

Package ‘simer’

January 30, 2025

Title Data Simulation for Life Science and Breeding

Version 0.9.0.6

Date 2025-1-29

Description Data simulator including genotype, phenotype, pedigree, selection and reproduction in R. It simulates most of reproduction process of animals or plants and provides data for GS (Genomic Selection), GWAS (Genome-Wide Association Study), and Breeding.
For ADI model, please see Kao C and Zeng Z (2002) <[doi:10.1093/genetics/160.3.1243](https://doi.org/10.1093/genetics/160.3.1243)>.
For build.cov, please see B. D. Ripley (1987) <ISBN:9780470009604>.

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URL <https://github.com/xiaolei-lab/SIMER>

BugReports <https://github.com/xiaolei-lab/SIMER/issues>

Imports utils, stats, Matrix, methods, MASS, bigmemory, jsonlite, Rcpp

LinkingTo Rcpp, RcppArmadillo, RcppProgress, BH, bigmemory

Depends R (>= 3.5.0)

Suggests knitr, igraph

RoxygenNote 7.3.2

Encoding UTF-8

NeedsCompilation yes

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Repository CRAN

Date/Publication 2025-01-30 09:50:03 UTC

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annotation *Annotation simulation*

Description

Generating a map with annotation information

Usage

annotation(SP, verbose = TRUE)

Arguments

SP a list of all simulation parameters.
verbose whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Jul 10, 2022

Value

the function returns a list containing

\$map\$pop.map the map data with annotation information.

\$map\$species the species of genetic map, which can be "arabidopsis", "cattle", "chicken", "dog", "horse", "human", "maize", "mice", "pig", and "rice".

\$map\$pop.marker the number of markers.

\$map\$num.chr the number of chromosomes.

\$map\$len.chr the length of chromosomes.

\$map\$qtn.model the genetic model of QTN such as 'A + D'.

\$map\$qtn.index the QTN index for each trait.

\$map\$qtn.num the QTN number for (each group in) each trait.

\$map\$qtn.dist the QTN distribution containing 'norm', 'geom', 'gamma' or 'beta'.

\$map\$qtn.var the variances for normal distribution.

\$map\$qtn.prob the probability of success for geometric distribution.

\$map\$qtn.shape the shape parameter for gamma distribution.

\$map\$qtn.scale the scale parameter for gamma distribution.

\$map\$qtn.shape1 the shape1 parameter for beta distribution.

\$map\$qtn.shape2 the shape2 parameter for beta distribution.

\$map\$qtn.ncp the ncp parameter for beta distribution.

\$map\$qtn.spot the QTN distribution probability in each block.

\$map\$len.block the block length.

\$map\$maf the maf threshold, markers less than this threshold will be exclude.

\$map\$recom.spot whether to generate recombination events.

\$map\$range.hot the recombination times range in the hot spot.

\$map\$range.cold the recombination times range in the cold spot.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))

# Run annotation simulation
SP <- annotation(SP)
```

bigt

Genotype transportor

Description

Transport genotype matrix.

Usage

```
bigt(pop.geno, ncpus = 0)
```

Arguments

pop.geno genotype matrix of (0, 1).
 ncpus the number of threads used, if NULL, (logical core number - 1) is automatically used.

Details

Build date: Jan 28, 2025 Last update: Jan 29, 2025

Value

genotype matrix of (0, 1, 2).

Author(s)

Dong Yin

Examples

```
library(bigmemory)
options(bigmemory.typecast.warning=FALSE)
pop.geno <- matrix(sample(c(0, 1), 16, replace = TRUE), 4, 4)
pop.geno[]
bigmat <- bigt(pop.geno)
bigmat[]
pop.geno <- as.big.matrix(pop.geno, type = 'char')
bigmat <- bigt(pop.geno)
bigmat[]
```

build.cov

Correlation building

Description

To build correlation of variables.

Usage

```
build.cov(df = NULL, mu = rep(0, nrow(Sigma)), Sigma = diag(2), tol = 1e-06)
```

Arguments

df a data frame needing building correlation.
 mu means of the variables.
 Sigma covariance matrix of variables.
 tol tolerance (relative to largest variance) for numerical lack of positive-definiteness in Sigma.

Details

Build date: Oct 10, 2019 Last update: Apr 28, 2022

Value

a data frame with expected correlation

Author(s)

Dong Yin and R

References

B. D. Ripley (1987) Stochastic Simulation. Wiley. Page 98

Examples

```
df <- data.frame(tr1 = rnorm(100), tr2 = rnorm(100))
df.cov <- build.cov(df)
var(df.cov)
```

cal.eff

QTN genetic effects

Description

Calculate for genetic effects vector of selected markers.

Usage

```
cal.eff(  
  qtn.num = 10,  
  qtn.dist = "norm",  
  qtn.var = 1,  
  qtn.prob = 0.5,  
  qtn.shape = 1,  
  qtn.scale = 1,  
  qtn.shape1 = 1,  
  qtn.shape2 = 1,  
  qtn.ncp = 0  
)
```

Arguments

qtn.num	integer: the QTN number of single trait; vector: the multiple group QTN number of single trait; matrix: the QTN number of multiple traits.
qtn.dist	the QTN distribution containing 'norm', 'geom', 'gamma' or 'beta'.
qtn.var	the standard deviations for normal distribution.
qtn.prob	the probability of success for geometric distribution.
qtn.shape	the shape parameter for gamma distribution.
qtn.scale	the scale parameter for gamma distribution.
qtn.shape1	the shape1 parameter for beta distribution.
qtn.shape2	the shape2 parameter for beta distribution.
qtn.ncp	the ncp parameter for beta distribution.

Details

Build date: Nov 14, 2018 Last update: Apr 28, 2022

Value

a vector of genetic effect.

Author(s)

Dong Yin

Examples

```
eff <- cal.eff(qtn.num = 10)
str(eff)
```

checkEnv *Environmental factor checking*

Description

Check the levels of environmental factors.

Usage

```
checkEnv(data, envName, verbose = TRUE)
```

Arguments

data	data needing check.
envName	the environmental factor name within the data.
verbose	whether to print detail.

Details

Build date: Sep 10, 2021 Last update: Apr 28, 2022

Value

data without environmental factors of wrong level.

Author(s)

Dong Yin

Examples

```
data <- data.frame(a = c(1, 1, 2), b = c(2, 2, 3), c = c(3, 3, 4))
envName <- c("a", "b", "c")
data <- checkEnv(data = data, envName = envName)
```

generate.map

Marker information

Description

Generate map data with marker information.

Usage

