

A Compact and Accurate Gaussian Variate Generator

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Abstract—A compact, fast, and accurate realization of a digital Gaussian variate generator (GVG) based on the Box–Muller algorithm is presented. The proposed GVG has a faster Gaussian sample generation rate and higher tail accuracy with a lower hardware cost than published designs. The GVG design can be readily configured to achieve arbitrary tail accuracy (i.e., with a proposed 16-bit datapath up to ± 15 times the standard deviation σ) with only small variations in hardware utilization, and without degrading the output sample rate. Polynomial curve fitting is utilized along with a hybrid (i.e., combination of logarithmic and uniform) segmentation and a scaling scheme to maintain accuracy. A typical instantiation of the proposed GVG occupies only 534 configurable slices, two on-chip block memories, and three dedicated multipliers of the Xilinx Virtex-II XC2V4000-6 field-programmable gate array (FPGA) and operates at 248 MHz, generating 496 million Gaussian variates (GVs) per second within a range of $\pm 6.66\sigma$. To accurately achieve a range of $\pm 9.4\sigma$, the GVG uses 852 configurable slices, three block memories, and three on-chip dedicated multipliers of the same FPGA while still operating at 248 MHz, generating 496 million GV's per second. The core area and performance of a GVG implemented in a 90-nm CMOS technology are also given. The statistical characteristics of the GVG are evaluated and confirmed using multiple standard statistical goodness-of-fit tests.

Index Terms—Box–Muller (BM) algorithm, field-programmable gate array (FPGA), Gaussian noise generator (GNG), low bit-error rate simulation, random number generation.

I. MOTIVATION

SEQUENCES of random variates with a Gaussian probability density function (PDF) are frequently required to model noisy natural phenomena. For example, in communications it is standard practice to use additive white Gaussian noise channels (AWGNs) and multipath Rayleigh fading channels. One practical application of Gaussian variates (GVs) is to evaluate the performance of communication systems. A communication system can be characterized through the bit-error rate (BER) versus signal-to-noise ratio (SNR) relationship, which for high SNR regions (and hence very low BERs) requires long-running Monte Carlo (MC) simulations. Although techniques such as importance sampling have been used to reduce the simulation time, these schemes are not general enough to be applicable to many problems [1]. For example, consider a digital communication system that is designed to achieve a BER of no more than 10^{-14} . This means that on average, 10^{14} symbols must be processed for

each erroneous symbol in a MC simulation of the system. One usually requires at least 100 such error events if the BER is to be estimated with reasonable statistical significance. In addition, approximately 10 samples per symbol interval are typically required to represent waveforms in the simulation. Thus, roughly 10^{17} samples must be processed. The generation of 10^9 GV's, using an optimized software simulator written in C, takes about 2.5 h on a dual core Pentium processor running at 3.0 GHz with a 1-MB L2 cache. By extrapolation, the generation of 10^{17} GV's would take about 27 000 years. Note that some recent standards give maximum allowable BERs of only 10^{-12} for specified SNRs (e.g., IEEE 802.3 10 Gbit/s Ethernet). Thus, maximizing the achievable GV generation rate is crucial for validating the BER performance of upcoming systems, and software-based GV generation is no longer adequate.

On the other hand, evaluating a communication system at a BER of 10^{-14} implies that approximately one in every 10^{14} received bits should be errored. In a typical binary phase-shift keying (BPSK) communication system, the inverse of the Gaussian error function for 10^{-14} [2] is a value that approaches 7.0. This means that random variates near the center of the distribution do not contribute significantly to the probability of error since the small corresponding signal deviations are readily tolerated by any system with that low a BER. Rather, it is the GV's with a value of 7σ or larger, where σ is the standard deviation of the Gaussian distribution, that will be the dominant source of errors. Therefore, for an MC simulation of a low BER system, the PDF of generated random numbers must be especially close to the Gaussian PDF at the high σ regions (the tails of the PDF).

Hardware-based Gaussian variate generators (GVGs) using analog devices [3]–[5] and digital components [6]–[20] have shown significant speedups compared to software implementations. However, digital implementations tend to be more desirable due to their predictable and controllable behavior, and because they can reproduce exactly the same pseudorandom sequence of variates in successive runs. Field-programmable gate arrays (FPGAs) are increasingly used as a prototyping platform for digital designs. In addition to their low cost compared to custom integrated circuits, the user-programmability of FPGAs facilitates debugging and design characterization, which can reduce the total design time significantly.

The fast and accurate GVG that we propose below utilizes a relatively small proportion of a typical contemporary FPGA. Various objectives have been considered when designing the GVG, as shown in the following.

- **Tail Accuracy:** The normal distribution is an open-ended distribution in which values of increasing magnitude occur with increasingly small probabilities. A GVG for low BER characterization must be able to generate GV's accurately especially at the high σ values. The proposed design uses

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polynomial curve fitting [21] and a hybrid (i.e., logarithmic and linear) domain segmentation and scaling scheme. This approach can efficiently achieve accurate Gaussian statistics deep into the distribution tails.

- *Hardware Efficiency*: Ideally, a hardware realization should minimize the number of required FPGA resources while achieving an acceptably high variate generation rate. Compactness is an important factor when the available hardware resources are especially limited, such as in a low-density FPGA.
- *Flexibility*: The design should be parameterizable to generate GVs within multiple standard deviations of the mean (right into the tails of the distribution). It is also important that the GVG be scalable to increase the output rate with multiple instantiations of the GVG.
- *Portability*: The design should preferably avoid exploiting device-specific features of FPGAs. The resulting more portable register transfer level design should be synthesizable for different hardware platforms and semicustom silicon technologies.
- *Statistical Accuracy*: The quality of the output variates should be supported by various standard goodness-of-fit statistical tests, such as statistical independence and small deviation from the ideal Gaussian PDF.

The key contribution of this paper is a compact and accurate synthesizable GVG core that can be configured to generate 16-bit GVs in two's complement fixed-point format for arbitrary maximum deviations into the tails of the distribution. The rest of this paper is organized as follows. Section II reviews several typical algorithms for generating GVs and compares related work on digital GVG implementations. Section III evaluates different pseudorandom number generators (PNGs) used in recently disclosed GVG designs and compares their statistical properties. Section IV describes tradeoffs involved in implementations of the Box-Muller (BM) algorithm. Section V presents the realization of the proposed scheme and discusses implementation results. Section VI is dedicated to the analysis of goodness-of-fit tests and simulation results. Concluding remarks appear in Section VII.

II. GVG ALGORITHMS AND RELATED WORK

Among the many proposed algorithms [2], [6], [9], [22]–[28] for generating GVs, the digital approaches [7], [13] are typically based on transformations of uniform distributions [29]. These algorithms include various rejection-acceptance methods [24], [27], [30]–[32], the BM method [22], and the inversion method [28].

The rejection-acceptance methods, including the polar method [31] and the Ziggurat algorithm [27], have been generally used in software implementations (e.g., the random number generator libraries in MATLAB). Due to conditional *if-then-else* assignment instructions, the output rate is not constant and hence such methods have received less attention as the basis for efficient hardware implementation. Implementation of the polar method in [11] uses a first-in first-out (FIFO) buffer to provide GVs at a constant output rate. A hardware-based implementation of a GVG based on the Ziggurat method is described in [16].

