

# The battle of total-order sensitivity estimators

R code

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```

# PRELIMINARY FUNCTIONS -----
# 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)
      library(i, character.only = TRUE)
    }
  }
}

# Install development version of sensobol
remotes::install_github("arnaldpuy/sensobol")

# Load the packages
loadPackages(c("Rcpp", "RcppArmadillo", "tidyverse", "parallel", "foreach",
              "doParallel", "Rfast", "data.table", "scales", "cowplot",
              "benchmarkme", "logitnorm", "sensobol", "ggrepel"))

# Create custom theme
theme_AP <- function() {
  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))
}

# Set checkpoint

dir.create(".checkpoint")
library("checkpoint")

checkpoint("2020-04-21",
           R.version ="3.6.3",
           checkpointLocation = getwd())

```

# 1 Functions

## 1.1 Savage scores

```
# SAVAGE SCORES -----  
  
savage_scores <- function(x) {  
  true.ranks <- rank(-x)  
  p <- sort(1 / true.ranks)  
  mat <- matrix(rep(p, length(p)), nrow = length(p), byrow = TRUE)  
  mat[upper.tri(mat)] <- 0  
  out <- sort(rowSums(mat), decreasing = TRUE)[true.ranks]  
  return(out)  
}
```

## 1.2 Sensitivity indices

```
# VARS FUNCTIONS -----  
  
vars_matrices <- function(star.centers, params, h, method = "QRN") {  
  out <- center <- sections <- A <- B <- AB <- X <- out <- list()  
  if(method == "QRN") {  
    mat <- randtoolbox::sobol(n = star.centers, dim = length(params))  
  } else if(method == "R") {  
    mat <- replicate(length(params), stats::runif(star.centers))  
  } else if(method == "LHS") {  
    mat <- lhs::randomLHS(star.centers, length(params))  
  } else {  
    stop("method should be either QRN, R or LHS")  
  }  
  for(i in 1:nrow(mat)) {  
    center[[i]] <- mat[i, ]  
    sections[[i]] <- sapply(center[[i]], function(x) {  
      all <- seq(x %% h, 1, h)  
      non.zeros <- all[all != 0] # Remove zeroes  
    })  
    B[[i]] <- sapply(1:ncol(mat), function(x)  
      sections[[i]][, x][!sections[[i]][, x] %in% center[[i]][x]])  
    A[[i]] <- matrix(center[[i]], nrow = nrow(B[[i]]),  
                      ncol = length(center[[i]]), byrow = TRUE)  
    X[[i]] <- rbind(A[[i]], B[[i]])  
    for(j in 1:ncol(A[[i]])) {  
      AB[[i]] <- A[[i]]  
      AB[[i]][, j] <- B[[i]][, j]  
      X[[i]] <- rbind(X[[i]], AB[[i]])  
    }  
    AB[[i]] <- X[[i]][(2 * nrow(B[[i]])) + 1]:nrow(X[[i]]), ]  
    out[[i]] <- rbind(unname(center[[i]]), AB[[i]])  
  }
```

```

    }
    return(do.call(rbind, out))
}

# Function to cut by size
CutBySize <- function(m, block.size, nb = ceiling(m / block.size)) {
  int <- m / nb
  upper <- round(1:nb * int)
  lower <- c(1, upper[-nb] + 1)
  size <- c(upper[1], diff(upper))
  cbind(lower, upper, size)
}

# Function to compute VARS-TI
vars_ti <- function(Y, star.centers, params, h) {
  n.cross.points <- length(params) * ((1 / h) - 1) + 1
  index.centers <- seq(1, length(Y), n.cross.points)
  mat <- matrix(Y[-index.centers], ncol = star.centers)
  indices <- CutBySize(nrow(mat), nb = length(params))
  out <- list()
  for(i in 1:nrow(indices)) {
    out[[i]] <- mat[indices[i], "lower":indices[i, "upper"], ]
  }
  d <- lapply(1:length(params), function(x)
    lapply(1:ncol(out[[x]]), function(j) {
      da <- c(out[[x]][, j][1],
              rep(out[[x]][, j][-c(1, length(out[[x]][, j]))], each = 2),
              out[[x]][, j][length(out[[x]][, j])])
    }))
  out <- lapply(d, function(x) lapply(x, function(y) matrix(y, nrow = length(y) / 2, byrow = TRUE)))
  variogr <- unlist(lapply(out, function(x) lapply(x, function(y)
    mean(0.5 * (y[, 1] - y[, 2]) ^ 2))) %>%
    lapply(., function(x) do.call(rbind, x)) %>%
    lapply(., mean)))
  covariogr <- unlist(lapply(out, function(x)
    lapply(x, function(y) cov(y[, 1], y[, 2]))) %>%
    lapply(., function(x) Rfast::colmeans(do.call(rbind, x))))
  VY <- var(Y[index.centers])
  Ti <- (variogr + covariogr) / VY
  output <- data.table::data.table(Ti)
  output[, `:=` (parameters = params)]
  return(output)
}

# COMPUTATION OF SOBOL' Ti INDICES -----
sobel_Ti <- function(d, N, params, total) {

```

```

m <- matrix(d, nrow = N)
k <- length(params)
if(total == "jansen" | total == "homma" | total == "sobol" | total == "monod" |
  total == "glen") {
  Y_A <- m[, 1]
  Y_AB <- m[, -1]
  f0 <- (1 / length(Y_A)) * sum(Y_A)
  VY <- 1 / length(Y_A) * sum((Y_A - f0) ^ 2)
  #  $VY \leftarrow 1 / \text{length}(Y_A) * (\sum(Y_A^2) -$ 
  #  $(1 / N * \sum(Y_A^2)))$  ((Variance used by Becker))
}
if(total == "jansen") {
  Ti <- (1 / (2 * N) * Rfast::colsums((Y_A - Y_AB) ^ 2)) / VY
} else if(total == "homma") {
  Ti <- (VY - (1 / N) * Rfast::colsums(Y_A * Y_AB) + f0 ^ 2) / VY
} else if(total == "sobol") {
  Ti <- ((1 / N) * Rfast::colsums(Y_A * (Y_A - Y_AB))) / VY
} else if(total == "monod") {
  Ti <- 1 - (1 / N * Rfast::colsums(Y_A * Y_AB) -
    (1 / N * Rfast::colsums((Y_A + Y_AB) / 2)) ^ 2) /
    (1 / N * Rfast::colsums((Y_A ^ 2 + Y_AB ^ 2) / 2) -
    (1 / N * Rfast::colsums((Y_A + Y_AB) / 2)) ^ 2)
} else if(total == "glen") {
  Ti <- 1 - (1 / (N - 1) *
    Rfast::colsums(((Y_A - mean(Y_A)) * (Y_AB - Rfast::colmeans(Y_AB))) /
    sqrt(var(Y_A) * Rfast::colVars(Y_AB))))
}
if(total == "azzini" | total == "lambtoni") {
  Y_A <- m[, 1]
  Y_B <- m[, 2]
  Y_AB <- m[, 3:(3 + k - 1)]
  Y_BA <- m[, (ncol(m) - k + 1):ncol(m)]
  f0 <- 1 / (2 * N) * sum(Y_A + Y_B)
  VY <- 1 / (2 * N - 1) * sum((Y_A - f0) ^ 2 + (Y_B - f0) ^ 2)
}
if(total == "azzini") {
  Ti <- 1 - abs(Rfast::colsums((Y_A - Y_BA) * (Y_B - Y_AB)) /
    (1 / 2 * Rfast::colsums((Y_A - Y_B) ^ 2 + (Y_AB - Y_BA) ^ 2)))
} else if(total == "lambtoni") {
  Ti <- (1 / (4 * N) * colSums((Y_A - Y_AB) ^ 2 + (Y_B - Y_BA) ^ 2, na.rm = TRUE)) / VY
}
if(total == "owen") {
  Y_A <- m[, 1]
  Y_B <- m[, 2]
  Y_BA <- m[, 3:(3 + k - 1)]
  Y_CB <- m[, (ncol(m) - k + 1):ncol(m)]
  VY <- sapply(1:k, function(j)

```

```

    mean(Rfast::rowmeans(m[,c(1, 2, 2 + j, 2 + j + k)]^ 2)) -
    mean(Rfast::rowmeans(m[,c(1, 2, 2 + j, 2 + j + k)]))^ ^2)
  Ti <- (VY - (1 / N * Rfast::colsums((Y_B - Y_CB) * (Y_BA - Y_A)))) / VY
}
output <- data.table(Ti)
output[, `:=` (parameters = paste("X", 1:k, sep = ""))]
return(output)
}

# CHECK THAT ALL TI ESTIMATORS WORK -----
# Settings
estimators <- c("jansen", "sobol", "homma", "azzini", "monod", "lamboni", "glen", "owen")
test_functions <- c("Ishigami", "Sobol'G", "Morris")
N <- 2 ^ 11

# Run model
ind <- Y <- mt <- list()
for(i in estimators) {
  for(j in test_functions) {
    if(i == "jansen" | i == "sobol" | i == "homma" | i == "monod" | i == "glen") {
      matrices <- c("A", "AB")
    } else if(i == "azzini" | i == "lamboni"){
      matrices <- c("A", "B", "AB", "BA")
    } else if(i == "owen") {
      matrices <- c("A", "B", "BA", "CB")
    }
    if(j == "Ishigami") {
      k <- 3
      modelRun <- sensobol::ishigami_Fun
    } else if(j == "Sobol'G") {
      k <- 8
      modelRun <- sensobol::sobol_Fun
    } else if(j == "Morris") {
      k <- 20
      modelRun <- sensitivity::morris.fun
    }
    mt[[i]][[j]] <- sobol_matrices(N = N, params = paste("X", 1:k, sep = ""), matrices = matrices)
    Y[[i]][[j]] <- modelRun(mt[[i]][[j]])
    ind[[i]][[j]] <- sobol_Ti(d = Y[[i]][[j]], params = paste("X", 1:k, sep = ""),
                                N = N, total = i)
  }
}

## Registered S3 method overwritten by 'sensitivity':
##   method     from
##   print.src dplyr

```

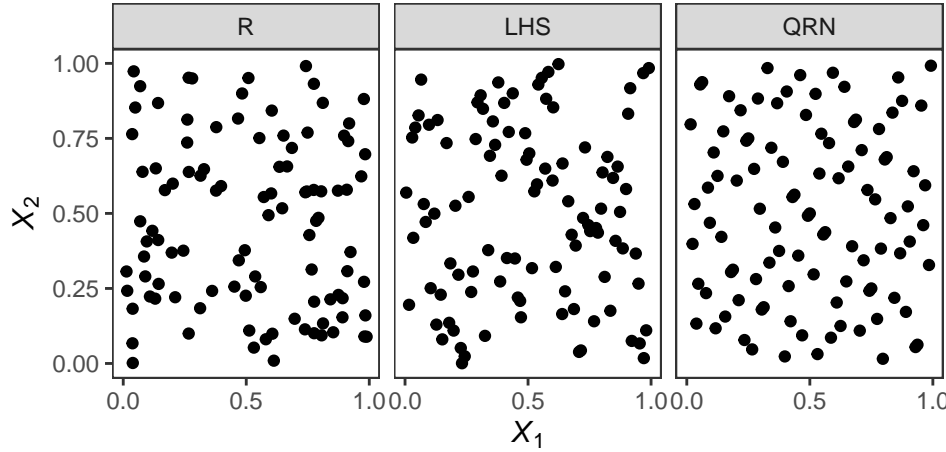
```