```
generate.map(  
  species = NULL,  
  pop.marker = NULL,  
  num.chr = 18,  
  len.chr = 1.5e+08  
)
```

Arguments

species	the species of genetic map, which can be "arabidopsis", "cattle", "chicken", "dog", "horse", "human", "maize", "mice", "pig", and "rice".
pop.marker	the number of markers.
num.chr	the number of chromosomes.
len.chr	the length of chromosomes.

Details

Build date: Mar 19, 2022 Last update: Apr 28, 2022

Value

a data frame with marker information.

Author(s)

Dong Yin

Examples

```
pop.map <- generate.map(pop.marker = 1e4)
str(pop.map)
```

<code>generate.pop</code>	<i>Population generator</i>
---------------------------	-----------------------------

Description

Generate population according to the number of individuals.

Usage

```
generate.pop(pop.ind = 100, from = 1, ratio = 0.5, gen = 1)
```

Arguments

<code>pop.ind</code>	the number of the individuals in a population.
<code>from</code>	initial index of the population.
<code>ratio</code>	sex ratio of males in a population.
<code>gen</code>	generation ID of the population.

Details

Build date: Nov 14, 2018 Last update: Apr 28, 2022

Value

a data frame of population information.

Author(s)

Dong Yin

Examples

```
pop <- generate.pop(pop.ind = 100)
head(pop)
```

`geno.cvt1`*Genotype code convertor 1*

Description

Convert genotype matrix from (0, 1) to (0, 1, 2).

Usage

```
geno.cvt1(pop.geno, ncpus = 0)
```

Arguments

<code>pop.geno</code>	genotype matrix of (0, 1).
<code>ncpus</code>	the number of threads used, if NULL, (logical core number - 1) is automatically used.

Details

Build date: Nov 14, 2018 Last update: Jan 30, 2025

Value

genotype matrix of (0, 1, 2).

Author(s)

Dong Yin

Examples

```
library(bigmemory)
options(bigmemory.typecast.warning=FALSE)
pop.geno <- matrix(sample(c(0, 1), 16, replace = TRUE), 4, 4)
pop.geno[]
bigmat <- geno.cvt1(pop.geno)
bigmat[]
pop.geno <- as.big.matrix(pop.geno, type = 'char')
bigmat <- geno.cvt1(pop.geno)
bigmat[]
```

`geno.cvt2`*Genotype code convertor 2*

Description

Convert genotype matrix from (0, 1, 2) to (0, 1).

Usage

```
geno.cvt2(pop.geno, ncpus = 0)
```

Arguments

<code>pop.geno</code>	genotype matrix of (0, 1, 2).
<code>ncpus</code>	the number of threads used, if NULL, (logical core number - 1) is automatically used.

Details

Build date: Jul 11, 2020 Last update: Jan 29, 2025

Value

genotype matrix of (0, 1).

Author(s)

Dong Yin

Examples

```
library(bigmemory)
options(bigmemory.typecast.warning=FALSE)
pop.geno <- matrix(sample(c(0, 1, 2), 8, replace = TRUE), 2, 4)
pop.geno[]
bigmat <- geno.cvt2(pop.geno)
bigmat[]
pop.geno <- as.big.matrix(pop.geno, type = 'char')
bigmat <- geno.cvt2(pop.geno)
bigmat[]
```

genotype

*Genotype simulation***Description**

Generating and editing genotype data.

Usage

```
genotype(SP = NULL, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Jan 28, 2025

Value

the function returns a list containing

\$geno\$pop.geno the genotype data.

\$geno\$inrows '1': one-row genotype represents an individual; '2': two-row genotype represents an individual.

\$geno\$pop.marker the number of markers.

\$geno\$pop.ind the number of individuals in the base population.

\$geno\$prob the genotype code probability.

\$geno\$rate.mut the mutation rate of the genotype data.

\$geno\$cld whether to generate a complete LD genotype data when 'inrows == 2'.

Author(s)

Dong Yin

Examples

```
# Generate genotype simulation parameters
SP <- param.geno(pop.marker = 1e4, pop.ind = 1e2)

# Run genotype simulation
SP <- genotype(SP)
```

getfam *Family index and within-family index*

Description

Get indice of family and within-family

Usage

```
getfam(sir, dam, fam.op, mode = c("pat", "mat", "pm"))
```

Arguments

sir	the indice of sires.
dam	the indice of dams.
fam.op	the initial index of family indice.
mode	"pat": paternal mode; "mat": maternal mode; "pm": paternal and maternal mode.

Details

Build date: Nov 14, 2018 Last update: Apr 30, 2022

Value

a matrix with family indice and within-family indice.

Author(s)

Dong Yin

Examples

```
s <- c(0, 0, 0, 0, 1, 3, 3, 1, 5, 7, 5, 7, 1, 3, 5, 7)
d <- c(0, 0, 0, 0, 2, 4, 4, 2, 6, 8, 8, 6, 6, 8, 4, 8)
fam <- getfam(sir = s, dam = d, fam.op = 1, mode = "pm")
fam
```

GxG.network

Genetic interaction network

Description

Generate genetic interaction effect combination network.

Usage

```
GxG.network(pop.map = NULL, qtn.pos = 1:10, qtn.model = "A:D")
```

Arguments

pop.map	the map data with annotation information.
qtn.pos	the index of QTNs in the map data.
qtn.model	the genetic model of QTN such as 'A:D'.

Details

Build date: Mar 19, 2022 Last update: Apr 28, 2022

Value

a data frame of genetic interaction effect.

Author(s)

Dong Yin

Examples

```
pop.map <- generate.map(pop.marker = 1e4)
GxG.net <- GxG.network(pop.map)
head(GxG.net)
```

IndPerGen

Individual number per generation

Description

Calculate the individual number per generation.

Usage

```
IndPerGen(  
  pop,  
  pop.gen = 2,  
  ps = c(0.8, 0.8),  
  reprod.way = "randmate",  
  sex.rate = 0.5,  
  prog = 2  
)
```

Arguments

pop	the population information containing environmental factors and other effects.
pop.gen	the generations of simulated population.
ps	if $ps \leq 1$, fraction selected in selection of males and females; if $ps > 1$, ps is number of selected males and females.
reprod.way	reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.
sex.rate	the sex ratio of simulated population.
prog	the progeny number of an individual.

Details

Build date: Apr 12, 2022 Last update: Apr 30, 2022

Value

the vector containing the individual number per generation.

Author(s)

Dong Yin

Examples

```
pop <- generate.pop(pop.ind = 100)  
count.ind <- IndPerGen(pop)
```

logging.initialize *Logging initialization*

Description

Initialize the logging process.

