

More Random Sampling Methods

This version of the document is dated 2021-04-11.

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1.1 Specific Distributions

Requires random real numbers. This section shows algorithms to sample several popular non-uniform distributions. The algorithms are exact unless otherwise noted, and applications should choose algorithms with either no error (including rounding error) or a user-settable error bound. See the **appendix** for more information.

1.1.1 Normal (Gaussian) Distribution

The [normal distribution](#) (also called the Gaussian distribution) takes the following two parameters:

- μ (μ) is the mean (average), or where the peak of the distribution's "bell curve" is.
- σ (σ), the standard deviation, affects how wide the "bell curve" appears. The probability that a number sampled from the normal distribution will be within one standard deviation from the mean is about 68.3%; within two standard deviations (2 times σ), about 95.4%; and within three standard deviations, about 99.7%. (Some publications give σ^2 , or variance, rather than standard deviation, as the second parameter. In this case, the standard deviation is the variance's square root.)

There are a number of methods for sampling the normal distribution. An application can combine some or all of these.

1. The ratio-of-uniforms method (given as `NormalRatioOfUniforms` below).
2. In the *Box-Muller transformation*, $\mu + \text{radius} * \cos(\text{angle})$ and $\mu + \text{radius} * \sin(\text{angle})$, where $\text{angle} = \text{RNDRANGEMaxExc}(0, 2 * \pi)$ and $\text{radius} = \sqrt{\text{Expo}(0.5)}$ * σ , are two independent values sampled from the normal distribution. The polar method (given as `NormalPolar` below) likewise produces two independent values

sampled from that distribution at a time.

3. Karney's algorithm to sample from the normal distribution, in a manner that minimizes approximation error and without using floating-point numbers (Karney 2014)⁽¹⁾.

For surveys of Gaussian samplers, see (Thomas et al. 2007)⁽²⁾, and (Malik and Hemani 2016)⁽³⁾.

```
METHOD NormalRatioOfUniforms(mu, sigma)
  while true
    a=RNDRANGEMinExc(0,1)
    b=RNDRANGE(0,sqrt(2.0/exp(1.0)))
    if b*b <= -a * a * 4 * ln(a)
      return (RNDINT(1) * 2 - 1) *
        (b * sigma / a) + mu
    end
  end
end
END METHOD

METHOD NormalPolar(mu, sigma)
  while true
    a = RNDRANGEMinExc(0,1)
    b = RNDRANGEMinExc(0,1)
    if RNDINT(1) == 0: a = 0 - a
    if RNDINT(1) == 0: b = 0 - b
    c = a * a + b * b
    if c != 0 and c <= 1
      c = sqrt(-ln(c) * 2 / c)
      return [a * sigma * c + mu, b * sigma * c + mu]
    end
  end
end
END METHOD
```

Notes:

1. The *standard normal distribution* is implemented as `Normal(0, 1)`.
2. Methods implementing a variant of the normal distribution, the *discrete Gaussian distribution*, generate *integers* that closely follow the normal distribution. Examples include the one in (Karney 2014)⁽¹⁾, an improved version in (Du et al. 2020)⁽⁴⁾, as well as so-called "constant-time" methods such as (Micciancio and Walter 2017)⁽⁵⁾ that are used above all in *lattice-based cryptography*.
3. The following are some approximations to the normal distribution that papers have suggested:
 - The sum of twelve `RNDRANGEMaxExc(0, sigma)` numbers, subtracted by $6 * \text{sigma}$. (Kabal 2000/2019)⁽⁶⁾ "warps" this sum in the following way (before adding the mean μ) to approximate the normal distribution better: $\text{ssq} = \text{sum} * \text{sum}$; $\text{sum} = (((0.0000001141 * \text{ssq} - 0.0000005102) * \text{ssq} + 0.00007474) * \text{ssq} + 0.0039439) * \text{ssq} + 0.98746) * \text{sum}$. See also "[Irwin-Hall distribution](#)", namely the sum of n many `RNDRANGE(0, 1)` numbers, on Wikipedia. D. Thomas (2014)⁽⁷⁾, describes a more general approximation called CLT_k , which combines k numbers in $[0, 1]$ sampled from the uniform distribution as follows: `RNDRANGE(0, 1) - RNDRANGE(0, 1) + RNDRANGE(0, 1) -`
 - Numerical **inversions** of the normal distribution's cumulative distribution function (CDF), including those by Wichura, by Acklam, and by Luu (Luu 2016)⁽⁸⁾. See also "[A literate program to compute](#)

[the inverse of the normal CDF](#)". Notice that the normal distribution's inverse CDF has no closed form.

