directory `examples` you will find some example programs. These sources can be compiled by

```
bauke@hal:~/trng-4.3/examples$ make examples
```

6 Examples

6.1 Hello world!

In listing 6.1 we present the simplest nontrivial C++ program that produces pseudo-random numbers by TRNG. Whenever one generates random numbers with TRNG at least two header files have to be included, one for a random number engine and one for a distribution function, see lines 4 and 5 in listing 6.1. In lines 9 and 11 respectively a random number engine and a random number distribution are declared. The parameters of a random number distribution object have to be specified by its declaration. In our example random numbers with a normal distribution with mean 6 and standard deviation of 2 are generated. Distribution parameters can be changed at run-time, if necessary. In the loop in lines 13 and 14 the random number engine object `R` and the random number distribution object `normal` are used to generate 1000 random numbers.

The program `hello_world.cc` has to be linked to the TRNG library. Using the GNU C++ compiler we transform the sources by

```
bauke@hal:~$ g++ -o hello_world hello_world.cc -ltrng4
```

into an executable.

In a second example we want to calculate an approximate value for π by a parallel Monte Carlo calculation. The general idea of this calculation is to choose random points in a square with edge length R . Some of these points fall into a sector of a circle in the square, see Figure 6.1. The value of π can be approximated by considering the fraction of points that fall into the

Listing 6.1: A simple TRNG sample program `hello_world.cc` that generates 1000 random variables with normal distribution.

```
1  #include <cstdlib>
2  #include <iostream>
3  // include TRNG header files
4  #include <trng/yarn2.hpp>
5  #include <trng/normal_dist.hpp>
6
7  int main() {
8      // random number engine
9      trng::yarn2 R;
10     // normal distribution with mean 6 and standard deviation 2
11     trng::normal_dist normal(6.0, 2.0);
12     // generate 1000 normal distributed random numbers
13     for (int i=0; i<1000; ++i)
14         std::cout << normal(R) << '\n';
15     return EXIT_SUCCESS;
16 }
```


6 Examples

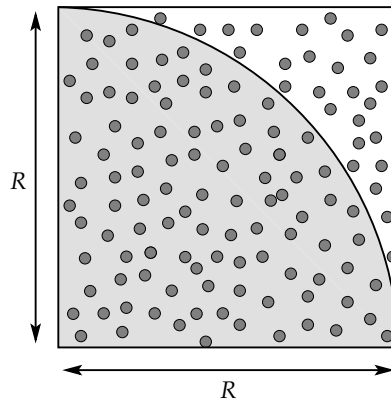


Figure 6.1: The numerical value of π can be estimated by throwing random points into a square.

circle. From the relation

$$\frac{\text{number of points in circle}}{\text{number of points in square}} \approx \frac{\pi R^2/4}{R^2} = \frac{\pi}{4}$$

we conclude

$$\pi \approx 4 \frac{\text{number of points in circle}}{\text{number of points in square}}.$$

In listing 6.2 we use this equation to estimate π . In the for-loop in lines 12 to 16 a random x -coordinate and a random y -coordinate are chosen. Both coordinates are independently uniformly distributed in $[0, 1)$. If $\sqrt{x^2 + y^2} < 1$, or equivalently $x^2 + y^2 < 1$, the point (x, y) lies within the circle. The program draws a huge number of points from the square and counts the number of points lying within the circle and at the end of the program the fraction $4 \cdot (\text{points in circle}) / (\text{points in square})$ is shown as an estimate for π .

Listing 6.2: Sequential Monte Carlo calculation of π .

```

1 #include <cstdlib>
2 #include <iostream>
3 #include <trng/yarn2.hpp>
4 #include <trng/uniform01_dist.hpp>
5
6 int main(int argc, char *argv[]) {
7     const long samples=10000001;           // total number of points in square
8     long in=0;                             // no points in circle
9     trng::yarn2 r;                          // random number engine
10    trng::uniform01_dist u;                 // random number distribution
11    // throw random points into square
12    for (long i=0; i<samples; ++i) {
13        double x=u(r), y=u(r);              // choose random x- and y-coordinates
14        if (x*x+y*y<=1.0)                   // is point in circle?
15            ++in;                           // increase counter
16    }
17    std::cout << "pi = " << 4.0*in/samples << std::endl;
18    return EXIT_SUCCESS;
19 }
```

6.2 Hello parallel world!

TRNG is designed as a random number generator library for sequential as well as for parallel applications. The library does not depend on any particular communication library, it may be utilized with Message Passing Interface (MPI), OpenMP, and as well as with POSIX threads, or any other communication library. This section gives a short tutorial on writing parallel Monte Carlo applications with TRNG and the MPI or OpenMP. Here we cannot give an introduction to MPI or OpenMP, readers, who are not familiar with parallel programming, may consult [44, 2, 47] instead.

How can we parallelize the Monte Carlo calculation of π ? A striking feature of the Monte Carlo π calculation algorithm from the previous section is, that the placement of some point in the square does not affect the placement of other points. In other words: throwing N points into a square is an embarrassingly parallel process. Everything that matters, is the fraction of points in the square that placed into the circle. Keeping this fact in mind the Monte Carlo calculation of π can be parallelized via block splitting or leapfrog method.

6.2.1 Block splitting

Here we choose a parallelization strategy that is based on the block splitting technique, introduced in section 2. A total of N points has to be selected by p processes. We number the

Listing 6.3: Parallel Monte Carlo calculation of π using block splitting and MPI.

```

1  #include <iostream>
2  #include "mpi.h"
3  #include <trng/yarn2.hpp>
4  #include <trng/uniform01_dist.hpp>
5
6  int main(int argc, char *argv[]) {
7      const long samples=10000001;           // total number of points in square
8      trng::yarn2 r;                          // random number engine
9      MPI::Init(argc, argv);                 // initialise MPI environment
10     int size=MPI::COMM_WORLD.Get_size();    // get total number of processes
11     int rank=MPI::COMM_WORLD.Get_rank();    // get rank of current process
12     long in=01;                             // number of points in circle
13     trng::uniform01_dist u;                 // random number distribution
14     r.jump(2*(rank*samples/size));          // jump ahead
15     // throw random points into square and distribute workload over all processes
16     for (long i=rank*samples/size; i<(rank+1)*samples/size; ++i) {
17         double x=u(r), y=u(r);              // choose random x- and y-coordinates
18         if (x*x+y*y<=1.0)                   // is point in circle?
19             ++in;                           // increase counter
20     }
21     // calculate sum of all local variables 'in' and store result in 'in_all' on process 0
22     long in_all;
23     MPI::COMM_WORLD.Reduce(&in, &in_all, 1, MPI::LONG, MPI::SUM, 0);
24     if (rank==0)                           // print result
25         std::cout << "pi = " << 4.0*in_all/samples << std::endl;
26     MPI::Finalize();                        // quit MPI
27     return EXIT_SUCCESS;
28 }
```