Usage

```
logging.initialize(module, outpath)
```

Arguments

module	the module name.
outpath	the path of output files, Simer writes files only if outpath is not 'NULL'.

Details

Build date: Jul 11, 2020 Last update: Apr 28, 2022

Value

none.

Author(s)

Dong Yin

logging.log *Logging*

Description

Print or write log.

Usage

```
logging.log(  
    ...,  
    file = NULL,  
    sep = " ",  
    fill = FALSE,  
    labels = NULL,  
    verbose = TRUE  
)
```


Arguments

...	R objects.
file	a connection or a character string naming the file to print to. If "" (the default), cat prints to the standard output connection, the console unless redirected by sink. If it is "lcmd", the output is piped to the command given by 'cmd', by opening a pipe connection.
sep	a character vector of strings to append after each element.
fill	a logical or (positive) numeric controlling how the output is broken into successive lines.
labels	a character vector of labels for the lines printed. Ignored if fill is FALSE.
verbose	whether to print detail.

Details

Build date: Jul 11, 2020 Last update: Apr 28, 2022

Value

none.

Author(s)

Dong Yin

Examples

```
logging.log('simer')
```

logging.print	<i>Logging printer</i>
---------------	------------------------

Description

Print R object information into file.

Usage

```
logging.print(x, file = NULL, append = TRUE, verbose = TRUE)
```

Arguments

x	a matrix or a list.
file	the filename of output file.
append	logical. If TRUE, output will be appended to file; otherwise, it will overwrite the contents of file.
verbose	whether to print details.

Details

Build date: Feb 7, 2020 Last update: Apr 28, 2022

Value

none.

Author(s)

Dong Yin

Examples

```
x <- list(a = "a", b = "b")
logging.print(x)
```

mate

Mate

Description

Mating according to the indice of sires and dams.

Usage

```
mate(pop.geno, index.sir, index.dam, ncpus = 0)
```

Arguments

pop.geno	the genotype data.
index.sir	the indice of sires.
index.dam	the indice of dams.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.

Details

Build date: Nov 14, 2018 Last update: Jan 28, 2025

Value

a genotype matrix after mating

Author(s)

Dong Yin

Examples

```
# Generate the genotype data
SP <- param.geno(pop.marker = 1e4, pop.ind = 1e2)
SP <- genotype(SP)
pop.geno <- SP$geno$pop.geno$gen1

# The mating design
index.sir <- rep(1:50, each = 2)
index.dam <- rep(51:100, each = 2)

# Mate according to mating design
geno.curr <- mate(pop.geno = pop.geno, index.sir = index.sir,
                 index.dam = index.dam)
geno.curr[1:5, 1:5]
```

mate.2waycro	<i>Two-way cross</i>
--------------	----------------------

Description

Produce individuals by two-way cross.

Usage

```
mate.2waycro(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Apr 30, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "2waycro")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Two different breeds are cut by sex
SP$pheno$pop$gen1$sex <- rep(c(1, 2), c(50, 50))
# Run selection
SP <- selects(SP)
# Run two-way cross
SP <- mate.2waycro(SP)
```

mate.3waycro

Three-way cross

Description

Produce individuals by three-way cross.

Usage

```
mate.3waycro(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Apr 11, 2022 Last update: Jan 28, 2025

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "3waycro")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Three different breeds are cut by sex
SP$pheno$pop$gen1$sex <- rep(c(1, 2, 1), c(30, 30, 40))
# Run selection
SP <- selects(SP)
```

```
# Run three-way cross
SP <- mate.3waycro(SP)
```

```
mate.4waycro
```

Four-way cross process

Description

Produce individuals by four-way cross.

Usage

```
mate.4waycro(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Apr 11, 2022 Last update: Jan 28, 2025

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```

# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "4waycro")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Four different breeds are cut by sex
SP$pheno$pop$gen1$sex <- rep(c(1, 2, 1, 2), c(25, 25, 25, 25))
# Run selection
SP <- selects(SP)
# Run four-way cross
SP <- mate.4waycro(SP)

```

mate.assort

Assortative mating

Description

Produce individuals by assortative mating.

Usage

```
mate.assort(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Sep 30, 2022 Last update: Sep 30, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "assort")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
# Run random mating
SP <- mate.assort(SP)
```

mate.backcro

Back cross

Description

Produce individuals by back cross.

Usage

```
mate.backcro(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Apr 12, 2022 Last update: Jan 28, 2025

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "backcro")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
```

```
# Two different breeds are cut by sex
SP$pheno$pop$gen1$sex <- rep(c(1, 2), c(50, 50))
# Run selection
SP <- selects(SP)
# Run back cross
SP <- mate.backcro(SP)
```

mate.clone

Clone

Description

Produce individuals by clone.

Usage

```
mate.clone(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Jan 28, 2025

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "clone")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
# Run clone
SP <- mate.clone(SP)
```

mate.dh

Doubled haploid

Description

Produce individuals by doubled haploid.

Usage

```
mate.dh(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Jan 28, 2025

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "dh")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
# Run doubled haploid
SP <- mate.dh(SP)
```

mate.disassort

Disassortative mating

Description

Produce individuals by disassortative mating.

Usage

```
mate.disassort(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Sep 30, 2022 Last update: Sep 30, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "disassort")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
```

```
# Run selection
SP <- selects(SP)
# Run random mating
SP <- mate.assort(SP)
```

mate.randexself *Random mating excluding self-pollination*

Description

Produce individuals by random mating excluding self-pollination.

Usage

```
mate.randexself(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Apr 30, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```

# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "randexself")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
# Run random mating excluding self-pollination
SP <- mate.randexself(SP)

```

mate.randmate	<i>Random mating</i>
---------------	----------------------

Description

Produce individuals by random-mating.

Usage

```
mate.randmate(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Apr 30, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "randmate")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
# Run random mating
SP <- mate.randmate(SP)
```

mate.selfpol

Self-pollination

Description

Produce individuals by self-pollination.

Usage

```
mate.selfpol(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Apr 30, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "selfpol")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
```

```
# Run selection
SP <- selects(SP)
# Run self-pollination
SP <- mate.selfpol(SP)
```

mate.userped	<i>User-specified pedigree mating</i>
--------------	---------------------------------------

Description

Produce individuals by user-specified pedigree mating.

