Efficient computation of Sobol' sensitivity indices based on ranking: R code

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1 Introduction

Global Sensitivity Analysis (SA) is the study of how the uncertainty in a given model output is apportioned to the uncertainty in the model inputs (Sobol' and Kucherenko 2005; A. Saltelli 2002b). Over the last two decades, Sobol' variance-based sensitivity indices have become a method of choice for practitioners. These methods typically require a large number of function evaluations to achieve acceptable convergence and can become impractical for large engineering problems. Although the efficiency of the computation of first-order effects (S_i) has improved significantly (A. Saltelli 2002a; Sobol' and Myshetskaya 2008; Owen 2013; Kucherenko and Song 2017), fewer advances have been made in the case of total-order (T_i) indices (Sobol' 2001; Saltelli et al. 2010; Kucherenko et al. 2015; Lo Piano et al. 2020).

Here we propose to increase the efficiency of the computation of T_i by monitoring their ranking. Being discrete variables, ranks may converge significantly quicker than the very sensitivity indices. Our approach is inspired by the ideas presented by Kreinin and Iscoe (2018), and also takes advantage of the fact that T_i converges faster than S_i : this is because the estimate for S_i depends upon elementary effects, related to output function values f as products $ff' - f_0^2$, where only one factor X_i is kept fixed between f and f', and f_0 is the sample output mean. Instead, for the estimation of the total sensitivity index T_i , we use differences f - f', but in this case between f and f' all values are fixed but one (Saltelli et al. 2010).

An elementary effect for S_i is therefore hardly ever zero, as f and f' are generally different, and many f and f' need to be averaged to show that, once f_0^2 is substracted, the effect is small or zero. However, an elementary effect for T_i is normally different from zero since only one factor has been shifted. Therefore, if there is a difference between f and f', this can only be attributed to the moved parameter X_i . If there is no difference when the factor is influential it is just because the step separating f from f' is too small or the step is across a point of non-monotonicity.

2 Materials and methods

Hereafter we present the R code upon which our paper is based.

2.1 Reproducibility

Let us first create a wrapper function that allows to load all the required R libraries in one go: data.table (Dowle and Srinivasan 2019), ggplot2 (Wickham 2016), sensobol (Puy 2019), scales (Wickham 2018), parallel (R Core Team 2019), cowplot (Wilke 2019), gridExtra(Auguie 2017), sensitivity (Iooss, Janon, and Pujol 2019), wesanderson (Ram and Wickham 2018) and RColorBrewer (Neuwirth 2014). We then load the package checkpoint (Microsoft Corporation 2018), which installs in a local directory the same package versions used in the study. This allows anyone that runs our code to fully reproduce our results anytime.

```
# LOAD PACKAGES ------
# Function to read in all required packages in one go:
loadPackages <- function(x) {
  for(i in x) {
   if(!require(i, character.only = TRUE)) {
    install.packages(i, dependencies = TRUE)</pre>
```

```
library(i, character.only = TRUE)
    }
 }
}
loadPackages(c("sensobol", "tidyverse", "parallel", "grid",
               "cowplot", "gridExtra", "sensitivity", "wesanderson", "RColorBrewer",
               "tikzDevice", "data.table"))
# SET CHECKPOINT -----
dir.create(".checkpoint")
library("checkpoint")
checkpoint("2020-07-02",
           R.version ="3.6.3",
           checkpointLocation = getwd())
# CUSTOM FUNCTION TO DEFINE THE PLOT THEMES -----
theme_AP <- function() {</pre>
  theme bw() +
  theme(panel.grid.major = element_blank(),
        panel.grid.minor = element_blank(),
        legend.background = element_rect(fill = "transparent",
                                         color = NA),
        legend.key = element_rect(fill = "transparent",
                                  color = NA))
}
```

2.2 Functions

2.2.1 Test functions

Here we define the test functions that we will use later on to benchmark our approach against the normal computation procedure: the Sobol' G (Sobol' 1993), the Bratley, Fox, and Niederreiter (1992), the Oakley and O'Hagan (2004) and the Morris (1991) functions.

```
# TEST FUNCTIONS -----

# Sobol G' function

G_Fun <- function(X) {
   a <- c(0, 0.5, 1, 1.5, 2, 2.5, 3, 3.5)
   y <- 1
   for (j in 1:8) {
      y <- y * (abs(4 * X[, j] - 2) + a[j])/(1 + a[j])
   }
}</pre>
```

```
return(y)
}
# Special G' function
G_Fun_mod <- function(X) {</pre>
  a \leftarrow c(0, 0, 0, 0, 0, 0, 0, 0)
  y <- 1
  for (j in 1:8) {
    y \leftarrow y * (abs(4 * X[, j] - 2) + a[j])/(1 + a[j])
  return(y)
}
# B1 function (in Kucherenko et al. 2011)
B1 <- function(X) {
  v <- 1
  for(j in 1:ncol(X)) {
    y \leftarrow y * (ncol(X) - X[, j]) / (ncol(X) - 0.5)
  return(y)
}
# C1 function (in Kucherenko et al. 2011)
C1 <- function(X) {
  y <- 1
  for (j in 1:ncol(X)) {
    y \leftarrow y * (abs(4 * X[, j] - 2))
  return(y)
}
# The function by Bratley et al. (1992) is defined in the sensobol package
# (Function A1 in Kucherenko et al. 2011)
# The function by Oakley and O'Hagan is defined in the sensobol package
# The function by Morris 1991 is included in the sensitivity package
```

