Package 'Ease'

July 21, 2025

Type Package

Title Simulating Explicit Population Genetics Models

Version 0.1.2

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Description Implementation in a simple and efficient way of fully customisable population genet-

ics simulations,

considering multiple loci that have epistatic interactions. Specifically suited to the modelling of multilocus nucleocytoplasmic systems (with both diploid and haploid loci), it is nevertheless possible

to simulate purely diploid (or purely haploid) genetic models.

Examples of models that can be simulated with Ease are numerous, for example models of genetic

incompatibilities as presented by Marie-

Orleach et al. (2022) <doi:10.1101/2022.07.25.501356>.

Many others are conceivable, although few are actually explored, Ease having been developed in particular to provide a solution so that these kinds of models can be simulated simply.

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Encoding UTF-8

Imports methods, Rcpp (>= 1.0.7), RcppProgress (>= 0.1), stats

LinkingTo Rcpp, RcppProgress

RoxygenNote 7.2.1

Collate RcppExports.R ToolFunctions.R ModelFunctions.R GenomeClass.R MutationMatrixClass.R SelectionClass.R PopulationClass.R MetapopulationClass.R UserFunctions.R

Suggests rmarkdown, knitr, testthat (>= 3.0.0)

VignetteBuilder knitr

Config/testthat/edition 3

NeedsCompilation yes

Repository CRAN

Date/Publication 2022-11-07 14:40:08 UTC

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alleleFreqMatGeneration

Generation of the matrix for calculating allelic frequencies

Description

Generates a matrix that allows to go from genotypic frequencies to allelic frequencies.

Usage

alleleFreqMatGeneration(genomeObj)

Arguments

genomeObj a Genome object

Details

An allele frequency matrix is a matrix with rows equal to the number of genotypes and columns equal to the number of alleles. By multiplying a row matrix of genotype frequencies we obtain a row matrix of associated allele frequencies.

Value

A matrix for calculating allelic frequencies from genotypes frequencies.

Author(s)

Ehouarn Le Faou

areThereHomoz Are there any allelic combinations including homozygosity

Description

Test if there are homozygotes in the specified allelic combinations of a selection formula

Usage

areThereHomoz(formula)

Arguments

formula a selection formula

Value

logical indicating if there are homozygotes

Author(s)

Ehouarn Le Faou

catn

Concatenate, print and line break

Description

Object output in the same way as the function cat but adding a line break at the end.

Usage

```
catn(..., file = "", sep = " ", fill = FALSE, labels = NULL, append = FALSE)
```

Arguments

| • • • | see cat. |
|--------|----------|
| file | see cat. |
| sep | see cat. |
| fill | see cat. |
| labels | see cat. |
| append | see cat. |

check.genome

Details

See cat.

Value

None (invisible NULL).

Author(s)

Ehouarn Le Faou

check.genome

The validity check for the Genome class

Description

The validity check for the Genome class

Usage

check.genome(object)

Arguments

object a Genome object

Value

A logical corresponding to whether the object is a correct Genome object.

Author(s)

check.metapopulation The validity check for the Metapopulation class

Description

The validity check for the Metapopulation class

Usage

check.metapopulation(object)

Arguments

object a Metapopulation object

Value

a boolean corresponding to whether the object is a correct Metapopulation object.

Author(s)

Ehouarn Le Faou

check.mutationMatrix The validity check associated with the MutationMatrix class

Description

The validity check associated with the MutationMatrix class

Usage

```
check.mutationMatrix(object)
```

Arguments

object an object of class MutationMatrix

Value

A logical corresponding to whether x is a correct MutationMatrix object.

Author(s)

check.population The validity check for the Population class

Description

The validity check for the Population class

Usage

```
check.population(object)
```

Arguments

object a Population object

Value

a boolean corresponding to whether the object is a correct Population object.

Author(s)

Ehouarn Le Faou

check.selection The validity check for the Selection class

Description

The validity check for the Selection class

Usage

```
check.selection(object)
```

Arguments

object Selection object

Value

A logical corresponding to whether the object is a correct Selection object.

Author(s)

extractAlleleComb Extract the allele combination

Description

Conversion of an allelic combination defined in a selection formula into the vector listing the alleles present (alleles that must be in the homozygous state appear 2 times, 1 time for heterozygous).

Usage

```
extractAlleleComb(xVect)
```

Arguments

xVect allelic combination extracted from a selection formula.

Value

the list of alleles that must be present in the genotype to match the input allelic combination

Author(s)

Ehouarn Le Faou

Genome-class Genome class

Description

The Genome class allows to define all the characteristics of the genome which will be used as a basis for the construction of transition matrices from one generation to another in simulations of the model.

Details

A genome includes the list of all possible haplotypes and genotypes resulting from the combination of the alleles defined in input. As the Ease package was originally built for population genetics simulations including both diploid and haploid loci, it is necessary that both types of loci are defined. Despite this, the user can define only diploid or only haploid loci if they wish. If no diploid locus is defined, one is automatically generated with only one allele, thus not influencing the simulation. The same applies if no haploid locus is defined.

Each locus is described by a vector of factors which are the names of the possible alleles at that locus. All diploid (resp. haploid) loci thus defined are grouped in a list, called listDipLoci (resp. listHapLoci). Therefore, a Genome class object has two lists of loci defined in this way, one for diploid loci, one for haploid loci. The alleles and loci (diploid and haploid) must all have different names so that no ambiguity can persist.

genotyping

If several are defined, the order of the diploid loci in the list is not trivial. The rates of two-toone combinations between them must indeed be defined by the vector recRate. For example, if three diploid loci are defined, recRate must be of length 2, the first of its values defining the recombination rate between the first and second loci, the second of its values the recombination rate between the second and third loci. For example, if we want to define two groups of two loci that are linked to each other but are on two different chromosomes, we can define a recRate = c(0.1, 0.5, 0.1). The first two loci are thus relatively linked (recombination rate of 0.1), as are the last two loci. On the other hand, the recombination rate of 0.5 between the second and third loci ensures that the two groups are independent.

