OpenTURNS and its graphical interface

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Abstract

OpenTURNS is an open source library for uncertainty propagation by probabilistic methods. Developed in a partnership of five industrial companies (EDF, Airbus, Phimeca, IMACS and ONERA), it benefits from a strong practical feedback. Classical algorithms of UQ are available: central dispersion, probability of exceedance, sensitivity analysis, metamodels and stochastic processes. Developed in C++, OpenTURNS is also available as a Python module and has gained maturity thanks to more than 10 years of development.

However, there are situations where the engineer in charge of performing an uncertainty study does not want to use a programming language such as C++ or Python. In this context, providing a graphical user interface (GUI) may allow to greatly increase the use of OpenTURNS and, more generally, of the UQ methodology.

In this paper, we present a basic tutorial of OpenTURNS in Python and will review the new features in the library, which include new incremental statistical estimators. In the second part, we review the new features in the open source GUI will be presented, including the management of stochastic processes.

Contents

1	Introduction	2
2	OpenTurns	2
	A tutorial example: the flooding model 3.1 Introduction	
	Estimating the mean with an incremental algorithm 4.1 Theory	

5	Estimate sensitivity indices with an incremental algorithm						
	5.1	Theory	7				
		5.1.1 Overview	8				
		5.1.2 Asymptotic distribution	8				
		5.1.3 Stopping criteria	9				
	5.2	Tutorial	9				
6	Nev	v features in the graphical user interface	10				
	6.1	Introduction	10				
	6.2	Dependency structures	11				
	6.3	Screening with the Morris method	11				
	6.4	Perspectives: one-dimensional stochastic processes	12				

1 Introduction

OpenTURNS is a C++ library for uncertainty propagation by probabilistic methods. OpenTURNS is also available as a Python module and has gained maturity thanks to more than 10 years of development. However, there are situations where the engineer in charge of performing an uncertainty study does not want to use a programming language such as C++, Python (e.g. OpenTURNS) or Matlab. In this context, providing a graphical user interface (GUI) may allow to increase the use of OpenTURNS and, more generally, of the UQ methodology.

2 OpenTurns

OpenTURNS[2, 5, 3] is an open source software, available as a C++ library and a Python interface. It works under the Linux and Windows environments. The key features of OpenTURNS are the following:

- open source initiative to secure the transparency of the approach,
- generic to the physical or industrial domains for treating of multi-physical problems,
- high performance computing,
- includes a variety of qualified algorithms in order to manage uncertainties in several situations.
- contains complete documentation (Reference Guide, Use Cases Guide, User manual, Examples Guide, and Developers' Guide).

OpenTURNS is available under the LGPL license.

The main features of OpenTURNS are uncertainty quantification, uncertainty propagation, sensitivity analysis and metamodeling.

Moreover generic wrappers allows to link OpenTURNS to any external code G.

OpenTURNS can be downloaded from its dedicated website www.openturns.org which offers different pre-compiled packages specific to several Windows and Linux environments. It is also possible to download the source files from the SourceForge server and to compile them within another environment: the OpenTURNS Developer's Guide provides advices to help compiling the source files.

3 A tutorial example: the flooding model

3.1 Introduction

In this paper, we illustrate our discussion with a simple application model that simulates the height of a river. The figure 1 presents the dyke that protects industrial facilities. When the

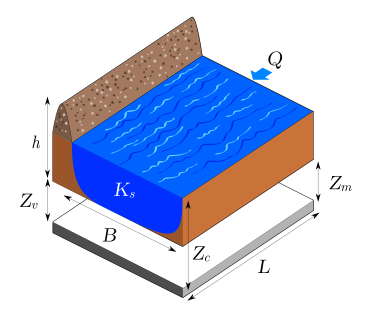


Figure 1: The flood example: simplified model of a river.

river height exceeds the one of the dyke, flooding occurs. This academic model is used as a pedagogical example in [7]. The model is based on a crude simplification of the 1D hydrodynamical equations of SaintVenant under the assumptions of uniform and constant flowrate and large rectangular sections. It consists of an equation that involves the characteristics of the river stretch:

$$S = Z_v + H \quad \text{with} \quad H = \left(\frac{Q}{BK_s\sqrt{\frac{Z_m - Z_v}{L}}}\right)^{0.6},\tag{1}$$

where S is the maximal annual overflow, H is the maximal annual height of the river, B is the river width and L is the length of the river stretch. In this paper, we set the values of L and B parameters :

$$L = 5000$$
 $B = 300$.

