

Fast Normal Random Number Generators for Vector Processors*

Richard P. Brent¹
Computer Sciences Laboratory
Australian National University
Canberra, ACT 0200

Abstract

We consider pseudo-random number generators suitable for vector processors. In particular, we describe vectorised implementations of the Box-Muller and Polar methods, and show that they give good performance on the Fujitsu VP2200. We also consider some other popular methods, e.g. the Ratio method and the method of Von Neumann and Forsythe, and show why they are unlikely to be competitive with the Polar method on vector processors.

1 Introduction

Several recent papers [2, 4, 16] have considered the generation of uniformly distributed pseudo-random numbers on vector and parallel computers. In many applications, random numbers from specified non-uniform distributions are required. These distributions may be continuous (e.g. normal, exponential) or discrete (e.g. Poisson). A common requirement is for the normal distribution.

The most efficient methods for generating normally distributed random variables on sequential machines [1, 3, 6, 7, 10, 11] involve the use of different approximations on different intervals, and/or the use of “rejection” methods, so they often do not vectorise well. Simple, “old-fashioned” methods may be preferable. In Section 2 we describe two such methods, and in Sections 3–4 we consider their efficient implementation on vector processors, and give the results of implementations on a Fujitsu VP2200/10. In Sections 5–6 we consider some other methods which are popular on serial machines, and show that they are unlikely to be competitive on vector processors.

2 Some Normal Generators

Assume that a good uniform random number generator which returns uniformly distributed numbers in the interval $[0, 1)$ is available, and that we wish to sample the normal distribution with mean μ and variance σ^2 . We can generate two independent, normally distributed numbers x, y by the following old algorithm due to Box and Muller [14] (*Algorithm B1*):

1. Generate independent uniform numbers u and v .
2. Set $r \leftarrow \sigma \sqrt{-2 \ln(1 - u)}$.
3. Set $x \leftarrow r \sin(2\pi v) + \mu$ and $y \leftarrow r \cos(2\pi v) + \mu$.

Two minor points:

- a. We have written $(1 - u)$ instead of u at step 2 because of our assumption that the uniform random number generator may return exactly 0 but never exactly 1. If the uniform generator never returns exactly 0, then $(1 - u)$ can be replaced by u . Similar comments apply below.
- b. The argument of \sin and \cos in step 3 is in the interval $[-\pi, \pi)$, but any interval of length 2π would be satisfactory.

The proof that the algorithm is correct follows on considering the distribution of (x, y) transformed to polar coordinates, and is similar to the proof of correctness of the Polar method, given in [10].

Algorithm B1 is a reasonable choice if vectorised square root, logarithm and trigonometric function routines are available. Each normally distributed number requires 1 uniformly distributed number, 0.5 square roots, 0.5 logarithms, and 1 \sin or \cos evaluation. Vectorised implementations of the Box-Muller method are discussed in Section 3.

A variation of Algorithm B1 is the *Polar* method of Box, Muller and Marsaglia (1958) (*Algorithm P* of Knuth [10]):

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¹E-mail address: rpb@cs1ab.anu.edu.au
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1. Generate independent uniform random numbers x and y in the interval $[-1, 1)$.
2. Compute $s \leftarrow x^2 + y^2$.
3. If $s \geq 1$ then go to step 1 (i.e. *reject* x and y) else go to step 4.
4. Set $r \leftarrow \sigma\sqrt{-2\ln(s)/s}$, and return $rx + \mu$ and $ry + \mu$.

It is easy to see that, at step 4, (x, y) is uniformly distributed in the unit circle, so s is uniformly distributed in $[0, 1)$. To avoid the remote possibility of division by zero at step 4, we could replace $\ln(s)/s$ by $\ln(1-s)/(1-s)$.

A proof that the values returned by Algorithm P are independent, normally distributed random numbers (with mean μ and variance σ^2) is given in Knuth [10]. On average, step 1 is executed $4/\pi$ times, so each normally distributed number requires $4/\pi \simeq 1.27$ uniform random numbers, 0.5 divisions, 0.5 square roots, and 0.5 logarithms. Compared to Algorithm B1, we have avoided the sin and cos computation at the expense of more uniform random numbers, 0.5 divisions, and the cost of implementing the acceptance/rejection process. This can be done using a vector gather. Vectorised implementations of the Polar method are discussed in Section 4.

