# Package 'BitBreedingSim'

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Type Package

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### Description

This function adds a Population object to an existing VCF object and creates a new VCF object that includes the population information.

### Usage

```
add_pop_to_VCF(vcf, pop)
```

### **Arguments**

vcf	A VCF object. This should be an object created using the relevant VCF func-
	tions in the package.
pop	A Population object. This should be an object created using the relevant Population functions in the package.

#### Value

A new VCF object that includes the added Population information.

```
# Assume 'vcf' is a valid VCF object and 'pop' is a valid Population object
new_vcf <- add_pop_to_VCF(vcf, pop)
summary(new_vcf) # Inspect the new VCF object</pre>
```

add\_trait\_A

add\_trait\_A

Add Trait with Additive Effects to BaseInfo

### Description

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

### Usage

```
add_trait_A(
  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
a	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 &lt;- c(3, 5, 10) marker &lt;- c(1, 2, 1000) loci &lt;- data.frame(chrom, marker)</pre>
num_loci	Optional. Number of loci (default is 1)

```
# Assume 'info' is a valid BaseInfo object
add_trait_A(info, "Trait1", mean=100.0, h2=0.6, sd=10.0, num_loci = 2)
trait <- get_trait(info, 1)
summary(trait)</pre>
```

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 $add\_trait\_AD$ 

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

```
add_trait_AD(
  info,
  name,
  mean,
  sd = NULL,
  h2 = NULL,
  H2 = 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 &lt;- c(3, 5, 10) marker &lt;- c(1, 2, 1000) loci &lt;- data.frame(chrom, marker)</pre>
num_loci	Optioal. Number of loci (default is 1)

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#### **Examples**

```
# Assume 'info' is a valid BaseInfo object
add_trait_AD(info, "Trait1", mean=100.0, h2=0.6, H2=0.7, sd=10.0, num_loci = 2)
trait <- get_trait(info, 1)
summary(trait)</pre>
```

check\_dataframe

Validate the structure and content of the data frame used for crossing information

#### **Description**

This function checks if the input data frame df meets the required structure for the cross\_by\_table function. It ensures that df is a valid data. frame, contains the required columns (mat, pat, num), and verifies that the num column has positive numeric values. If any of these checks fail, the function stops execution and displays an error message.

#### Usage

```
check_dataframe(df)
```

#### Arguments

df

A data frame. Must include the following columns: - mat: Names of the maternal parents for each cross. - pat: Names of the paternal parents for each cross. - num: Number of offspring to generate for each cross. Must contain positive numeric values.

#### Value

No return value. If the validation is successful, the function proceeds silently. If the validation fails, the function stops execution and displays an error message.

```
# Valid input example
df <- data.frame(mat = c("mat1", "mat2"), pat = c("pat1", "pat2"), num = c(1, 2))
check_dataframe(df) # Passes silently

# Invalid input example: Missing 'num' column
df_invalid <- data.frame(mat = c("mat1", "mat2"), pat = c("pat1", "pat2"))
check_dataframe(df_invalid) # Throws an error</pre>
```

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```
check_parent_existance
```

Check Parent Existence in Population

#### **Description**

This function checks whether the maternal and paternal names in the given cross table are present in the specified maternal and paternal populations.

#### Usage

```
check_parent_existance(df, mat_pop, pat_pop)
```

#### **Arguments**

df	A data.frame representing the cross table containing 'mat' (maternal names) and 'pat' (paternal names) columns.
mat_pop	An external pointer to the maternal Population object.
pat_pop	An external pointer to the paternal Population object.

#### Value

This function does not return a value. It outputs messages if any maternal or paternal names in the cross table are not found in the respective populations.

### **Examples**

```
# Assuming 'df', 'mat_pop', and 'pat_pop' are valid objects
check_parent_existance(df, mat_pop, pat_pop)
```

create\_base\_info

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.

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#### Usage

