Package 'MoBPS'

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Type Package
Title Modular Breeding Program Simulator
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Description Framework for the simulation framework for the simulation of complex breeding programs and compare their economic and genetic impact. The package is also used as the background simulator for our a web-based interface http://www.mobps.de . Associated publication: Pook et al. (2020) doi:10.1534/g3.120.401193 .
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add.array

Add a genotyping array

Description

Function to add a genotyping array for the population

Usage

```
add.array(population, marker.included = TRUE, array.name = NULL)
```

Arguments

population population list
marker.included

Vector with number of SNP entries coding if each marker is on the array (TRUE/FALSE)

array.name Name of the added array

Value

Population list

Examples

```
data(ex_pop)
population <- add.array(ex_pop, marker.included = c(TRUE, FALSE), array.name="Half-density")</pre>
```

add.combi 5

add.combi	Add a trait as a linear combination of other traits	

Description

Function to create an additional trait that is the results of a linear combination of the other traits

Usage

```
add.combi(population, trait, combi.weights, trait.name = NULL)
```

Arguments

population population list

trait trait nr. for which to implement a combination of other traits combi.weights Weights (only linear combinations of other traits are allowed!)

trait.name Name of the trait generated

Value

Population list Population list

Examples

```
data(ex_pop)
population <- creating.trait(ex_pop, n.additive = 100)
population <- add.combi(population, trait = 3, combi.weights = c(1,5))</pre>
```

add.diag

Add something to the diagonal

Description

Function to add numeric to the diagonal of a matrix

Usage

```
add.diag(M, d)
```

Arguments

M Matrix

d Vector to add to the diagonal of the matrix

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Value

Matrix with increased diagonal elements

Matrix with modified diagonal entries

Examples

```
A <- matrix(c(1,2,3,4), ncol=2)
B <- add.diag(A, 5)
```

add.founder.kinship

Add a relationship matrix for founder individuals

Description

Function to relationship matrix for founder individuals that is used for any calculation of the pedigree

Usage

```
add.founder.kinship(population, founder.kinship = "vanRaden", gen = 1)
```

Arguments

population population list

founder.kinship

Default is to use vanRaden relationship. Alternative is to enter a pedigree-matrix

(order of individuals is first male then female)

gen Generation for which to enter the pedigree-matrix

Value

Population list

Examples

```
data(ex_pop)
population <- add.founder.kinship(ex_pop)</pre>
```

alpha_to_beta 7

			1 .
aТ	pna	to	_beta

Moore-Penrose-Transfomration

Description

Internal transformation using Moore-Penrose

Usage

```
alpha_to_beta(alpha, G, Z)
```

Arguments

alpha alpha

G kinship-matrix

Z genomic information matrix

Value

Vector with single marker effects

analyze.bv

Analyze genomic values

Description

Function to analyze correlation between bv/bve/pheno

Usage

```
analyze.bv(
  population,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  bvrow = "all",
  advanced = FALSE
)
```

Arguments

population Population list

gen Quick-insert for database (vector of all generations to export)

database Groups of individuals to consider for the export

cohorts Quick-insert for database (vector of names of cohorts to export)

bvrow Which traits to display

advanced Set to TRUE to also look at offspring pheno

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Value

[1] Correlation between BV/BVE/Phenotypes [[2]] Genetic variance of the traits

Examples

```
data(ex_pop)
analyze.bv(ex_pop,gen=1)
```

analyze.population

Analyze allele frequency of a single marker

Description

Analyze allele frequency of a single marker

Usage

```
analyze.population(
  population,
  chromosome = NULL,
  snp = NULL,
  snp.name = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL
```

Arguments

population Population list

Chromosome Number of the

chromosome Number of the chromosome of the relevant SNP

snp Number of the relevant SNP snp.name Name of the SNP to analyze

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

Value

Frequency of AA/AB/BB in selected gen/database/cohorts

Examples

```
data(ex_pop)
analyze.population(ex_pop, snp=1, chromosome=1, gen=1:5)
```

bit.snps 9

bit.snps

Decoding of bitwise-storing in R

Description

Function for decoding in bitwise-storing in R (only 30 of 32 bits are used!)

