Package 'malan'

November 28, 2023

```
Type Package
Title MAle Lineage ANalysis
Version 1.0.3
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Description MAle Lineage ANalysis by simulating
     genealogies backwards and imposing short tandem repeats (STR) mutations forwards.
     Intended for forensic Y chromosomal STR (Y-STR) haplotype analyses.
     Numerous analyses are possible, e.g. number of matches and meiotic distance to matches.
     Refer to papers mentioned in citation(`malan") (DOI's:
     <doi:10.1371/journal.pgen.1007028>,
     <doi:10.21105/joss.00684> and
     <doi:10.1016/j.fsigen.2018.10.004>).
License GPL-2 | file LICENSE
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Description

Simulating genealogies backwards and imposing STR mutations forwards.

Details

See vignettes and manual for documentation.

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References

Andersen MM, Balding DJ (2017) How convincing is a matching Y-chromosome profile? PLoS Genet 13(11): e1007028. doi:10.1371/journal.pgen.1007028.

See Also

Useful links:

• Report bugs at https://github.com/mikldk/malan/issues

analyse_mixture_result 5

```
analyse_mixture_result
```

Analyse mixture results

Description

Calculate LR-like quantities by haplotype counts.

Usage

```
analyse_mixture_result(
  mix_res,
  unique_haps_in_mixture,
  unique_haps_in_mixture_counts
)
```

Arguments

Details

NOTE: Only takes up to 9 contributors!

Value

A list with numeric quantities

```
analyse_mixture_results
```

Analyse mixture results in a vectorised fashion

Description

Refer to analyse_mixture_result() for details. Essentially, analyse_mixture_result() is run on each element of mixture_results.

Usage

```
analyse_mixture_results(
  mixture_results,
  unique_haps_in_mixture_list,
  unique_haps_in_mixture_counts_list
)
```

Arguments

```
mixture_results
```

List of n mixture results from mixture_info_by_individuals_2pers(), mixture_info_by_individuals_ture_info_by_individuals_5pers()
mixture_info_by_individuals_5pers()

unique_haps_in_mixture_list

List of n included unique haplotypes, one for each element in mix_res

unique_haps_in_mixture_counts_list

List of n population counts of the included unique haplotypes

Details

NOTE: Only takes up to 9 contributors!

Value

A list with lists of numeric quantities

```
as_tbl_graph.malan_pedigreelist

Get tidy graph object
```

Description

```
Get tidy graph object tbl_graph(), e.g. to plot with ggraph().
```

Usage

```
## S3 method for class 'malan_pedigreelist'
as_tbl_graph(x, ...)
```

Arguments

```
x malan_pedigreelist
... Ignored
```

Value

```
tbl_graph() object
```

brothers_matching 7

brothers_matching

Number of brothers with matching haplotype

Description

Get individual's number of brothers that matches individual's haplotype

Usage

```
brothers_matching(individual)
```

Arguments

individual individual

Value

Number of brothers that matches individual's haplotype

```
build_haplotype_hashmap
```

Build hashmap of haplotype to individuals

Description

Makes it possible to find all individuals' pid with a certain haplotype. Must be used with e.g. get_matching_pids_from_hashmap().

Usage

```
build_haplotype_hashmap(individuals, progress = TRUE)
```

Arguments

individuals List of individuals to build hashmap of

progress Show progress?

Value

External pointer to hashmap with haplotype as keys and vector of individuals' pid as value

See Also

```
get_matching_pids_from_hashmap().
```

build_pedigrees

Build pedigrees from (individuals in) a population.

Description

In a newly simulated population, each individual only knows its father and children. Using this information, this function builds pedigrees. This makes it easier to e.g. population haplotypes, find path between two individuals (if they are not in the same pedigree, they are not connected).

Usage

```
build_pedigrees(population, progress = TRUE)
```

Arguments

population Population generated by sample_geneology() or sample_geneology_varying_size().

progress Show progress.

Value

An object with class malan_pedigreelist (an internal list of external pointers to pedigrees).

See Also

```
sample_geneology() and sample_geneology_varying_size() for simulating populations.
```

Examples

```
sim <- sample_geneology(100, 10)
str(sim, 1)
sim$population
peds <- build_pedigrees(sim$population)
peds</pre>
```

```
{\tt calc\_autosomal\_genotype\_conditional\_cumdist}
```

Calculate conditional genotype cumulative probabilities with theta

Description

Calculate conditional genotype cumulative probabilities with theta

Usage

```
calc_autosomal_genotype_conditional_cumdist(allele_dist, theta)
```

Arguments

allele_dist Allele distribution (probabilities) – gets normalised theta Theta correction between 0 and 1 (both included)

Value

Matrix: row i: conditional cumulative distribution of alleles given allele i

calc_autosomal_genotype_probs

Calculate genotype probabilities with theta

Description

Calculate genotype probabilities with theta

Usage

```
calc_autosomal_genotype_probs(allele_dist, theta)
```

Arguments

allele_dist Allele distribution (probabilities) – gets normalised theta Theta correction between 0 and 1 (both included)

construct_M

Construct M matrix

Description

Construct M matrix

Usage

```
construct_M(meioses, mu_dw, mu_up)
```

Arguments

meioses number of meioses separating the two individuals

mu_dw mutation rate for 1-step down-mutation mu_up mutation rate for 1-step up-mutation count_brothers

Number of brothers

Description

Get individual's number of brothers

Usage

```
count_brothers(individual)
```

Arguments

individual

individual

Value

Number of brothers

See Also

```
get_brothers()
```

count_haplotype_near_matches_individuals

Count near haplotype matches in list of individuals

Description

Counts the number of types close to haplotype in individuals.

Usage

```
count_haplotype_near_matches_individuals(individuals, haplotype, max_dist)
```

Arguments

individuals List of individuals to count occurrences in.

haplotype Haplotype to count near-matches occurrences of.

max_dist Maximum distance (0 = match, 1 = 1 STR allele difference, ...)

Value

Number of times that a haplotype within a radius of max_dist of haplotype occurred amongst individuals.

See Also

 $count_haplotype_occurrences_individuals(), pedigree_haplotype_matches_in_pedigree_meiosis_L1_dists()$

Description

Counts the number of types haplotype appears in individuals.

Usage

```
count_haplotype_occurrences_individuals(individuals, haplotype)
```

Arguments

individuals List of individuals to count occurrences in.

haplotype Haplotype to count occurrences of.

Value

 $Number\ of\ times\ that\ haplotype\ occurred\ amongst\ individuals.$

See Also

```
pedigree_haplotype_matches_in_pedigree_meiosis_L1_dists(), count_haplotype_near_matches_individuals(
```

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes(peds, 2, c(0, 0))
count_haplotype_occurrences_individuals(sim$end_generation_individuals, c(0, 0))</pre>
```

12 count_uncles

```
count_haplotype_occurrences_pedigree

Count haplotypes occurrences in pedigree
```

Description

Counts the number of types haplotype appears in pedigree.