```
create_base_info(
  positions = NULL,
  num_markers = 1000,
  bp = 1e+08,
  chrom_maps = NULL,
  num_chroms = 10,
  cM = 100,
  seed = -1
)
```

#### **Arguments**

positions Optional. A list of numeric vectors, where each vector represents the marker

positions for a chromosome in base pairs. The number of elements in positions defines the number of chromosomes and overrides num\_chroms. If positions

is provided, num\_markers is ignored.

num\_markers Optional. An integer. Number of markers per chromosome. Ignored if positions

is provided. Default is 1000.

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

positions is provided. Default is 100000000.

chrom\_maps Optional. 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.). For a given cM value, the corresponding position can be interpolated or extrapolated, and vice versa. This effectively represents the mapping as a piecewise linear approximation of the chromosome's relationship between cM and base pair positions. If chrom\_maps

is provided, the parameter num\_chroms is ignored.

num\_chroms Optional. An integer. Number of chromosomes. Ignored if either chrom\_maps

or positions is provided. Default is 10.

cM Optional. A numeric. Length of each chromosome in centiMorgans. Ignored if

chrom\_maps is provided. Default is 100.

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

which generates a random seed.

#### Value

An external pointer to a BaseInfo object.

### **Examples**

```
# Create a BaseInfo object with a random seed
base_info_random <- create_base_info()
get_info(base_info_random)</pre>
```

# Create a BaseInfo object with a specific seed for reproducible results

```
base_info_reproducible <- create_base_info(seed = 123)
get_info(base_info_reproducible)

# Create marker positions manually
num_chr <- 10
position <- sapply(1:100, function(i) i * 10000000)
positions <- replicate(10, position, simplify = FALSE)

# Create a chromosome map with 100 cM and 100 Mbp
cM <- c(14.0, 27.9, 40.2, 48.3, 50.0, 51.7, 59.8, 72.1, 86.0, 100.0)
chr_position <- sapply(1:10, function(i) i * 100000000)
chrom_map <- data.frame(cM, chr_position)
chrom_maps <- replicate(10, chrom_map, simplify = FALSE)
names(chrom_maps) <- paste0("chr", 1:10)
info <- create_base_info(positions = positions, chrom_maps = chrom_maps, seed = 123)
get_info(info)</pre>
```

create\_HaploArray\_from\_pop

Create a 3-dimensional HaploArray from a Population object

#### **Description**

This function converts a Population object into a 3-dimensional array (HaploArray). The resulting HaploArray contains dimensions representing individuals, markers, and alleles (Maternal and Paternal). The HaploArray has the following dimensions:

dim1 Number of individuals.

dim2 Number of markers.

**dim3** Size is 2, representing Maternal and Paternal alleles.

The array also includes dimension names such as individual names, marker names, and allele labels (Maternal and Paternal).

#### Usage

```
create_HaploArray_from_pop(pop)
```

#### **Arguments**

рор

An external pointer to a Population object. This object must be properly initialized.

#### **Details**

Note: This function is the reverse of create\_pop\_from\_HaploArray, which creates a Population object from a HaploArray.

#### Value

A 3-dimensional array (HaploArray) containing the following dimensions:

**Individuals** Individual names as rows.

Markers Marker names as columns.

Alleles Maternal and Paternal alleles as the third dimension.

#### **Examples**

```
info <- create_base_info(num_chroms=2, num_markers=10, cM=100, bp=1e6, seed=2)
add_trait_A(info, "Trait1", mean=100.0, h2=0.6, sd=10.0, num_loci = 2)
trait <- get_trait(info, 1)

haploArray <- array(
    data = rbinom(n = 120, size = 1, prob = 0.3),
    dim = c(3, 20, 2),
    dimnames = list(
        paste0("Ind_", 1:3),
        paste0("Mrk_", 1:20),
        c("Maternal", "Paternal")
    )
    )
    pop <- create_pop_from_HaploArray(haploArray, info)
    prog <- cross_randomly(pop, pop, num_inds = 5, name_base = "prog_")
    haploArrayProg <- create_HaploArray_from_pop(prog)</pre>
```

create\_info\_pop\_from\_VCF

Create BaseInfo and Population from a VCF file

#### **Description**

This function takes a VCF object and a seed value, and returns both a Population object and its associated BaseInfo object. It reads the input VCF and initializes the data accordingly. The seed value is used to initialize the pseudo-random number generator for the BaseInfo object. If the seed is set to -1, an appropriate value will be automatically chosen.

#### **Usage**