# Run model for VARS
star.centers <- 200
h <- 0.2
vars.ind <- Y <- list()
for(j in test_functions) {
  if(j == "Ishigami") {
    k <- 3
    modelRun <- sensobol::ishigami_Fun
  } else if(j == "Sobol'G") {
    k <- 8
    modelRun <- sensobol::sobol_Fun
  } else if(j == "Morris") {
    k <- 20
    modelRun <- sensitivity::morris.fun
  }
  mt[[j]] <- vars_matrices(star.centers = star.centers,
                           params = paste("X", 1:k, sep = ""),
                           h = h)
  Y[[j]] <- modelRun(mt[[j]])
  vars.ind[[j]] <- vars_ti(Y = Y[[j]], star.centers = star.centers,
                           params = paste("X", 1:k, sep = ""), h = h)
}
vars.ind <- rbindlist(vars.ind, idcol = "Function")[
  , estimator:= "vars"] %>%
  setcolororder(., c("estimator", "Function", "Ti", "parameters"))

# PLOT SENSITIVITY INDICES -----
lapply(ind, function(x) rbindlist(x, idcol = "Function")) %>%
  rbindlist(., idcol = "estimator") %>%
  rbind(vars.ind) %>%
  .[, parameters:= factor(parameters, levels = paste("X", 1:20, sep = ""))] %>%
  .[, Function:= factor(Function, levels = test_functions)] %>%
  ggplot(., aes(parameters, Ti, fill = estimator)) +
  geom_bar(stat = "identity",
            position = position_dodge(0.7),
            color = "black") +
  scale_fill_discrete(name = expression(paste("Sobol' ", T[italic(i)]))),
            labels = c("Azzini", "Glen and Isaacs", "Homma and Saltelli",
                      "Jansen", "Lamboni", "Janon / Monod", "Owen",
                      "Sobol", "Razavi and Gupta")) +
  facet_grid(~Function,
            scales = "free_x",
            space = "free_x") +
  labs(x = "",
       y = expression(T[italic(i)])) +

```





## 2 The metaproblem

```
# CREATE METAPROBLEM -----
function_list <- list(
  Linear = function(x) x,
  Quadratic = function(x) x ^ 2,
  Cubic = function(x) x ^ 3,
  Exponential = function(x) exp(1) ^ x / (exp(1) - 1),
  Periodic = function(x) sin(2 * pi * x) / 2,
  Discontinuous = function(x) ifelse(x > 0.5, 1, 0),
  Non.monotonic = function(x) 4 * (x - 0.5) ^ 2,
  Inverse = function(x) (10 - 1 / 1.1) ^ -1 * (x + 0.1) ^ -1,
  No.effect = function(x) x * 0,
  Trigonometric = function(x) cos(x)
)

# PLOT METAPROBLEM -----
a <- ggplot(data.frame(x = runif(100)), aes(x)) +
  map(1:length(function_list), function(nn) {
    stat_function(fun = function_list[[nn]],
                  geom = "line",
                  aes_(color = factor(names(function_list[nn])),
                       linetype = factor(names(function_list[nn]))))
  }) +
  labs(color= "Function", linetype = "Function",
       x = expression(italic(x)),
       y = expression(italic(y))) +
  theme_AP() +
  theme(legend.position = "right")

## Warning: `mapping` is not used by stat_function()
```

```

## Warning: `mapping` is not used by stat_function()

a

```

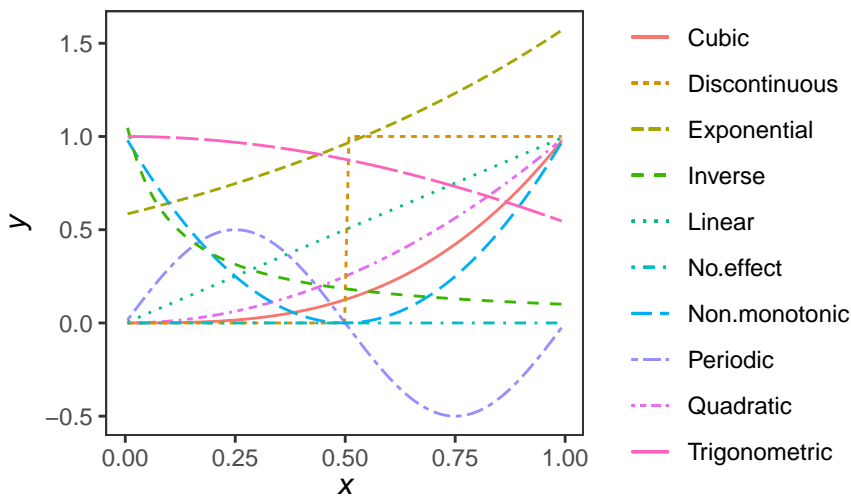


Figure 1: Functions used in the metafunction of Becker (2019).

```

# CREATE FUNCTION FOR RANDOM DISTRIBUTIONS -----
sample_distributions <- list(
  "uniform" = function(x) x,
  "normal" = function(x) qnorm(x, 0.5, 0.2),
  "beta" = function(x) qbeta(x, 8, 2),
  "beta2" = function(x) qbeta(x, 2, 8),
  "beta3" = function(x) qbeta(x, 2, 0.5),
  "beta4" = function(x) qbeta(x, 0.5, 2),
  "logitnormal" = function(x) qlogitnorm(x, 0, 3.16)
  # Logit-normal, Bates too?
)

```

```

random_distributions <- function(X, phi) {
  names_ff <- names(sample_distributions)
  if(!phi == length(names_ff) + 1) {
    out <- sample_distributions[[names_ff[phi]]](X)
  } else {
    temp <- sample(names_ff, ncol(X), replace = TRUE)
    out <- sapply(seq_along(temp), function(x) sample_distributions[[temp[x]]](X[, x]))
  }
  return(out)
}

# PLOT DISTRIBUTIONS -----
names_ff <- names(sample_distributions)
prove <- randtoolbox::sobol(n = 1000, dim = length(names_ff))

out <- data.table(sapply(seq_along(names_ff), function(x)
  sample_distributions[[names_ff[x]]])(prove[, x]))

b <- data.table::melt(out) %>%
  ggplot(., aes(value, group = variable, colour = variable)) +
  geom_density() +
  scale_color_discrete(labels = c("U(0, 1)",
                                    "N(0.5, 0.2)",
                                    "Beta(8, 2)",
                                    "Beta(2, 8)",
                                    "Beta(2, 0.5)",
                                    "Beta(0.5, 2)",
                                    "Logitnormal(0, 3.16)"),
                        name = ""))
  labs(x = expression(italic(x)),
       y = "Density") +
  theme_AP()

## Warning in melt.data.table(out): id.vars and measure.vars are internally
## guessed when both are 'NULL'. All non-numeric/integer/logical type columns are
## considered id.vars, which in this case are columns []. Consider providing at
## least one of 'id' or 'measure' vars in future.

b

# MERGE METAFUNCTION PLOT AND DISTRIBUTIONS PLOT -----
plot_grid(b, a, ncol = 1, labels = "auto", align = "hv")

```

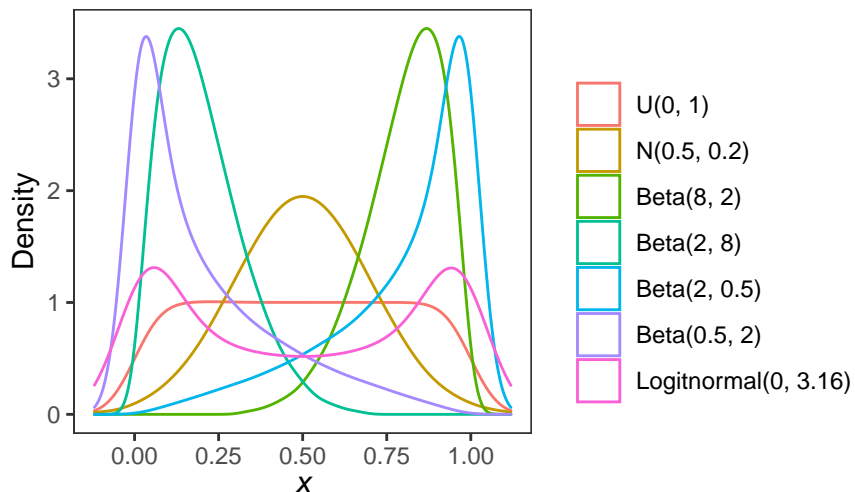
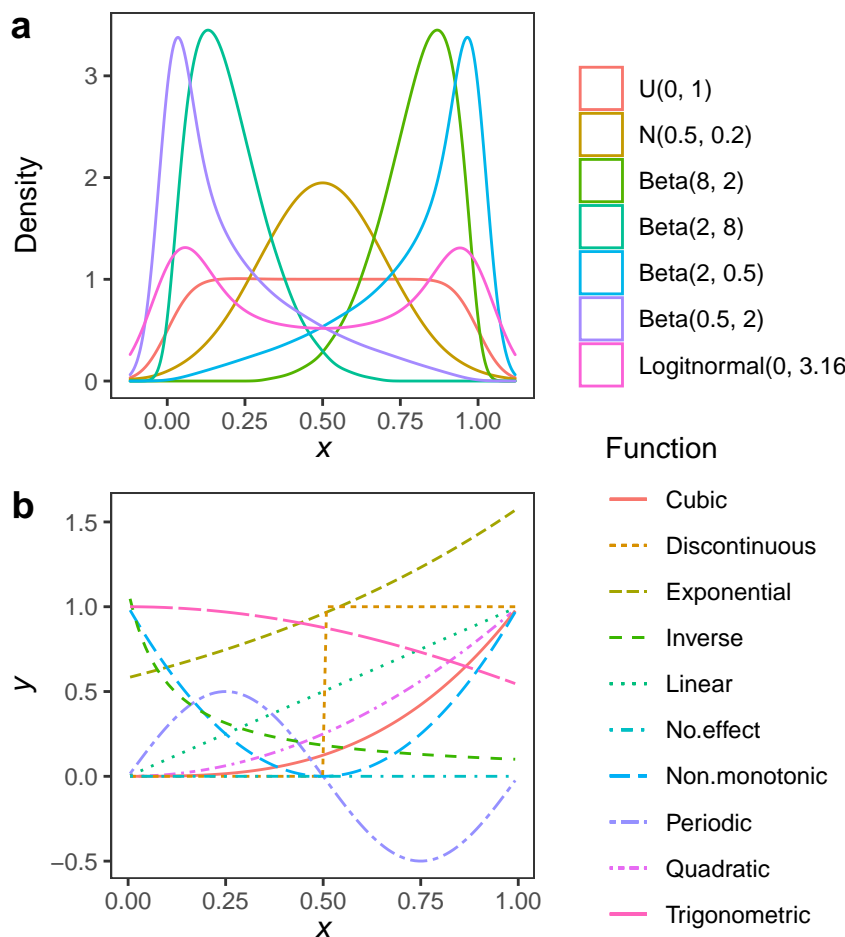


Figure 2: Distributions used in the metafunction of Becker (2019).




---

# EXEMPLIFY THE METAFUNCTION WITH AN EXAMPLE -----