Usage

```
count_haplotype_occurrences_pedigree(
  pedigree,
  haplotype,
  generation_upper_bound_in_result = -1L
)
```

Arguments

pedigree Pedigree to count occurrences in.
haplotype Haplotype to count occurrences of.

generation_upper_bound_in_result

Only consider matches in generation 0, 1, ... generation_upper_bound_in_result. -1 means disabled, consider all generations. End generation is generation 0. Second last generation is 1. And so on.

Value

Number of times that haplotype occurred in pedigree.

See Also

```
pedigree_haplotype_matches_in_pedigree_meiosis_L1_dists().
```

count_uncles

Number of uncles

Description

Get individual's number of uncles

Usage

```
count_uncles(individual)
```

Arguments

individual individual

Value

Number of uncles

See Also

```
get_uncles()
```

delete_haplotypeids_hashmap

Delete haplotype hashmap

Description

Delete hashmap made by build_haplotype_hashmap().

Usage

```
delete_haplotypeids_hashmap(hashmap)
```

Arguments

hashmap

Hashmap made by build_haplotype_hashmap()

See Also

```
get_matching_pids_from_hashmap() and build_haplotype_hashmap().
```

estimate_autotheta_1subpop_genotypes

Estimate autosomal theta from genotypes

Description

Estimate autosomal theta for one subpopulation given a sample of genotypes.

Usage

```
estimate_autotheta_1subpop_genotypes(genotypes, return_estimation_info = FALSE)
```

Arguments

genotypes Matrix of genotypes: two columns (allele1 and allele2) and a row per individual return_estimation_info

Whether to return the quantities used to estimate theta

Details

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Value

List:

- theta
 - estimate: Vector of length 1 containing estimate of theta or NA if it could not be estimated
 - error: true if an error happened, false otherwise
 - details: contains description if an error happened
 - estimation_info: If return_estimation_info = true: a list with information used to estimate theta. Else NULL.

estimate_autotheta_1subpop_individuals

Estimate autosomal theta from individuals

Description

Estimate autosomal theta for one subpopulation given a list of individuals.

Usage

```
estimate_autotheta_1subpop_individuals(
  individuals,
  return_estimation_info = FALSE
)
```

Arguments

```
individuals Individuals to get haplotypes for. return_estimation_info
```

Whether to return the quantities used to estimate theta

Details

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Value

List:

- theta
 - estimate: Vector of length 1 containing estimate of theta or NA if it could not be estimated
 - error: true if an error happened, false otherwise
 - details: contains description if an error happened
 - estimation_info: If return_estimation_info = true: a list with information used to estimate theta. Else NULL.

estimate_autotheta_subpops_genotypes

Estimate autosomal F, theta, and f from subpopulations of genotypes

Description

Estimates autosomal F, theta, and f for a number of subpopulations given a list of genotypes.

Usage

```
estimate_autotheta_subpops_genotypes(subpops, subpops_sizes)
```

Arguments

subpops List of subpopulations, each a list of individuals

subpops_sizes Size of each subpopulation

Details

 $Assumes \ that \ pedigrees_all_populate_autosomal() \ was \ used \ first \ to \ populate \ autosomal \ genotypes.$

Based on Bruce S Weir, Genetic Data Analysis 2, 1996. (GDA2).

Value

Estimates of autosomal F, theta, and f as well as additional information

estimate_autotheta_subpops_individuals

Estimate autosomal F, theta, and f from subpopulations of individuals

Description

Estimates autosomal F, theta, and f for a number of subpopulations given a list of individuals.

Usage

```
estimate_autotheta_subpops_individuals(subpops, subpops_sizes)
```

Arguments

subpops List of subpopulations, each a list of individuals

subpops_sizes Size of each subpopulation

Details

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Based on Bruce S Weir, Genetic Data Analysis 2, 1996. (GDA2).

Value

Estimates of autosomal F, theta, and f as well as additional information

```
estimate_autotheta_subpops_pids
```

Estimate autosomal F, theta, and f from subpopulations of individual

Description

Estimates autosomal F, theta, and f for a number of subpopulations given a list of pids (individual ids).

Usage

```
estimate_autotheta_subpops_pids(population, subpops, subpops_sizes)
```

Arguments

population Population obtain from simulation

subpops List of individual pids subpops_sizes Size of each subpopulation

Details

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Based on Bruce S Weir, Genetic Data Analysis 2, 1996. (GDA2).

Value

Estimates of autosomal F, theta, and f as well as additional information

estimate_autotheta_subpops_unweighted_genotypes

Unweighted estimate of autosomal theta from subpopulations of genotypes

Description

Estimates unweighted autosomal theta for a number of subpopulations given a list of subpopulations of genotypes.

Usage

estimate_autotheta_subpops_unweighted_genotypes(subpops, assume_HWE)

Arguments

subpops List of individual genotypes

assume_HWE if the alleles themselves are used instead of genotypes

Details

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Based on Weir and Goudet, Genetics 2017: http://www.genetics.org/content/early/2017/05/26/genetics.116.198424

Value

Estimate of autosomal theta

18 father_matches

estimate_autotheta_subpops_unweighted_pids

Unweighted estimate of autosomal theta from subpopulations of individual ids

Description

Estimates unweighted autosomal theta for a number of subpopulations given a list of pids (individual ids).

Usage

estimate_autotheta_subpops_unweighted_pids(population, subpops, assume_HWE)

Arguments

population Population obtain from simulation

subpops List of individual pids

assume_HWE if the alleles themselves are used instead of genotypes

Details

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Based on Weir and Goudet, Genetics 2017: http://www.genetics.org/content/early/2017/05/26/genetics.116.198424

Value

Estimate of autosomal theta

father_matches Father matches

Description

Does the father have the same profile as individual?