```
create_info_pop_from_VCF(vcf, seed = -1)
```

#### **Arguments**

vcf An external pointer to a VCF object.

An integer. The seed value for initializing the BaseInfo object's pseudo-random number generator. Defaults to -1, which automatically selects a suitable seed.

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#### Value

A list containing two elements:

info An external pointer to a BaseInfo object.

**pop** An external pointer to a Population object.

#### **Examples**

```
# Assuming 'vcf_file' is a valid VCF file
vcf <- read_VCF(vcf_file)
result <- create_info_pop_from_VCF(vcf, seed = 42)
summary(result$info)
summary(result$pop)</pre>
```

create\_origins

Create origins for a Population object

#### **Description**

Create origins for a Population object

#### Usage

```
create_origins(info, names = NULL, num_inds = NULL, name_base = NULL)
```

### Arguments

info An external pointer to a BaseInfo object.

num\_inds An integer. The number of individuals.

name\_base A string. The base name for individuals.

#### Value

An external pointer to a Population object.

```
# Example 1: Specify individual names directly
pop <- create_origins(info, names = c("p1", "p2"))
get_individual_names(pop)

# Example 2: Generate individual names using name_base and num_inds
pop2 <- create_origins(info, num_inds = 2, name_base = "q")
get_individual_names(pop2)</pre>
```

```
create_pop_from_HaploArray
```

Create a Population object from a HaploArray

#### **Description**

This function takes a 3-dimensional array (HaploArray) and a BaseInfo object to create a Population object. The HaploArray should have dimensions representing individuals, markers, and alleles (Maternal and Paternal). Allele values must be binary (0 or 1). A value of 0 indicates that the allele matches the reference genome, while a value of 1 indicates the alternative allele. The function validates the input data and initializes the Population object accordingly.

#### Usage

```
create_pop_from_HaploArray(haploArray, info)
```

#### **Arguments**

haploArray A 3-dimensional array where:

dim1 Number of individuals.

dim2 Number of markers.

**dim3** Size must be 2, representing Maternal and Paternal alleles.

Allele values must be binary (0 or 1), based on the reference genome.

info An external pointer to a BaseInfo object.

#### Value

An external pointer to a Population object.

```
# Create a HaploArray
haploArray <- array(
  data = rbinom(n = 30, size = 1, prob = 0.3),
  dim = c(3, 5, 2),
  dimnames = list(
    paste0("Ind_", 1:3),
    paste0("Mrk_", 1:5),
    c("Maternal", "Paternal")
)

# Assuming 'info' is a valid BaseInfo object
pop <- create_pop_from_HaploArray(haploArray, info)
summary(pop)</pre>
```

12 cross\_by\_table

c. cost_sy_table cross populations according to a label	cross_by_table	Cross populations according to a table	
---	----------------	--	--

### Description

This function performs crossing of individuals from two Population objects (maternal and paternal parents) based on specified crossing information provided in a table. Each row of the table defines a specific cross, including the number of offspring to generate for that cross. You can specify either name\_base or provide a complete names vector for the offspring.

### Usage

```
cross_by_table(
   df,
   mat_pop,
   pat_pop,
   names = NULL,
   name_base = NULL,
   num_threads = 0
)
```

### Arguments

df	A data frame containing crossing information. It should include the following columns:
	mat Names of the maternal parents for each cross.
	pat Names of the paternal parents for each cross.
	num Number of offspring to generate for each cross.
mat_pop	A Population object representing the maternal parents. This should be an object created using the relevant Population functions in the package.
pat_pop	A Population object representing the paternal parents. This should be an object created using the relevant Population functions in the package.
names	Optional. A character vector containing specific names for individuals. If provided, the length of names should match the total number of offspring specified in df.
name_base	Optional. A character string used as the base name for generating individual names. For example, if name_base = "prog_", the generated names will follow the format c("prog_1", "prog_2",).
num_threads	Optional. A positive integer representing the number of threads to use. If not specified or set to 0, the maximum number of available threads is used.

### Value

A Population object with offspring individuals.

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#### **Examples**

