

# Package ‘CAISer’

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**Type** Package

**Title** Comparison of Algorithms with Iterative Sample Size Estimation

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**Description** Functions for performing experimental comparisons of algorithms using adequate sample sizes for power and accuracy. Implements the methodology originally presented in Campelo and Takahashi (2019)  [<doi:10.1007/s10732-018-9396-7>](https://doi.org/10.1007/s10732-018-9396-7) for the comparison of two algorithms, and later generalised in Campelo and Wanner (Submitted, 2019)  [<doi:10.48550/arXiv.1908.01720>](https://doi.org/10.48550/arXiv.1908.01720).

**License** GPL-2

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**VignetteBuilder** knitr

**URL** <https://fcampelo.github.io/CAISer/>

**BugReports** <https://github.com/fcampelo/CAISer/issues>

**NeedsCompilation** no

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boot_sdm	<i>Bootstrap the sampling distribution of the mean</i>
----------	--

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Description

Bootstraps the sampling distribution of the means for a given vector of observations

Usage

```
boot_sdm(x, boot.R = 999, ncpus = 1, seed = NULL)
```

Arguments

- x                      vector of observations
- boot.R                number of bootstrap resamples
- ncpus                 number of cores to use
- seed                  seed for the PRNG

Value

vector of bootstrap estimates of the sample mean

## References

- A.C. Davison, D.V. Hinkley: Bootstrap methods and their application. Cambridge University Press (1997)
- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. Journal of Heuristics 25(2):305-338, 2019.

## Author(s)

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

## Examples

```
x <- rnorm(15, mean = 4, sd = 1)
my.sdm <- boot_sdm(x)
hist(my.sdm, breaks = 30)
qqnorm(my.sdm, pch = 20)

x <- runif(12)
my.sdm <- boot_sdm(x)
qqnorm(my.sdm, pch = 20)

# Convergence of the SDM to a Normal distribution as sample size is increased
X <- rchisq(1000, df = 3)
x1 <- rchisq(10, df = 3)
x2 <- rchisq(20, df = 3)
x3 <- rchisq(40, df = 3)
par(mfrow = c(2, 2))
plot(density(X), main = "Estimated pop distribution");
hist(boot_sdm(x1), breaks = 25, main = "SDM, n = 10")
hist(boot_sdm(x2), breaks = 25, main = "SDM, n = 20")
hist(boot_sdm(x3), breaks = 25, main = "SDM, n = 40")
par(mfrow = c(1, 1))
```

---

calc_instances	<i>Calculates number of instances for the comparison of multiple algorithms</i>
----------------	---

---

## Description

Calculates either the number of instances, or the power(s) of the comparisons of multiple algorithms.

## Usage

```
calc_instances(
  ncomparisons,
  d,
  ninstances = NULL,
```

```

power = NULL,
sig.level = 0.05,
alternative.side = "two.sided",
test = "t.test",
power.target = "mean"
)

```

### Arguments

ncomparisons	number of comparisons planned
d	minimally relevant effect size (MRES, expressed as a standardized effect size, i.e., "deviation from H0" / "standard deviation")
ninstances	the number of instances to be used in the experiment.
power	target power for the comparisons (see Details)
sig.level	desired family-wise significance level (alpha) for the experiment
alternative.side	type of alternative hypothesis to be performed ("two.sided" or "one.sided")
test	type of test to be used ("t.test", "wilcoxon" or "binomial")
power.target	which comparison should have the desired power? Accepts "mean", "median", or "worst.case" (this last one is equivalent to the Bonferroni correction).

### Details

The main use of this routine uses the closed formula of the t-test to calculate the number of instances required for the comparison of pairs of algorithms, given a desired power and standardized effect size of interest. Significance levels of each comparison are adjusted using Holm's step-down correction (the default). The routine also takes into account whether the desired statistical power refers to the mean power (the default), median, or worst-case (which is equivalent to designing the experiment for the more widely-known Bonferroni correction). See the reference by Campelo and Wanner for details.

### Value

a list object containing the following items:

- ninstances - number of instances
- power - the power of the comparison
- d - the effect size
- sig.level - significance level
- alternative.side - type of alternative hypothesis
- test - type of test

### Sample Sizes for Nonparametric Methods

If the parameter `test` is set to either `Wilcoxon` or `Binomial`, this routine approximates the number of instances using the ARE of these tests in relation to the paired `t.test`, using the formulas (see reference by Campelo and Takahashi for details):

$$n.wilcox = n.ttest / 0.86 = 1.163 * n.ttest$$

$$n.binom = n.ttest / 0.637 = 1.570 * n.ttest$$

### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

### References

- P. Mathews. Sample size calculations: Practical methods for engineers and scientists. Mathews Malnar and Bailey, 2010.
- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. *Journal of Heuristics* 25(2):305-338, 2019.
- F. Campelo, E. Wanner: Sample size calculations for the experimental comparison of multiple algorithms on multiple problem instances. Submitted, *Journal of Heuristics*, 2019.