The other four input variables Q, K_s, Z_v and Z_m are defined in Table 1 with their probability distribution. The randomness of these variables is due to their spatio-temporal variability, our ignorance of their true value or some inaccuracies of their estimation. We make the hypothesis that the input variables are independent.

Input	Description	Unit	Probability distribution
\overline{Q}	Maximal annual flowrate	m^3/s	Gumbel $\mathcal{G}(scale = 558, mode = 1013)$
K_s	Strickler coefficient	-	Normal $\mathcal{N}(30, 7.5)$
Z_v	River downstream level	\mathbf{m}	Uniform $\mathcal{U}(49,51)$
Z_m	River upstream level	\mathbf{m}	Uniform $\mathcal{U}(54,56)$

Table 1: Input variables of the flood model and their probability distributions.

The goal of this study is twofold:

• we want to estimate the mean river height E(S),

• we want to perform the sensitivity analysis of the model, i.e. we want to rank the inputs Q, K_s , Z_v and Z_m with respect to their contributions to the variability of the output S.

3.2 Define the random vector

In this section, we present the Python script which allows to define the output random vector in OpenTURNS.

We begin by importing the required modules.

```
from openturns.viewer import View
import openturns as ot
from math import sqrt
import pylab as pl
```

We first define the function through which we want to propagate the uncertainties with the def operator.

```
def functionFlood(X) :
    Hd = 3.0
    Zb = 55.5
    L = 5.0e3
    B = 300.0
    Zd = Zb + Hd
    Q, Ks, Zv, Zm = X
    alpha = (Zm - Zv)/L
    H = (Q/(Ks*B*sqrt(alpha)))**(3.0/5.0)
    S = H + Zv
    return [S]
```

Then we convert this Python function into an OpenTURNS function with the Python-Function class.

```
\begin{array}{l} input\_dimension \, = \, 4 \\ g \, = \, ot \, . \, PythonFunction (input\_dimension \, , \, \, 1 \, , \, \, functionFlood ) \end{array}
```

Now we create the distributions for the input variables.

- ullet There are several ways to set the parameters of the Gumbel distribution for the Q variable. Here the parameters are defined with the scale and mode parameters, which corresponds to the GumbelAB class.
- The Q and K_s variables must remain positive (a negative value is not compatible with the physical model). For this reason, we must truncate the distribution with TruncatedDistribution.

```
\label{eq:myParam} \begin{split} & myParam = ot.GumbelAB(1013., 558.)\\ & Q = ot.ParametrizedDistribution(myParam)\\ & otLOW = ot.TruncatedDistribution.LOWER\\ & Q = ot.TruncatedDistribution(Q, 0, otLOW)\\ & Ks = ot.Normal(30.0, 7.5)\\ & Ks = ot.TruncatedDistribution(Ks, 0, otLOW)\\ & Zv = ot.Uniform(49.0, 51.0)\\ & Zm = ot.Uniform(54.0, 56.0) \end{split}
```

We set the descriptions of the random variables: they are used for the graphics.

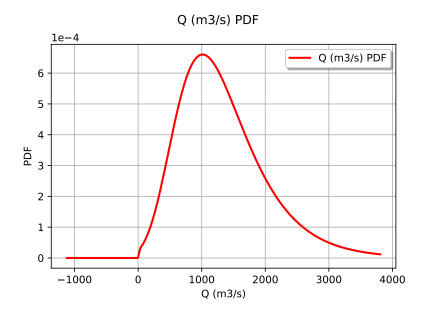


Figure 2: The probability density function of the variable Q.

```
Q. setDescription (["$Q_{\sqcup}(m^3/s)$"]) Ks. setDescription (["$Ks_{\sqcup}(m^{1/3})/s)$"]) Zv. setDescription (["Zv_{\sqcup}(m)"]) Zm. setDescription (["Zm_{\sqcup}(m)"])
```

The drawPDF method plots the the probability distribution function of the variable.

```
Q. drawPDF()
```

The previous session produces the figure 2. When we closely look at the PDF of Q, we see a small increase of the density for Q = 0, because of the truncation of the distribution.

