Package 'BitBreedingSim'

December 18, 2024

Type Package	
Title Fast Breeding Simulation	
Version 0.1.0	
Author Minoru Inamori	
Maintainer Minoru Inamori <inamori@ut-biomet.org></inamori@ut-biomet.org>	
Description Use bit operations to speed up breeding simulations.	
License MIT License	
Imports Rcpp (>= 1.0.5)	
LinkingTo Rcpp	
RoxygenNote 7.3.2	
Encoding UTF-8	
Suggests roxygen2	
Roxygen list(markdown = TRUE)	
Contents	
addTraitA	
createBaseInfo	

 createOrigins
 5

 cross
 5

 getGenotypes
 6

 getInfo
 7

 getMap
 7

 getPhasedGenotypes
 8

 getPhasedIntGenotypes
 8

 getPhenotypes
 9

 getPopInfo
 9

 getPopNames
 10

 getTrait
 10

 joinPops
 11

 selectPop
 11

 writeVCF
 12

2 addTraitAD

Index 13

addTraitA	Add Trait with Additive Effects to BaseInfo	

Description

This function adds a trait with additive effects to the BaseInfo object.

Usage

```
addTraitA(info, name, mean, h2, sd = NULL, a = NULL, loci = NULL, num_loci = 1)
```

Arguments

info	External pointer to BaseInfo object
name	Name of the trait
mean	Phenotype mean
h2	Heritability
sd	Optional. Phenotype standard deviation
а	Optional. Numeric vector of additive effects
loci	Optional. List of loci in the form of a data frame with two columns: 'chrom' and 'marker'. For example:
	<pre>chrom <- c(3, 5, 10) marker <- c(1, 2, 1000) loci <- data.frame(chrom, marker)</pre>
num_loci	Optional. Number of loci (default is 1)

addTraitAD Add Trait	vith Additive and Dominance	Effects to BaseInfo
----------------------	-----------------------------	---------------------

Description

This function adds a trait with additive and dominance effects to the BaseInfo object.

createBaseInfo 3

Usage

```
addTraitAD(
  info,
  name,
  mean,
  sd = NULL,
  h2 = NULL,
  H2 = NULL,
  a = NULL,
  d = NULL,
  loci = NULL,
  num_loci = 1
)
```

Arguments

info	External pointer to BaseInfo object
name	Name of the trait
mean	Phenotype mean
sd	Optional. Phenotype standard deviation
h2	Optional. Narrow-sense heritability (proportion of variance due to additive genetic effects)
H2	Optional. Broad-sense heritability (proportion of variance due to all genetic effects, can be NULL)
а	Optional. Numeric vector of additive effects
d	Optional. Numeric vector of dominance effects
loci	Optional. List of loci in the form of a data frame with two columns: 'chrom' and 'marker'. For example:
	<pre>chrom <- c(3, 5, 10) marker <- c(1, 2, 1000) loci <- data.frame(chrom, marker)</pre>
num_loci	Optioal. Number of loci (default is 1)

 ${\tt createBaseInfo}$

Create a BaseInfo object

Description

This function creates a BaseInfo object. If the seed is set to -1, a random seed is generated, resulting in different outcomes each time the function is called. If a specific seed is provided, the random number generation will be based on that seed, ensuring reproducible results.

4 createBaseInfo

Usage

```
createBaseInfo(
  chrom_maps = NULL,
  num_chroms = 10,
  num_markers = 1000,
  cM = 100,
  bp = 1e+06,
  seed = -1
)
```

Arguments

chrom_maps A list of data.frames, each representing a chromosome map. Each data.frame should have two columns: 'cM' for centiMorgans and 'position' for base pair positions. The list should be named, with each name corresponding to a chromosome identifier (e.g., "chr1", "chr2", etc.). If chrom_maps is provided, the parameters num_chroms, num_markers, cM, and bp are ignored. num_chroms Optional. An integer. Number of chromosomes. Ignored if chrom_maps is provided. Default is 10. Optional. An integer. Number of markers per chromosome. Ignored if chrom_maps num_markers is provided. Default is 1000. cM A numeric. Optional. Length of each chromosome in centiMorgans. Ignored if chrom_maps is provided. Default is 100. An integer. Optional. Length of each chromosome in base pairs. Ignored if bp

chrom_maps is provided. Default is 1000000.

