

# Discrepancy measures for sensitivity analysis

R code

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# 1 Preliminary

```
# PRELIMINARY #####
```

```
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.margin=margin(0, 0, 0, 0),
          legend.box.margin=margin(-5,-5,-5,-5),
          legend.key = element_rect(fill = "transparent",
                                     color = NA),
          strip.background = element_rect(fill = "white"))
}
```

```
# Load the packages
```

```
sensobol::load_packages(c("sensobol", "data.table", "tidyverse", "parallel",
                           "RcppAlgos", "scales", "doParallel", "benchmarkme",
                           "cowplot", "wesanderson", "microbenchmark"))
```

```
# CHECKPOINT -----
```

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

checkpoint("2023-01-13", R.version ="4.2.1", checkpointLocation = getwd())
```

```
# C++ CODE -----
```

```
# Source cpp code -----
```

```
cpp_functions <- c("cpp_functions.cpp", "L2star_functions.cpp",
                   "L2_functions.cpp", "L2centered_functions.cpp",
                   "L2wraparound_functions.cpp", "L2modified_functions.cpp")
```

```
for(i in 1:length(cpp_functions)) {
  Rcpp::sourceCpp(cpp_functions[i])
}
```

## 2 Define functions

```
# DEFINE FUNCTIONS #####
# SAVAGE SCORES -----
savage_scores <- function(x, type) {

  if (type == "ersatz") {

    true.ranks <- rank(x)

  } else {

    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)
}

# SALTELLI ERSATZ DISCREPANCY -----
saltelli_ersatz <- function(mat) {

  N <- nrow(mat)

  s <- ceiling(sqrt(N))

  # Create the zero matrix
  mat_zeroes <- matrix(0, s, s)

  # Compute index for x_i
  m <- ceiling(mat[, 1] * s)

  # Compute index for y
  x <- mat[, 2]
  n_norm <- (x-min(x))/(max(x)-min(x)) # Scale y to 0, 1
  n <- ceiling(n_norm * s)

  # Turn y==0 to y == 1
  n <- ifelse(n == 0, 1, n)
```

```

# Merge and identify which cells are occupied by points
ind <- cbind(m, n)
mat_zeroes[ind] <- 1

# Compute discrepancy
out <- sum(mat_zeroes==1) / N

return(out)
}

# DISCREPANCY FUNCTION WRAP-UP -----
# Function to rescale -----
rescale_fun <- function(x) (x - min(x)) / (max(x) - min(x))

# Wrap up function -----
discrepancy_fun <- function (design, type) {

  X <- as.matrix(design)
  dimension <- ncol(X)
  n <- nrow(X)

  # Reescalate if needed-----
  if (min(X) < 0 || max(X) > 1) {

    X <- apply(X, 2, rescale_fun)
  }

  # Compute discrepancy
  if (type == "symmetric") {

    P <- 1 + 2 * X - 2 * X^2
    s1 <- DisS2_Rowprod(t(P), dimension)
    s2 <- DisS2_Crossprod(c(t(X)), dimension)
    R <- sqrt(((4/3)^dimension) - ((2/n) * s1) + ((2^dimension/n^2) * s2))

  } else if (type == "star") {

    dL2 <- DisL2star_Crossprod(t(X), dimension)
    R <- sqrt(3^(-dimension) + dL2)

  } else if (type == "L2") {

    P <- X * (1 - X)
    s1 <- DisL2_Rowprod(t(P), dimension)
  }
}

```

```

s2 <- DisL2_Crossprod(c(t(X)), dimension)
R <- sqrt(12^(-dimension) - (((2^(1 - dimension))/n) * s1) + ((1/n^2) * s2))

} else if (type == "centered") {

P <- 1 + 0.5 * abs(X - 0.5) - 0.5 * (abs(X - 0.5)^2)
s1 <- DisC2_Rowprod(t(P), dimension)
s2 <- DisC2_Crossprod(c(t(X)), dimension)
R <- sqrt(((13/12)^dimension) - ((2/n) * s1) + ((1/n^2) * s2))

} else if (type == "wraparound") {

s1 <- DisW2_Crossprod(t(X), dimension)
R <- sqrt(-(4/3)^dimension + (1/n^2) * s1)

} else if (type == "modified") {

P <- 3 - X^2
s1 <- DisM2_Rowprod(t(P), dimension)
s2 <- DisM2_Crossprod(c(t(X)), dimension)
R <- sqrt(((4/3)^dimension) - (((2^(1 - dimension))/n) * s1) + ((1/n^2) * s2))

} else if (type == "ersatz") {

R <- saltelli_ersatz(X)

}
return(R)
}

# Final discrepancy function -----
discrepancy <- function(mat, y, params, type) {

value <- sapply(1:ncol(mat), function(j) {
  design <- cbind(mat[, j], y)
  value <- discrepancy_fun(design = design, type = type)

})
return(value)
}

# FUNCTION TO COMPUTE JANSEN T_I -----
jansen_ti <- function(d, N, params) {

m <- matrix(d, nrow = N)
k <- length(params)

```

```

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)
value <- (1 / (2 * N) * Rfast::colsums((Y_A - Y_AB) ^ 2)) / VY

return(value)
}

```

### 3 Metafunction

```

# DISTRIBUTIONS OF THE METAFUNCTIONS #####
# Functions -----
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),
  Piecewise.large = function(x) ((-1) ^ as.integer(4 * x) *
    (0.125 - (x %% 0.25)) + 0.125),
  Piecewise.small = function(x) ((-1) ^ as.integer(32 * x) *
    (0.03125 - 2 * (x %% 0.03125)) + 0.03125) / 2,
  Oscillation = function(x) x ^ 2 - 0.2 * cos(7 * pi * x)
)

# Truncated distributions -----
truncated_normal <- function(x, min, max, mean, sd) {

  a <- pnorm(min, mean, sd)
  b <- pnorm(max, mean, sd)

  uniform <- qunif(x, a, b)
  out <- qnorm(uniform, mean, sd)

  return(out)
}

truncated_beta <- function(x, min, max, shape1, shape2) {

```

```

a <- pbeta(min, shape1, shape2)
b <- pbeta(max, shape1, shape2)

uniform <- qunif(x, a, b)
out <- qbeta(uniform, shape1, shape2)

return(out)
}

truncated_logitnorm <- function(x, min, max, mu, sigma) {

  a <- logitnorm::plogitnorm(min, mu = mu, sigma = sigma)
  b <- logitnorm::plogitnorm(max, mu = mu, sigma = sigma)

  uniform <- qunif(x, a, b)
  out <- logitnorm::qlogitnorm(uniform, mu = mu, sigma = sigma)

  return(out)
}

# Random distributions -----
sample_distributions <- list(

  "uniform" = function(x) x,
  "normal" = function(x) truncated_normal(x, 0, 1, 0.5, 0.15),
  "beta" = function(x) truncated_beta(x, 0, 1, 8, 2),
  "beta2" = function(x) truncated_beta(x, 0, 1, 2, 8),
  "beta3" = function(x) truncated_beta(x, 0, 1, 2, 0.8),
  "beta4" = function(x) truncated_beta(x, 0, 1, 0.8, 2),
  "logitnormal" = function(x) truncated_logitnorm(x, 0, 1, 0, 3.16)

)

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 #####

```

```

# Density function
sample_distributions_PDF <- list(
  "uniform" = function(x) dunif(x, 0, 1),
  "normal" = function(x) dnorm(x, 0.5, 0.15),
  "beta" = function(x) dbeta(x, 8, 2),
  "beta2" = function(x) dbeta(x, 2, 8),
  "beta3" = function(x) dbeta(x, 2, 0.8),
  "beta4" = function(x) dbeta(x, 0.8, 2),
  "logitnormal" = function(x) logitnorm::dlogitnorm(x, 0, 3.16)
)

names_ff <- names(sample_distributions)
x <- seq(0, 1, .001)
out <- matrix(rep(x, length(names_ff)), ncol = length(names_ff))

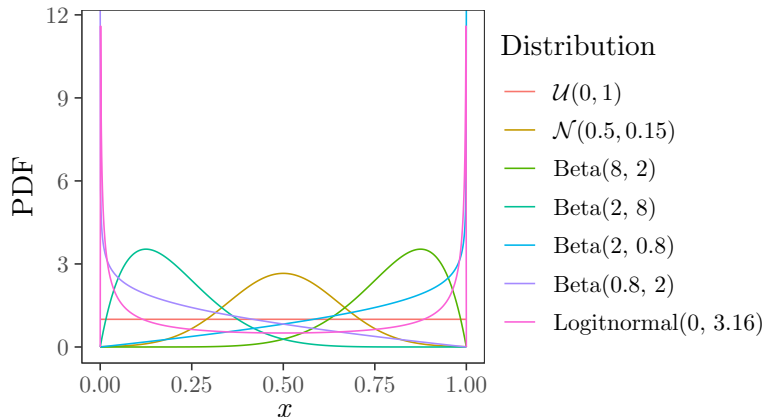
dt <- data.table(sapply(seq_along(names_ff), function(x)
  sample_distributions_PDF[[names_ff[x]]](out[, x])))

dt <- setnames(dt, paste("V", 1:length(names_ff), sep = ""), names_ff) %>%
  .[, x:= x] %>%
  melt(., measure.vars = names_ff)

plot.distributions <- ggplot(dt, aes(x = x, y = value, group = variable)) +
  geom_line(aes(color = variable)) +
  scale_color_discrete(labels = c("$\\mathcal{U}(0, 1)$",
                                    "$\\mathcal{N}(0.5, 0.15)$",
                                    "Beta(8, 2)",
                                    "Beta(2, 8)",
                                    "Beta(2, 0.8)",
                                    "Beta(0.8, 2)",
                                    "Logitnormal(0, 3.16)"),
                                    name = "Distribution") +
  labs(x = expression(italic(x)),
       y = "PDF") +
  theme_AP() +
  theme(legend.text.align = 0)