```
N <- 10000 # Sample size
R <- 100 # Number of bootstrap replications
k <- 17 # Number of model inputs
```

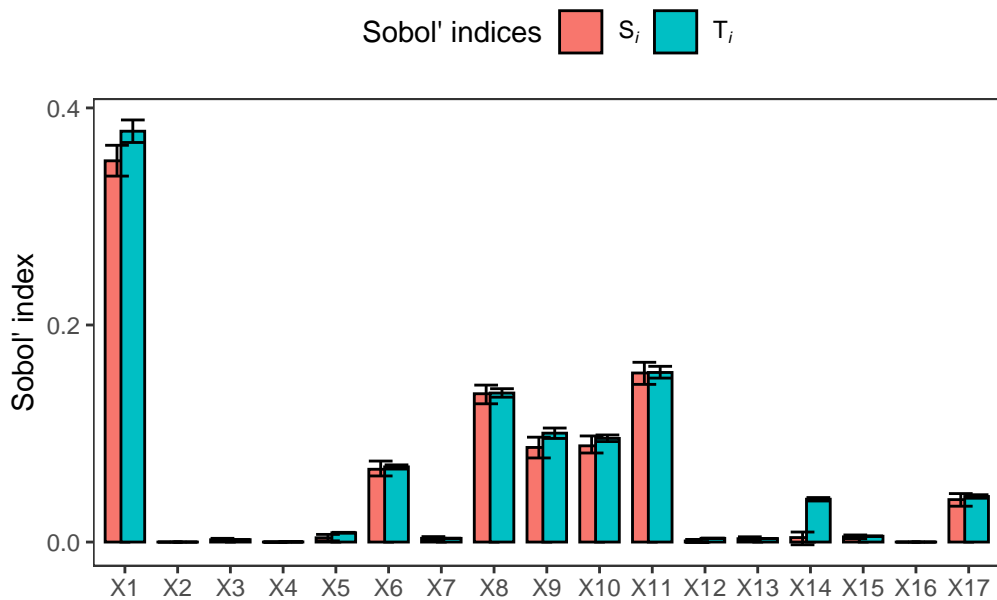
```

k_2 <- 0.5 # Fraction of active pairwise interactions
k_3 <- 0.2 # Fraction of active three-wise interactions
epsilon <- 666 # to reproduce the results
params <- paste("X", 1:k, sep = "")
A <- sobol_matrices(N = N, params = params)

# Compute
Y <- sensobol::metafunction(data = A, k_2 = k_2, k_3 = k_3, epsilon = epsilon)
indices <- data.table(sensobol::sobol_indices(Y = Y, N = N, params = params, boot = TRUE, R = 1000))
.[, parameters:= factor(parameters, levels = paste("X", 1:k, sep = ""))]

ggplot(indices, aes(parameters, original, fill = sensitivity)) +
  geom_bar(stat = "identity",
            position = position_dodge(0.6),
            color = "black") +
  geom_errorbar(aes(ymin = low.ci,
                     ymax = high.ci),
                position = position_dodge(0.6)) +
  scale_x_discrete(labels = ggplot2:::parse_safe) +
  scale_y_continuous(breaks = pretty_breaks(n = 3)) +
  labs(x = "",
       y = "Sobol' index") +
  scale_fill_discrete(name = "Sobol' indices",
                      labels = c(expression(S[italic(i)]),
                                expression(T[italic(i)]))) +
  theme_AP() +
  theme(legend.position = "top")

```



### 3 The model

#### 3.1 Settings

```
# DEFINE SETTINGS ----

N <- 2 ^ 11 # Sample size of sample matrix
h <- 0.2 # step for VARS
R <- 500 # Number of bootstrap replicas
n_cores <- ceiling(detectCores() * 0.5)
order <- "second"
params <- c("k_2", "k_3", "epsilon", "phi", "delta", "tau")
N.high <- 2 ^ 11 # Maximum sample size of the large sample matrix
```

#### 3.2 Sample matrix

```
# CREATE SAMPLE MATRIX ----

# Main matrix without Nt and k
tmp <- sobol_matrices(N = N, params = c(params, "N_t", "k"), order = order, matrices = c("A",
A <- tmp[1:N, ]
B <- tmp[(N + 1):(2 * N), ]

first <- 1:length(params)
loop <- c(first, utils::combn(1:length(params), 2, simplify = FALSE))

X <- rbind(A, B)
for(i in loop) {
  AB <- A
  AB[, i] <- B[, i]
  X <- rbind(X, AB)
}
mat <- X

# Matrix with clusters (Nt,k)
AB <- AB2 <- AB3 <- A
AB[, c("N_t", "k")] <- B[, c("N_t", "k")] # (Nt,k)
AB2[, c("k_2", "k_3", "epsilon", "phi", "N_t", "k")] <- B[, c("k_2", "k_3", "epsilon", "phi",
AB3[, c("delta", "tau")] <- B[, c("delta", "tau")] # Manageable uncertainties

# Final matrix
mat <- rbind(mat, AB, AB2, AB3)

# TRANSFORM MATRIX ----

mat[, 1] <- round(qunif(mat[, 1], 0.3, 0.5), 2) # k_2
mat[, 2] <- round(qunif(mat[, 2], 0.1, 0.3), 2) # k_3
```

```

mat[, 3] <- floor(qunif(mat[, 3], 1, 200)) # Epsilon
mat[, 4] <- floor(mat[, 4] * (8 - 1 + 1)) + 1 # Phi
mat[, 5] <- floor(mat[, 5] * (3 - 1 + 1)) + 1 # Delta
mat[, 6] <- floor(mat[, 6] * (2 - 1 + 1)) + 1 # Tau
mat[, 7] <- floor(qunif(mat[, 7], 10, 1000)) # Nt
mat[, 8] <- floor(qunif(mat[, 8], 3, 100)) # k

# Define the base sample matrix
N.all <- apply(mat, 1, function(x) ceiling(x["N_t"] / (x["k"] + 1)))
N.azzini <- apply(mat, 1, function(x) ceiling(x["N_t"] / (2 * x["k"] + 2)))
N.vars <- apply(mat, 1, function(x) floor(x["N_t"] / ((x["k"]) * ((1 / h) - 1) + 1)))

tmp <- data.table(cbind(mat, N.all, N.azzini, N.vars))

# Now constrain N as a function of vars
tmp <- tmp[, N.all:= ifelse(N.vars == 1 | N.vars == 0, ceiling(2 * (4 * k + 1) / (k + 1)), N.all)]
tmp <- tmp[, N.azzini:= ifelse(N.vars == 1 | N.vars == 0, ceiling(2 * (4 * k + 1) / (2 * k + 2)), N.azzini)]
tmp <- tmp[, N.vars:= ifelse(N.vars == 1 | N.vars == 0, 2, N.vars)]

C.all <- apply(tmp, 1, function(x) x["N.all"] * (x["k"] + 1))
C.azzini <- apply(tmp, 1, function(x) x["N.azzini"] * (2 * x["k"] + 2))
C.vars <- apply(tmp, 1, function(x) x["N.vars"] * (x["k"] * ((1 / h) - 1) + 1))

mat <- cbind(tmp, C.all, C.azzini, C.vars)

```

### 3.3 Define the model

```

# DEFINE MODEL ----

model_Ti <- function(k, N.all, N.azzini, N.vars, h,
                      N.high, k_2, k_3, epsilon, phi, delta, tau) {
  ind <- list()
  estimators <- c("jansen", "sobol", "homma", "monod", "azzini",
                  "lamboni", "glen", "owen", "vars")
  if(tau == 1) {
    method <- "R"
  } else if(tau == 2) {
    method <- "QRN"
  }
  all.but.azzini <- sobol_matrices(N = N.all, params = paste("X", 1:k, sep = ""),
                                    matrices = c("A", "AB"), method = method)
  azzini <- sobol_matrices(N = N.azzini, params = paste("X", 1:k, sep = ""),
                           matrices = c("A", "B", "AB", "BA"), method = method)
  owen.matrix <- sobol_matrices(N = N.azzini, params = paste("X", 1:k, sep = ""),
                                 matrices = c("A", "B", "BA", "CB"), method = method)
  vars.matrix <- vars_matrices(star.centers = N.vars, params = paste("X", 1:k, sep = ""),
                                h = h, method = method)

```

```

large.matrix <- sobol_matrices(N = N.high, params = paste("X", 1:k, sep = ""),
                                matrices = c("A", "AB"), method = method)
set.seed(epsilon)
all.matrices <- random_distributions(X = rbind(all.but.azzini, azzini,
                                                owen.matrix, vars.matrix,
                                                large.matrix),
                                       phi = phi)
output <- sensobol::metafunction(data = all.matrices,
                                   k_2 = k_2,
                                   k_3 = k_3,
                                   epsilon = epsilon)
full.ind <- sobol_Ti(d = tail(output, nrow(large.matrix)),
                      N = N.high,
                      params = paste("X", 1:k, sep = ""),
                      total = "jansen")
full.ind[, sample.size:= "N"]

# Define indices of Y for estimators
Nt.all.but.azzini <- N.all * (k + 1)
Nt.azzini.owen <- N.azzini * ((2 * k) + 2)
Nt.vars <- N.vars * (k * ((1 / h) - 1) + 1)
lg.all.but.azzini <- 1:Nt.all.but.azzini
lg.azzini <- (length(lg.all.but.azzini) + 1):(length(lg.all.but.azzini) + Nt.azzini.owen)
lg.owen <- (max(lg.azzini) + 1):(max(lg.azzini) + Nt.azzini.owen)
lg.vars <- (max(lg.owen) + 1): (max(lg.owen) + Nt.vars)

for(i in estimators) {
  if(i == "jansen" | i == "sobel" | i == "homma" | i == "monod" | i == "glen") {
    y <- output[lg.all.but.azzini]
    n <- N.all
  } else if(i == "azzini" | i == "lamboni") {
    y <- output[lg.azzini]
    n <- N.azzini
  } else if(i == "owen") {
    y <- output[lg.owen]
    n <- N.azzini
  } else if(i == "vars") {
    y <- output[lg.vars]
    star.centers <- N.vars
  }
  if(!i == "vars") {
    ind[[i]] <- sobol_Ti(d = y, N = n, params = paste("X", 1:k, sep = ""), total = i)
  }
  if(i == "vars") {
    ind[[i]] <- vars_ti(Y = y, star.centers = star.centers,
                           params = paste("X", 1:k, sep = ""), h = h)
  }
}

```

```

    ind[[i]][, sample.size:= "n"]
    ind[[i]] <- rbind(ind[[i]], full.ind)
}
# Arrange data
out <- rbindlist(ind, idcol = "estimator")
out.wide <- dcast(out, estimator + parameters ~ sample.size, value.var = "Ti")
# Replace NaN
for (i in seq_along(out.wide))
  set(out.wide, i=which(is.nan(out.wide[[i]])), j = i, value = 0)
# Replace Inf
for (i in seq_along(out.wide))
  set(out.wide, i=which(is.infinite(out.wide[[i]])), j = i, value = 0)
# Replace Na
for (i in seq_along(out.wide))
  set(out.wide, i=which(is.na(out.wide[[i]])), j = i, value = 0)
# CHECK DELTA
if(delta == 1) { # Regular Pear
  final <- out.wide[, .(correlation = cor(N, n)), estimator]
} else if(delta == 2) { # kendall tau
  final <- out.wide[, .(correlation = pcaPP::cor.fk(N, n)), estimator]
} else { # Savage ranks
  final <- out.wide[, lapply(.SD, savage_scores), .SDcols = c("N", "n"), estimator][
    , .(correlation = cor(N, n)), estimator]
}
return(final)
}

```

### 3.4 Run the model