Then we create the input random vector **inputvector**: by default, the copula is independent. Finally, we create the output random vector **S**.

```
\begin{array}{lll} X = ot.ComposedDistribution([Q, Ks, Zv, Zm]) \\ inputRV = ot.RandomVector(X) \\ S = ot.RandomVector(g, inputRV) \end{array}
```

These steps are typical of *probabilistic programming*: we have defined the random variables involved in the problem *without* having generating a sample so far.

4 Estimating the mean with an incremental algorithm

4.1 Theory

In this section, we present the principles that are used in a new incremental algorithm in OpenTURNS 1.12; the goal of this algorithm is to estimate the mean of a random variable.

Assume that the output $Y \in \mathbb{R}^{n_Y}$ is a random vector and that we want to estimate the mean $E(Y_i)$ for $i = 1, ..., n_Y$.

The Monte Carlo algorithm is based on the the sample mean:

$$\mu_i = \frac{1}{n} \sum_{j=1}^n Y_j^{(i)}$$

for $i = 1, ..., n_Y$ where n is the sample size and $Y_j^{(i)}$ are i.i.d. outcomes of the random output.

The algorithm is based on the fact that the sample mean is asymptotically gaussian:

$$\mu_i \sim \mathcal{N}\left(E(Y_i), \frac{V(Y_i)}{n}\right).$$

for $i = 1, ..., n_Y$ where $V(Y_i)$ is the variance of the i-th output and n is the sample size.

In general, most users set the sample size n in advance and estimate the precision afterwards. Let s_i be the (unbiased) sample standard deviation of the output Y_i :

$$s_i = \sqrt{\frac{1}{n-1} \sum_{j=1}^{N} (y_j^{(i)} - \mu_i)^2}$$

for $i = 1, ..., n_Y$. The absolute precision of the estimate μ_i can be estimated based on the sample standard deviation of the estimator:

$$\sigma_i = \frac{s_i}{\sqrt{n}}$$

for $i=1,...,n_Y$. If $\mu_i \neq 0$ and if $E(Y_i) \neq 0$, the relative precision can be estimated based on the coefficient of variation σ_i/μ_i for $i=1,...,n_Y$.

Instead, suppose that we set the absolute precision in advance and wish to determine the smallest sample size n that achieves this precision. If the variance $V(Y_i)$ is known (which rarely happens in practice), we can set the value of n so that the standard deviation $\sqrt{V(Y_i)}$ is small enough. If we want to set the relative precision, we can consider the coefficient of variation of the estimator $\frac{\sqrt{V(Y_i)}}{E(Y_i)\sqrt{n}}$ as a criterion (if $E(Y_i) \neq 0$). However, we generally do not know the values of neither $E(Y_i)$ nor $V(Y_i)$.

The purpose of the algorithm is to increase the sample size n incrementally until a stopping criteria is met. At each iteration, we approximate the values of $E(Y_i)$ and $V(Y_i)$ by their empirical estimators.

In order to get the best possible performance on distributed supercomputers and multicore workstations, the size of the sample increases by block. For exemple, if the block size is equal to 100, then the sample size will be equal to 0, 100, 200, etc... On each block, the evaluation of the outputs can be parallelized, which allows to improve the performance of the algorithm.

Since there are in general several outputs, i.e. $n_Y \ge 1$, we use a stopping criteria which is based on a operator. There are three mathematical stopping criteria available:

- through an operator on the coefficient of variation $\frac{\sigma_i}{\mu_i}$ (a relative criterion)
- through an operator on the standard deviation σ_i (absolute criterion)
- on the maximum standard deviation per component: $\sigma_i \leq \max_{i=1,...,n_Y} \sigma_i$ (absolute criterion)