The BM algorithm [22] has been widely used to transform pairs of uniformly distributed numbers into samples from a 2-D bivariate normal distribution. The inputs to the BM algorithm are two independent uniformly-distributed pseudorandom numbers (PNs), $u_1, u_2 \in (0, 1)$. The outputs are two independent samples n_1 and n_2 , from a zero-mean, unit-variance Gaussian distribution $\mathcal{N}(0, 1)$. The transformation involves multiplying

$$f(u_1) = \sqrt{-2 \ln(u_1)}$$

by $g_1(u_2) = \sin(2\pi u_2)$ and $g_2(u_2) = \cos(2\pi u_2)$ to yield n_1 and n_2 , respectively. We will use $g(\cdot)$ to denote the implemented sinusoidal function, $g_1(\cdot)$ or $g_2(\cdot)$. Digital implementations of high-quality PNGs and accurate realizations of the required logarithm, square root, and trigonometric functions have been investigated extensively over the last three decades [33]. Consequently, the BM transformation has been used in many FPGA [8], [10], [12], [14]–[20] and parallel processor GVGs [34].

The inversion method is a standard approach for generating random variates with arbitrary probability density distributions [28]. To generate GVs, it transforms uniform random variables $u \in [0, 1]$ into Gaussian variates n by approximating the non-decreasing inverse of the Gaussian cumulative distribution function (CDF) as $n = F^{-1}(u)$. Since there is no closed-form approximation for $F^{-1}(\cdot)$, the GVG in [35] uses a lookup table to store the CDF inverse. This method requires a large memory to generate accurate GVGs at the tails of the Gaussian distribution. The GVG in [36] uses linear interpolation to approximate the inverse of the Gaussian CDF while requiring less memory. A more efficient approach is proposed in [20] that uses nonuniform segmentation of $F^{-1}(\cdot)$ to more accurately approximate the Gaussian distribution at the tails.

Another scheme is based on the Wallace method [26], where new GVs are generated by applying linear transformations to a pool of Gaussian samples. Due to the inherent feedback in this method, unwanted correlations can occur between successive transformations [15]. Using proper parameter selection to minimize the correlation effects, a hardware-based GVG-based is proposed in [15].

To choose an appropriate algorithm for digital hardware implementation, compromises must be made between the simplicity of the algorithm, the maximum error of the approximations, the robustness with respect to the distribution parameters, and the efficiency (minimum resource requirements, maximum output rate, and latency). For example, the output rate of the inversion-based scheme is half that of the BM scheme; however, it can exploit the symmetry of $F^{-1}(\cdot)$ to implement a more compact GVG without the need to implement trigonometric functions. On the other hand, trigonometric functions, such as sine and cosine, can be implemented relatively accurately in a small fraction of an FPGA. One may choose the BM algorithm and implement $f(u_1)$ efficiently, along with trigonometric functions, to increase the output sample rate. A more detailed comparison of various GVG algorithms can be found in [29] and [37].

Characteristics of different hardware-based GVGs are given in Table I. Table II summarizes the major characteristics of various disclosed GVG implementations based on the BM method.

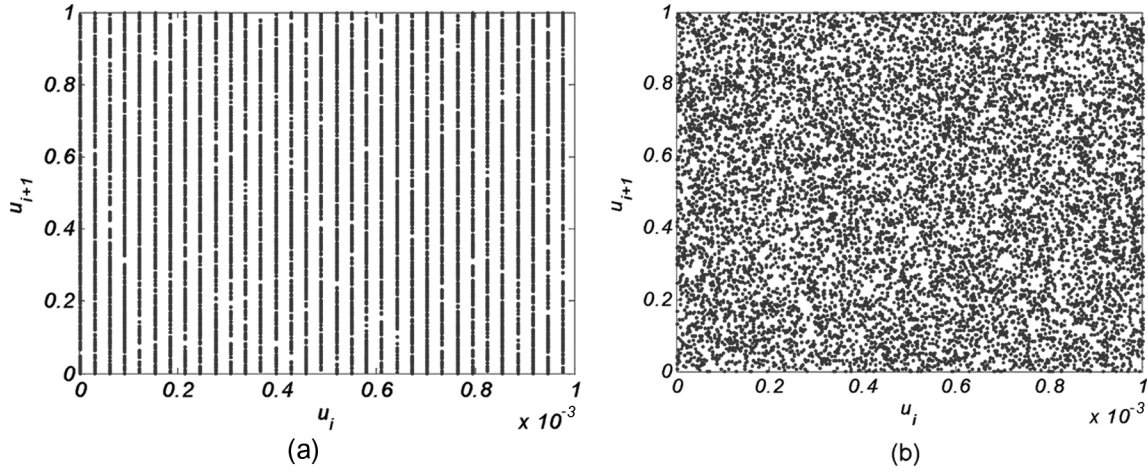


Fig. 1. 2-D scatter plot of 10^7 PN pairs (u_i, u_{i+1}) generated with (a) a 52-bit LFSR with primitive polynomial $p(x) = x^{52} + x^3 + 1$ and (b) a CTG with $\rho = 2^{258}$ [48], when a small portion of the u_i -axis is magnified.

TABLE I
POLAR-BASED [11], ZIGGURAT-BASED [16], WALLACE-BASED [15], AND
INVERSION-BASED [20] GVG DESIGNS

Design ^a	[11]	[16]	[15]	[20]
Maximum deviation	N/A	N/A	7σ	8.2σ
Clock freq. (MHz)	73	170	155	232
Output rate (MGVs/sec)	146	168	155	232
Number of logic cells	336	891	770	548
Resource utilization	7.0%	3.8%	3.3%	2.3%
On-chip memory blocks	1.4	4	6	2
18 × 18-bit Mults.	N/A	2	4	2
Bit-width of GV's	19	32	24	16

^aReference [11] used the Altera Mercury EP1M120F484C7 FPGA; the three other designs used a Xilinx Virtex-II XC2V4000-6 FPGA.

TABLE II
SOME PUBLISHED FPGA IMPLEMENTATIONS OF
GVGS BASED ON THE BM METHOD

Design ^a	[10]	[8]	[12]	[17]	[19]
Maximum deviation	4.0σ	4.8σ	6.7σ	5.0σ	8.2σ
Clock freq. (MHz)	98	245	133	165	233
Output rate (MGVs/sec)	24.5	245	133	165	466
Number of logic cells	437	480	2514	702	1528
Resource utilization	8.0%	2.0%	10.9%	3.0%	6.6%
On-chip memory blocks	0.5	5	2	5	3
18 × 18-bit Mults.	N/A	5	8	1	12
Bit-width of GV's	12	16	32	16	16

^aReference [10] used the Altera Flex10KE FPGA; the four other designs used a Xilinx Virtex-II XC2V4000-6 FPGA.

III. CLOSER LOOK AT PNGS

Using the BM algorithm, the normality of the resulting distribution depends on the statistical properties of the PNGs (how closely they resemble truly random sequences) and on the accuracy of the $f(\cdot)$ and $g(\cdot)$ computations. The class of linear PNGs has been used widely in many applications for several decades. However, many linear PNGs that perform well in standard statistical tests for randomness are known to be inadequate for MC simulations [38], [39]. Linear PNGs have two main deficiencies. First, a regular lattice structure may be present in the d -tuples constructed from successive random numbers [40]–[42]. Second, they tend to produce correlated low-order bits as well as

TABLE III
PNGS USED IN SOME PUBLISHED GVG DESIGNS

Design	Type	Period
[10]	LFSR	2^{60}
[8]	LF-LFSR	2^{190}
[12]	LFSRs	2^{60}
[17]	LFSR	2^{52}
[15], [16], [19], [20]	CTG	2^{88}

long-range correlations, especially for intervals that are a power of two [43]–[45].