Usage

```
bit.snps(bit.seq, nbits, population = NULL, from.p.bit = 1)
```

Arguments

bit.seq bitweise gespeicherte SNP-Sequenz
nbits Number of usable bits (default: 30)

population Population list from.p.bit Bit to start on

Value

De-coded marker sequence

bit.storing

Bitwise-storing in R

Description

Function for bitwise-storing in R (only 30 of 32 bits are used!)

Usage

```
bit.storing(snpseq, nbits)
```

Arguments

snpseq SNP sequence

nbits Number of usable bits (default: 30)

Value

Bit-wise coded marker sequence

breeding.diploid

Breeding function

Description

Function to simulate a step in a breeding scheme

Usage

```
breeding.diploid(
  population,
 mutation.rate = 10^-8,
  remutation.rate = 10^-8,
  recombination.rate = 1,
  selection.m = NULL,
  selection.f = NULL,
  new.selection.calculation = TRUE,
  selection.function.matrix = NULL,
  selection.size = 0,
  ignore.best = 0,
  breeding.size = 0,
  breeding.sex = NULL,
  breeding.sex.random = FALSE,
  relative.selection = FALSE,
  class.m = 0,
  class.f = 0,
  add.gen = 0,
  recom.f.indicator = NULL,
  duplication.rate = 0,
  duplication.length = 0.01,
  duplication.recombination = 1,
  new.class = 0L,
  bve = FALSE,
  sigma.e = NULL,
  sigma.g = 100,
  new.bv.child = NULL,
  phenotyping.child = NULL,
  relationship.matrix = "vanRaden",
  relationship.matrix.ogc = "kinship",
  computation.A = NULL,
  computation.A.ogc = NULL,
  delete.haplotypes = NULL,
  delete.individuals = NULL,
  fixed.breeding = NULL,
  fixed.breeding.best = NULL,
 max.offspring = Inf,
 max.litter = Inf,
```

```
store.breeding.totals = FALSE,
forecast.sigma.g = TRUE,
multiple.bve = "add",
store.bve.data = FALSE,
fixed.assignment = FALSE,
reduce.group = NULL,
reduce.group.selection = "random",
selection.highest = c(TRUE, TRUE),
selection.criteria = NULL,
same.sex.activ = FALSE,
same.sex.sex = 0.5,
same.sex.selfing = FALSE,
selfing.mating = FALSE,
selfing.sex = 0.5,
praeimplantation = NULL,
heritability = NULL,
repeatability = NULL,
save.recombination.history = FALSE,
martini.selection = FALSE,
BGLR.bve = FALSE,
BGLR.model = "RKHS",
BGLR.burnin = 500,
BGLR.iteration = 5000,
BGLR.print = FALSE,
copy.individual = FALSE,
copy.individual.m = FALSE,
copy.individual.f = FALSE,
dh.mating = FALSE,
dh.sex = 0.5,
n.observation = NULL,
bve.0isNA = FALSE,
phenotype.bv = FALSE,
delete.same.origin = FALSE,
remove.effect.position = FALSE,
estimate.u = FALSE,
new.phenotype.correlation = NULL,
new.residual.correlation = NULL,
new.breeding.correlation = NULL,
estimate.add.gen.var = FALSE,
estimate.pheno.var = FALSE,
best1.from.group = NULL,
best2.from.group = NULL,
best1.from.cohort = NULL,
best2.from.cohort = NULL,
add.class.cohorts = TRUE,
store.comp.times = TRUE,
store.comp.times.bve = TRUE,
store.comp.times.generation = TRUE,
```

```
import.position.calculation = NULL,
BGLR.save = "RKHS",
BGLR.save.random = FALSE,
ogc = FALSE,
ogc.target = "min.sKin",
ogc.uniform = NULL,
ogc.ub = NULL,
ogc.1b = NULL,
ogc.ub.sKin = NULL,
ogc.lb.BV = NULL,
ogc.ub.BV = NULL,
ogc.eq.BV = NULL,
ogc.ub.sKin.increase = NULL,
ogc.lb.BV.increase = NULL,
emmreml.bve = FALSE,
rrblup.bve = FALSE,
sommer.bve = FALSE,
sommer.multi.bve = FALSE,
nr.edits = 0,
gene.editing.offspring = FALSE,
gene.editing.best = FALSE,
gene.editing.offspring.sex = c(TRUE, TRUE),
gene.editing.best.sex = c(TRUE, TRUE),
gwas.u = FALSE,
approx.residuals = TRUE,
sequenceZ = FALSE,
maxZ = 5000,
maxZtotal = 0,
delete.sex = 1:2,
gwas.group.standard = FALSE,
y.gwas.used = "pheno",
gen.architecture.m = 0,
gen.architecture.f = NULL,
add.architecture = NULL,
ncore = 1,
ncore.generation = 1,
Z.integer = FALSE,
store.effect.freq = FALSE,
backend = "doParallel",
randomSeed = NULL,
randomSeed.generation = NULL,
Rprof = FALSE,
miraculix = NULL,
miraculix.cores = 1,
miraculix.mult = NULL,
miraculix.chol = TRUE,
best.selection.ratio.m = 1,
best.selection.ratio.f = NULL,
```

```
best.selection.criteria.m = "bv",
best.selection.criteria.f = NULL,
best.selection.manual.ratio.m = NULL,
best.selection.manual.ratio.f = NULL,
best.selection.manual.reorder = TRUE,
bve.class = NULL,
parallel.generation = FALSE,
name.cohort = NULL,
display.progress = TRUE,
combine = FALSE,
repeat.mating = NULL,