3 Implementation of the Box-Muller Method

We have implemented the Box-Muller method (Algorithm B1 above) and several refinements (B2, B3) on a Fujitsu VP 2200/10 vector processor at the Australian National University. The implementations all return double-precision real results, and in cases where approximations to sin, cos, sqrt and/or ln have been made, the absolute error is considerably less than 10^{-10} . Thus, statistical tests using less than about 10^{20} random numbers should not be able to detect any bias due to the approximations. The calling sequences allow for an array of random numbers to be returned. This permits vectorisation and amortises the cost of a subroutine call over the cost of generating many random numbers.

Our method B2 is the same as B1, except that we replace calls to the Fortran library sin and cos by an inline computation, using a fast, but sufficiently accurate, approximation. Let $y = (2\pi v - \pi)/16$, where $0 \leq v < 1$, so $|y| \leq \pi/16$. We approximate $\sin y$ by a polynomial of the form $s_1y + s_3y^3 + s_5y^5 + s_7y^7$, and

component	B1	B2	B3	P1	P2	R1
ln	13.1	13.1	7.1	13.1	7.1	0.3
sqrt	8.8	8.8	1.0	8.8	1.0	0.0
sin/cos	13.8	6.6	6.6	0.0	0.0	0.0
other	5.9	5.6	11.6	11.9	13.8	35.1
total	41.6	34.1	26.3	33.8	21.9	35.4

Table 1: Cycles per normal random number

$\cos y$ by a polynomial of the form $c_0 + c_2y^2 + c_4y^4 + c_6y^6$. Then, using the identities

$$\sin 2y = 2 \sin y \cos y, \quad \cos 2y = 1 - 2 \sin^2 y$$

four times, we can compute $\sin 16y$ and $\cos 16y$ with a small number of multiplications and additions. The computation is vectorizable.

Times, in machine cycles per normally distributed number, for methods B1, B2 (and other methods described below) are given in Table 1. In all cases the generalised Fibonacci random number generator RANU4 (described in [4]) was used to generate the required uniform random numbers, and a large number of random numbers were generated, so that vector lengths were long. RANU4 generates a uniformly distributed random number in 2.2 cycles on the VP 2200/10. (The cycle time of the VP 2200/10 at ANU is 3.2 nsec, and two multiplies and two adds can be performed per clock cycle, so the peak speed is 1.25 Gflop.)

The Table gives the total times and also the estimated times for the four main components:

1. ln computation (actually 0.5 times the cost of one ln computation since the times are per normal random number generated).
2. sqrt computation (actually 0.5 times).
3. sin or cos computation.
4. other, including uniform random number generation.

The results for method B1 show that the sin/cos and ln computations are the most expensive (65% of the total time). Method B2 is successful in reducing the sin/cos time from 33% of the total to 19%.

In Method B2, 64% of the time is consumed by the computation of $\sqrt{-\ln(1-u)}$. An obvious way to reduce this time is to use a fast approximation to the function

$$f(u) = \sqrt{-\ln(1-u)},$$

just as we used a fast approximation to sin and cos to speed up method B1. However, this is difficult to

accomplish with sufficient accuracy, because the function $f(u)$ has singularities at both endpoints of the unit interval. Method B3 overcomes this difficulty in the following way.

1. We approximate the function

$$g(u) = u^{-1/2}f(u) = \sqrt{\frac{-\ln(1-u)}{u}},$$

rather than $f(u)$. Using the Taylor series for $\ln(1-u)$, we see that $g(u) = 1 + u/4 + \dots$ is well-behaved near $u = 0$.