Even though linear PNGs have inferior statistics, linear feedback shift registers (LFSRs) have been widely used in many hardware applications. For example, the GVG in [12] uses 50 60-bit LFSRs operating in parallel with different initial seeds to generate PN's within (0,1). The hardware-based GVG in [8] also uses a multiple-bit leap-forward LFSR (LF-LFSR) [46]. Fig. 1(a) shows the 2-D scatter plot of 10^7 PN's pairs (u_i, u_{i+1}) generated using a 52-bit LFSR with the primitive polynomial $p(x) = x^{52} + x^3 + 1$ used in [17] when a small portion of the u_i -axis is magnified. A lattice structure is clearly visible for this LFSR. Hence, LFSRs and parallel LFSR implementations may not be adequate for MC simulation [47].

One general recommendation for using linear PNGs is that the period length ρ should be chosen to be much larger than the number of random numbers required by the MC simulation [39], [43]. In fact, based on empirical experiences, the square root of the period length of the linear PNG has been reported to be a prudent upper bound for the usable sample size [44]. Rather than implementing a single linear PNG component with a very long period (e.g., $\rho \approx 2^{200}$ or more), several simple PNGs that are chosen properly and allow for very fast implementation, such as Tausworthe generators (TGs) [49], can be combined. The period of a suitably combined generator will be the product of the periods of its components [44]. Moreover, a combined generator yields sequences that have much less regular structure than the corresponding sequences of their individual component generators [48], while running almost as fast as the component generators. Table III gives the characteristics of the LFSRs and combined TGs (CTGs) used in different recently published GVGs.

TABLE IV
MAXIMUM ABSOLUTE VALUE OF A GV FOR VARIOUS PRECISIONS OF u_1

WL	8	16	24	32	48	64
Maximum $ f(u_1) $	3.33	4.70	5.76	6.66	8.15	9.41

Even though a lattice structure is not visible in the measured scatter plot in Fig. 1(b), in general, regularities exist in every output sequence from PNs [50], including CTGs [42]. Since hidden correlations and lattice structures between PNs can compromise simulation results [51], the important issue is to determine if an application is sensitive to the particular regularities and would thus yield biased results. If the structure of the output sequence of a linear PNG is too regular, then one may use nonlinear PNGs [41] or combine a linear PNG with a long period with a nonlinear PNG to reduce the regularity. Empirical tests have shown that combined linear-nonlinear PNGs are more robust than linear ones [52] and remain useful for longer sequences than linear PNGs [39], [45]. However, the main drawback of nonlinear PNGs is that they are considerably slower than linear ones, which may make them unacceptable for very long-running MC simulations.

While the randomness of the PNs impacts the normality of the generated Gaussian distribution, the precisions of u_1 and $f(u_1)$ limit the range of the generated GVs. Since $|g(\cdot)| \leq 1$, the maximum absolute value of a GV beyond 1.0 is determined by the function $f(\cdot)$. We will show in Section IV that the precision of u_2 is less critical and so the $g(\cdot)$ values can use a smaller bitwidth. Assume that $u_1 \in (0, 1)$ is represented in the unsigned fixed-point format $Q(WL)$, where WL is the word length. The closer the value of u_1 is to 0, the larger is the precision that is required for u_1 (i.e., the larger the WL), the greater is the value of $f(u_1)$ and, therefore, the greater is the range of the generated GVs. One way to ensure that the absolute value $|n|$ of the generated GVs is greater than or equal to $\varsigma\sigma$, for some desired factor ς , is to choose the precision WL of u_1 to be large enough so that $\sqrt{2WL \ln(2)} \geq \varsigma\sigma$. One can readily verify that for 32-bit precision in u_1 , $|n| \leq 6.66\sigma$ can be obtained. Table IV presents the maximum representable value of $f(u_1)$, namely $\sqrt{2WL \ln(2)}$, for various precisions of u_1 .

IV. APPROXIMATING $f(\cdot)$ AND $g(\cdot)$ FUNCTIONS

The logarithm and square root operations required in $f(u_1)$ can be computed using various techniques such as iterative procedures and series expansions [33], [53]. The execution times of the iterative procedures grow logarithmically or linearly with the precision of u_1 [33]. Consequently, for the high precision PNs that are required to generate GVs well into the tails of the Gaussian distribution, the output rate can be relatively slow [18]. On the other hand, using a series expansion to calculate logarithms may result in unacceptable PDF accuracy if an insufficient number of terms of the Taylor series is included [6]. To avoid these limitations, we adopted a polynomial curve fitting approach [21] to approximate $f(u_1)$ between (0,1). In this application, a polynomial approximation represents the continuous function $f(u_1)$ with a polynomial $p(u_1) = \sum_{v=0}^V a_v u_1^v$ of finite degree V over an interval $[\alpha, \beta]$. The various common curve fitting algorithms, such as linear or cubic interpolations or

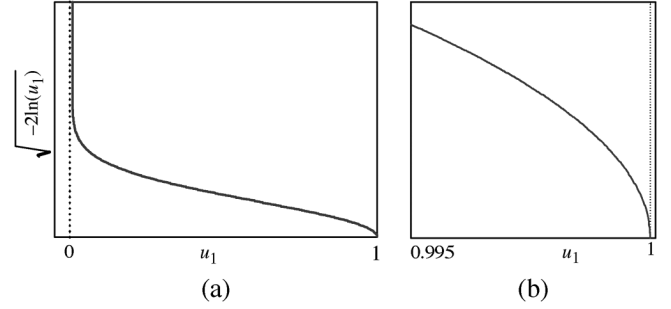


Fig. 2. (a) Plot of $f(u_1) = \sqrt{-2 \ln(u_1)}$. (b) Nonlinear behavior of $f(u_1)$ in the vicinity of $u_1 = 1$.

rational polynomial interpolations, differ from each other with respect to the computational complexity and the residual error, which is given by

$$\xi = \left\| f(u_1) - \sum_{v=0}^V a_v u_1^v \right\| \quad (1)$$

where $\|\cdot\|$ denotes a suitable norm [21]. A maximum likelihood estimate of the polynomial coefficients can be obtained using the orthogonal least squares fit (OLSF) method [54], which minimizes the value of $\sum_{v=1}^V d_v^2$, where d_v denotes the maximum perpendicular distance from the v th point on the polynomial to the point on $f(u_1)$. Due to the nonlinear shape of $f(\cdot)$, a relatively high-degree single polynomial is required to approximate $f(\cdot)$ accurately over the full interval (0,1). Instead, to increase the speed of the computation, the (0,1) interval can be divided into several segments bounded at a set of points called joints. The value of $f(\cdot)$ within each segment can then be approximated more efficiently using separate spline functions. The fewer the segments, the higher the degree of the polynomial that is required to approximate $f(\cdot)$ within each segment. One can thus trade off the order of the polynomial against the number of segments. The polynomial degree directly affects the residual error (including the approximation error and computation error), hardware complexity, memory and functional resource utilization (e.g., a degree V polynomial requires V multipliers and V adders), and latency. A more compact implementation could be obtained using time-shared hardware, but the latency would be increased.