4. A pair of *q-Gaussian* random variates with parameter q less than 3 can be generated using the Box-Muller transformation, except radius is $\text{radius} = \sqrt{-2 * (\text{pow}(u, 1 - qp) - 1) / (1 - qp)}$ (where $qp = (1 + q) / (3 - q)$ and $u = \text{RNDRANGE}(0, 1)$), and the two variates are not independent of each other (Thistleton et al. 2007)⁽⁹⁾.

1.1.2 Gamma Distribution

The following method samples a number from a *gamma distribution* and is based on Marsaglia and Tsang's method from 2000⁽¹⁰⁾ and (Liu et al. 2015)⁽¹¹⁾. Usually, the number expresses either—

- the lifetime (in days, hours, or other fixed units) of a random component with an average lifetime of `meanLifetime`, or
- a random amount of time (in days, hours, or other fixed units) that passes until as many events as `meanLifetime` happen.

Here, `meanLifetime` must be an integer or noninteger greater than 0, and `scale` is a scaling parameter that is greater than 0, but usually 1 (the random gamma number is multiplied by `scale`).

```
METHOD GammaDist(meanLifetime, scale)
  // Needs to be greater than 0
  if meanLifetime <= 0 or scale <= 0: return error
  // Exponential distribution special case if
  // `meanLifetime` is 1 (see also (Devroye 1986), p. 405)
  if meanLifetime == 1: return Expo(1.0 / scale)
  if meanLifetime < 0.3 // Liu, Martin, Syring 2015
    lamda = (1.0/meanLifetime) - 1
    w = meanLifetime / (1-meanLifetime) * exp(1)
    r = 1.0/(1+w)
    while true
      z = 0
      x = RNDRANGE(0, 1)
      if x <= r: z = -ln(x/r)
      else: z = -Expo(lamda)
      ret = exp(-z/meanLifetime)
      eta = 0
      if z>=0: eta=exp(-z)
      else: eta=w*lamda*exp(lamda*z)
      if RNDRANGE(0, eta) < exp(-ret-z): return ret * scale
    end
  end
  d = meanLifetime
  v = 0
  if meanLifetime < 1: d = d + 1
  d = d - (1.0 / 3) // NOTE: 1.0 / 3 must be a fractional number
  c = 1.0 / sqrt(9 * d)
  while true
    x = 0
    while true
      x = Normal(0, 1)
      v = c * x + 1;
      v = v * v * v
      if v > 0: break
    end
    u = RNDRANGEMinExc(0,1)
```

```

        x2 = x * x
        if u < 1 - (0.0331 * x2 * x2): break
        if ln(u) < (0.5 * x2) + (d * (1 - v + ln(v))): break
    end
    ret = d * v
    if meanLifetime < 1
        ret = ret * pow(RNDRANGE(0, 1), 1.0 / meanLifetime)
    end
    return ret * scale
END METHOD

```

Note: The following is a useful identity for the gamma distribution: $\text{GammaDist}(a) = \text{BetaDist}(a, b - a) * \text{GammaDist}(b)$ (Stuart 1962)⁽¹²⁾.

1.1.3 Beta Distribution

The beta distribution is a bounded-domain probability distribution; its two parameters, a and b , are both greater than 0 and describe the distribution's shape. Depending on a and b , the shape can be a smooth peak or a smooth valley.

The following method samples a number from a *beta distribution*, in the interval $[0, 1]$.

```

METHOD BetaDist(a, b)
    if b==1 and a==1: return RNDRANGE(0, 1)
    // Min-of-uniform
    if a==1: return 1.0-pow(RNDRANGE(0, 1),1.0/b)
    // Max-of-uniform. Use only if a is small to
    // avoid accuracy problems, as pointed out
    // by Devroye 1986, p. 675.
    if b==1 and a < 10: return pow(RNDRANGE(0, 1),1.0/a)
    x=GammaDist(a,1)
    return x/(x+GammaDist(b,1))
END METHOD

```

I give an [error-bounded sampler](#) for the beta distribution (when a and b are both 1 or greater) in a separate page.