2.2.2 Sample matrices

The following code chunk allows to create r replicas of the \mathbf{A} , \mathbf{B} , and $\mathbf{A}_{\mathbf{B}}^{i}$ matrices using Sobol' quasi-random number sequences (Bratley and Fox 1988). In this report we will stick with just one replica to save model runs in the model execution stage.

```
# SAMPLE MATRICES -----
CutBySize <- function(m, block.size, nb = ceiling(m / block.size)) {
  int <- m / nb
  upper <- round(1:nb * int)</pre>
```

```
lower \leftarrow c(1, upper[-nb] + 1)
  size <- c(upper[1], diff(upper))</pre>
  cbind(lower, upper)
}
scrambled_sobol <- function(A, B) {</pre>
  X <- rbind(A, B)</pre>
  for(i in 1:ncol(A)) {
    AB <- A
    AB[, i] \leftarrow B[, i]
    X <- rbind(X, AB)</pre>
  }
  AB <- X
  return(AB)
}
scrambled_replicas <- function(N, k, version) {</pre>
  X <- A <- B <- out <- list()</pre>
  df <- randtoolbox::sobol(n = N * version, dim = k * 2)</pre>
  indices <- CutBySize(nrow(df), nb = version)</pre>
  for(i in 1:nrow(indices)) {
    lower <- indices[i, "lower"]</pre>
    upper <- indices[i, "upper"]</pre>
    X[[i]] <- df[lower:upper, ]</pre>
  }
  for(i in seq_along(X)) {
    A[[i]] \leftarrow X[[i]][, 1:k]
    B[[i]] \leftarrow X[[i]][, (k + 1) : (k * 2)]
  }
  for(i in seq_along(A)) {
    out[[i]] <- scrambled_sobol(A[[i]], B[[i]])</pre>
  }
  return(out)
}
```

2.2.3 Sobol' indices

We use the estimators by Saltelli et al. (2010) and Jansen (1999) for S_i and T_i respectively.

The following code has been adapted from (???). The setting boot = TRUE in sobol_compute allows the function to randomly sample with replacement the row indices of the A, B, and A_B^i matrices, and compute S_i and T_i in the resulting sample. This setting is required to create 2n bootstrap replicas of the ranks during the execution of the smart_rank algorithm (see below). If boot = FALSE, sobol_compute directly calculates S_i and T_i on the original matrices, without bootstrapping. Finally, the function returns the sensitivity indices or the ranks depending on whether ranks = FALSE or ranks = TRUE respectively.

```
# SOBOL' INDICES -----
sobol_computeS <- function(Y_A, Y_B, Y_AB) {</pre>
  n <- length(Y_A[!is.na(Y_A)])</pre>
  f0 \leftarrow (1 / (2 * n)) * sum(Y_A + Y_B, na.rm = TRUE)
  VY \leftarrow 1 / n * sum((Y_A - f0) ^ 2, na.rm = TRUE)
  Si \leftarrow (1 / n) * sum(Y_B * (Y_AB - Y_A), na.rm = TRUE) / VY
  STi \leftarrow ((1 / (2 * n)) * sum((Y_A - Y_AB) ^ 2, na.rm = TRUE)) / VY
  return(c(Si, STi))
}
sobol_MapplyS <- function(d) {</pre>
  return(mapply(sobol_computeS,
                  d[, "Y_A"],
                  d[, "Y_B"],
                  d[, "Y_AB"]))
}
sobol_compute <- function(Y, params, boot = FALSE, ranks = FALSE) {</pre>
  k <- length(params)</pre>
  p \leftarrow length(1:(length(Y) / (k + 2)))
  Y_A \leftarrow Y[1:p]
  Y_B \leftarrow Y[(p + 1) : (2 * p)]
  Y_AB \leftarrow Y[(2 * p + 1):((length(Y) / (k + 2)) * (k + 2))]
  parameters <- rep(params, each = length(Y_A))</pre>
  vec <- cbind(Y_A, Y_B, Y_AB)</pre>
  out <- data.table::data.table(vec, parameters)</pre>
  if(boot == TRUE) {
    indices <- out[, sample(.I, replace = TRUE), parameters][, V1]</pre>
    output <- out[indices, sobol_MapplyS(.SD), by = parameters][</pre>
       , sensitivity:= rep(c("Si", "STi"), times = k)]
  if(boot == FALSE) {
    output <- out[, sobol_MapplyS(.SD), by = parameters][
       , sensitivity:= rep(c("Si", "STi"), times = k)]
  if(ranks == FALSE) {
    final <- output[, V1, sensitivity][, V1]</pre>
  if(ranks == TRUE) {
    final <- output[, rank(-V1), sensitivity][, V1]</pre>
  }
  return(final)
}
```

2.3 The new algorithm

Here we describe the new algorithm to increase the efficiency of the computation of T_i . Its behaviour is graphically represented in Fig. 1. It works as follows:

- 1. It runs sequentially at sample sizes $N = 2^2, 2^3, ..., 2^m$, where m is defined by the user. After constructing a (N, 2k) Sobol' matrix, for i = 1, 2, ...k parameters, it computes the model output, the sensitivity indices S_{Ti} and T_i and the ranks of the parameters T_{Ri} .
- 2. At each sample size $N \neq 2^2$, it checks the following statement:
 - (a) Condition 1: $T_{Ri}^{(N)}, T_{Rj}^{(N)}, \dots \equiv T_{Ri}^{(N-1)}, T_{Rj}^{(N-1)}, \dots$
- 3. If true, the algorithm considers that the ranks have converged and computes the set-sensitivity of the parameters with $T_i < 0.05$. The total cost C' of the analysis at convergence is therefore C' = N'(k+3), where N' is the size of the sample matrix at convergence. At rank convergence, and for the next sample sizes, the algorithm computes the T_i of the influential parameters only, as well as the T_s , leading to (N N')(k k' + 3) model runs, where k' is the number of non-influential parameters. Overall, the total cost C is C = N'(k+3) + (N N')(k k' + 3) = N(k k' + 3) + N'k'.
- 4. If $T_{Ri}^{(N)}, T_{Rj}^{(N)}, ... \neq T_{Ri}^{(N-1)}, T_{Rj}^{(N-1)}, ...$, the algorithm checks for two more conditions before considering that the ranks have not converged, and that is necessary to increase the sample size. This is to ensure that the lack of convergence at N' is not caused by small scillations between non-important parameters, i.e. with $T_i < 0.05$, which are negligible for ranking purposes.
 - (a) Condition 2: let $T_i^{(N)}, T_j^{(N)}, \ldots$, and $T_i^{(N-1)}, T_j^{(N-1)}, \ldots$ be the total sensitivity indices of the parameters whose ranks at N and N-1 do not converge. The algorithm then checks for the following condition: $|(T_i^{(N)} T_j^{(N)} \ldots)|/(T_i^{(N)} + T_j^{(N)} + \ldots) < 0.1$, and $|(T_i^{(N-1)} T_j^{(N-1)} \ldots)|/(T_i^{(N-1)} + T_j^{(N-1)} + \ldots) < 0.1$. If the condition is true for both vectors, the algorithm considers that the oscillation of ranks is negligible from a factor fixing standpoint. The ranks have thus converged, and it applies the computations described in point 3 above. If the condition is false for at least one vector, it checks condition 3.
 - (b) Condition 3: the algorithm checks whether $T_i^{(N)}, T_j^{(N)}, \ldots < 0.02$ and $T_i^{(N-1)}, T_j^{(N-1)}, \ldots < 0.02$. If true for both vectors, it considers that the swapping of ranks is due to insignificant oscillations between parameters the ranks have thus converged, and applies the computations described in point 3 above. If false, it definetely considers that the sample size is not enough to guarantee a robust ranking, and the algorithm jumps to the next sample size.

```
# DEFINE ALGORITHM -----

# Compute normal

compute_normal <- function(N, params, test_F) {
    A <- sobol_matrices(n = N, k = length(params))
    Y <- test_F(A)
    indices <- sobol_compute(Y, params = params)
    ranks <- sobol_compute(Y, params = params, ranks = TRUE)
    dt <- data.table(cbind(indices, ranks))</pre>
```