2. The approximation to $g(u)$ is only used in the interval $0 \leq u \leq \tau$, where $\tau < 1$ is suitably chosen. For $\tau < u < 1$ we use the slow but accurate Fortran `ln` and `sqrt` routines.
3. We make a change of variable of the form $v = (\alpha u + \beta)/(\gamma u + \delta)$, where α, \dots, δ are chosen to map $[0, \tau]$ to $[-1, 1]$, and the remaining degrees of freedom are used to move the singularities of the function $h(v) = g(u)$ as far away as possible from the region of interest (which is $-1 \leq v \leq 1$). To be more precise, let ρ be a positive parameter. Then we can choose

$$\tau = 1 - \left(\frac{\rho}{\rho + 2}\right)^2,$$

$$v = (\rho + 1) \left(\frac{(\rho + 2)u - 2}{2(\rho + 1) - (\rho + 2)u}\right),$$

and the singularities of $h(v)$ are at $\pm(\rho + 1)$.

For simplicity, we choose $\rho = 1$, which experiment shows is close to optimal on the VP 2200/10. Then $\tau = 8/9$, $v = (6u - 4)/(4 - 3u)$, and $h(v)$ has singularities at $v = \pm 2$, corresponding to the singularities of $g(u)$ at $u = 1$ and $u = \infty$. A polynomial of the form $h_0 + h_1v + \dots + h_{15}v^{15}$ can be used to approximate $h(v)$ with absolute error less than 2×10^{-11} on $[-1, 1]$. About 30 terms would be needed if we attempted to approximate $g(u)$ to the same accuracy by a polynomial on $[0, \tau]$. We use polynomial approximations which are close to minimax approximations. These may easily be obtained by truncating Chebyshev series, as described in [5].

It appears that this approach requires the computation of a square root, since we really want $f(u) = u^{1/2}g(u)$, not $g(u)$. However, a trick allows this square root computation to be avoided, at the expense of an additional uniform random number generation (which is cheap) and a few arithmetic operations. Recall that

u is a uniformly distributed random variable on $[0, 1)$. We generate *two* independent uniform variables, say u_1 and u_2 , and let $u \leftarrow \max(u_1, u_2)^2$. It is easy to see that u is in fact uniformly distributed on $[0, 1)$. However, $u^{1/2} = \max(u_1, u_2)$ can be computed without calling the library `sqrt` routine. To summarise, a non-vectorised version of method B3 is:

1. Generate uniform random numbers u_1, u_2 and u_3 on $[0, 1)$.
2. Set $m \leftarrow \max(u_1, u_2)$ and $u \leftarrow m^2$.
3. If $u > 8/9$ then
 - 3.1. set $r \leftarrow \sigma \sqrt{-\ln(1-u)}$ using Fortran library routines, else
 - 3.2. set $v \leftarrow (6u - 4)/(4 - 3u)$, evaluate $h(v)$ as described above, and set $r \leftarrow \sigma m h(v)$.
4. Evaluate $s \leftarrow \sin(2\pi u_3 - \pi)$ and $c \leftarrow \cos(2\pi u_3 - \pi)$ as described above.
5. Return $\mu + cr\sqrt{2}$ and $\mu + sr\sqrt{2}$, which are independent, normal random numbers with mean μ and standard deviation σ .

Vectorization of method B3 is straightforward, and can take advantage of the ‘‘list vector’’ technique on the VP2200. The idea is to gather those $u > 8/9$ into a contiguous array, call the vectorised library routines to compute an array of $\sqrt{-\ln(1-u)}$ values, and scatter these back. The gather and scatter operations do introduce some overhead, as can be seen from the row labelled ‘‘other’’ in the Table. Nevertheless, on the VP2200, method B3 is about 23% faster than method B2, and about 37% faster than the straightforward method B1. These ratios could be different on machines with more (or less) efficient implementations of scatter and gather.

Petersen [16] gives times for normal and uniform random number generators on a NEC SX-3. His implementation *normalen* of the Box-Muller method takes 55.5 nsec per normally distributed number, i.e. it is 2.4 times faster than our method B1, and 1.51 times faster than our method B3. The model of SX-3 used by Petersen has an effective peak speed of 2.75 Gflop, which is 2.2 times the peak speed of the VP 2200/10. Considering the relative speeds of the two machines and the fact that the SX-3 has a hardware square root function, our results are quite encouraging.