### Examples

```
# Calculate sample size for mean-case power
K      <- 10  # number of comparisons
alpha  <- 0.05 # significance level
power  <- 0.9  # desired power
d      <- 0.5  # MRES

out <- calc_instances(K, d,
                      power      = power,
                      sig.level = alpha)

# Plot power of each comparison to detect differences of magnitude d
plot(1:K, out$power,
     type = "b", pch = 20, las = 1, ylim = c(0, 1), xlab = "comparison",
     ylab = "power", xaxs = "i", xlim = c(0, 11))
grid(11, NA)
points(c(0, K+1), c(power, power), type = "l", col = 2, lty = 2, lwd = .5)
text(1, 0.93, sprintf("Mean power = %2.2f for N = %d",
                      out$mean.power, out$ninstances), adj = 0)

# Check sample size if planning for Wilcoxon tests:
calc_instances(K, d,
               power      = power,
               sig.level = alpha,
               test = "wilcoxon")$ninstances
```

```

# Calculate power profile for predefined sample size
N <- 45
out2 <- calc_instances(K, d, ninstances = N, sig.level = alpha)

points(1:K, out2$power, type = "b", pch = 19, col = 3)
text(6, .7, sprintf("Mean power = %2.2f for N = %d",
                    out2$mean.power, out2$ninstances), adj = 0)

# Sample size for worst-case (Bonferroni) power of 0.8, using Wilcoxon
out3 <- calc_instances(K, d, power = 0.9, sig.level = alpha,
                      test = "wilcoxon", power.target = "worst.case")
out3$ninstances

# For median power:
out4 <- calc_instances(K, d, power = 0.9, sig.level = alpha,
                      test = "wilcoxon", power.target = "median")
out4$ninstances
out4$power

```

---

calc_nreps	<i>Determine sample sizes for a set of algorithms on a single problem instance</i>
------------	--

---

## Description

Iteratively calculates the required sample sizes for K algorithms on a given problem instance, so that the standard errors of the estimates of the pairwise differences in performance is controlled at a predefined level.

## Usage

```

calc_nreps(
  instance,
  algorithms,
  se.max,
  dif = "simple",
  comparisons = "all.vs.all",
  method = "param",
  nstart = 20,
  nmax = 1000,
  seed = NULL,
  boot.R = 499,
  ncpus = 1,
  force.balanced = FALSE,
  load.folder = NA,
  save.folder = NA
)

```

**Arguments**

instance	a list object containing the definitions of the problem instance. See Section Instance for details.
algorithms	a list object containing the definitions of all algorithms. See Section Algorithms for details.
se.max	desired upper limit for the standard error of the estimated difference between pairs of algorithms. See Section Pairwise Differences for details.
dif	type of difference to be used. Accepts "perc" (for percent differences) or "simple" (for simple differences)
comparisons	type of comparisons being performed. Accepts "all.vs.first" (in which cases the first object in algorithms is considered to be the reference algorithm) or "all.vs.all" (if there is no reference and all pairwise comparisons are desired).
method	method to use for estimating the standard errors. Accepts "param" (for parametric) or "boot" (for bootstrap)
nstart	initial number of algorithm runs for each algorithm. See Section Initial Number of Observations for details.
nmax	maximum <b>total</b> allowed number of runs to execute. Loaded results (see load.folder below) do not count towards this total.
seed	seed for the random number generator
boot.R	number of bootstrap resamples to use (if method == "boot")
ncpus	number of cores to use
force.balanced	logical flag to force the use of balanced sampling for the algorithms on each instance
load.folder	name of folder to load results from. Use either "" or "/" for the current working directory. Accepts relative paths. Use NA for not saving. calc_nreps() will look for a .RDS file with the same name
save.folder	name of folder to save the results. Use either "" or "/" for the current working directory. Accepts relative paths. Use NA for not saving.

**Value**

a list object containing the following items:

- instance - alias for the problem instance considered
- Xk - list of observed performance values for all algorithms
- Nk - vector of sample sizes generated for each algorithm
- Diffk - data frame with point estimates, standard errors and other information for all algorithm pairs of interest
- seed - seed used for the PRNG
- dif - type of difference used
- method - method used ("param" / "boot")
- comparisons - type of pairings ("all.vs.all" / "all.vs.first")

### Instance

Parameter instance must be a named list containing all relevant parameters that define the problem instance. This list must contain at least the field `instance$FUN`, with the name of the function implementing the problem instance, that is, a routine that calculates  $y = f(x)$ . If the instance requires additional parameters, these must also be provided as named fields.

### Algorithms

Object `algorithms` is a list in which each component is a named list containing all relevant parameters that define an algorithm to be applied for solving the problem instance. In what follows `algorithm[[k]]` refers to any algorithm specified in the `algorithms` list.

`algorithm[[k]]` must contain an `algorithm[[k]]$FUN` field, which is a character object with the name of the function that calls the algorithm; as well as any other elements/parameters that `algorithm[[k]]$FUN` requires (e.g., stop criteria, operator names and parameters, etc.).

The function defined by the routine `algorithm[[k]]$FUN` must have the following structure: supposing that the list in `algorithm[[k]]` has fields `algorithm[[k]]$FUN = "myalgo"`, `algorithm[[k]]$par1 = "a"` and `algorithm$par2 = 5`, then:

```
myalgo <- function(par1, par2, instance, ...){
  # do stuff
  # ...
  return(results)
}
```

That is, it must be able to run if called as:

```
# remove '$FUN' and '$alias' fields from list of arguments
# and include the problem definition as field 'instance'
myargs      <- algorithm[names(algorithm) != "FUN"]
myargs      <- myargs[names(myargs) != "alias"]
myargs$instance <- instance

# call function
do.call(algorithm$FUN,
        args = myargs)
```

The `algorithm$FUN` routine must return a list containing (at least) the performance value of the final solution obtained, in a field named `value` (e.g., `result$value`) after a given run.

### Initial Number of Observations

In the **general case** the initial number of observations per algorithm (`nstart`) should be relatively high. For the parametric case we recommend between 10 and 20 if outliers are not expected, or between 30 and 50 if that assumption cannot be made. For the bootstrap approach we recommend using at least 20. However, if some distributional assumptions can be made - particularly low



skewness of the population of algorithm results on the test instances), then `nstart` can in principle be as small as 5 (if the output of the algorithms were known to be normal, it could be 1).

In general, higher sample sizes are the price to pay for abandoning distributional assumptions. Use lower values of `nstart` with caution.