Appropriate segmentation is crucial to the quality of the resulting GVG. The optimal partitioning into segments primarily depends on the nonlinear shape of the function and also the maximum required polynomial degree for each segment. Although any choice of segment boundaries can easily be implemented in a software simulation, some choices imply overly complex decoding circuitry and are thus undesirable for high-speed GVG realization. As shown in Fig. 2, the function $f(u_1)$ has two high-slope regions: in the vicinity of $u_1 = 0$ (which impacts the value of GVs in the tail) and close to $u_1 = 1$ (which impacts the value of GVs in the middle of the distribution). Since a small input change may lead to a (very) large output change, the input domains near 0 and 1 need smaller segments than the relatively linear regions in the middle of the domain.

Different nonuniform segmentation schemes were already proposed in [10] and [12]. In [10], only the nonlinear region close to 0 is segmented nonuniformly and it utilizes on-chip

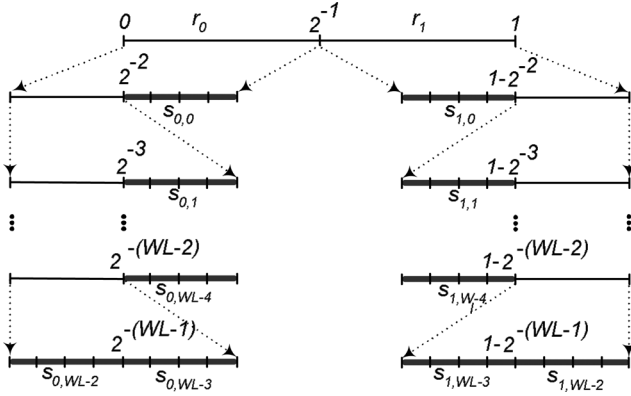


Fig. 3. Hybrid (logarithmic and uniform) segmentation of $u_1 \in (0, 1)$.

memory to store the precomputed values of $f(\cdot)$ in memory. The GVG released by Xilinx [8] appears to be based on the GVG design in [10]. However, the analysis in [17] shows that the accuracy of this GVG degrades at the tails of the PDF for $|n| \geq 4.8\sigma$. The method in [12] uses nonuniform segmentation in both the regions close to $u_1 = 0$ and $u_1 = 1$ and uses a piecewise linear approximation for more accurately computing $f(u_1)$ within each segment. The central limit theorem (CLT) [29], [37] was exploited in both [10] and [12] to improve the statistics of the resulting distribution by averaging the output of multiple GVs. The quality of the GVG in [12] was improved in [19] by utilizing a more accurate approximation scheme that increases the quality of the generated GVs further into the tails. In [19], rather than approximating the nonlinear function $f(u_1) = \sqrt{-2\ln(u_1)}$, the square root and logarithm functions are evaluated separately.

We propose a hybrid segmentation scheme in which (similar to [55] but without requiring the CLT stage) both logarithmic and uniform segmentations are utilized. First, the domain $(0, 1)$ of u_1 is divided into two subintervals $r_0 \in (0, 0.5)$ and $r_1 \in [0.5, 1)$. Let u_1 be represented as an unsigned fixed-point number with $WL = 32$ bits of precision. The value, 0 or 1, of the most significant bit (MSB) of u_1 indicates whether a particular u_1 resides in subinterval r_0 or r_1 , respectively. Subinterval r_0 is then segmented logarithmically into $WL - 1$ segments from $u_1 = 0.5$ down to 0, as shown in Fig. 3. Subinterval r_1 is segmented similarly from $u_1 = 0.5$ up to 1. Clearly, the number of logarithmic segments $W = 2(WL - 1)$ is limited by the precision of u_1 . Each segment will be denoted by $s_{r,w}$, where binary subscript r specifies the half range (r_0 or r_1) and w denotes the logarithmic segment number $w = 0, \dots, WL - 2$. Then, each segment $s_{r,w}$ is subdivided uniformly into $L = 2^\ell$ subsegments where each subsegment is denoted by $s_{r,w,l}$. Note that if one of the $s_{r,w}$ segments, where $w = WL - \ell - 1$ to $w = WL - 2$ is addressed, then u_1 does not have sufficient bits to address the uniform segments within $s_{r,w}$. To support this scheme, we need to use $WL + \ell$ -bit precision for u_1 . The number of uniform segments depends on the desired accuracy and on the size of memory that is available to store the coefficients of $2^{\ell+1}(WL - 1)$ polynomials. Note that since approximating $f(u_1)$ close to $u_1 = 1$ impacts the error in the center of the generated distribution and approximating $f(u_1)$ close to

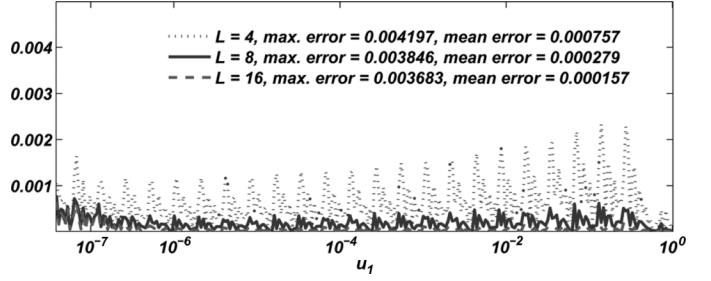


Fig. 4. Absolute approximation error of $f(\cdot)$ for different numbers of uniform segments.

$u_1 = 0$ impacts the error at the tails of the distribution, the hybrid segmentation scheme considers both regions carefully by logarithmic and uniform segmentation to achieve an accurate overall approximation for $f(\cdot)$. Fig. 4 plots on a logarithmic scale the resulting absolute error when approximating $f(\cdot)$ using $W = 62$ for three different numbers of uniform segments $L = 4, 8, 16$. Fig. 4 shows that for $L = 8$ and $L = 16$ there is relatively little degradation in the accuracy of $f(u_1)$.

After segmentation, the coefficients of the polynomials for approximating $f(u_1)$ within $2^{\ell+1}(WL - 1)$ different segments are calculated and optimized using the OLSF method to minimize the residual error. An important point to note is that as u_1 approaches 2^{-WL} from above (for the nonlinear region just above $u_1 = 0$) or when u_1 approaches $1 - 2^{-WL}$ from below (for the nonlinear region just below $u_1 = 1$), the slope $df(u_1)/du_1$ tends toward infinity and, therefore, the coefficients of $p(u_1)$ become very large. However, the value of $p(u_1)$ lies within $(0, \sqrt{2WL \ln 2})$. For example, for a $WL = 32$ -bit representation of u_1 , the coefficients of $p(u_1)$ are of the order of 2^{32} but the value of $|p(u_1)|$ lies in $[0, 6.66]$, which can be represented relatively accurately in two's complement 16-bit fixed-point format with a 12-bit fraction, i.e., in $Q(16, 12)$. Storing the large coefficient values of $p(u_1)$ on-chip requires large memories, increases the hardware complexity and may slow down the variate generation rate. To overcome these problems, [12] proposed to trade off precision for range and store scaling factors (multiples of two) along with the coefficients into an on-chip memory. While scaling is performed dynamically in [12], we propose a different scaling method that can also be used statically (i.e., at design synthesis time). Using this method, only the adjusted coefficients of $p(u_1)$, for all $2L(WL - 1)$ segments are stored in on-chip memory. This scheme reduces the memory requirements, decreases the hardware complexity, and maintains the accuracy of the computation.