plot.distributions

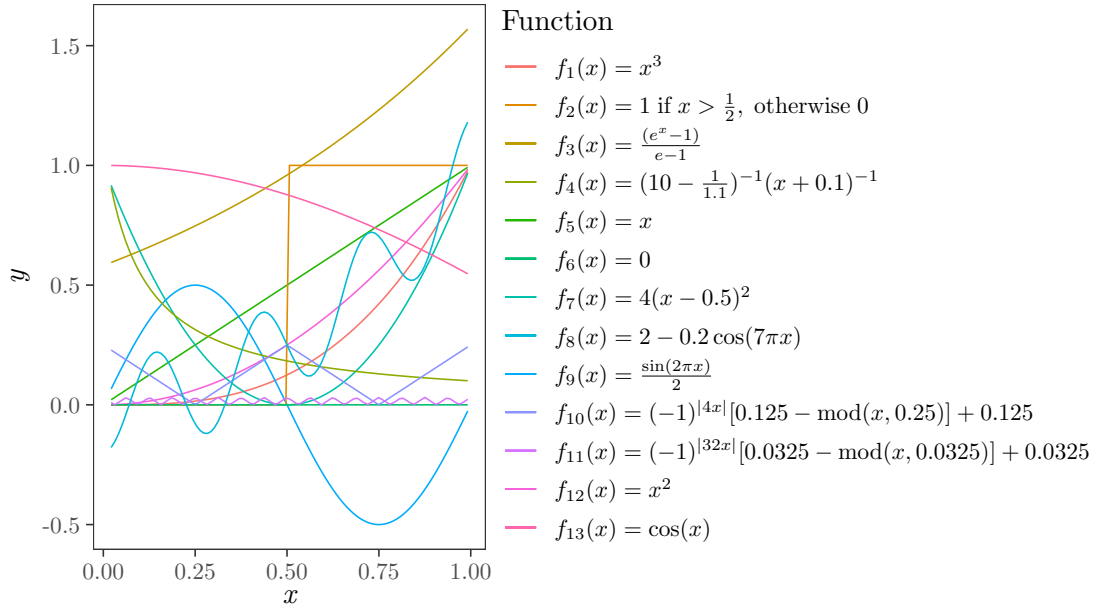
```



```
# PLOT METAFUNCTION #####
plot.metafunction <- 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]))))
  }) +
  labs(color= "Function", linetype = "Function",
       x = expression(italic(x)),
       y = expression(italic(y))) +
  scale_color_discrete(labels = c("$f_1(x) = x^3$",
                                    "$f_2(x) = 1 \\\hspace{1mm} \\mbox{if} \\\hspace{1mm} x > \\\frac{1}{2}$",
                                    "$f_3(x) = \\\frac{(e^x - 1)}{e-1}$",
                                    "$f_4(x) = (10-\\frac{1}{1.1})^{-1}(x + 0.1)^{-1}$",
                                    "$f_5(x) = x$",
                                    "$f_6(x) = 0$",
                                    "$f_7(x) = 4(x - 0.5)^2$",
                                    "$f_8(x) = 2 - 0.2 \\\cos(7 \\\pi x)$",
                                    "$f_9(x) = \\\frac{\\\sin(2 \\\pi x)}{2}$",
                                    "$f_{10}(x) = (-1)^{|4x|} [0.125- \\\mbox{mod}(x, 0.25)] + 0.125$",
                                    "$f_{11}(x) = (-1)^{|32x|} [0.0325- \\\mbox{mod}(x, 0.0325)] + 0.0325$",
                                    "$f_{12}(x) = x^2$",
                                    "$f_{13}(x) = \\\cos(x)$"))
  +
  theme_AP() +
  theme(legend.text.align = 0)

## Warning: `aes_()` was deprecated in ggplot2 3.0.0.
## i Please use tidy evaluation ideoms with `aes()``
```

```
plot.metafunction
```



## 4 Plots to show discrepancy

```
# LIST OF FUNCTIONS #####
# NEW FUNCTIONS -----
f1_fun <- function(x) 10 * x[, 1] + 0.2 * x[, 2]^3
f2_fun <- function(x) 2 * x[, 1] - x[, 2]^2
f3_fun <- function(x) x[, 1]^2 + x[, 2]^4 + x[, 1] * x[, 2] + x[, 2] * x[, 3]^4
f4_fun <- function(x) 0.2 * exp(x[, 1] - 3) + 2.2 * abs(x[, 2]) + 1.3 * x[, 2]^6 -
  2 * x[, 2]^2 - 0.5 * x[, 2]^4 - 0.5 * x[, 1]^4 + 2.5 * x[, 1]^2 + 0.7 * x[, 1]^3 +
  3 / ((8 * x[, 1] - 2)^2 + (5 * x[, 2] - 3)^2 + 1) + sin(5 * x[, 1]) * cos(3 * x[, 1]^2)
f6_fun <- function(x) 0.2 * exp(x[, 1] + 2 * x[, 2])
fun_vec <- paste("f", 1:4, "_fun", sep = "")
f_list <- list(f1_fun, f2_fun, f3_fun, f4_fun)
names(f_list) <- fun_vec
# RUN FUNCTIONS -----
output <- ind <- list()
for (i in names(f_list)) {
  if(i == "f3_fun") {
```

```

k <- 3

} else {

  k <- 2
}
params <- paste("$x_",
  1:k, "$",
  sep = "")
N <- 100
mat <- sobol_matrices(N = N, params = params, scrambling = 1)
y <- f_list[[i]](mat)
y <- rescale_fun(y)
ind[[i]] <- sobol_indices(Y = y, N = N, params = params)
output[[i]] <- plot_scatter(data = mat, N = N, Y = y, params = params) +
  labs(x = "$x$",
    y = "$y$") +
  scale_x_continuous(breaks = c("0" = 0, "0.5" = 0.5, "1" = 1)) +
  scale_y_continuous(breaks = c("0" = 0, "0.5" = 0.5, "1" = 1)) +
  theme(plot.margin = unit(c(0, 0.2, 0, 0), "cm"))
}

## Warning in randtoolbox::sobol(n = N, dim = k * n.matrices, ...): scrambling is
## currently disabled.

## Warning in randtoolbox::sobol(n = N, dim = k * n.matrices, ...): scrambling is
## currently disabled.

## Warning in randtoolbox::sobol(n = N, dim = k * n.matrices, ...): scrambling is
## currently disabled.

## Warning in randtoolbox::sobol(n = N, dim = k * n.matrices, ...): scrambling is
## currently disabled.

ind

## $f1_fun
##
## First-order estimator: saltelli | Total-order estimator: jansen
##
## Total number of model runs: 400
##
## Sum of first order indices: 1.062445
##          original sensitivity parameters
## 1: 1.062355e+00      Si      $x_1$
## 2: 8.943357e-05      Si      $x_2$ 
## 3: 1.064639e+00      Ti      $x_1$ 
## 4: 3.840524e-04      Ti      $x_2$ 

## $f2_fun
##

```

```

## First-order estimator: saltelli | Total-order estimator: jansen
##
## Total number of model runs: 400
##
## Sum of first order indices: 1.037818
##      original sensitivity parameters
## 1: 0.8303711      Si      $x_1$
## 2: 0.2074472      Si      $x_2$
## 3: 0.8311775      Ti      $x_1$
## 4: 0.2078450      Ti      $x_2$


## $f3_fun
##
## First-order estimator: saltelli | Total-order estimator: jansen
##
## Total number of model runs: 500
##
## Sum of first order indices: 0.9844601
##      original sensitivity parameters
## 1: 0.47789926      Si      $x_1$ 
## 2: 0.45561260      Si      $x_2$ 
## 3: 0.05094827      Si      $x_3$ 
## 4: 0.46752749      Ti      $x_1$ 
## 5: 0.51394544      Ti      $x_2$ 
## 6: 0.04775163      Ti      $x_3$ 

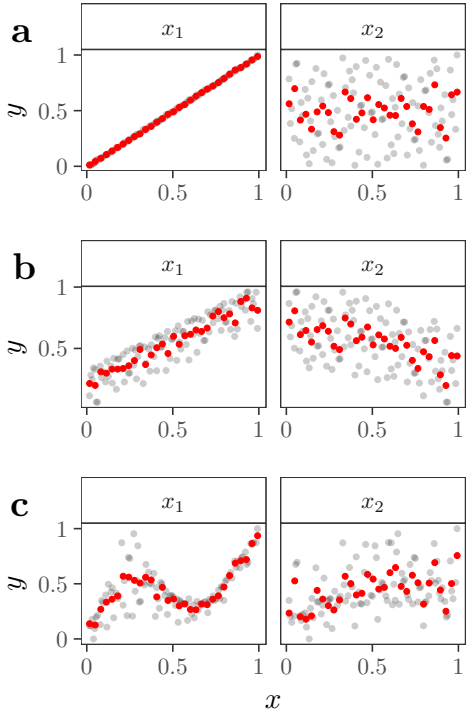
## $f4_fun
##
## First-order estimator: saltelli | Total-order estimator: jansen
##
## Total number of model runs: 400
##
## Sum of first order indices: 1.11431
##      original sensitivity parameters
## 1: 0.7858947      Si      $x_1$ 
## 2: 0.3284152      Si      $x_2$ 
## 3: 0.7622537      Ti      $x_1$ 
## 4: 0.3619071      Ti      $x_2$ 

# PLOT LIST OF FUNCTIONS #####
#####

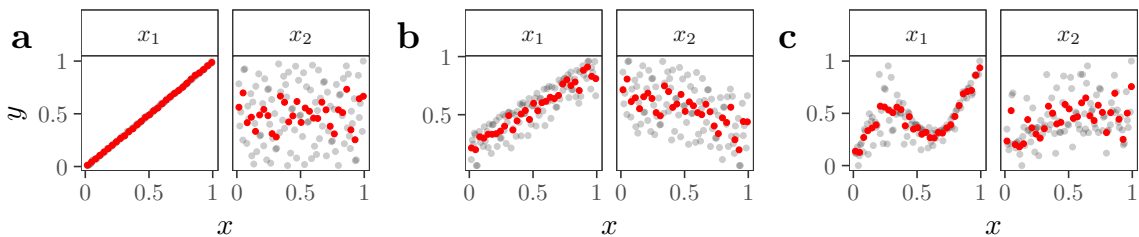
plot_list <- list(output[[1]] + labs(x = "", y = "$y$"),
                    output[[2]] + labs(x = "", y = "$y$"),
                    output[[4]])

scat_plot <- plot_grid(plotlist = plot_list, ncol = 1, labels = "auto")
scat_plot

```



```
# PLOT LIST OF FUNCTIONS #####
plot_list <- list(output[[1]] + labs(x = "$x$", y = "$y$"),
                    output[[2]] + labs(x = "$x$", y = ""),
                    output[[4]] + labs(x = "$x$", y = ""))
scat_plot <- plot_grid(plotlist = plot_list, ncol = 3, labels = "auto")
scat_plot
```



## 5 Plot sampling points in grid

```
# FUNCTION TO DRAW SAMPLING POINTS IN GRID #####
plot_grid_fun <- function(N, type, output = "plot") {
  set.seed(2)
  mat <- sobol_matrices(N = N, params = paste("$x_",$, 1:2, "$", sep = ""),
                        matrices = "A", type = type)
}
```

```

n.ersatz <- saltelli_ersatz(mat)

s <- ceiling(sqrt(N))

out <- mat %>%
  data.table() %>%
  ggplot(., aes(`$x_1`, `$x_2`)) +
  geom_point(color = "red") +
  theme_bw() +
  labs(x = "$x$", y = "y") +
  scale_x_continuous(breaks = seq(0, 1, 1/s), expand = c(0, 0), limits = c(0,1)) +
  scale_y_continuous(breaks = seq(0, 1, 1/s), expand = c(0, 0), limits = c(0,1)) +
  theme(panel.grid.minor.x = element_blank(),
        panel.grid.minor.y = element_blank(),
        axis.ticks = element_blank(),
        axis.text = element_blank(),
        plot.margin = unit(c(0.1, 0.1, 0.1, 0.1), "cm"))

if (output == "plot") {

  return(out)

} else {

  return(n.ersatz)
}

}

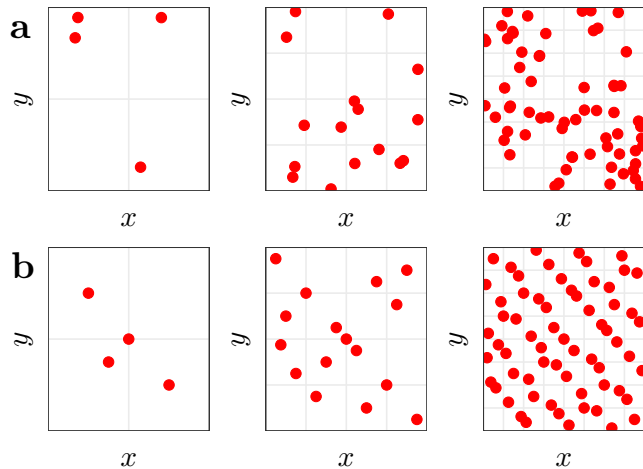
# PLOT #####
# Define loop -----
type <- c("R", "QRN")
N <- 2^seq(2, 6, 2)

# Plot -----
pt <- lapply(type, function(type) lapply(N, function(N) plot_grid_fun(N = N, type = type)))
out <- list()

for (i in 1:length(type)) {
  out[[i]] <- plot_grid(plotlist = pt[[i]], ncol = length(N), labels = ""))
}

all.grids <- plot_grid(out[[1]], out[[2]], ncol = 1, labels = "auto")
all.grids

```