```

# RUN MODEL -----
# Define parallel computing
cl <- makeCluster(n_cores)
registerDoParallel(cl)

# Compute
Y.ti <- foreach(i=1:nrow(mat),
  .packages = c("sensobol", "data.table", "pcaPP",
    "logitnorm", "dplyr")) %dopar%
{
  model_Ti(k = mat[[i, "k"]],
    k_2 = mat[[i, "k_2"]],
    k_3 = mat[[i, "k_3"]],
    epsilon = mat[[i, "epsilon"]],
    phi = mat[[i, "phi"]],
    delta = mat[[i, "delta"]],
    tau = mat[[i, "tau"]]],

```

```

        N.all = mat[[i, "N.all"]],
        N.azzini = mat[[i, "N.azzini"]],
        N.vars = mat[[i, "N.vars"]],
        N.high = N.high,
        h = h)
    }

# Stop parallel cluster
stopCluster(cl)

```

### 3.5 Arrange output

```

# ARRANGE OUTPUT -----
out_cor <- rbindlist(Y.ti, idcol = "row")

mt.dt <- data.table(mat) %>%
  .[, row:= 1:.N]

full_output <- merge(mt.dt, out_cor) %>%
  .[, Nt:= ifelse(estimator == "azzini" | estimator == "lamboni"
    | estimator == "owen", N.azzini * (2 * k + 2),
    ifelse(estimator == "vars", N.vars * (k * ((1 / h) -1) + 1), N.all * (k + 1))
  .[, estimator:= ifelse(estimator %in% "azzini", "Azzini and Rosati",
    ifelse(estimator %in% "homma", "Homma and Saltelli",
    ifelse(estimator %in% "monod", "Janon/Monod",
    ifelse(estimator %in% "jansen", "Jansen",
    ifelse(estimator %in% "glen", "Glen and Isaac",
    ifelse(estimator %in% "lamboni", "Lamboni"
    ifelse(estimator %in% "owen", "Owen
    ifelse(estimator %in% "vars"
      "Sobol'")))))))] %>%
  .[!estimator == "Lamboni"]

# Define A matrix (but drop Lamboni)
A <- full_output[!estimator == "Lamboni", .SD[1:N], estimator]

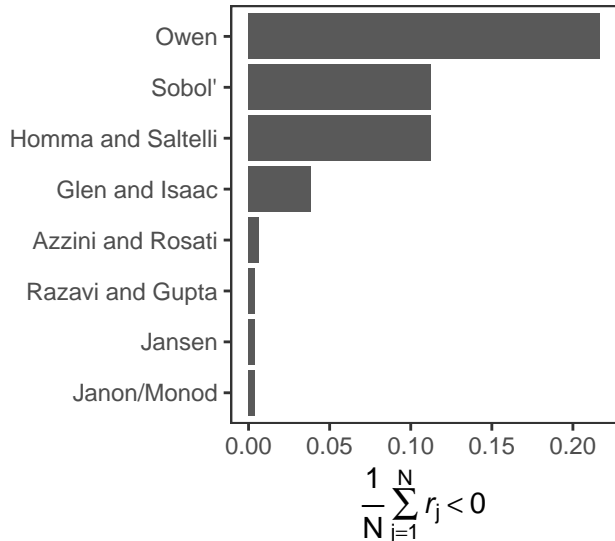
# EXPORT OUTPUT -----
fwrite(A, "A.csv")
fwrite(full_output, "full_output.csv")

```

## 4 Uncertainty analysis

```
# PLOT PROPORTION OF NEGATIVE VALUES -----
```

```
A[, sum(correlation < 0) / .N, estimator] %>%
  ggplot(., aes(reorder(estimator, V1), V1)) +
  geom_bar(stat = "identity") +
  coord_flip() +
  labs(y = expression(frac(1, N) ~ sum(italic(r)[j] < 0, j == 1, N)),
       x = "") +
  theme_AP()
```

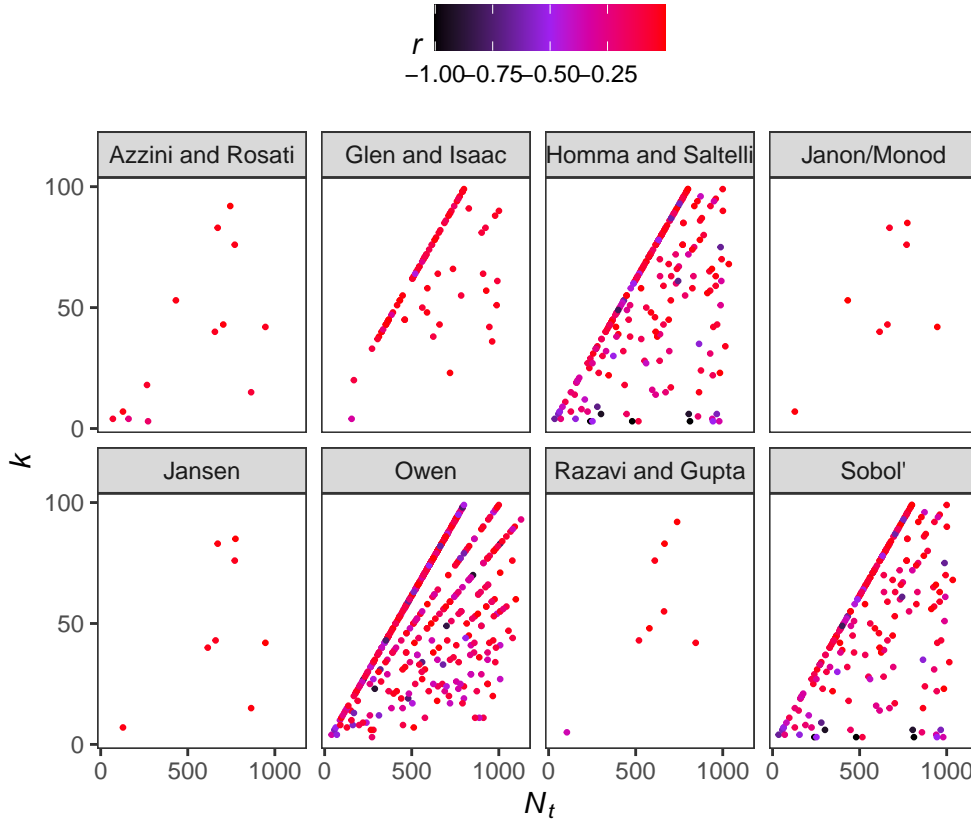



---

```
# MAP VALUES WITH NEGATIVE R ----
```

```
index.neg <- A[, .I[correlation < 0]]

A[index.neg] %>%
  ggplot(., aes(Nt, k, color = correlation)) +
  geom_point(size = 0.5) +
  scale_colour_gradientn(colours = c("black", "purple", "red"),
                         name = expression(italic(r))) +
  scale_x_continuous(breaks = pretty_breaks(n = 3)) +
  scale_y_continuous(breaks = pretty_breaks(n = 3)) +
  labs(x = expression(italic(N[t])),
       y = expression(italic(k))) +
  facet_wrap(~estimator,
             ncol = 4) +
  theme_AP() +
  theme(legend.position = "top")
```



```

# PLOT OUTPUT -----
# Compute median and quantiles
dt_median <- A[, .(median = median(correlation),
                  low.ci = quantile(correlation, 0.25),
                  high.ci = quantile(correlation, 0.75)), estimator]

A[, .(median = median(correlation),
      low.ci = quantile(correlation, 0.25),
      high.ci = quantile(correlation, 0.75)), estimator][order(median)]

##          estimator     median    low.ci   high.ci
## 1:           Owen 0.2283349 0.02561785 0.4918428
## 2: Homma and Saltelli 0.3335354 0.11428571 0.5854888
## 3:           Sobol' 0.3335354 0.11428571 0.5854888
## 4: Glen and Isaac 0.4059889 0.19739556 0.6444691
## 5: Azzini and Rosati 0.7790664 0.61657561 0.9081717
## 6: Janon/Monod 0.8694113 0.75568980 0.9503799
## 7:           Jansen 0.8736469 0.75963825 0.9551326
## 8: Razavi and Gupta 0.9156334 0.81183826 0.9717718

a <- ggplot(A, aes(correlation)) +
  geom_rect(data = dt_median,
            aes(xmin = low.ci,
                xmax = high.ci,

```

```

        ymin = -Inf,
        ymax = Inf),
    fill = "blue",
    color = "white",
    alpha = 0.1,
    inherit.aes = FALSE) +
geom_histogram() +
geom_vline(data = dt_median, aes(xintercept = median),
    lty = 2,
    color = "red") +
facet_wrap(~estimator,
    ncol = 4) +
scale_x_continuous(breaks = pretty_breaks(n = 3)) +
scale_y_continuous(breaks = pretty_breaks(n = 3)) +
labs(x = expression(italic(r)),
    y = "Counts") +
theme_AP() +
theme(axis.text.x = element_text(angle = 45, hjust = 1))

# Scatterplot
b <- ggplot(A, aes(Nt, k, color = correlation)) +
    geom_point(size = 0.1) +
    scale_colour_gradientn(colours = c("black", "purple", "red", "orange", "lightgreen"),
        name = expression(italic(r))) +
    scale_x_continuous(breaks = pretty_breaks(n = 3)) +
    scale_y_continuous(breaks = pretty_breaks(n = 3)) +
    labs(x = expression(italic(N[t])),
        y = expression(italic(k))) +
    facet_wrap(~estimator,
        ncol = 4) +
    theme_AP() +
    theme(legend.position = "top")

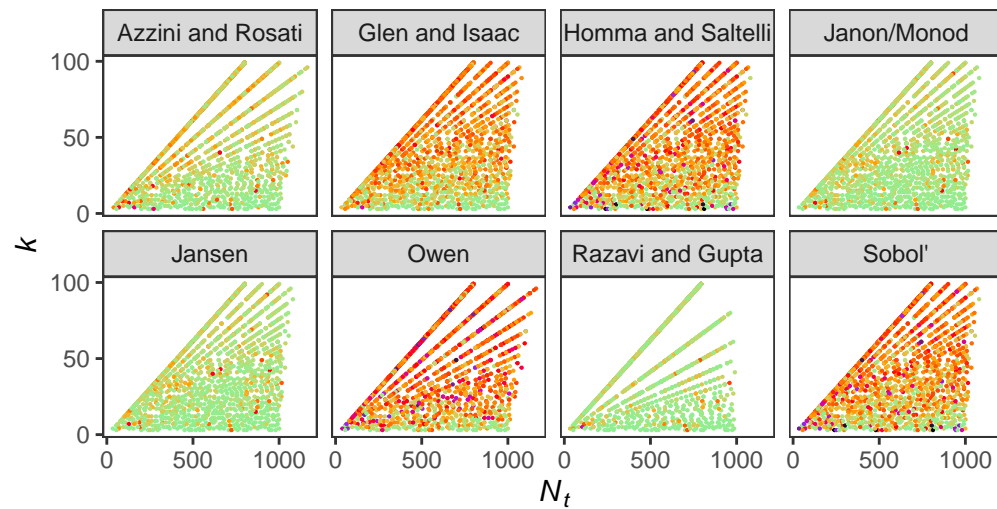
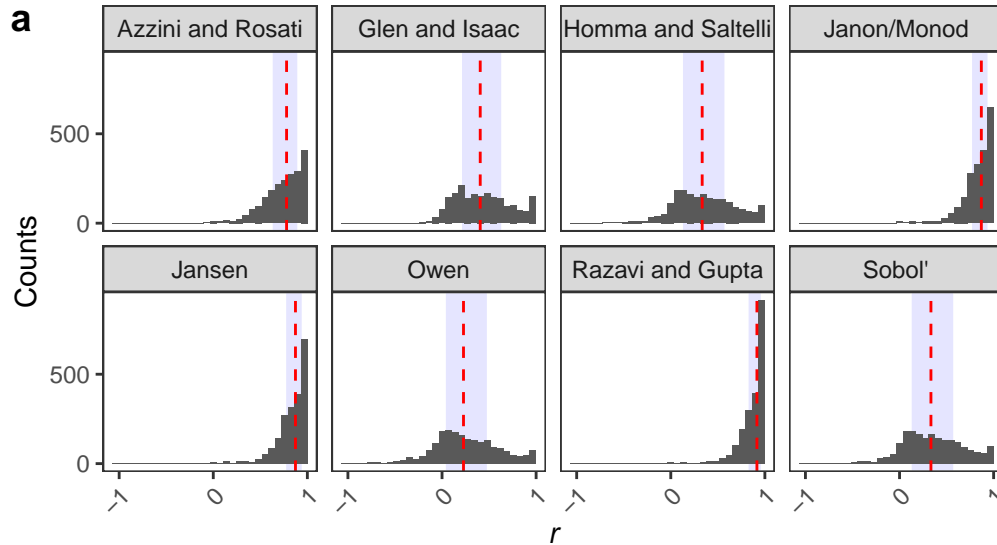
# Ratio
c <- A[, ratio:= Nt / k] %>%
    ggplot(., aes(ratio, correlation)) +
    geom_point(alpha = 0.1, size = 0.2) +
    facet_wrap(~estimator,
        ncol = 4) +
    geom_smooth() +
    labs(x = expression(italic(N[t]/k)),
        y = expression(italic(r))) +
    scale_x_log10() +
    scale_y_continuous(breaks = pretty_breaks(n = 3)) +
    theme_AP()