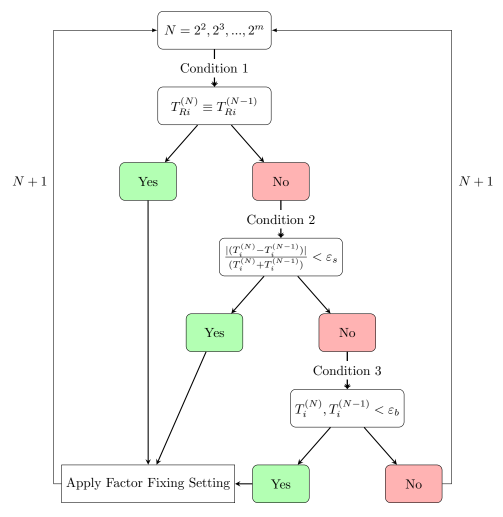


Figure 1: Diagram of the new algorithm

```
dt[, parameters:= rep(params, times = 2)][
    , sensitivity:= rep(c("Si", "STi"), each = length(params))][
    , method:= "Normal.approach"][
    , N := N][
    , model.runs:= N * (length(params) + 2)]
 return(dt)
}
# Algorithm to save runs
compute_saving <- function(dt, params, N, test_F, eps = 0.05) {</pre>
  dt2 \leftarrow copy(dt)
  set <- dt[sensitivity == "STi" & indices <= 0.05][, parameters]
  if(length(set) > 1) {
    Ti.set <- match(set, params)</pre>
    Ti.important <- match(setdiff(params, set), params)</pre>
    A.important <- sobol_matrices(N, k = length(params), cluster = Ti.important)
    A.set <- sobol_matrices(N, k = length(params), cluster = list(Ti.set))
    A.full <- rbind(A.important, A.set[(2 * N + 1):nrow(A.set), ])
    Y <- test_F(A.full)
    indices <- sobol_compute(Y, params = c(params[Ti.important], "set"))</pre>
    ranks <- sobol_compute(Y, params = c(params[Ti.important], "set"), ranks = TRUE)</pre>
    dt <- data.table(cbind(indices, ranks))</pre>
    dt[, parameters:= rep(c(params[Ti.important], "set"), times = 2)][
      , sensitivity:= rep(c("Si", "STi"), each = length(c(params[Ti.important], "set")))][
      , method:= "New.approach"][
      , N := N
      , model.runs:= N * (length(params) - length(set) + 3)]
    final <- rbind(dt2, dt)</pre>
  } else {
   final <- dt
  }
 return(final)
}
# Full algorithm
full_algorithm <- function(max.exponent, params, epsilon_1, epsilon_2, test_F) {</pre>
 N <- sapply(2:max.exponent, function(x) 2 ^ x)</pre>
 dt <- list()</pre>
  for(i in seq_along(N)) {
    dt[[i]] <- compute_normal(N[i], params = params, test_F = test_F)</pre>
    if(i > 1) {
      a <- dt[[i]][sensitivity == "STi" & method == "Normal.approach"]</pre>
      b <- dt[[i-1]][sensitivity == "STi" & method == "Normal.approach"]
      # CHECK CONDITION 1-----
      condition1 <- identical(a[, ranks], b[, ranks])</pre>
      # CHECK CONDITION 2-----
      ind <- which(!a[, ranks] == b[, ranks])</pre>
```

```
condition2 <- condition2 <- all(abs(a[ind][, indices] - b[ind][, indices]) /</pre>
                           (a[ind][, indices] + b[ind][, indices]) < epsilon_1)</pre>
      # CHECK CONDITION 3 -----
      condition3 <- all(c(a[ind][, indices], b[ind][, indices]) < epsilon_2)</pre>
      if(condition1 == TRUE |
         condition1 == FALSE & condition2 == TRUE |
         condition1 == FALSE & condition2 == FALSE & condition3 == TRUE) {
        dt[[i]] <- compute_saving(dt[[i]], params = params, N = N[i], test_F = test_F)
      } else {
        dt[[i]] <- compute_normal(N[i], params = params, test_F = test_F)
    }
  }
  # Compute total number of model runs for the new algorithm
  final <- rbindlist(dt)</pre>
  N_convergence <- final[method == "New.approach" & sensitivity == "STi", min(N)]
  a <- final[N == N_convergence & method == "Normal.approach"
             & sensitivity == "STi"][, parameters]
 b <- final[N == N_convergence & !parameters == "set" &
               method == "New.approach" & sensitivity == "STi"][, parameters]
 k_noninfluential <- length(setdiff(a, b))</pre>
  final[, model.runs:= ifelse(method %in% "New.approach",
                               model.runs + N_convergence * k_noninfluential,
                               model.runs)]
 return(final)
}
# Full algorithm in a function
fun_algorithm <- function(max.exponent, epsilon_1, epsilon_2) {</pre>
  test_functions <- c("G_Fun", "G_Fun_mod", "bratley1992_Fun",</pre>
                       "oakley_Fun", "morris_Fun", "B1", "C1")
  out <- list()
  for(i in test functions) {
    if(i == "G_Fun") {
      params <- paste("X", 1:8, sep = "")</pre>
      test_F <- G_Fun</pre>
    } else if(i == "G_Fun_mod") {
      params <- paste("X", 1:8, sep = "")</pre>
      test_F <- G_Fun_mod</pre>
    } else if(i == "bratley1992_Fun") {
      params <- paste("X", 1:8, sep = "")</pre>
     test_F <- bratley1992_Fun
    } else if(i == "oakley_Fun") {
      params <- paste("X", 1:15, sep = "")</pre>
      test_F <- oakley_Fun</pre>
    } else if(i == "B1") {
      params <- paste("X", 1:8, sep = "")</pre>
```

```
test_F <- B1
    } else if(i == "C1") {
      params <- paste("X", 1:8, sep = "")</pre>
      test_F <- C1
    } else {
      params <- paste("X", 1:20, sep = "")</pre>
      test_F <- sensitivity::morris.fun</pre>
    out[[i]] <- full_algorithm(max.exponent = max.exponent,</pre>
                                  params = params,
                                  epsilon_1 = epsilon_1,
                                  epsilon_2 = epsilon_2,
                                  test_F = test_F)
  }
  final <- rbindlist(out, idcol = "model")</pre>
  return(final)
}
```

We test the performance of the algorithm with the Sobol' G (Sobol' 1993), the Bratley, Fox, and Niederreiter (1992) the Oakley and O'Hagan (2004) and the Morris (1991) functions at each N, for $N = 2^2, 2^3, \dots, 2^{16}$.

```
# RUN THE MODEL -----

# Define epsilons and max exponent
epsilon_1 <- seq(0.05, 0.1, 0.05)
epsilon_2 <- seq(0.01, 0.02, 0.005)
max.exponent <- 16

# Run model
out <- mclapply(epsilon_1, function(x)
    lapply(epsilon_2, function(y)
    fun_algorithm(max.exponent = max.exponent, epsilon_1 = x, epsilon_2 = y)),
    mc.cores = detectCores())</pre>
```