Usage

father_matches(individual)

Arguments

individual individual

Value

Whether father has the same profile as individual or not

from_igraph 19

from_igraph

Convert igraph to population

Description

Convert igraph to population

Usage

```
from_igraph(x, ...)
```

Arguments

x igraph, must be a forest of directed trees with unique positive integer names (as they will be pid's)

.. Ignored

Value

A population

Examples

```
g <- igraph::graph_from_literal( 2 +- 1 -+ 3, 4 -+ 5 )
plot(g)
pop <- from_igraph(g)
peds <- build_pedigrees(pop, progress = FALSE)
plot(peds)
infer_generations(peds)
get_generation(get_individual(pop, 1))
get_generation(get_individual(pop, 2))
get_generation(get_individual(pop, 3))
get_generation(get_individual(pop, 4))
get_generation(get_individual(pop, 5))</pre>
```

from_igraph_rcpp

Generate paternal brothers population

Description

Generate paternal brothers population

Usage

```
from_igraph_rcpp(vertices, edges)
```

Arguments

vertices vector of vertices edges matrix with edges

Value

An external pointer to the population.

```
generate_get_founder_haplotype_db
```

Generate a function to simulate pedigree founder haplotype based on a haplotype databasep

Description

Generate a function to simulate pedigree founder haplotype based on a haplotype databasep

Usage

```
generate_get_founder_haplotype_db(db)
```

Arguments

db

data frame or matrix with haplotypes from which the founder is randomly simulated

```
generate_get_founder_haplotype_ladder
```

Generate a function to simulate pedigree founder haplotype based on ladder information

Description

Generate a function to simulate pedigree founder haplotype based on ladder information

Usage

```
generate_get_founder_haplotype_ladder(ladder_min, ladder_max)
```

Arguments

ladder_min vector of minimum alleles; ladder_min[i] is the minimum allele at locus i vector of minimum alleles; ladder_max[i] is the maximum allele at locus i

```
get_allele_counts_genotypes
```

Get autosomal allele counts from subpopulations of genotypes

Description

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Usage

```
get_allele_counts_genotypes(subpops)
```

Arguments

subpops

List of individual genotypes

Value

Matrix with allele counts

```
get_allele_counts_pids
```

Get autosomal allele counts from subpopulations given by pids

Description

Assumes that pedigrees_all_populate_autosomal() was used first to populate autosomal genotypes.

Usage

```
get_allele_counts_pids(population, subpops)
```

Arguments

population Population obtain from simulation

subpops List of individual pids

Value

Matrix with allele counts

22 get_children

get_brothers

Get brothers

Description

Get individual's brothers

Usage

```
get_brothers(individual)
```

Arguments

individual

individual

Value

List with brothers

See Also

```
get_father(), get_uncles(), get_children(), get_cousins()
```

get_children

Get children

Description

Get individual's children

Usage

```
get_children(individual)
```

Arguments

individual

individual

Value

List with children

See Also

```
get_father(), get_brothers(), get_uncles(), get_cousins()
```

get_cousins 23

get_cousins

Get cousins

Description

Get individual's cousins

Usage

```
get_cousins(individual)
```

Arguments

individual

individual

Value

List with cousins

See Also

```
get_brothers(), get_uncles(), get_children()
```

get_family_info

Get individual's family information

Description

Get individual's family information

Usage

```
get_family_info(individual)
```

Arguments

individual

individual

Value

List with family information

24 get_generation

get_father

Get father

Description

Get individual's father

Usage

```
get_father(individual)
```

Arguments

individual individual

Value

Father

See Also

```
get_brothers(), get_uncles(), get_children(), get_cousins()
```

get_generation

Get individual's generation number

Description

Note that generation 0 is final, end generation. 1 is second last generation etc.

Usage

```
get_generation(individual)
```

Arguments

individual

Individual

Value

generation

Examples

```
sim <- sample_geneology(100, 10)
indv <- get_individual(sim$population, 1)
get_generation(indv)</pre>
```

get_haplotype 25

get_haplotype

Get haplotype from an individual

Description

Requires that haplotypes are first populated, e.g. with pedigrees_all_populate_haplotypes(), pedigrees_all_populate_haplotypes_custom_founders(), or pedigrees_all_populate_haplotypes_ladder_bounders().

Usage

```
get_haplotype(individual)
```

Arguments

individual Individual to get haplotypes for.

Value

Haplotype for individual.

See Also

```
get_haplotypes_individuals() and get_haplotypes_pids().
```

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes(peds, 2, c(1, 1))
get_haplotype(sim$end_generation_individuals[[1]])</pre>
```

```
get_haplotypes_individuals
```

Get haplotype matrix from list of individuals

Description

Requires that haplotypes are first populated, e.g. with pedigrees_all_populate_haplotypes(), pedigrees_all_populate_haplotypes_custom_founders(), or pedigrees_all_populate_haplotypes_ladder_bounders().

Usage

```
get_haplotypes_individuals(individuals)
```

Arguments

individuals Individuals to get haplotypes for.

Value

Matrix of haplotypes where row i is the haplotype of individuals[[i]].

See Also

```
get_haplotypes_pids().
```

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes(peds, 2, c(1, 1))
get_haplotypes_individuals(sim$end_generation_individuals)</pre>
```

```
{\tt get\_haplotypes\_in\_pedigree}
```

Get haplotypes in pedigree

Description

Get haplotypes in pedigree

Usage

```
get_haplotypes_in_pedigree(ped)
```

Arguments

ped

Pedigree

Value

List with haplotypes

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes(peds, 2, c(1, 1))
get_haplotypes_in_pedigree(peds[[1]])</pre>
```

get_haplotypes_pids 27

get_haplotypes_pids Get haplotypes from a vector of pids.

Description

Requires that haplotypes are first populated, e.g. with pedigrees_all_populate_haplotypes(), pedigrees_all_populate_haplotypes_custom_founders(), or pedigrees_all_populate_haplotypes_ladder_bounders().

Usage

```
get_haplotypes_pids(population, pids)
```

Arguments

population Population

pids Vector of pids to get haplotypes for.

Value

Matrix of haplotypes where row i is the haplotype of individuals[[i]].

See Also

```
get_haplotypes_individuals().
```

get_individual

Get individual by pid

Description

Get individual by pid

Usage

```
get_individual(population, pid)
```

Arguments

population Population pid pid

Value

Individual

Examples

```
sim <- sample_geneology(100, 10)
indv <- get_individual(sim$population, 1)
get_pid(indv)</pre>
```

get_individuals

Get all individuals in population

Description

Get all individuals in population

Usage

```
get_individuals(population)
```

Arguments

population Population

get_matching_pids_from_hashmap

Get individuals with a certain haplotype id by hashmap lookup

Description

By using hashmap made by build_haplotype_hashmap(), it is easy to get all individuals with a certain haplotype id.

Usage

```
get_matching_pids_from_hashmap(hashmap, haplotype)
```

Arguments

hashmap to make lookup in, made by build_haplotype_hashmap()

haplotype to get individuals that has this haplotype id

Value

List of individuals with a given haplotype id

See Also