Slots

listHapLoci a list of haploid loci

listDipLoci a list of diploid loci

recRate a two-by-two recombination rate vector

nbHL the number of haploid loci

nbDL the number of diploid loci

listLoci the list of all loci

haplotypesHL haplotypes of haploid loci only

haplotypesDL haplotypes of diploid loci only

haplotypes haplotypes of all loci

alleles the vector of all the alleles

nbAlleles the number of alleles

nbHaplo the number of haplotypes

IDhaplotypes IDs of haplotypes

genotypes the list of genotypes

nbGeno the number of genotypes

IDgenotypes IDs of genotypes

IDgenome ID of the genome

Author(s)

Ehouarn Le Faou

genotyping

Genotyping

Description

Generation of genotypes associated with a Genome object.

Usage

genotyping(genomeObj)

Arguments

genomeObj a Genome object.

Details

The output genotypes are described as a list of three matrices. A genotype consists of two diploid haplotypes (first two matrices) and one haploid haplotype (third matrix), which are read at the same row number on all three matrices.

Value

A list of matrices describing genotypes in rows.

Author(s)

Ehouarn Le Faou

getCustomOutput Getting the custom output

Description

Getting the custom output

Usage

```
getCustomOutput(metapop)
```

Arguments

metapop a Metapopulation objects

Value

The list generated through the custom result function, if at least it was specified during the simulation of the Metapopulation.

Author(s)

Ehouarn Le Faou

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getRecords

Description

Getting the simulation results

Usage

getRecords(metapop)

Arguments

metapop a Metapopulation objects

Value

A list where each item is associated with a simulation. Each of these elements consists of a list of data.frames, one per population. These data.frames consist of the same columns as the results (see getResults documentation), except that they do not include the stop conditions.

Author(s)

Ehouarn Le Faou

getResults Getting the simulation results

Description

Getting the simulation results

Usage

getResults(metapop)

Arguments

metapop a Metapopulation objects

Value

A data.frame where each line corresponds to a simulation. The results include : - the last generation reached (the threshold or the generation that first verified at least one of the stopping conditions) - the final population size - the genotype frequencies - allelic frequencies - the reason(s) for the stop (either the threshold was reached, i.e. unstopped or the stop condition(s) that was (were) reached, in the form of boolean values - Average fitness (individual, gamete production and gametic)

Author(s)

Ehouarn Le Faou

haploCrossMatrix Haplotype crossing matrix generation

Description

Generation of the haplotype crossing matrix associated to a Genome object.

Usage

haploCrossMatrix(genomeObj)

Arguments

genomeObj a Genome object

Details

A crossover matrix is a square matrix of size equal to the number of haplotypes. It describes for each combination of two gametic haplotypes the genotype index resulting from their syngamy. In the general case it is not a symmetrical matrix (it is if a single haploid locus with a single allele is defined), because the transmission of haploid loci is only maternal, therefore non-symmetrical as is the transmission of diploid loci. It is therefore necessary to enter the haplotype frequencies of male gametes in the columns and the haplotype frequencies of female gametes in the rows during the calculations (this is done in the simulations).

Value

An haplotype crossing matrix.

Author(s)

haplotyping

Description

Generation of haplotypes associated with a Genome object.

Usage

```
haplotyping(genomeObj)
```

Arguments

genomeObj a Genome object

Details

The generated haplotypes are output as a list of three enumeration in the form of matrices of alleles (each row corresponding to an haplotype, each column to a locus). The first enumeration corresponds to haplotypes considering only haploid loci, the second only diploid loci and the third all loci (with two matrices, 1 for haploid loci, 1 for diploid loci).

Value

A list of matrices describing haplotypes in rows.

Author(s)

Ehouarn Le Faou

IDgenomeGeneration Genome identifier

Description

Generation of the input genome ID, i.e. the concatenation in string form of the names of the loci and alleles constituting this genome.

Usage

IDgenomeGeneration(listLoci, alleles)

Arguments

| listLoci | the list of all loci |
|----------|-------------------------------|
| alleles | the vector of all the alleles |

Value

The genome ID as a character string.

Author(s)

Ehouarn Le Faou

IDgenotypeGeneration Genotype identifier

Description

Generation of the input genotype ID, i.e. the concatenation in string form of the names of the alleles constituting these haplotypes (the two from the diploid genome and the one from the haploid genome).

Usage

```
IDgenotypeGeneration(dl1, dl2, hl = NULL)
```

Arguments

| dl1 | the first diploid haplotype as a character (or factors) vector. |
|-----|------------------------------------------------------------------|
| dl2 | the second diploid haplotype as a character (or factors) vector. |
| hl | the haploid haplotype as a character (or factors) vector. |

Value

The genotype ID as a character string.

Author(s)

IDhaplotypeGeneration Haplotype identifier

Description

Generation of the input haplotype ID, i.e. the concatenation in string form of the names of the alleles constituting this haplotype.

Usage

```
IDhaplotypeGeneration(dl, hl)
```

Arguments

| dl | the diploid haplotype as a character (or factors) vector. |
|----|-----------------------------------------------------------|
| hl | the haploid haplotype as a character (or factors) vector. |

Value

The haplotype ID as a character string.

Author(s)

Ehouarn Le Faou

initialize, Genome-method

Initialize method for the Genome class

Description

Initialize method for the Genome class

Usage

```
## S4 method for signature 'Genome'
initialize(.Object, listHapLoci, listDipLoci, recRate)
```

Arguments

| .Object | a Genome object |
|-------------|----------------------------------------|
| listHapLoci | a list of haploid loci |
| listDipLoci | a list of diploid loci |
| recRate | a two-by-two recombination rate vector |

Value

A Genome object

Author(s)

Ehouarn Le Faou

Description

Initialize method for the Metapopulation class

Usage

```
## S4 method for signature 'Metapopulation'
initialize(.Object, populations, migMat)
```

Arguments

| .Object | a Metapopulation object |
|-------------|--------------------------------|
| populations | list of Population $object(s)$ |
| migMat | migration matrix |

Value

a Metapopulation object

Author(s)

Description

Initialize method for the MutationMatrix class

Usage

```
## S4 method for signature 'MutationMatrix'
initialize(.Object, genomeObj, mutHapLoci, mutDipLoci)
```

Arguments

| .Object | a MutationMatrix object |
|------------|-------------------------------------------------------------|
| genomeObj | a Genome object |
| mutHapLoci | a list of haploid locus by locus allelic mulation matrices. |
| mutDipLoci | a list of diploid locus by locus allelic mulation matrices. |