Usage

```
mate.userped(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Apr 12, 2022 Last update: Apr 30, 2022

Value

the function returns a list containing

\$reprod\$pop.sel the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the sex ratio of simulated population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```

# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "userped")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run user-specified pedigree mating
SP <- mate.userped(SP)

```

param.annot	<i>Annotation parameters generator</i>
-------------	--

Description

Generate parameters for annotation data simulation.

Usage

```
param.annot(SP = NULL, ...)
```

Arguments

SP	a list of all simulation parameters.
...	one or more parameter(s) for map simulation.

Details

Build date: Feb 24, 2022 Last update: Jul 10, 2022

Value

the function returns a list containing

\$map\$pop.map the map data with annotation information.

\$map\$species the species of genetic map, which can be "arabidopsis", "cattle", "chicken", "dog", "horse", "human", "maize", "mice", "pig", and "rice".

\$map\$pop.marker the number of markers.

\$map\$num.chr the number of chromosomes.

\$map\$len.chr the length of chromosomes.

\$map\$qtn.model the genetic model of QTN such as 'A + D'.

\$map\$qtn.index the QTN index for each trait.

\$map\$qtn.num the QTN number for (each group in) each trait.

\$map\$qtn.dist the QTN distribution containing 'norm', 'geom', 'gamma' or 'beta'.

\$map\$qtn.var the standard deviations for normal distribution.

\$map\$qtn.prob the probability of success for geometric distribution.

\$map\$qtn.shape the shape parameter for gamma distribution.

\$map\$qtn.scale the scale parameter for gamma distribution.

\$map\$qtn.shape1 the shape1 parameter for beta distribution.

\$map\$qtn.shape2 the shape2 parameter for beta distribution.

\$map\$qtn.ncp the ncp parameter for beta distribution.

\$map\$qtn.spot the QTN distribution probability in each block.

\$map\$len.block the block length.

\$map\$maf the maf threshold, markers less than this threshold will be exclude.

\$map\$recom.spot whether to generate recombination events.

\$map\$range.hot the recombination times range in the hot spot.

\$map\$range.cold the recombination times range in the cold spot.

Author(s)

Dong Yin

Examples

```
SP <- param.annot(qtn.num = list(tr1 = 10))
str(SP)
```

param.geno

Genotype parameters generator

Description

Generate parameters for genotype data simulation.

Usage

```
param.geno(SP = NULL, ...)
```

Arguments

SP a list of all simulation parameters.
 ... one or more parameter(s) for genotype simulation.

Details

Build date: Feb 21, 2022 Last update: Jan 27, 2025

Value

the function returns a list containing

\$geno\$pop.geno the genotype data.

\$geno\$inrows '1': one-row genotype represents an individual; '2': two-row genotype represents an individual.

\$geno\$pop.marker the number of markers.

\$geno\$pop.ind the number of individuals in the base population.

\$geno\$prob the genotype code probability.

\$geno\$rate.mut the mutation rate of the genotype data.

\$geno\$cld whether to generate a complete LD genotype data when 'inrows == 2'.

Author(s)

Dong Yin

Examples

```
SP <- param.geno(pop.marker = 1e4, pop.ind = 1e2)
str(SP)
```

param.global

Global parameters generator

Description

Generate parameters for global options.

Usage

```
param.global(SP = NULL, ...)
```

Arguments

SP a list of all simulation parameters.
 ... one or more parameter(s) for global options.

Details

Build date: Apr 16, 2022 Last update: Jul 4, 2022

Value

the function returns a list containing

\$replication the replication times of simulation.

\$seed.sim simulation random seed.

\$out the prefix of output files.

\$outpath the path of output files, Simer writes files only if outpath is not 'NULL'.

\$out.format 'numeric' or 'plink', the data format of output files.

\$pop.gen the generations of simulated population.

\$out.geno.gen the output generations of genotype data.

\$out.pheno.gen the output generations of phenotype data.

\$useAllGeno whether to use all genotype data to simulate phenotype.

\$ncpus the number of threads used, if NULL, (logical core number - 1) is automatically used.

\$verbose whether to print detail.

Author(s)

Dong Yin

Examples

```
SP <- param.global(out = "simer")
str(SP)
```

param.pheno

Phenotype parameters generator

Description

Generate parameters for phenotype data simulation.

Usage

```
param.pheno(SP = NULL, ...)
```

Arguments

SP a list of all simulation parameters.
 ... one or more parameter(s) for phenotype simulation.

Details

Build date: Feb 21, 2022 Last update: Jul 4, 2022

Value

the function returns a list containing

\$pheno\$pop the population information containing environmental factors and other effects.

\$pheno\$pop.ind the number of individuals in the base population.

\$pheno\$pop.rep the repeated times of repeated records.

\$pheno\$pop.rep.bal whether repeated records are balanced.

\$pheno\$pop.env a list of environmental factors setting.

\$pheno\$phe.type a list of phenotype types.

\$pheno\$phe.model a list of genetic model of phenotype such as "T1 = A + E".

\$pheno\$phe.h2A a list of additive heritability.

\$pheno\$phe.h2D a list of dominant heritability.

\$pheno\$phe.h2GxG a list of GxG interaction heritability.

\$pheno\$phe.h2GxE a list of GxE interaction heritability.

\$pheno\$phe.h2PE a list of permanent environmental heritability.

\$pheno\$phe.var a list of phenotype variance.

\$pheno\$phe.corA the additive genetic correlation matrix.

\$pheno\$phe.corD the dominant genetic correlation matrix.

\$pheno\$phe.corGxG the GxG genetic correlation matrix.

\$pheno\$phe.corPE the permanent environmental correlation matrix.

\$pheno\$phe.corE the residual correlation matrix.

Author(s)

Dong Yin

Examples

```
SP <- param.pheno(phe.model = list(tr1 = "T1 = A + E"))
str(SP)
```

param.reprod

Reproduction parameters generator

Description

Generate parameters for reproduction.

Usage

```
param.reprod(SP = NULL, ...)
```

Arguments

SP	a list of all simulation parameters.
...	one or more parameter(s) for reproduction.

Details

Build date: Apr 6, 2022 Last update: Jul 4, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the male rate in the population.

\$reprod\$prog the progeny number of an individual.

Author(s)

Dong Yin

Examples

```
SP <- param.reprod(reprod.way = "randmate")
str(SP)
```

param.sel *Selection parameters generator*

Description

Generate parameters for selection.

Usage

```
param.sel(SP = NULL, ...)
```

Arguments

SP a list of all simulation parameters.
 ... one or more parameter(s) for selection.

Details

Build date: Apr 6, 2022 Last update: Jul 4, 2022

Value

the function returns a list containing

\$sel\$pop.sel the selected males and females.