```
# Example using name_base
df <- data.frame(mat = c("mat1", "mat2"), pat = c("pat1", "pat2"), num = c(1, 2))
new_population <- cross_by_table(df, mat_pop, pat_pop, name_base = "prog_")
# Example using predefined names
names_vector <- c("child1", "child2", "child3")
new_population <- cross_by_table(df, mat_pop, pat_pop, names = names_vector)
# Summary of the new Population object
summary(new_population)</pre>
```

cross\_randomly

Cross Population randomly

#### **Description**

This function performs random crossing of individuals from two Population objects (maternal and paternal parents) to generate a new Population object. For each offspring, a parent is randomly selected from each Population (one from the maternal Population and one from the paternal Population). You can specify either name\_base and num\_inds or provide a complete names vector for the offspring.

#### Usage

```
cross_randomly(
  mat_pop,
  pat_pop,
  names = NULL,
  num_inds = NULL,
  name_base = NULL,
  num_threads = 0
)
```

#### **Arguments**

mat_pop	A Population object representing the maternal parents. This should be an object created using the relevant Population functions in the package.
pat_pop	A Population object representing the paternal parents. This should be an object created using the relevant Population functions in the package.
names	Optional. A character vector containing specific names for individuals. If provided, the length of names determines the number of offspring.
num_inds	Optional. A positive integer representing the number of offspring. Required if name_base is specified.

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name\_base Optional. A character string used as the base name for generating individ-

ual names. For example, if name\_base = "p" and num\_inds = 3, the generated names will be c("p1", "p2", "p3"). This parameter must be used in combina-

tion with num\_inds.

num\_threads Optional. A positive integer representing the number of threads to use. If not

specified or set to 0, the maximum number of available threads is used.

#### Value

A Population object with offspring individuals.

### **Examples**

```
# Example using name_base and num_inds
new_population <- cross_randomly(mothers, fathers, num_inds = 100, name_base = "offspring_")
# Example using predefined names
names_vector <- c("child1", "child2", "child3")
new_population <- cross_randomly(mothers, fathers, names = names_vector)
# Summary of the new Population object
summary(new_population)</pre>
```

extract\_map\_data

Extract chromosome and position data from a Map object

#### **Description**

This function retrieves detailed data from a Map object, including chromosome names and marker positions, and returns the data as a data.frame for easier manipulation.

#### Usage

```
extract_map_data(map)
```

#### **Arguments**

map

An external pointer to a Map object. This object must be properly initialized and created using appropriate functions.

#### Value

A data.frame containing the following columns:

**chrom** A character vector containing the names of chromosomes.

**position** An integer vector specifying the positions of markers along the chromosomes.

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#### **Examples**

```
# Assuming 'map' is a valid Map object
map_data <- extract_map_data(map)
head(map_data) # Display the first few rows of the data.frame</pre>
```

generate_names	Generate names for individuals based on a base name and number of
	individuals

### Description

This function creates a vector of names for individuals when names is not provided. It uses a base name (name\_base) and a specified number of individuals (num\_inds) to generate names in the format name\_base1, name\_base2, ..., up to the total number.

#### Usage

```
generate_names(name_base, num_inds)
```

### **Arguments**

name\_base A character string used as the base for generating names. For example, name\_base

= "p" will generate names like "p1", "p2", ....

num\_inds A positive integer indicating the number of names to generate.

### Value

A character vector of generated names.

#### **Examples**

```
# Generate 3 names with base "p"
generated_names <- generate_names("p", 3)
print(generated_names) # Output: "p1", "p2", "p3"</pre>
```

get\_genotypes 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 individuals and columns correspond to markers.

#### Usage

```
get_genotypes(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 individuals and columns are markers.

### **Examples**

```
info <- create_base_info()
pop <- create_origins(info, num_inds = 2, names = c("p1", "p2"))
geno <- get_genotypes(pop)
geno[, 1:5]</pre>
```

get\_individual\_names

Get name data from a Population object

#### **Description**

Get name data from a Population object

### Usage

```
get_individual_names(pop)
```

#### **Arguments**

pop

An external pointer to a Population object.