```
build_haplotype_hashmap().
```

get_nodes_edges 29

get_nodes_edges

Get nodes and edges

Description

Get nodes and edges in malan_pedigreelist. For example to plot via as_tbl_graph().

Usage

```
get_nodes_edges(x, ...)
```

Arguments

x malan_pedigreelist

... Ignored

Value

List with entries nodes and edges

get_pedigrees_tidy

Get pedigrees information in tidy format

Description

Get pedigrees information in tidy format

Usage

```
get_pedigrees_tidy(pedigrees)
```

Arguments

pedigrees

Pedigrees

get_pedigree_as_graph Get pedigree information as graph (mainly intended for plotting)

Description

Get pedigree information as graph (mainly intended for plotting)

Usage

```
get_pedigree_as_graph(ped)
```

Arguments

ped

Pedigree

```
get_pedigree_from_individual
```

Get pedigree from individual

Description

Get pedigree from individual

Usage

```
get_pedigree_from_individual(individual)
```

Arguments

individual

Individual

Value

pedigree

get_pedigree_id 31

get_pedigree_id

Get pedigree id

Description

Get pedigree id

Usage

```
get_pedigree_id(ped)
```

Arguments

ped

Pedigree

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
get_pedigree_id(peds[[1]])</pre>
```

```
get_pedigree_id_from_pid
```

Get pedigree ids from pids

Description

Get pedigree ids from pids

Usage

```
get_pedigree_id_from_pid(population, pids)
```

Arguments

population Population

pids Pids

Value

Vector with pedigree ids

get_pid

Get pid from individual

Description

Get pid from individual

Usage

```
get_pid(individual)
```

Arguments

individual

Individual to get pid of

Value

pid

Examples

```
sim <- sample_geneology(100, 10)</pre>
indv <- get_individual(sim$population, 1)</pre>
get_pid(indv)
```

```
get_pids_in_pedigree Get pids in pedigree
```

Description

Get pids in pedigree

Usage

```
get_pids_in_pedigree(ped)
```

Arguments

ped

Pedigree

Examples

```
sim <- sample_geneology(100, 10)</pre>
peds <- build_pedigrees(sim$population)</pre>
get_pids_in_pedigree(peds[[1]])
```

get_uncles 33

get_uncles

Get uncles

Description

Get individual's uncles

Usage

```
get_uncles(individual)
```

Arguments

individual

individual

Value

List with uncles

See Also

```
get_brothers(), get_children(), get_cousins()
```

```
get_zero_haplotype_generator
```

Generate a function to generate the zero haplotype

Description

Generate a function to generate the zero haplotype

Usage

```
get_zero_haplotype_generator(loci)
```

Arguments

loci

Number of loci

grandfather_matches

Description

Does the frandfather have the same profile as individual?

Usage

```
grandfather_matches(individual)
```

Arguments

individual individual

Value

Whether grandfather has the same profile as individual or not

Description

Individuals with the same haplotype will have the same hash (integer) and individuals with different haplotypes will have different hashes (integers).

Usage

haplotypes_to_hashes(population, pids)

Arguments

population Population obtained from simulation

pids Vector of individual pids

Details

This can be useful if for example using haplotypes to define groups and the haplotype itself is not of interest.

Value

Integer vector with haplotype hashes

```
haplotype_matches_individuals
```

Get individuals matching from list of individuals

Description

Get the indvididuals that matches haplotype in individuals.

Usage

```
haplotype_matches_individuals(individuals, haplotype)
```

Arguments

```
individuals List of individuals to count occurrences in. haplotype Haplotype to count occurrences of.
```

Value

List of individuals that matches haplotype amongst individuals.

See Also

```
pedigree_haplotype_matches_in_pedigree_meiosis_L1_dists().
```

```
haplotype_partially_matches_individuals
```

Get individuals partially matching from list of individuals

Description

Get the indvididuals that partially matches haplotype in individuals.

Usage

```
haplotype_partially_matches_individuals(
  individuals,
  haplotype,
  ignore_loci = as.integer(c())
)
```

Arguments

```
individuals List of individuals to count occurrences in.
```

haplotype Haplotype to count occurrences of.

ignore_loci Vector of loci to ignore (1 = ignore first locus etc.)

36 infer_generations

Value

List of individuals that partially matches haplotype amongst individuals.

infer_generation

Infer individual's generation number

Description

Takes as input final generation, then moves up in pedigree and increments generation number.

Usage

```
infer_generation(final_generation)
```

Arguments

final_generation

Individuals in final generation

Details

Note: Only works when all final generation individuals are provided.

 $infer_generations$

Infer generation numbers from pedigrees

Description

Infer generation numbers from pedigrees

Usage

```
infer_generations(peds)
```

Arguments

peds

Pedigrees infered by build_pedigrees()

Value

Nothing

load_haplotypes 37

load_haplotypes

Load haplotypes to individuals

Description

Note that individuals loaded this way does not have information about generation.

Usage

```
load_haplotypes(population, pid, haplotypes, progress = TRUE)
```

Arguments

population of individuals pid ID of male

haplotypes • row i has pid[i] ID

progress Show progress.

load_individuals

Construct a population from data

Description

Note that individuals loaded this way does not have information about generation.

Usage

```
load_individuals(pid, pid_dad, progress = TRUE, error_on_pid_not_found = TRUE)
```

Arguments

pid ID of male

pid_dad ID of male's father, 0 if not known

progress Show progress.

error_on_pid_not_found

Error if pid not found

38 meiotic_dist

```
{\it meioses\_generation\_distribution} \\ {\it Meiotic~distribution}
```

Description

Get the distribution of number of meioses from individual to all individuals in individual's pedigree. Note the generation_upper_bound_in_result parameter.

Usage

```
meioses_generation_distribution(
  individual,
  generation_upper_bound_in_result = -1L
)
```

Arguments

individual Individual to calculate all meiotic distances from generation_upper_bound_in_result

Limit on distribution; -1 means no limit. 0 is the final generation. 1 second last generation etc.

meiotic_dist

Meiotic distance between two individuals

Description

Get the number of meioses between two individuals. Note, that pedigrees must first have been inferred by build_pedigrees().

Usage

```
meiotic_dist(ind1, ind2)
```

Arguments

ind1 Individual 1 ind2 Individual 2

Value

Number of meioses between ind1 and ind2 if they are in the same pedigree, else -1.

meiotic_dist_threshold

meiotic_dist_threshold

Meiotic distance between two individuals (with threshold)

Description

Get the number of meioses between two individuals. Note, that pedigrees must first have been inferred by build_pedigrees().

