Package ‘isqg’

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Type Package

Title In Silico Quantitative Genetics

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Description Accomplish high performance simulations in quantitative genetics.
The molecular genetic components are represented by R6/C++ classes and methods.
Mimic the meiosis recombination and de novo genetic variability by means a
The core computational algorithm is implemented using 'Boost' dynamic bitsets
(Schaling, 2014) [ISBN:978-1937434366]. A mix between low and high level interfaces
provides great flexibility and allows user defined extensions and a wide range of
applications.

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Encoding UTF-8

NeedsCompilation yes

SystemRequirements C++11

Imports Rcpp (>= 0.12.15), R6

LinkingTo Rcpp, BH

Collate 'ISQG.R' 'Mating.R' 'R6Classes.R' 'Functions.R' 'Trait.R'
'RcppExports.R' 'Hooks.R'

LazyData true

RoxygenNote 6.0.1

Repository CRAN

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- isqg-package  
  isqg: A package to perform in silico quantitative genetics

Description

isqg provides R6/C++ classes for in silico quantitative genetics. Mimic the meiosis recombination. Allows user defined extensions which provides great flexibility.

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- fitness  
  Simulated Trait for Individuals According to Basic Models

Description

Constructor of instances of the Trait class given the focal specie and the parameters to define infinitesimal or quantitative fitness.

Usage

set_infty(specie, m = 0, a = 1, d = 0, genes = NULL)

set_quant(specie, m, data)
founder

Constructor of a Founder Instances of the Specimen Class

Description

Constructor of instances of the Specimen class given the Specie from which the individual will belong where all loci will equal to the provided genotype.

Usage

founder(specie, code)

Arguments

specie an instance of the R6 class Specie with the genome’s parameters.
m, a, d a length-one numeric vector with respectively the mean, the additive and the dominant effects.
genes a character vector with the putative genes.
data a data frame with the genes (snp) and their additive and dominant effects

Details

Infinitesimal traits need the mean, the additive and the dominant effect and optionally the vector of the putative genes. Quantitative traits are defined given the mean and a data frame with the putative genes and their additive and dominant effects.

Value

Objects of R6 class with methods to mimic in silico Traits.

Examples

data(ToyMap)
spc <- set_specie(ToyMap)
AA <- founder(spc, "AA")
aa <- spc$founder("aa")

F1 <- cross(n = 1, AA, aa) # the hybrid

## set a infinitesimal & a quantitative fitness
infity <- set_infty(spc, m = 0, a = 1, d = .5) # partial dominance
gen <- data.frame(snp = sample(ToyMap$snp, 10), add = rnorm(10), dom = rnorm(10))quant <- set_quant(spc, m = 0, data = genes)

## evaluating the breeding value
infity$alpha(AA)
quant$alpha(F1)
Arguments

specie an instance of the R6 class Specie with the genome’s parameters.

code a length one character vector with one of the genotype codes: "AA", "Aa", "aA" or "aa".

Details

Genotypes can be coded as AA, Aa, aA or aa, that meant to represent both homozigous (AA and aa) as well as both heterozigous (Aa and aA).

Value

Objects of R6 class with methods to mimic in silico Specimens.

Examples

```r
data(ToyMap)
spc <- set_specie(ToyMap)

## through standalone function
AA <- founder(spc, "AA")
aa <- founder(spc, "aa")

## or by the Specie’s method
Aa <- spc$founder("Aa")
aA <- spc$founder("aA")
```

<table>
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<th>Codify Specimens` Genotypes</th>
</tr>
</thead>
</table>

Description

Codify Specimens’ genotypes instances as numeric codes [-1/0/1] or as character vector that keeps the phase information.

Usage

```r
genotype(pop, phase = FALSE)
```

Arguments

pop a list with instances of the R6 class Specimen.

phase logical should the codes keep the phase.

Value

A numeric or character matrix with the codified Specimens’ genotypes.
import

Examples

data(ToyMap)
spc <- set_specie(ToyMap)

Aa <- founder(spc, "Aa")
aA <- spc$founder("aA")

Both <- list(Aa = Aa, aA = aA)

## different ways
genotype(Both) # as numeric
genotype(Both, phase = TRUE) # as character

import Constructor of a Custom Instances of the Specimen Class

Description

Constructor of instances of the Specimen class given the Specie from which the individual will belong where the loci will equal to the provided genotype from two strings one for each homologous.