Listing 6.4: Parallel Monte Carlo calculation of π using block splitting and OpenMP.

```

1  #include <iostream>
2  #include <omp.h>
3  #include <trng/yarn2.hpp>
4  #include <trng/uniform01_dist.hpp>
5
6  int main(int argc, char *argv[]) {
7      const long samples=10000001;          // total number of points in square
8      long in=01;                          // number of points in circle
9      // distribute workload over all processes
10 #pragma omp parallel
11 {
12     trng::yarn2 r;                        // random number engine
13     int size=omp_get_num_threads();       // get total number of processes
14     int rank=omp_get_thread_num();        // get rank of current process
15     trng::uniform01_dist u;              // random number distribution
16     r.jump(2*(rank*samples/size));        // jump ahead
17     // throw random points into square
18     for (long i=rank*samples/size; i<(rank+1)*samples/size; ++i) {
19         double x=u(r), y=u(r);           // choose random x- and y-coordinates
20         if (x*x+y*y<=1.0)                 // is point in circle?
21 #pragma omp critical
22             ++in;                        // increase counter
23     }
24 }
25 // print result
26 std::cout << "pi = " << 4.0*in/samples << std::endl;
27 return EXIT_SUCCESS;
28 }

```

points from 0 to $N - 1$ and the processes from 0 to $p - 1$ respectively. The number of a process is called its rank. To distribute the workload equally, we split the entire set of N points into p consecutive blocks of about N/p points. To be specific, a process with rank r selects the points with numbers

$$\lfloor N \cdot r/p \rfloor \quad \text{to} \quad \lfloor N \cdot (r+1)/p \rfloor - 1,$$

where $\lfloor \cdot \rfloor$ denotes rounding to zero. Each point is determined by two coordinates and a process with rank r consumes

$$2(\lfloor N \cdot (r+1)/p \rfloor - \lfloor N \cdot r/p \rfloor)$$

random numbers, which are generated by the same random number engine.

All concurrent processes generate random points by their own local copy of the same random number engine. Of course, if all these engines start from the same initial state, they will produce the same sequence of random numbers. For that reason each process jumps $2\lfloor N \cdot r/p \rfloor$ steps ahead, before any random numbers are consumed. This ensures that sequences of random numbers of two different processes never overlap, and furthermore, the outcome of the parallelized program is the same as for the sequential in the previous section, even in its statistical errors.

Listing 6.3 presents an implementation of the parallel Monte Carlo computation of π by MPI, while in listing 6.4 an implementation presented that is based on OpenMP. Note the parenthesis within the argument of the jump method in lines 15 and 17 respectively. Together

with the C++ rounding rules they are the C++ equivalent to the $\lfloor \cdot \rfloor$ function.

There is one important conceptual difference between the MPI version and the OpenMP implementation. While MPI is based on a distributed memory model, OpenMP can utilize shared memory. For that reason the MPI program counts how many points lie in the circle for each process in a process local variable `in`. At the end of the computation the process local variables have to be summed up by `MPI::COMM_WORLD.Reduce` to the (process local) variable `in_all` on the process with rank zero. In a OpenMP program this global reduction can be avoided by using a shared memory variable. But here concurrent write accesses to `in` have to be prevented by the `pragma omp critical` in lines 23 to 24.

6.2.2 Leapfrog

Leapfrog is a convenient approach to derive p non overlapping streams of pseudo-random numbers from a single base stream. As defined in section 3.1 each parallel random number engine provides a `split` method for leapfrog. If `split(p, s)` is called, the internal parameters of the random number engine are changed in such a way, that future calls to `operator()` will generate the s th sub-stream of p sub-streams. Sub-streams are numbered from 0 to $p - 1$.

Listing 6.5: Parallel Monte Carlo calculation of π using leapfrog and MPI.

```

1  #include <iostream>
2  #include "mpi.h"
3  #include <trng/yarn2.hpp>
4  #include <trng/uniform01_dist.hpp>
5
6  int main(int argc, char *argv[]) {
7      const long samples=10000001;          // total number of points in square
8      trng::yarn2 rx, ry;                   // random number engines for x- and y-coordinates
9      MPI::Init(argc, argv);                // initialize MPI environment
10     int size=MPI::COMM_WORLD.Get_size();  // get total number of processes
11     int rank=MPI::COMM_WORLD.Get_rank();  // get rank of current process
12     // split PRN sequences by leapfrog method
13     rx.split(2, 0);                        // choose sub-stream no. 0 out of 2 streams
14     ry.split(2, 1);                        // choose sub-stream no. 1 out of 2 streams
15     rx.split(size, rank);                  // choose sub-stream no. rank out of size streams
16     ry.split(size, rank);                  // choose sub-stream no. rank out of size streams
17     long in=0;                             // number of points in circle
18     trng::uniform01_dist u;                // random number distribution
19     // throw random points into square and distribute workload over all processes
20     for (long i=rank; i<samples; i+=size) {
21         double x=u(rx), y=u(ry);           // choose random x- and y-coordinates
22         if (x*x+y*y<=1.0)                  // is point in circle?
23             ++in;                          // increase counter
24     }
25     // calculate sum of all local variables 'in' and store result in 'in_all' on process 0
26     long in_all;
27     MPI::COMM_WORLD.Reduce(&in, &in_all, 1, MPI::LONG, MPI::SUM, 0);
28     if (rank==0)                          // print result
29         std::cout << "pi = " << 4.0*in_all/samples << std::endl;
30     MPI::Finalize();                      // quit MPI
31     return EXIT_SUCCESS;
32 }
```

Listing 6.6: Parallel Monte Carlo calculation of π using leapfrog and OpenMP.

```

1  #include <iostream>
2  #include <omp.h>
3  #include <trng/yarn2.hpp>
4  #include <trng/uniform01_dist.hpp>
5
6  int main(int argc, char *argv[]) {
7      const long samples=10000001;          // total number of points in square
8      long in=01;                          // no points in circle
9      // distribute workload over all processes
10     #pragma omp parallel
11     {
12         trng::yarn2 rx, ry;                // random number engines for x- and y-coordinates
13         int size=omp_get_num_threads();    // get total number of processes
14         int rank=omp_get_thread_num();     // get rank of current process
15         // split PRN sequences by leapfrog method
16         rx.split(2, 0);                    // choose sub-stream no. 0 out of 2 streams
17         ry.split(2, 1);                    // choose sub-stream no. 1 out of 2 streams
18         rx.split(size, rank);              // choose sub-stream no. rank out of size streams
19         ry.split(size, rank);              // choose sub-stream no. rank out of size streams
20         trng::uniform01_dist u;           // random number distribution
21         // throw random points into square
22         for (long i=rank; i<samples; i+=size) {
23             double x=u(rx), y=u(ry);       // choose random x- and y-coordinates
24             if (x*x+y*y<=1.0)               // is point in circle?
25             #pragma omp critical
26                 ++in;                      // increase counter
27         }
28     }
29     // print result
30     std::cout << "pi = " << 4.0*in/samples << std::endl;
31     return EXIT_SUCCESS;
32 }

```

Changing line 15 or line 17 in listing 6.3 or listing 6.4 respectively, which reads

```
r.jump(2*(rank*samples/size)); // jump ahead
```

into

```
r.split(size, rank);           // choose sub-stream no. rank out of size streams
```

provides different statistically independent sub-streams of pseudo-random numbers to each process.

But note, the pseudo-random numbers of the base stream are now utilized in a completely different fashion. The sequential program and also the two on block splitting based programs from section 6.2.1 determine the position of a point (its x - and y -coordinate) by two consecutive pseudo-random numbers of the base sequence. After calling `split(size, rank)` consecutive calls to `operator()` will return pseudo-random numbers, that are no longer neighboring numbers of the base sequence. In fact they have a distance of `size` with respect to the original sequence of pseudo-random numbers. For that reason the proposed replacement of the call of the `jump` method to a call to the `split` method will result in another value for the approximation of π with another statistical error.

