Stochastic Formal Methods for Rare Failure Events due to the Accumulation of Errors

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Abstract—This paper provides an accurate bound on the number of numeric operations (fixed or floating point) that can safely be performed before accuracy is lost based on the assumption that accumulated errors are uniformly distributed in $\pm \frac{1}{2}$ unit in the last place. This work has important implications for control systems with safety-critical software, as these systems are now running fast enough and long enough for their errors to impact on their functionality. Furthermore, worst-case analysis would blindly advise the replacement of existing systems that have been successfully running for years and that will continue running before software development practices evolve. We present here new theorems that we are currently validating with the PVS proof assistant. This theory will allow code analyzing tools to produce formal certificates of accurate behavior. FAA regulations require that the probability of an error be below $10^{-9}$ for a 10 hour flight \([1]\). Such a low failure rate is stretching the limits of generic calculations solely based on the standard deviation of random variables for the intermediate sums. We need many individual errors for the Central Limit Theorem approximation to be sufficiently accurate (distance well below $10^{-10}$). The precise bound presented here enhances the number of bits of the result that can safely be regarded as correct.

I. INTRODUCTION

Formal proof assistants are used in areas where errors can cause loss of life or significant financial damage as well as in areas where common misunderstandings can falsify key assumptions. For this reason, formal proof assistants have been much used for floating point arithmetic \([2], [3], [4], [5], [6]\) and probabilistic or randomized algorithms \([7], [8]\). Previous references just link to a few projects using proof assistants such as ACL2, HOL \([9]\), Coq \([10]\) and PVS \([11]\).

All the above projects that deal with floating point arithmetic aim at containing worst case behavior. Recent work has shown that worst case analysis may be meaningless for applications that run for a long time. For example, a process adds numbers in $\pm 1$ with a measure error of $\pm 2^{-25}$. If this process adds $2^{25}$ items, then the accumulated error is $\pm 1$, and note that 10 hours of flight time at operating frequency of 1 kHz is approximately $2^{25}$ operations. Yet we easily agree that provided the measure errors are not correlated, the actual accumulated error will be much smaller.

We recall in Section I the model that we are using. Section II recalls our formal developments in probability and presents our new theorems. The Doobs-Kolmogorov inequality was used in \([12]\) but precise definitions of measure and round-off errors used in Section IV provide a much more precise way to compute the probability that a piece of software will successfully run within an acceptable error bound. We use the result of Section IV for intermediate sums before approximations based on the Central Limit Theorem are sufficiently accurate. As this new bound is long to obtain, it might not be required for ordinary pieces of software. Yet, failure rate as defined in aeronautics is very low and this new bound is necessary whatever the computational price.

We cannot use the Central Limit Theorem before the approximation to normal distribution is sufficiently close so that the sum including the additional approximation errors is bounded by $10^{-9}$ (there are potentially $2^{25}$ additional errors). Our suggestion is to carry on precise computing as long as possible and revert to results linked to Doobs-Kolmogorov inequality \([12]\) and the Central Limit Theorem when the approximation error becomes negligible compared to $10^{-9}$.

In the rest of this text, we assume that the created round-off and measure errors are unbiased independent random variables or that their expectation conditional to the previous errors is zero.

II. STOCHASTIC MODEL AND STATE OF THE ART

A. Individual round-off and measure errors

We are dealing with fixed or floating point numbers. A floating point number represents $v = m \times 2^e$ where $e$ is an integer and $m$ is a fixed point number \([13]\). IEEE 754 standard \([14]\) uses sign-magnitude notation for the mantissa and the first bit $b_0$ of the mantissa is implicit in most cases ($b_0 = 1$) leading to the following definition where $s$ and all the $b_i$ are either 0 or 1 (bits).

$$v = (-1)^s \times b_0 \cdot b_1 \cdots b_{p-1} \times 2^e$$

Some circuits such as the TMS320 use two’s complement notation for $m$ leading to the following definition \([15]\).