By default, the maximum coefficient of variation is used, i.e. the operator is the maximum so that the algorithm stops when:

$$\max_{i=1,\dots,n_Y} \frac{\sigma_i}{\mu_i} \le \max_{COV}.$$

4.2 Tutorial

In this section, we present how to use the ExpectationSimulationAlgorithm class in the tutorial flooding example. We set the maximum number of iterations with the setMaximum-OuterSampling so that we use at most 1000 iterations. In order to evaluate the function with blocks of size 10, we use the setBlockSize. In this simulation, we use a relative stopping criteria and configure the maximum coefficient of variation to be equal to 0.001.

```
\label{eq:algo} \begin{array}{ll} algo \ = \ ot. \ Expectation Simulation Algorithm (S) \\ algo \ . \ set Maximum Outer Sampling (1000) \\ algo \ . \ set Block Size (10) \\ algo \ . \ set Maximum Coefficient Of Variation (0.001) \end{array}
```

The computationnaly intensive part of the simulation is associated with the run method.

```
algo.run()
```

Once the simulation is done, the getResult method allows to access to the results.

```
\label{eq:continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous_continuous
```

The previous session prints the following output.

```
Mean = 52.520729
Number of calls to G = 500
Coef. of var.=0.000994
```

The estimate of the mean has a known asymptotical gaussian distribution, which can be retrieved with the getExpectationDistribution method.

```
\begin{array}{ll} expectation Distribution = result.getExpectation Distribution () \\ \textbf{print} (expectation Distribution) \\ View (expectation Distribution .drawPDF()) \end{array}
```

We emphasize that the output of the getExpectationDistribution method is a Distribution in the OpenTURNS sense: the whole information is available, not just a part of it, making the output as programmatically meaningful as possible.

```
expectationDistribution = result.getExpectationDistribution()
expectationDistribution.drawPDF()
```

The previous script produces the figure 3. The figure shows that we have an accurate estimate of the mean, up to approximately 2 significant digits.

5 Estimate sensitivity indices with an incremental algorithm

5.1 Theory

In this section, we present the principles that are used in a new incremental algorithm in OpenTURNS 1.12 which computes the Sobol' sensitivity indices.

In [8] the authors derive a new method to estimate the Sobol' sensitivity indices; one of the advantages of the new estimator is that it is associated with an asymptotic distribution, which is derived thanks to the so called "delta"-method. Based on a suggestion by R.Lebrun, A. Dumas [4] used the same theoretical method in order to derive the asymptotic distribution of Sobol' sensitivity indices already available in OpenTURNS.

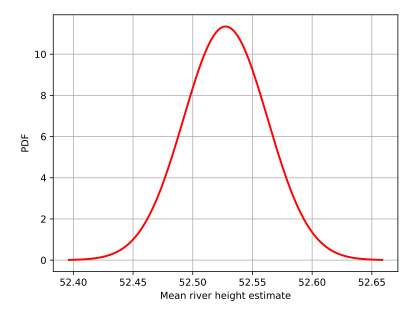


Figure 3: The probability density function of the estimate of the mean of the river height.

5.1.1 Overview

Let us denote by $X \in \mathbb{R}^{n_X}$ a random vector. This algorithm works in the general case where the output $Y \in \mathbb{R}^{n_Y}$ is a random vector: in this case it operates on aggregated indices. In order to simplify the discussion, let us make the hypothesis that $n_Y = 1$.

The Sobol' first order sensitivity indices are defined by

$$S_i = \frac{V(E(Y|X_i))}{V(Y)}$$

for $i=1,...,n_X$. The total order sensitivity indices are defined by :

$$T_i = 1 - \frac{V\left(E\left(Y|X_{-i}\right)\right)}{V(Y)}$$

where -i is the set of indices which are different from i. In the remaining of this section, we focus on the first order sensitivity indice and let the reader consider [1] for the total order indices. Moreover, the derivation is the same for all input variables so that we omit the indice i in order to simplify the notations.

5.1.2 Asymptotic distribution

The algorithm is based on the fact that the estimators of the first and total order Sobol' sensitivity indices asymptotically have the gaussian distribution. This gaussian distribution can be derived from the so called "delta"-method.

Assume that the Sobol' estimator is

$$\overline{S} = \Psi\left(\overline{U}\right)$$

where Ψ is a multivariate function, U is a multivariate sample and \overline{U} is its sample mean. Each Sobol' estimator can be associated with a dedicated choice of function Ψ and vector U.

Let us denote by Φ_j^F (resp. Φ_j^T) the cumulated distribution function of the gaussian distribution of the first (resp. total) order sensitivity indice of the j-th input variable.