6 Examples

To prevent this issue, we use the fact that the leapfrog method can be applied several times to a sequence of pseudo-random numbers by successive calls to `split`. Each time `split` is invoked the sequence is split into further sub-sequences. In listing 6.5 and listing 6.6 it is shown how this works. Both programs start with two random number engines of the same kind.

```
trng::yarn2 rx, ry; // random number engines for x- and y-coordinates
```

Later all x - and y -coordinates will be determined exclusively by one of these random number engines. But without any manipulations of the internal status via `jump` or `split` method, both engines will return the same sequences of pseudo-random numbers. Therefore, if the coordinates of each point are chosen by calling `operator()` of `rx` and `ry` once, all points will lie on the diagonal of the square. For that reason the sequences are split by

```
rx.split(2, 0); // choose sub-stream no. 0 out of 2 streams  
ry.split(2, 1); // choose sub-stream no. 1 out of 2 streams
```

into two non overlapping sequences. Now successive calls to `operator()` will return different sequences of pseudo-random numbers and the points are uniformly distributed over the square. But still each process consumes the same two sequences of random numbers. However, this can be solved by calling the `split` method a second time.

```
rx.split(size, rank); // choose sub-stream no. rank out of size streams  
ry.split(size, rank); // choose sub-stream no. rank out of size streams
```

6.2.3 Block splitting or leapfrog?

TRNG provides two powerful techniques for parallelizing streams of pseudo-random numbers, namely block splitting and leapfrog. Which one to choose, depends highly on the structure of your Monte Carlo algorithm and your needs.

In the simplest case, each process of a parallel Monte Carlo application with a fixed number of processes p (that does not change at run time) has just to be equipped with some source of pseudo-random numbers and the only requirement on the p streams of pseudo-random numbers is, that they do not overlap with any stream of pseudo-random numbers on any other process. In this case it is sufficient to use a single random number engine of the same type for each of the p process. Different streams are deviated by the leapfrog method and calling the `split` method of a pseudo-random number engine object after these random number engines have been initialized with the same parameters and the same seed. Of course with this simple minded approach the outcome of a Monte Carlo application (and the actual statistical errors) will depend on the number of processes.

On the other hand it is often desirable to design a parallel Monte Carlo algorithm in such a way that its outcome is independent of the number of processes. That means, that the Monte Carlo algorithm plays fair, see also section 2.3. Usually this additional constraint can be fulfilled by a creative combination of block splitting, leapfrog method and using more than one random number engine per processor. The previous sections gave already some elementary examples, how this can be achieved. But in general this can be quite intricate. Therefore we give some general guidelines.

- Identify the inherently parallel parts of the Monte Carlo algorithm. Which steps of the Monte Carlo algorithm cannot be parallelized?

6 Examples

- Break the parallelizable tasks into p (p number of processes) smaller sub-parts of approximately equal size.
- Is the number of pseudo-random numbers, that is consumed by a parallelizable task (before it is divided into subparts), constant or does it change at runtime? If it is constant, break up the sequence of a single pseudo-random number engine into sub-streams in such a way, that it mimics the way in which the parallelizable task is split into independent sub-problems. This can always be achieved by calling the `split` or the `jump` method of a random number engine object.
- If the number of pseudo-random numbers, that is consumed by a parallelizable task, is not constant or cannot be determined a priori, e. g. because this number itself is a function of the random number sequence, an upper bound for this number may be estimated. With this number a Monte Carlo algorithm can often be parallelized as if the number of consumed random numbers was fixed.

To make these advises somewhat more clear, we give a further example. Imagine the simulation of a site percolation process [52] on a two-dimensional square lattice of size $N = N_x \times N_y$. In site percolation each site of the lattice is occupied with probability P independently of the other sites and clusters of neighboring occupied sites are constructed afterward. Once these clusters are known, one can answer for a particular realization of occupied sites a lot of questions, that arise in percolation theory. Is there a spanning cluster, that connects the lower line of the grid and its upper line? What is the size of the largest cluster? And so on. How can we parallelize such a Monte Carlo simulation for site percolation?

The easiest way to do this, is not to parallelize at all. At least not the analysis of a single realization of occupied sites itself. Usually one is not interested in the analysis of a single realization of occupied sites by itself, but one wants to know statistical properties of site percolation (or another problem) that arise after averaging over many, lets say M , realizations of systems of the same kind. Its is quite natural to spread the workload over p processors in such a way, that each process analyzes each p th lattice of the M lattices. If we number the processes by its rank from 0 to $p - 1$ and the lattices from 0 to $M - 1$, each process starts with a lattice which number equals the process' rank. Thereafter each process can skip $p - 1$ lattices, because these are handled by other processes, and continue with the next lattice. Of course each process has not only to skip the work, that is done by other process, but also the pseudo-random numbers, that would be consumed by analyzing the skipped lattices. Listing 6.7 gives a sketch of such a parallelized site percolation program.

Unfortunately it is not always possible to parallelize a Monte Carlo simulation in such a coarse-grained fashion like in the last example. Sometimes (e. g. in the Swendsen-Wang-cluster-algorithm [53, 42]) the generation and the analysis of a single lattice has to be parallelized by itself. For that reason we split the lattice into $p_x \times p_y$ sub-lattices in such a way that the number of parallel processes p equals $p_x \times p_y$ and $p_x \approx p_y$. Each process is responsible for one of the sub-lattices and uses the same random number engine. This generic parallelization paradigm is also known as domain decomposition.

To make the site percolation lattice generation independent of the number processes and thus independent of the details of the lattice partition, some numbers within the stream of pseudo-random numbers of the random number engine have to be skipped by the `jump` method. If we determine the state (occupied or not occupied) of the sites in a row-major fashion, the `jump` method has to be called, whenever a process has filled a row of its sub-lattice. Of course each

Listing 6.7: Sketch of a coarse-grained parallel Monte Carlo simulation of site percolation via MPI. The program creates many realizations of lattices with randomly occupied sites. Each realization is generated by a single process.

```

1  #include <cstdlib>
2  #include <trng/yarn2.hpp>
3  #include <trng/uniform01_dist.hpp>
4  #include "mpi.h"
5
6  const int number_of_realizations=1000;
7  const int Nx=250, Ny=200;           // grid size
8  const int number_of_PRNs_per_sweep=Nx*Ny;
9  int site[Nx][Ny];                  // lattice
10 const double P=0.46;                // occupation probability
11
12 int main(int argc, char *argv[]) {
13     MPI::Init(argc, argv);           // initialize MPI environment
14     int size=MPI::COMM_WORLD.Get_size(); // get total number of processes
15     int rank=MPI::COMM_WORLD.Get_rank(); // get rank of current process
16     trng::yarn2 R;                   // random number engine
17     trng::uniform01_dist u;          // random number distribution
18     // skip random numbers that are consumed by other processes
19     R.jump(rank*number_of_PRNs_per_sweep);
20     for (int i=rank; i<number_of_realizations; i+=size) {
21         // consume Nx * Ny pseudo-random numbers
22         for (int x=0; x<Nx; ++x)
23             for (int y=0; y<Ny; ++y)
24                 if (u(R)<P)
25                     site[x][y]=1;    // site is occupied
26                 else
27                     site[x][y]=0;    // site is not occupied
28         // skip random numbers that are consumed by other processes
29         R.jump((size-1)*number_of_PRNs_per_sweep);
30         // analyze lattice
31         // ... source omitted
32     }
33     MPI::Finalize();                 // quit MPI
34     return EXIT_SUCCESS;
35 }

```

process has to skip a certain amount of pseudo-random numbers at the start of the simulation, too.