\$sel\$ps if ps <= 1, fraction selected in selection of males and females; if ps > 1, ps is number of selected males and females.

\$sel\$decr whether the sort order is decreasing.

\$sel\$sel.crit the selection criteria, it can be 'TBV', 'TGV', and 'pheno'.

\$sel\$sel.single the single-trait selection method, it can be 'ind', 'fam', 'infam', and 'comb'.

\$sel\$sel.multi the multiple-trait selection method, it can be 'index', 'indcul', and 'tmd'.

\$sel\$index.wt the weight of each trait for multiple-trait selection.

\$sel\$index.tdm the index of tandem selection for multiple-trait selection.

\$sel\$goal.perc the percentage of goal more than the mean of scores of individuals.

\$sel\$pass.perc the percentage of expected excellent individuals.

Author(s)

Dong Yin

Examples

```
SP <- param.sel(sel.single = "ind")
str(SP)
```

param.simer

Parameter generator

Description

Generate parameters for Simer.

Usage

```
param.simer(SP = NULL, ...)
```

Arguments

SP	a list of all simulation parameters.
...	one or more parameter(s) for simer.

Details

Build date: Apr 17, 2022 Last update: Jul 4, 2022

Value

the function returns a list containing

\$global a list of global parameters.

\$map a list of marker information parameters.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

\$sel a list of selection parameters.

\$reprod a list of reproduction parameters.

Author(s)

Dong Yin

Examples

```
SP <- param.simer(out = "simer")  
str(SP)
```

phenotype *Phenotype simulation*

Description

Generate single-trait or multiple-trait phenotype by mixed model.

Usage

```
phenotype(SP = NULL, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Jan 28, 2025

Value

the function returns a list containing

\$pheno\$pop the population information containing environmental factors and other effects.

\$pheno\$pop.ind the number of individuals in the base population.

\$pheno\$pop.rep the repeated times of repeated records.

\$pheno\$pop.rep.bal whether repeated records are balanced.

\$pheno\$pop.env a list of environmental factors setting.

\$pheno\$phe.type a list of phenotype types.

\$pheno\$phe.model a list of genetic model of phenotype such as "T1 = A + E".

\$pheno\$phe.h2A a list of additive heritability.

\$pheno\$phe.h2D a list of dominant heritability.

\$pheno\$phe.h2GxG a list of GxG interaction heritability.

\$pheno\$phe.h2GxE a list of GxE interaction heritability.

\$pheno\$phe.h2PE a list of permanent environmental heritability.

\$pheno\$phe.var a list of phenotype variance.

\$pheno\$phe.corA the additive genetic correlation matrix.

\$pheno\$phe.corD the dominant genetic correlation matrix.

\$pheno\$phe.corGxG the GxG genetic correlation matrix.

\$pheno\$phe.corPE the permanent environmental correlation matrix.

\$pheno\$phe.corE the residual correlation matrix.

Author(s)

Dong Yin

ReferencesKao C and Zeng Z (2002) <<https://www.genetics.org/content/160/3/1243.long>>**Examples**

```

# Prepare environmental factor list
pop.env <- list(
  F1 = list( # fixed effect 1
    level = c("1", "2"),
    effect = list(tr1 = c(50, 30), tr2 = c(50, 30))
  ),
  F2 = list( # fixed effect 2
    level = c("d1", "d2", "d3"),
    effect = list(tr1 = c(10, 20, 30), tr2 = c(10, 20, 30))
  ),
  C1 = list( # covariate 1
    level = c(70, 80, 90),
    slope = list(tr1 = 1.5, tr2 = 1.5)
  ),
  R1 = list( # random effect 1
    level = c("l1", "l2", "l3"),
    ratio = list(tr1 = 0.1, tr2 = 0.1)
  )
)

# Generate genotype simulation parameters
SP <- param.annot(qtn.num = list(tr1 = c(2, 8), tr2 = 10),
  qtn.model = "A + D + A:D")
# Generate annotation simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(
  SP = SP,
  pop.ind = 100,
  pop.rep = 2, # 2 repeated record
  pop.rep.bal = TRUE, # balanced repeated record
  pop.env = pop.env,
  phe.type = list(
    tr1 = "continuous",
    tr2 = list(case = 0.01, control = 0.99)
  ),
  phe.model = list(
    tr1 = "T1 = A + D + A:D + F1 + F2 + C1 + R1 + A:F1 + E",
    tr2 = "T2 = A + D + A:D + F1 + F2 + C1 + R1 + A:F1 + E"
  ),
  phe.var = list(tr1 = 100, tr2 = 100)
)

```

```
# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
```

pop.geno	<i>Raw genotype matrix from outside in simdata</i>
----------	--

Description

Raw genotype matrix from outside in simdata

Usage

```
data(simdata)
```

Format

matrix

Examples

```
data(simdata)
dim(pop.geno)
head(pop.geno)
```

pop.map	<i>Map file from outside in simdata</i>
---------	---

Description

Map file from outside in simdata

Usage

```
data(simdata)
```

Format

list

Examples

```
data(simdata)
dim(pop.map)
head(pop.map)
```

remove_bigmatrix *Big.matrix removing*

Description

Remove big.matrix safely.

Usage

```
remove_bigmatrix(x, desc_suffix = ".geno.desc", bin_suffix = ".geno.bin")
```

Arguments

x the filename of big.matrix.
desc_suffix the suffix of description file of big.matrix.
bin_suffix the suffix of binary file of big.matrix.

Details

Build date: Aug 8, 2019 Last update: Apr 30, 2022

Value

TRUE or FALSE

Author(s)

Haohao Zhang and Dong Yin

Examples

```
library(bigmemory)
mat <- filebacked.big.matrix(
  nrow = 10,
  ncol = 10,
  init = 0,
  type = 'char',
  backingpath = ".",
  backingfile = 'simer.geno.bin',
  descriptorfile = 'simer.geno.desc')

remove_bigmatrix(x = "simer")
```

reproduces

*Reproduction***Description**

Population reproduction by different mate design.

Usage

```
reproduces(SP, ncpus = 0, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Nov 14, 2018 Last update: Apr 29, 2022

Value

the function returns a list containing

\$reprod\$pop.gen the generations of simulated population.

\$reprod\$reprod.way reproduction method, it consists of 'clone', 'dh', 'selfpol', 'randmate', 'randexself', 'assort', 'disassort', '2waycro', '3waycro', '4waycro', 'backcro', and 'userped'.

\$reprod\$sex.rate the male rate in the population.