Value

A MutationMatrix object

Author(s)

Ehouarn Le Faou

initialize, Population-method

Initialize method for the Population *class*

Description

Initialize method for the Population class

Usage

```
## S4 method for signature 'Population'
initialize(
   .Object,
   name,
   size,
   dioecy,
   selfRate,
```

```
demography,
growthRate,
initGenoFreq,
genomeObj,
initPopSize,
selectionObj,
mutMatrixObj
```

)

Arguments

| a Population object |
|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| the name of the population. |
| the size of the population. |
| logical indicating whether the population is dioecious or not (hermaphrodite). |
| the selfing rate of the population |
| logical indicating whether the population has stochastic demography (this does not include migration), i.e. non-constant size and potentially population growth or decay, depending on the situation it is in. |
| growth rate of the population. |
| A row matrix of the size of the genotype number describing the initial allele frequencies common to all simulations |
| a Genome object |
| initial population size, knowing that if the demography is extinct, the initial population size will automatically be set equal to the population size. |
| a Selection object |
| a MutationMatrix object |
| |

Value

a Population object

Author(s)

Ehouarn Le Faou

initialize, Selection-method

Initialize method for the Selection class

Description

Initialize method for the Selection class

Usage

S4 method for signature 'Selection'
initialize(.Object, genomeObj)

Arguments

| .Object | a Selection object |
|-----------|--------------------|
| genomeObj | a Genome object |

Value

A Selection object

Author(s)

Ehouarn Le Faou

is.correct.transition.matrix

Test if a matrix is a correct transition matrix

Description

Test if a matrix is a correct transition matrix

Usage

```
is.correct.transition.matrix(x, type, name)
```

Arguments

| Х | a matrix. |
|------|----------------------------------------------------------------|
| type | type of the matrice (mutation matrix ? recombination matrix ?) |
| name | the name of the matrix. |

Value

A logical corresponding to whether x is a correct transition matrix, i.e. a square matrix with dimensions greater than 0 and whose rows sum to 1.

Author(s)

is.default.matrix *Test if a matrix is a default matrix*

Description

Test if a matrix is a default matrix

Usage

is.default.matrix(x)

Arguments ×

a matrix.

Value

A logical corresponding to whether x is a default matrix (matrix of dimension 0x0).

Author(s)

Ehouarn Le Faou

is.probability.matrix Test if a matrix is of probability

Description

Test if a matrix is of probability

Usage

is.probability.matrix(x)

Arguments

x a matrix.

Value

A logical corresponding to whether x is a probability matrix (sum of rows equal to 1).

Author(s)

isAffected

Description

Determination for a given genotype or haplotype whether it contains the allelic combination under selection

Usage

isAffected(refDNAtype, selDNAtype)

Arguments

| refDNAtype | the reference allelic combination (that of a genotype or a haplotype |
|------------|----------------------------------------------------------------------|
| selDNAtype | the selected allelic combination |

Value

a logic indicating whether the reference genotype or haplotype is affected by the allelic combination under selection

Author(s)

Ehouarn Le Faou

isHaploSelectFormula Are there any allelic combinations including homozygosity

Description

Test if there are homozygotes in the specified allelic combinations of a list of selection formulas

Usage

```
isHaploSelectFormula(selectFormula)
```

Arguments

selectFormula a list of selection formula

Value

logical indicating if there are homozygotes

listing

Description

Listing from the elements of a vector by producing a string, with comma separation between each element and the word "and" between the last two elements.

Usage

listing(vect)

Arguments

vect a vector of any class.

Value

A listing of the elements of the input vector as a string.

Author(s)

Ehouarn Le Faou

meiosisMatrix Meiosis matrix generation

Description

Generation of the meiosis matrix associated to a Genome object.

Usage

```
meiosisMatrix(genomeObj)
```

Arguments

genomeObj a Genome object

Details

A meiosis matrix is a matrix where the number of rows is equal to the number of genotypes and the number of columns to the number of haplotypes. It is a matrix that allows to pass from parental genotypes to gametic haplotypes by meiosis. It is a probability matrix in that the sum of the values in each row is equal to 1. For a given genotype, the row associated with it describes the probabilistic proportions that lead by meiosis to the production of the other genotypes (and of itself if there are no mutations).

Metapopulation-class

Value

A meiosis matrix (probability matrix that associates to each genotype in a row the probability of producing each of the possible haplotypes).

Author(s)

Ehouarn Le Faou

Metapopulation-class Metapopulation

Description

The class Metapopulation is used to centralise the information relating to the populations that we want to simulate, as well as to define the migration conditions between them if there are several. This class is thus defined by a list of objects Population and a migration matrix.

Slots

populations list of objects Population

nbPop number of populations

names names of the populations

migMat migration matrix between population (if there is more than one)

sizes sizes of the populations

dioecies sexual systems of the populations (they must all be the same)

selfRates selfing rates of the populations

demographies demography parameter of the populations

growthRates growth rates of the populations

initPopSizes initial population sizes of the populations

initGenoFreqs initial genotypic frequencies of the populations

genome a Genome object

genomeIDs ID of the Genome object

mutMat a MutationMatrix object

selection a Selection object

recMat recombination matrix

meiosisMat a meiosis matrix

haploCrossMat an haplotype crossing matrix

haploCrossMatNamed an haplotype crossing matrix with names of genotypes instead of their indices

gametogenesisMat a gametogenesis matrix

- alleleFreqMat a matrix for calculating allelic frequencies
- rawOutputSimul raw output of the simulation function, its refinement is done directly afterwards in the simulate method
- stopCondition list of stop conditions for the simulation (if required)
- IDstopCondition names of stop conditions. They are given an arbitrary name if none is given by the user.

results data.frame.

records list.

customOutput list.