Listing 6.8 shows the outline of a fine-grained parallel Monte Carlo simulation of site percolation via MPI, where each single lattice generation is done in parallel via domain decomposition. This program shows two noteworthy implementation details. First the program uses a runtime generated Cartesian communicator rather than the standard communicator `MPI::COMM_WORLD` as seen in the MPI examples so far. Such a communicator reflects the special topology of the domain decomposition and eases its implementation significantly. The number of sub-lattices in each dimension, p_x and p_y respectively, is determined by `MPI::Compute_dims`, see [44, 2] for details. Its result (returned in the field `dims`) determines the topology of the Cartesian communicator `Comm`. Another nice feature of the example code in listing 6.8 is, that it does not assume, that the number of sites in any dimension is a multiple of the number of sub-lattices

6 Examples

Listing 6.8: Sketch of a fine-grained parallel Monte Carlo simulation of site percolation via MPI. The program creates many realizations of lattices with randomly occupied sites. Each realization is generated by all processes together, workload is distributed by domain decomposition.

```

1  #include <cstdlib>
2  #include <new>
3  #include <trng/yarn2.hpp>
4  #include <trng/uniform01_dist.hpp>
5  #include "mpi.h"
6
7  const int number_of_realizations=1000;
8  const int Nx=250, Ny=200;           // grid size
9  const double P=0.46;                // occupation probability
10
11 int main(int argc, char *argv[]) {
12     MPI::Init(argc, argv);           // initialize MPI environment
13     int size=MPI::COMM_WORLD.Get_size(); // get total number of processes
14     // create a two-dimensional Cartesian communicator
15     int dims[2] = {0, 0};            // number of processes in each dimension
16     int coords[2];                   // coordinates of current process within the grid
17     bool periods[2] = { false, false }; // no periodic boundary conditions
18     // calculate a balanced grid partitioning such that size = dims[0]*dims[1]
19     MPI::Compute_dims(MPI::COMM_WORLD.Get_size(), 2, dims);
20     MPI::Cartcomm Comm=MPI::COMM_WORLD.Create_cart(2, dims, periods, true);
21     int rank=Comm.Get_rank();         // get rank of current process
22     Comm.Get_coords(rank, 2, coords); // get coordinates of current process
23     // determine section of current process
24     int x0=coords[0]*Nx/dims[0], x1=(coords[0]+1)*Nx/dims[0], Nx1=x1-x0,
25         y0=coords[1]*Ny/dims[1], y1=(coords[1]+1)*Ny/dims[1], Ny1=y1-y0;
26     int *site=new int[Nx1*Ny1];      // allocate memory to store a sublattice
27     trng::yarn2 R;                   // random number engine
28     trng::uniform01_dist u;          // random number distribution
29     // skip random numbers that are consumed by other processes
30     R.jump(Nx*y0+x0);
31     for (int i=0; i<number_of_realizations; ++i) {
32         // consume Nx1 * Ny1 pseudo-random numbers
33         int *s=site;
34         for (int y=y0; y<y1; ++y) {
35             for (int x=x0; x<x1; ++x) {
36                 if (u(R)<P)
37                     *s=1;              // site is occupied
38                 else
39                     *s=0;              // site is not occupied
40                 ++s;
41             }
42             // skip random numbers that are consumed by other processes
43             R.jump(Nx-Nx1);
44         }
45         // skip random numbers that are consumed by other processes
46         R.jump(Nx*(Ny-Ny1));
47         // analyze lattice
48         // ... source omitted
49     }
50     MPI::Finalize();                 // quit MPI
51     return EXIT_SUCCESS;
52 }

```

in this dimension. So the sizes of the sub-lattices can vary slightly from process to process. The precise range of coordinates, that each process is responsible for, is calculated in lines 24 and 25.

Skipping numbers in a pseudo-random number sequence via `jump` is not for free. Of course it is so smart, that it can jump ahead without actually generating the numbers that have to be skipped. But the complexity of `jump` grows logarithmically in its argument. If the domain decomposition is coarse-grained enough, the overhead introduced by skipping numbers via `jump` can be neglected. But if the number of processes, that generate a site percolation lattice, becomes larger and larger, at a certain point this overhead can no longer be ignored and starts to limit the speedup, that is achievable by parallelization. Finding the right level of granularity is a general problem in parallel computing. On one hand one wants to use a large number of processes to attain a large speedup, on the other hand, the relative portion of the inherent sequential part of a program and the overhead introduced by the parallelization grow with the number of processes as well. This fact is also known as Amdahl's law.

6.3 Using TRNG with STL and Boost

Whenever large scale Monte Carlo applications are written, they will not base on TRNG solely, but also on other libraries, e. g. the C++ Standard Template Library (STL) or Boost [4]. In this section we show, how to use TRNG in combination with the STL, especially its containers and algorithms and the `bind` facility of Boost¹. We assume you are familiar with the concepts of the C++ STL, otherwise we suggest to read [41].

Imagine a C++ array or an STL container like a vector or a list of integers that has to be populated by random numbers with a given distribution. This can be achieved by a simple loop.

```
trng::yarn2 R;           // random number engine
trng::uniform_int_dist U(0, 100); // random number distribution
std::vector<long> v(10); // vector of long with 10 elements
for (std::vector<long>::iterator i(v.begin()), end(v.end()); i!=end; ++i)
    *i=U(R);             // generate a random number from distribution U by engine R
```

This loop looks innocent, but it is not. It is error-prone and it is not obvious what is actually effected by the loop. The loop is error-prone because the programmer has to take care that the type of the iterator `i` fits to the container. Things become much more handy, if STL algorithms like `std::generate` are used.

The template function `std::generate` takes an iterator range and a function object that takes no arguments as its arguments. The prototype of this function reads

```
namespace std {

    template <class ForwardIterator, class Generator>
    void generate(ForwardIterator first, ForwardIterator last, Generator gen);

}
```

and it assigns the result of invoking `gen` to each element in the range `[first, last)`. Random number distributions as introduced in section 3.2 do not meet the requirements of `std::`

¹The `bind` facility of Boost will be part of future versions of the STL.