$$v = (b_0 \cdot b_1 \cdots b_{p-1} - 2 \times s) \times 2^e$$

In fixed point notation $e$ is a constant provided by the data type and the first bit is no longer implicitly equal to 1.
For all the previous notations, we define for any representable number \( v \), the unit in the last place function where \( e \) is the exponent of \( v \) as above.

\[
\text{ulp}(v) = 2^e - p + 1
\]

A variable \( v \) is set either by an external sensor or by an operation. Trailing digits of numbers randomly chosen from a logarithmic distribution \([16, p. 254-264]\) are approximately uniformly distributed. So we can assume that if \( v \) is a data obtained by an accurate sensor, the difference between \( v \) and the actual value \( \overline{v} \) is uniformly distributed in the range \( \pm \text{ulp}(v)/2 \). We can model the error \( v - \overline{v} \) by a random variable \( X \) with expectation \( \mathbb{E}(X) = 0 \) and variance \( \text{Var}(X) = \text{ulp}(v)^2/12 \). The sensor may be less accurate leading to a larger variance but we assume that it is not biased.

Round-off errors created by operators are discrete and they are not necessarily distributed uniformly \([13]\). The distribution is very specific but as soon as we verify that the expectation is \( \mathbb{E}(X) = 0 \) and the error is bounded, we may safely use a replacement variable \( X' \) such that

\[
\forall \epsilon \quad \mathbb{P}(|X'| > \epsilon) \geq \mathbb{P}(|X| > \epsilon).
\]

### B. Error of an accumulation loop

We will use only one example for very long accumulations as calculations carried on at the end of Section \([\text{X}]\) heavily rely on the fact that all the errors introduced are uniformly distributed over the same interval. In other cases such as the second example of \([13]\), one may either revert to the exponential-time formula stated in Section \([\text{IV}-\text{A}]\) or use the magnitude of the largest possible error for all the random variables.

The example given in listing 1 sums data produced by a fixed point sensor \( x_i \) with a measure error \( X_i \).

**Listing 1. Simple discrete integration from [19]**

```plaintext
1. a_0 = 0
2. for (i = 0; ; i < n; i = i + 1)
3. a_{i+1} = a_i + x_i
```

We can safely assume that \( X_i \) are independent identical uniformly distributed random variables over \( \pm \text{ulp}(x_i)/2 \). Data are fixed point meaning that the sum \( a_i + x_i \) does not introduce any rounding error and the weight of one unit in the last place does not depend on \( x_i \) so we write ulp instead of ulp(\( x_i \)). After \( n \) iterations, we want the probability that the accumulated measure error has always been constrained into user specified bounds \( \epsilon \). Using the Doobs-Kolmogorov inequality \([13]\) where \( S_i = \sum_{j=1}^{i} X_j \), we have that

\[
\mathbb{P} \left( \max_{1 \leq i \leq n} |S_i| \leq \epsilon \right) \leq 1 - \frac{\text{ulp}^2}{12\epsilon^2}.
\]

We will see that we can exhibit a tighter bound using \([23]\). This bound can be produced using a time- and space-polynomial algorithm provided all the random variables are uniformly distributed over the same interval. Complexity is a key issue as typically \( n \lesssim 2^{25} \) meaning that the quadratic algorithm proposed Section \([\text{IV}]\) should be considered as the biggest amenable algorithm.

### III. Probability Distribution of Being Safe

#### A. Probability

We presented in \([12]\) an account of probability with an informal approach while taking foundational matters seriously. The PVS system underlying these results is built on the firm foundations for probability theory (using measure theory) \([21], [23]\). A middle way between extreme formality and an accessible level of informality is to be found in \([23]\).

We now begin by recalling in Figure 2 definitions implemented in PVS \([12]\). The PVS development of probability spaces in Figure 3 takes three parameters: \( T \), the sample space, \( S \), a \( \sigma \)-algebra of permitted events, and, \( \mathbb{P} \), a probability measure, which assigns to each permitted event in \( S \), a probability between 0 and 1. Properties of probability that are independent of the particular details of \( T \), \( S \) and \( \mathbb{P} \) are then provided in this file.