### Value

A data.frame containing the following columns:

name Names from the Population objectmat Maternal names from the Population objectpat Paternal names from the Population object

get\_info

#### **Examples**

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

get\_info

Retrieve key values from a BaseInfo object

#### **Description**

This function extracts essential information from a BaseInfo object. A BaseInfo object stores metadata related to genetic simulations, such as the number of chromosomes, traits, and markers. This function is useful for summarizing or verifying the content of a BaseInfo object during analysis.

#### Usage

```
get_info(info)
```

#### **Arguments**

info

An external pointer to a BaseInfo object. This object must be properly initialized and created using the appropriate BaseInfo constructor or relevant functions.

#### Value

A list containing the following elements:

**num\_chroms** An integer representing the total number of chromosomes contained in the BaseInfo object.

num\_traits An integer indicating the number of traits managed by the BaseInfo object.

**num\_markers** An integer specifying the total number of markers tracked in the BaseInfo object.

```
# Assume 'info' is a valid BaseInfo object
values <- get_info(info)
print(values$num_chroms) # Print the number of chromosomes
print(values$num_traits) # Print the number of traits
print(values$num_markers) # Print the number of markers</pre>
```

get\_map

Retrieve Genetic Map Information

#### Description

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

#### Usage

```
get_map(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 <- get_map(info)</pre>
  print(map)
## End(Not run)
```

get\_phased\_genotypes 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 individuals.

#### Usage

```
get_phased_genotypes(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 individuals and columns are markers.

#### **Examples**

```
info <- create_base_info()
pop <- create_origins(info, num_inds = 2, names = c("p1", "p2"))
geno <- get_phased_genotypes(pop)
geno[, 1:5]</pre>
```

```
get_phased_int_genotypes
```

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 individuals and columns correspond to markers. Each individual has two rows: the first row represents the maternal allele and the second row represents the paternal allele.

#### Usage

```
get_phased_int_genotypes(pop)
```

### **Arguments**

pop

An external pointer to a Population object.

#### **Details**

Genotype is represented as integers: 0 or 1

### Value

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

```
info <- create_base_info()
pop <- create_origins(info, num_inds = 2, names = c("p1", "p2"))
geno <- get_phased_int_genotypes(pop)
geno[, 1:5]</pre>
```

get\_pop\_info

get	nhenotypes

Get phenotypes from a Population object

### **Description**

Get phenotypes from a Population object

### Usage

```
get_phenotypes(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 <- get_phenotypes(pop, 1)
print(phenotypes)</pre>
```

get\_pop\_info

Get information for a Population object

### Description

This function retrieves detailed information about a Population object, including the number of individuals, the number of chromosomes, and the number of markers in the population.

### Usage

```
get_pop_info(pop)
```

#### **Arguments**

pop

A Population object. This should be an object created using the relevant Population functions in the package.

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#### Value

A list containing: - num\_individuals: The number of individuals in the population. - num\_chromosomes: The number of chromosomes in the population. - num\_markers: The number of markers in the population.

### **Examples**

```
# Create base information
info <- create_base_info()
# Create a population with 2 individuals
pop <- create_origins(info, num_inds = 2, names = c("p1", "p2"))
# Retrieve population information
get_pop_info(pop)</pre>
```

get\_trait

Get a trait from a BaseInfo object

### Description

Get a trait from a BaseInfo object

#### Usage

```
get_trait(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.

```
# Assume 'info' is a valid BaseInfo object
trait <- get_trait(info, 1)
summary(trait)</pre>
```

join\_pops

get\_VCF\_info

Get information from a VCF object

#### Description

This function retrieves detailed information about a VCF object, including the number of individuals and the number of markers contained within the VCF.

### Usage

```
get_VCF_info(vcf)
```

#### **Arguments**

vcf

A VCF object. This should be an object created using the relevant VCF functions in the package.

#### Value

A list containing: - num\_individuals: The number of individuals in the VCF object. - num\_markers: The number of markers in the VCF object.