### Pairwise Differences

Parameter `dif` informs the type of difference in performance to be used for the estimation ( $\mu_a$  and  $\mu_b$  represent the mean performance of any two algorithms on the test instance, and  $\mu$  represents the grand mean of all algorithms given in `algorithms`):

- If `dif == "perc"` and `comparisons == "all.vs.first"`, the estimated quantity is  $\phi_{1b} = (\mu_1 - \mu_b)/\mu_1 = 1 - (\mu_b/\mu_1)$ .
- If `dif == "perc"` and `comparisons == "all.vs.all"`, the estimated quantity is  $\phi_{ab} = (\mu_a - \mu_b)/\mu$ .
- If `dif == "simple"` it estimates  $\mu_a - \mu_b$ .

### Author(s)

Felipe Campelo (<fcampelo@gmail.com>)

### References

- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. *Journal of Heuristics* 25(2):305-338, 2019.
- P. Mathews. Sample size calculations: Practical methods for engineers and scientists. Mathews Malnar and Bailey, 2010.
- A.C. Davison, D.V. Hinkley: Bootstrap methods and their application. Cambridge University Press (1997)
- E.C. Fieller: Some problems in interval estimation. *Journal of the Royal Statistical Society. Series B (Methodological)* 16(2), 175–185 (1954)
- V. Franz: Ratios: A short guide to confidence limits and proper use (2007). <https://arxiv.org/pdf/0710.2024v1.pdf>
- D.C. Montgomery, C.G. Runger: Applied Statistics and Probability for Engineers, 6th ed. Wiley (2013)

### Examples

```
# Example using dummy algorithms and instances. See ?dummyalgo for details.
# We generate dummy algorithms with true means 15, 10, 30, 15, 20; and true
# standard deviations 2, 4, 6, 8, 10.
algorithms <- mapply(FUN = function(i, m, s){
  list(FUN = "dummyalgo",
       alias = paste0("algo", i),
       distribution.fun = "rnorm",
       distribution.pars = list(mean = m, sd = s)),
  i = c(alg1 = 1, alg2 = 2, alg3 = 3, alg4 = 4, alg5 = 5),
  m = c(15, 10, 30, 15, 20),
  s = c(2, 4, 6, 8, 10),
```

```

SIMPLIFY = FALSE)

# Make a dummy instance with a centered (zero-mean) exponential distribution:
instance = list(FUN = "dummyinstance", distr = "rexp", rate = 5, bias = -1/5)

# Explicitate all other parameters (just this one time:
# most have reasonable default values)
myreps <- calc_nreps(instance = instance,
                    algorithms = algorithms,
                    se.max     = 0.05,      # desired (max) standard error
                    dif        = "perc",    # type of difference
                    comparisons = "all.vs.all", # differences to consider
                    method     = "param",   # method ("param", "boot")
                    nstart     = 15,        # initial number of samples
                    nmax       = 1000,      # maximum allowed sample size
                    seed       = 1234,      # seed for PRNG
                    boot.R     = 499,       # number of bootstrap resamples (unused)
                    ncpus      = 1,         # number of cores to use
                    force.balanced = FALSE, # force balanced sampling?
                    load.folder = NA,       # file to load results from
                    save.folder = NA)       # folder to save results

summary(myreps)
plot(myreps)

```

---

calc\_se

---

*Calculates the standard error for simple and percent differences*


---

## Description

Calculates the sample standard error for the estimator differences between multiple algorithms on a given instance.

## Usage

```

calc_se(
  Xk,
  dif = "simple",
  comparisons = "all.vs.all",
  method = "param",
  boot.R = 999
)

```

## Arguments

Xk	list object where each position contains a vector of observations of algorithm k on a given problem instance.
dif	name of the difference for which the SEs are desired. Accepts "perc" (for percent differences) or "simple" (for simple differences)

comparisons	standard errors to be calculated. Accepts "all.vs.first" (in which cases the first object in <code>algorithms</code> is considered to be the reference algorithm) or "all.vs.all" (if there is no reference and all pairwise SEs are desired).
method	method used to calculate the interval. Accepts "param" (using parametric formulas based on normality of the sampling distribution of the means) or "boot" (for bootstrap).
boot.R	(optional) number of bootstrap resamples (if <code>method == "boot"</code> )

### Details

- If `dif == "perc"` it returns the standard errors for the sample estimates of pairs  $(\mu_2 - \mu_1)/\mu$ , where  $\mu_1, \mu_2$  are the means of the populations that generated sample vectors  $x_1, x_2$ , and
- If `dif == "simple"` it returns the SE for sample estimator of  $(\mu_2 - \mu_1)$

### Value

a list object containing the following items:

- `Phi.est` - estimated values of the statistic of interest for each pair of algorithms of interest (all pairs if `comparisons == "all.vs.all"`, or all pairs containing the first algorithm if `comparisons == "all.vs.first"`).
- `se` - standard error estimates

### References

- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. *Journal of Heuristics* 25(2):305-338, 2019.

### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

### Examples

```
# three vectors of normally distributed observations
set.seed(1234)
Xk <- list(rnorm(10, 5, 1), # mean = 5, sd = 1,
          rnorm(20, 10, 2), # mean = 10, sd = 2,
          rnorm(50, 15, 5)) # mean = 15, sd = 3

calc_se(Xk, dif = "simple", comparisons = "all.vs.all", method = "param")
calc_se(Xk, dif = "simple", comparisons = "all.vs.all", method = "boot")

calc_se(Xk, dif = "perc", comparisons = "all.vs.first", method = "param")
calc_se(Xk, dif = "perc", comparisons = "all.vs.first", method = "boot")

calc_se(Xk, dif = "perc", comparisons = "all.vs.all", method = "param")
calc_se(Xk, dif = "perc", comparisons = "all.vs.all", method = "boot")
```

---

 consolidate\_partial\_results

*Consolidate results from partial files*


---

### Description

Consolidates results from a set of partial files (each generated by an individual call to `calc_nreps()`) into a single output structure, similar (but not identical) to the output of `run_experiment()`. This is useful e.g., to consolidate the results from interrupted experiments.