Each available estimator in the library provides its own distribution, namely the Saltelli, Mauntz-Kucherenko, Jansen and Martinez estimators.

5.1.3 Stopping criteria

We set $\alpha \in [0,1]$ the level of the confidence interval and $\epsilon \in (0,1]$ the length of the confidence interval. The algorithms stops when, on all components, one of the two following conditions are satisfied:

- first and total order indices haved been estimated with enough precision or
- the first order indices are separable from the total order indices.

The precision is said to be sufficient if the $1-2\alpha$ confidence interval is smaller than ϵ :

$$(\Phi_{j}^{F})^{-1}(1-\alpha) - (\Phi_{j}^{F})^{-1}(\alpha) \le \epsilon$$

and

$$(\Phi_{i}^{T})^{-1}(1-\alpha) - (\Phi_{i}^{T})^{-1}(\alpha) \le \epsilon$$

for $j = 1, ..., n_X$. The first order indices are separable from the total order indices if

$$\Phi_i^F(1-\alpha) \leq \Phi_i^T(\alpha)$$

for $j = 1, ..., n_X$. This criteria allows to stop when the algorithm has detected an interaction between input variables with sufficient precision.

5.2 Tutorial

In this section, we present how to use the SaltelliSensitivityAlgorithm classe in the tutorial flooding example.

We first set the parameters of the algoritms. We set the alpha variable so that a 90% confidence interval is used. In order to get confidence intervals which are not greater than 0.2, we set the variable epsilon variable accordingly. The block size corresponds to the size of the Sobol' design of experiment generated at each iteration. Finally, the batchsize variable contains the number of points evaluated simultaneously by the model.

Then we create the algorithm and configure the algorithm so that it uses the previous variables. Moreover, we configure the algorithm so that at most 100 iterations are used.

```
estimator = ot.SaltelliSensitivityAlgorithm()
estimator.setUseAsymptoticDistribution(True)
algo = ot.SobolSimulationAlgorithm(X, g, estimator)
algo.setMaximumOuterSampling(100) # number of iterations
algo.setBlockSize(blocksize)
algo.setBatchSize(batchsize)
algo.setIndexQuantileLevel(alpha) # alpha
algo.setIndexQuantileEpsilon(epsilon) # epsilon
algo.run()
```

Once that the algorithm has run, the results can be retrieved and estimates of first and total order indices can be printed.

```
 \begin{array}{lll} result &= algo.getResult() \\ fo &= result.getFirstOrderIndicesEstimate() \\ to &= result.getTotalOrderIndicesEstimate() \\ print("First\_order\_=\_\%s" \% (str(fo))) \\ print("Total\_order\_=_L\%s" \% (str(to))) \\ \end{array}
```

The previous script produces the following output.

```
First order = [0.529059, 0.205321, 0.381215, 0.0316592]
Total order = [0.481355, 0.130565, 0.362292, 0.011206]
```

We can obtain the asymptotic distribution of the first and total order indices. For example, the following script extracts the first component of the asymptotic distribution of the first order indice (which corresponds to the variable Q) and plots it.

```
dist_fo = result.getFirstOrderIndicesDistribution()
dist_fo_i = dist_fo.getMarginal(0)
graph = dist_fo_i.drawPDF()
graph.setTitle("S0")
graph.setXTitle("S0")
```

The previous script produces the figure 4.

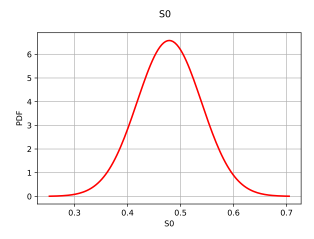


Figure 4: Asymptotic distribution of the first order Sobol' indices for the Q variable.

In order to get a more compact view of the first and total order indices along with their confidence intervals, we often represent the 90% confidence intervals with a vertical bar. The figure 5 presents the Sobol' indices with asymptotic confidence intervals. We observe that the the confidence intervals are relatively small, as expected.