Author(s)

Ehouarn Le Faou

METAPOP_SIMULATION Simulation of a metapopulation

Description

Simulation of a metapopulation

Usage

METAPOP_SIMULATION(nbPop, ids, migMat, nsim, verbose, recording, recordGenGap, drift, nbHaplo, nbGeno, idGeno, nbAlleles, idAlleles, nbLoci, initGenoFreq, meiosisMat, gametogenesisMat, popSize, threshold, dioecy, selfRate,

```
stopCondition,
IDstopCondition,
haploCrossMat,
alleleFreqMat,
gamFit,
indFit,
gamProdFit,
demography,
growthRate,
initPopSize,
nameOutFunct
```

Arguments

)

| nbPop | number of populations in the metapopulation |
|-------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| ids | population IDs |
| migMat | migration matrix |
| nsim | number of simulations |
| verbose | boolean determining if the progress of the simulations should be displayed or not (useful in case of many simulations) |
| recording | a boolean indicating whether to record all mutations, i.e. to record allelic and genotypic frequencies along the simulations |
| recordGenGap | the number of generations between two records during simulation, if the record parameter is TRUE. Whatever the value of this parameter, both the first and the last generation will be included in the record |
| drift | a boolean indicating whether genetic drift should be considered (i.e. whether deterministic simulations are performed or not) |
| nbHaplo | number of haplotypes |
| nbGeno | number of genotypes |
| idGeno | genotypes ID |
| nbAlleles | number of alleles for each loci |
| idAlleles | alleles ID |
| nbLoci | number of loci |
| initGenoFreq | list of initial genotype frequencies in the populations |
| meiosisMat gametogenesisMa | meiosis matrix t |
| | gametogenesis matrix |
| popSize | list population sizes |
| threshold | threshold for simulations |
| dioecy | whether the population(s) is dioecious or not (hermaphrodism) |
| selfRate | list of the selfing rate in populations (only for hermaphroditic population) |

mutation

| <pre>stopCondition IDstopCondition</pre> | list of stop conditions |
|------------------------------------------|--------------------------------------------|
| | vector of stop condition ID |
| haploCrossMat | haplotypes crossing matrix |
| alleleFreqMat | matrix for calculating allelic frequencies |
| gamFit | fitness of gametes |
| indFit | fitness of individuals |
| gamProdFit | fitness for gamete production |
| demography | list of population demographies |
| growthRate | list of population growth rates |
| initPopSize | list of initial population |
| nameOutFunct | name of the custom output function |

Author(s)

Ehouarn Le Faou

mutation

Definition of a mutation

Description

Utility function to easily generate a mutation matrix (see setMutationMatrix).

Usage

mutation(from, to, rate)

Arguments

| from | name of the original allele |
|------|-----------------------------------|
| to | name of the mutant allele |
| rate | rate at which the mutation occurs |

Details

Mutation occurs from one allele to another at a specific rate. Please take care to define alleles as traits, that these alleles are present in the genome you are using and that the alleles are associated with the same locus.

Value

A standardised list of input parameters that will be used by the function setMutationMatrix to generate the mutation matrix.

MutationMatrix-class

Examples

Example with two loci, each with two alleles

```
# Definition of the genome
DL <- list(dl = c("A", "a"))
HL <- list(hl = c("B", "b"))
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)
# The mutation function allows each transition from one allele to
# another to be defined individually, to produce the mutation matrix
# as follows:
mutMatrixObj <- setMutationMatrix(genomeObj,
mutations = list(
    mutation(from = "A", to = "a", rate = 0.1),
    mutation(from = "B", to = "b", rate = 0.1)
)
```

MutationMatrix-class Mutation matrix

Description

A mutation matrix is used to simulate mutations that affect loci. An object of the class MutationMatrix does not only contain a (haplotypic) mutation matrix. It also contains the attributes necessary for the construction and easy-to-read display of this matrix.

Details

The mutation matrix itself is a square matrix of size equal to the number of haplotypes. It is a probability matrix in that the sum of the values in each row is equal to 1. For a given haplotype, the row associated with it describes the probabilistic proportions that lead by mutation of this haplotype to the production of the other haplotypes (and of itself if there are no mutations).

Slots

mutHapLoci a list of haploid locus by locus allelic mulation matrices. mutDipLoci a list of diploid locus by locus allelic mulation matrices. mutLoci a list concatenating mutHapLoci and mutDipLoci nbAlDL a vector of the number(s) of alleles at each haploid locus nbAlHL a vector of the number(s) of alleles at each diploid locus mutationMatrix the haplotypic mutation matrix nbHaplo the number of haplotypes nbDL the number of haploid loci nbHL the number of haploid loci haplotypes the enumeration of haplotypes IDgenome ID of the associated genome

Author(s)

Ehouarn Le Faou

mutMatFriendly Individual mutation definition to allelic mutation matrices

Description

Translation of the list of individually defined mutations into allelic mutation matrices which are then used to generate the genotypic mutation matrix.

Usage

mutMatFriendly(genomeObj, mutations)

Arguments

| genomeObj | a Genome object |
|-----------|-------------------------------------------------------------------|
| mutations | list of mutations defined individually with the function mutation |

Value

A list of the two list of allelic mutation matrices, for haploid and diploid loci respectively.

Author(s)

Ehouarn Le Faou

mutMatRates

Mutation matrix from rates

Description

Generation of a mutation matrix from the allele enumeration vector of a loci and the forward and backward mutation rates.

Usage

mutMatRates(alleles, forwardMut, backwardMut)

Arguments

| alleles | allele enumeration vector of a locus |
|-------------|--------------------------------------|
| forwardMut | forward mutation rate |
| backwardMut | backward mutation rate |

outFunct

Details

See MutationMatrix for more details on mutation matrices.

Value

An allelic mutation matrix (probability matrix which associates to each allele in a row the probability of mutating or not to the other alleles of the locus in question).

Author(s)

Ehouarn Le Faou

outFunct

Custom output function

Description

Allow to produce a custom output for a simulation.

Usage

outFunct(pop)

Arguments

pop

list of some characteristics of the population : - customOutput : list of all previous savings - gen : generation - freqGeno : list of genotypic frequency matrices (matrix 1 x # genotypes). The list is constructed as follows: if the population is hermaphroditic it has only one element "ind", if the population is dioecious it has three elements, "female", "male" and "ind" which correspond respectively to the genotypic frequencies of the females, the males and the average of the two (assuming a sex ratio of 50:50). - freqHaplo : list of genotypic frequency matrices (matrix 1 x # haplotypes). The list is constructed in the same way as for genotypic frequencies (see above). - freqAlleles : list of allelic frequency matrices (matrix 1 x # alleles). The list is constructed in the same way as for genotypic frequencies (see above).