If \( T \) is countable – as it is for discrete random variables – then we may take \( \sigma = \wp(T) \). As we wish to discuss continuous random variables then we partially instantiate this PVS file with \( T = \text{real} \), and \( S = \text{borel set} \) (the Borel sets). If we go further and also specify \( \mathbb{P} \), we will have described the random variable distributions as well. Of particular interest later is the fact that the sum of two random variables is itself a random variable, and consequently any finite sum of random variables will be a random variable.

Note that the product probability \( \mathbb{P}_{\mathbb{R}} \) has the effect of declaring that the experiments carried out in probability spaces \((T_1, \sigma_1, \mathbb{P}_1) \) and \((T_2, \sigma_2, \mathbb{P}_2) \) are independent. Obviously, the process of forming products can be extended to any finite product of finitely many probability spaces.

In Figure 3, we define the conditional probability \( \mathbb{P}(A; B) \) (written \( \mathbb{P}(A \mid B) \) as PVS will not permit the use of “;” as an operator). We take the opportunity to prove Bayes’ Theorem along the way.

#### B. Continuous Uniform Random Variables

If \( X \) is a continuous random variable distributed uniformly over the interval \([a, b]\), then informally it takes any value within the interval \([a, b]\) with equal probability.

To make this more formal, we define the characteristic function of a set \( S \) as the function \( \chi_S \), which takes the values 1 or 0 depending on whether it is applied to a member of \( S \).

**Definition 1:**

\[
\chi_S(x) = \begin{cases} 
1 & x \in S \\
0 & x \notin S
\end{cases}
\]

Now the probability density function \( f \) of the uniform random variable over the closed interval \([a, b]\) is \( \frac{1}{b-a} \chi_{[a,b]} \). From this we can calculate the distribution function:

\[
F(x) = \int_{-\infty}^{x} f(x)dx,
\]
probabilty_space[T:TYPE+, (IMPORTING finite_measure@subset_algebra_def[T]) % sample space
S:sigma_algebra, (IMPORTING probability_measure[T,S]) % permitted events
P:probability_measure % probability measure
]: THEORY

BEGIN

IMPORTING finite_measure@sigma_algebra[T,S],probability_measure[T,S],continuous_functions_aux[real]

A,B: VAR (S)
x,y: VAR real
n0z: VAR nzreal
t: VAR T
n: VAR nat

null?(A) :bool = P(A) = 0
non_null?(A) :bool = NOT null?(A)
independent?(A,B):bool = P(intersection(A,B)) = P(A) * P(B) % Note that it DOES NOT say = 0
random_variable?(X:[T->real]):bool = FORALL x: member({ t | X(t) <= x},S)
zero: (random_variable?) = (LAMBDA t: 0)
random_variable: TYPE+ = (random_variable?) CONTAINING zero

X,Y: VAR random_variable
XS: VAR [nat->random_variable]

<= (X,x): (S) = { t | X(t) <= x}; % Needed for syntax purposes! < > /= >= omitted
complement_le1: LEMMA complement(X <= x) = (x < X)
complement_lt1: LEMMA complement(x < X) = (X <= x)
complement_eq: LEMMA complement(X = x) = (X /= x)
complement_le2: LEMMA complement(x <= X) = (X < x)
complement_ne: LEMMA complement(X /= x) = (X = x)