```

# EXTRACT ERSATZ MEASURE #####
N <- 2^seq(2, 14, 2)

ersatz.measure <- lapply(type, function(type) lapply(N, function(N)
  plot_grid_fun(N = N, type = type, output = "ersatz")))

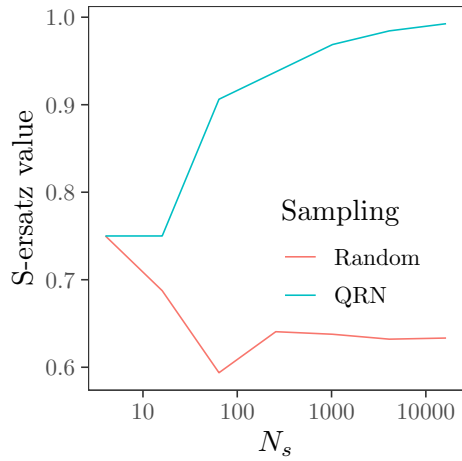
col_names <- c("Random", "QRN")
dt.ersatz <- data.table(do.call(cbind, ersatz.measure))
dt.ersatz <- setnames(dt.ersatz, paste("V", 1:2, sep = ""), col_names)
dt.ersatz <- dt.ersatz[, sample.size:= N]

dt.ersatz[, (col_names):= lapply(.SD, as.numeric), .SDcols = (col_names)]

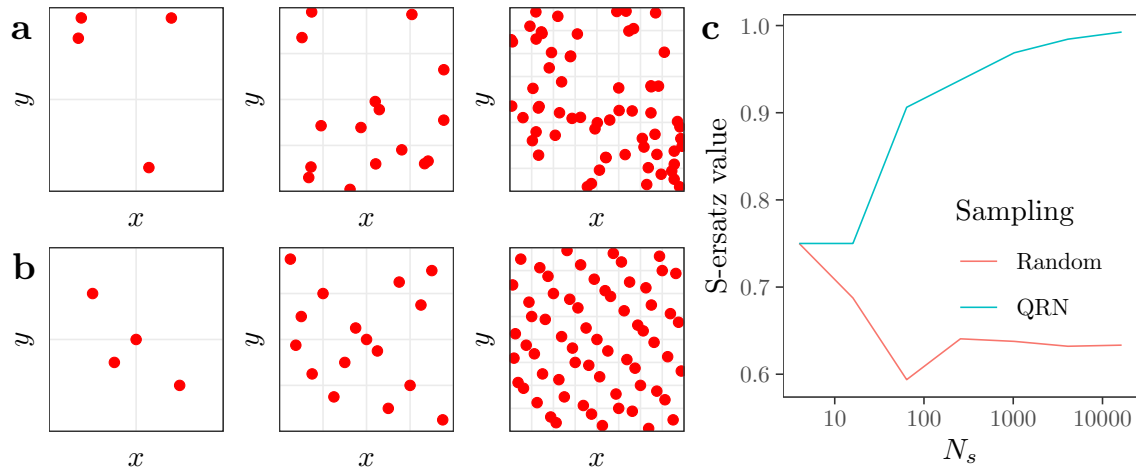
plot.ersatz <- melt(dt.ersatz, measure.vars = c("Random", "QRN")) %>%
  ggplot(., aes(sample.size, value, color = variable, group = variable)) +
  geom_line() +
  scale_x_log10() +
  labs(x = "$N_s$", y = "S-ersatz value") +
  scale_color_discrete(name = "Sampling") +
  theme_AP() +
  theme(legend.position = c(0.7, 0.35))

plot.ersatz

```



```
plot_grid(all.grids, plot.ersatz, ncol = 2, labels = c("", "c"),
          rel_widths = c(0.6, 0.4))
```



## 6 The model

```
# DEFINE MODEL #####
model_fun <- function(tau, epsilon, base.sample.size, cost.discrepancy, phi, k) {
  params <- paste("X", 1:k, sep = "")
  # Define vectors to loop through -----
  simulations <- c("discrepancy", "jansen")
  disc.type <- c("symmetric", "star", "L2", "centered", "wraparound",
                 "modified", "ersatz")
  # Select the sampling method -----
  if (tau == 1) {
```

```

type <- "R"

} else if (tau == 2) {

  type <- "QRN"
}

# If statements to select matrices and N as a function
# of the estimator used -----
#-----#
mat <- y <- ind <- disc <- output <- jansen.results <- list()

for (i in simulations) {

  if (i == "discrepancy") {

    matrices <- "A"
    N <- cost.discrepancy

  } else if (i == "jansen") {
    matrices <- c("A", "AB")
    N <- base.sample.size
  }

  # Construct the sample matrix, randomly transform it
  # according to phi and run the metafunction -----
  #-----#
}

if (i == "discrepancy") {

  set.seed(epsilon)
  mat.uniform <- sobol_matrices(matrices = matrices, N = N, params = params,
                                 type = type)

  set.seed(epsilon)
  mat[[i]] <- random_distributions(sobol_matrices(matrices = matrices,
                                                    N = N, params = params,
                                                    type = type), phi = phi)

} else if (i == "jansen") {

  set.seed(epsilon)
  mat[[i]] <- random_distributions(sobol_matrices(matrices = matrices,
                                                    N = N, params = params,
                                                    type = type), phi = phi)
}

```

```

set.seed(epsilon)
y[[i]] <- sensobol::metafunction(data = mat[[i]], epsilon = epsilon)

# Calculate first (saltelli) and total order (jansen) indices
# and discrepancy values -----
#-----#
if (i == "jansen") {

  ind[[i]] <- jansen_ti(d = y[[i]], N = base.sample.size, params = params)

}

if (i == "discrepancy") {

  discrepancy.val <- lapply(disc.type, function(x)
    discrepancy(mat = mat.uniform, y = y[[i]], params = params, type = x))
}

# Arrange output -----
names(discrepancy.val) <- disc.type
all.simulations <- c(ind, discrepancy.val)

# Savage scores -----
all.simulations.savage <- list()

for (i in names(all.simulations)) {

  all.simulations.savage[[i]] <- savage_scores(all.simulations[[i]], type = i)
}

# Correlation between indices and discrepancy measures -----
#-----#
for (i in disc.type) {

  jansen.results[[i]] <- cor(all.simulations.savage$jansen, all.simulations.savage[[i]])

}

# Arrange output -----
output <- unlist(jansen.results)

return(output)

```

```
}
```

## 7 Sample matrix

```
# CREATE SAMPLE MATRIX #####
# DEFINE SETTINGS -----
N <- 2^9
params <- c("epsilon", "phi", "k", "tau", "base.sample.size")
mat <- sobol_matrices(matrices = "A", N = N, params = params)

# DEFINE DISTRIBUTIONS -----
mat[, "epsilon"] <- floor(qunif(mat[, "epsilon"], 1, 200))
mat[, "phi"] <- floor(mat[, "phi"] * 8) + 1
mat[, "k"] <- floor(qunif(mat[, "k"], 3, 50))
mat[, "tau"] <- floor(mat[, "tau"] * 2) + 1
mat[, "base.sample.size"] <- floor(qunif(mat[, "base.sample.size"], 10, 100))

# RE-ARRANGE COST OF ANALYSIS -----
cost.jansen <- mat[, "base.sample.size"] * (mat[, "k"] + 1)
cost.saltelli <- mat[, "base.sample.size"] * (mat[, "k"] + 2)
cost.discrepancy <- cost.jansen

final.mat <- cbind(mat, cost.jansen, cost.saltelli, cost.discrepancy)
```

## 8 Run simulations

```
# RUN SIMULATIONS #####
y <- mclapply(1:nrow(final.mat), function(i) {
  model_fun(tau = final.mat[i, "tau"],
             epsilon = final.mat[i, "epsilon"],
             base.sample.size = final.mat[i, "base.sample.size"],
             cost.discrepancy = final.mat[i, "cost.discrepancy"],
             phi = final.mat[i, "phi"],
             k = final.mat[i, "k"]),
  mc.cores = floor(detectCores() * 0.75))
```

## 9 Arrange output

```
# ARRANGE OUTPUT #####
final.dt <- do.call(rbind, y) %>%
  cbind(final.mat, .) %>%
```

```

data.table() %>%
  .[, id:= .I]

# EXPORT RESULTS -----
fwrite(final.dt, "final.dt.csv")

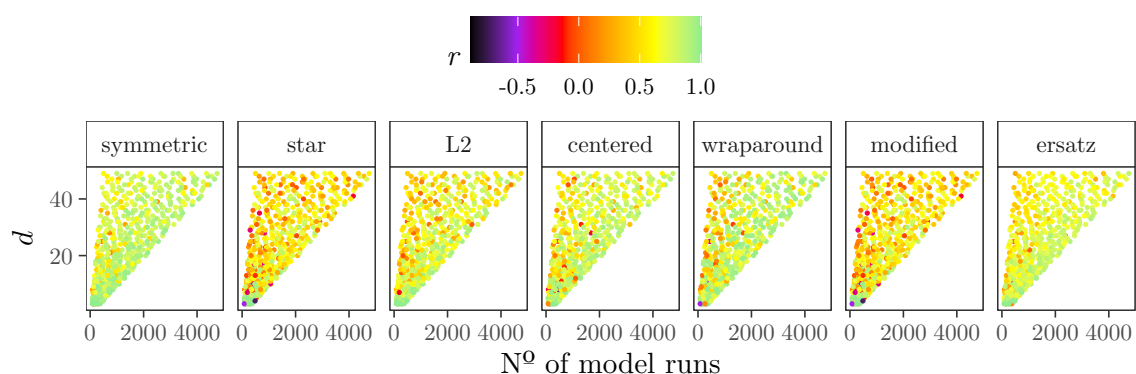
# PLOT #####
disc.type <- c("symmetric", "star", "L2", "centered",
              "wraparound", "modified", "ersatz")

# SCATTERPLOT -----
scatter <- melt(final.dt, measure.vars = disc.type) %>%
  ggplot(., aes(cost.discrepancy, k, color = value)) +
  geom_point(size = 0.4) +
  scale_colour_gradientn(colours = c("black", "purple", "red", "orange",
                                      "yellow", "lightgreen"),
                         name = expression(italic(r)),
                         breaks = pretty_breaks(n = 3)) +
  facet_grid(~variable) +
  labs(x = "Nº of model runs", y = "$d$") +
  scale_x_continuous(breaks = pretty_breaks(n = 3)) +
  scale_y_continuous(breaks = pretty_breaks(n = 3)) +
  theme_AP() +
  theme(legend.position = "none")

legend <- get_legend(scatter + theme(legend.position = "top"))

scatter.plot <- plot_grid(legend, scatter, ncol = 1, rel_heights = c(0.25, 0.75))
scatter.plot

```



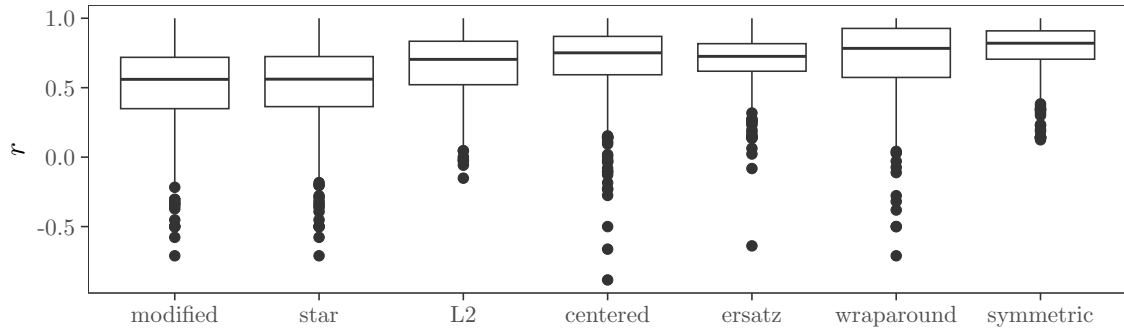
```