1.1.4 von Mises Distribution

The *von Mises distribution* describes a distribution of circular angles and uses two parameters: mean is the mean angle and kappa is a shape parameter. The distribution is uniform at $\text{kappa} = 0$ and approaches a normal distribution with increasing kappa .

The algorithm below samples a number from the von Mises distribution, and is based on the Best-Fisher algorithm from 1979 (as described in (Devroye 1986)⁽¹³⁾ with errata incorporated).

```

METHOD VonMises(mean, kappa)
    if kappa < 0: return error
    if kappa == 0
        return RNDRANGEMinMaxExc(mean-pi, mean+pi)
    end
    r = 1.0 + sqrt(4 * kappa * kappa + 1)
    rho = (r - sqrt(2 * r)) / (kappa * 2)
    s = (1 + rho * rho) / (2 * rho)
    while true
        u = RNDRANGEMaxExc(-pi, pi)
        v = RNDRANGEMinMaxExc(0, 1)
        z = cos(u)

```

```

w = (1 + s*z) / (s + z)
y = kappa * (s - w)
if y*(2 - y) - v >= 0 or ln(y / v) + 1 - y >= 0
  if angle < -1: angle = -1
  if angle > 1: angle = 1
  // NOTE: Inverse cosine replaced here
  // with `atan2` equivalent
  angle = atan2(sqrt(1-w*w),w)
  if u < 0: angle = -angle
  return mean + angle
end
end
END METHOD

```

1.1.5 Stable Distribution

As more and more numbers, sampled independently at random in the same way, are added together, their distribution tends to a [stable distribution](#), which resembles a curve with a single peak, but with generally "fatter" tails than the normal distribution. (Here, the stable distribution means the "alpha-stable distribution".) The pseudocode below uses the Chambers-Mallows-Stuck algorithm. The `Stable` method, implemented below, takes two parameters:

- `alpha` is a stability index in the interval (0, 2].
- `beta` is an asymmetry parameter in the interval [-1, 1]; if `beta` is 0, the curve is symmetric.

```

METHOD Stable(alpha, beta)
  if alpha <= 0 or alpha > 2: return error
  if beta < -1 or beta > 1: return error
  halfpi = pi * 0.5
  unif=RNDRANGEMinMaxExc(-halfpi, halfpi)
  c=cos(unif)
  if alpha == 1
    s=sin(unif)
    if beta == 0: return s/c
    expo=Expo(1)
    return 2.0*((unif*beta+halfpi)*s/c -
      beta * ln(halfpi*expo*c/(unif*beta+halfpi)))/pi
  else
    z=-tan(alpha*halfpi)*beta
    ug=unif+atan2(-z, 1)/alpha
    cpow=pow(c, -1.0 / alpha)
    return pow(1.0+z*z, 1.0 / (2*alpha))*
      (sin(alpha*ug)*cpow)*
      pow(cos(unif-alpha*ug)/expo, (1.0 - alpha) / alpha)
  end
END METHOD

```

Methods implementing the strictly geometric stable and general geometric stable distributions are shown below (Kozubowski 2000)⁽¹⁴⁾. Here, `alpha` is in (0, 2], `lamda` is greater than 0, and `tau`'s absolute value is not more than $\min(1, 2/\alpha - 1)$. The result of `GeometricStable` is a symmetric Linnik distribution if `tau` = 0, or a Mittag-Leffler distribution if `tau` = 1 and `alpha` < 1.

```

METHOD GeometricStable(alpha, lamda, tau)
  rho = alpha*(1-tau)/2
  sign = -1

```

```

if tau==1 or RNDINT(1)==0 or RNDRange(0, 1) < tau
    rho = alpha*(1+tau)/2
    sign = 1
end
w = 1
if rho != 1
    rho = rho * pi
    cotparam = RNDRange(0, rho)
    w = sin(rho)*cos(cotparam)/sin(cotparam)-cos(rho)
end
return Expo(1) * sign * pow(lamda*w, 1.0/alpha)
END METHOD

METHOD GeneralGeoStable(alpha, beta, mu, sigma)
    z = Expo(1)
    if alpha == 1: return mu*z+Stable(alpha, beta)*sigma*z+
        sigma*z*beta*2*pi*ln(sigma*z)
    else: return mu*z+
        Stable(alpha, beta)*sigma*pow(z, 1.0/alpha)
END METHOD