-(X) :random_variable = (LAMBDA t: -X(t)); % Needed for syntax purposes! + - * / omitted
+(X,Y) :random_variable = (LAMBDA t: X(t) + Y(t));
-(X,Y) :random_variable = (LAMBDA t: X(t) - Y(t));

partial_sum_is_random_variable:
  LEMMA random_variable?(LAMBDA t: sigma(0,n,LAMBDA n: XS(n)(t)))

distribution_function?(F:[real->probability]):bool
  = EXISTS X: FORALL x: F(x) = P(X <= x)
distribution_function: TYPE+ = (distribution_function?) CONTAINING
  (LAMBDA x: IF x < 0 THEN 0 ELSE 1 ENDIF)
distribution_function(X)(x):probability = P(X <= x)
F: VAR distribution_function

convergence_in_distribution?(XS,X):bool
  = FORALL x: continuous((distribution_function(X),x) IMPLIES
  convergence((LAMBDA n: distribution_function(XS(n))(x)),
  distribution_function(X)(x))

invert_distribution: LEMMA LET F = distribution_function(X) IN
  P(x < X) = 1 - F(x) % Lemma 2.1.11-a (G&S)
interval_distribution: LEMMA LET F = distribution_function(X) IN
  x <= y IMPLIES
  P(intersection(x < X, X <= y)) = F(y) - F(x) % Lemma 2.1.11-b (G&S)
limit_distribution: LEMMA LET F = distribution_function(X) IN
  P(X = x) = F(x) - limit(LAMBDA n: F(x-1/(n+1))) % Lemma 2.1.11-c (G&S)
distribution_0: LEMMA convergence(F o (lambda (n:nat): -n),0) % Lemma 2.1.6-a0 (G&S)
distribution_1: LEMMA convergence(F,1) % Lemma 2.1.6-a1 (G&S)
distribution_increasing: LEMMA increasing?(F) % Lemma 2.1.6-b (G&S)
distribution_right_continuous: LEMMA right_continuous(F) % Lemma 2.1.6-c (G&S)

END probability_space

Fig. 2. Abbreviated probability space file in PVS
from which we can calculate the probability

\[ P(x < X \leq y) = F(y) - F(x). \]

In the case where \( X \) is distributed \( U_{[0,1]} \), and because – for any \( f(x) \) with \( \int f = \mathcal{F} \) – we have

\[ \int_{-\infty}^{\infty} f(x)\chi_{[a,b]}(x)dx = \int_{-\infty}^{\infty} f(x) - F(a)\chi_{[a,b]}(x) + (F(b) - F(a))\chi_{[b,\infty]}(x). \]

We also observe that if \( X \) is distributed \( U_{[a,b]} \), then \( \mathbb{E}(X) = \frac{a+b}{2} \), and \( \text{Var}(X) = \frac{(a-b)^2}{12} \). So, with \( a = 0, b = 1 \) we get:

\[ \mu = \frac{1}{2}, \quad \sigma^2 = \frac{1}{12}. \]

**Definition 2:** If we have a sequence of continuous random variables \( \{X_n\} \), then we define their partial sums as a sequence of continuous random variables \( \{S_n\} \) with the property

\[ S_n = \sum_{i=1}^{n} X_i. \]

**Theorem 1:** If continuous random variables \( X \) and \( Y \) have joint probability density functions \( f \), then \( Z = X + Y \) has probability density function:

\[ f_Z(z) = \int_{-\infty}^{\infty} f(x, z-x)dx. \]

In the special case where \( X \) and \( Y \) are independent, then (because the joint probability density function \( f(x, y) \) can be expressed as the product \( f_X(x)f_Y(y) \)) we have the **Continuous Convolution Theorem**:

**Theorem 2:** If continuous random variables \( X \) and \( Y \) are independent and have probability density functions \( f_X \) and \( f_Y \) respectively, then \( Z = X + Y \) has probability density function:

\[ f_Z(z) = \int_{-\infty}^{\infty} f_X(x) f_Y(z-x)dx = \int_{-\infty}^{\infty} f_X(z-x) f_Y(x)dx. \]

**C. Continuous random vectors**

We want to generalize Theorem 2 in a probability space \( (T, \sigma, P) \) to a large number of random variables \( \{X_n\} \) to produce a bound for each intermediate sum in \( \{S_n\} \). We introduce random vectors.

**Definition 3:** A random vector of dimension \( n \) is a collection of \( n \) random variables

\[ X = (X_1, \cdots, X_n) \]

In this definition, the components need not to be independent variables meaning that the information on the distribution function of the components does not fully characterize the probabilistic behavior of random vector \( X \).