For clarity, we explain the scaling method assuming that each segment is approximated using a first-degree polynomial $p(u_1) = a_{r,w,l}u_1 + b_{r,w,l}$. However, the scaling scheme is independent of the number of logarithmic segments and the order of the polynomial. When u_1 lies within r_0 and in segment $s_{0,w}$, a new scaled variable $\tilde{u}_{1,w}$ is defined as $\tilde{u}_{1,w} = 2^{wu_1}$. To compensate for the scaling of u_1 , the $a_{0,w,l}$ slope of segment $s_{0,w,l}$ is shifted to the right w bit positions as $\tilde{a}_{0,w,l} = 2^{-w}a_{0,w,l}$. Thus, the largest slope values are scaled down by a factor of 2^{-WL+2} . When u_1 lies within $s_{1,w}$, to scale the slopes and intercepts of $p(\cdot)$ accurately, the variable u_1 is transformed to a new variable $\hat{u}_1 = 1 - u_1$. Thus, as $u_1 \rightarrow 1^-$, $\hat{u}_1 \rightarrow 0^-$. Similar to the procedure for region r_0 , when u_1 resides in segment $s_{1,w,l}$, the

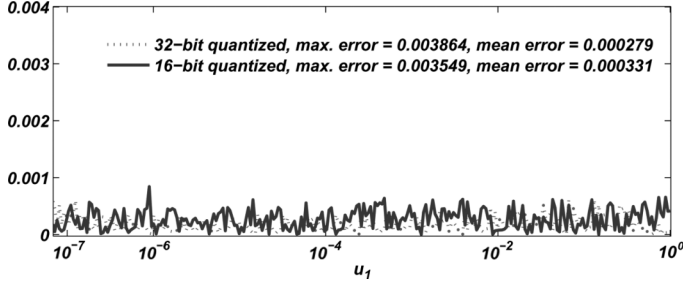


Fig. 5. Absolute value of the approximation error of $f(\cdot)$ for two different quantization formats.

large value of $a_{1,w,l}$ can be scaled down by 2^{-w} and $b_{1,w,l}$ can be adjusted as $a_{1,w,l} + b_{1,w,l}$.

According to the value of u_1 , an addressing unit (AU) is required to calculate the scaled value of u_1 , namely \tilde{u}_1 . The AU also identifies the subinterval r_i , the segment number w , and the sub-segment number l . Then the scaled coefficients of $p(u_1)$ can be addressed and read directly from memory to approximate $p(\tilde{u}_1)$. When u_1 resides in segment $s_{0,w,l}$, $p(\tilde{u}_1)$ will be computed as

$$p(\tilde{u}_1) = \tilde{a}_{0,w,l}\tilde{u}_1 + b_{0,w,l} = 2^w \tilde{a}_{0,w,l}u_1 + b_{0,w,l}. \quad (2)$$

When u_1 resides in segment $s_{1,w,l}$, then $p(\tilde{u}_1)$ will be computed as

$$p(\tilde{u}_1) = -2^{-w}a_{1,w,l}\tilde{u}_1 + (a_{1,w,l} + b_{1,w,l}). \quad (3)$$

Using the scaled coefficients and \tilde{u}_1 , the computations in (2) and (3) are representable using 16-bit fixed-point numbers. For example, an experimental range analysis of the values of $\tilde{a}_{r,w,l}$ and $\tilde{b}_{r,w,l}$ for a first-degree polynomial approximation within each segment shows that these values can be represented within 16 bits with $Q(16, 14)$ and $Q(16, 12)$, respectively. Fig. 5 plots over a logarithmic scale the absolute value of the approximation error for $f(u_1)$ using $W = 62$ and $L = 8$ for two different quantization formats. The relatively similar values of the mean and maximum errors demonstrates the effectiveness of our segmentation scheme in equally limiting the approximation error over the range of $u_1 \in (0, 1)$.

There are various standard approaches for approximating trigonometric functions such as CORDIC algorithms [56], polynomial approximations, and various table lookup schemes (e.g., direct table lookup and multipartite techniques [57]). The choice of a method and a particular implementation depends on such requirements as throughput, latency, and area as well as the required accuracy. The accuracy is determined by the error of the approximation and by the roundoff errors that occur during the evaluation of the approximation.

Storing and later retrieving quantized values of the trigonometric function in an on-chip memory, so-called table lookup, is relatively fast, but the approximation accuracy is limited by the size of the on-chip memory. What is more, the size of the memory grows exponentially with the size of the input word, which confines this solution to relatively small input precisions (i.e., ≤ 12 bits). In the present application the table size can be reduced by exploiting the symmetry properties of the $\sin(\cdot)$ and $\cos(\cdot)$ functions and then storing only a quarter cycle of the trigonometric function in memory.

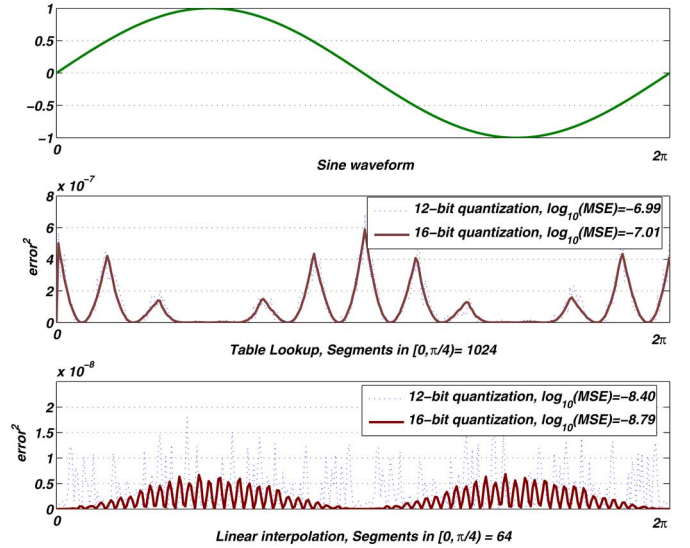


Fig. 6. Sine approximation errors using table lookup and linear interpolation.

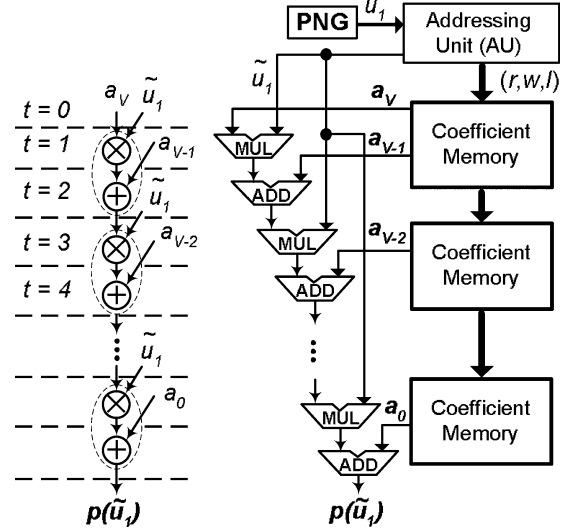


Fig. 7. Dataflow diagram for computing $p(\tilde{u}_1)$.

Another scheme that may require less memory, compared to table lookup but with similar accuracy, is to use linear (or higher order) interpolation. For example, Synopsys Inc. provides a trigonometric Intellectual Property (IP) core to compute $\sin(2\pi u_2)$ by segmenting the quarter cycle into 64 straight-line segments [58]. Fig. 6 plots the squared error of the table lookup method with 1024 segments over the quarter cycle along with the one for the linear interpolation method with 64 segments over the quarter cycle. Fig. 6 shows that the mean-squared errors (MSEs) of these two schemes, when implemented using 16-bit fixed-point format, are relatively close.