Usage

```
meiotic_dist_threshold(ind1, ind2, threshold)
```

Arguments

ind1 Individual 1 ind2 Individual 2

threshold Max search radius, if exceeding, return -1

Value

Number of meioses between ind1 and ind2 if they are in the same pedigree, else -1.

Description

Get all individual IDs within a meiotic radius Note, that pedigrees must first have been inferred by build_pedigrees().

Usage

```
meiotic_radius(ind, radius)
```

Arguments

ind Individual radius Max radius

Value

Matrix with ID and meiotic radius

```
mixture_info_by_individuals_2pers
```

Mixture information about 2 persons' mixture of donor1 and donor2.

Description

Mixture information about 2 persons' mixture of donor1 and donor2.

Usage

```
mixture_info_by_individuals_2pers(
  individuals,
  donor1,
  donor2,
  include_genealogy_info = TRUE
)
```

Arguments

individuals Individuals to consider as possible contributors and thereby get information

from.

donor1 Contributor1/donor 1 donor2 Contributor2/donor 2 include_genealogy_info

Include information about meiotic distances and family info

Value

A list with mixture information about the mixture donor1+donor2+donor3 from individuals

See Also

```
mixture_info_by_individuals_3pers, mixture_info_by_individuals_4pers, mixture_info_by_individuals_5pe
```

```
mixture_info_by_individuals_3pers
```

Mixture information about 3 persons' mixture of donor1, donor2 and donor3.

Description

Mixture information about 3 persons' mixture of donor1, donor2 and donor3.

Usage

```
mixture_info_by_individuals_3pers(individuals, donor1, donor2, donor3)
```

Arguments

individuals	Individuals to	consider as	s possible	contributors	and	thereby	get information
	C						

from.

donor1Contributor1/donor 1donor2Contributor2/donor 2donor3Contributor3/donor 3

Value

A list with mixture information about the mixture donor1+donor2+donor3 from individuals

See Also

mixture_info_by_individuals_2pers, mixture_info_by_individuals_4pers, mixture_info_by_individuals_5pe

mixture_info_by_individuals_4pers

Mixture information about 4 persons' mixture of donor1, donor2, donor3 and donor4.

Description

Mixture information about 4 persons' mixture of donor1, donor2, donor3 and donor4.

Usage

```
mixture_info_by_individuals_4pers(individuals, donor1, donor2, donor3, donor4)
```

Arguments

individuals	Individuals to consider as possible contributors and thereby get information from.
donor1	Contributor1/donor 1
donor2	Contributor2/donor 2
donor3	Contributor3/donor 3
donor4	Contributor4/donor 4

Value

A list with mixture information about the mixture donor1+donor2+donor3 from individuals

See Also

mixture_info_by_individuals_2pers, mixture_info_by_individuals_3pers, mixture_info_by_individuals_5pers

```
mixture_info_by_individuals_5pers

Mixture information about 5 persons' mixture of donor1, donor2, donor3, donor4 and donor5.
```

Mixture information about 5 persons' mixture of donor1, donor2, donor3, donor4 and donor5.

Usage

```
mixture_info_by_individuals_5pers(
  individuals,
  donor1,
  donor2,
  donor3,
  donor4,
  donor5
)
```

Arguments

individuals	Individuals to consider as possible contributors and thereby get information from.
donor1	Contributor1/donor 1
donor2	Contributor2/donor 2
donor3	Contributor3/donor 3
donor4	Contributor4/donor 4
donor5	Contributor5/donor 5

Value

A list with mixture information about the mixture donor1+donor2+donor3 from individuals

See Also

 $\verb|mixture_info_by_individuals_2pers|, \verb|mixture_info_by_individuals_3pers|, \verb|mixture_info_by_individuals_4pers|, \verb|mixture_info_by_info$

```
pedigrees_all_populate_autosomal

Populate 1-locus autosomal DNA profile in pedigrees with single-step
mutation model.
```

Populate 1-locus autosomal DNA profile from founder and down in all pedigrees. Note, that only alleles from ladder is assigned and that all founders draw type randomly.

Usage

```
pedigrees_all_populate_autosomal(
  pedigrees,
  allele_dist,
  theta,
  mutation_rate,
  progress = TRUE
)
```

Arguments

pedigrees Pedigree list in which to populate genotypes

allele_dist Allele distribution (probabilities) – gets normalised
theta Theta correction between 0 and 1 (both included)

mutation_rate Mutation rate between 0 and 1 (both included)

progress Show progress

Details

Note, that pedigrees must first have been inferred by build_pedigrees().

See Also

 $\verb|pedigrees_all_populate_haplotypes_custom_founders()| and | pedigrees_all_populate_haplotypes_ladder_bounders()| and | pedigrees_all_populate_haplotypes_la$

```
pedigrees_all_populate_haplotypes

Populate haplotypes in pedigrees (0-founder/unbounded).
```

Populate haplotypes from founder and down in all pedigrees. Note, that haplotypes are unbounded and that all founders get haplotype rep(0L, loci).

Usage

```
pedigrees_all_populate_haplotypes(
  pedigrees,
  loci,
  mutation_rates,
  prob_two_step = 0,
  prob_genealogical_error = 0,
  progress = TRUE
)
```

Arguments

```
pedigrees Pedigree list in which to populate haplotypes

loci Number of loci

mutation_rates Vector with mutation rates, length loci

prob_two_step Given a mutation happens, this is the probability that the mutation is a two-step mutation

prob_genealogical_error

Probability that a genealogical error happens: if so, give individual haplotype rep(0L, loci) instead of father's

progress Show progress
```

Details

Note, that pedigrees must first have been inferred by build_pedigrees().

See Also

```
pedigrees_all_populate_haplotypes_custom_founders() and pedigrees_all_populate_haplotypes_ladder_bounders()
```

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes(peds, 2, c(1, 1))
get_haplotype(sim$end_generation_individuals[[1]])</pre>
```

```
pedigrees_all_populate_haplotypes_custom_founders

Populate haplotypes in pedigrees (custom founder/unbounded).
```

Populate haplotypes from founder and down in all pedigrees. Note, that haplotypes are unbounded. All founders get a haplotype from calling the user provided function get_founder_haplotype().

Usage

```
pedigrees_all_populate_haplotypes_custom_founders(
  pedigrees,
  mutation_rates,
  get_founder_haplotype = NULL,
  prob_two_step = 0,
  prob_genealogical_error = 0,
  progress = TRUE
)
```

Arguments

Details

Note, that pedigrees must first have been inferred by build_pedigrees().

See Also

```
pedigrees_all_populate_haplotypes() and pedigrees_all_populate_haplotypes_ladder_bounded().
```

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes_custom_founders(
   peds, c(1, 1), function(x) c(10, 10))
get_haplotype(sim$end_generation_individuals[[1]])</pre>
```

```
pedigrees_all_populate_haplotypes_ladder_bounded

Populate haplotypes in pedigrees (custom founder/bounded).
```

Description

Populate haplotypes from founder and down in all pedigrees. Note, that haplotypes are bounded by ladder_min and ladder_max. All founders get a haplotype from calling the user provided function get_founder_haplotype().