Details

This function is called each generation in each population of a simulation and systematically returns a list with the first element being a logic that indicates whether something should be saved. If so, the second element of this list will be saved.

By default the save nothing function, but it can be changed by the user as an argument in the simulate method of the Metapopulation class.

Author(s)

Population-class Population

Description

The Population class allows for the collection of the parameters necessary to characterise a biological population. It is an essentially useful class in that no method associated with the population class can simulate its dynamics. To do this, it is necessary to use the Metapopulation class, which takes as input a list of populations (from one). The Population class is also used to check that each of these parameters is compatible with each other.

Details

Thus to build an object of class Ease, it is necessary to have defined an object Genome, as well as an object MutationMatrix and an object Selection (even if it is neutral, see setSelectNeutral).

Slots

name the name of the population.

size the size of the population.

dioecy logical indicating whether the population is dioecious or not (hermaphrodite).

selfRate the selfing rate of the population

demography logical indicating whether the population has stochastic demography (this does not include migration), i.e. non-constant size and potentially population growth or decay, depending on the situation it is in.

- growthRate growth rate of the population.
- initGenoFreq A row matrix of the size of the genotype number describing the initial allele frequencies common to all simulations
- genome a Genome object
- initPopSize initial population size, knowing that if the demography is extinct, the initial population size will automatically be set equal to the population size.
- selection a Selection object

mutMat a MutationMatrix object

Author(s)

print, Genome-method *Print method for the* Genome *class*

Description

Print method for the Genome class

Usage

S4 method for signature 'Genome'
print(x, ...)

Arguments

| х | a Genome object |
|---|-----------------|
| | Ignored. |

Value

No return value, only a display.

Author(s)

Ehouarn Le Faou

print,Metapopulation-method

Print method for the Metapopulation class

Description

Print method for the Metapopulation class

Usage

```
## S4 method for signature 'Metapopulation'
print(x, ...)
```

Arguments

| Х | a Metapopulation object |
|---|-------------------------|
| | Ignored. |

Author(s)

print,MutationMatrix-method

Print method for the MutationMatrix class

Description

Print method for the MutationMatrix class

Usage

```
## S4 method for signature 'MutationMatrix'
print(x, ...)
```

Arguments

| Х | a MutationMatrix object |
|---|-------------------------------|
| | there are no more parameters. |

Value

No return value, only a display.

Author(s)

Ehouarn Le Faou

print, Population-method

Print method for the Population class

Description

Print method for the Population class

Usage

```
## S4 method for signature 'Population'
print(x, ...)
```

Arguments

... the other parameter is frame, which is a logic indicating whether the frame surrounding the display of the population characteristics should be displayed or not.

Author(s)

Ehouarn Le Faou

print,Selection-method

Print method for the Selection class

Description

Print method for the Selection class

Usage

S4 method for signature 'Selection'
print(x, ...)

Arguments

| х | a Selection object |
|---|-------------------------------|
| | there are no more parameters. |

Value

No return value, only a display.

Author(s)

Ehouarn Le Faou

recombinationMatrix Recombination matrix generation

Description

Generation of the recombination matrix associated to a Genome object.

Usage

recombinationMatrix(genomeObj)

Arguments

genomeObj a Genome object

Details

A recombination matrix is a square matrix of size equal to the number of genotypes. It is a probability matrix in that the sum of the values in each row is equal to 1. For a given genotype, the row associated with it describes the probabilistic proportions that lead by recombination between diploid loci to the production of the other genotypes (and of itself if there are no mutations).

Value

A recombination matrix (probability matrix which associates to each genotype in a row the probability of recombining or not and of becoming another genotype or remaining the same).

Author(s)

Ehouarn Le Faou

rowResultGen

Processing a result (or record) list

Description

Processing a result (or record) list

Usage

rowResultGen(x)

Arguments

x list of result or record

Value

Merges the column names of the matrices making up the list with the names of the matrices, then merges the matrices together.

Author(s)

Ehouarn Le Faou

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selectFormIntoVect Conversion of selection formulas

Description

Conversion of a list of selection formulas into a genotypic (or haplotypic) fitness vector associated with a Genome object.

Usage

```
selectFormIntoVect(selectFormula, genomeObj, haplo = FALSE)
```

Arguments

| selectFormula | a list of selection formulas |
|---------------|--------------------------------------------------------------------------------------------------------------------|
| genomeObj | a Genome object |
| haplo | logical indicating whether the selection should apply to haplotypes (in the case of gametic selection for example) |

Value

a vector of fitness values

Author(s)

Ehouarn Le Faou

selectInputTreatment Treatment of selection formulas

Description

Determines whether an entry for the selection is a list of selection formulas or just a vector. If it is a list of formulas, turns them into a vector. If it is a vector, does nothing.

Usage

```
selectInputTreatment(selectInput, genomeObj, haplo = FALSE)
```

Arguments

| selectInput | a selection input |
|-------------|--------------------------------------------------------------------------------------------------------------------|
| genomeObj | a Genome object |
| haplo | logical indicating whether the selection should apply to haplotypes (in the case of gametic selection for example) |

Value

a vector of fitness values

Author(s)

Ehouarn Le Faou

Selection-class Selection class

Description

Class used to generate objects that manage the selection in the simulations.

Details

An object of type Selection is an object which describes the set of fitnesses which will be taken into account in the simulations. The selection according to these fitnesses can be applied at three levels: at the level of the individual, at the level of the production of gametes and at the level of the gametes themselves. Selection is therefore genotypic in the first two cases (each genotype is associated with a fitness value) and haplotypic in the third (each haplotype is associated with a fitness value).