### Usage

```
consolidate_partial_results(Configuration, folder = "./nreps_files")
```

### Arguments

`Configuration` a named list containing all parameters required in a call to `run_experiment()` except instances and algorithms. See the parameter list and default values in `run_experiment()`. Notice that this is always returned as part of the output structure of `run_experiment()`, so it generally easier to just retrieve it from previously saved results.

`folder` folder where the partial files are located.

### Value

a list object containing the following fields:

- `data.raw` - data frame containing all observations generated
- `data.summary` - data frame summarizing the experiment.
- `N` - number of instances sampled
- `total.runs` - total number of algorithm runs performed
- `instances.sampled` - names of the instances sampled

---

 dummyalgo

*Dummy algorithm routine to test the sampling procedures*


---

### Description

This is a dummy algorithm routine to test the sampling procedures, in combination with `dummyinstance()`. `dummyalgo()` receives two parameters that determine the distribution of performances it will exhibit on a hypothetical problem class: `distribution.fun` (with the name of a random number generation function, e.g. `rnorm`, `runif`, `rexp` etc.); and `distribution.pars`, a named list of parameters to be passed on to `distribution.fun`. The third parameter is an instance object (see `calc_nreps()` for details), which is a named list with the following fields:

- FUN = "dummyinstance" - must always be "dummyinstance" (will be ignored otherwise).
- distr - the name of a random number generation function.
- ... - other named fields with parameters to be passed down to the function in distr.

### Usage

```
dummyalgo(
  distribution.fun = "rnorm",
  distribution.pars = list(mean = 0, sd = 1),
  instance = list(FUN = "dummyinstance", distr = "rnorm", mean = 0, sd = 1)
)
```

### Arguments

distribution.fun	name of a function that generates random values according to a given distribution, e.g., "rnorm", "runif", "rexp" etc.
distribution.pars	list of named parameters required by the function in distribution.fun. Parameter n (number of points to generate) is unnecessary (this routine always forces n = 1).
instance	instance parameters (see Details).

### Details

distribution.fun and distribution.pars regulate the mean performance of the dummy algorithm on a given (hypothetical) problem class, and the between-instances variance of performance. The instance specification in instance regulates the within-instance variability of results. Ideally the distribution parameters passed to the instance should result in a within-instance distribution of values with zero mean, so that the mean of the values returned by dummyalgo is regulated only by distribution.fun and distribution.pars.

The value returned by dummyalgo is sampled as follows:

```
offset <- do.call(distribution.fun, args = distribution.pars)
y <- offset + do.call("dummyinstance", args = instance)
```

### Value

a list object with a single field \$value, containing a scalar numerical value distributed as described at the end of Details.

### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

### See Also

[dummyinstance\(\)](#)

## Examples

```
# Make a dummy instance with a centered (zero-mean) exponential distribution:
instance = list(FUN = "dummyinstance", distr = "rexp", rate = 5, bias = -1/5)

# Simulate a dummy algorithm that has a uniform distribution of expected
# performance values, between -25 and 50.
dummyalgo(distribution.fun = "runif",
          distribution.pars = list(min = -25, max = 50),
          instance = instance)
```

---

dummyinstance

*Dummy instance routine to test the sampling procedures*

---

## Description

This is a dummy instance routine to test the sampling procedures, in combination with [dummyalgo\(\)](#). `dummyinstance()` receives a parameter `distr` containing the name of a random number generation function (e.g. `rnorm`, `runif`, `rexp` etc.), plus a variable number of arguments to be passed down to the function in `distr`.

## Usage

```
dummyinstance(distr, ..., bias = 0)
```

## Arguments

<code>distr</code>	name of a function that generates random values according to a given distribution, e.g., "rnorm", "runif", "rexp" etc.
<code>...</code>	additional parameters to be passed down to the function in <code>distr</code> . Parameter <code>n</code> (number of points to generate) is unnecessary (this routine always forces <code>n = 1</code> ).
<code>bias</code>	a bias term to add to the results of the distribution function (e.g., to set the mean to zero).

## Value

a single numeric value sampled from the desired distribution.

## Author(s)

Felipe Campelo (<[fcampelo@ufmg.br](mailto:fcampelo@ufmg.br)>, <[f.campelo@aston.ac.uk](mailto:f.campelo@aston.ac.uk)>)

## See Also

[dummyalgo\(\)](#)

**Examples**

```
dummyinstance(distr = "rnorm", mean = 10, sd = 1)

# Make a centered (zero-mean) exponential distribution:
lambda = 4

# 10000 observations
set.seed(1234)
y <- numeric(10000)
for (i in 1:10000) y[i] <- dummyinstance(distr = "rexp", rate = lambda,
                                         bias = -1/lambda)

mean(y)
hist(y, breaks = 50, xlim = c(-0.5, 2.5))
```

example\_SANN

*Simulated annealing (for testing/examples)***Description**

Adapted from stats::optim(). Check their documentation / examples for details.

**Usage**

```
example_SANN(Temp, budget, instance)
```

**Arguments**

Temp	controls the "SANN" method. It is the starting temperature for the cooling schedule.
budget	stop criterion: number of function evaluations to execute
instance	an instance object (see <a href="#">calc_nreps()</a> for details)

**Examples**

```
## Not run:
instance <- list(FUN = "TSP.dist", mydist = datasets::eurodist)

example_SANN(Temp = 2000, budget = 10000, instance = instance)

## End(Not run)
```

---

get_observations	<i>Run an algorithm on a problem.</i>
------------------	---------------------------------------

---

## Description

Call algorithm routine for the solution of a problem instance

## Usage

```
get_observations(algo, instance, n = 1)
```

## Arguments

algo	a list object containing the definitions of the algorithm. See <a href="#">calc_nreps()</a> for details.
instance	a list object containing the definitions of the problem instance. See <a href="#">calc_nreps()</a> for details.
n	number of observations to generate.