# Merge plot

```

```
plot_grid(a, b, ncol = 1, labels = "auto", rel_heights = c(0.85, 1))
```

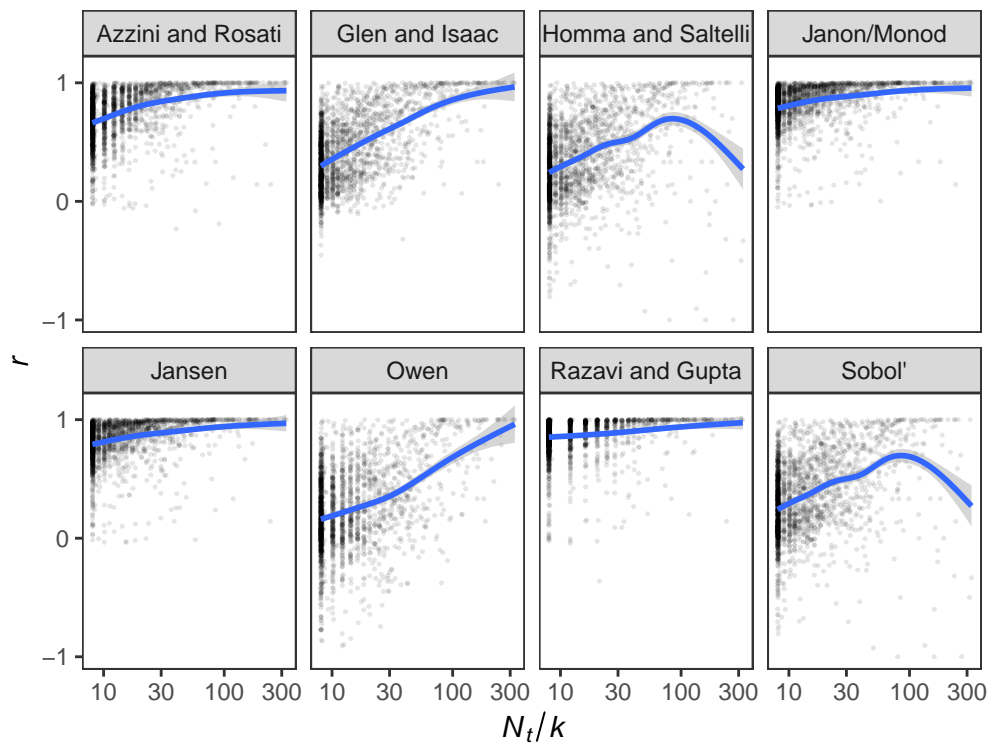
## `stat\_bin()` using `bins = 30` . Pick better value with `binwidth` .



# DISPLAY THE RATIO  $N_t/k$  FOR EACH SIMULATION AND ESTIMATOR -----

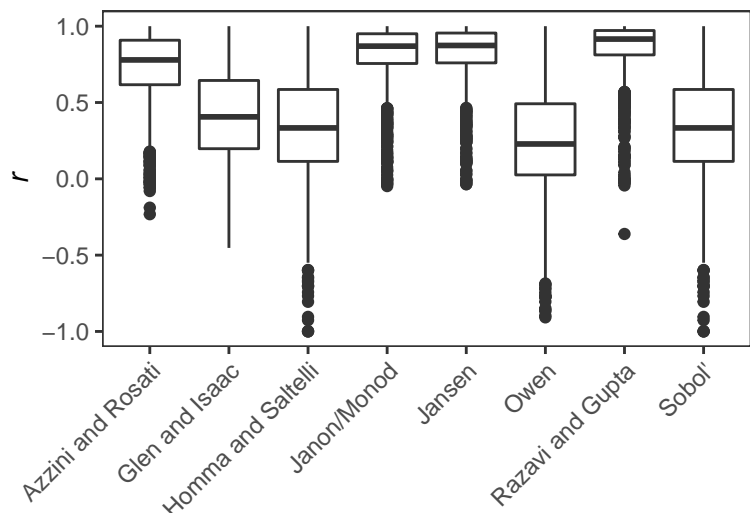
**c**

```
## `geom_smooth()` using method = 'gam' and formula 'y ~ s(x, bs = "cs")'
```



# PLOT BOXPLOT -----

```
ggplot(A, aes(estimator, correlation)) +
  geom_boxplot() +
  labs(x = "",
       y = expression(italic(r))) +
  theme_AP() +
  theme(axis.text.x = element_text(angle = 45, hjust = 1))
```



# PLOT MEDIANS -----

```
vv <- seq(5, 100, 5)
```

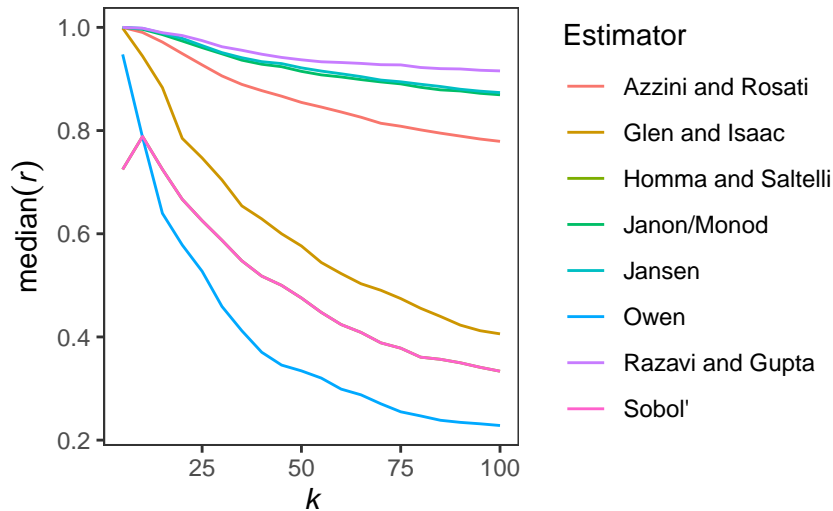
```

dt <- lapply(vv, function(x) A[k <= x, median(correlation), estimator])

names(dt) <- vv

rbindlist(dt, idcol = "k") %>%
  .[, k:= as.numeric(k)] %>%
  ggplot(., aes(k, V1, color = estimator)) +
  scale_color_discrete(name = "Estimator") +
  labs(x = expression(italic(k)),
       y = expression(median(italic(r)))) +
  geom_line() +
  theme_AP()

```



```

# Median Nt/k
dt.tmp <- A[, .(min = min(ratio), max = max(ratio))]

v <- seq(0, ceiling(dt.tmp$max), 20)
a <- c(v[1], rep(v[-c(1, length(v))], each = 2), v[length(v)])
indices <- matrix(a, ncol = 2 ,byrow = TRUE)

out <- list()
for(i in 1:nrow(indices)) {
  out[[i]] <- A[ratio > indices[i, 1] & ratio < indices[i, 2]]
}

names(out) <- rowmeans(indices)

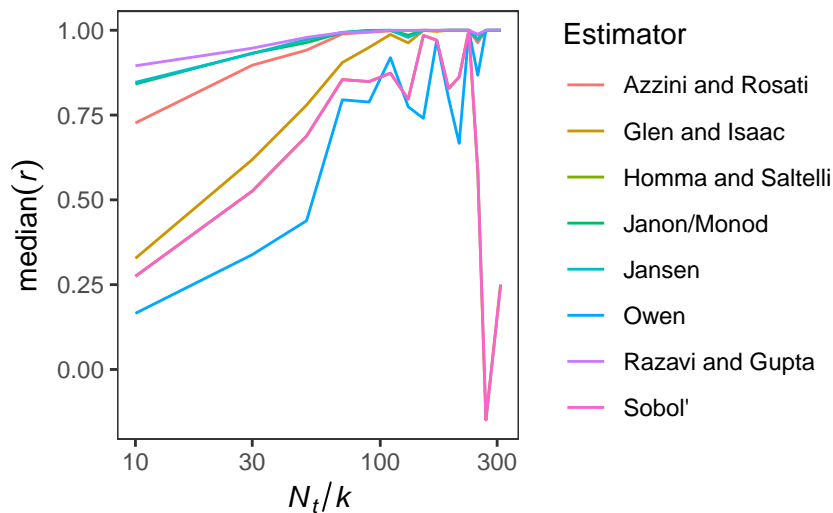
# Plot
lapply(out, function(x) x[, median(correlation, na.rm = TRUE), estimator]) %>%
  rbindlist(., idcol = "N") %>%
  .[, N:= as.numeric(N)] %>%
  ggplot(., aes(N, V1, group = estimator, color = estimator)) +

```

```

geom_line() +
  labs(x = expression(italic(N[t])/k)),
  y = expression(median(italic(r)))) +
  scale_color_discrete(name = "Estimator") +
  scale_x_log10() +
  theme_AP()

```



## 5 Sensitivity analysis

### 5.1 Scatterplots

```

# SCATTERPLOTS OF MODEL OUTPUT AGAINST PARAMETERS ----

A <- setnames(A, c("N_t", "k_2", "k_3"), c("N[t]", "k[2]", "k[3]"))

scatter.dt <- melt(A, measure.vars = c("k", "N[t]", "k[2]", "k[3]",
                                         "epsilon", "phi", "delta", "tau")) %>%
  split(., .$estimator)