```

1.1.6 Multivariate Normal (Multinormal) Distribution

The following pseudocode calculates a random vector (list of numbers) that follows a [***multivariate normal \(multinormal\) distribution***](#). The method MultivariateNormal takes the following parameters:

- A list, mu (μ), which indicates the means to add to the random vector's components. mu can be nothing, in which case each component will have a mean of zero.
- A list of lists cov, that specifies a *covariance matrix* (Σ , a symmetric positive definite $N \times N$ matrix, where N is the number of components of the random vector).

```

METHOD Decompose(matrix)
    numrows = size(matrix)
    if size(matrix[0])!=numrows: return error
    // Does a Cholesky decomposition of a matrix
    // assuming it's positive definite and invertible
    ret=NewList()
    for i in 0...numrows
        submat = NewList()
        for j in 0...numrows: AddItem(submat, 0)
        AddItem(ret, submat)
    end
    s1 = sqrt(matrix[0][0])
    if s1==0: return ret // For robustness
    for i in 0...numrows
        ret[0][i]=matrix[0][i]*1.0/s1
    end
    for i in 0...numrows
        msum=0.0
        for j in 0...i: msum = msum + ret[j][i]*ret[j][i]
        sq=matrix[i][i]-msum
        if sq<0: sq=0 // For robustness
        ret[i][i]=math.sqrt(sq)
    end
    for j in 0...numrows
        for i in (j + 1)...numrows
            // For robustness
            if ret[j][j]==0: ret[j][i]=0

```

```

        if ret[j][j]!=0
            msum=0
            for k in 0...j: msum = msum + ret[k][i]*ret[k][j]
            ret[j][i]=(matrix[j][i]-msum)*1.0/ret[j][j]
        end
    end
end
return ret
END METHOD

```

```

METHOD MultivariateNormal(mu, cov)
    mulen=size(cov)
    if mu != nothing
        mulen = size(mu)
        if mulen!=size(cov): return error
        if mulen!=size(cov[0]): return error
    end
    // NOTE: If multiple random points will
    // be generated using the same covariance
    // matrix, an implementation can consider
    // precalculating the decomposed matrix
    // in advance rather than calculating it here.
    cho=Decompose(cov)
    i=0
    ret=NewList()
    vars=NewList()
    for j in 0...mulen: AddItem(vars, Normal(0, 1))
    while i<mulen
        nv=Normal(0,1)
        msum = 0
        if mu == nothing: msum=mu[i]
        for j in 0...mulen: msum=msum+vars[j]*cho[j][i]
        AddItem(ret, msum)
        i=i+1
    end
    return ret
end

```

Note: The [Python sample code](#) contains a variant of this method for generating multiple random vectors in one call.

Examples:

1. A vector that follows a **binormal distribution** (two-variable multinormal distribution) is a vector of two numbers from the normal distribution, and can be sampled using the following idiom: `MultivariateNormal([mu1, mu2], [[s1*s1, s1*s2*rho], [rho*s1*s2, s2*s2]])`, where `mu1` and `mu2` are the means of the vector's two components, `s1` and `s2` are their standard deviations, and `rho` is a *correlation coefficient* greater than -1 and less than 1 (0 means no correlation).
2. **Log-multinormal distribution:** Generate a multinormal random vector, then apply `exp(n)` to each component `n`.
3. A **Beckmann distribution**: Generate a random binormal vector `vec`, then apply `Norm(vec)` to that vector.
4. A **Rice (Rician) distribution** is a Beckmann distribution in which the binormal random pair is generated with `m1 = m2 = a / sqrt(2)`, `rho = 0`, and `s1 = s2 = b`, where `a` and `b` are the parameters to the Rice distribution.
5. **Rice-Norton distribution:** Generate `vec = MultivariateNormal([v,v,v], [[w,0,0],[0,w,0],[0,0,w]])` (where `v = a/sqrt(m*2)`, `w = b*b/m`, and `a`, `b`, and `m` are the parameters to the Rice-Norton distribution), then apply `Norm(vec)` to

that vector.