**Definition 4:** A random vector \( X \) has distribution function \( F \), if

\[ P(X_1 \leq x_1, \cdots, X_n \leq x_n) = F(x_1, \cdots, x_n). \]

Once we have defined the distribution function of a random vector we want to define it’s probability density.

**Definition 5:** A random vector \( X \) is continuous if its distribution function can be expressed as

\[ F(x_1, \cdots, x_n) = \int_{-\infty}^{x_1} \cdots \int_{-\infty}^{x_n} f(x_1, \cdots, x_n)dx_1 \cdots dx_n \]

for some \( \sigma \)-integrable function \( f: \mathbb{R}^n \rightarrow [0, \infty) \). We call the function \( f \) the probability density for random vector \( X \).
A random variable $X$ has **distribution function** $F$, if $\mathbb{P}(X \leq x) = F(x)$.

A random variable $X$ is **continuous** if its distribution function can be expressed as

$$F(x) = \int_{-\infty}^{x} f(x) \, dx$$

for some integrable function $f : \mathbb{R} \to [0, \infty)$. We call the function $f$ the **probability density function** for the random variable $X$.

We define the probability of “$A$ given $B$” (written $\mathbb{P}(A; B)$) as:

$$\mathbb{P}(A; B) = \frac{\mathbb{P}(A \cap B)}{\mathbb{P}(B)}$$

whenever $\mathbb{P}(B) > 0$.

A $\sigma$-**algebra** over a type $T$, is a subset of the power-set of $T$, which includes the empty set $\emptyset$, and is closed under the operations of complement, countable union and countable intersection.

A **Measurable Space** $(T, \sigma)$ is a set (or in PVS a type) $T$, and a $\sigma$-algebra over $T$.

A function $\mu : \sigma \rightarrow \mathbb{R}_{\geq 0}$ is a Measure over the $\sigma$-algebra $\sigma$, when $\mu(\emptyset) = 0$, and for a sequence of disjoint elements $\{E_n\}$ of $\sigma$:

$$\mu \left( \bigcup_{n=0}^{\infty} E_n \right) = \sum_{n=0}^{\infty} \mu(E_n).$$

A **Measure Space** $(T, \sigma, \mu)$ is a measurable space $(T, \sigma)$ equipped with a measure $\mu$.

A **Probability Space** $(T, \sigma, \mathbb{P})$ is a measure space $(T, \sigma, \mathbb{P})$ in which the measure $\mathbb{P}$ is finite for any set in $\sigma$, and in which:

$$\mathbb{P}(T^c) = 1 - \mathbb{P}(T).$$

If $(T_1, \sigma_1, \mathbb{P}_1)$ and $(T_2, \sigma_2, \mathbb{P}_2)$ are probability spaces then we can construct a **product probability space** $(T_3, \sigma_3, \mathbb{P}_3)$, where:

- $T_3 = T_1 \times T_2$
- $\sigma_3 = \sigma_1 \times \sigma_2$
- $\mathbb{P}_3(a, b) = \mathbb{P}_1(a) \mathbb{P}_2(b)$

where $\mathbb{P}_3$ is the extension of $\mathbb{P}_3$ that has the whole of $\sigma_3$ as its domain.

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We compute simple density functions through the following theorem that is proved by induction using the definition of independent random variables.

**Theorem 3:** A random vector $X$ of continuous random variables $(X_1, \ldots, X_n)$ with respective density functions $f_1, \ldots, f_n$ is continuous and has a density function

$$f(x_1, \ldots, x_n) = f_1(x_1) \times \cdots \times f_n(x_n)$$

if and only if $X_1, \ldots, X_n$ are independent.

During the proof, we show by Fubini’s theorem that for independent random variables

$$\mathbb{P}(X_1 \in A_1, \ldots, X_n \in A_n) = \mathbb{P}(X_1 \in A_1) \times \cdots \times \mathbb{P}(X_n \in A_n).$$

More complex density functions are handled through a smooth change of variables.