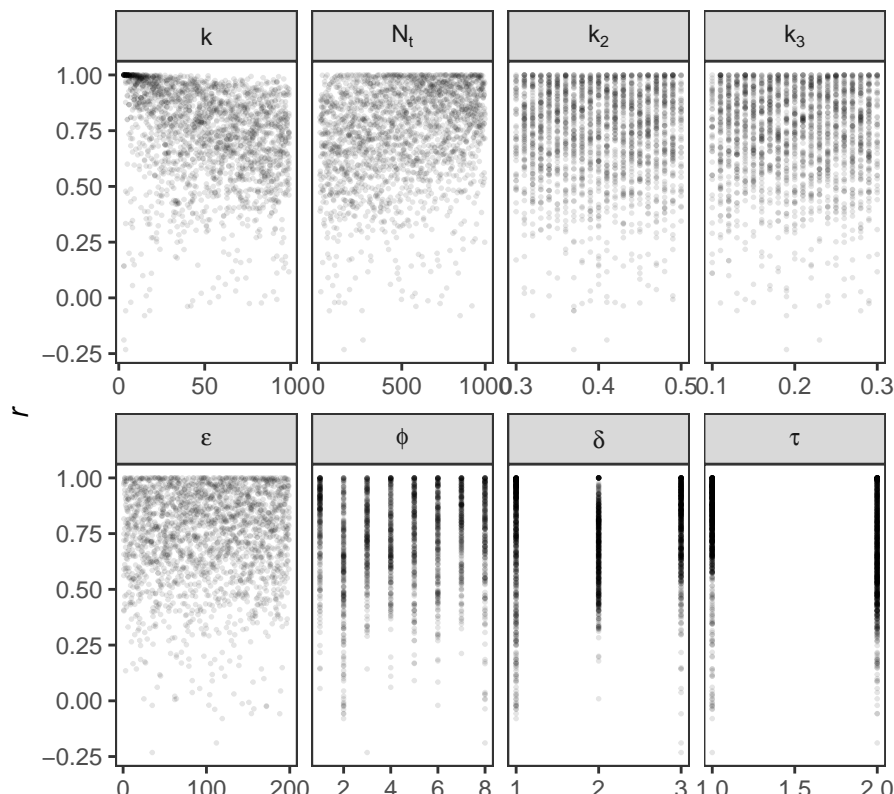
gg <- list()
for(i in names(scatter.dt)) {
  gg[[i]] <- ggplot(scatter.dt[[i]], aes(value, correlation)) +
    geom_point(alpha = 0.1, size = 0.3) +
    facet_wrap(~ variable,
               scales = "free_x",
               labeller = label_parsed,
               ncol = 4) +
    scale_color_manual(values = c("#00BFC4", "#F8766D")) +
    scale_x_continuous(breaks = pretty_breaks(n = 3)) +
    theme_AP() +
    labs(x = "", y = expression(italic(r))) +
    theme()
}

```

```
    ggttitle(names(scatter.dt[i]))  
}  
  
gg
```

```
## $`Azzini and Rosati`
```

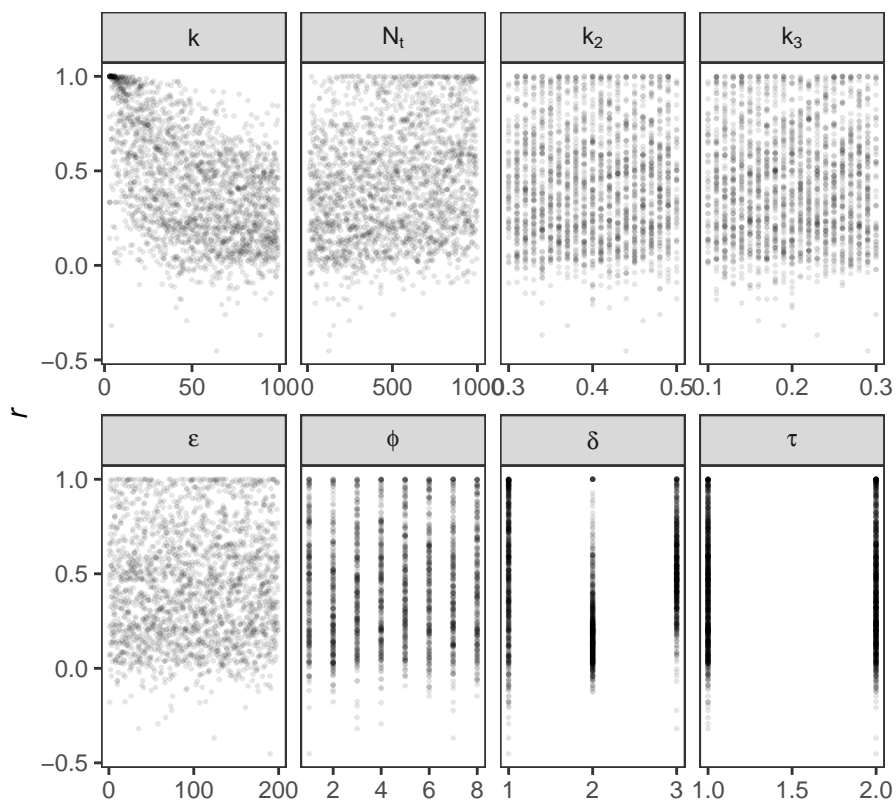
Azzini and Rosati