Slots

genome a Genome object

IDhaplotypes IDs of haplotypes

IDgenotypes IDs of genotypes

IDgenome ID of the associated genome

nbHaplo the number of haplotypes

nbGeno the number of genotypes

 $\mathsf{gamFit}\ \mathsf{the}\ \mathsf{list}\ \mathsf{of}\ \mathsf{gametes'}\ \mathsf{fitness}$

indFit the list of individuals' fitness

gamProdFit the list of gamete production fitness

s0nInds a logical indicating whether a selection on individuals has been configured by the user

sOnGams a logical indicating whether a selection on gametes has been configured by the user

sOnGamsProd a logical indicating whether a selection on gamete production has been configured by the user

Author(s)

selection.form.treatment

Treatment of a selection formula

Description

Conversion of the factors of a selection formula into the list of corresponding allelic combinations

Usage

```
selection.form.treatment(factors, genomeObj = NULL, checking = FALSE)
```

Arguments

| factors | formula factors (right-hand members) |
|-----------|------------------------------------------------------------------------------------------------|
| genomeObj | a Genome object for the test (see checking parameter) |
| checking | logical indicating whether a test verifying the compatibility of input factors with the genome |

Value

A list of vectors enumerating the allelic combinations that correspond to the factors

Author(s)

Ehouarn Le Faou

setGenome

Setting the genome

Description

Generation of a genome class object from the list of haploid loci and diploid loci. Each loci is defined by a factor vector that enumerates its alleles.

Usage

```
setGenome(listHapLoci = list(), listDipLoci = list(), recRate = numeric())
```

Arguments

| listHapLoci | a list of haploid loci |
|-------------|----------------------------------------|
| listDipLoci | a list of diploid loci |
| recRate | a two-by-two recombination rate vector |

Details

A genome includes the list of all possible haplotypes and genotypes resulting from the combination of the alleles defined in input. As the Ease package was originally built for population genetics simulations including both diploid and haploid loci, it is necessary that both types of loci are defined. Despite this, the user can define only diploid or only haploid loci if they wish. If no diploid locus is defined, one is automatically generated with only one allele, thus not influencing the simulation. The same applies if no haploid locus is defined.

Each locus is described by a vector of factors which are the names of the possible alleles at that locus. All diploid (resp. haploid) loci thus defined are grouped in a list, called listDipLoci (resp. listHapLoci). Therefore, a Genome class object has two lists of loci defined in this way, one for diploid loci, one for haploid loci. The alleles and loci (diploid and haploid) must all have different names so that no ambiguity can persist.

If several are defined, the order of the diploid loci in the list is not trivial. The rates of two-toone combinations between them must indeed be defined by the vector recRate. For example, if three diploid loci are defined, recRate must be of length 2, the first of its values defining the recombination rate between the first and second loci, the second of its values the recombination rate between the second and third loci. For example, if we want to define two groups of two loci that are linked to each other but are on two different chromosomes, we can define a recRate = c(0.1, 0.5, 0.1). The first two loci are thus relatively linked (recombination rate of 0.1), as are the last two loci. On the other hand, the recombination rate of 0.5 between the second and third loci ensures that the two groups are independent.

Value

a Genome object

Author(s)

Ehouarn Le Faou

Examples

```
DL <- list(dl = c("A", "a"))
HL <- list(hl = c("B", "b"))
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)</pre>
```

setMetapopulation Setting a metapopulation

Description

A metapopulation is a set of population(s) (from 1) that are simulated with potential migration between them. Only genotypes can migrate, i.e. adult individuals.

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setMetapopulation

Usage

setMetapopulation(populations, migMat = matrix(1))

Arguments

| populations | a list of Population objects |
|-------------|------------------------------|
| migMat | a migration matrix |

Details

The construction of a Metapopulation object requires only two arguments (one optional). The first is a population(s) list, defined from the population class. The second is a migration matrix, which connects the populations together. This matrix is a probability matrix (square with the sum of the rows equal to 1, whose size is equal to the number of populations) where each value corresponds to the proportion of individuals (genotypes) that disperse from their source population (row) to their target population (column).

Value

a Metapopulation object

Author(s)

Ehouarn Le Faou

```
# Definition of a population in its simplest form:
DL <- list(dl = c("A", "a"))</pre>
HL <- list(hl = c("B", "b"))
mutations <- list(</pre>
  mutation(from = "A", to = "a", rate = 1e-3),
  mutation(from = "B", to = "b", rate = 1e-3)
)
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)</pre>
pop <- setPopulation(</pre>
  name = "A",
  size = 1000,
  dioecy = TRUE,
  genomeObj = genomeObj,
  selectionObj = setSelectNeutral(genomeObj),
  mutMatrixObj = setMutationMatrix(genomeObj, mutations = mutations)
)
metapop <- setMetapopulation(populations = list(pop))</pre>
metapop <- simulate(metapop, nsim = 10, seed = 123)</pre>
# Other examples available in the documentation of the package
```

setMutationMatrix Setting the mutation matrix

Description

Generation of the mutation matrix associated with the genome given as input. A mutation matrix is used to simulate mutations that affect loci. An object of the class MutationMatrix does not only contain a (genotypic) mutation matrix. It also contains the attributes necessary for the construction and easy-to-read display of this matrix. The mutation matrix itself is a square matrix of size equal to the number of genotypes. It is a probability matrix in that the sum of the values in each row is equal to 1. For a given genotype, the row associated with it describes the probabilistic proportions that lead by mutation of this genotype to the production of the other genotypes (and of itself if there are no mutations).

Usage

```
setMutationMatrix(genomeObj, ...)
```

Arguments

genomeObj a Genome object ... see details.

Details

There are three ways to define the mutation matrix associated with a Genome class object.

1) By giving two lists of allelic mutation matrices mutHapLoci and mutDipLoci, for haploid and diploid loci respectively. Each of these lists contains as many matrices as there are loci. These matrices are transition matrices (squares, with the sum of the rows equal to 1) of size equal to the number of alleles at the locus concerned.

2) By giving a forward and a backward allelic mutation rate (forwardMut and backwardMut respectively). The generated mutation matrices will thus be defined with the same rates for all loci. A forward mutation rate means that the transition from one allele to another is done in the order in which they were defined when the Genome class object was created, and in the other direction for the backward rate.

3) By giving a list of mutations generated through the mutation function.