## Value

vector of observed performance values

## Author(s)

Felipe Campelo (<[fcampelo@ufmg.br](mailto:fcampelo@ufmg.br)>, <[f.campelo@aston.ac.uk](mailto:f.campelo@aston.ac.uk)>)

## See Also

[calc\\_nreps\(\)](#)

## Examples

```
# Make a dummy instance with a centered (zero-mean) exponential distribution:
instance <- list(FUN = "dummyinstance", distr = "rexp", rate = 5, bias = -1/5)

# Simulate a dummy algorithm that has a uniform distribution of expected
# performance values, between -25 and 50.
algorithm <- list(FUN = "dummyalgo",
                  distribution.fun = "runif",
                  distribution.pars = list(min = -25, max = 50))
x <- get_observations(algorithm, instance, n = 1000)
hist(x)
```



plot.CAISEr

plot.CAISEr

**Description**

S3 method for plotting *CAISEr* objects output by `run_experiment()`.

**Usage**

```
## S3 method for class 'CAISEr'
plot(
  x,
  y = NULL,
  ...,
  latex = FALSE,
  reorder = FALSE,
  show.text = TRUE,
  digits = 3,
  layout = NULL
)
```

**Arguments**

<code>x</code>	list object of class <i>CAISEr</i> .
<code>y</code>	unused. Included for consistency with generic plot method.
<code>...</code>	other parameters to be passed down to specific plotting functions (currently unused)
<code>latex</code>	logical: should labels be formatted for LaTeX? (useful for later saving using library <i>TikzDevice</i> )
<code>reorder</code>	logical: should the comparisons be reordered alphabetically?
<code>show.text</code>	logical: should text be plotted?
<code>digits</code>	how many significant digits should be used in text?
<code>layout</code>	optional parameter to override the layout of the plots (see <code>gridExtra::arrangeGrob()</code> for details. The default layout is <code>lay = rbind(c(1,1,1,1,1,1), c(1,1,1,1,1,1), c(2,2,2,3,3,3))</code>

**Value**

list containing (1) a list of *ggplot2* objects generated, and (2) a list of data frames used for the creation of the plots.

plot.nreps

*plot.nreps***Description**

S3 method for plotting *nreps* objects output by `calc_nreps()`.

**Usage**

```
## S3 method for class 'nreps'
plot(
  x,
  y = NULL,
  ...,
  instance.name = NULL,
  latex = FALSE,
  show.SE = TRUE,
  show.CI = TRUE,
  sig.level = 0.05,
  show.text = TRUE
)
```

**Arguments**

<code>x</code>	list object of class <i>nreps</i> (generated by <code>calc_nreps()</code> ) or of class <i>CAISER</i> (in which case an <code>instance.name</code> must be provided).
<code>y</code>	unused. Included for consistency with generic <code>plot</code> method.
<code>...</code>	other parameters to be passed down to specific plotting functions (currently unused)
<code>instance.name</code>	name for instance to be plotted if object is of class <i>CAISER</i> . Ignored otherwise.
<code>latex</code>	logical: should labels be formatted for LaTeX? (useful for later saving using library <i>TikzDevice</i> )
<code>show.SE</code>	logical: should standard errors be plotted?
<code>show.CI</code>	logical: should confidence intervals be plotted?
<code>sig.level</code>	significance level for the confidence interval. $0 < \text{sig.level} < 1$
<code>show.text</code>	logical: should text be plotted?

**Value**

ggplot object (invisibly)

print.CAISEr

*print.CAISEr***Description**

S3 method for printing *CAISEr* objects (the output of `run_experiment()`).

**Usage**

```
## S3 method for class 'CAISEr'
print(x, ..., echo = TRUE, digits = 4, right = TRUE, breakrows = FALSE)
```

**Arguments**

<code>x</code>	list object of class <i>CAISEr</i> (generated by <code>run_experiment()</code> )
<code>...</code>	other parameters to be passed down to specific summary functions (currently unused)
<code>echo</code>	logical flag: should the print method actually print to screen?
<code>digits</code>	the minimum number of significant digits to be used. See <code>print.default()</code> .
<code>right</code>	logical, indicating whether or not strings should be right-aligned.
<code>breakrows</code>	logical, indicating whether to "widen" the output table by placing the bottom half to the right of the top half.

**Value**

data frame object containing the summary table (invisibly)

**Examples**

```
# Example using four dummy algorithms and 100 dummy instances.
# See [dummyalgo()] and [dummyinstance()] for details.
# Generating 4 dummy algorithms here, with means 15, 10, 30, 15 and standard
# deviations 2, 4, 6, 8.
algorithms <- mapply(FUN = function(i, m, s){
  list(FUN = "dummyalgo",
        alias = paste0("algo", i),
        distribution.fun = "rnorm",
        distribution.pars = list(mean = m, sd = s)),
    i = c(alg1 = 1, alg2 = 2, alg3 = 3, alg4 = 4),
    m = c(15, 10, 30, 15),
    s = c(2, 4, 6, 8),
    SIMPLIFY = FALSE)

# Generate 100 dummy instances with centered exponential distributions
instances <- lapply(1:100,
  function(i) {rate <- runif(1, 1, 10)
    list(FUN = "dummyinstance",
```

```

        alias = paste0("Inst.", i),
        distr = "rexp", rate = rate,
        bias = -1 / rate))

my.results <- run_experiment(instances, algorithms,
                            d = 1, se.max = .1,
                            power = .9, sig.level = .05,
                            power.target = "mean",
                            dif = "perc", comparisons = "all.vs.all",
                            seed = 1234, ncpus = 1)

my.results

```

---

run_experiment	<i>Run a full experiment for comparing multiple algorithms using multiple instances</i>
----------------	---

---

## Description

Design and run a full experiment - calculate the required number of instances, run the algorithms on each problem instance using the iterative approach based on optimal sample size ratios, and return the results of the experiment. This routine builds upon [calc\\_instances\(\)](#) and [calc\\_nreps\(\)](#), so refer to the documentation of these two functions for details.