An integer. Optional. A seed for random number generation. Default is -1,

which generates a random seed.

Value

seed

An external pointer to a BaseInfo object.

Examples

```
# Create a BaseInfo object with a random seed
base_info_random <- createBaseInfo()
getInfo(base_info_random)

# Create a BaseInfo object with a specific seed for reproducible results
base_info_reproducible <- createBaseInfo(seed = 123)
getInfo(base_info_reproducible)

# Create a chromosome map with 100 cM and 1 Mbp, containing 1000 markers
f <- function(x) { (x^3 / (1 + x^2) + 8/5) * 500 / 16 }
cM <- sapply(1:1000, function(i) f(i/250 - 2))
position <- sapply(1:1000, function(i) i * 1000)
chrom_map <- data.frame(cM, position)
chrom_maps <- replicate(10, chrom_map, simplify = FALSE)</pre>
```

createOrigins 5

```
names(chrom_maps) <- paste0("chr", 1:10)
info <- createBaseInfo(chrom_maps, seed = 123)
getInfo(info)</pre>
```

createOrigins

Create origins for a Population object

Description

Create origins for a Population object

Usage

```
createOrigins(num_inds, info, name_base)
```

Arguments

num_inds An integer. The number of individuals.

info An external pointer to a BaseInfo object.

name_base A string. The base name for individuals.

Value

An external pointer to a Population object.

cross

Cross two Population

Description

Cross two Population

Usage

```
cross(num_inds, mothers, fathers, name_base, num_threads = 0)
```

Arguments

num_inds An integer. The number of individuals.

mothers An external pointer to a Population object.

fathers An external pointer to a Population object.

name_base A string. The base name for individuals.

num_threads Optional. An integer. The number of threads to be used. If not specified, the

function will use the maximum number of available threads.

6 getGenotypes

Value

An external pointer to a Population object.

Examples

```
# Assuming 'mothers' and 'fathers' are valid Population objects
new_population <- cross(100, mothers, fathers, "prog_")
summary(new_population)</pre>
```

getGenotypes

Get genotypes from a Population object

Description

This function retrieves the genotypes from a given Population object. The genotypes are represented in a matrix format where rows correspond to samples and columns correspond to markers.

Usage

```
getGenotypes(pop)
```

Arguments

pop

An external pointer to a Population object.

Details

Genotype encoding:

- 0/0 is encoded as -1
- 0/1 is encoded as 0
- 1/1 is encoded as 1

Value

A matrix of genotypes where rows are samples and columns are markers.

getInfo 7

getInfo

Get the values of a BaseInfo object

Description

Get the values of a BaseInfo object

Usage

```
getInfo(info)
```

Arguments

info

An external pointer to a BaseInfo object

Value

The list of the values of a BaseInfo object

getMap

Retrieve Genetic Map Information

Description

This function retrieves the genetic map information from a BaseInfo object.

Usage

```
getMap(info)
```

Arguments

info

An object of class BaseInfo. This object contains the genetic map information.

Value

A list of data frames, each representing a chromosome. Each data frame contains two columns:

- cM: The centiMorgan positions of the markers.
- position: The base pair positions of the markers.

Examples

```
## Not run:
    # Assuming `info` is a valid BaseInfo object
    map <- getMap(info)
    print(map)
## End(Not run)</pre>
```

 ${\tt getPhasedGenotypes}$

Get phased genotypes from a Population object

Description

This function retrieves the genotypes from a given Population object. The genotypes are represented in a matrix format where rows correspond to markers and columns correspond to samples.

Usage

getPhasedGenotypes(pop)

Arguments

pop

An external pointer to a Population object.