Value

a MutationMatrix object

Author(s)

setPopulation

Examples

```
### Example with two loci, each with two alleles ###
# Definition of the genome
DL <- list(dl = c("A", "a"))</pre>
HL <- list(hl = c("B", "b"))
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)</pre>
# Three ways to define the same mutation matrix associated with the
# genome defined above:
# 1) Mutation matrix from matrices
mutHapLoci <- list(matrix(c(0.99, 0.01, 0.01, 0.99), 2))</pre>
mutDipLoci <- list(matrix(c(0.99, 0.01, 0.01, 0.99), 2))</pre>
# One can then define the MutationMatrix class object:
setMutationMatrix(genomeObj,
  mutHapLoci = mutHapLoci,
  mutDipLoci = mutDipLoci
)
# 2) Mutation matrix from mutation rates
mutMatrixObj <- setMutationMatrix(genomeObj, forwardMut = 0.1)</pre>
# or by adding a backward mutation rate:
mutMatrixObj <- setMutationMatrix(genomeObj,</pre>
  forwardMut = 1e-3,
  backwardMut = 1e-4
)
# 3) Mutation matrix from single mutation definition
mutMatrixObj <- setMutationMatrix(genomeObj,</pre>
  mutations = list(
    mutation(from = "A", to = "a", rate = 0.1),
    mutation(from = "B", to = "b", rate = 0.1)
  )
)
```

setPopulation Setting a population

Description

Generation of a population by providing all the necessary ingredients for its definition, including a genome, a mutation matrix and a selection regime.

Usage

```
setPopulation(
    name,
```

```
size,
dioecy,
genomeObj,
mutMatrixObj,
selectionObj,
selfRate = 0,
demography = F,
growthRate = 0,
initPopSize = NULL,
initGenoFreq = NULL)
```

Arguments

| name | the name of the population |
|--------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| size | the population size |
| dioecy | logical indicating whether the simulated population is dioecious or hermaphroditic |
| genomeObj | a Genome object |
| mutMatrixObj | a MutationMatrix object |
| selectionObj | a Selection object |
| selfRate | the selfing rate |
| demography | a logic indicating whether the population should have a demography (stochas- ticity in the number of individuals present in the population + logistic growth with carrying capacity equal to the size parameter) |
| growthRate | a Genome object |
| initPopSize | the initial size of the population. It is necessarily equal to size if the population has no demography. |
| initGenoFreq | a vector of the size of the genotype number describing the initial allele frequen- cies common to all simulations |

Details

A population is defined strictly by a name, a size, a sexual system (dioecy or hermaphodite), and the three objects defined previously: genome, mutation matrix and selection. In addition to that, it is possible to define - a selfing rate (by default equal to 0) - a vector of initial genotypic frequencies - a demography

Two demographic regimes are possible: no demography, i.e. a fixed population size, or demography, i.e. a population where the size fluctuates stochastically. The boolean argument 'demography' is used to define whether there should be stochasticity. For a fixed population size, it is therefore sufficient to define that 'demography = FALSE' (default) and to set the desired population size with the 'popSize' parameter.

For a fluctuating demography, 'demography' must be 'TRUE' and three other parameters are then needed: the initial population size ('initPopSize'), the population growth rate ('growthRate') and the carrying capacity of the population (the population size, 'popSize').

It is also possible to avoid defining a population size altogether, by setting off the genetic drift ('drift' parameter). This will allow the model to be simulated deterministically.

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setSelectNeutral

Value

a Population object

Author(s)

Ehouarn Le Faou

Examples

```
# Definition of a population in its simplest form:
DL <- list(dl = c("A", "a"))</pre>
HL <- list(hl = c("B", "b"))
mutations <- list(</pre>
 mutation(from = "A", to = "a", rate = 1e-3),
  mutation(from = "B", to = "b", rate = 1e-3)
)
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)</pre>
pop <- setPopulation(</pre>
  name = "A",
  size = 1000,
  dioecy = TRUE,
  genomeObj = genomeObj,
  selectionObj = setSelectNeutral(genomeObj),
  mutMatrixObj = setMutationMatrix(genomeObj, mutations = mutations)
)
```

setSelectNeutral Setting the selection

Description

Generation of a neutral class Selection object. It can be used as a basis for adding selection layers with the setSelectOnInds, setSelectOnGametes or setSelectOnGametesProd functions, or if the model is neutral.

Usage

```
setSelectNeutral(genomeObj)
```

Arguments

genomeObj a Genome object

Details

An object of type Selection is an object which describes the set of fitnesses which will be taken into account in the simulations. The selection according to these fitnesses can be applied at three levels: at the level of the individual, at the level of the production of gametes and at the level of the gametes themselves. Selection is therefore genotypic in the first two cases (each genotype is associated with a fitness value) and haplotypic in the third (each haplotype is associated with a fitness value).

Value

a Selection object

Author(s)

Ehouarn Le Faou

Examples

```
### Example with two loci, each with two alleles ###
# Definition of the diploid locus
DL <- list(dl = c("A", "a"))</pre>
# Definition of the haploid locus
HL <- list(hl = c("B", "b"))
# Definition of the object of Genome class
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)</pre>
genomeObj
### Exemple with more diploid loci ###
# Definition of the diploid loci
DL <- list(
  dl1 = c("A", "a"),
  dl2 = c("B", "b"),
  dl3 = c("C", "c")
)
# Definition of the haploid locus
HL <- list(hl = c("D", "d"))
# Definition of the object of Genome class, with in addition the necessary
# definition of recombination rates between loci:
genomeObj <- setGenome(</pre>
  listHapLoci = HL, listDipLoci = DL,
  recRate = c(0.1, 0.5)
)
# Here we have a 0.1 recombination rate between dl1 and dl2 and a 0.5
# recombination rate between dl2 and dl3. It is as if dl1 and dl2 were linked,
# for example on the same chromosome, and that dl2 (and dl1 by consequence)
# and dl3 were independent, for example on different chromosomes.
```

genomeObj

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setSelectOnGametes Setting the selection on gametes

Description

Generation of an object of the Selection class which defines a selection among the individuals either by adding this type of selection to an already existing SelectionObj object (parameter selectionObj) or by creating one.

Usage

```
setSelectOnGametes(
  genomeObj = NULL,
  gamFit = c(),
  femaleFit = c(),
  maleFit = c(),
  selectionObj = NULL
)
```

Arguments

| genomeObj | a Genome object |
|--------------|--------------------------------------------------------------------------------------------------------------------|
| gamFit | an haplotypic fitness vector for all individuals |
| femaleFit | an haplotypic fitness vector for females only |
| maleFit | an haplotypic fitness vector for males only |
| selectionObj | a Selection object (in the case where the selection on individuals is overlaid on an existing Selection object) |

Value

a Selection object

Author(s)

Ehouarn Le Faou

```
DL <- list(dl = c("A", "a"))
HL <- list(hl = c("B", "b"))
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)
selectionObj <- setSelectOnGametes(
  genomeObj = genomeObj,
  gamFit = c(1, 1, 0.5, 0)
)</pre>
```

```
setSelectOnGametesProd
```

Setting the selection on gamete production

Description

Generation of an object of the Selection class which defines a selection on the gamete production either by adding this type of selection to an already existing SelectionObj object (parameter selectionObj) or by creating one.