\$reprod\$prog the progeny number of an individual.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
```

```

SP <- param.sel(SP = SP, sel.single = "ind")
# Generate reproduction parameters
SP <- param.reprod(SP = SP, reprod.way = "randmate")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
# Run reproduction
SP <- reproduces(SP)

```

selects

Selection

Description

Select individuals by combination of selection method and criterion.

Usage

```
selects(SP = NULL, verbose = TRUE)
```

Arguments

SP	a list of all simulation parameters.
verbose	whether to print detail.

Details

Build date: Sep 8, 2018 Last update: Apr 30, 2022

Value

the function returns a list containing

\$sel\$pop.sel the selected males and females.

\$sel\$ps if $ps \leq 1$, fraction selected in selection of males and females; if $ps > 1$, ps is number of selected males and females.

\$sel\$decr whether the sort order is decreasing.

\$sel\$sel.crit the selection criteria, it can be 'TBV', 'TGV', and 'pheno'.

\$sel\$sel.single the single-trait selection method, it can be 'ind', 'fam', 'infam', and 'comb'.

\$sel\$sel.multi the multiple-trait selection method, it can be 'index', 'indcul', and 'tmd'.

\$sel\$index.wt the weight of each trait for multiple-trait selection.

\$sel\$index.tdm the index of tandem selection for multiple-trait selection.

\$sel\$goal.perc the percentage of goal more than the mean of scores of individuals.

\$sel\$pass.perc the percentage of expected excellent individuals.

Author(s)

Dong Yin

Examples

```
# Generate annotation simulation parameters
SP <- param.annot(qtn.num = list(tr1 = 10))
# Generate genotype simulation parameters
SP <- param.geno(SP = SP, pop.marker = 1e4, pop.ind = 1e2)
# Generate phenotype simulation parameters
SP <- param.pheno(SP = SP, pop.ind = 100)
# Generate selection parameters
SP <- param.sel(SP = SP, sel.single = "ind")

# Run annotation simulation
SP <- annotation(SP)
# Run genotype simulation
SP <- genotype(SP)
# Run phenotype simulation
SP <- phenotype(SP)
# Run selection
SP <- selects(SP)
```

simer

Simer

Description

Main function of Simer.

Usage

```
simer(SP)
```

Arguments

SP a list of all simulation parameters.

Details

Build date: Jan 7, 2019 Last update: Apr 29, 2022

Value

the function returns a list containing

\$global a list of global parameters.

\$map a list of marker information parameters.

\$geno a list of genotype simulation parameters.

\$pheno a list of phenotype simulation parameters.

\$sel a list of selection parameters.

\$reprod a list of reproduction parameters.

Author(s)

Dong Yin, Lilin Yin, Haohao Zhang, and Xiaolei Liu

Examples

```
# Generate all simulation parameters
SP <- param.simer(out = "simer")

# Run Simer
SP <- simer(SP)
```

simer.Data

Data handling

Description

Make data quality control for genotype, phenotype, and pedigree.

Usage

```
simer.Data(jsonList = NULL, out = "simer.qc", ncpus = 0, verbose = TRUE)
```

Arguments

jsonList	a list of data quality control parameters.
out	the prefix of output files.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: May 26, 2021 Last update: Apr 28, 2022

Value

the function returns a list containing

\$genotype the path of genotype data.

\$pedigree the filename of pedigree data.

\$selection_index the selection index for all traits.

\$breeding_value_index the breeding value index for all traits.

\$quality_control_plan a list of parameters for data quality control.

\$breeding_plan a list of parameters for genetic evaluation.

Author(s)

Dong Yin

Examples

```
# Read JSON file
jsonFile <- system.file("extdata", "04breeding_plan", "plan1.json", package = "simer")
jsonList <- jsonlite::fromJSON(txt = jsonFile, simplifyVector = FALSE)

## Not run:
# It needs 'plink' and 'hiblup' software
jsonList <- simer.Data(jsonList = jsonList)

## End(Not run)
```

`simer.Data.Bfile2MVP` *simer.Data.Bfile2MVP: To transform plink binary data to MVP package*

Description

transforming plink binary data to MVP package.

Usage

```
simer.Data.Bfile2MVP(  
  bfile,  
  out = "simer",  
  maxLine = 10000,  
  type.geno = "char",  
  threads = 10,  
  verbose = TRUE  
)
```

Arguments

bfile	Genotype in binary format (.bed, .bim, .fam).
out	the name of output file.
maxLine	the max number of line to write to big matrix for each loop.
type.geno	the type of genotype elements.
threads	number of thread for transforming.
verbose	whether to print the reminder.

Details

Build date: Sep 12, 2018 Last update: Dec 28, 2024

Value

number of individuals and markers. Output files: genotype.desc, genotype.bin: genotype file in bigmemory format phenotype.phe: ordered phenotype file, same taxa order with genotype file map.map: SNP information

Author(s)

Haohao Zhang and Dong Yin

Examples

```
# Get bfile path
bfilePath <- file.path(system.file("extdata", "02plinkb", package = "simer"), "demo")

# Data converting
simer.Data.Bfile2MVP(bfilePath, tempfile("outfile"))
```

simer.Data.cHIBLUP *Genetic evaluation*

Description

The function of calling HIBLUP software of C version.

Usage

```
simer.Data.cHIBLUP(
  jsonList = NULL,
  hiblupPath = "",
  mode = "A",
  vc.method = "AI",
  ncpus = 10,
  verbose = TRUE
)
```

Arguments

jsonList	the list of genetic evaluation parameters.
hiblupPath	the path of HIBLUP software.
mode	'A' or 'AD', Additive effect model or Additive and Dominance model.
vc.method	default is 'AI', the method of calculating variance components in HIBLUP software.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: June 28, 2021 Last update: Apr 28, 2022

Value

the function returns a list containing

\$randList a list of estimated random effects.

\$varList a list of variance components.

\$covA the genetic covariance matrix for all traits.

\$corA the genetic correlation matrix for all traits.

Author(s)

Dong Yin

Examples

```
# Read JSON file
jsonFile <- system.file("extdata", "04breeding_plan", "plan1.json", package = "simer")
jsonList <- jsonlite::fromJSON(txt = jsonFile, simplifyVector = FALSE)

## Not run:
# It needs 'hiblup' software
gebvs <- simer.Data.cHIBLUP(jsonList = jsonList)

## End(Not run)
```

simer.Data.Env *Environmental factor selection*

Description

To find appropriate fixed effects, covariates, and random effects.

Usage