# BOXPLOT -----
boxplots <- melt(final.dt, measure.vars = disc.type) %>%
  ggplot(., aes(reorder(variable, value), value)) +
  geom_boxplot() +
  labs(y = "$r$", x = "") +
  scale_y_continuous(breaks = pretty_breaks(n = 3)) +

```

```
theme_AP()
```

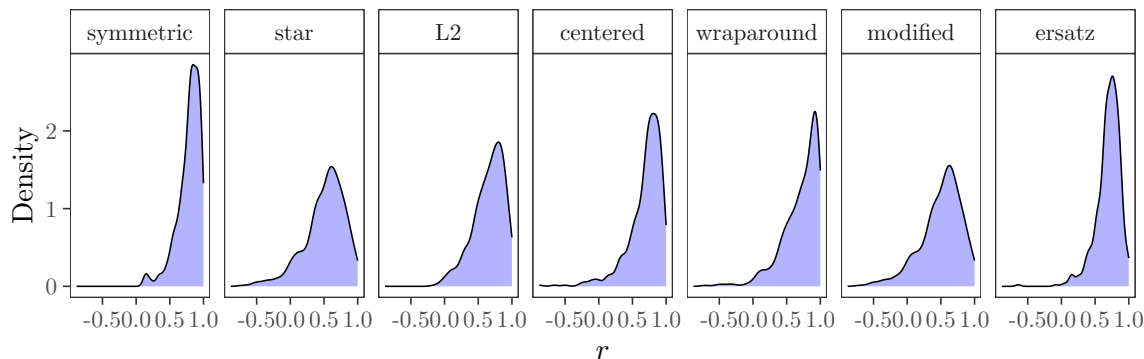
```
boxplots
```



```
# DENSITY PLOT #####
```

```
density.plot <- melt(final.dt, measure.vars = disc.type) %>%
  ggplot(., aes(value)) +
  geom_density(alpha = 0.3, color = "black", fill = "blue") +
  facet_wrap(~variable, ncol = 7) +
  scale_x_continuous(breaks = pretty_breaks(n = 3)) +
  labs(x = "$r$", y = "Density") +
  theme_AP()
```

```
density.plot
```



```
# REFINE BOXPLOTS #####
```

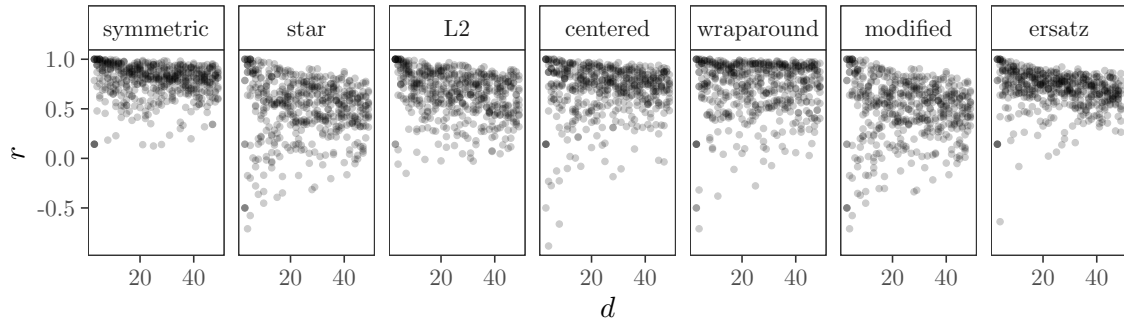
```
scatter_and_boxplots <- plot_grid(scatter.plot, density.plot, ncol = 1,
                                    labels = "auto", rel_heights = c(0.56, 0.44))
```

```
# PLOT SCATTERPLOTS OF Y AGAINST X_i #####
```

```
scatter.d <- melt(final.dt, measure.vars = disc.type) %>%
  ggplot(., aes(k, value)) +
  geom_point(alpha = 0.2, size = 0.8) +
  labs(x = "$d$ ", y = "$r$") +
  scale_x_continuous(breaks = pretty_breaks(n = 3)) +
```

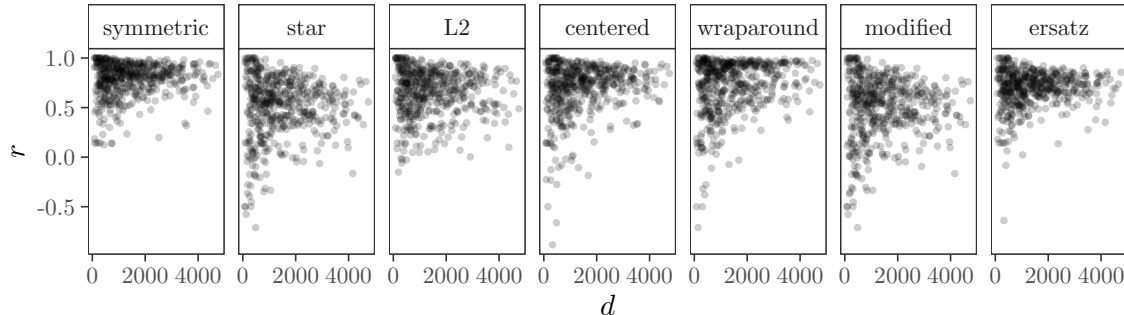
```
facet_wrap(~variable, ncol = 7) +
theme_AP()
```

scatter.d



```
scatter.n <- melt(final.dt, measure.vars = disc.type) %>%
ggplot(., aes(cost.discrepancy, value)) +
geom_point(alpha = 0.2, size = 0.8) +
labs(x = "$d$", y = "$r$") +
scale_x_continuous(breaks = pretty_breaks(n = 3)) +
facet_wrap(~variable, ncol = 7) +
theme_AP()
```

scatter.n



```
plot_grid(scatter.plot, scatter.d, ncol = 1,
          labels = "auto", rel_heights = c(0.55, 0.45))
```

```
## Warning in (function (texString, cex = 1, face = 1, engine =
## getOption("tikzDefaultEngine")), : Attempting to calculate the width of a
## Unicode string using the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.
```

```
## Warning in (function (texString, cex = 1, face = 1, engine =
## getOption("tikzDefaultEngine")), : Attempting to calculate the width of a
## Unicode string using the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.
```

```
## Warning in (function (texString, cex = 1, face = 1, engine =
## getOption("tikzDefaultEngine")), : Attempting to calculate the width of a
```

```

## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

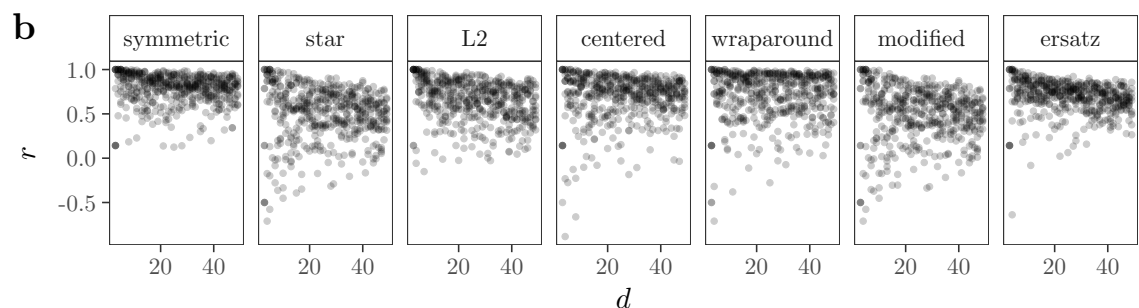
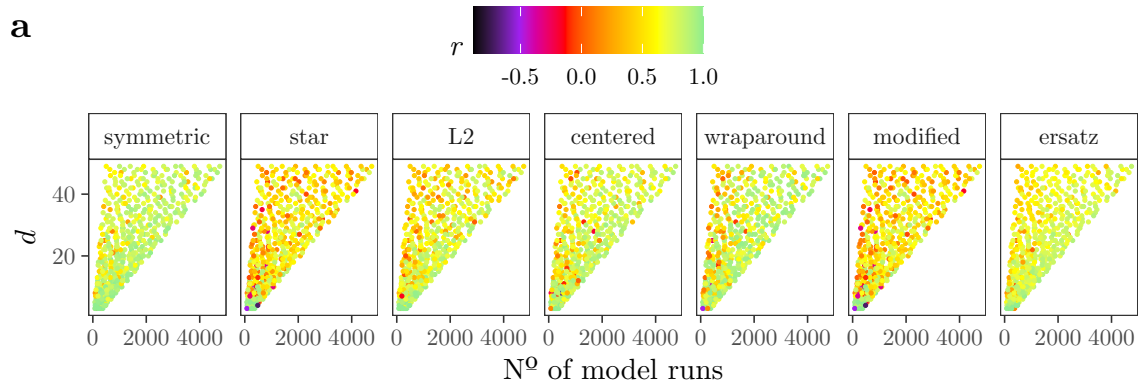
## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicode section
## of ?tikzDevice for more information.

```



```

# PAIRWISE CORRECTED MOOD TEST #####
mood.test.dt <- melt(final.dt, measure.vars = disc.type)

moot.output <- RVAideMemoire::pairwise.mood.medtest(resp = mood.test.dt$value,
                                                    fact = mood.test.dt$variable)$p.value

pvalue.plot <- ggcorrplot::ggcorrplot(moot.output, type = "lower") +
  scale_fill_gradient(low = "white", high = "red",
                      name = "$p$-value") +
  theme_AP() +
  labs(x = "", y = "") +
  scale_x_discrete(guide = guide_axis(n.dodge = 2))

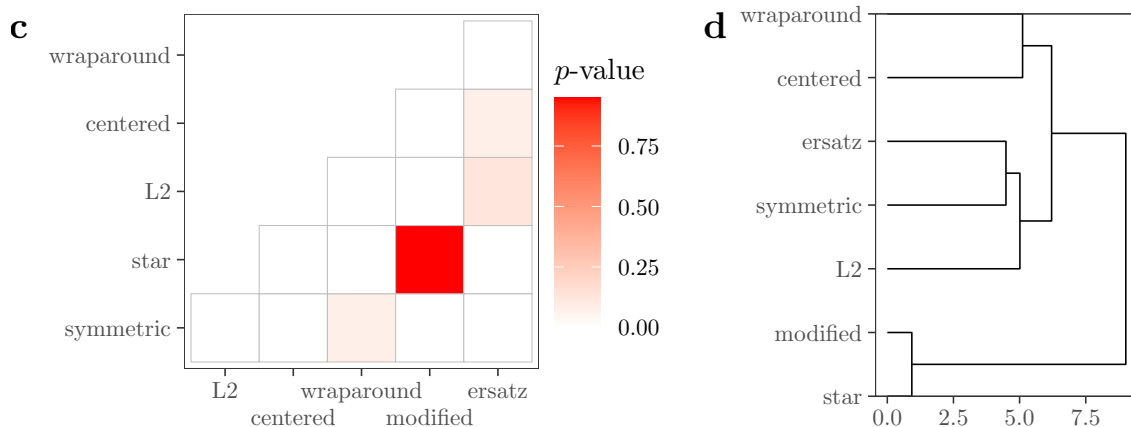
## Scale for fill is already present.
## Adding another scale for fill, which will replace the existing scale.
# HIERARCHICAL CLUSTER #####
df <- final.dt[, ..disc.type] %>%
  data.frame()

colclus <- hclust(dist(t(df))) #cluster the columns

tree.plot <- ggplot() + ggdendroplot::geom_dendro(colclus) +
  theme_AP() +
  labs(x = "", y = "") +
  coord_flip()

mood.tree.plot <- plot_grid(pvalue.plot, tree.plot, ncol = 2, labels = c("c", "d"),
                             rel_widths = c(0.6, 0.4))
mood.tree.plot

```