**Theorem 4:** Let $X = (X_1, \ldots, X_n)$ be a continuous random vector with density function $f_X$ and $g$ a transformation from $\mathbb{R}^n$ to $\mathbb{R}^n$ to a random vector $Y = (Y_1, \ldots, Y_n)$. Provided that

- $g$ is continuously differentiable on an open subset $U$ of $\mathbb{R}^n$,
- the Jacobian matrix

$$J_g = \left[ \frac{\partial g_i}{\partial x_j} \right]_{1 \leq i, j \leq n}$$

is never singular,
- $g$ admits an inverse on $U$ with the same properties, in particular its Jacobian matrix is never singular and it is the inverse of the Jacobian matrix of $g$,
- and finally $P(X \in U) = 1$

then $Y$ is a continuous random vector and with density function

$$f_Y(y) = f_X(g^{-1}(y)) \left| \text{det} J_g(g^{-1}(y)) \right|^{-1} \chi_U(y).$$

Notice that we obtain $\{S_n\}$ from $\{X_n\}$ through the following transformation:

$$g(x_1, \ldots, x_n) = \left( x_1, x_1 + x_2, \ldots, \sum_{i=1}^{n} x_i \right).$$

Its inverse $g^{-1}$ on $U = \mathbb{R}^n$ is

$$g(y_1, \ldots, y_n) = \left( y_1, y_2 - y_1, \ldots, y_n - \sum_{i=1}^{n-1} y_i \right),$$

its Jacobian matrix is an upper triangular matrix with all components equal to 1 and its determinant is also equal to 1. Finally $V = \mathbb{R}^n$.

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### IV. Corners of Hypercubes

What we are actually interested in is whether a series of calculations might accumulate a sufficiently large error to become meaningless. In the language we have developed, we are computing the probability that a sequence of $n$ calculations has failed because it has exceeded the error-bound somewhere.

$$\mathbb{P} \left( \max_{1 \leq i \leq n} (|S_i|) \geq \epsilon \right)$$
A. Sums of Arbitrary Continuous Uniform Random Variables

For a little while, we assume that each random variable $X_i$ is uniform on a given interval $[-\lambda_i, \lambda_i]$. Theorems 3 and 4 imply that the probability of failure is given by the ratio

$$
P \left( \max_{1 \leq i \leq n} (|S_i|) \geq \epsilon \right) = \frac{V \left( P_n^c \cap H_n \right)}{V(H_n)} = 1 - 2^{-n}V(S_n)
$$

where $V$ is the volume operator applied to the following sets

$$
P_n = \left\{ (x_1, \ldots, x_n) \mid \forall i \ |x_i| \leq 1 \land \sum_{j=1}^{i} \lambda_j x_j \leq \epsilon \right\},
$$

$$
H_n = \left\{ (x_1, \ldots, x_n) \mid \forall i \ |x_i| \leq 1 \right\},
$$

and $P_n^c$ is the complement of polyhedron $P_n$.

There is no direct formula to compute $V(P_n)$ due to linear dependencies between the constraints.

$$
\left\{ \begin{array}{l}
|x_i| \leq 1 \\
\sum_{j=1}^{i} \lambda_j x_j \leq \epsilon
\end{array} \right.
$$

The authors of this work and the authors of [20] agree that small or elementary perturbations are certainly the key to compute accurately $V(P_n)$ for small values of $n$. Yet perturbation are not amenable for large values of $n$.