6. A **standard complex normal distribution** is a binormal distribution in which the binormal random pair is generated with $s_1 = s_2 = \sqrt{0.5}$ and $\mu_1 = \mu_2 = 0$ and treated as the real and imaginary parts of a complex number.
7. **Multivariate Linnik distribution:** Generate a multinormal random vector, then multiply each component by $\text{GeometricStable}(\alpha/2.0, 1, 1)$, where α is a parameter in $(0, 2]$ (Kozubowski 2000)⁽¹⁴⁾.

1.1.7 Gaussian and Other Copulas

A *copula* is a way to describe the dependence between randomly sampled numbers.

One example is a *Gaussian copula*; this copula is sampled by sampling from a **multinormal distribution**, then converting the resulting numbers to *dependent* uniform random values. In the following pseudocode, which implements a Gaussian copula:

- The parameter *covar* is the covariance matrix for the multinormal distribution.
- $\text{erf}(v)$ is the **error function** of the number v (see the appendix).

```
METHOD GaussianCopula(covar)
  mvn=MultivariateNormal(nothing, covar)
  for i in 0...size(covar)
    // Apply the normal distribution's CDF
    // to get uniform numbers
    mvn[i] = (erf(mvn[i]/(sqrt(2)*sqrt(covar[i][i])))+1)*0.5
  end
  return mvn
END METHOD
```

Each of the resulting uniform random values will be in the interval $[0, 1]$, and each one can be further transformed to any other probability distribution (which is called a *marginal distribution* or *marginal* here) by taking the quantile of that uniform number for that distribution (see "**Inverse Transform Sampling**", and see also (Cario and Nelson 1997)⁽¹⁵⁾.)

Note: The Gaussian copula is also known as the *normal-to-anything* method.

Examples:

1. To generate two correlated uniform random values with a Gaussian copula, generate $\text{GaussianCopula}([[1, \rho], [\rho, 1]])$, where ρ is the Pearson correlation coefficient, in the interval $[-1, 1]$. (Other correlation coefficients besides ρ exist. For example, for a two-variable Gaussian copula, the **Spearman correlation coefficient** srho can be converted to ρ by $\rho = \sin(\text{srho} * \pi / 6) * 2$. Other correlation coefficients, and other measures of dependence between randomly sampled numbers, are not further discussed in this document.)
2. The following example generates a two-dimensional random vector that follows a Gaussian copula with exponential marginals (ρ is the Pearson correlation coefficient, and rate1 and rate2 are the rates of the two exponential marginals).

```
METHOD CorrelatedExpo(rho, rate1, rate2)
```



```

        copula = GaussianCopula([[1, rho], [rho, 1]])
        // Transform to exponentials using that
        // distribution's quantile function
        return [-log1p(-copula[0]) / rate1,
                -log1p(-copula[1]) / rate2]
    END METHOD

```

3. The **T-Poisson hierarchy** (Knudson et al. 2021)⁽¹⁶⁾ is a way to generate N-dimensional Poisson-distributed random vectors via copulas. Each of the N dimensions is associated with a parameter `lamda` and a marginal that must be a *continuous non-negative* probability distribution (one that takes on any of an uncountable number of non-negative values, such as any number 0 or greater). To sample from the T-Poisson hierarchy—

1. sample an N-dimensional random vector via a copula (such as `GaussianCopula`), producing an N-dimensional vector of correlated uniform numbers; then
2. for each component in the vector, take that component's quantile for the corresponding marginal; then
3. for each component in the vector, replace it with `Poisson(lamda * c)`, where `c` is that component and `lamda` is the `lamda` parameter for the corresponding dimension.

The following example implements the T-Poisson hierarchy using a Gaussian copula and exponential marginals.

```

METHOD PoissonH(rho, rate1, rate2, lambda1, lambda2)
    vec = CorrelatedExpo(rho, rate1, rate2)
    return [Poisson(lambda1*vec[0]),Poisson(lambda2*vec[1])]
END METHOD

```

Other kinds of copulas describe different kinds of dependence between randomly sampled numbers. Examples of other copulas are—

- the **Fréchet-Hoeffding upper bound copula** $[x, x, \dots, x]$ (e.g., $[x, x]$), where $x = \text{RNDRange}(0, 1)$,
- the **Fréchet-Hoeffding lower bound copula** $[x, 1.0 - x]$ where $x = \text{RNDRange}(0, 1)$,
- the **product copula**, where each number is a separately generated `RNDRange(0, 1)` (indicating no dependence between the numbers), and
- the **Archimedean copulas**, described by M. Hofert and M. Mächler (2011)⁽¹⁷⁾.