6 Examples

generate, because their overloaded call operator requires at least one argument, namely a random number engine, see Table 3.2. For that reason we need a function adapter, that makes random number distributions compatible with `std::generate`. The following template class `binder_cl` is such a function adapter.

```
template<typename PRN_dist_t, typename PRN_engine_t>
class binder_cl {
    PRN_dist_t &dist;
    PRN_engine_t &engine;
public:
    binder_cl(PRN_dist_t &dist, PRN_engine_t &engine) : dist(dist), engine(engine) {
    }
    typename PRN_dist_t::result_type operator()() {
        return dist(engine);
    }
};
```

It holds a reference to a random number engine and a reference to a random number distribution respectively as private data members. Its call operator calls the call operator of the random number distribution with the random number engine as its argument. With this template class an STL container `v` can be filled by

```
trng::yarn2 R; // random number engine
trng::uniform_int_dist U(0, 100); // random number distribution
std::vector<long> v(10); // vector of long with 10 elements
std::generate(v.begin(), v.end(), binder_cl<trng::uniform_int_dist, trng::yarn2>(U, R));
```

The statement

```
binder_cl<trng::uniform_int_dist, trng::yarn2>(U, R)
```

creates a temporary anonymous object of the class `binder_cl<trng::uniform_int_dist, trng::yarn2>`, which is a instantiation of the template class `binder_cl`. Up to now we have not gained very much. Now we can replace an explicit loop by a template function `std::generate`, but the syntax is clumsy and as error-prone as the explicit loop, because the types, that specify the template class `binder_cl` have to be given explicitly. This is a common obstacle of generic programming in C++ but this can be avoided by a further helper function `make_binder`.

```
template<typename PRN_dist_t, typename PRN_engine_t>
inline
binder_cl<PRN_dist_t, PRN_engine_t> make_binder(PRN_dist_t &dist, PRN_engine_t &engine) {
    return binder_cl<PRN_dist_t, PRN_engine_t>(dist, engine);
}
```

With this little helper function the line

```
std::generate(v.begin(), v.end(), binder_cl<trng::uniform_int_dist, trng::yarn2>(U, R));
```

can be simplified to

```
std::generate(v.begin(), v.end(), make_binder(U, R));
```

Adapting function objects to functions and algorithms is a common task in generic programming. The C++ STL is equipped with some adapter functions like `std::bind1st` or `std::bind2nd`, but they are of limited use and from time to time further adapter functions

6 Examples

have to be created, as shown in the preceding paragraphs. The bind facility of the Boost library generalizes the STL function adapters and we do not have to write our own function adapters. Here we can give only a glimpse of the bind facility, everyone how wants to explore the full capabilities of `boost::bind` should read the Boost documentation.

The boost equivalent to

```
std::generate(v.begin(), v.end(), make_binder(U, R));
```

reads

```
std::generate(v.begin(), v.end(), boost::bind(U, boost::ref(R)));
```

In this example the function `boost::bind` returns a temporary function object whose call operator requires no arguments. The function `boost::ref` assures that the temporary function object holds a reference to the random number engine `R`, otherwise it would contain a copy of `R`. Omitting `boost::ref` may have unexpected side effects, e. g. the loop

```
for (int i(0); i<10; ++i)
    std::generate(v.begin(), v.end(), boost::bind(U, R));
```

would fill the vector `v` ten times with random numbers, each time with the same set of random numbers. Because `boost::bind` generates at each call to `std::generate` a copy of the random number engine `R` and this copy determines the random values in `v`, but not the random number engine `R` itself. As a consequence of this copy process `std::generate` generates random numbers by a random number engine, that starts with the same internal state in each cycle of the loop.

Listing 6.9 demonstrates all the techniques for binding function arguments that have been discussed in this section. Additionally it shows that TRNG random number engine meet the requirements of the STL function `std::random_shuffle` directly, no function adaption via `boost::bind` is needed.

Listing 6.9: This demo program demonstrates the interplay of TRNG, the C++ STL and the bind facility of Boost.

```
1  #include <cstdlib>
2  #include <iostream>
3  #include <vector>
4  #include <algorithm>
5  #include <trng/config.hpp>
6  #include <trng/yarn2.hpp>
7  #include <trng/uniform_int_dist.hpp>
8  #if defined HAVE_BOOST
9      #include <boost/bind.hpp>
10 #else
11
12     // helper class
13     template<typename PRN_dist_t, typename PRN_engine_t>
14     class binder_cl {
15     public:
16         PRN_dist_t &dist;
17         PRN_engine_t &engine;
18         binder_cl(PRN_dist_t &dist, PRN_engine_t &engine) : dist(dist), engine(engine) {
19         }
20         typename PRN_dist_t::result_type operator()() {
21             return dist(engine);
22         }
23     };
```

6 Examples

```
23 };
24
25 // convenience function
26 template<typename PRN_dist_t, typename PRN_engine_t>
27 inline
28 binder_cl<PRN_dist_t, PRN_engine_t> make_binder(PRN_dist_t &dist, PRN_engine_t &engine) {
29     return binder_cl<PRN_dist_t, PRN_engine_t>(dist, engine);
30 }
31
32 #endif
33
34
35 // print an iterator range to stdout
36 template<typename iter>
37 void print_range(iter i1, iter i2) {
38     while (i1!=i2) std::cout << (*(i1++)) << '\t';
39     std::cout << "\n\n";
40 }
41
42 int main() {
43     trng::yarn2 R;
44     trng::uniform_int_dist U(0, 100);
45     std::vector<long> v(10);
46
47     std::cout << "random number generation by call operator\n";
48     for (std::vector<long>::size_type i=0; i<v.size(); ++i)
49         v[i]=U(R);
50     print_range(v.begin(), v.end());
51     std::vector<long> w(12);
52     #if defined HAVE_BOOST
53     std::cout << "random number generation by std::generate\n";
54     std::generate(w.begin(), w.end(), boost::bind(U, boost::ref(R)));
55     print_range(w.begin(), w.end());
56     std::cout << "random number generation by std::generate\n";
57     std::generate(w.begin(), w.end(), boost::bind(U, boost::ref(R)));
58     print_range(w.begin(), w.end());
59     #else
60     std::cout << "random number generation by std::generate\n";
61     std::generate(w.begin(), w.end(), make_binder(U, R));
62     print_range(w.begin(), w.end());
63     std::cout << "random number generation by std::generate\n";
64     std::generate(w.begin(), w.end(), make_binder(U, R));
65     print_range(w.begin(), w.end());
66     #endif
67     std::cout << "same sequence as above, but in a random shuffled order\n";
68     std::random_shuffle(w.begin(), w.end(), R);
69     print_range(w.begin(), w.end());
70     return EXIT_SUCCESS;
71 }
```

7 Implementation details and efficiency

Random number engines `trng::mrng`, `trng::mrgns`, `trng::yarnn`, and `trng::yarnns` utilize LFSR sequences

$$r_i = a_1 \cdot r_{i-1} + a_2 \cdot r_{i-2} + \dots + a_n \cdot r_{i-n} \bmod m \quad (7.1)$$

over a prime field \mathbb{F}_m . The modulus m may be any prime. But LFSR sequences over \mathbb{F}_2 have found much more proliferation in the random number generation business than LFSR sequences over other prime fields. LFSR sequences over general prime fields have been proposed in the literature [16, 23, 20] as PRNGs. But so far, they found less attention by practitioners because it is not straight forward to implement LFSR sequences over \mathbb{F}_m efficiently, if m is a large prime, especially if m of the order of the largest in a single computer word representable integer. For that reason, we present some implementation techniques.

We assume that all integer arithmetic is done in w -bit registers and $m < 2^{w-1}$. Under this condition addition of modulo m can be done without overflow problems. But multiplying two $(w-1)$ -bit integers modulo m is not straightforward because the intermediate product has $2(w-1)$ significant bits and cannot be stored in a w -bit register. For the special case $a_k < \sqrt{m}$ Schrage [50] showed how to calculate $a_k \cdot r_{i-k} \bmod m$ without overflow. Based on this technique a portable implementation of LFSR sequences with coefficients $a_k < \sqrt{m}$ is presented in [24]. For parallel PRNGs this methods do not apply because the leapfrog method may yield coefficients that violate this condition. Knuth [20, section 3.2.1.1] proposed a generalization of Schrage's method for arbitrary positive factors less than m , but this method requires up to twelve multiplications and divisions and is therefore not very efficient.