So we have to revert to the bound $V(P_n)$ using the following sets with only one constraint which denotes no failure only at iteration $n$

$$
P_t(\epsilon) = \left\{ (x_1, \ldots, x_i) \mid \forall j \ |x_j| \leq 1 \land \sum_{j=1}^{i} \lambda_j x_j \leq \epsilon \right\}.
$$

We could have used

$$
P_n = \bigcap_{i=1}^{n} P_t(\epsilon) \times H_n - i
$$

where $P_t(\epsilon) \times H_n - i$ is the Cartesian product of sets but we obtain a better bound with

$$
P_n = \bigcap_{i=1}^{n} \left( P_t(\epsilon) \cup P_t^c(\epsilon + \lambda_i) \times H_1 \right) \times H_n - i
$$

that is easily proved since

$$
P_{t-1}(\epsilon) \subset P_{t-1}(\epsilon + \lambda_i)
$$

with the convention that $P_0 = \{\}$. Using property of the complement and volume operators, we deduce that

$$
V \left( P_n^c \cap H_n \right) \leq \sum_{i=1}^{n} 2^{n-i} \left( 2V(P_{t-1}(\epsilon + \lambda_i)) - V(P_t(\epsilon)) \right)
$$

The approximation is reduced due to the fact that

$$
P_t(\epsilon) \subset P_{t-1}(\epsilon + \lambda_i) \times H_1
$$

and so the volume of the set difference is the difference of the volumes of the sets with the convention that $V(\{\}) = 1$.

In the rest of the text, we heavily use notations and results of [20] to obtain $V(P_t(n, \mu))$ for appropriate $n$ and $\mu$ (we change summation index from $i$ to $n$ for this reason). For a given vector

$$
A = (a_1, a_2, \ldots, a_n),
$$

where $a_j = \lambda_j / \mu$, we define the $n \times (n+1)$ matrix $S = (I_n, A)$ and we deduce $1 \times (n+1)$ matrix

$$
P = (1, a_1, a_2 \ldots a_n).
$$

We want to compute $V(P_t(n, \mu)) = \nu(P) = \frac{2^n}{n!} \sigma(S)$ and we obtain $\sigma(S)$ by a formula that require the definition of

- $m = 1$
- $i = \{\kappa\}$ where $\kappa = (1)$, $S_{\kappa} = (1)$ and $det(S_{\kappa}) = 1$
- $\kappa^c = (2, 3, \ldots (n + 1))$, $det(\kappa^c) = a_j - 1$ and

$$
\alpha_{\kappa} = \frac{n}{\prod_{j=1}^{n} a_j^{-1}}
$$

- for $\gamma \in \Gamma$,

$$
(s_{\kappa} \gamma ; 1) = s_{\gamma} = 1 + \sum_{j=1}^{n} \gamma_j a_j,
$$

The formula becomes

$$
\nu(P) = \frac{1}{n!} \prod_{\gamma \in \Gamma} \left( a_j \gamma \right)^{-1} \epsilon_{\gamma} s_{\gamma}^n \text{sgn}(s_{\gamma})
$$

where all the quantities have been instantiacted and the cardinal of $\Gamma$ is $2^n$ (remember that $n \leq 2^{25}$).

B. Sums of Independent Identical Continuous Uniformly Distributed Random Variables

In the rest of this section we assume that all the $a_j$ are equal to $a$ (and all the $\lambda_i$ are equal to $\lambda$). For a $\gamma \in \Gamma$, we define $n_{\gamma}$ as the number of positions where $\gamma_j = -1$ for $j$ between $2$ and $n + 1$.

The formula becomes

$$
\nu(P) = \frac{1}{a^n n!} \sum_{\gamma \in \Gamma} (-1)^{n_{\gamma}} (1 + (n - 2n_{\gamma})a)^n \text{sgn}(1 + (n - 2n_{\gamma})a),
$$

or equivalently

$$
\nu(P) = \frac{1}{n!} \sum_{\gamma \in \Gamma} (-1)^{n_{\gamma}} (a^{-1} + n - 2a_{\gamma})^n \text{sgn}(a^{-1} + n - 2a_{\gamma}),
$$

and we change the summation index $p = n - n_{\gamma}$ or $n - 2n_{\gamma} = 2p - n$ to obtain

$$
\frac{1}{n!} \sum_{p=0}^{n} \binom{n}{p} (-1)^p (n - a^{-1} - 2p)^n \text{sgn}(a^{-1} - n + 2p).
$$