repeat.mating.copy = NULL,
repeat.mating.fixed = NULL,
repeat.mating.overwrite = TRUE,
time.point = 0,
creating.type = 0,
multiple.observation = FALSE,
new.bv.observation = NULL,
new.bv.observation.gen = NULL,
new.bv.observation.cohorts = NULL,
new.bv.observation.database = NULL,
phenotyping = NULL,
phenotyping.gen = NULL,
phenotyping.cohorts = NULL,
phenotyping.database = NULL,
bve.gen = NULL,
bve.cohorts = NULL,
bve.database = NULL,
sigma.e.gen = NULL,
sigma.e.cohorts = NULL,
sigma.e.database = NULL,
sigma.g.gen = NULL,
sigma.g.cohorts = NULL,
sigma.g.database = NULL,
gwas.gen = NULL,
gwas.cohorts = NULL,
gwas.database = NULL,
bve.insert.gen = NULL,
bve.insert.cohorts = NULL,
bve.insert.database = NULL,
reduced.selection.panel.m = NULL,
reduced.selection.panel.f = NULL,
breeding.all.combination = FALSE,
depth.pedigree = 7,
depth.pedigree.ogc = 7,
copy.individual.keep.bve = TRUE,
copy.individual.keep.pheno = TRUE,
bve.avoid.duplicates = TRUE,
```

```
report.accuracy = TRUE,
share.genotyped = 1,
singlestep.active = FALSE,
remove.non.genotyped = TRUE,
added.genotyped = 0,
fast.uhat = TRUE,
offspring.bve.parents.gen = NULL,
offspring.bve.parents.database = NULL,
offspring.bve.parents.cohorts = NULL,
offspring.bve.offspring.gen = NULL,
offspring.bve.offspring.database = NULL,
offspring.bve.offspring.cohorts = NULL,
culling.gen = NULL,
culling.database = NULL,
culling.cohort = NULL,
culling.time = Inf,
culling.name = "Not_named",
culling.bv1 = 0,
culling.share1 = 0,
culling.bv2 = NULL,
culling.share2 = NULL,
culling.index = 0,
culling.single = TRUE,
culling.all.copy = TRUE,
calculate.reliability = FALSE,
selection.m.gen = NULL,
selection.f.gen = NULL,
selection.m.database = NULL,
selection.f.database = NULL,
selection.m.cohorts = NULL,
selection.f.cohorts = NULL,
selection.m.miesenberger = FALSE,
selection.f.miesenberger = NULL,
selection.miesenberger.reliability.est = "estimated",
miesenberger.trafo = 0,
multiple.bve.weights.m = 1,
multiple.bve.weights.f = NULL,
multiple.bve.scale.m = "bv_sd",
multiple.bve.scale.f = NULL,
verbose = TRUE,
bve.parent.mean = FALSE,
bve.grandparent.mean = FALSE,
bve.mean.between = "bvepheno",
bve.direct.est = TRUE,
bve.pseudo = FALSE,
bve.pseudo.accuracy = 1,
miraculix.destroyA = TRUE,
mas.bve = FALSE,
```

```
mas.markers = NULL,
      mas.number = 5,
      mas.effects = NULL,
      threshold.selection = NULL,
      threshold.sign = ">",
      input.phenotype = "own",
      bve.ignore.traits = NULL,
      bv.ignore.traits = NULL,
      genotyped.database = NULL,
      genotyped.gen = NULL,
      genotyped.cohorts = NULL,
      genotyped.share = 1,
      genotyped.array = 1,
      sex.s = NULL,
      bve.imputation = TRUE,
      bve.imputation.errorrate = 0,
      share.phenotyped = 1,
      avoid.mating.fullsib = FALSE,
      avoid.mating.halfsib = FALSE,
      max.mating.pair = Inf,
      bve.per.sample.sigma.e = TRUE,
      bve.solve = "exact"
    )
Arguments
    population
                     Population list
    mutation.rate
                     Mutation rate in each marker (default: 10^-8)
    remutation.rate
                     Remutation rate in each marker (default: 10^-8)
    recombination.rate
                     Average number of recombination per 1 length unit (default: 1M)
    selection.m
                     Selection criteria for male individuals (Set to "random" to randomly select indi-
                     viduals - this happens automatically when no the input in selection.criteria has
                     no input ((usually breeding values)))
    selection.f
                     Selection criteria for female individuals (default: selection.m , alt: "random",
                     function")
    new.selection.calculation
                     If TRUE recalculate breeding values obtained by selection.function.matrix
    selection.function.matrix
                     Manuel generation of a temporary selection function (Use BVs instead!)
    selection.size Number of selected individuals for breeding (default: c(0,0) - alt: positive num-
                     Not consider the top individuals of the selected individuals (e.g. to use 2-10 best
    ignore.best
                     individuals)
```