```
##
```

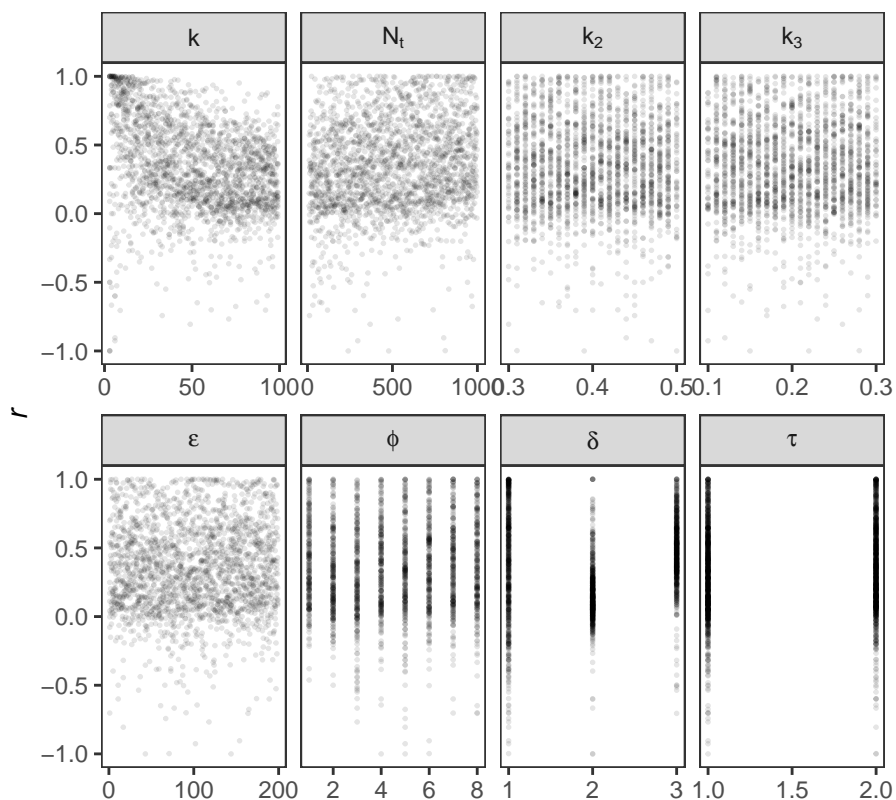
```
## $`Glen and Isaac`
```

## Glen and Isaac



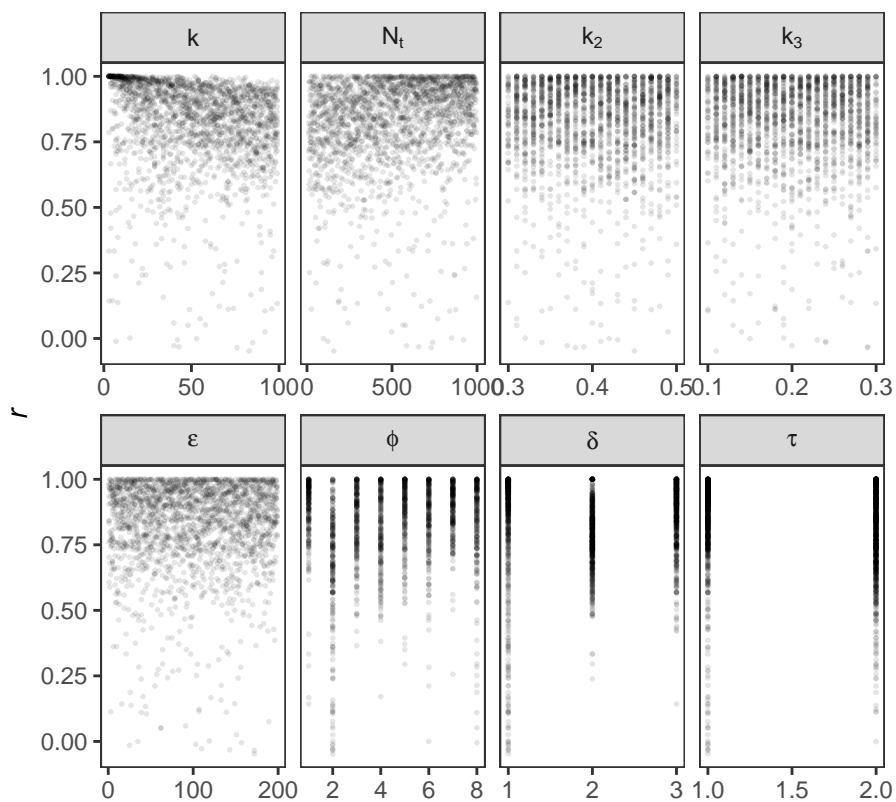
```
##  
## $`Homma and Saltelli`
```

## Homma and Saltelli



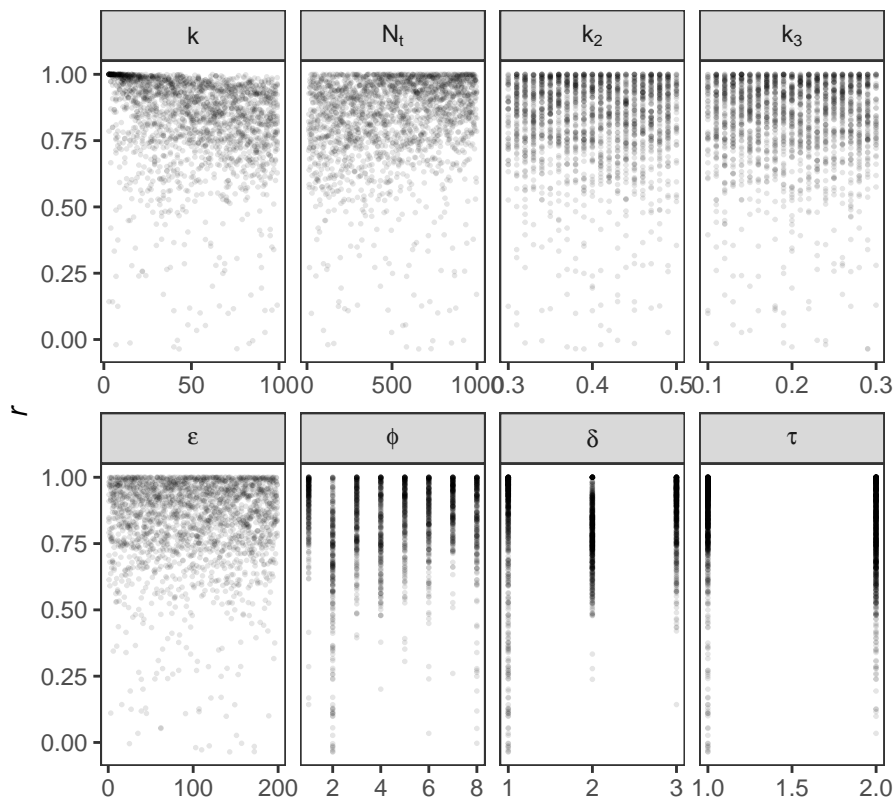
```
##  
## $`Janon/Monod`
```

## Janon/Monod



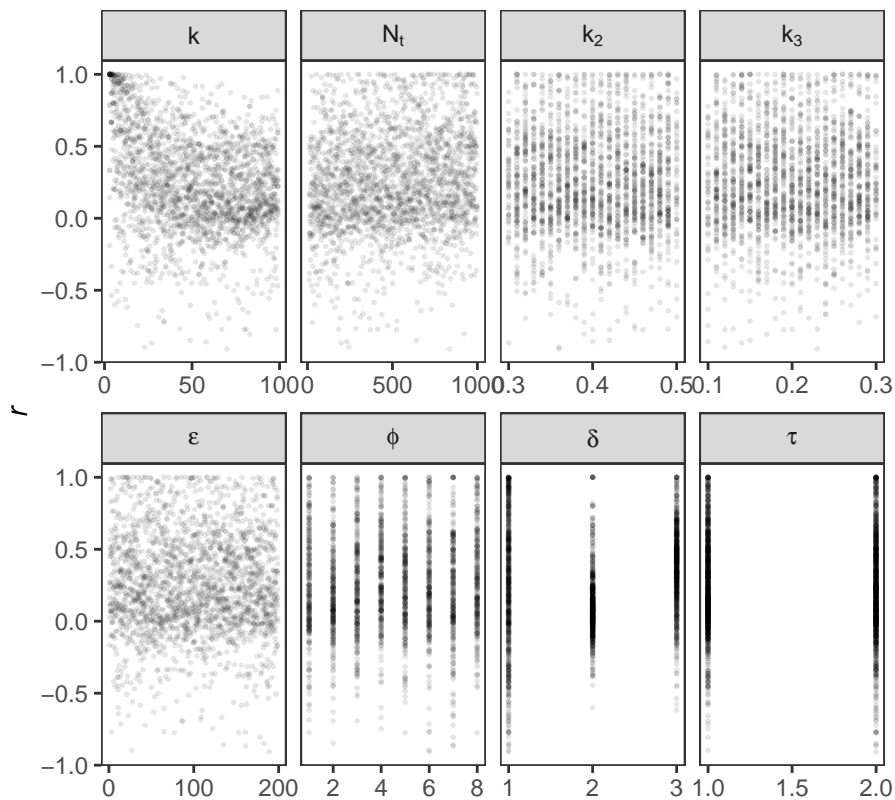
```
##  
## $Jansen
```

Jansen



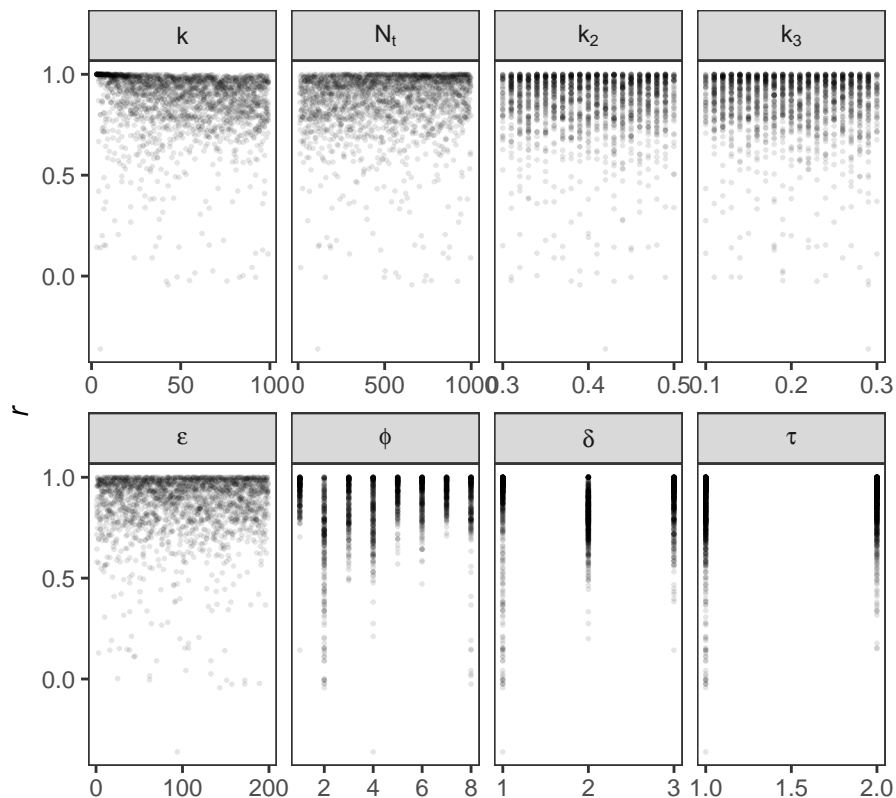
```
##  
## $Owen
```

Owen

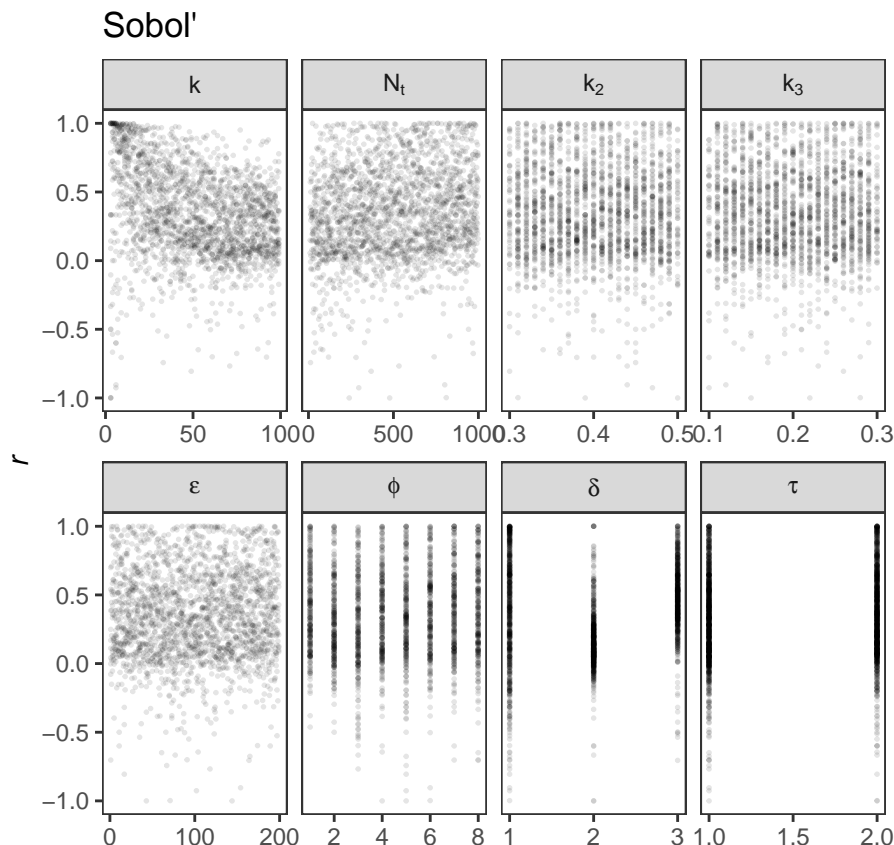


```
##  
## $`Razavi and Gupta`
```

## Razavi and Gupta



```
##  
## $`Sobol`
```



## 5.2 Sobol' indices

```
# SENSITIVITY ANALYSIS -----
# Show rows with NA
full_output[is.na(correlation), ]

## Empty data.table (0 rows and 18 cols): row,k_2,k_3,epsilon,phi,delta...
# Substitute NA by 0
full_output <- full_output[, correlation:= ifelse(is.na(correlation) == TRUE, 0, correlation)]

# New vector of parameters to plot better
params.new <- c("k[2]", "k[3]", "epsilon", "phi", "delta", "tau")

# Compute Sobol' indices except for the cluster Nt,k
indices <- full_output[!row %in% mat[, tail(.I, N)]][
  , sobol_indices(Y = correlation,
    N = N,
    params = params.new,
    first = "jansen",
    boot = TRUE,
    R = R,
    order = order),
```

```

estimator]

# Compute Sobol' indices for cluster
index.clusters <- mat[, tail(.I, 3 * N)]

indicesC <- rbind(full_output[row %in% 1:(2 * N)], full_output[row %in% index.clusters])[,
  sobol_indices(Y = correlation,
    N = N,
    params = c("N[t]~k", "f(x)", "delta~tau"),
    first = "jansen",
    boot = TRUE,
    R = R,
    order = "first"),
  estimator]

# Final
indices <- rbind(indices, indicesC[parameters == "N[t]~k"])

# Export
fwrite(indices, "indices.csv")

# PLOT SOBOL' INDICES -----
all.indices <- list(indices, indicesC)

gg <- list()
for(i in seq_along(all.indices)) {
  gg[[i]] <- ggplot(all.indices[[i]][sensitivity == "Si" |
                                         sensitivity == "Ti"],
    aes(parameters, original, fill = sensitivity)) +
    geom_bar(stat = "identity",
      position = position_dodge(0.6),
      color = "black") +
    geom_errorbar(aes(ymin = low.ci,
                      ymax = high.ci),
      position = position_dodge(0.6)) +
    scale_x_discrete(labels = ggplot2:::parse_safe) +
    scale_y_continuous(breaks = pretty_breaks(n = 3)) +
    facet_wrap(~estimator,
      ncol = 4) +
    labs(x = "",
      y = "Sobol' index") +
    scale_fill_discrete(name = "Sobol' indices",
      labels = c(expression(S[italic(i)]),
                expression(T[italic(i)]))) +
    theme_AP() +
    theme(legend.position = "top")
}

