Package 'BitBreedingSim'

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Type Package

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License MIT License
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addTraitA A

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
num_loci	Number of loci (default is 1)

addTraitAD

Add Trait with Additive and Dominance Effects to BaseInfo

Description

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

Usage

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

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Arguments

info	External pointer to BaseInfo object
name	Name of the trait
mean	Phenotype mean
a	Optional numeric vector of additive effects
d	Optional numeric vector of dominance effects
loci	Optional list of loci
num_loci	Number of loci (default is 1)
Optional	H2 Broad-sense heritability (proportion of variance due to all genetic effects,

createBaseInfo

Create a BaseInfo object

can be NULL)

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.

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	An integer. Number of chromosomes. Ignored if chrom_maps is provided. Default is 10.
num_markers	An integer. Number of markers per chromosome. Ignored if chrom_maps is provided. Default is 1000.
сМ	A numeric. Length of each chromosome in centiMorgans. Ignored if chrom_maps is provided. Default is 100.

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bp An integer. Length of each chromosome in base pairs. Ignored if chrom_maps

is provided. Default is 1000000.

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

ates a random seed.

Value

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)
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.

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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.

Value

An external pointer to a Population object.

Examples

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

Get genotypes from a Population object getGenotypes

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

An external pointer to a Population object. pop

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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.

getGenotypesNaive

Get genotypes from a Population object (slow version)

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

```
getGenotypesNaive(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.

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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

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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.

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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.

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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.

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Examples

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

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