4 Implementation of the Polar Method

The times given in Table 1 for methods B1–B3 can be used to predict the best possible performance of the Polar method (Section 2). The Polar method avoids the computation of \sin and \cos , so could gain up to 6.6 cycles per normal random number over method B3. However, we would expect the gain to be less than this because of the overhead of a vector gather caused by use of a rejection method. A straightforward vectorised implementation of the Polar method, called method P1, was written to test this prediction. The results are shown in Table 1. 13.8 cycles are saved by avoiding the \sin and \cos function evaluations, but the overhead increases by 6.0 cycles, giving an overall saving of 7.8 cycles or 19%. Thus, method P1 is about the same speed as method B2, but not as fast as method B3.

Encouraged by our success in avoiding most \ln and $\sqrt{}$ computations in the Box-Muller method (see method B3), we considered if a similar idea would work for the Polar method. In fact, it does. Step 4 of the Polar method (Section 2) involves the computation of $\sqrt{-2\ln(s)/s}$, where $0 < s < 1$. The function has a singularity at $s = 0$, but we can approximate it quite well on an interval such as $[1/9, 1]$, using a method similar to that used to approximate the function $g(u)$ of Section 3.

Inspection of the proof in Knuth [10] shows that step 4 of the Polar method can be replaced by

- 4a. Set $r \leftarrow \sigma\sqrt{-2\ln(u)/s}$,
and return $rx + \mu$ and $ry + \mu$

where u is any uniformly distributed variable on $(0, 1]$, provided u is independent of $\arctan(y/x)$. In particular, we can take $u = 1 - s$. Thus, omitting the constant factor $\sigma\sqrt{2}$, we need to evaluate $\sqrt{-\ln(1-s)/s}$, but this is just $g(s)$, and we can use exactly the same approximation as in Section 3. This gives us method P2. To summarise, a non-vectorised version of method P2 is:

1. Generate independent uniform random numbers x and y in the interval $[-1, 1]$.
2. Compute $s \leftarrow x^2 + y^2$.
3. If $s \geq 1$ then go to step 1 (i.e. *reject* x and y) else go to step 4.
4. If $s > 8/9$ then
 - 4.1. set $r \leftarrow \sigma\sqrt{-\ln(1-s)/s}$ using Fortran library routines, else

4.2. set $v \leftarrow (6s - 4)/(4 - 3s)$, evaluate $h(v)$ as described in Section 3, and set $r \leftarrow \sigma h(v)$.

5. Return $xr\sqrt{2} + \mu$ and $yr\sqrt{2} + \mu$, which are independent, normal random numbers with mean μ and standard deviation σ .

To vectorise steps 1–3, we simply generate vectors of x_j and y_j values, compute $s_j = x_j^2 + y_j^2$, and compress by omitting any triple (x_j, y_j, s_j) for which $s_j \geq 1$. This means that we can not predict in advance how many normal random numbers will be generated, but this problem is easily handled by introducing a level of buffering. The vectorised version of method P2 is called RANN3B, and the user-friendly routine which performs the buffering and calls RANN3B is called RANN3.

The second-last column of Table 1 gives results for method P2 (actually for RANN3, since the buffering overhead is included). There is a saving of 11.9 cycles or 35% compared to method P1, and the method is 17% faster than the fastest version of the Box-Muller method (method B3). The cost of logarithm and square root computations is only 37% of the total, the remainder being the cost of generating uniform random numbers (about 13%) and the cost of the rejection step and other overheads (about 50%). On the VP2200/10 we can generate more than 14 million normally distributed random numbers per second (one per 70 nsec).

5 The Ratio Method

The Polar method is one of the simplest of a class of rejection methods for generating random samples from the normal (and other) distributions. Other examples are given in [1, 3, 6, 10]. It is possible to implement some of these methods in a manner similar to our implementation of method P2. For example, a popular method is the Ratio Method of Kinderman and Monahan [9] (also described in [10], and improved in [11]). In its simplest form, the Ratio Method is given by *Algorithm R*:

1. Generate independent uniform random numbers u and v in $[0, 1]$.
2. Set $x \leftarrow \sqrt{8/e}(v - \frac{1}{2})/(1 - u)$.
3. If $-x^2 \ln(1 - u) > 4$ then go to step 1 (i.e. *reject* x) else go to step 4.
4. Return $\sigma x + \mu$.