Details

Genotype is 010, 011, 110, or 111

Value

A matrix of genotypes where rows are samples and columns are markers.

getPhasedIntGenotypes Get phased integer genotypes from a Population object

Description

This function retrieves the genotypes from a given Population object. The genotypes are represented in a matrix format where rows correspond to samples and columns correspond to markers. Each sample has two rows: the first row represents the maternal allele and the second row represents the paternal allele.

Usage

getPhasedIntGenotypes(pop)

Arguments

pop

An external pointer to a Population object.

Details

Genotype is represented as integers: 0 or 1

getPhenotypes 9

Value

A matrix of genotypes where rows are samples and columns are markers. Each sample has two rows: the first row is the maternal allele and the second row is the paternal allele.

getPhenotypes

Get phenotypes from a Population object

Description

Get phenotypes from a Population object

Usage

```
getPhenotypes(pop, i)
```

Arguments

pop An external pointer to a Population object.

i An integer index representing the trait for which phenotypes are to be retrieved.

The index should be between 1 and the total number of traits available in the

Population object.

Value

A vector of phenotypes.

Examples

```
# Assuming 'pop' is a valid Population object and trait index 1 is valid
phenotypes <- getPhenotypes(pop, 1)
print(phenotypes)</pre>
```

getPopInfo

Get information for a Population object

Description

Get information for a Population object

Usage

```
getPopInfo(pop)
```

Arguments

pop

An external pointer to a BaseInfo object.

10 getTrait

Value

A list containing the number of individuals and the number of chromosomes in the population.

getPopNames

Get name data from a Population object

Description

Get name data from a Population object

Usage

```
getPopNames(pop)
```

Arguments

pop

An external pointer to a Population object.

Value

A data frame containing the names, maternal names, and paternal names from the Population object.

Examples

```
# Assuming 'pop' is a valid Population object
name_data <- getPopNames(pop)
print(name_data)</pre>
```

getTrait

Get a trait from a BaseInfo object

Description

Get a trait from a BaseInfo object

Usage

```
getTrait(info, i)
```

Arguments

info An external pointer to a BaseInfo object.i An integer. The index of the trait to retrieve.

Value

The trait at the specified index.

joinPops 11

joinPops

Join multiple Population objects

Description

Join multiple Population objects

Usage

```
joinPops(...)
```

Arguments

External pointers to Population objects.

Value

An external pointer to a new Population object containing the combined individuals.

Examples

```
# Assuming 'pop1', 'pop2', and 'pop3' are valid Population objects
combined_pop <- joinPops(pop1, pop2, pop3)
print(combined_pop)</pre>
```

selectPop

Select individuals from a Population object

Description

Select individuals from a Population object

Usage

```
selectPop(pop, indices_R)
```

Arguments

pop

An external pointer to a Population object.

indices

A vector of integer indices representing the individuals to be selected.

Value

An external pointer to a new Population object containing the selected individuals.

12 writeVCF

Examples

```
# Assuming 'pop' is a valid Population object and indices are valid
selected_pop <- selectPop(pop, c(1, 2, 3))
print(selected_pop)
```

writeVCF

Write Population to VCF file

Description

This function writes a Population object to a VCF file.

Usage

```
writeVCF(pop, filename)
```

Arguments

pop An external pointer to a Population object.

filename A string specifying the path to the output VCF file.

Examples

```
# Assuming 'pop' is a valid Population object
writeVCF(pop, "output.vcf")
```

Index

```
addTraitA, 2
addTraitAD, 2
{\tt createBaseInfo, 3}
createOrigins, 5
cross, 5
getGenotypes, 6
getInfo, 7
getMap, 7
{\tt getPhasedGenotypes}, \textcolor{red}{8}
{\tt getPhasedIntGenotypes}, \\ 8
getPhenotypes, 9
getPopInfo, 9
{\tt getPopNames}, \\ 10
getTrait, 10
joinPops, 11
selectPop, 11
writeVCF, 12
```