```

plot_grid(scatter_and_boxplots, mood.tree.plot, ncol = 1,
          rel_heights = c(0.6, 0.4))

```

```

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

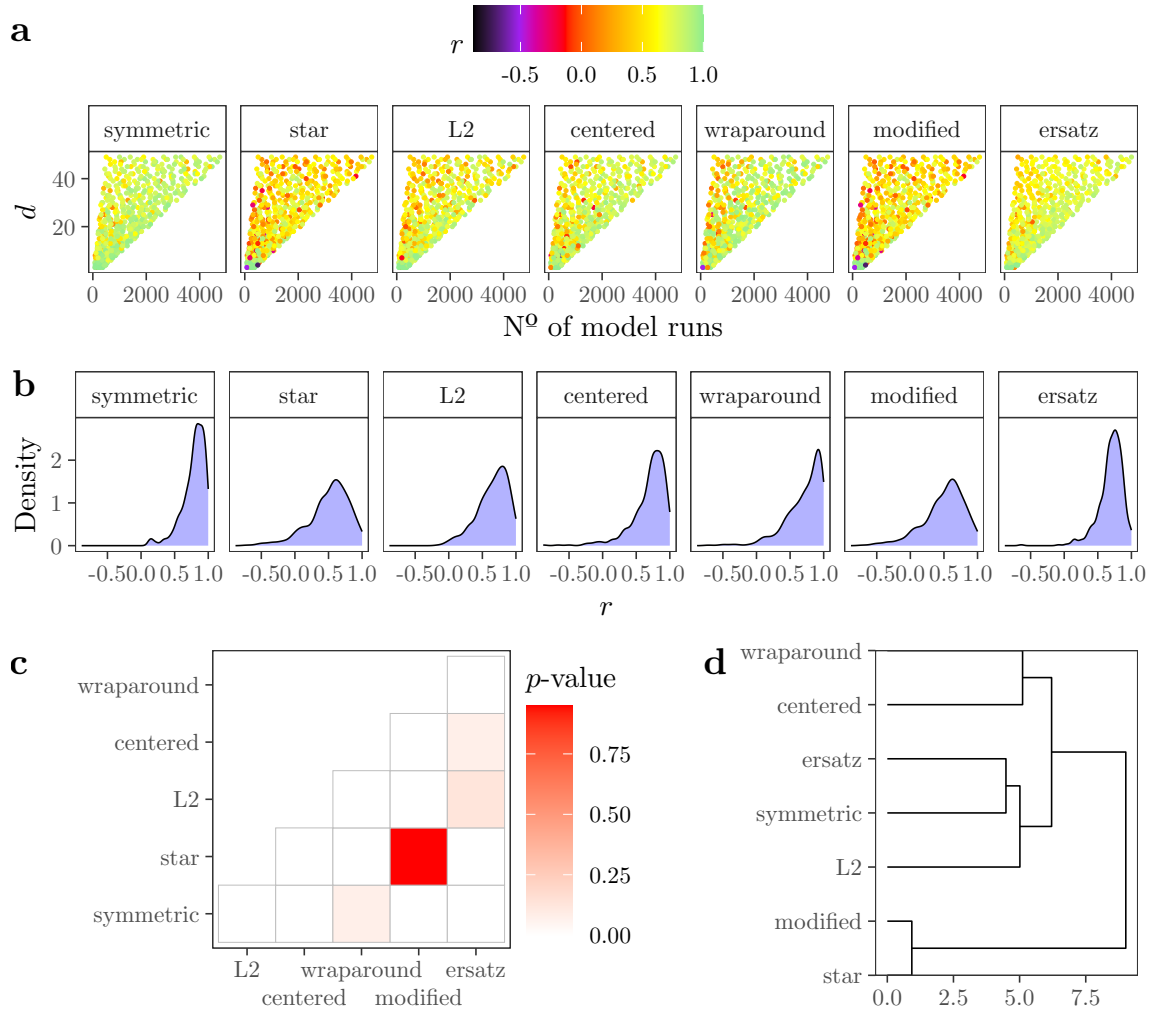
## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

## Warning in (function (texString, cex = 1, face = 1, engine =
##getOption("tikzDefaultEngine"), : Attempting to calculate the width of a
## Unicode stringusing the pdftex engine. This may fail! See the Unicodesection
## of ?tikzDevice for more information.

```



## 10 Check negative values

```
# NEGATIVE VALUES -----
negative.dt <- melt(final.dt, measure.vars = disc.type) %>%
  .[value < 0]

# DISCREPANCY -----
negative_fun <- function(cost.discrepancy, base.sample.size,
                         tau, k, epsilon, phi, disc.measure) {

  params <- paste("X", 1:k, sep = "")

  # If for the sampling method -----
  if (tau == 1) {
```

```

type <- "R"

} else if (tau == 2) {

  type <- "QRN"
}

# For discrepancy ----

set.seed(epsilon)
mat.disc <- sobol_matrices(N = cost.discrepancy, params = params, matrices = "A",
                           type = type)
set.seed(epsilon)
mat.disc2 <- random_distributions(mat.disc, phi = phi)
Y <- metafunction(mat.disc2, epsilon = epsilon)
discrepancy <- discrepancy(mat = mat.disc, y = Y, params = params, type = disc.measure)

# For Jansen ----

set.seed(epsilon)
mat.jansen <- sobol_matrices(matrices = c("A", "AB"), N = base.sample.size,
                             params = params, type = type)

set.seed(epsilon)
mat.jansen <- random_distributions(mat.jansen, phi = phi)
y <- metafunction(mat.jansen, epsilon = epsilon)
jansen <- jansen_ti(d = y, N = base.sample.size, params = params)

# Merge and create dataset ----

output <- data.table(cbind(discrepancy, jansen)) %>%
  .[, `:=` (params = factor(params, levels = params),
            ranks.jansen = rank(jansen),
            ranks.discrepancy = rank(discrepancy),
            savage.jansen = savage_scores(jansen, type = type),
            savage.discrepancy = savage_scores(discrepancy, type))]

# Plots

scatter.plot <- data.table(cbind(mat.disc, Y)) %>%
  melt(., measure.vars = params) %>%
  ggplot(., aes(value, Y)) +
  geom_hex() +
  stat_summary_bin(fun = "mean", geom = "point", colour = "red", size = 0.7) +
  facet_wrap(~variable, ncol = 4) +
  theme_AP()

```

```

bar.plot <- ggplot(output, aes(params, jansen)) +
  geom_bar(stat = "identity") +
  labs(x = "", y = "Sobol' indices") +
  theme_AP()

full.output <- list(output, scatter.plot, bar.plot)
names(full.output) <- c("output", "scatter", "bar")

return(full.output)
}

# RUN FUNCTION #####
#####

out.dt <- mclapply(1:nrow(negative.dt), function(i) {

  out <- negative_fun(cost.discrepancy = negative.dt[i]$cost.discrepancy,
                        base.sample.size = negative.dt[i]$base.sample.size,
                        tau = negative.dt[i]$tau,
                        k = negative.dt[i]$k,
                        epsilon = negative.dt[i]$epsilon,
                        phi = negative.dt[i]$phi,
                        disc.measure = negative.dt[i]$variable)

}, mc.cores = detectCores())

# ARRANGE AND PLOT NEGATIVE VALUES #####
#####

da <- final.dt[negative.dt[, id]] %>%
  .[, row:= .I]

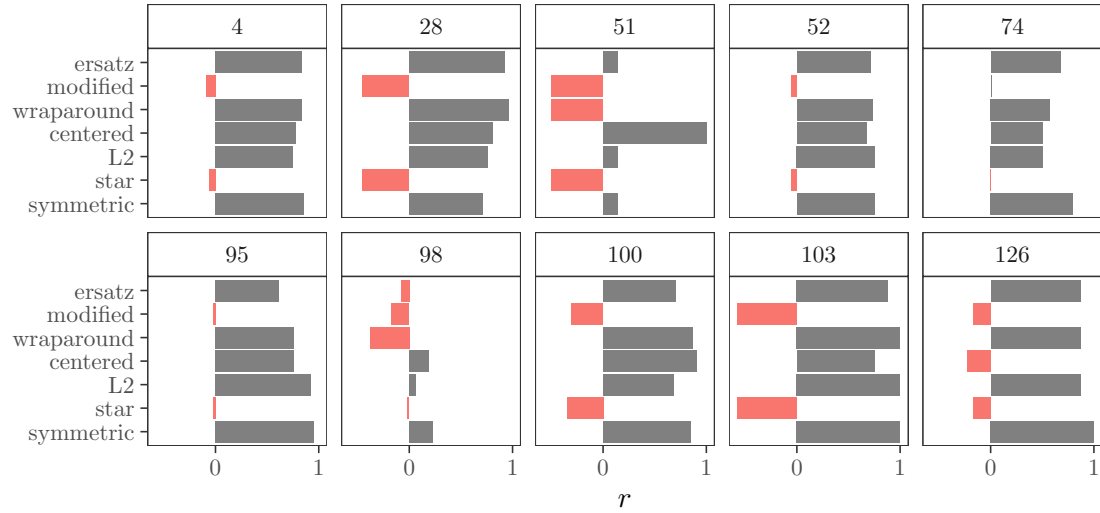
vec.split <- split(da$row, ceiling(seq_along(da$row)/10))

lapply(vec.split, function(id) {

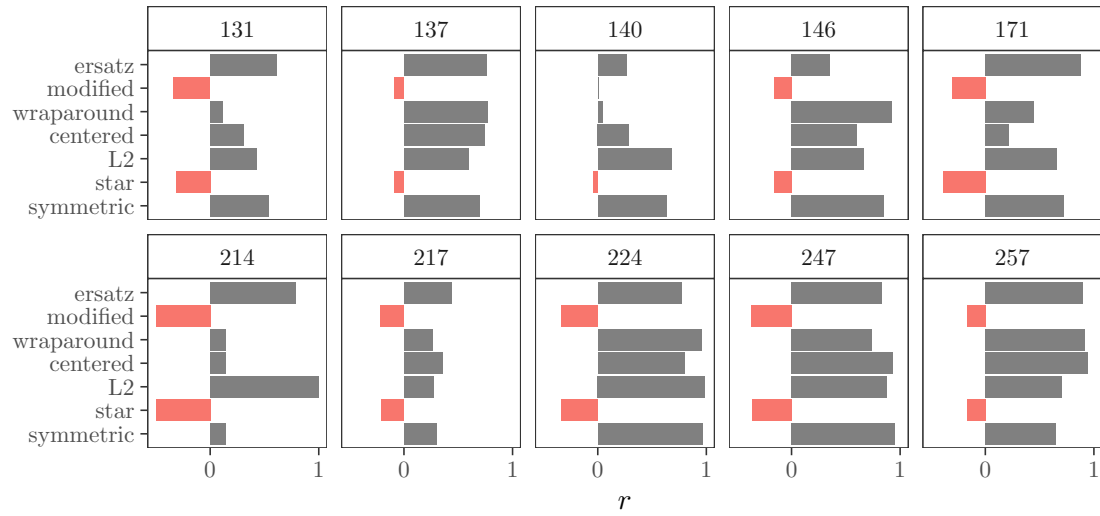
  da[id] %>%
    melt(. , measure.vars = disc.type) %>%
    ggplot(. , aes(variable, value, fill = ifelse(value < 0, "red", NA))) +
    geom_bar(stat = "identity") +
    facet_wrap(~id, ncol = 5) +
    scale_y_continuous(breaks = pretty_breaks(n = 2)) +
    coord_flip() +
    theme_AP() +
    labs(x = "", y = "$r$") +
    theme(legend.position = "none")