Algorithm R returns a normally distributed random number using (on average) $8/\sqrt{\pi e} \simeq 2.74$ uniform random numbers and 1.37 logarithm evaluations. The proof of correctness, and various refinements which reduce the number of logarithm evaluations, are given in [9, 10, 11]. The idea of the proof is that x is normally distributed if the point (u, v) lies inside a certain closed curve C which in turn is inside the rectangle $[0, 1] \times [-\sqrt{2/e}, +\sqrt{2/e}]$. Step 3 rejects (u, v) if it is outside C .

The function $\ln(1 - u)$ occurring at step 3 has a singularity at $u = 1$, but it can be evaluated using a polynomial or rational approximation on some interval $[0, \tau]$, where $\tau < 1$, in much the same way as the function $g(u)$ of Section 3.

The refinements added by Kinderman and Monahan [9] and Leva [11] avoid most of the logarithm evaluations. The following step is added:

- 2.5. If $P_1(u, v)$ then go to step 4
 else if $P_2(u, v)$ then go to step 1
 else go to step 3.

Here $P_1(u, v)$ and $P_2(u, v)$ are easily-computed conditions. Geometrically, P_1 corresponds to a region R_1 which lies inside C , and P_2 corresponds to a region R_2 which encloses C , but R_1 and R_2 have almost the same area. Step 3 is only executed if (u, v) lies in the borderline region $R_2 \setminus R_1$.

Step 2.5 can be vectorised, but at the expense of several vector scatter/gather operations. Thus, the saving in logarithm evaluations is partly cancelled out by an increase in overheads. The last column (R1) of Table 1 gives the times for our implementation on the VP2200. As expected, the time for the logarithm computation is now negligible, and the overheads dominate. In percentage terms the times are:

- 1% logarithm computation (using the library routine),
- 17% uniform random number computation,
- 23% scatter and gather to handle borderline region,
- 59% step 2.5 and other overheads.

Although disappointing, the result for the Ratio method is not surprising, because the computations and overheads are similar to those for method P2 (though with less logarithm computations), but only half as many normal random numbers are produced. Thus, we would expect the Ratio method to be slightly better than half as fast as method P2, and this is what Table 1 shows.

6 GRAND

On serial machines GRAND [3] is competitive with the Ratio method. In fact, GRAND is the fastest of the methods compared by Leva [11]. GRAND is based on an idea of Von Neumann and Forsythe for generating samples from a distribution with density function $c \exp(-h(x))$, where $0 \leq h(x) \leq 1$:

1. Generate a uniform random number x , and set $u_0 \leftarrow h(x)$.
2. Generate independent uniform random numbers u_1, u_2, \dots
 until the first $k > 0$ such that $u_{k-1} < u_k$.
3. If k is odd then return x ,
 else reject x and go to step 1.

A proof of correctness is given in Knuth [10].

It is hard to see how to implement GRAND efficiently on a vector processor. There are two problems –

1. k is not bounded, even though its expected value is small. Thus, a sequence of gather operations seems to be required. The result would be similar to Petersen's implementation [16] of a generator for the Poisson distribution (much slower than his implementation for the normal distribution).
2. Because of the restriction $0 \leq h(x) \leq 1$, the area under the normal curve $\exp(-x^2/2)/\sqrt{2\pi}$ has to be split into different regions from which samples are drawn with probabilities proportional to their areas. This complicates the implementation of the rejection step.

For these reasons we would expect a vectorised implementation of GRAND to be even slower than our implementation of the Ratio method. Similar comments apply to other rejection methods which use an iterative rejection process and/or several different regions.

7 Conclusion

We have shown that both the Box-Muller and Polar methods vectorise well, and that it is possible to avoid and/or speed up the evaluation of the functions (sin, cos, ln, sqrt) which appear necessary. On the VP2200/10 our best implementation of the Polar method takes 21.9 machine cycles per normal random number, slightly faster than our best implementation of the Box-Muller method (26.3 cycles).

We also considered the vectorisation of some other popular methods for generating normally distributed random numbers, such as the Ratio method and the method of Von Neumann and Forsythe, and showed why such methods are unlikely to be faster than the Polar method on a vector processor.

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