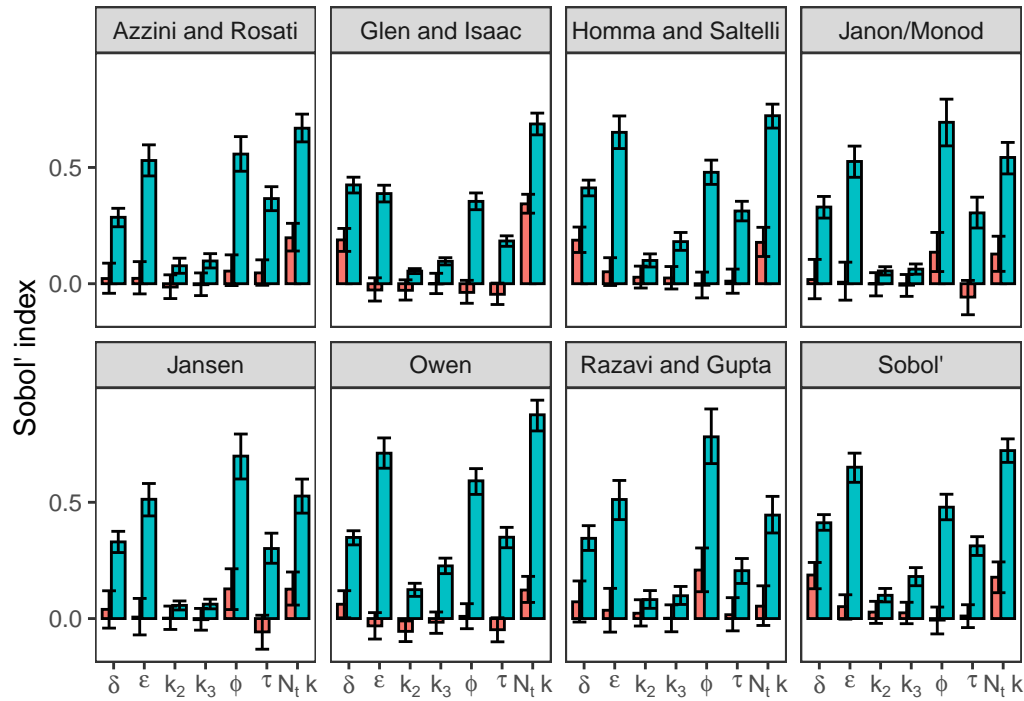
```

```
}
```

```
gg
```

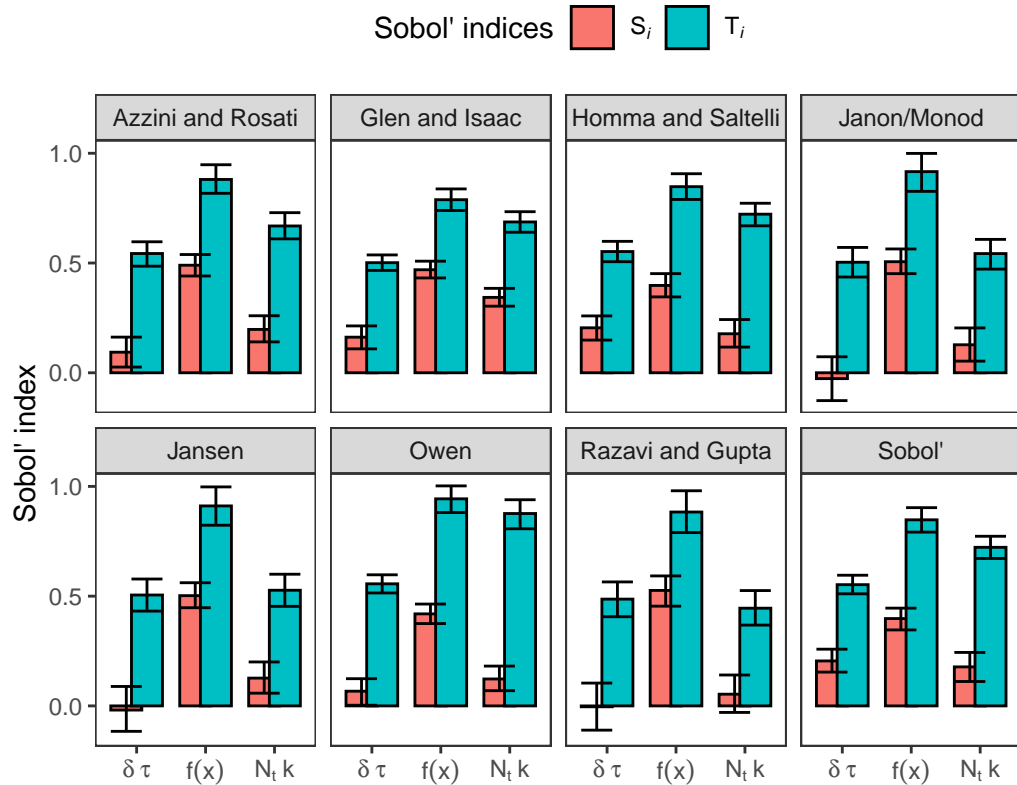
```
## [[1]]
```

Sobol' indices   $S_i$    $T_i$



```
##
```

```
## [[2]]
```



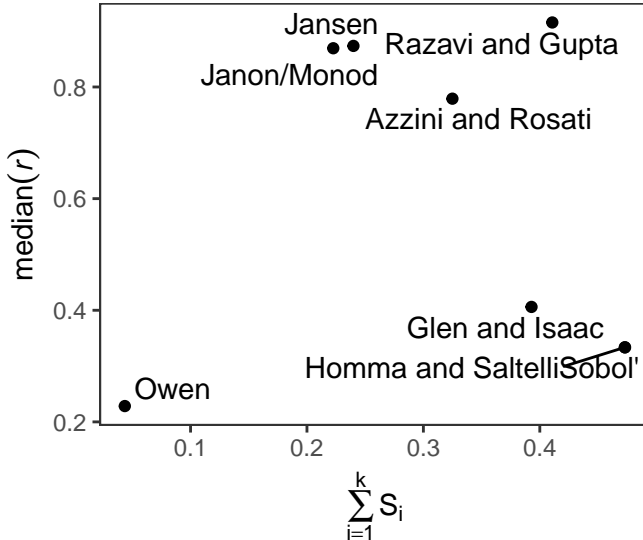
# SUM OF FIRST-ORDER INDICES -----

```
indices[sensitivity == "Si", sum(original), estimator]
```

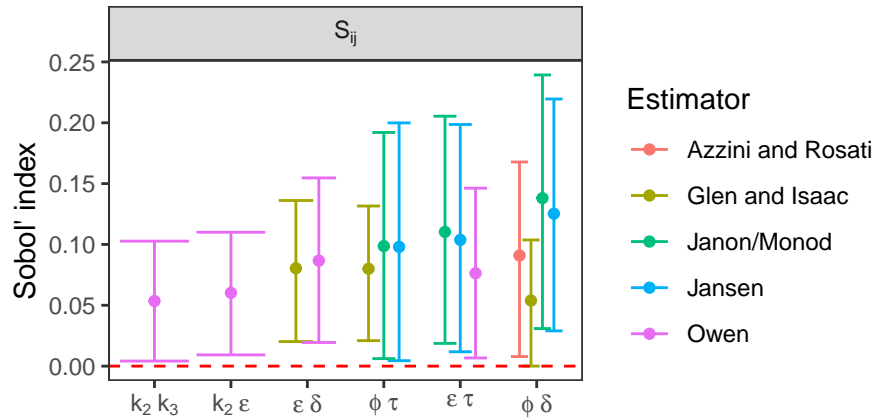
```
##             estimator      V1
## 1: Azzini and Rosati 0.32492667
## 2: Glen and Isaac 0.39285762
## 3: Homma and Saltelli 0.47299950
## 4: Jansen 0.23989969
## 5: Janon/Monod 0.22259432
## 6: Owen 0.04353922
## 7: Sobol' 0.47320580
## 8: Razavi and Gupta 0.41072467
```

# Plot

```
merge(indices[sensitivity == "Si", sum(original), estimator],
      dt_median, by = "estimator") %>%
  ggplot(., aes(V1, median)) +
  geom_point() +
  labs(x = expression(sum(S[i], i==1, k)),
       y = expression(median(italic(r)))) +
  geom_text_repel(aes(label = estimator)) +
  theme_AP()
```



```
# PLOT SECOND-ORDER EFFECTS -----
indices[sensitivity == "Sij"] %>%
  .[low.ci > 0] %>%
  .[, sensitivity:= ifelse(sensitivity %in% "Sij", "S[ij]", sensitivity)] %>%
  .[, parameters:= gsub(parameters,
    pattern = ".",
    replacement = "~",
    fixed = TRUE)] %>%
ggplot(., aes(reorder(parameters, original), original, color = estimator)) +
  geom_point(position = position_dodge(0.6)) +
  geom_errorbar(aes(ymax = high.ci, ymin = low.ci),
    position = position_dodge(0.6)) +
  geom_hline(yintercept = 0,
    lty = 2,
    color = "red") +
  facet_grid(~sensitivity,
    scales = "free_x",
    space = "free_x",
    labeller = label_parsed) +
  scale_x_discrete(labels = ggplot2:::parse_safe) +
  scale_color_discrete(name = "Estimator") +
  labs(x = "",
    y = "Sobol' index") +
  theme_AP()
```



```
# EXPORT SOBOL' INDICES -----
```

```
fwrite(dt_median, "dt_median.csv")
```

## 6 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 10.15.4  
##  
## 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] parallel stats      graphics grDevices utils      datasets  methods  
## [8] base  
##  
## other attached packages:  
## [1] pcaPP_1.9-73           checkpoint_0.4.9  
## [3] ggrepel_0.8.2          sensobol_0.3  
## [5] logitnorm_0.8.37       benchmarkme_1.0.3  
## [7] cowplot_1.0.0          scales_1.1.0  
## [9] data.table_1.12.8       Rfast_1.9.9  
## [11] RcppZiggurat_0.1.5     doParallel_1.0.15  
## [13] iterators_1.0.12        foreach_1.5.0  
## [15]forcats_0.5.0          stringr_1.4.0  
## [17] dplyr_0.8.5            purrr_0.3.4  
## [19] readr_1.3.1            tidyverse_1.3.0  
## [21] tibble_3.0.1           ggplot2_3.3.0  
## [23] tidyverse_1.3.0         RcppArmadillo_0.9.860.2.0  
## [25] Rcpp_1.0.4.6  
##  
## loaded via a namespace (and not attached):  
## [1] httr_1.4.1              jsonlite_1.6.1      modelr_0.1.6  
## [4] Rdpack_0.11-1           assertthat_0.2.1    cellranger_1.1.0  
## [7] yaml_2.2.1              remotes_2.1.1       pillar_1.4.3  
## [10] backports_1.1.6          lattice_0.20-41    glue_1.4.0  
## [13] digest_0.6.25           rvest_0.3.5        colorspace_1.4-1  
## [16] htmltools_0.4.0          Matrix_1.2-18      pkgconfig_2.0.3  
## [19] bibtex_0.4.2.2          broom_0.5.6        haven_2.2.0  
## [22] mvtnorm_1.1-0           farver_2.0.3       generics_0.0.2  
## [25] ellipsis_0.3.0          withr_2.2.0        cli_2.0.2  
## [28] magrittr_1.5             crayon_1.3.4       readxl_1.3.1  
## [31] evaluate_0.14            fs_1.4.1           fansi_0.4.1
```

```

## [34] nlme_3.1-147           xml2_1.3.1          tools_3.6.3
## [37] hms_0.5.3                gbRd_0.4-11        lifecycle_0.2.0
## [40] munsell_0.5.0             reprex_0.3.0       compiler_3.6.3
## [43] rlang_0.4.5               grid_3.6.3         rstudioapi_0.11
## [46] labeling_0.3              rmarkdown_2.1      gtable_0.3.0
## [49] codetools_0.2-16          DBI_1.1.0         curl_4.3
## [52] benchmarkmeData_1.0.3     R6_2.4.1          lubridate_1.7.8
## [55] knitr_1.28                stringi_1.4.6     vctrs_0.2.4
## [58] dbplyr_1.4.3              tidyselect_1.0.0  xfun_0.13

## Return the machine CPU
cat("Machine:    "); print(get_cpu()$model_name)

## Machine:
## [1] "Intel(R) Core(TM) i9-9900K CPU @ 3.60GHz"

## Return number of true cores
cat("Num cores:   "); print(detectCores(logical = FALSE))

## Num cores:
## [1] 8

## Return number of threads
cat("Num threads: "); print(detectCores(logical = TRUE))

## Num threads:
## [1] 16

## Return the machine RAM
cat("RAM:        "); print (get_ram()); cat("\n")

## RAM:
## 34.4 GB

```

## References

Becker, William. 2019. “Sensitivity analysis on a shoestring : screening model inputs at low sample size.”