})

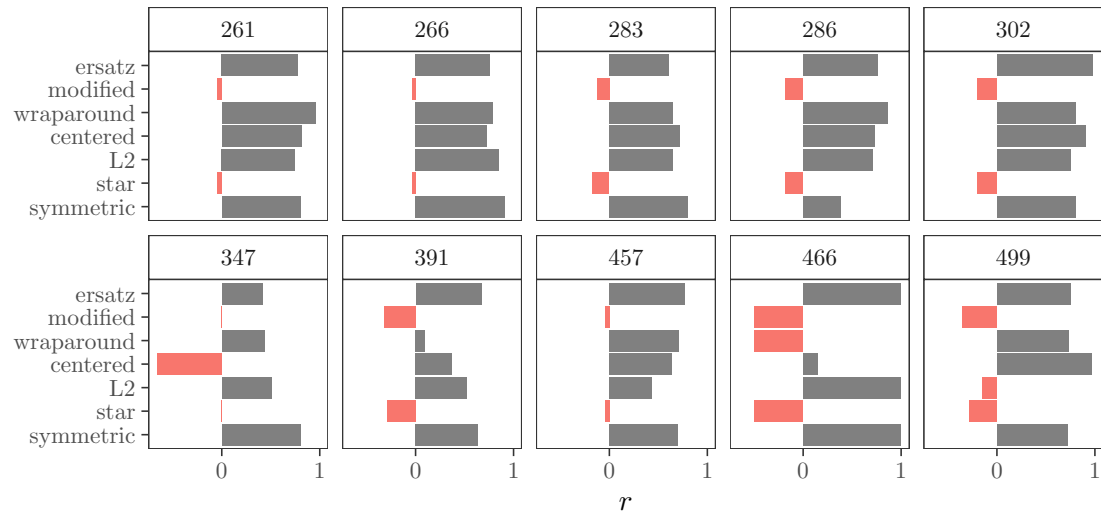
## $`1`
```



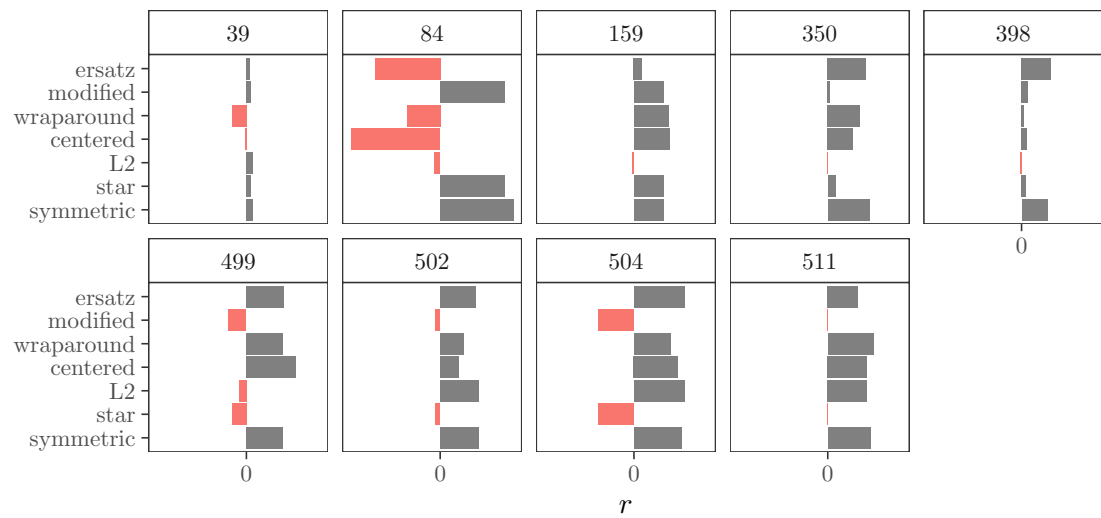
```
##  
## $`2`
```



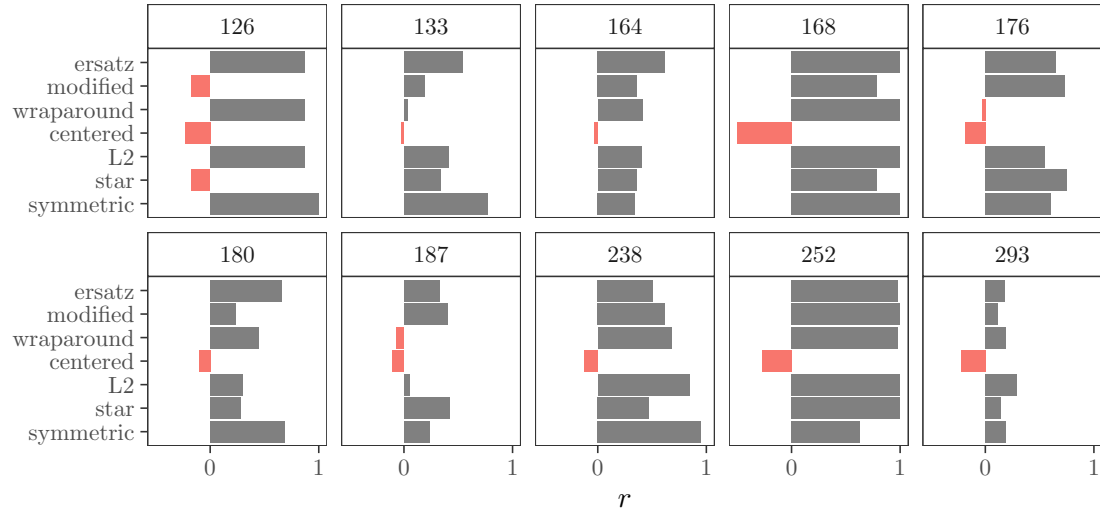
```
##  
## $`3`
```



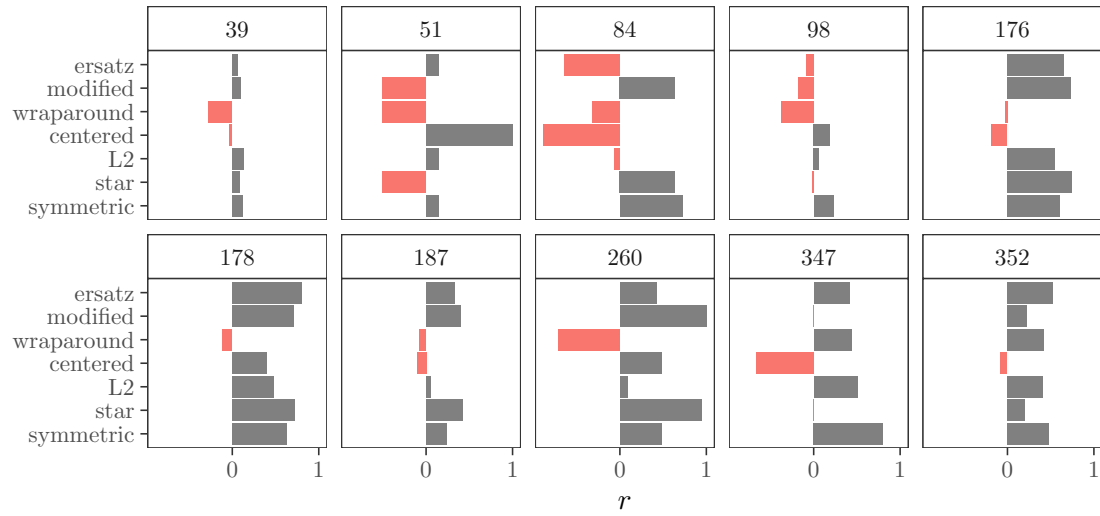
```
## $`4`
```



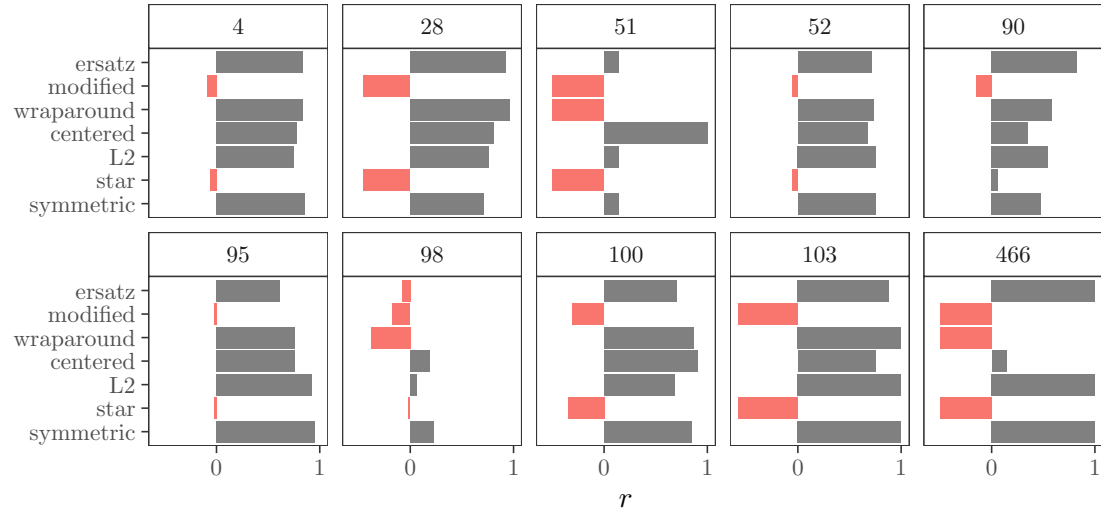
```
## $`5`
```



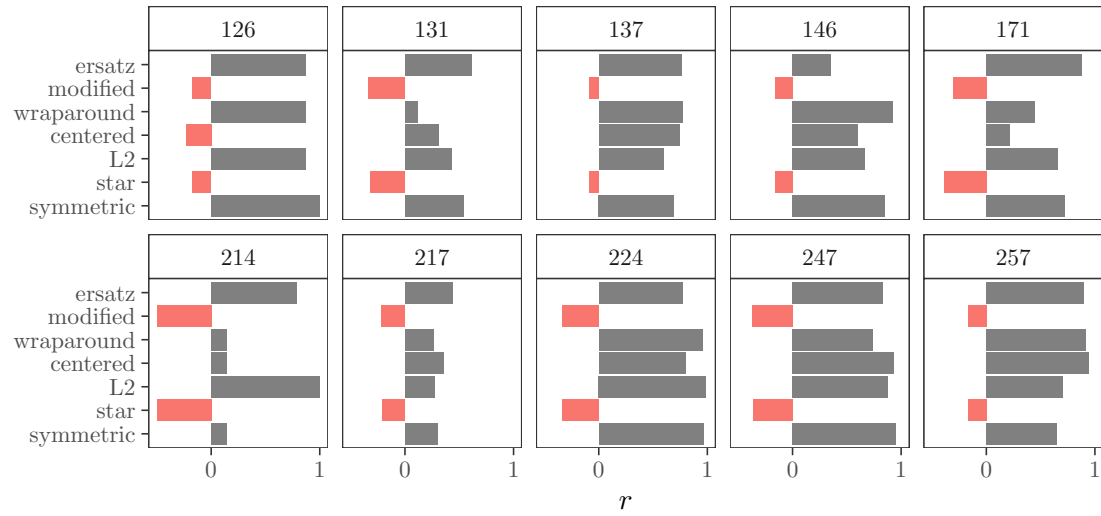
```
##  
## $`6`
```



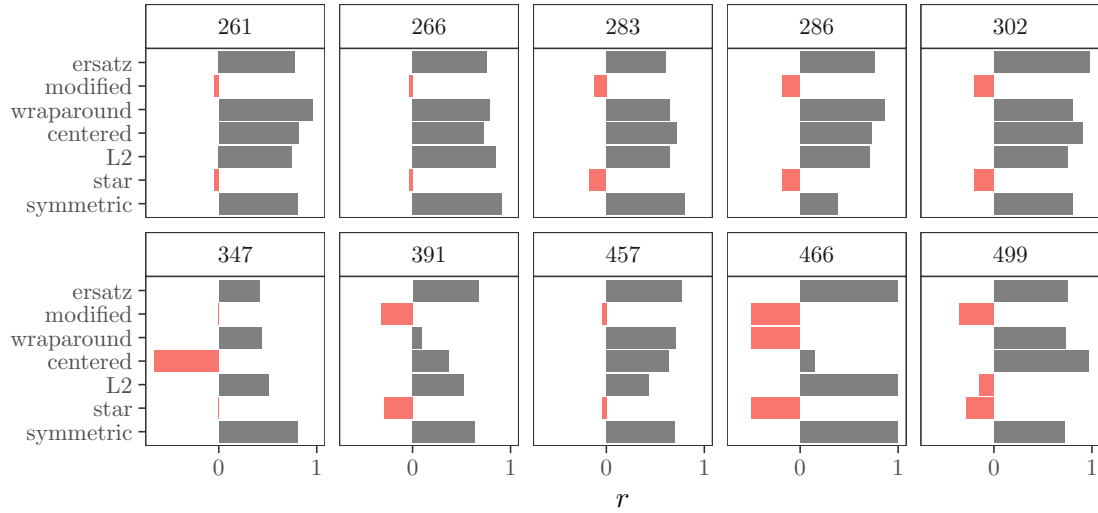
```
##  
## $`7`
```



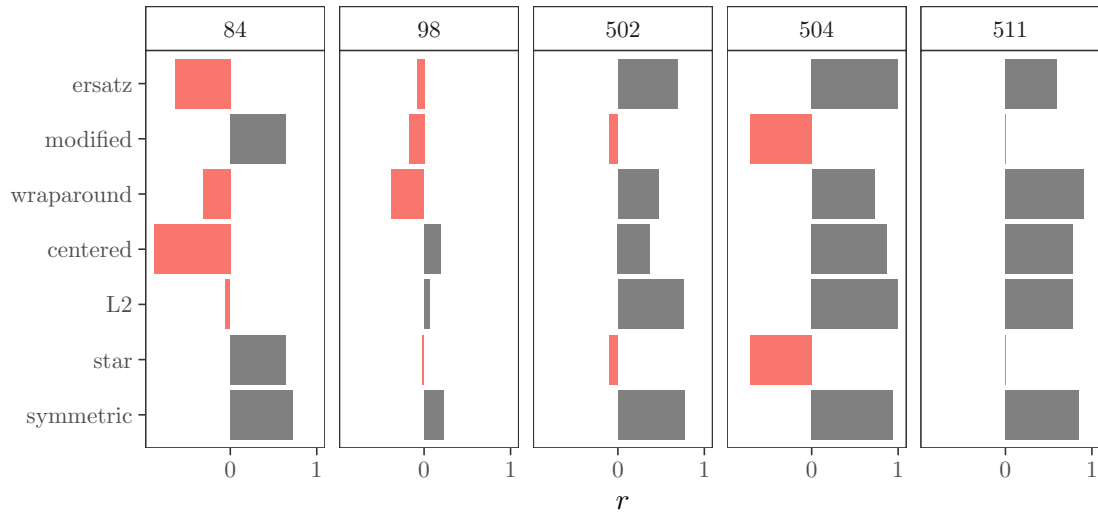
```
##  
## $`8`
```