Usage

import(specie, genotype)

Arguments

specie an instance of the R6 class Specie with the genome’s parameters.
genotype a named character vector with the coded/phased genotypes.

Value

Objects of R6 class with methods to mimic in silico Specimens.

Examples

data(ToyMap)
spc <- set_specie(ToyMap)

## simulating what is very close to your real genotypes
Real <- sample(c(’2 2’, ’2 1’, ’1 2’, ’1 1’), size = nrow(ToyMap), replace = TRUE)
names(Real) <- ToyMap$snp # ensure snp names!

## now you can play _in silico_
Virtual <- import(spc, Real)
SI <- Virtual$selfcross(n = 10)
Breed Simulated Individuals According to Basic Mating Schemes

Description

Performs the simple mating schemes bi-parental cross, self-cross and haploid duplication, respectively through the functions `cross`, `selfcross` and `dh` and return the respective size `n` progeny involving the parental individuals belonging to the same specie.

Usage

```r
cross(n = 1, p1, p2)
selfcross(n = 1, gid)
dh(n = 1, gid)
```

Arguments

- `n` a length-one integer vector with the size of the progeny.
- `p1, p2, gid` are instances of the class specimen which will be used as the parents.

Details

Basically this family of functions take simulated individuals belonging to the same simulated specie, performs the meiosis that generates individual’s gametes. According to the scheme applied the gametes are merged into new simulated individuals. These are wrap functions to the C++ class that mimic the meiosis recombination process.

Value

a size `n` list with instances of the class Specimen that represent new individuals belonging to the progeny of the respective mating scheme.

Examples

```r
data(ToyMap)
spc <- set_specie(ToyMap)
AA <- founder(spc, "AA")
aa <- founder(spc, "aa")

## Mather Design
F1 <- cross(n = 1, AA, aa)
BC1 <- cross(n = 5, F1, AA)
BC2 <- F1 %*% cross(n = 5, aa) # using R6 methods
F2 <- selfcross(n = 10, F1)
RIL <- dh(n = 10, F1)
## chainable R6 methods
```
set_specie

F3 <- F1$selfcross(n = 1, replace = TRUE)$selfcross(n = 1, replace = TRUE)

---

**Description**

Constructor of instances of the Specie class given the map of the genome and optionally a pointer to a C++ function which will drive the meiosis process.

**Usage**

```r
set_specie(data, meiosis = NULL)
```

**Arguments**

- `data`  A data frame with the map of the Genome to be simulates.
- `meiosis`  A pointer to a C++ function of the meiosis process.

**Value**

Objects of R6 class with methods to mimic in silico Genomes.

**Examples**

```r
data(toymap)
spc_standard <- set_specie(toymap)

## generate standard de novo variability
spc_standard$gamete(n = 100)

## Not run:
## write your function is C++ and then wrap it as a pointer
Meiosis <- "
// [[Rcpp::depends(isqg)]]

#include <isqg.h> // loading headers of the package
#include <vector>
#include <algorithm>

// NOTE:
// loci are independent to each other
Map meiosis(Chromosome * group) {

    Map map(group->get_map()) ;
    for (auto it = 0; it < map.size(); it++)
        if (static_cast<bool>(R::rbinom(1., .5))) map.at(it) = 2. + 1. ;
```
Specie

Class providing object with methods to mimic in silico Genomes

Description

Mean to mimic a in silico Genomes. It is the machine instances of the simulator.

Details

Object of R6 class that points to C++ objects.

Value

Objects of R6 class with methods to mimic in silico Genomes.

Fields

.ptr External pointer to the instance of the C++ class Specie.
Specimen

Class providing object with methods to mimic in silico Specimens

Description

Mean to mimic a in silico Specimens. It is the working instances of the simulator.

Details

Object of R6 class that points to C++ objects.

Value

Objects of R6 class with methods to mimic in silico Specimens.

Fields

.ptr External pointer to the instance of the C++ class Specimen.

ToyMap

Toy Example of a Map for in silico Quantitative Genetics

Description

This data comprise 2 chromosomes with 2cM each and 21 monitored loci in each chromosome

Usage

data(ToyMap)

Format

a data.frame, 42 rows and 3 columns (snp, chr, pos).
**Description**

Mean to mimic a in silico Trait. It is the working instances of the simulator.

**Details**

Object of R6 class that points to C++ objects.

**Value**

Objects of R6 class with methods to mimic in silico Traits.

**Fields**

\.ptr  External pointer to the instance of the C++ class Trait.
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