We now arrange the results for better plotting:

```
results <- lapply(out, function(x) lapply(x, function(y) {
  tmp <- y[, parameters:= factor(parameters,</pre>
                            levels = c(paste("X", 1:20, sep = ""), "set"))] %>%
    .[, Type:= ifelse(parameters %in% "set", "Set", "Individual")] %>%
    .[, model:= ifelse(model == "G_Fun", "Sobol G",
                       ifelse(model == "G_Fun_mod", "Sobol G modified",
                               ifelse(model == "bratley1992_Fun", "Bratley et al. 1994",
                                      ifelse(model == "oakley_Fun", "Oakley and O'Hagan 2004",
                                             ifelse(model == "morris_Fun", "Morris 1991",
                                                    ifelse(model == "B1", "B1", "C1"))))))
}))
# Compare number of model runs and savings
tmp <- lapply(out, function(x) lapply(x, function(y) {</pre>
  tmp <- y[, .(model.runs = unique(model.runs)), .(model, N, method)] %>%
    dcast(., model + N ~ method, value.var = "model.runs")
}))
savings.dt <- lapply(tmp, function(x) lapply(x, function(y) {</pre>
  tmp <- y[, New.approach:= ifelse(New.approach %in% NA, Normal.approach, New.approach)] %>%
    .[, saving:= 1 - (New.approach / Normal.approach)] %>%
    .[, model:= ifelse(model == "G_Fun", "Sobol G",
                       ifelse(model == "G_Fun_mod", "Sobol G modified",
                               ifelse(model == "bratley1992_Fun", "Bratley et al. 1994",
                                      ifelse(model == "oakley_Fun", "Oakley and O'Hagan 2004",
                                             ifelse(model == "morris_Fun", "Morris 1991",
                                                    ifelse(model == "B1", "B1", "C1"))))))
}))
```

3 Results

Fig. 2a shows the evolution of the T_i and the T_s along different sample sizes. The onset of the lines marks the sample size at which our algorithm kicks in, i.e. the sample size at which the ranks, according to the conditions set in the previous chapter, are considered converged. The first dot thus indicates the smallest sample size needed to obtain a robust screening of the parameters: note that, once the ranks converge, they do not "disconverge" and the algorithm is activated at every sample size. The computation of the T_s shows that grouping non-important factors might not have a dramatic effect in the model output if they are fixed to reduce model complexity. All $T_s \approx 0.05$ except in the case of the Oakley and O'Hagan (2004) function, for which $T_s \approx 0.15$.

It is also worth stressing that the cardinality of the T_s , i.e. the number of elements in the set, is constant throughout the range of sample sizes tested (Fig. 2b), and that the T_s always includes the same set of parameters regardless of the sample size used (not shown in the plot but checked in R). This suggests that, once the algorithm kicks in, there is no need to increase the sample size in order to achieve more precise results for a factor fixing setting. However, some influential parameters with

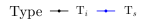
very similar T_i values might still swap ranks, and a larger sample size might be needed to ensure their full stabilization.

Under this scenario, our algorithm allows to search for a full robust ranking of all parameters through different sample sizes while saving a significant number of model runs in the process. This is shown in Fig. 2c: the x axis presents the total number of model runs once the algorithm is activated, whereas the y axis the number of model runs saved as a percentage over the total N(k+2). Note that there is a drop in the savings at rank convergence compared to the traditional approach: this is because the algorithm adds N' model runs to compute the T_s . Once converged, the total number of runs is N(k-k'+3) + N'k', which yields to significant savings in the computation cost, i.e. up to 40% in the case of the Bratley, Fox, and Niederreiter (1992) and the Morris (1991) functions.

4 Preliminary conclusions

- Our algorithm finds the minimum sample size needed to ensure a robust screening of parameters. This will allow researchers to save computer time when searching for an ideal sample size to ensure consistent results in a factor fixing setting.
- If what is aimed at is at ensuring full rank convergence, our algorithm allows to explore different sample sizes while providing significant and incremental benefits in terms of savings over the total cumulative number of runs.
- A necessary pre-condition for the algorithm to allow to save model runs is the existence of at least two parameters with $T_i < 0.05$. This is not uncommon in real case studies, where sensitivity indices tend to follow a Pareto distribution (i.e. the indices of 80% of the parameters are responsible for less than 20% of the variance and those of the remainder form more than 80%).
- It should be noted that the results are fully dependent on some arbitrarily selected values: 1) 0.05 as a threshold to distinguish influential from non-influential parameters, 2) 0.1 to define Sergei's 'condition 2' (see code in page 13 and 14), and 3) 0.02 in 'condition 3' (see code in page 14). The value of 0.05 might be the most disputed choice as other thresholds and screening methods have been proposed in the literature.

\$`0.05` ## \$`0.05`\$`0.01`



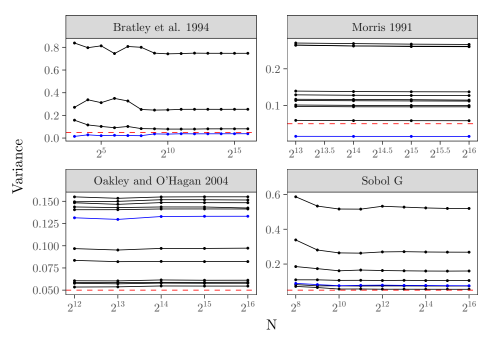
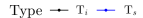


Figure 2: Evolution of the T_i and the T_s along different sample sizes. The horizontal, red dotted line is at 0.05.

```
##
## $`0.05`$`0.015`
##
## $`0.05`$`0.02`
##
##
## $`0.1`
##
## $`0.1`$`0.01`
```