Number of individuals to generate

breeding.size

Share of female animals (if single value is used for breeding size; default: 0.5) breeding.sex breeding.sex.random If TRUE randomly chose sex of new individuals (default: FALSE - use expected relative.selection Use best.selection.ratio instead! class.m Migrationlevels of male individuals to consider for mating process (default: 0) class.f Migrationlevels of female individuals to consider for mating process (default: add.gen Generation you want to add the new individuals to (default: New generation) recom.f.indicator Use step function for recombination map (transform snp.positions if possible instead) duplication.rate Share of recombination points with a duplication (default: 0 - DEACTIVATED) duplication.length Average length of a duplication (Exponentially distributed) duplication.recombination Average number of recombinations per 1 length uit of duplication (default: 1) new.class Migration level of newly generated individuals (default: 0) If TRUE perform a breeding value estimation (default: FALSE) bve sigma.e Environmental variance (default: 100) Genetic variance (default: 100 - only used if not computed via estimate.sigma.g^2 sigma.g in der Zuchtwertschaetzung (Default: 100) new.bv.child (OLD! - use phenotyping.child) Starting phenotypes of newly generated individuals (default: "mean" of both parents, "obs" - regular observation, "zero" -0) phenotyping.child Starting phenotypes of newly generated individuals (default: "mean" of both parents, "obs" - regular observation, "zero" - 0) relationship.matrix Method to calculate relationship matrix for the breeding value estimation (Default: "vanRaden", alt: "kinship", "CE", "non_stand", "CE2", "CM") relationship.matrix.ogc Method to calculate relationship matrix for OGC (Default: "kinship", alt: "van-Raden", "CE", "non_stand", "CE2", "CM") (OLD! - use relationship.matrix) Method to calculate relationship matrix for the computation.A breeding value estimation (Default: "vanRaden", alt: "kinship", "CE", "non_stand", "CE2", "CM") computation.A.ogc (OLD! use relationship.matrix.ogc) Method to calculate pedigree matrix in OGC (Default: "kinship", alt: "vanRaden", "CE", "non_stand", "CE2", "CM") delete.haplotypes Generations for with haplotypes of founders can be deleted (only use if storage

problem!)