## Usage

```

run_experiment(
  instances,
  algorithms,
  d,
  se.max,
  power = 0.8,
  sig.level = 0.05,
  power.target = "mean",
  dif = "simple",
  comparisons = "all.vs.all",
  alternative = "two.sided",
  test = "t.test",
  method = "param",
  nstart = 20,
  nmax = 100 * length(algorithms),
  force.balanced = FALSE,
  ncpus = 2,
  boot.R = 499,
  seed = NULL,
  save.partial.results = NA,
  load.partial.results = NA,

```

```

    save.final.result = NA
)

```

## Arguments

instances	list object containing the definitions of the <i>available</i> instances. This list may (or may not) be exhausted in the experiment. To estimate the number of required instances, see <a href="#">calc_instances()</a> . For more details, see Section Instance List.
algorithms	a list object containing the definitions of all algorithms. See Section Algorithms for details.
d	minimally relevant effect size (MRES), expressed as a standardized effect size, i.e., "deviation from H0" / "standard deviation". See <a href="#">calc_instances()</a> for details.
se.max	desired upper limit for the standard error of the estimated difference between pairs of algorithms. See Section Pairwise Differences for details.
power	(desired) test power. See <a href="#">calc_instances()</a> for details. Any value equal to or greater than one will force the method to use all available instances in Instance.list.
sig.level	family-wise significance level (alpha) for the experiment. See <a href="#">calc_instances()</a> for details.
power.target	which comparison should have the desired power? Accepts "mean", "median", or "worst.case" (this last one is equivalent to the Bonferroni correction).
dif	type of difference to be used. Accepts "perc" (for percent differences) or "simple" (for simple differences)
comparisons	type of comparisons being performed. Accepts "all.vs.first" (in which cases the first object in algorithms is considered to be the reference algorithm) or "all.vs.all" (if there is no reference and all pairwise comparisons are desired).
alternative	type of alternative hypothesis ("two.sided" or "less" or "greater"). See <a href="#">calc_instances()</a> for details.
test	type of test to be used ("t.test", "wilcoxon" or "binomial")
method	method to use for estimating the standard errors. Accepts "param" (for parametric) or "boot" (for bootstrap)
nstart	initial number of algorithm runs for each algorithm. See Section Initial Number of Observations for details.
nmax	maximum number of runs to execute on each instance (see <a href="#">calc_nreps()</a> ). Loaded results (see load.partial.results below) do not count towards this maximum.
force.balanced	logical flag to force the use of balanced sampling for the algorithms on each instance
ncpus	number of cores to use
boot.R	number of bootstrap resamples to use (if method == "boot")
seed	seed for the random number generator

**save.partial.results**

should partial results be saved to files? Can be either NA (do not save) or a character string pointing to a folder. File names are generated based on the instance aliases. **Existing files with matching names will be overwritten.** run\_experiment() uses **.RDS** files for saving and loading.

**load.partial.results**

should partial results be loaded from files? Can be either NA (do not save) or a character string pointing to a folder containing the file(s) to be loaded. run\_experiment() will use **.RDS** file(s) with a name(s) matching instance aliases. run\_experiment() uses **.RDS** files for saving and loading.

**save.final.result**

should the final results be saved to file? Can be either NA (do not save) or a character string pointing to a folder where the results will be saved on a **.RDS** file starting with CAISer\_results\_ and ending with 12-digit datetime tag in the format YYYYMMDDhhmmss.

**Value**

a list object containing the following fields:

- Configuration - the full input configuration (for reproducibility)
- data.raw - data frame containing all observations generated
- data.summary - data frame summarizing the experiment.
- N - number of instances sampled
- N.star - number of instances required
- total.runs - total number of algorithm runs performed
- instances.sampled - names of the instances sampled
- Underpowered - flag: TRUE if  $N < N.star$

**Instance List**

Parameter instances must contain a list of instance objects, where each field is itself a list, as defined in the documentation of function [calc\\_nreps\(\)](#). In short, each element of instances is an instance, i.e., a named list containing all relevant parameters that define the problem instance. This list must contain at least the field instance\$FUN, with the name of the problem instance function, that is, a routine that calculates  $y = f(x)$ . If the instance requires additional parameters, these must also be provided as named fields. An additional field, "instance\$alias", can be used to provide the instance with a unique identifier (e.g., when using an instance generator).

**Algorithm List**

Object algorithms is a list in which each component is a named list containing all relevant parameters that define an algorithm to be applied for solving the problem instance. In what follows algorithms[[k]] refers to any algorithm specified in the algorithms list.

algorithms[[k]] must contain an algorithms[[k]]\$FUN field, which is a character object with the name of the function that calls the algorithm; as well as any other elements/parameters that algorithms[[k]]\$FUN requires (e.g., stop criteria, operator names and parameters, etc.).