6 New features in the graphical user interface

6.1 Introduction

There are situations where the engineer in charge of performing an uncertainty study does not want to use a programming language such as C++ or Python. In this context, providing

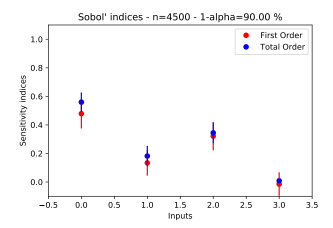


Figure 5: Sobol' indices with asymptotic confidence intervals.

a graphical user interface (GUI) may allow to greatly increase the use of OpenTURNS and, more generally, of the UQ methodology.

This is why we develop since 2016 a graphical user interface (GUI) of OpenTURNS, which is integrated within SALOME [6]. This GUI is developed with the OpenSource LGPL license, which is the same as OpenTURNS and SALOME. SALOME binaries for the Linux platform are provided at the following URL:

https://www.salome-platform.org/contributions/edf_products

Details on the main features and the internal architecture of the GUI were already presented in [3], this is why this paper focuses on the new features.

6.2 Dependency structures

The GUI allows to define advanced dependency structures, based on copulas. The figure 6 presents the dialog box in which the copulas can be selected and configured.

The principle is to create sub-groups within the input variables. Within a given sub-group, we can select the copula and configure its parameters. Seven copulas are available: independent, Gaussian, Ali-Mikhail-Hak, Clayton, Farlie-Gumbel-Morgensten, Frank or an inference result.

For example, the figure 6 considers the situation in which the model has five inputs named X0, X1, X2, X3 and X4. In this particular model, the sub-group [X0,X1] is associated with the Gaussian copula while the sub-group [X3,X4] is associated with the Gumbel copula. The variable X2 remains independent from the others in this model.

Moreover, any multivariate sample can be used to estimate the parameters of a copula. This is case, the results of an inference can be reused in a dependency model.

6.3 Screening with the Morris method

The qualitative sensitivity analysis based on Morris's method [10] aims at selecting the significant input variables in a costly computer code which may have a large number of inputs. The GUI performs the screening analysis based on the OpenTURNS otmorris module [1]. If we note e_i^k the k-th computed elementary effects associated to the i-th input marginal, we get μ_i^* and σ_i respectively the absolute mean and the standard deviation of

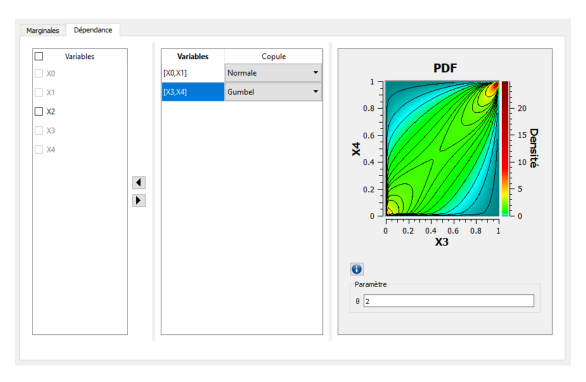


Figure 6: Managing a copula in the OpenTURNS GUI.

the elementary effects. The goal of this method is to spread the inputs variables into three classes, based on μ_i^* and the $\rho_i = \frac{\mu_i^*}{\sigma_i}$ factors:

- 1. if μ_i^* is close to zero, the i-th variable has no effect,
- 2. if $\rho_i \leq 0.5$ the i-th variable has almost linear effects,
- 3. if $\rho_i \geq 1$ the i-th variable has non-linear and non-monotonic effects

The figure 7 presents the dialog box which contains the parameters of the algorithm. The user can set the number of trajectories and the number of levels for each variable. The dialog box automatically computes the corresponding number of simulations and prints it in the bottom of the dialog box.

Once the simulations are performed, the figure 8 presents the results. The main figure presents the mean and standard deviations of the elementary effects. A table (not shown in the figure) containing the list of input variables allows to see in which category fall each variable. A default classification is done by the GUI, but can be modified by the user.

6.4 Perspectives: one-dimensional stochastic processes

In this section, we present the current developments of the GUI, which focuses on the management on stochastic fields.

Indeed, there are various situations in which the simulator through which we propagate the uncertainties produces a stochastic process. This happens for example in the case where the simulator produces a time series or a one-dimensional spatial field.

The figure 9 presents a set of trajectories in the GUI. In general, the sample size is large and this graphics does not convey much information, because the trajectories overlap and hide each other.