The only way to implement (2.6) without additional measures to circumvent overflow problems is to restrict m to $m < 2^{w/2}$. On machines with 32-bit registers, 16 random bits per number is not enough for some applications. Fortunately today's C compiler provide fast 64-bit-arithmetic even on 32-CPU's and genuine 64-CPU's become more and more common. This allows us to increase m to 32.

7.1 Efficient modular reduction

Since the modulo operation in (2.6) is usually slower than other integer operations like addition, multiplication, Boolean operations or shifting, it has a significant impact on the total performance of PRNGs based on LFSR sequences. If the modulus is a Mersenne Prime $m = 2^e - 1$, however, the modulo operation can be done using only a few additions, Boolean operations and shift operations [45].

A summand $s = a_k \cdot r_{i-k}$ in (2.6) will never exceed $(m-1)^2 = (2^e - 2)^2$ and for each positive integer $s \in [0, (2^e - 1)^2]$ there is a unique decomposition of s into

$$s = r \cdot 2^e + q \quad \text{with} \quad 0 \leq q < 2^e. \quad (7.2)$$

From this decomposition we conclude

$$\begin{aligned} s - r \cdot 2^e &= q \\ s - r(2^e - 1) &= q + r \\ s \bmod (2^e - 1) &= q + r \bmod (2^e - 1) \end{aligned}$$

and r and q are bounded from above by

$$q < 2^e \quad \text{and} \quad r \leq \lfloor (2^e - 2)^2 / 2^e \rfloor < 2^e - 2$$

respectively, and therefore

$$q + r < 2^e + 2^e - 2 = 2m.$$

So if $m = 2^e - 1$ and $s \leq (m - 1)^2$, $x = s \bmod m$ can be calculated solely by shift operations, Boolean operations and addition, viz

$$x = (s \bmod 2^e) + \lfloor s / 2^e \rfloor. \quad (7.3)$$

If (7.3) yields a value $x \geq m$ we simply subtract m .

From a computational point of view Mersenne Prime moduli are optimal and we propose to choose the modulus $m = 2^{31} - 1$. This is the largest positive integer that can be represented by a signed 32-bit integer variable, and it is also a Mersenne Prime. On the other hand our theoretical considerations favor Sophie-Germain Prime moduli, for which (7.3) does not apply directly. But one can generalize (7.3) to moduli $2^e - k$ [34]. Again we start from a decomposition of s into

$$s = r \cdot 2^e + q \quad \text{with} \quad 0 \leq q < 2^e, \quad (7.4)$$

and conclude

$$\begin{aligned} s - r \cdot 2^e &= q \\ s - r(2^e - k) &= q + kr \\ s \bmod (2^e - k) &= q + kr \bmod (2^e - k). \end{aligned}$$

The sum $s' = q + kr$ exceeds the modulus at most by a factor $k + 1$, because by applying

$$q < 2^e \quad \text{and} \quad r \leq \lfloor (2^e - k - 1)^2 / 2^e \rfloor < 2^e - k - 1$$

we get the bound

$$q + kr < 2^e + k(2^e - k - 1) = (k + 1)m.$$

In addition by the decomposition of $s' = q + kr$

$$s' = r' \cdot 2^e + q' \quad \text{with} \quad 0 \leq q' < 2^e,$$

it follows

$$s \bmod (2^e - k) = s' \bmod (2^e - k) = q' + kr' \bmod (2^e - k),$$

and this time the bounds

$$q' < 2^e \quad \text{and} \quad r' \leq \lfloor (k + 1)(2^e - k) / 2^e \rfloor < k + 1$$

and

$$q' + kr' < 2^e + k(k+1) = m + k(k+2).$$

hold. Therefore if $m = 2^e - k$, $s \leq (m - k)^2$ and $k(k+2) \leq m$, $x = s \bmod m$ can be calculated solely by shift operations, Boolean operations and addition, viz

$$\begin{aligned} s' &= (s \bmod 2^e) + k \lfloor s/2^e \rfloor \\ x &= (s' \bmod 2^e) + k \lfloor s'/2^e \rfloor. \end{aligned} \tag{7.5}$$

If (7.5) yields a value $x \geq m$, a single subtraction of m will complete the modular reduction. To carry out (7.5) twice as many operations as for (7.3) are needed. But (7.5) applies for all moduli $m = 2^e - k$ with $k(k+2) \leq m$.

7.2 Fast delinearization

YARN generators hide linear structures of LFSR sequences q_i by raising a generating element g to the power $g^{q_i} \bmod m$. This can be done efficiently by binary exponentiation, which takes $\mathcal{O}(\log m)$ steps. But considering LFSR sequences with only a few feedback taps ($n \leq 6$) and $m \approx 2^{31}$ even fast exponentiation is significantly more expensive than a single iteration of (2.6). Therefore we propose to implement exponentiation by table look up. If m is a $2e'$ -bit number we apply the decomposition

$$\begin{aligned} q_i &= q_{i,1} \cdot 2^{e'} + q_{i,0} \quad \text{with} \\ q_{i,1} &= \lfloor q_i/2^{e'} \rfloor, \quad q_{i,0} = q_i \bmod 2^{e'} \end{aligned} \tag{7.6}$$

and use the identity

$$r_i = g^{q_i} \bmod m = (g^{2^{e'}})^{q_{i,1}} \cdot g^{q_{i,0}} \bmod m \tag{7.7}$$

to calculate $g^{q_i} \bmod m$ by two table look-ups and one multiplication modulo m . If $m < 2^{31}$ the tables for $(g^{2^{e'}})^{q_{i,1}} \bmod m$ and $g^{q_{i,0}} \bmod m$ have 2^{16} and 2^{15} entries respectively and fit easily into the cache of modern CPUs.

7.3 Performance

By TRNG we provide an optimized PRNG library. The implementation uses 64-bit-arithmetic, fast modular reduction (7.3) and (7.5) and exponentiation by table look-up (7.7) to implement PRNGs based on LFSR sequences over prime fields, with Mersenne or Sophie-Germain Prime modulus. PRNGs of TRNG are able to compete with other sequential PRNGs in terms of speed and statistical properties but do support block splitting and leapfrog, too. Table 7.1 shows some benchmark results. For this benchmark 2^{26} PRNs were generated and the execution time was measured to compute how many PRNs each PRNG is able to generate per second. Apparently the performance of the PRNGs of TRNG compete quite well with popular PRNGs like the Mersenne Twister (boost : : mt19937 and mt19937), lagged Fibonacci generators (LFSR sequences over \mathbb{F}_2) or RANLUX that can be found in the Boost library [4] or the GNU Scientific Library [13].