Usage

```
pedigrees_all_populate_haplotypes_ladder_bounded(
  pedigrees,
  mutation_rates,
  ladder_min,
  ladder_max,
  get_founder_haplotype = NULL,
  prob_two_step = 0,
  prob_genealogical_error = 0,
  progress = TRUE
)
```

Arguments

```
pedigrees
                 Pedigree list in which to populate haplotypes
mutation_rates Vector with mutation rates
ladder_min
                 Lower bounds for haplotypes, same length as mutation_rates
ladder_max
                  Upper bounds for haplotypes, same length as mutation_rates; all entries must
                  be strictly greater than ladder_min
get_founder_haplotype
                 Function taking no arguments returning a haplotype of length(mutation_rates)
                 Given a mutation happens, this is the probability that the mutation is a two-
prob_two_step
                  step mutation; refer to details for information about behaviour around ladder
                  boundaries
prob_genealogical_error
                  Probability that a genealogical error happens: if so, give individual haplotype
                  get_founder_haplotype() instead of father's
                  Show progress
progress
```

pedigrees_count 47

Details

Given that a two step mutation should happen (probability specified by prob_two_step): With distances >= 2 to ladder bounds, mutations happen as usual. At distance = 0 or 1 to a ladder bound, the mutation is forced to move away from the boundary.

Note, that pedigrees must first have been inferred by build_pedigrees().

See Also

pedigrees_all_populate_haplotypes() and pedigrees_all_populate_haplotypes_custom_founders().

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_all_populate_haplotypes_ladder_bounded(
   peds, c(1, 1), c(0L, 0L), c(10L, 10L),
   function(x) c(10, 10))
get_haplotype(sim$end_generation_individuals[[1]])</pre>
```

pedigrees_count

Get number of pedigrees

Description

Get number of pedigrees

Usage

```
pedigrees_count(pedigrees)
```

Arguments

pedigrees

Pedigrees

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_count(peds)</pre>
```

48 pedigree_as_igraph

pedigrees_table

Get distribution of pedigree sizes

Description

Get distribution of pedigree sizes

Usage

```
pedigrees_table(pedigrees)
```

Arguments

pedigrees

Pedigrees

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigrees_table(peds)</pre>
```

pedigree_as_igraph

Convert pedigree to igraph

Description

Convert pedigree to igraph

Usage

```
pedigree_as_igraph(x, ...)
```

Arguments

x Pedigree ... ignored

Value

igraph object

Gives information about all individuals in pedigree that matches an individual. Just as count_haplotype_occurrences_indicounts the number of occurrences amongst a list of individuals, this gives detailed information about matching individuals in the pedigree, e.g. meiotic distances and maximum L1 distance on the path as some of these matches may have (back)mutations between in between them (but often this will be 0).

Usage

```
pedigree_haplotype_matches_in_pedigree_meiosis_L1_dists(
   suspect,
   generation_upper_bound_in_result = -1L,
   error_on_no_haplotype = TRUE
)
```

Arguments

```
suspect Individual that others must match the profile of.

generation_upper_bound_in_result

Only consider matches in generation 0, 1, ... generation_upper_bound_in_result.

-1 means disabled, consider all generations. End generation is generation 0.

Second last generation is 1. And so on.

error_on_no_haplotype

raise error or silently ignore individuals with no haplotype
```

Value

Matrix with information about matching individuals. Columns in order: meioses (meiotic distance to suspect), max_L1 (on the path between the matching individual and suspect, what is the maximum L1 distance between the suspect's profile and the profiles of the individuals on the path), pid (pid of matching individual)

See Also

```
count_haplotype_occurrences_individuals().
```

50 pedigree_size

Description

Gives information about all individuals in pedigree that almost matches an individual. Just as count_haplotype_near_matches_individuals() counts the number of occurrences amongst a list of individuals, this gives detailed information about almost matching individuals in the pedigree: for now, the meiotic distances.

Usage

```
pedigree_haplotype_near_matches_meiosis(
   suspect,
   max_dist,
   generation_upper_bound_in_result = -1L
)
```

Arguments

suspect Individual that others must match the profile of.

max_dist Maximum distance (0 = match, 1 = 1 STR allele difference, ...)

generation_upper_bound_in_result

Only consider matches in generation 0, 1, ... generation_upper_bound_in_result. -1 means disabled, consider all generations. End generation is generation 0. Second last generation is 1. And so on.

Value

Matrix with information about matching individuals. Columns in order: 1) meioses (meiotic distance to suspect), 2) haplotype distance, 3) pid (pid of matching individual)

See Also

```
count_haplotype_near_matches_individuals().
```

pedigree_size

Get pedigree size

Description

Get pedigree size

```
pedigree_size_generation
```

Usage

```
pedigree_size(ped)
```

Arguments

ped

Pedigree

Examples

```
sim <- sample_geneology(100, 10)
peds <- build_pedigrees(sim$population)
pedigree_size(peds[[1]])</pre>
```

```
pedigree_size_generation
```

Size of pedigree

Description

Get the size of the pedigree. Note the generation_upper_bound_in_result parameter.

Usage

```
pedigree_size_generation(pedigree, generation_upper_bound_in_result = -1L)
```

Arguments

```
pedigree Pedigree to get size of
```

generation_upper_bound_in_result

Limit on generation to include in count; -1 means no limit. 0 only include the final generation. 1 only second last generation etc.

Description

Plot malan_pedigree.

Usage

```
## S3 method for class 'malan_pedigree'
plot(
    X,
    ids = TRUE,
    haplotypes = FALSE,
    locus_sep = " ",
    mark_pids = NULL,
    label_color = "black",
    node_color = "lightgray",
    mark_color = "orange",
    ...
)
```

Arguments

```
Pedigree
Х
                 Show pids
ids
                 Show haplotypes
haplotypes
                 Locus separator in haplotypes
locus_sep
                 Vector of pids to highlight
mark_pids
label_color
                 Label color
node_color
                 Node color
mark_color
                 Highlight color
                 Passed to igraph::plot.igraph
```