```
##  
## $`9`
```



```
##  
## $`10`
```



## 11 Computing time

```
##### COMPUTING TIME #####
# SAMPLE SIZE-----
N <- 2^9
params <- c("N", "k")

# SAMPLE MATRIX -----
dt <- sobol_matrices(matrices = "A", N = N, params = params)
dt[, 1] <- floor(qunif(dt[, 1], 10, 10^3 + 1))
dt[, 2] <- floor(qunif(dt[, 2], 3, 50))
```

```

# RUN SIMULATIONS -----
y <- mclapply(1:nrow(dt), function(i) {

  params <- paste("x", 1:dt[i, "k"], sep = "")
  N <- dt[i, "N"]

  output[[i]] <- lapply(disc.type, function(x) {

    mat <- sobol_matrices(matrices = "A", N = N, params = params)
    y <- metafunction(data = mat)
    da <- microbenchmark(discrepancy(mat = mat, y = y, params = params, type = x),
                          times = 1)$time

  })
}

}, mc.cores = detectCores())

# ARRANGE DATA #####
da <- list()

for (i in 1:length(y)) {

  names(y[[i]]) <- disc.type
  da[[i]] <- data.frame(do.call(rbind, y[[i]])) %>%
    rownames_to_column(., var = "discrepancy") %>%
    data.table() %>%
    setnames(., colnames(.), c("discrepancy", "time")) %>%
    .[, time:= time / 10^6] # From nano to milliseconds

}

timing.dt <- rbindlist(da, idcol = "row")
fwrite(timing.dt, "timing.dt.csv")
fwrite(dt, "dt.csv")

## x being coerced from class: matrix to data.table
# PLOT RESULTS #####
# SCATTERPLOT -----
scatter.timing <- dcast(timing.dt, row ~ discrepancy, value.var = "time") %>%
  cbind(dt, .) %>%
  melt(., measure.vars = disc.type) %>%
  ggplot(., aes(N, k, color = value)) +
  geom_point(size = 0.8) +
  scico::scale_colour_scico(palette = "lajolla",

```

```

            breaks = c(1, 10^2, 10^4),
            trans = "log",
            name = "Time \n (Milliseconds)") +
facet_grid(~variable) +
scale_x_continuous(breaks = pretty_breaks(n = 3)) +
labs(x = "Nº of model runs", y = "$d$") +
theme_AP() +
theme(legend.position = "none")

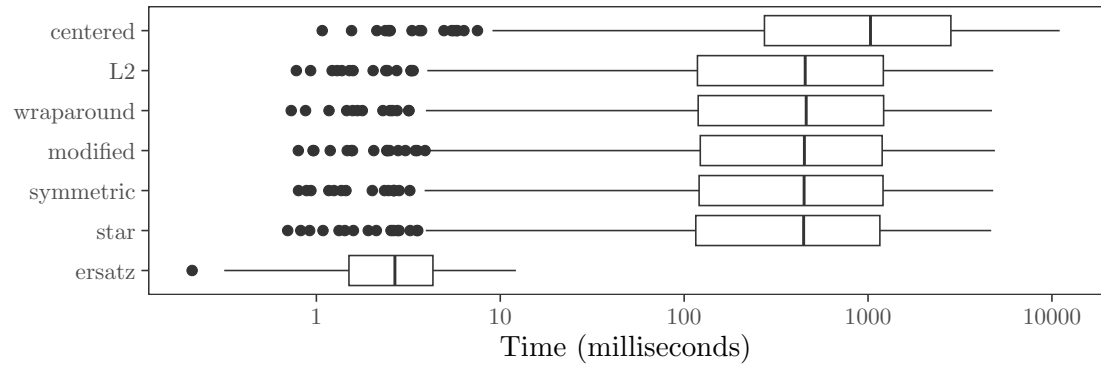
legend <- get_legend(scatter.timing + theme(legend.position = "top"))

time.plot <- plot_grid(legend, scatter.timing, ncol = 1, rel_heights = c(0.19, 0.81))

# BOXPLOT -----
boxplot.timing <- ggplot(timing.dt, aes(reorder(discrepancy, time), time)) +
  geom_boxplot() +
  coord_flip() +
  labs(x = "", y = "Time (milliseconds)") +
  scale_y_log10() +
  theme_AP()

boxplot.timing

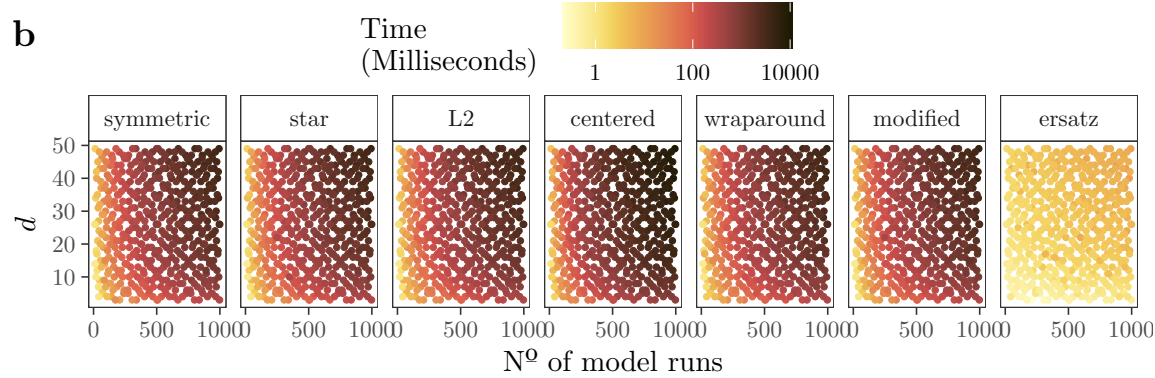
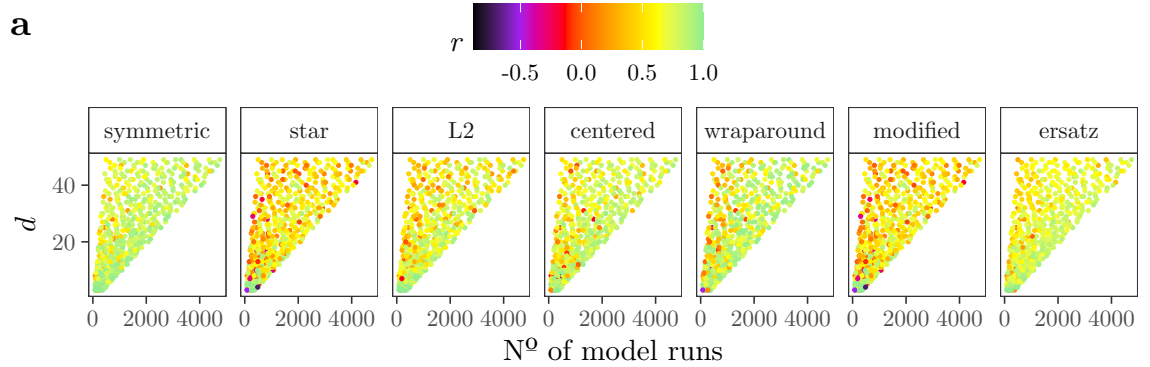
```



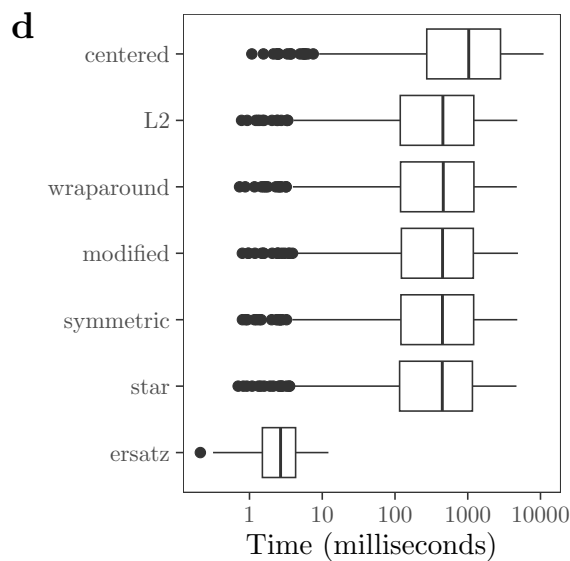
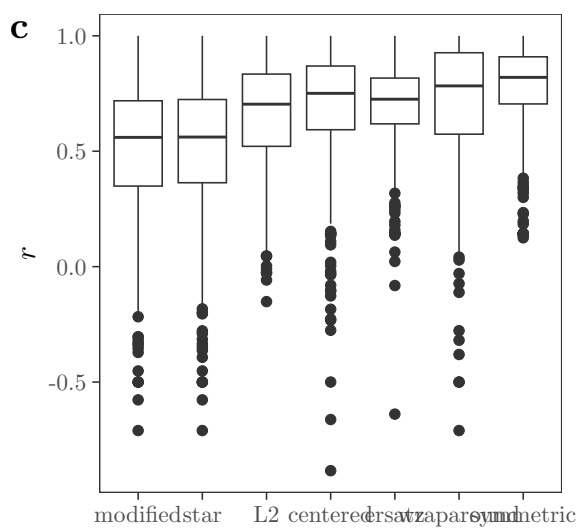
```