```
simer.Data.Env(
  jsonList = NULL,
  hiblupPath = "",
  header = TRUE,
  sep = "\t",
  ncpus = 10,
  verbose = TRUE
)
```

Arguments

jsonList	the list of environmental factor selection parameters.
hiblupPath	the path of HIBLUP software.
header	the header of file.
sep	the separator of file.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: July 17, 2021 Last update: Apr 28, 2022

Value

the function returns a list containing

\$genotype the path of genotype data.

\$pedigree the filename of pedigree data.

\$selection_index the selection index for all traits.

\$breeding_value_index the breeding value index for all traits.

\$quality_control_plan a list of parameters for data quality control.

\$breeding_plan a list of parameters for genetic evaluation.

Author(s)

Dong Yin

Examples

```
# Read JSON file
jsonFile <- system.file("extdata", "04breeding_plan", "plan1.json", package = "simer")
jsonList <- jsonlite::fromJSON(txt = jsonFile, simplifyVector = FALSE)

## Not run:
# It needs 'hiblup' software
jsonList <- simer.Data.Env(jsonList = jsonList)

## End(Not run)
```

simer.Data.Geno	<i>Genotype data quality control</i>
-----------------	--------------------------------------

Description

Data quality control for genotype data in MVP format and PLINK format.

Usage

```
simer.Data.Geno(  
  fileMVP = NULL,  
  fileBed = NULL,  
  filePlinkPed = NULL,  
  filePed = NULL,  
  filePhe = NULL,  
  out = "simer.qc",  
  genoType = "char",  
  filter = NULL,  
  filterGeno = NULL,  
  filterHWE = NULL,  
  filterMind = NULL,  
  filterMAF = NULL,  
  ncpus = 0,  
  verbose = TRUE  
)
```

Arguments

fileMVP	genotype in MVP format.
fileBed	genotype in PLINK binary format.
filePlinkPed	genotype in PLINK numeric format.
filePed	the filename of pedigree data.
filePhe	the filename of phenotype data, it can be a vector.
out	the prefix of output files.

genoType	type parameter in bigmemory, genotype data. The default is char, it is highly recommended *NOT* to modify this parameter.
filter	filter of genotyped individual.
filterGeno	threshold of sample miss rate.
filterHWE	threshold of Hardy-Weinberg Test.
filterMind	threshold of variant miss rate.
filterMAF	threshold of Minor Allele Frequency.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: May 26, 2021 Last update: Apr 28, 2022

Value

the function returns files

<out>.bed the .bed file of PLINK binary format.

<out>.bim the .bim file of PLINK binary format.

<out>.fam the .fam file of PLINK binary format.

Author(s)

Dong Yin

Examples

```
# Get the prefix of genotype data
fileBed <- system.file("extdata", "02plinkb", "demo", package = "simer")

## Not run:
# It needs 'plink' software
simer.Data.Geno(fileBed=fileBed)

## End(Not run)
```

simer.Data.Impute *Genotype data imputation*

Description

Impute the missing value within genotype data.

Usage

```
simer.Data.Impute(  
  fileMVP = NULL,  
  fileBed = NULL,  
  out = NULL,  
  maxLine = 10000,  
  ncpus = 0,  
  verbose = TRUE  
)
```

Arguments

fileMVP	genotype in MVP format.
fileBed	genotype in PLINK binary format.
out	the name of output file.
maxLine	number of SNPs, only used for saving memory when calculate kinship matrix.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: May 26, 2021 Last update: Apr 28, 2022

Value

the function returns files

<out>.geno.desc the description file of genotype data.

<out>.geno.bin the binary file of genotype data.

<out>.geno.ind the genotyped individual file.

<out>.geno.map the marker information data file.

Author(s)

Dong Yin

Examples

```
# Get the prefix of genotype data
fileMVP <- system.file("extdata", "02plinkb", "demo", package = "simer")

## Not run:
# It needs 'beagle' software
fileMVPimp <- simer.Data.Impute(fileBed = fileBed)

## End(Not run)
```

simer.Data.Json *Data quality control*

Description

Make data quality control by JSON file.

Usage

```
simer.Data.Json(
  jsonFile,
  hiblupPath = "",
  out = "simer.qc",
  dataQC = TRUE,
  buildModel = TRUE,
  buildIndex = TRUE,
  ncpus = 10,
  verbose = TRUE
)
```

Arguments

jsonFile	the path of JSON file.
hiblupPath	the path of HIBLUP software.
out	the prefix of output files.
dataQC	whether to make data quality control.
buildModel	whether to build EBV model.
buildIndex	whether to build Selection Index.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Oct 19, 2020 Last update: Apr 28, 2022

Value

the function returns a list containing

\$genotype the path of genotype data.

\$pedigree the filename of pedigree data.

\$selection_index the selection index for all traits.

\$breeding_value_index the breeding value index for all traits.

\$quality_control_plan a list of parameters for data quality control.

\$breeding_plan a list of parameters for genetic evaluation.

Author(s)

Dong Yin

Examples

```
# Get JSON file
jsonFile <- system.file("extdata", "04breeding_plan", "plan1.json", package = "simer")

## Not run:
# It needs 'plink' and 'hiblup' software
jsonList <- simer.Data.Json(jsonFile = jsonFile)

## End(Not run)
```

simer.Data.Map

simer.Data.Map: To check map file

Description

checking map file.

Usage

```
simer.Data.Map(
  map,
  out = "simer",
  cols = 1:5,
  header = TRUE,
  sep = "\t",
  verbose = TRUE
)
```

Arguments

map	the name of map file or map object(data.frame or matrix).
out	the name of output file.
cols	selected columns.
header	whether the file contains header.
sep	separator of the file.
verbose	whether to print detail.

Details

Build date: Sep 12, 2018 Last update: July 25, 2022

Value

Output file: <out>.map

Author(s)

Haohao Zhang and Dong Yin

Examples

```
# Get map path
mapPath <- system.file("extdata", "01bigmemory", "demo.geno.map", package = "simer")

# Check map data
simer.Data.Map(mapPath, tempfile("outfile"))
```

simer.Data.MVP2Bfile *simer.Data.MVP2Bfile: To transform MVP data to binary format*

Description

transforming MVP data to binary format.

Usage

```
simer.Data.MVP2Bfile(
  bigmat,
  map,
  pheno = NULL,
  out = "simer",
  threads = 1,
  verbose = TRUE
)
```

Arguments

bigmat	Genotype in bigmatrix format (0,1,2).
map	the map file.
pheno	the phenotype file.
out	the name of output file.
threads	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print the reminder.

Details

Build date: Sep 12, 2018 Last update: Jan 29, 2025

Value

NULL Output files: .bed, .bim, .fam

Author(s)

Haohao Zhang and Dong Yin

Examples