The function defined by the routine `algorithms[[k]]$FUN` must have the following structure: supposing that the list in `algorithms[[k]]` has fields `algorithm[[k]]$FUN = "myalgo"`, `algorithms[[k]]$par1 = "a"` and `algorithms[[k]]$par2 = 5`, then:

```
myalgo <- function(par1, par2, instance, ...){
  #
  # <do stuff>
  #
  return(results)
}
```

That is, it must be able to run if called as:

```
# remove '$FUN' and '$alias' field from list of arguments
# and include the problem definition as field 'instance'
myargs      <- algorithm[names(algorithm) != "FUN"]
myargs      <- myargs[names(myargs) != "alias"]
myargs$instance <- instance

# call function
do.call(algorithm$FUN,
        args = myargs)
```

The `algorithm$FUN` routine must return a list containing (at least) the performance value of the final solution obtained, in a field named `value` (e.g., `result$value`) after a given run. In general it is easier to write a small wrapper function around existing implementations.

### Initial Number of Observations

In the *general case* the initial number of observations / algorithm / instance (`nstart`) should be relatively high. For the parametric case we recommend 10~15 if outliers are not expected, and 30~40 (at least) if that assumption cannot be made. For the bootstrap approach we recommend using at least 15 or 20. However, if some distributional assumptions can be made - particularly low skewness of the population of algorithm results on the test instances), then `nstart` can in principle be as small as 5 (if the output of the algorithm were known to be normal, it could be 1).

In general, higher sample sizes are the price to pay for abandoning distributional assumptions. Use lower values of `nstart` with caution.

### Pairwise Differences

Parameter `dif` informs the type of difference in performance to be used for the estimation ( $\mu_a$  and  $\mu_b$  represent the mean performance of any two algorithms on the test instance, and  $\mu$  represents the grand mean of all algorithms given in `algorithms`):

- If `dif == "perc"` and `comparisons == "all.vs.first"`, the estimated quantity is:  $\phi_{1b} = (\mu_1 - \mu_b) / \mu_1 = 1 - (\mu_b / \mu_1)$ .

- If `dif == "perc"` and `comparisons == "all.vs.all"`, the estimated quantity is:  $\phi_{ab} = (\mu_a - \mu_b) / \mu$ .
- If `dif == "simple"` it estimates  $\mu_a - \mu_b$ .

### Sample Sizes for Nonparametric Methods

If the parameter `"` is set to either Wilcoxon or 'Binomial', this routine approximates the number of instances using the ARE of these tests in relation to the paired t.test:

- `n.wilcox = n.ttest / 0.86 = 1.163 * n.ttest`
- `n.binom = n.ttest / 0.637 = 1.570 * n.ttest`

### Author(s)

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

### References

- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. *Journal of Heuristics* 25(2):305-338, 2019.
- P. Mathews. Sample size calculations: Practical methods for engineers and scientists. Mathews Malnar and Bailey, 2010.
- A.C. Davison, D.V. Hinkley: Bootstrap methods and their application. Cambridge University Press (1997)
- E.C. Fieller: Some problems in interval estimation. *Journal of the Royal Statistical Society. Series B (Methodological)* 16(2), 175–185 (1954)
- V. Franz: Ratios: A short guide to confidence limits and proper use (2007). <https://arxiv.org/pdf/0710.2024v1.pdf>
- D.C. Montgomery, C.G. Runger: Applied Statistics and Probability for Engineers, 6th ed. Wiley (2013)
- D.J. Sheskin: Handbook of Parametric and Nonparametric Statistical Procedures, 4th ed., Chapman & Hall/CRC, 1996.

### Examples

```
# Example using four dummy algorithms and 100 dummy instances.
# See [dummyalgo()] and [dummyinstance()] for details.
# Generating 4 dummy algorithms here, with means 15, 10, 30, 15 and standard
# deviations 2, 4, 6, 8.
algorithms <- mapply(FUN = function(i, m, s){
  list(FUN = "dummyalgo",
       alias = paste0("algo", i),
       distribution.fun = "rnorm",
       distribution.pars = list(mean = m, sd = s))},
  i = c(alg1 = 1, alg2 = 2, alg3 = 3, alg4 = 4),
  m = c(15, 10, 30, 15),
  s = c(2, 4, 6, 8),
  SIMPLIFY = FALSE)
```



```
# Generate 100 dummy instances with centered exponential distributions
instances <- lapply(1:100,
  function(i) {rate <- runif(1, 1, 10)
    list(FUN = "dummyinstance",
      alias = paste0("Inst.", i),
      distr = "rexp", rate = rate,
      bias = -1 / rate)})

my.results <- run_experiment(instances, algorithms,
  d = .5, se.max = .1,
  power = .9, sig.level = .05,
  power.target = "mean",
  dif = "perc", comparisons = "all.vs.all",
  ncpus = 1, seed = 1234)

# Take a look at the results
summary(my.results)
plot(my.results)
```

se\_boot

*Bootstrap standard errors***Description**

Calculates the standard errors of a given statistic using bootstrap

**Usage**

```
se_boot(Xk, dif = "simple", comparisons = "all.vs.all", boot.R = 999, ...)
```

**Arguments**

Xk	list object where each position contains a vector of observations of algorithm k on a given problem instance.
dif	name of the difference for which the SEs are desired. Accepts "perc" (for percent differences) or "simple" (for simple differences)
comparisons	standard errors to be calculated. Accepts "all.vs.first" (in which cases the first object in <code>algorithms</code> is considered to be the reference algorithm) or "all.vs.all" (if there is no reference and all pairwise SEs are desired).
boot.R	(optional) number of bootstrap resamples (if <code>method == "boot"</code> )
...	other parameters (used only for compatibility with calls to <code>se_param()</code> , unused in this function)

**Value**

Data frame containing, for each pair of interest, the estimated difference (column "Phi") and the sample standard error (column "SE")

References

- A.C. Davison, D.V. Hinkley: Bootstrap methods and their application. Cambridge University Press (1997)
- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. Journal of Heuristics 25(2):305-338, 2019.