```
{\tt plot.malan\_pedigreelist}
```

Plot pedigree list

Description

Plot malan_pedigreelist generated by build_pedigrees().

Usage

```
## S3 method for class 'malan_pedigreelist' plot(x, ...)
```

Arguments

```
x Pedigree list ignored
```

```
population_populate_autosomal_infinite_alleles
```

Populate 1-locus autosomal DNA profile in pedigrees with infinite alleles mutation model.

Description

Populate 1-locus autosomal DNA profile from founder and down in all pedigrees. Note, that all founders have type 0 to begin with.

Usage

```
population_populate_autosomal_infinite_alleles(
  population,
  mutation_rate,
  progress = TRUE
)
```

Arguments

population Population in which to populate genotypes mutation_rate Mutation rate between 0 and 1 (both included)

progress Show progress

Details

The maternal allele is taken by random from the 2*N[g] alleles in the previous generation consisting of N[g] males with descendants in the live population.

This is also why this is not using pedigrees but instead the population.

Note, that pedigrees need not be inferred.

See Also

```
pedigrees_all_populate_haplotypes_custom_founders() and pedigrees_all_populate_haplotypes_ladder_bounders
```

```
population_size_generation
Size of population
```

Description

Get the size of the population. Note the generation_upper_bound_in_result parameter.

Usage

```
population\_size\_generation(population, generation\_upper\_bound\_in\_result = -1L)
```

Arguments

```
population Population to get size of generation_upper_bound_in_result
```

Limit on generation to include in count; -1 means no limit. 0 only include the final generation. 1 only second last generation etc.

Description

Print pedigree

Usage

```
## S3 method for class 'malan_pedigree' print(x, ...)
```

Arguments

x Pedigree ... ignored

```
print.malan_pedigreelist
```

Print pedigree list

Description

Print malan_pedigreelist generated by build_pedigrees().

Usage

```
## S3 method for class 'malan_pedigreelist' print(x, ...)
```

Arguments

x Pedigrees (malan_pedigreelist)

... ignored

print.malan_population 55

```
print.malan_population
```

 $Print\ population$

Description

Print malan_population generated by sample_geneology() or sample_geneology_varying_size().

Usage

```
## S3 method for class 'malan_population' print(x, ...)
```

Arguments

```
x Population (malan_population)
... ignored
```

```
\label{lem:population_abort} Print. \verb|malan_population_abort| \\ Print. \verb|malan_population_abort| \\
```

Description

Print malan_population_abort

Usage

```
## S3 method for class 'malan_population_abort' print(x, \ldots)
```

Arguments

```
x malan_population_abort
... ignored
```

print_individual

Print individual

Description

Print individual

Usage

```
print_individual(individual)
```

Arguments

individual Individual

Examples

```
sim <- sample_geneology(100, 10)
indv <- get_individual(sim$population, 1)
print_individual(indv)</pre>
```

```
relationship_allele_diff_dist
```

Calculate distribution of allele difference

Description

Calculate distribution of allele difference after m meioses.

Usage

```
relationship_allele_diff_dist(meioses, mu_dw, mu_up, method = "explicit")
```

Arguments

meioses number of meioses separating the two individuals

mu_dw mutation rate for 1-step down-mutation mu_up mutation rate for 1-step up-mutation

method "explicit" (default): use known formulas for eigenvalues and eigenvectors.

Can cause numerical problems. "matmult": do matrix multiplication instead of diagonalisation. "matmult_mpfr": as "matmult" but with the Rmpfr library (note that this returns list instead of data.frame). "r_eigen": use R's

eigen() function to find eigen values. Mostly for debugging.

Value

data.frame with columns d (allele difference) and p (prob)

```
relationship_allele_diff_dist_sym
```

Calculate distribution of allele difference for symmetric mutation rates

Description

Calculate distribution of allele difference after m meioses.

Usage

```
relationship_allele_diff_dist_sym(meioses, mu_updw, method = "explicit")
```

Arguments

meioses number of meioses separating the two individuals

mu_updw mutation rate for 1-step down- and up-mutations, i.e. total mutation rate is

2*mu_updw

method "explicit" (default): use known formulas for eigenvalues and eigenvectors.

Can cause numerical problems. "matmult": do matrix multiplication instead of diagonalisation. "matmult_mpfr": as "matmult" but with the Rmpfr library (note that this returns list instead of data.frame). "r_eigen": use R's

eigen() function to find eigen values. Mostly for debugging.

Value

data.frame with columns d (allele difference) and p (prob)

```
sample_autosomal_genotype
```

Sample genotype with theta

Description

Sample genotype with theta

Usage

```
sample_autosomal_genotype(allele_dist, theta)
```

Arguments

allele_dist Allele distribution (probabilities) – gets normalised theta Theta correction between 0 and 1 (both included)

58 sample_geneology

sample_geneology

Simulate a geneology with constant population size.

Description

This function simulates a geneology where the last generation has population_size individuals.

Usage

```
sample_geneology(
  population_size,
  generations,
  generations_full = 1L,
  generations_return = 3L,
  enable_gamma_variance_extension = FALSE,
  gamma_parameter_shape = 5,
  gamma_parameter_scale = 1/5,
  progress = TRUE,
  verbose_result = FALSE
)
```

Arguments

```
population_size
```

The size of the population.

generations

The number of generations to simulate:

- -1 for simulate to 1 founder
- else simulate this number of generations.

generations_full

Number of full generations to be simulated.

generations_return

How many generations to return (pointers to) individuals for.

enable_gamma_variance_extension

Enable symmetric Dirichlet (and disable standard Wright-Fisher).

gamma_parameter_shape

Parameter related to symmetric Dirichlet distribution for each man's probability to be father. Refer to details.

gamma_parameter_scale

Parameter realted to symmetric Dirichlet distribution for each man's probability to be father. Refer to details.

progress Show progress.

verbose_result Verbose result.

sample_geneology 59

Details

By the backwards simulating process of the Wright-Fisher model, individuals with no descendants in the end population are not simulated. If for some reason additional full generations should be simulated, the number can be specified via the generations_full parameter. This can for example be useful if one wants to simulate the final 3 generations although some of these may not get (male) children.

Let α be the parameter of a symmetric Dirichlet distribution specifying each man's probability to be the father of an arbitrary male in the next generation. When $\alpha=5$, a man's relative probability to be the father has 95\ constant 1 under the standard Wright-Fisher model and the standard deviation in the number of male offspring per man is 1.10 (standard Wright-Fisher = 1).

This symmetric Dirichlet distribution is implemented by drawing father (unscaled) probabilities from a Gamma distribution with parameters gamma_parameter_shape and gamma_parameter_scale that are then normalised to sum to 1. To obtain a symmetric Dirichlet distribution with parameter α , the following must be used: ' $gamma_parameter_shape$ ' = α and ' $gamma_parameter_scale$ ' = $1/\alpha$.

Value

A malan_simulation / list with the following entries:

- population. An external pointer to the population.
- generations. Generations actually simulated, mostly useful when parameter generations = -1.
- founders. Number of founders after the simulated generations.
- growth_type. Growth type model.
- sdo_type. Standard deviation in a man's number of male offspring. StandardWF or GammaVariation depending on enable_gamma_variance_extension.
- end_generation_individuals. Pointers to individuals in end generation.
- individuals_generations. Pointers to individuals in last generations_return generation (if generations_return = 3, then individuals in the last three generations are returned).

If verbose_result is true, then these additional components are also returned:

- individual_pids. A matrix with pid (person id) for each individual.
- father_pids. A matrix with pid (person id) for each individual's father.
- father_indices. A matrix with indices for fathers.

See Also