with

$$
\text{sgn}(a^{-1} - n + 2p) = \left\{ \begin{array}{ll}
1 & \text{if} \ 2p > n - a^{-1} \\
0 & \text{if} \ 2p = n - a^{-1} \\
-1 & \text{if} \ 2p < n - a^{-1}
\end{array} \right.
$$
We observe that $\nu$ is trivial when $n\alpha < 1$ and so we consider only the case where $n - \alpha^{-1} \geq 0$. When $n = \alpha^{-1}$, we obtain the volume of the full cube

$$\frac{1}{n!} \sum_{p=1}^{n} \binom{n}{p} (-1)^p (-2p)^n = 2^n$$

deriving a known formula [24].

So the probability of failure is bounded by

$$B_n(\epsilon, \lambda) = \sum_{i=1}^{n} 2^{-i} \left(2V_{i-1}\left(\frac{\epsilon}{\lambda} + 1\right) - V_i\left(\frac{\epsilon}{\lambda}\right)\right)$$

with

$$V_i(\eta) = \frac{1}{i!} \sum_{p=0}^{i} \binom{i}{p} (-1)^p (i - \eta - 2p) \text{sgn}(\eta - i + 2p)$$

for $\eta \geq i$ and $V_i(\eta) = 2^i$ otherwise.

We obtain the equivalent formula below starting at the first index where $i > \epsilon/\lambda$

$$B_n(\epsilon, \lambda) = 1 - 2^{-n} V_n\left(\frac{\epsilon}{\lambda}\right) + \sum_{i=n+1}^{\infty} 2^{-i} \left(V_i\left(\frac{\epsilon}{\lambda} + 1\right) - V_i\left(\frac{\epsilon}{\lambda}\right)\right)$$

V. Future work

At the time that we are submitting this work, results of Sections III-C and IV-A are not fully certified using PVS proof assistant. But we anticipate no problem has these results are generalizations to vector of our earlier work [12] and set theory except that our results are based on a formula obtained through Fourier Analysis. As Fourier Analysis has not been implemented in any proof checking environment, we are using the formula as a assumption to our theory. We hope that Fourier Analysis will be available in the future so we can remove any assumption from the theory we have developed. Yet certifying the results of [24] will be challenging.

This work will be continued in two directions. The first direction is to modify Fluctuat to generate theorems that can be checked automatically by PVS using ProofLite as proposed in [3, 4]. This work will be carried in collaboration with the developers of Fluctuat within the EVA-Flo project of the ANR. The software will conservatively estimate the final effect of the errors introduced by each individual operations and compute upper bounds of their magnitudes.

The second direction is to develop and check accurate proofs the errors of individual operations. A uniformly distributed random variable whose variance depends only on the operation and the computed result might provide a too pessimistic bound. For example the floating point addition of a large number with a small number absorbs the small number meaning that the round-off error may be far below half an ulp of the computed result.

Two’s complement operation of the TMS320 circuit can either round or truncate the result. If truncation is used, it introduces a drift and our development cannot be used. Should we wish to extend this work to account for drifts (non-zero means for the random variables $\{X_n\}$), we will also have to consider higher-order error terms that also introduce a drift.

This library and future work will be included into NASA Langley PVS library as soon as it becomes stable.

VI. Conclusions

To the best of our knowledge this paper presents the first application of the volume of the corner of a $n$-dimension hypercube to software reliability with $n$ large enough for exponential-time solution to be impractical. In addition, we are finishing certification of our results with PVS. The major restriction lies in the fact that we have been forced to insist that individual errors have no drift, and are independent. Notice that even with a high tolerance of error, and with independent errors, we will still eventually fail. Our results permit the development of safe upper limits on the number of operations that a piece of numeric software should be permitted to undertake.

It is worth pointing out that violating our assumptions (independence of errors, and zero drift) would lead to worse results, so one should treat the limits we have deduced with caution, should these assumptions not be met.

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