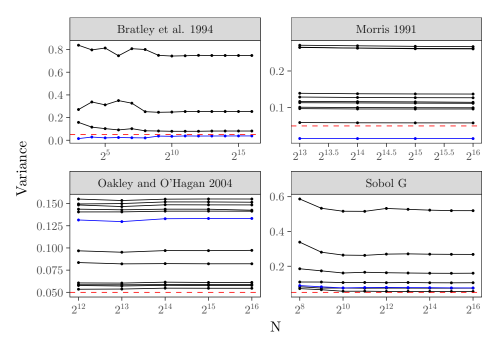


Figure 3: Evolution of the T_i and the T_s along different sample sizes. The horizontal, red dotted line is at 0.05.

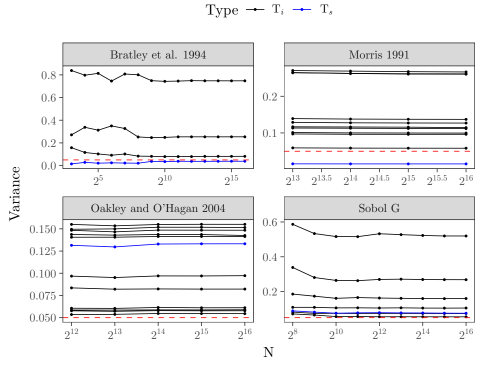
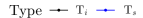


Figure 4: Evolution of the T_i and the T_s along different sample sizes. The horizontal, red dotted line is at 0.05.



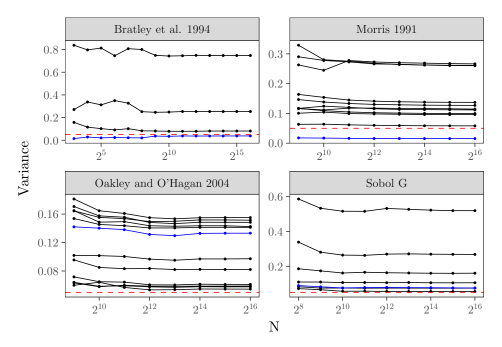


Figure 5: Evolution of the T_i and the T_s along different sample sizes. The horizontal, red dotted line is at 0.05.

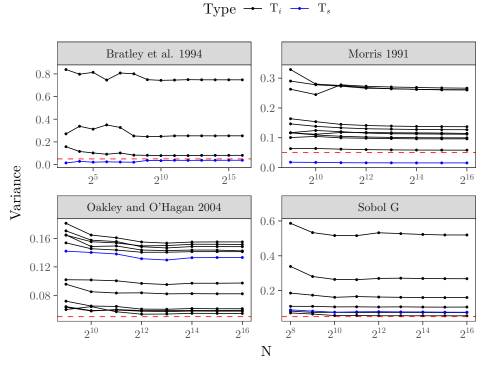
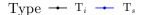


Figure 6: Evolution of the T_i and the T_s along different sample sizes. The horizontal, red dotted line is at 0.05.

```
##
## $`0.1`$`0.02`
```



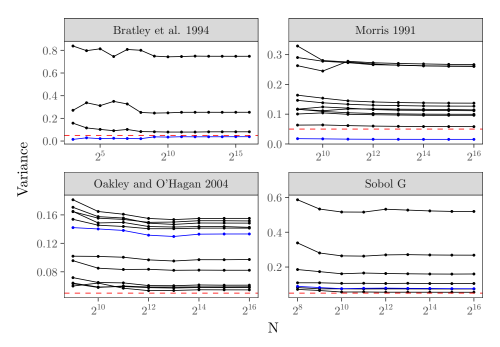


Figure 7: Evolution of the T_i and the T_s along different sample sizes. The horizontal, red dotted line is at 0.05.

```
# PLOT SAVINGS -----
lapply(savings.dt, function(x) lapply(x, function(y)
 melt(y, measure.vars = c("New.approach", "Normal.approach")) %>%
    .[, saving:= saving * 100] %>%
    setnames(., "model", "Model")) %>%
   rbindlist(., idcol = "epsilon_2")) %>%
 rbindlist(., idcol = "epsilon_1") %>%
  .[, epsilon_1:= paste("$\\varepsilon_s= ", epsilon_1, "$", sep = "")] %>%
  .[, epsilon_2:= paste("\varepsilon_b= ", epsilon_2, "\s, sep = "")] %>%
  ggplot(., aes(N, saving, color = Model, shape = Model, group = Model)) +
  geom_point() +
  geom_line() +
  scale x continuous(trans="log",
                    breaks = trans_breaks("log2", function(x) 2 ^ x),
                    labels = trans_format("log2", math_format(2^.x))) +
 labs(x = "Number of model runs",
       y = "Saving (\\)") +
 facet_grid(epsilon_1 ~ epsilon_2) +
  theme_AP() +
  theme(legend.position = "top") +
  guides(color = guide_legend(nrow = 3, byrow = TRUE))
```

Warning: The shape palette can deal with a maximum of 6 discrete values because
more than 6 becomes difficult to discriminate; you have 7. Consider
specifying shapes manually if you must have them.