#### **Examples**

```
# Assume 'vcf' is a valid VCF object
vcf_info <- get_VCF_info(vcf)
print(vcf_info) # Displays the number of individuals and markers</pre>
```

join\_pops

Join multiple Population objects

### **Description**

Join multiple Population objects

#### Usage

```
join_pops(...)
```

#### **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 <- join_pops(pop1, pop2, pop3)
print(combined_pop)</pre>
```

modify\_trait\_parameters

Modify Trait Parameters

#### **Description**

This function allows modifying the parameters of a specified trait in a BaseInfo object. At least one of h2, a, or additive\_multiplier must be provided. The function ensures that only valid parameter combinations are specified.

#### Usage

```
modify_trait_parameters(
  info,
  i,
  h2 = NULL,
  a = NULL,
  additive_multiplier = NULL)
```

#### **Arguments**

info A BaseInfo object that contains the trait information.

An integer index of the trait to be modified.

(Optional) Narrow-sense heritability. If only h2 is specified, the additive effect remains unchanged, and the error variance is adjusted to match the specified h2.

(Optional) A numeric vector specifying the additive effects for each QTL. If a is provided with h2, the error variance is adjusted to match the specified h2, while the additive effects remain as specified.

additive\_multiplier

(Optional) A scalar to scale the current additive effect. If provided with h2, the error variance is adjusted to match the specified h2, while the additive effects remain as specified.

**Note:** Only one of a or additive\_multiplier can be specified at the same time.

- If only h2 is specified, it adjusts the error variance to match the specified h2 while keeping the additive effects unchanged.
- If a or additive\_multiplier is specified alongside h2, both the additive effects and the error variance are adjusted.
- If only a or additive\_multiplier is specified, only the additive effects are modified.

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#### Value

No return value. The function modifies the parameters of the specified trait directly.

print\_trait

Print detailed information about a Trait object

### Description

This function prints the detailed characteristics of a Trait object in a human-readable format. A Trait object represents a characteristic or feature in genetic simulations, and this function displays its metadata such as its name, type, mean, standard deviation, heritability (h2), associated loci, additive effects, and dominant effects (if applicable).

#### Usage

```
print_trait(trait, ...)
```

#### **Arguments**

trait

A Trait object containing the following elements:

**name** The name of the trait as a character string.

**type** The type of the trait (e.g., "quantitative" or "binary").

mean The mean value of the trait as a numeric.

sd The standard deviation of the trait as a numeric.

**h2** The narrow-sense heritability of the trait as a numeric (between 0 and 1).

**H2** (Optional) The broad-sense heritability of the trait as a numeric (if applicable)

**hasdominants** A logical value indicating whether the trait includes dominant effects.

**loci** A list of loci associated with the trait, formatted as coordinate pairs.

additives A numeric vector representing the additive effects of the trait.

**dominants** (Optional) A numeric vector representing the dominant effects of the trait (if applicable).

... Additional arguments (not currently used).

#### **Details**

The loci are formatted as coordinate pairs, representing their positions. Additive and dominant effects are printed as space-separated values for clarity.

#### Value

Prints the detailed information of the Trait object directly to the console.

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### **Examples**

```
info <- create_base_info(num_chroms=2, num_markers=10, cM=100, bp=1e6, seed=2)
addTraitA(info, "Trait1", mean=100.0, h2=0.6, sd=10.0, num_loci = 2)
trait <- getTrait(info, 1)
print_trait(trait)

# Alternatively, use the generic print() function
print(trait)</pre>
```

read\_VCF

Read a VCF file and return a VCF object

#### **Description**

This function reads a VCF file from the specified filename and returns a VCF object as an external pointer. If the input is not a valid character string, a message is displayed and NULL is returned.

#### Usage

```
read_VCF(filename)
```

#### **Arguments**

filename

A character string specifying the path to the VCF file.

#### Value

An external pointer to a VCF object, or NULL if an error occurs.