V. REALIZATION OF THE GVG

Fig. 7 shows (on the left) a dataflow diagram illustrating the evaluation of $p(\tilde{u}_1)$ in its factored form and (on the right) the corresponding hardware datapath. The core of the GVG contains pipelined fixed-point multipliers, adders, registers, and routing resources. The operations are fully pipelined and scheduled to maximize the output rate. To form the PNG, we used 32- and 64-bit CTGs [59], [60] that can be implemented

TABLE V
PERFORMANCE OF THREE CTGs IMPLEMENTED ON THE XILINX VIRTEX-II XC2V4000-6

Design	Output bitwidth	Approx. period	Logic cells	Clock freq. (MHz)
CTG88	32	2^{88}	72	400
CTG133	32	2^{133}	114	400
CTG258	64	2^{258}	254	400

using only shift and bitwise logical operations [48], [61]. The characteristics of three CTGs implemented on the Xilinx Virtex-II XC2V4000-6 are compared in Table V. To determine the segment number w of a given PN input u_1 , the AU uses a small leading ones detector (LOD) circuit [62]. The LOD requires 57 and 122 slices for a 32- and 64-bit u_1 , respectively, on a Virtex-II XC2V4000-6 FPGA. For a 32-bit u_1 , the AU requires 226 configurable slices while it requires 511 slices for a 64-bit u_1 . The AU operates at 275 MHz for both the 32- and 64-bit u_1 . The generated triple (r, w, ℓ) from the AU is used to address the coefficient memory. For $WL = 32$ (thus $31 \times 2 = 62$ logarithmic segments in both the r_0 and r_1 regions) and $L = 8$ uniform segments within each logarithmic segment, one BlockRAM is required to store the $2 \times 62 \times 8 = 992$ scaled coefficients $\tilde{a}_{r,w,\ell}$ and $\tilde{b}_{r,w,\ell}$ in the 16-bit fixed-point formats $Q(16, 14)$ and $Q(16, 12)$, respectively.

The important feature of the proposed approximation strategy is that due to the efficiency of the hybrid segmentation and scaling scheme, the dataflow diagram for computing $p(\tilde{u}_1)$ can be implemented using only 16-bit fixed-point variables, almost independent of the desired tail accuracy. In fact, the precision of u_1 and the size of the coefficient memory can always be adjusted to generate GVs with various tail accuracies using the same 16-bit datapath. To do this, only the PNG and AU must be modified slightly, and the coefficient memories configured correspondingly. While five bits are required to represent the integer part of generated GVs up to 15σ , 11 bits are dedicated to the fractional part to represent GVs in two's complement format. Fortunately, the 16-bit GV format tends to be a common precision for many digital signal processing applications. The Xilinx AWGN core [8] also uses the 16-bit fixed-point format.

To approximate $g_1(u_2)$ and $g_2(u_2)$, the quarter cycle of the sine function is partitioned into 1024 uniform segments. The precomputed values in fixed-point format $Q(16, 15)$ are stored in a dual-port memory block. Using the symmetry of the sine and cosine functions, $g_1(u_2)$ and $g_2(u_2)$ can then be approximated compactly. The table lookup scheme not only provides smaller and faster implementation (as shown in Table VI), it also has relatively small MSE, as discussed in Section IV. The calculated $f(u_1) = p(\tilde{u}_1)$ is then multiplied by $g_1(u_2)$ and $g_2(u_2)$ to generate two GVs in the fixed-point format $Q(16, 11)$.

Table VII summarizes the implementation results for the new GVG on three different FPGAs, utilizing first-degree approximations. The results in Table VII show that the output rate of the proposed GVG is independent of the tail accuracy. In fact, the design can be configured to achieve higher tail accuracy with only a small cost in extra hardware. Moreover, if there are sufficient resources available on the FPGA beyond those required

TABLE VI
CHARACTERISTICS OF TWO DIFFERENT IMPLEMENTATIONS OF BOTH $g_1(u_2)$ AND $g_2(u_2)$ ON A XILINX VIRTEX-II XC2V4000-6 FPGA

Design ^a	Clock freq. (MHz)	Logic cells	Memory blocks	18×18 -bit Mults.
I	272	60	1	0
II ^b	142	107	1	2

^aDesign I uses a table lookup scheme. To approximate $\sin(2\pi u_2)$ and $\cos(2\pi u_2)$, the quarter cycle is partitioned into 1024 uniform segments. Design II segments sine and cosine waveforms each into 256 uniform segments and use linear interpolation to approximate their values over $(0, 2\pi)$. The coefficients of each segment in fixed-point format $Q(16, 12)$ are stored in one dual-port memory block.

^bA faster implementation of this design can run at 250 MHz while requires 144 slices.

TABLE VII
TYPICAL REALIZATIONS OF THE PROPOSED GVG

Device ^a	I	II	III	IV
Bitwidth of u_1	32	32	64	32
Period of the PNG	$\approx 2^{113}$	$\approx 2^{88}$	$\approx 2^{258}$	$\approx 2^{133}$
Max. deviation	6.66σ	6.66σ	9.4σ	6.66σ
Clock freq. (MHz)	269	248	248	336
Output rate (MGVs/sec)	538	496	496	672
Number of logic cells	576	534	852	668
Resource utilization	1.3%	2.3%	3.6%	0.4%
On-chip memory blocks	2	2	3	2
18×18 -bit Mults.	3	3	3	3

^aDesign I was synthesized for a Xilinx Virtex-II Pro XC2VP100-6 FPGA. Designs II and III were synthesized for a Xilinx Virtex-II XC2V4000-6 FPGA. Design IV was synthesized for an Altera StratixII EP2S180F1508C4 FPGA.

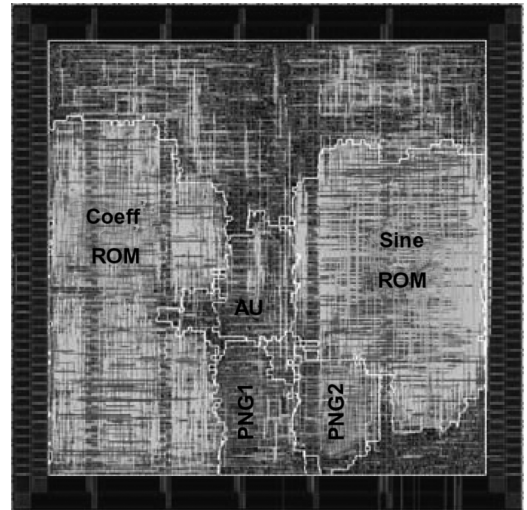


Fig. 8. 0.126-mm² GVG chip layout in 90-nm CMOS technology.

by a single GVG datapath, then multiple instances of the same GVG datapath, with different initial seeds for the PNGs, could be readily instantiated to speed up the total GV generation rate.