```
library(bigmemory)

# Generate bigmat and map
bigmat <- as.big.matrix(matrix(1:6, 3, 2))
map <- generate.map(pop.marker = 3)

# Data converting
simer.Data.MVP2Bfile(bigmat, map, out=tempfile("outfile"))
```

simer.Data.MVP2MVP *Genotype data conversion*

Description

Convert genotype data from MVP format to MVP format.

Usage

```
simer.Data.MVP2MVP(fileMVP, genoType = "char", out = "simer", verbose = TRUE)
```

Arguments

fileMVP	the prefix of MVP file.
genoType	type parameter in bigmemory data. The default is 'char', it is highly recommended *NOT* to modify this parameter.
out	the prefix of output files.
verbose	whether to print detail.

Details

Build date: May 26, 2021 Last update: Apr 28, 2022

Value

the function returns files

<out>.geno.desc the description file of genotype data.

<out>.geno.bin the binary file of genotype data.

<out>.geno.ind the genotyped individual file.

<out>.geno.map the marker information data file.

Author(s)

Dong Yin

Examples

```
# Get the prefix of genotype data
fileMVP <- system.file("extdata", "01bigmemory", "demo", package = "simer")

# Convert genotype data from MVP to MVP
simer.Data.MVP2MVP(fileMVP, out = tempfile("outfile"))
```

simer.Data.Ped

Pedigree data quality control

Description

Data quality control for pedigree data.

Usage

```

simer.Data.Ped(
  filePed,
  fileMVP = NULL,
  out = NULL,
  standardID = FALSE,
  fileSir = NULL,
  fileDam = NULL,
  exclThres = 0.1,
  assignThres = 0.05,
  header = TRUE,
  sep = "\t",
  ncpus = 0,
  verbose = TRUE
)

```

Arguments

filePed	the filename of pedigree need correcting.
fileMVP	genotype in MVP format.
out	the prefix of output file.
standardID	whether kid id is 15-character standard.
fileSir	the filename of candidate sires.
fileDam	the filename of candidate dams.
exclThres	if conflict ratio is more than exclThres, exclude this parent.
assignThres	if conflict ratio is less than assignThres, assign this parent to the individual.
header	whether the file contains header.
sep	separator of the file.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: May 6, 2021 Last update: Apr 28, 2022

Value

the function returns files

<out>.ped.report the report file containing correction condition.

<out>.ped.error the file containing pedigree error.

<out>.ped the pedigree file after correction.

Author(s)

Lilin Yin and Dong Yin

Examples

```
# Get the filename of pedigree data
filePed <- system.file("extdata", "05others", "pedigree.txt", package = "simer")

# Get the prefix of genotype data
fileMVP <- system.file("extdata", "01bigmemory", "demo", package = "simer")

# Run pedigree correction
simer.Data.Ped(filePed = filePed, fileMVP = fileMVP, out = tempfile("outfile"))
```

simer.Data.Pheno *Phenotype data quality control*

Description

Data quality control for phenotype data.

Usage

```
simer.Data.Pheno(
  filePhe = NULL,
  filePed = NULL,
  out = NULL,
  planPhe = NULL,
  pheCols = NULL,
  header = TRUE,
  sep = "\t",
  missing = c(NA, "NA", "Na", ".", "-", "NAN", "nan", "na", "N/A", "n/a", "<NA>", "",
    "-9", 9999),
  verbose = TRUE
)
```

Arguments

filePhe	the phenotype files, it can be a vector.
filePed	the pedigree files, it can be a vector.
out	the prefix of output file.
planPhe	the plans for phenotype quality control.
pheCols	the column needing extracting.
header	the header of file.
sep	the separator of file.
missing	the missing value.
verbose	whether to print detail.

Details

Build date: June 13, 2021 Last update: Apr 28, 2022

Value

the function returns files

<out>.phe the phenotype file after correction.

Author(s)

Haohao Zhang and Dong Yin

Examples

```
# Get the filename of phenotype data
filePhe <- system.file("extdata", "05others", "phenotype.txt", package = "simer")

# Run phenotype correction
simer.Data.Pheno(filePhe = filePhe, out = tempfile("outfile"))
```

simer.Data.SELIND *Selection index construction*

Description

The function of General Selection Index.

Usage

```
simer.Data.SELIND(jsonList = NULL, hiblupPath = "", ncpus = 10, verbose = TRUE)
```

Arguments

jsonList	the list of selection index construction parameters.
hiblupPath	the path of HIBLUP software.
ncpus	the number of threads used, if NULL, (logical core number - 1) is automatically used.
verbose	whether to print detail.

Details

Build date: Aug 26, 2021 Last update: Apr 28, 2022

Value

the function returns a list containing

\$genotype the path of genotype data.

\$pedigree the filename of pedigree data.

\$selection_index the selection index for all traits.

\$breeding_value_index the breeding value index for all traits.

\$quality_control_plan a list of parameters for data quality control.

\$breeding_plan a list of parameters for genetic evaluation.

Author(s)

Dong Yin

References

Y. S. Chen, Z. L. Sheng (1988) The Theory of General Selection Index. Genetic Report, 15(3): P185-P190

Examples

```
# Read JSON file
jsonFile <- system.file("extdata", "04breeding_plan", "plan1.json", package = "simer")
jsonList <- jsonlite::fromJSON(txt = jsonFile, simplifyVector = FALSE)

## Not run:
# It needs 'hiblup' software
jsonList <- simer.Data.SELIND(jsonList = jsonList)

## End(Not run)
```

simer.Version

Simer version

Description

Print simer version.

Usage

```
simer.Version(width = 60, verbose = TRUE)
```

Arguments

width the width of the message.

verbose whether to print detail.

Details

Build date: Aug 30, 2017 Last update: Apr 30, 2022

Value

version number.

Author(s)

Dong Yin, Lilin Yin, Haohao Zhang, and Xiaolei Liu

Examples

```
simer.Version()
```

write.file

File writing

Description

Write files of Simer.

Usage

```
write.file(SP)
```

Arguments

SP a list of all simulation parameters.

Details

Build date: Jan 7, 2019 Last update: Jan 28, 2025

Value

none.

Author(s)

Dong Yin

Examples

```
outpath <- tempdir()
SP <- param.simer(out = "simer")
SP <- simer(SP)
SP$global$outpath <- outpath
write.file(SP)
unlink(file.path(outpath, "180_Simer_Data_numeric"), recursive = TRUE)
```

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