```
# Assuming 'example.vcf' is a valid VCF file
vcf <- read_VCF("example.vcf")
if (is.null(vcf)) {
  cat("Failed to read the VCF file.\n")
}</pre>
```

set\_individual\_names

select\_pop

Select individuals from a Population object

#### **Description**

Select individuals from a Population object

#### Usage

```
select_pop(pop, indices)
```

#### **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.

#### **Examples**

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

### Description

This function assigns names to individuals in a Population object. The names argument must be a character vector, and pop must be an object of the "Population" class.

### Usage

```
set_individual_names(names, pop)
```

#### **Arguments**

names A character vector containing the names of individuals to assign.

pop A Population object where the individual names will be set.

#### Value

None. The function modifies the Population object directly.

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#### **Examples**

```
# Create a Population object and set individual names
info <- create_base_info()
pop <- create_origins(info, num_inds = 2, name_base = "ind")
set_individual_names(c("sample1", "sample2"), pop)

# Access the updated individual names
get_individual_names(pop)</pre>
```

summary\_map

Summarize the details of a Map object

### **Description**

This function extracts and displays key information from a Map object. A Map object represents the genetic map used in simulations, containing details about chromosomes and points (e.g., markers or loci) along the map. The summary provides an overview of the structure and size of the map.

#### Usage

```
summary_map(map, ...)
```

#### **Arguments**

An external pointer to a Map object. This object must be properly initialized and created using the appropriate functions.

... Additional arguments (not currently used).

### Value

Prints a formatted summary of the Map object directly to the console.

```
# Assuming 'map' is a valid Map object
summary_map(map)
```

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summary\_population

Summarize the details of a Population object

### **Description**

This function extracts and displays key information from a Population object. A Population object represents a collection of individuals, chromosomes, and genetic markers used in genetic simulations. The summary includes the number of individuals, chromosomes, and markers in the population, providing an overview of its structure and size.

#### Usage

```
summary_population(pop, ...)
```

#### **Arguments**

pop

An external pointer to a Population object. This object must be properly initial-

ized and created using the appropriate functions or constructors.

. . .

Additional arguments (not currently used).

#### Value

Prints a formatted summary of the Population object directly to the console.

#### **Examples**

```
# Assuming 'pop' is a valid Population object
summary_population(pop)
```

summary\_trait

Summarize the details of a Trait object

#### **Description**

This function provides a formatted summary of a Trait object. A Trait object represents a characteristic or feature in genetic simulations, and contains metadata such as its name, type, mean, standard deviation, heritability (h2), and information about associated loci. If the trait includes dominant effects, the broad-sense heritability (H2) is also displayed.

### Usage

```
summary_trait(trait)
```

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### **Arguments**

trait

A Trait object containing the following elements:

**name** The name of the trait as a character string.

**type** The type of the trait (e.g., "Additive Effect Only").

mean The mean value of the trait as a numeric.

sd The standard deviation of the trait as a numeric.

**h2** The narrow-sense heritability of the trait as a numeric (between 0 and 1).

**H2** (Optional) The broad-sense heritability of the trait as a numeric (if applicable)

hasdominants A logical value indicating whether the trait includes dominant

**loci** A list of loci associated with the trait.

#### Value

Prints a summary of the Trait object directly to the console.

#### **Examples**

```
info <- create_base_info(num_chroms=2, num_markers=10, cM=100, bp=1e6, seed=2)
addTraitA(info, "Trait1", mean=100.0, h2=0.6, sd=10.0, num_loci = 2)
trait <- getTrait(info, 1)
summary_trait(trait)

# Using the generic summary() function
summary(trait)</pre>
```

write\_VCF

Write VCF or Population object to a VCF file

#### **Description**

This function writes either a VCF object or a Population object to a VCF file. If the input is a VCF object, it directly writes the VCF data to the specified file. If the input is a Population object, it generates VCF data from the Population and writes it to the file.

#### Usage

```
write_VCF(obj, filename)
```

#### **Arguments**

obj An object of class VCF or Population. The data to write to the VCF file.

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

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```
# Example 1: Write a Population object to a VCF file
pop <- create_origins(info, num_inds = 2, names = c("p1", "p2"))
write_VCF(pop, "population_output.vcf")

# Example 2: Write a VCF object to a VCF file
vcf <- read_VCF("input.vcf")
write_VCF(vcf, "vcf_output.vcf")</pre>
```

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