Fig. 8 shows the layout of the 0.126-mm² GVG chip designed in a 90-nm CMOS technology using a dual-threshold

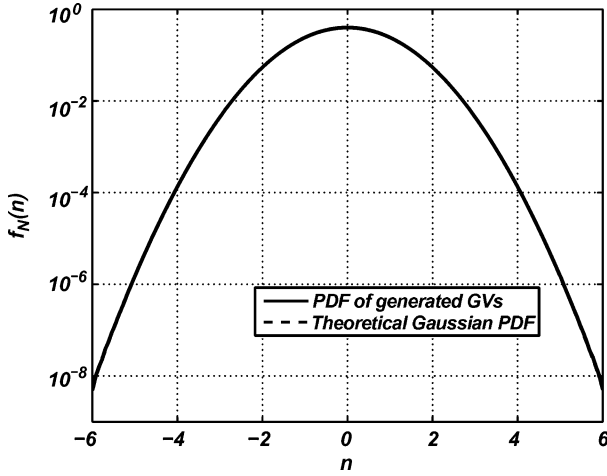


Fig. 9. Gaussian PDF compared with the PDF of 10^{11} generated GV's.

standard cell library. The core area is dominated by the coefficient ROM and the dual-port sine ROM. Without access to a commercial ROM core generator, we implemented the ROM array using standard cells. Custom ROM blocks would significantly reduce the required area. The unlabeled area in the core layout is occupied by two-stage pipelined multipliers, adders, registers and routing resources. The core operates at 537 MHz, generating over one billion GV's per second while dissipating 12.3 mW of dynamic power. The static power dissipation was estimated to be 9.91 mW. This core has been implemented successfully in silicon in a 500-Mb/s one-half rate low-density parity-check convolution decoder [63] to permit measurements of BER versus SNR at-speed and on-chip.

VI. GVG STATISTICAL TESTS

Normality tests are well-known statistical measures used to determine if generated GV's fit the expected normal distribution [64]. The power of statistical tests for different characteristics of the normal distribution varies depending on the nature of any deviations from ideal normality.

Since the tail of the Gaussian PDF is not visible on the linear scale, Fig. 9 superimposes the PDF of 10^{11} generated GV's on top of a PDF plot of the exact normal distribution using a logarithmic scale. The two plots are indistinguishable over $\pm 6.0\sigma$. To generate the GV's at the extreme tails of the distribution, say where $6.0\sigma < |n| < 9.4\sigma$, at least 10^{22} samples are required to accurately measure the PDF, which takes a prohibitively long time. Instead, the PDF of the Gaussian variable n , $f_N(n)$, can be expressed in terms of its CDF, $F_N(n)$. Note that the PDF can be written as

$$f_N(n) = \frac{d}{dn} F_N(n) = \frac{d}{dn} \Pr(N < n). \quad (4)$$

The “importance sampling” expression can be written using Bayes' law as

$$F_N(n) = F_N(n|U_1 < u_\tau) \Pr(U_1 < u_\tau) + F_N(n|U_1 \geq u_\tau) \Pr(U_1 \geq u_\tau). \quad (5)$$

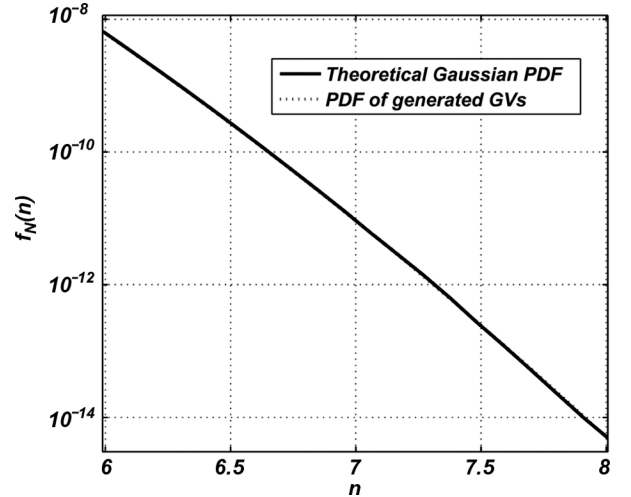


Fig. 10. Gaussian PDF compared with the PDF of the 5×10^8 generated GV's for $6 \leq |n| \leq 8$.

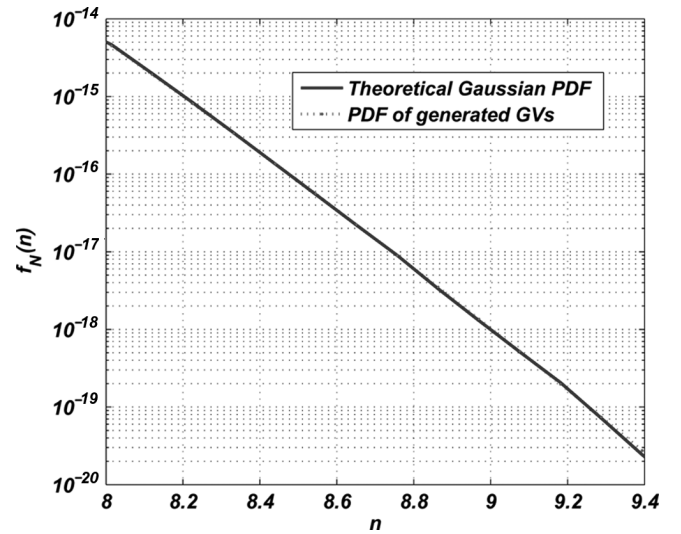


Fig. 11. Gaussian PDF compared with PDF of the 5×10^8 generated GV's for $8 \leq |n| \leq 9.4$.

To measure the PDF in the tails of the distribution, GV's such that $U_1 < u_\tau \ll 1$ are first generated. The PDF of the generated GV's can then be given as

$$f_N(n) = \frac{d}{dn} F_N(n|U_1 < u_\tau) \Pr(U_1 < u_\tau) \forall |n| > \sqrt{-2 \ln(u_\tau)}. \quad (6)$$

In this method, we do not generate $U_1 \geq u_\tau$ and, therefore, the second term of (5) approaches 0. Using this method, we divided the 6σ to 9.4σ region into two sub-regions, from 6σ to 8σ and from 8σ to 9.4σ , and then generated 5×10^{13} GV's. The theoretical Gaussian PDF is compared on a logarithmic scale in Figs. 10 and 11 with the PDF of generated GV's at the very end of the Gaussian distribution tails.

The Pearson Chi-Square test can be utilized to determine the normality of generated GV's with a desired significance level α [64]. The Chi-Square test involves quantizing the horizontal axis of the PDF into γ cells. The $\chi^2_{\alpha, \gamma-1}$ statistic can be calculated

TABLE VIII
 χ^2 -TEST SIMULATION RESULTS

Range	Calculated $\chi^2_{\alpha,97}$	Threshold				
		α	132 0.01	121 0.05	115 0.10	111 0.15
$ n \leq 4.5$	109		pass	pass	pass	pass
$4.5 < n < 6.0$	90		pass	pass	pass	pass
$6.0 < n < 7.5$	103		pass	pass	pass	pass
$7.5 < n < 9.4$	110		pass	pass	pass	pass

based on the actual and expected number of samples appearing in each cell and serves as an overall quality metric as follows:

$$\chi^2_{\alpha,\gamma-1} = \sum_{i=1}^{\gamma} \frac{(m_i - KP_i)^2}{KP_i}$$

where K is the number of generated GV's. $\chi^2_{\alpha,\gamma-1}$ is a Chi-Square random variable with $\gamma - 1$ degrees of freedom (DOF) [64]. For each cell i , m_i is the number of samples observed in the bin and KP_i is the expected number of samples according to the normal distribution. Since a normal distribution is completely specified by two measures, the mean and the standard deviation, the number of DOF is reduced by 2 from $\gamma - 1$ to $\gamma - 3$ [65]. To perform the Chi-Square test on the generated GV's, we divided the horizontal axis of the PDF into four regions, as given in Table VIII, and segmented each interval into 100 bins and ensured that at least 50 GV's fall into each bin. The Chi-Square test is passed if the calculated $\chi^2_{\alpha,\gamma-3}$ is smaller than a given threshold. Table VIII shows that the designed GVG successfully passes the Chi-Square test, even for stringent values of the significance level. For more details on the Chi-Square test please refer to [64]. The Anderson-Darling statistic test enhances the sensitivity of the statistic in the distribution tails. The \mathcal{A}^2 test statistic for a normal distribution can be calculated numerically as [64]

$$\mathcal{A}^2 = -K - 1/K \sum_{i=1}^K (2i - 1) [\ln(F_Z(z_i)) + \ln(1 - F_Z(z_{K+1-i}))]$$

where $F_Z(\cdot)$ is the standard normal CDF and $\{z_i\}$ are the ordered generated GV's. Since the measured parameter of $\mathcal{A}^2(1 + 0.75/K + 2.25/K^2)$ was less than the critical value of 0.752, we can again accept the normality hypothesis for our GVG.

To evaluate the correlation among subsequently generated GV's, a sequence containing 10^7 variates generated by our GVG was subjected to the linear Pearson's correlation test. No regular lattice structure was apparent, as shown in Fig. 12. Fig. 13 plots the autocorrelation values over the range of lags ± 2048 for 10^7 generated GV's. The approximated 95% confidence interval for the calculated autocorrelation was at $\pm 6.32 \times 10^{-4}$. The autocorrelation is evidently very low for all nonzero lags.

To evaluate the accuracy of the design in a BER simulation, the GVG core was instantiated in a communication system with BPSK signaling over an AWGN channel. Fig. 14 plots the theoretical BER against the generated simulation result using our GVG. For the sake of comparison, we implemented a GVG based on the CLT using a summation of 12 uniform PNs [37]. This plot shows that the use of an inaccurate GVG may lead to inaccurate simulation results. Specifically, at $\text{BER} = 10^{-10}$ the inaccurate

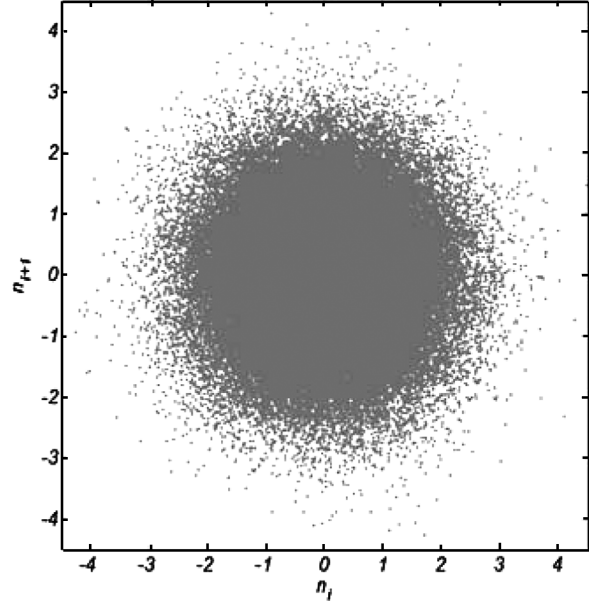
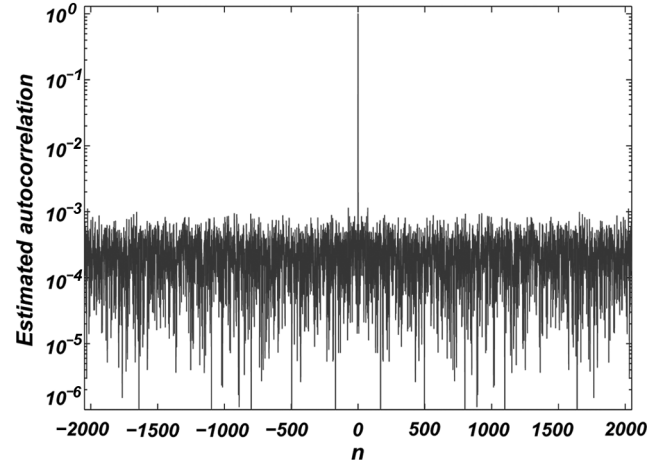
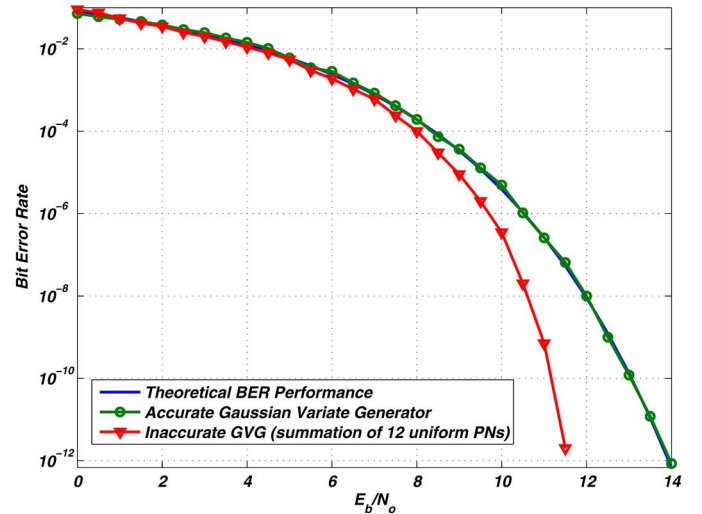
Fig. 12. 2-D scatter plot of 10^7 generated GV's.Fig. 13. Estimated autocorrelation for 10^7 generated GV's.

Fig. 14. Application of the GVG in BER simulation of a BPSK modulated communication system over an AWGN channel.

Gaussian noise results in an almost 2-dB shift in the measured BER, and the shift gets worse at higher values of E_b/N_o .

VII. CONCLUSION

In this paper, a fast and compact GVG was described that has a faster Gaussian variate generation rate, with a lower hardware cost, and generates Gaussian samples up to σ values larger than published designs (as summarized in Tables I, II, and VII). The functions required by the BM method were approximated using polynomial curve fitting, hybrid segmentation (i.e., combination of logarithmic and uniform segmentation), and a scaling scheme to maintain the accuracy of approximation. The scaling scheme is independent of the number of segments and the order of the polynomial. The polynomial coefficients are scaled at synthesis time to reduce the memory requirements, decrease the hardware complexity, and maintain the accuracy of the computation. The dataflow diagram of the GVG can be implemented using 16-bit fixed-point variables to generate GVs in $Q(16, 11)$ format for arbitrary tail ranges of up to $\pm 15\sigma$. The synthesizable GVG core can be easily configured to generate GVs in two's complement fixed-point format for larger tail values at a small cost in extra hardware, without degrading the sample rate (see Table VII). A typical instantiation of the proposed GVG uses only 1.3% of the Xilinx Virtex-II Pro XC2VP100-6 FPGA, two on-chip block memories, and three dedicated multipliers and operates at 269 MHz, generating 538 million GVs per second within up to $\pm 6.66\sigma$. The implementation costs and performance were illustrated by typical realizations for FPGAs and a 90-nm CMOS ASIC. The proposed GVG should therefore be of significant assistance in the characterization of low-BER systems.

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