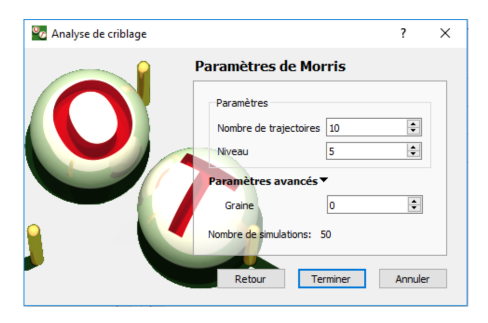


Figure 7: Performing screening with Morris's method in the GUI.

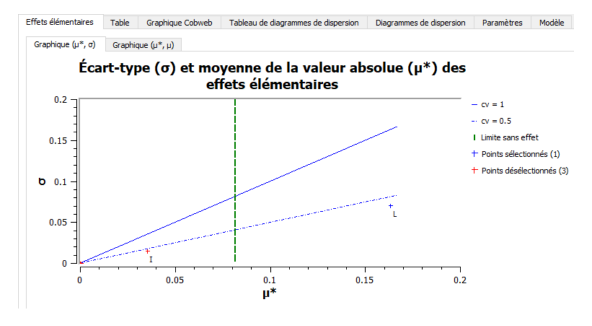


Figure 8: Results of the screening with Morris's method in the GUI.

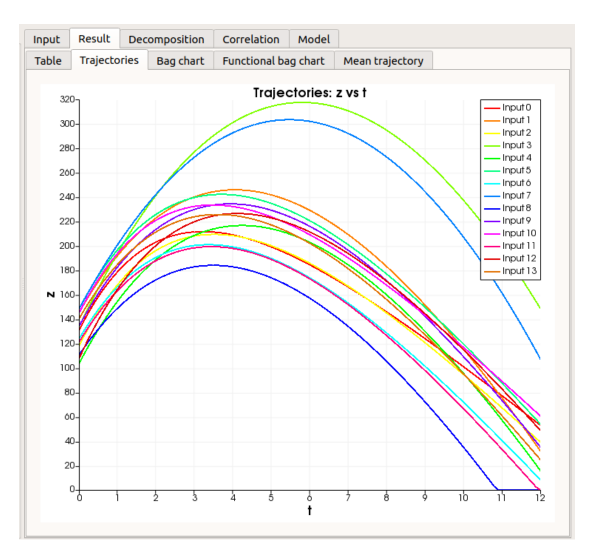


Figure 9: A sample set of trajectories in the GUI.

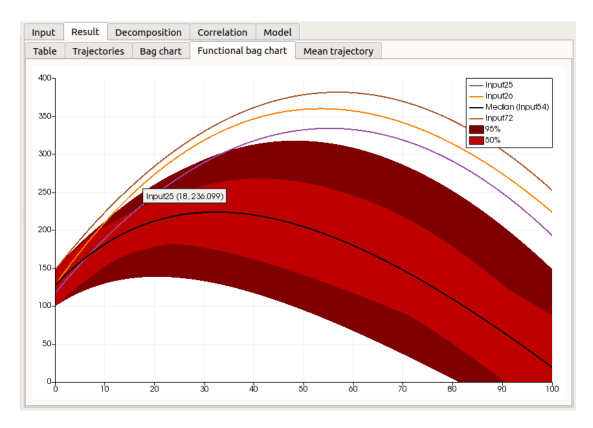


Figure 10: The functional bag chart of Paraview to plot a functional boxplot and detect outlier trajectories.

Obviously, this situation is more complex than the classical output random vector that many engineers are used to and require more advanced probabilistic methods. The most common way of managing such situation is to use a dimension reduction method such as the principal component analysis or the Karhunen-Loève decomposition.

This is why Ribes et al [11] developed a new visualization tool in [9], based on a work by Kitware. This tool is the *functional bag chart*, also known as the highest density region plot in the bibliography. The figure 10 presents the functional bag chart of a sample set of trajectories. This graphics allows to plot a functional boxplot in the sense that it plots a functional 95% confidence region. The graphics also allows to detect outlier trajectories, i.e. trajectories which achieve a low density in the reduced space.

The future version will extend these functional analyses to higher dimensions, including 2D stochastic fields.

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