2 Notes

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3 Appendix

3.1 Implementation of erf

The pseudocode below shows an approximate implementation of the [error function](#) erf, in case the programming language used doesn't include a built-in version of erf (such as JavaScript at the time of this writing). In the pseudocode, EPSILON is a very small number to end the iterative calculation.

```
METHOD erf(v)
  if v==0: return 0
  if v<0: return -erf(-v)
  if v==infinity: return 1
  // NOTE: For Java `double`, the following
  // line can be added:
  // if v>=6: return 1
  i=1
  ret=0
  zp=-(v*v)
  zval=1.0
  den=1.0
  while i < 100
    r=v*zval/den
    den=den+2
    ret=ret+r
    // NOTE: EPSILON can be pow(10,14),
    // for example.
    if abs(r)<EPSILON: break
    if i==1: zval=zp
    else: zval = zval*zp/i
    i = i + 1
```

```

end
return ret*2/sqrt(pi)
END METHOD

```

3.2 Exact, Error-Bounded, and Approximate Algorithms

There are three kinds of randomization algorithms:

1. An *exact algorithm* is an algorithm that samples from the exact distribution requested, assuming that computers—
 - can store and operate on real numbers (which have unlimited precision), and
 - can generate independent uniform random real numbers

(Devroye 1986, p. 1-2)⁽¹³⁾. However, an exact algorithm implemented on real-life computers can incur error due to the use of fixed precision, such as rounding and cancellations, especially when floating-point numbers are involved. An exact algorithm can achieve a guaranteed bound on accuracy (and thus be an *error-bounded algorithm*) using either arbitrary-precision or interval arithmetic (see also Devroye 1986, p. 2)⁽¹³⁾. All methods given on this page are exact unless otherwise noted. Note that the RNDRANGE method is exact in theory, but has no required implementation.

2. An *error-bounded algorithm* is a sampling algorithm with the following requirements:
 - If the ideal distribution is discrete (takes on a countable number of values), the algorithm samples exactly from that distribution.
 - If the ideal distribution is continuous, the algorithm samples from a distribution that is close to the ideal within a user-specified error tolerance (see below for details). The algorithm can instead sample a number from the distribution only partially, as long as the fully sampled number can be made close to the ideal within any error tolerance desired.
 - In sampling from a distribution, the algorithm incurs no approximation error not already present in the inputs (except errors needed to round the final result to the user-specified error tolerance).

Many error-bounded algorithms use random bits as their only source of randomness. An application should use error-bounded algorithms whenever possible.

3. An *inexact, approximate, or biased algorithm* is neither exact nor error-bounded; it uses "a mathematical approximation of sorts" to sample from a distribution that is close to the desired distribution (Devroye 1986, p. 2)⁽¹³⁾. An application should use this kind of algorithm only if it's willing to trade accuracy for speed.

Most algorithms on this page, though, are not *error-bounded*, but even so, they may still be useful to an application willing to trade accuracy for speed.

There are many ways to describe closeness between two distributions. One suggestion by Devroye and Gravel (2020)⁽¹⁸⁾ is Wasserstein distance (or "earth-mover distance"). Here, an algorithm has accuracy ϵ (the user-specified error tolerance) if it samples from a distribution that is close to the ideal distribution by a Wasserstein distance of not more than ϵ .

Examples:

1. Sampling from the exponential distribution via $-\ln(\text{RNDRANGE}(0, 1))$ is an *exact algorithm* (in theory), but not an *error-bounded* one for common floating-point number formats. The same is true of the Box-Muller transformation.
2. Sampling from the exponential distribution using the ExpoExact method from the section "[Exponential Distribution](#)" is an *error-bounded algorithm*. Karney's algorithm for the normal distribution (Karney 2014)⁽¹⁾ is also error-bounded because it returns a result that can be made to come close to the normal distribution within any error tolerance desired simply by appending more random digits to the end. See also (Oberhoff 2018)⁽¹⁹⁾.
3. Examples of *approximate algorithms* include sampling from a Gaussian-like distribution via a sum of $\text{RNDRANGE}(0, 1)$, or most cases of modulo reduction to produce uniform-like integers at random (see notes in the section "[RNDINT](#)").

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