Author(s)

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

Examples

```
# three vectors of normally distributed observations
set.seed(1234)
Xk <- list(rnorm(10, 5, 1), # mean = 5, sd = 1,
          rnorm(20, 10, 2), # mean = 10, sd = 2,
          rnorm(20, 15, 5)) # mean = 15, sd = 3

se_boot(Xk, dif = "simple", comparisons = "all.vs.all")
se_boot(Xk, dif = "perc", comparisons = "all.vs.first")
se_boot(Xk, dif = "perc", comparisons = "all.vs.all")
```

---

se_param	<i>Parametric standard errors</i>
----------	-----------------------------------

---

Description

Calculates the standard errors of a given statistic using parametric formulas

Usage

```
se_param(Xk, dif = "simple", comparisons = "all.vs.all", ...)
```

Arguments

Xk	list object where each position contains a vector of observations of algorithm k on a given problem instance.
dif	name of the difference for which the SEs are desired. Accepts "perc" (for percent differences) or "simple" (for simple differences)
comparisons	standard errors to be calculated. Accepts "all.vs.first" (in which cases the first object in algorithms is considered to be the reference algorithm) or "all.vs.all" (if there is no reference and all pairwise SEs are desired).
...	other parameters (used only for compatibility with calls to <code>se_boot()</code> , unused in this function)

**Value**

Data frame containing, for each pair of interest, the estimated difference (column "Phi") and the sample standard error (column "SE")

**References**

- E.C. Fieller: Some problems in interval estimation. Journal of the Royal Statistical Society. Series B (Methodological) 16(2), 175–185 (1954)
- V. Franz: Ratios: A short guide to confidence limits and proper use (2007). <https://arxiv.org/pdf/0710.2024v1.pdf>
- D.C. Montgomery, C.G. Runger: Applied Statistics and Probability for Engineers, 6th ed. Wiley (2013)
- F. Campelo, F. Takahashi: Sample size estimation for power and accuracy in the experimental comparison of algorithms. Journal of Heuristics 25(2):305-338, 2019.

**Author(s)**

Felipe Campelo (<fcampelo@ufmg.br>, <f.campelo@aston.ac.uk>)

**Examples**

```
# three vectors of normally distributed observations
set.seed(1234)
Xk <- list(rnorm(10, 5, 1), # mean = 5, sd = 1,
          rnorm(20, 10, 2), # mean = 10, sd = 2,
          rnorm(20, 15, 5)) # mean = 15, sd = 3

se_param(Xk, dif = "simple", comparisons = "all.vs.all")
se_param(Xk, dif = "perc", comparisons = "all.vs.first")
se_param(Xk, dif = "perc", comparisons = "all.vs.all")
```

---

summary.CAISEr	<i>summary.CAISEr</i>
----------------	-----------------------

---

**Description**

S3 method for summarizing *CAISEr* objects output by `run_experiment()`. Input parameters `test`, `alternative` and `sig.level` can be used to override the ones used in the call to `run_experiment()`.

**Usage**

```
## S3 method for class 'CAISEr'
summary(object, test = NULL, alternative = NULL, sig.level = NULL, ...)
```

**Arguments**

object	list object of class <i>CAISEr</i> (generated by <code>run_experiment()</code> )
test	type of test to be used ("t.test", "wilcoxon" or "binomial")
alternative	type of alternative hypothesis ("two.sided" or "less" or "greater"). See <code>calc_instances()</code> for details.
sig.level	desired family-wise significance level (alpha) for the experiment
...	other parameters to be passed down to specific summary functions (currently unused)

**Value**

A list object is returned invisibly, containing the details of all tests performed as well as information on the total number of runs dedicated to each algorithm.

**Examples**

```
# Example using four dummy algorithms and 100 dummy instances.
# See [dummyalgo()] and [dummyinstance()] for details.
# Generating 4 dummy algorithms here, with means 15, 10, 30, 15 and standard
# deviations 2, 4, 6, 8.
algorithms <- mapply(FUN = function(i, m, s){
  list(FUN = "dummyalgo",
        alias = paste0("algo", i),
        distribution.fun = "rnorm",
        distribution.pars = list(mean = m, sd = s)),
  i = c(alg1 = 1, alg2 = 2, alg3 = 3, alg4 = 4),
  m = c(15, 10, 30, 15),
  s = c(2, 4, 6, 8),
  SIMPLIFY = FALSE)

# Generate 100 dummy instances with centered exponential distributions
instances <- lapply(1:100,
  function(i) {rate <- runif(1, 1, 10)
    list(FUN = "dummyinstance",
          alias = paste0("Inst.", i),
          distr = "rexp", rate = rate,
          bias = -1 / rate)})

my.results <- run_experiment(instances, algorithms,
  d = 1, se.max = .1,
  power = .9, sig.level = .05,
  power.target = "mean",
  dif = "perc", comparisons = "all.vs.all",
  seed = 1234, ncpus = 1)

summary(my.results)

# You can override some defaults if you want:
summary(my.results, test = "wilcoxon")
```

---

summary.nreps	<i>summary.nreps</i>
---------------	----------------------

---

**Description**

S3 method for summarizing *nreps* objects output by [calc\\_nreps\(\)](#).

**Usage**

```
## S3 method for class 'nreps'  
summary(object, ...)
```

**Arguments**

object	list object of class <i>nreps</i> (generated by <a href="#">calc_nreps()</a> )
...	other parameters to be passed down to specific summary functions (currently unused)

---

TSP.dist	<i>TSP instance generator (for testing/examples)</i>
----------	--

---

**Description**

Adapted from stats::optim(). Check their documentation / examples for details.

**Usage**

```
TSP.dist(x, mydist)
```

**Arguments**

x	a valid closed route for the TSP instance
mydist	object of class <i>dist</i> defining the TSP instance

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