# MERGE PLOTS #####
all.scatter <- plot_grid(scatter.plot, time.plot, ncol = 1, labels = "auto")
all.scatter

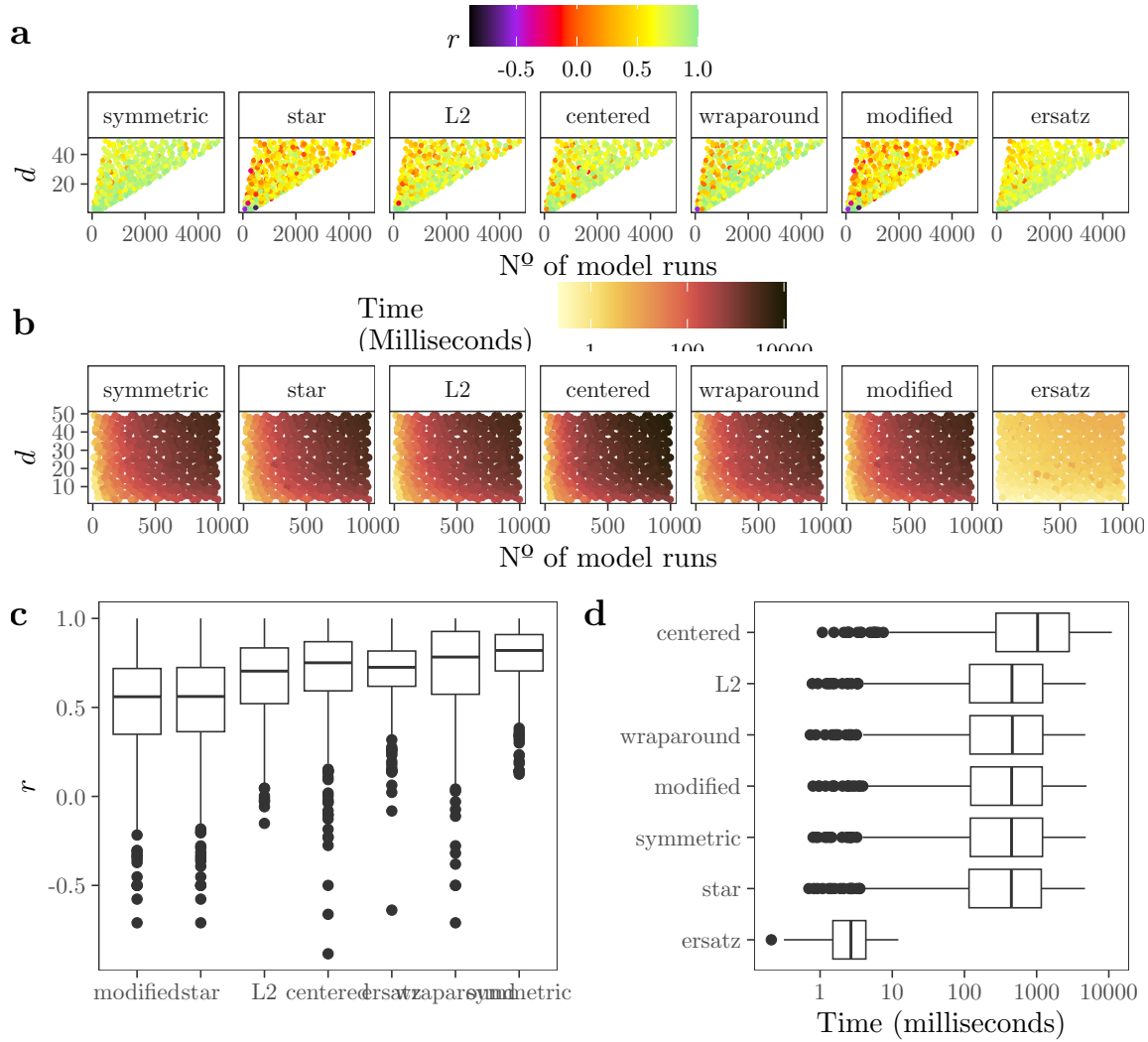
```



```
# PLOT BOXPLOTS #####
all.boxplots <- plot_grid(boxplots, boxplot.timing, ncol = 2, labels = c("c", "d"))
all.boxplots
```



```
plot_grid(all.scatter, all.boxplots, ncol = 1, rel_heights = c(0.5, 0.4))
```



```

# CALCULATE TIME COMPLEXITY #####
#####

# Setting -----
sample.size <- seq(100, 5000, 10)
dimensions <- seq(3, 200, 1)

# Function -----
timing_fun <- function(N, k) {

  params <- paste("X", 1:k, sep = "")
  mat <- sobol_matrices(matrices = "A", N = N, params = params)
  y <- metafunction(mat)
  out <- lapply(disc.type, function(x)
    microbenchmark(discrepancy(mat = mat, y = y, params = params, type = x),
                   times = 1)$time)
  names(out) <- disc.type

  return(out)
}

```

```

}

# Run model -----
timing.N <- mclapply(sample.size, function(N)
  timing_fun(N = N, k = 5), mc.cores = detectCores())

timing.dimensions <- mclapply(dimensions, function(k)
  timing_fun(N = 500, k = k), mc.cores = detectCores())

# ARRANGE RESULTS #####
names(timing.N) <- sample.size
names(timing.dimensions) <- dimensions

timing.N.dt <- lapply(timing.N, function(x) lapply(x, function(y) data.table(y))) %>%
  lapply(., function(x) rbindlist(x, idcol = "Discrepancy")) %>%
  rbindlist(., idcol = "approach") %>%
  .[, approach:= as.numeric(approach)] %>%
  .[, method:= "$N_s$"] %>%
  .[, y:= y / 10^6]

timing.dimensions.dt <- lapply(timing.dimensions, function(x) lapply(x, function(y) data.table(
  lapply(., function(x) rbindlist(x, idcol = "Discrepancy")) %>%
  rbindlist(., idcol = "approach") %>%
  .[, approach:= as.numeric(approach)] %>%
  .[, method:= "$d$"] %>%
  .[, y:= y / 10^6]

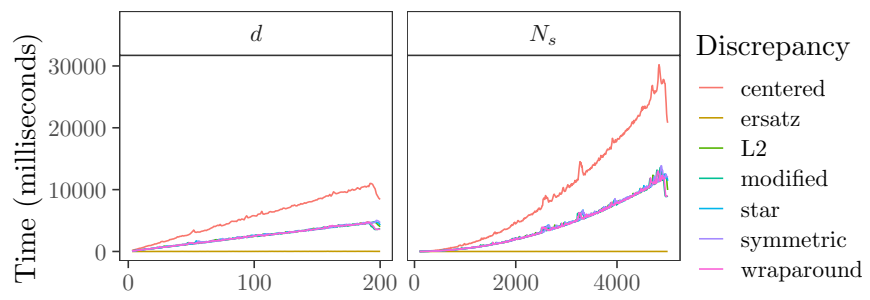
full.timing.dt <- rbind(timing.N.dt, timing.dimensions.dt)

fwrite(full.timing.dt, "full.timing.dt.csv")

# PLOT RESULTS #####
time.complexity <- ggplot(full.timing.dt, aes(approach, y,
                                                 group = Discrepancy,
                                                 color = Discrepancy)) +
  geom_line() +
  theme_AP() +
  facet_wrap(~method, scales = "free_x") +
  labs(x = "", y = "Time (milliseconds)") +
  scale_x_continuous(breaks = pretty_breaks(n = 3)) +
  theme(legend.key.width = unit(0.4, "cm"),
        legend.key.height = unit(0.4, "cm"))

time.complexity

```



## 12 Session information

```
# SESSION INFORMATION #####
sessionInfo()

## R version 4.2.0 (2022-04-22)
## Platform: aarch64-apple-darwin20 (64-bit)
## Running under: macOS Monterey 12.4
##
## Matrix products: default
## BLAS:    /Library/Frameworks/R.framework/Versions/4.2-arm64/Resources/lib/libRblas.0.dylib
## LAPACK:  /Library/Frameworks/R.framework/Versions/4.2-arm64/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] wesanderson_0.3.6 cowplot_1.1.1      benchmarkme_1.0.7 doParallel_1.0.17
## [5] iterators_1.0.14 foreach_1.5.2       scales_1.2.0      RcppAlgos_2.5.3
## [9]forcats_0.5.1   stringr_1.4.0       dplyr_1.0.9      purrr_0.3.4
## [13]readr_2.1.2    tidyverse_1.3.1     tibble_3.1.7      ggplot2_3.3.6
## [17]tidyverse_1.3.1 data.table_1.14.2 sensobol_1.1.1
##
## loaded via a namespace (and not attached):
## [1] Rcpp_1.0.8.3          lattice_0.20-45    lubridate_1.8.0
## [4] gmp_0.6-5             assertthat_0.2.1    digest_0.6.29
## [7] utf8_1.2.2            R6_2.5.1           cellranger_1.1.0
## [10]backports_1.4.1       reprex_2.0.1        evaluate_0.15
## [13]httr_1.4.3             pillar_1.7.0        Rdpack_2.3
## [16]rlang_1.0.2            readxl_1.4.0        rstudioapi_0.13
## [19]Matrix_1.4-1           tikzDevice_0.12.3.1 rmarkdown_2.14
## [22]munsell_0.5.0          broom_0.8.0         compiler_4.2.0
## [25]modelr_0.1.8           xfun_0.31          pkgconfig_2.0.3
## [28]htmltools_0.5.2        tidyselect_1.1.2    codetools_0.2-18
## [31]fansi_1.0.3            crayon_1.5.1       tzdb_0.3.0
## [34]dbplyr_2.1.1           withr_2.5.0         rbibutils_2.2.8
## [37]grid_4.2.0              jsonlite_1.8.0     gtable_0.3.0
## [40]lifecycle_1.0.1         DBI_1.1.2          magrittr_2.0.3
## [43]cli_3.3.0               stringi_1.7.6      fs_1.5.2
## [46]benchmarkmeData_1.0.4   xml2_1.3.3         ellipsis_0.3.2
## [49]generics_0.1.2          vctrs_0.4.1         tools_4.2.0
## [52]glue_1.6.2              hms_1.1.1          fastmap_1.1.0
## [55]yaml_2.3.5              colorspace_2.0-3    filehash_2.4-3
```

```
## [58] rvest_1.0.2           knitr_1.39          haven_2.5.0
## Return the machine CPU
cat("Machine:    "); print(get_cpu()$model_name)

## Machine:
## [1] "Apple M1 Max"
## Return number of true cores
cat("Num cores:   "); print(detectCores(logical = FALSE))

## Num cores:
## [1] 10
## Return number of threads
cat("Num threads: "); print(detectCores(logical = FALSE))

## Num threads:
## [1] 10
```