Warning: Removed 180 rows containing missing values (geom_point).

```
→ B1 → Bratley et al. 1994 → C1

Model → Morris 1991 → Oakley and O'Hagan 2004 → Sobol G

— Sobol G modified
```

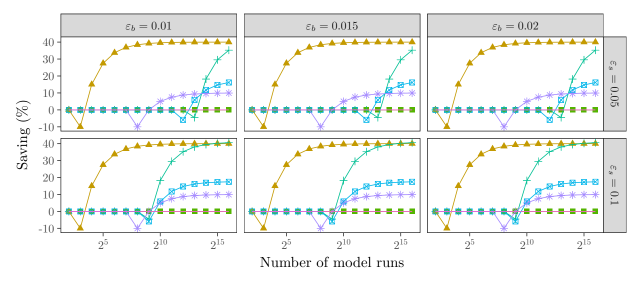


Figure 8: Percentage of savings over the total accumulated model runs.

```
# PLOT SAVINGS -----
lapply(savings.dt, function(x) lapply(x, function(y)
 melt(y, measure.vars = c("New.approach", "Normal.approach")) %>%
    .[, saving:= saving * 100] %>%
   setnames(., "model", "Model")) %>%
   rbindlist(., idcol = "epsilon_2")) %>%
 rbindlist(., idcol = "epsilon_1") %>%
  .[, epsilon_1:= paste("\varepsilon_s= ", epsilon_1, "\s", sep = "")] %>%
  .[, epsilon_2:= paste("$\\varepsilon_b= ", epsilon_2, "$", sep = "")] %>%
  .[!Model %in% c("C1", "B1", "Sobol G modified")] %>%
  ggplot(., aes(N, saving, color = Model, shape = Model, group = Model)) +
  geom_point() +
  geom_line() +
  scale_x_continuous(trans="log",
                     breaks = trans_breaks("log2", function(x) 2 ^ x),
                     labels = trans_format("log2", math_format(2^.x))) +
 labs(x = "Number of model runs",
       y = "Saving (\\)") +
 facet_grid(epsilon_1 ~ epsilon_2) +
```

```
theme_AP() +
theme(legend.position = "top") +
guides(color = guide_legend(nrow = 1, byrow = TRUE))
```

Model → Bratley et al. 1994 → Morris 1991 → Oakley and O'Hagan 2004 → Sobol G

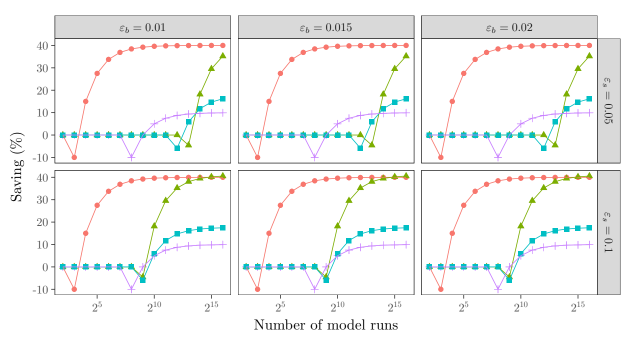


Figure 9: Percentage of savings over the total accumulated model runs.

```
# PLOT RANKS -----
results[[2]][[3]][method == "Normal.approach"] %>%
  .[, sensitivity:= ifelse(sensitivity %in% "Si", "$S_i$", "$T_i$")] %>%
  .[sensitivity == "$T_i$"] %>%
  ggplot(., aes(N, ranks, group = parameters,
                color = parameters)) +
  geom_point(size = 0.5) +
 geom_line() +
 labs(x = "Base sample size",
       y = "Rank") +
  scale_color_discrete(name = "Parameters") +
  scale_x_continuous(trans="log",
                     breaks = trans_breaks("log2", function(x) 2 ^ x),
                     labels = trans_format("log2", math_format(2^.x))) +
  scale_y_reverse() +
  facet_grid(model ~ sensitivity,
             scales = "free_y") +
  theme_bw() +
 theme(legend.position = "top",
        panel.grid.major = element_blank(),
```

```
.[!model %in% c("C1", "B1", "Sobol G modified")] %>%
ggplot(., aes(N, ranks, group = parameters,
              color = parameters)) +
geom_point(size = 0.5) +
geom_line() +
labs(x = "Base sample size",
     y = "Rank") +
scale_color_discrete(name = "Parameters") +
scale_x_continuous(trans="log",
                   breaks = trans breaks("log2", function(x) 2 ^ x),
                   labels = trans_format("log2", math_format(2^.x))) +
scale_y_reverse() +
facet_grid(model ~ sensitivity,
           scales = "free_y") +
theme_bw() +
theme(legend.position = "top",
      panel.grid.major = element_blank(),
      panel.grid.minor = element_blank(),
      legend.background = element_rect(fill = "transparent",
                                       color = NA),
      legend.key = element_rect(fill = "transparent",
                                color = NA))
```