Table 7.1: Performance of various random number engines from TRNG and Boost. Test program was compiled and executed on a Pentium IV 3 GHz with an Intel C++ compiler version 9.1 and option -O3.

generator	PRNs per second
TRNG	
trng::lcg64	$156 \cdot 10^6$
trng::mrg2	$48 \cdot 10^6$
trng::mrg3	$42 \cdot 10^6$
trng::mrg3s	$25 \cdot 10^6$
trng::mrg4	$38 \cdot 10^6$
trng::mrg5	$35 \cdot 10^6$
trng::mrg5s	$17 \cdot 10^6$
trng::yarn2	$26 \cdot 10^6$
trng::yarn3	$23 \cdot 10^6$
trng::yarn3s	$12 \cdot 10^6$
trng::yarn4	$20 \cdot 10^6$
trng::yarn5	$22 \cdot 10^6$
trng::yarn5s	$9 \cdot 10^6$
Boost	
boost::minstd_rand	$77 \cdot 10^6$
boost::ecuyer1988	$36 \cdot 10^6$
boost::kreutzer1986	$69 \cdot 10^6$
boost::hellekalek1995	$2 \cdot 10^6$
boost::mt11213b	$74 \cdot 10^6$
boost::mt19937	$74 \cdot 10^6$
boost::lagged_fibonacci607	$43 \cdot 10^6$
boost::lagged_fibonacci1279	$43 \cdot 10^6$
boost::lagged_fibonacci2281	$47 \cdot 10^6$
boost::lagged_fibonacci3217	$48 \cdot 10^6$
boost::lagged_fibonacci4423	$48 \cdot 10^6$
boost::lagged_fibonacci9689	$63 \cdot 10^6$
boost::lagged_fibonacci19937	$48 \cdot 10^6$
boost::lagged_fibonacci23209	$63 \cdot 10^6$
boost::lagged_fibonacci44497	$63 \cdot 10^6$
GNU Scientific Library	
mt19937	$41 \cdot 10^6$
r250	$85 \cdot 10^6$
gfsr4	$77 \cdot 10^6$
ran2	$27 \cdot 10^6$
ranlux	$8 \cdot 10^6$
ranlux389	$5 \cdot 10^6$

8 Quality

Sequences of PRNs are sequences of deterministic numbers, that try to mimic true random numbers, and one may wonder, how close sequences produced by TRNG can come to sequences of real random numbers? This question can be answered (at least partly) by statistical tests. One can apply a battery of tests on a generator, and the more tests a generator can pass, the better its quality. One distinguishes empirical and theoretical test procedures.

Empirical tests take a finite sequence of PRNs and compute certain statistics, e. g. chi-square or Kolmogorov-Smirnov statistics, to judge the generator as “random” or not. The test statistic is a random variate with a probability distribution, that can be calculated under the assumption that the test statistic is a function of true random numbers. This probability distribution is used to judge a finite sequence of PRNs as possibly random or non-random. For example in an actual test we may find a value of the test statistic that is so large (or small) that such a value or a larger (or smaller) value can be found by chance for true random numbers with a probability of 5 % only. In this case we assume the PRNG has failed the test and its sequence of PRNs behaves non-random. But note, we may be wrong, there is a 5 % probability that we have just seen normal statistical deviations. Therefore a statistical test should be applied several times. If the PRNG fails more often than it can be explained by normal statistical deviations, it has a serious flaw and should be rejected as non-random.

While empirical tests focus only on the statistical properties of a finite stream of PRNs and ignore all the details of the underlying PRNG algorithm, theoretical tests analyze the PRNG algorithm itself by number-theoretic methods and establish a priori characteristics of the PRN sequence. These a priori characteristics may be used to choose good parameter sets for a certain class of PRNGs, e. g. the coefficients of the LFSR sequences in the random number engines `trng::mrng` and `trng::yarn` (see section 4.1) have been found by an extensive computer search [24] and give good results in the spectral test [20], the most important theoretical test for this class of generators.

On one hand the more kinds of statistical test procedures a PRNG masters, the more we will trust its statistical properties. On the other hand statistical test can never prove that an finite sequence of numbers is “random” or not. Knuth writes in [20]:

“In practice, we apply about half a dozen different kinds of statistical tests on a sequence, and if it passes them satisfactorily we consider it to be random—it is then presumed innocent until proven guilty.”

All PRNGs of TRNG and sub-streams of them have been subject to different statistical tests. In respect of these tests the generator you find in TRNG are comparable to other well-known high quality generators like the mersenne twister generator [35]. The tables 8.1 to 8.14 present results of various statistical tests of streams of pseudo-random numbers, that are generated by PRNGs of TRNG with default parameters and no leapfrog splitting. Each test was applied eight times and the tables 8.1 to 8.14 show how often each test has been failed. Note, at a confidence level of 0.1 or 0.9 even a perfect random number generator will “fail” these tests on average in one of ten cases. All statistical tests are implemented by the Random Number

Generator Test Suite (RNGTS) [48]¹. A detailed descriptions of the statistical tests can be found on the RNGTS web site or in [20].

Table 8.1: Test results for random number engine `trng::lcg64`.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Chi-Square-Uniformity-Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
n-Block-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	0 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	1 of 8 failed	4 of 8 failed	2 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Poker-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	5 of 8 failed	3 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Random-Walk Test	2 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	1 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Squeeze-Test	3 of 8 failed	3 of 8 failed	3 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Craps-Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed

¹We had to apply some minor modifications to RNGTS in order to adapt this test suite to TRNG.

Table 8.2: Test results for random number engine trng: :mrg2.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	2 of 8 failed	3 of 8 failed	3 of 8 failed	1 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
n-Block-Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	1 of 8 failed	3 of 8 failed	3 of 8 failed
Serial correlation Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Random-Walk Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Squeeze-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	2 of 8 failed	4 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	1 of 8 failed	2 of 8 failed	4 of 8 failed	3 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed

Table 8.3: Test results for random number engine trng: :mrg3.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	1 of 8 failed	1 of 8 failed	5 of 8 failed	3 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
n-Block-Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	3 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Poker-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	2 of 8 failed
Serial correlation Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Random-Walk Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Serial Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Squeeze-Test	2 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	3 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	2 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	2 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed

Table 8.4: Test results for random number engine trng: :mrg3s.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	0 of 8 failed	1 of 8 failed	3 of 8 failed	2 of 8 failed
Chi-Square-Uniformity-Test	1 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Gap-Test	2 of 8 failed	3 of 8 failed	1 of 8 failed	0 of 8 failed
n-Block-Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	2 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Permutation-Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Poker-Test	2 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	1 of 8 failed
Serial correlation Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Random-Walk Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Serial Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	1 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Squeeze-Test	1 of 8 failed	1 of 8 failed	3 of 8 failed	3 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	2 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed

Table 8.5: Test results for random number engine trng: :mrg4.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
n-Block-Test	0 of 8 failed	1 of 8 failed	4 of 8 failed	2 of 8 failed
Ising-Model-Test (energy)	1 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	0 of 8 failed
Permutation-Test	2 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	2 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	4 of 8 failed	1 of 8 failed
Serial correlation Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Random-Walk Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	1 of 8 failed	1 of 8 failed	3 of 8 failed	3 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Squeeze-Test	3 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Birthday-Spacing-Test	3 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	3 of 8 failed	1 of 8 failed	1 of 8 failed

Table 8.6: Test results for random number engine trng::mrg5.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	2 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
n-Block-Test	2 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Permutation-Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Poker-Test	1 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	0 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Random-Walk Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Squeeze-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	2 of 8 failed	2 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed

Table 8.7: Test results for random number engine trng::mrg5s.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	1 of 8 failed	2 of 8 failed	2 of 8 failed	1 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Gap-Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
n-Block-Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	1 of 8 failed
Permutation-Test	0 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Random-Walk Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Squeeze-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	4 of 8 failed	4 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Craps-Test	1 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed

Table 8.8: Test results for random number engine trng: :yarn2.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Gap-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
n-Block-Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Ising-Model-Test (energy)	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Permutation-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Random-Walk Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Serial Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	1 of 8 failed	2 of 8 failed	2 of 8 failed	1 of 8 failed
Squeeze-Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed

Table 8.9: Test results for random number engine trng: :yarn3.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	1 of 8 failed	3 of 8 failed	2 of 8 failed	2 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
n-Block-Test	0 of 8 failed	0 of 8 failed	4 of 8 failed	2 of 8 failed
Ising-Model-Test (energy)	1 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	1 of 8 failed	3 of 8 failed	2 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Random-Walk Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Squeeze-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	2 of 8 failed	3 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	2 of 8 failed	3 of 8 failed	1 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed

Table 8.10: Test results for random number engine trng: :yarn3s.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	1 of 8 failed	4 of 8 failed	1 of 8 failed	0 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
n-Block-Test	0 of 8 failed	0 of 8 failed	4 of 8 failed	1 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Poker-Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	5 of 8 failed	3 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed
Random-Walk Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Serial Test	2 of 8 failed	4 of 8 failed	0 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	1 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Squeeze-Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	2 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	1 of 8 failed

Table 8.11: Test results for random number engine trng: :yarn4.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	2 of 8 failed	3 of 8 failed	1 of 8 failed	0 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Gap-Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
n-Block-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Ising-Model-Test (energy)	1 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Permutation-Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Poker-Test	0 of 8 failed	0 of 8 failed	4 of 8 failed	1 of 8 failed
Maximum-of-t Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Random-Walk Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Squeeze-Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	2 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Binary-Rank Test (K-S)	1 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Craps-Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed

Table 8.12: Test results for random number engine trng: :yarn5.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	1 of 8 failed
n-Block-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (energy)	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	4 of 8 failed	3 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Random-Walk Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Squeeze-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed

Table 8.13: Test results for random number engine trng: :yarn5s.

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Chi-Square-Uniformity-Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Gap-Test	0 of 8 failed	0 of 8 failed	3 of 8 failed	2 of 8 failed
n-Block-Test	1 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (energy)	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	1 of 8 failed	1 of 8 failed	2 of 8 failed	2 of 8 failed
Permutation-Test	1 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Poker-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	2 of 8 failed
Serial correlation Test	1 of 8 failed	3 of 8 failed	1 of 8 failed	0 of 8 failed
Random-Walk Test	0 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	0 of 8 failed	1 of 8 failed	0 of 8 failed
Collision-Test (Hash-Test)	0 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Squeeze-Test	1 of 8 failed	1 of 8 failed	4 of 8 failed	2 of 8 failed
Birthday-Spacing-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Binary-Rank Test (K-S)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Minimum-Distance Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed

Table 8.14: Test results for random number engine boost : :mt19937 (Mersenne Twister generator).

test	confidence level			
	0.05	0.1	0.9	0.95
KS-Uniformity-Test	2 of 8 failed	2 of 8 failed	3 of 8 failed	2 of 8 failed
Chi-Square-Uniformity-Test	1 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Gap-Test	1 of 8 failed	2 of 8 failed	2 of 8 failed	0 of 8 failed
n-Block-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Ising-Model-Test (energy)	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Ising-Model-Test (specific heat)	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
CouponCollector-Test	0 of 8 failed	1 of 8 failed	2 of 8 failed	2 of 8 failed
Permutation-Test	0 of 8 failed	0 of 8 failed	0 of 8 failed	0 of 8 failed
Poker-Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Maximum-of-t Test	1 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Serial correlation Test	0 of 8 failed	0 of 8 failed	2 of 8 failed	1 of 8 failed
Random-Walk Test	1 of 8 failed	1 of 8 failed	0 of 8 failed	0 of 8 failed
Serial Test	0 of 8 failed	1 of 8 failed	1 of 8 failed	1 of 8 failed
Collision-Test (Hash-Test)	4 of 8 failed	5 of 8 failed	4 of 8 failed	3 of 8 failed
Squeeze-Test	2 of 8 failed	2 of 8 failed	1 of 8 failed	0 of 8 failed
Birthday-Spacing-Test	2 of 8 failed	2 of 8 failed	1 of 8 failed	1 of 8 failed
Binary-Rank Test (K-S)	1 of 8 failed	1 of 8 failed	2 of 8 failed	1 of 8 failed
Minimum-Distance Test	0 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed
Craps-Test	0 of 8 failed	2 of 8 failed	0 of 8 failed	0 of 8 failed

9 Frequently asked questions

What license or licenses are you using for TRNG? TRNG is free software; you can redistribute it and/or modify it under the terms of the GNU General Public License in version 2 as published by the Free Software Foundation.

Why is the library called TRNG? Who is Tina? Tina is the name of a Linux cluster at the Institute of Theoretical Physics at the University Magdeburg in Germany. TRNG was written to carry out Monte Carlo simulations on this parallel computer. The name Tina is a self-referring acronym for “Tina is no acronym”. The abbreviation TRNG stands for “Tina’s Random Number Generator Library”. But sometimes it is used in the literature for “true random number generator” as well, which is a technical device, that generates random numbers by a physical process (e. g. radioactive decay or noise in a electric circuit).

I am confused, there are so many different PRNGs in TRNG. Which one is the best? There is nothing like the best PRNG. If a generator behaves as a good source of randomness or not can depend on your Monte Carlo application, and there are trade-offs between speed and quality. In general, it is a good idea to test if the outcome of a Monte Carlo simulation is independent of the underlying PRNG. Therefore TRNG offers so many of them.

But generally speaking, YARN generators are a good choice (see section). If the PRNG is the bottleneck of your Monte Carlo simulation you might try the linear congruential generator (see section 4.1.1) or in the case of a sequential simulation a lagged Fibonacci generator with four feedback taps (see section 4.1.4).

Why is TRNG written in C++? C++ provides a lot of advanced features like inline functions and static polymorphism via templates. These language features give us the power to implement a fast, portable and easy to use library of PRNGs. Other languages (like FORTRAN or C) do not offer these (or comparable) features, are significantly slower (like Java or scripting languages), or are supported by fewer platforms (like C#).

How can I use TRNG in my FORTRAN programs? Unfortunately this is not possible. TRNG makes heavy use of special C++ language features like classes, inline functions, and templates. All these concepts have no counterpart in the FORTRAN programming language. Large parts of TRNG even do not reside in the library that you link with `-ltrng4` to your object code. Template functions and inline functions are defined exclusively in the header files.

How can I use TRNG in my C programs? Unfortunately this is not possible. Here the same statements apply like in the last question. But in fact, it is much more easy to port a C program to C++ than porting a FORTRAN program to C++. Just comply with the following recipe.

9 Frequently asked questions

- Rename header files *foo.h* of the C standard library into *cfoo* but let other header files untouched, i. e., change

```
#include <stdio.h>
#include <math.h>
#include <unistd.h>
```

into

```
#include <cstdio>
#include <cmath>
#include <unistd.h>
```

Note, *unistd.h* is not part of the C standard library.

- Insert the line

```
using namespace std;
```

after the include directives of each source file.

- Do not use C++ function names that are C++ keywords, i. e., *class*, *new*, *public* or *private*.

This recipe will give you an ugly but valid C++ program, at least in the most cases. Now you have to compile your “C” program by a C++ compiler, but it can benefit the high quality parallel PRNGs of TRNG.

Where can I report bugs or make a feature request. Send bugs reports and feature requests to heiko.bauke@physics.ox.ac.uk.

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