```
sample_geneology_varying_size().
```

Examples

```
sim <- sample_geneology(100, 10)
str(sim, 1)
sim$population
peds <- build_pedigrees(sim$population)</pre>
```

peds

```
sample_geneology_varying_size
```

Simulate a geneology with varying population size.

Description

This function simulates a geneology with varying population size specified by a vector of population sizes, one for each generation.

Usage

```
sample_geneology_varying_size(
  population_sizes,
  generations_full = 1L,
  generations_return = 3L,
  enable_gamma_variance_extension = FALSE,
  gamma_parameter_shape = 5,
  gamma_parameter_scale = 1/5,
  progress = TRUE
)
```

Arguments

```
population_sizes
```

The size of the population at each generation, g. population_sizes[g] is the population size at generation g. The length of population_sizes is the number of generations being simulated.

generations_full

Number of full generations to be simulated.

generations_return

How many generations to return (pointers to) individuals for.

enable_gamma_variance_extension

Enable symmetric Dirichlet (and disable standard Wright-Fisher).

gamma_parameter_shape

Parameter related to symmetric Dirichlet distribution for each man's probability to be father. Refer to details.

gamma_parameter_scale

Parameter realted to symmetric Dirichlet distribution for each man's probability to be father. Refer to details.

progress Show progress.

Details

By the backwards simulating process of the Wright-Fisher model, individuals with no descendants in the end population are not simulated If for some reason additional full generations should be simulated, the number can be specified via the generations_full parameter. This can for example be useful if one wants to simulate the final 3 generations although some of these may not get (male) children.

Let α be the parameter of a symmetric Dirichlet distribution specifying each man's probability to be the father of an arbitrary male in the next generation. When $\alpha=5$, a man's relative probability to be the father has 95\ constant 1 under the standard Wright-Fisher model and the standard deviation in the number of male offspring per man is 1.10 (standard Wright-Fisher = 1).

This symmetric Dirichlet distribution is implemented by drawing father (unscaled) probabilities from a Gamma distribution with parameters gamma_parameter_shape and gamma_parameter_scale that are then normalised to sum to 1. To obtain a symmetric Dirichlet distribution with parameter α , the following must be used: ' $gamma_parameter_shape$ ' = α and ' $gamma_parameter_scale$ ' = $1/\alpha$.

Value

A malan_simulation / list with the following entries:

- population. An external pointer to the population.
- generations. Generations actually simulated, mostly useful when parameter generations
 -1.
- founders. Number of founders after the simulated generations.
- growth_type. Growth type model.
- sdo_type. Standard deviation in a man's number of male offspring. StandardWF or GammaVariation depending on enable_gamma_variance_extension.
- end_generation_individuals. Pointers to individuals in end generation.
- individuals_generations. Pointers to individuals in last generations_return generation (if generations_return = 3, then individuals in the last three generations are returned).

See Also

```
sample_geneology().
```

Examples

```
sim <- sample_geneology_varying_size(10*(1:10))
str(sim, 1)
sim$population
peds <- build_pedigrees(sim$population)
peds</pre>
```

62 split_by_haplotypes

set_generation

Set individual's generation number

Description

Note that generation 0 is final, end generation. 1 is second last generation etc.

Usage

```
set_generation(individual, generation)
```

Arguments

individual Individual

generation Generation to assign

Examples

```
sim <- sample_geneology(100, 10)
indv <- get_individual(sim$population, 1)
get_generation(indv)
set_generation(indv, 100)
get_generation(indv)</pre>
```

split_by_haplotypes

Split pids by haplotype

Description

Individuals with the same haplotype will be in the same group and individuals with different haplotypes will be in different groups.

Usage

```
split_by_haplotypes(population, pids)
```

Arguments

population Population obtained from simulation

pids Vector of individual pids

Value

List of integer vector, element i is an Integer Vector with all pids from pids with the same haplotype

test_create_population 63

test_create_population

Generate test population

Description

Generate test population

Usage

```
test_create_population()
```

Value

An external pointer to the population.

ystr_kits

Kit information about Y-STR markers

Description

A dataset containing information about the Y chromosomal short tandem repeat (Y-STR) markers that are present in the kit.

Usage

```
ystr_kits
```

Format

A data frame with 88 rows and 2 variables:

Marker name of Y-STR marker

Kit name of Y-STR kit

Source

```
https://www.yhrd.org
```

[[.malan_pedigreelist

ystr_markers

Mutational information about Y-STR markers

Description

A dataset from yhrd.org (and their sources) containing mutational information about Y chromosomal short tandem repeat (Y-STR) markers used in forensic genetics.

Usage

```
ystr_markers
```

Format

A data frame with 29 rows and 5 variables:

Marker name of Y-STR marker

Meioses number of meioses observed

Mutations number of mutations observed in the corresponding number of Meioses

MutProb point estimate of mutation probability, MutProb = Mutations/Meioses

Alleles observed alleles

Details

Note, that loci with duplications (DYS385a/b as well as DYF387S1a/b have been split into two loci).

Source

```
https://www.yhrd.org
```

[[.malan_pedigreelist Get pedigree from pedigree list

Description

Get pedigree from malan_pedigreelist generated by build_pedigrees().

Usage

```
## S3 method for class 'malan_pedigreelist' x[[...]]
```

[[.malan_population 65

Arguments

x Element id
... ignored

Value

Pedigree

[[.malan_population Get individual from population by pid

Description

Get individual from population by pid

Usage

```
## S3 method for class 'malan_population' x[[...]]
```

Arguments

x pid ... ignored

Value

Individual

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