Usage

```
setSelectOnGametesProd(
  genomeObj = NULL,
  indProdFit = c(),
  femProdFit = c(),
  maleProdFit = c(),
  selectionObj = NULL
)
```

Arguments

| genomeObj | a Genome object |
|--------------|-----------------------------------------------------------------------------------------------------------------|
| indProdFit | a genotypic fitness vector for all individuals |
| femProdFit | a genotypic fitness vector for females only |
| maleProdFit | a genotypic fitness vector for males only |
| selectionObj | a Selection object (in the case where the selection on individuals is overlaid on an existing Selection object) |

Value

a Selection object

Author(s)

Ehouarn Le Faou

```
DL <- list(dl = c("A", "a"))
HL <- list(hl = c("B", "b"))
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)
selectionObj <- setSelectOnGametesProd(
  genomeObj = genomeObj,
   indProdFit = c(1, 1, 1, 1, 0.5, 0)
)</pre>
```

setSelectOnInds

Description

Generation of an object of the Selection class which defines a selection among the individuals either by adding this type of selection to an already existing SelectionObj object (parameter selectionObj) or by creating one.

Usage

```
setSelectOnInds(
  genomeObj = NULL,
  indFit = c(),
  femaleFit = c(),
  maleFit = c(),
  selectionObj = NULL
)
```

Arguments

| genomeObj | a Genome object |
|--------------|--------------------------------------------------------------------------------------------------------------------|
| indFit | a genotypic fitness vector for all individuals (whether or not they are hermaphordite) |
| femaleFit | a genotypic fitness vector for females only (only if the population is dioecious) |
| maleFit | a genotypic fitness vector for males only (only if the population is dioecious) |
| selectionObj | a Selection object (in the case where the selection on individuals is overlaid on an existing Selection object) |

Value

a Selection object

Author(s)

Ehouarn Le Faou

```
DL <- list(dl = c("A", "a"))
HL <- list(hl = c("B", "b"))
genomeObj <- setGenome(listHapLoci = HL, listDipLoci = DL)
selectionObj <- setSelectOnInds(
  genomeObj = genomeObj,
  indFit = c(1, 1, 1, 1, 0.5, 0)
)</pre>
```

show, Genome-method Show method for the Genome class

Description

Show method for the Genome class

Usage

S4 method for signature 'Genome'
show(object)

Arguments

object a Genome object

Value

No return value, only a display.

Author(s)

Ehouarn Le Faou

 $\verb|show,Metapopulation-method||$

Show method for the Metapopulation class

Description

Show method for the Metapopulation class

Usage

S4 method for signature 'Metapopulation'
show(object)

Arguments

object a Metapopulation object

Author(s)

show,MutationMatrix-method

Show method for the MutationMatrix class

Description

Show method for the MutationMatrix class

Usage

```
## S4 method for signature 'MutationMatrix'
show(object)
```

Arguments

object a MutationMatrix object

Value

No return value, only a display.

Author(s)

Ehouarn Le Faou

show, Population-method

Show method for the Population class

Description

Show method for the Population class

Usage

```
## S4 method for signature 'Population'
show(object)
```

Arguments

object a Population object

Author(s)

show, Selection-method Show method for the Selection class

Description

Show method for the Selection class

Usage

S4 method for signature 'Selection'
show(object)

Arguments

object a Selection object

Value

No return value, only a display.

Author(s)

Ehouarn Le Faou

simulate,Metapopulation-method

Simulate method for the Metapopulation class

Description

Performing simulations of an Metapopulation object. The returned object is the same Metapopulation object completed with the results and records if they have been activated.

Usage

```
## S4 method for signature 'Metapopulation'
simulate(
   object,
   nsim = 1,
   seed = NULL,
   threshold = 500,
   includefreqGeno = TRUE,
   recording = FALSE,
   recordGenGap = 1,
   drift = TRUE,
   includeParams = TRUE,
```

```
includeFitness = TRUE,
verbose = FALSE,
stopCondition = list(),
nameOutFunct = "outFunct"
)
```

Arguments

| object | a Metapopulation object |
|-----------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| nsim | the number of simulation to perform |
| seed | the RNG seed to be fixed (allows exact reproduction of results) |
| threshold | maximum duration of a simulation (in generations) |
| includefreqGeno | |
| | a logical indicating whether to include genotype frequencies in the results |
| recording | a logical indicating whether to record all mutations, i.e. to record allelic and genotypic frequencies along the simulations |
| recordGenGap | the number of generations between two records during simulation, if the record parameter is TRUE. Whatever the value of this parameter, both the first and the last generation will be included in the record |
| drift | a logical indicating whether genetic drift should be considered (i.e. whether deterministic simulations are performed or not) |
| includeParams | a logical indicating whether the parameters should be included in the result data.frame (can be useful when compiling multiple result tables) |
| includeFitness | a logical indicating whether the mean fitness should be included in the result data.frame (can be useful when compiling multiple result tables) |
| verbose | logical determining if the progress of the simulations should be displayed or not (useful in case of many simulations) |
| stopCondition | list of vectors that each describe the allele(s) that must be fixed to define a stop condition. Each of these vectors will therefore be associated with a stop condition |
| nameOutFunct | name of the custom output function. This function is called each generation in each population of a simulation and systematically returns a list with the first element being a logic that indicates whether something should be saved. If so, the second element of this list will be saved. If the customOutFunct parameter is null (default), there will be no custom output. |

Value

An Metapopulation object from which we can now extract the results (or the records if recording = TRUE) with the getResults and getRecords functions.

Author(s)

whichHomoz

Description

Determine which alleles are at least once input as homozygous in the formula.

Usage

whichHomoz(formula)

Arguments

formula a selection formula

Value

the enumeration of alleles that appear at least once homozygous

Author(s)

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