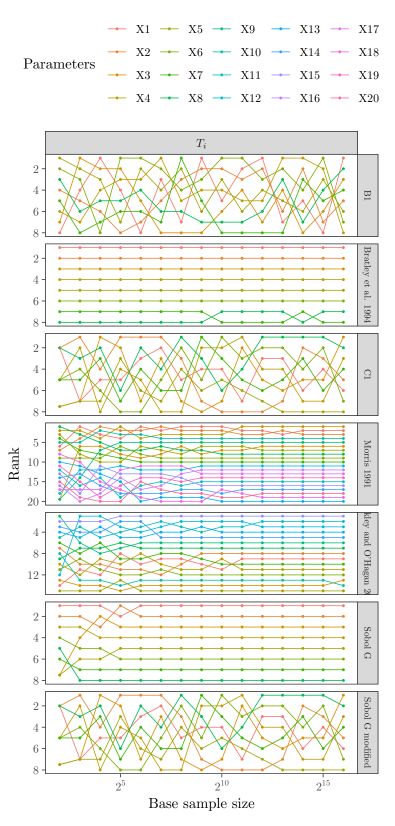


Figure 10: Evolution of ranks for total-order indices across different base sample sizes

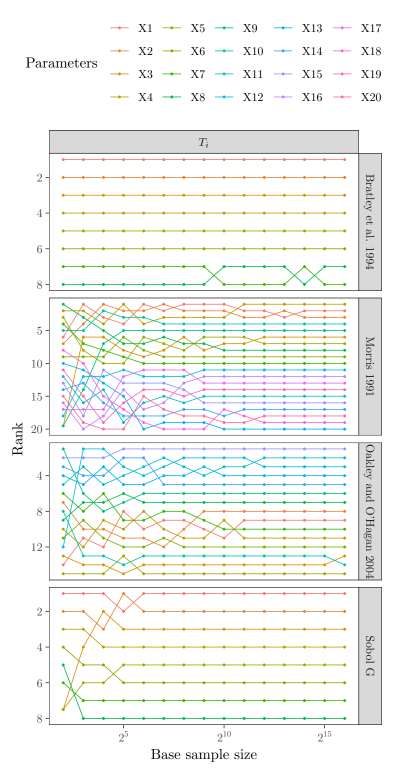


Figure 11: Evolution of ranks for total-order indices across different base sample sizes.

5 Session information

```
# SESSION INFORMATION ----
sessionInfo()
## R version 3.6.3 (2020-02-29)
## Platform: x86_64-apple-darwin15.6.0 (64-bit)
## Running under: macOS Catalina 10.15.5
##
## Matrix products: default
## BLAS:
           /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRblas.0.dylib
## LAPACK: /Library/Frameworks/R.framework/Versions/3.6/Resources/lib/libRlapack.dylib
##
## locale:
## [1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
## attached base packages:
## [1] grid
                 parallel stats
                                      graphics grDevices utils
                                                                    datasets
## [8] methods
                 base
##
## other attached packages:
## [1] scales_1.1.1
                           checkpoint_0.4.9
                                               data.table_1.12.8
                                                                  tikzDevice_0.12.3
## [5] RColorBrewer_1.1-2 wesanderson_0.3.6
                                               sensitivity_1.18.1 gridExtra_2.3
## [9] cowplot_1.0.0
                           forcats_0.5.0
                                               stringr_1.4.0
                                                                  dplyr_0.8.5
## [13] purrr_0.3.4
                           readr_1.3.1
                                                                  tibble_3.0.2
                                               tidyr_1.0.2
## [17] ggplot2_3.3.2
                           tidyverse_1.3.0
                                               sensobol_0.3
##
## loaded via a namespace (and not attached):
## [1] httr_1.4.1
                         jsonlite_1.6.1
                                           foreach_1.5.0
                                                            modelr_0.1.6
## [5] Rdpack_1.0.0
                         assertthat_0.2.1 highr_0.8
                                                            cellranger_1.1.0
## [9] yaml_2.2.1
                         pillar_1.4.6
                                                            lattice 0.20-41
                                           backports_1.1.8
## [13] glue_1.4.1
                         digest_0.6.25
                                           rvest_0.3.5
                                                            colorspace_1.4-1
## [17] htmltools 0.4.0
                         pkgconfig 2.0.3
                                           bibtex_0.4.2.2
                                                            broom 0.5.6
## [21] haven_2.2.0
                         generics_0.0.2
                                           farver_2.0.3
                                                            ellipsis_0.3.1
## [25] withr 2.2.0
                         cli_2.0.2
                                           magrittr_1.5
                                                            crayon_1.3.4
## [29] readxl_1.3.1
                         evaluate_0.14
                                           fs_1.4.1
                                                            fansi_0.4.1
## [33] nlme_3.1-147
                         xm12_1.3.1
                                           tools_3.6.3
                                                            hms_0.5.3
## [37] gbRd_0.4-11
                         lifecycle_0.2.0
                                           munsell_0.5.0
                                                            reprex_0.3.0
## [41] compiler_3.6.3
                         tinytex_0.22
                                           rlang_0.4.7
                                                            iterators_1.0.12
## [45] rstudioapi_0.11
                         filehash_2.4-2
                                           labeling_0.3
                                                            rmarkdown_2.1
## [49] boot_1.3-24
                         gtable_0.3.0
                                           codetools_0.2-16 DBI_1.1.0
## [53] R6_2.4.1
                         lubridate_1.7.8
                                           knitr_1.28
                                                            stringi_1.4.6
## [57] Rcpp_1.0.5
                         vctrs_0.3.1
                                           dbplyr_1.4.3
                                                            tidyselect_1.0.0
## [61] xfun_0.13
```

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