delete.individuals Generations for with individuals are completley deleted (only use if storage problem!) fixed.breeding Set of targeted matings to perform fixed.breeding.best Perform targeted matings in the group of selected individuals Maximum number of offspring per individual (default: c(Inf,Inf) - (m,w)) max.offspring Maximum number of offspring per individual (default: c(Inf,Inf) - (m,w)) max.litter store.breeding.totals If TRUE store information on selected animals in \$info\$breeding.totals forecast.sigma.g Set FALSE to not estimate sigma.g (Default: TRUE) multiple.bve Way to handle multiple traits in bv/selection (default: "add", alt: "ranking") store.bve.data If TRUE store information of bve in \$info\$bve.data fixed.assignment Set TRUE for targeted mating of best-best individual till worst-worst (of selected). set to "bestworst" for best-worst mating (OLD! - use culling modules) Groups of animals for reduce to a new size (by reduce.group changing class to -1) reduce.group.selection (OLD! - use culling modules) Selection criteria for reduction of groups (cf. selection.m / selection.f - default: "random") selection.highest If 0 individuals with lowest bve are selected as best individuals (default c(1,1) -(m,w)selection.criteria What to use in the selection proces (default: "bve", alt: "bv", "pheno") same.sex.activ If TRUE allow matings of individuals of same sex Probability to use female individuals as parents (default: 0.5) same.sex.sex same.sex.selfing Set to TRUE to allow for selfing when using same.sex matings selfing.mating If TRUE generate new individuals via selfing selfing.sex Share of female individuals used for selfing (default: 0.5) praeimplantation Only use matings the lead to a specific genotype in a specific marker heritability Use sigma.e to obtain a certain heritability (default: NULL) repeatability Set this to control the share of the residual variance (sigma.e) that is permanent (there for each observation) save.recombination.history If TRUE store the time point of each recombination event martini.selection If TRUE use the group of non-selected individuals as second parent BGLR.bve If TRUE use BGLR to perform breeding value estimation

BGLR.model Select which BGLR model to use (default: "RKHS", alt: "BRR", "BL", "BayesA",

"BayesB", "BayesC")

BGLR.burnin Number of burn-in steps in BGLR (default: 1000)

BGLR. iteration Number of iterations in BGLR (default: 5000)

BGLR.print If TRUE set verbose to TRUE in BGLR

copy.individual

If TRUE copy the selected father for a mating

copy.individual.m

If TRUE generate exactly one copy of all selected male in a new cohort (or more by setting breeding.size)

copy.individual.f

If TRUE generate exactly one copy of all selected female in a new cohort (or more by setting breeding.size)

dh.mating If TRUE generate a DH-line in mating process

dh. sex Share of DH-lines generated from selected female individuals

n.observation Number of phenotypes generated per individuals (influences environmental vari-

ance)

bve.@isNA Individuals with phenotype 0 are used as NA in breeding value estimation

phenotype.bv If TRUE use phenotype as estimated breeding value

delete.same.origin

If TRUE delete recombination points when genetic origin of adjacent segments is the same

remove.effect.position

If TRUE remove real QTLs in breeding value estimation

estimate.u If TRUE estimate u in breeding value estimation (Y = Xb + Zu + e)

new.phenotype.correlation

(OLD! - use new.residual.correlation!) Correlation of the simulated environmental variance

new.residual.correlation

Correlation of the simulated environmental variance

new.breeding.correlation

Correlation of the simulated genetic variance (child share! heritage is not influenced!)

estimate.add.gen.var

If TRUE estimate additive genetic variance and heritability based on parent model

estimate.pheno.var

If TRUE estimate total variance in breeding value estimation

best1.from.group

(OLD!- use selection.m.database) Groups of individuals to consider as First Parent / Father (also female individuals are possible)

best2.from.group

(OLD!- use selection.f.database) Groups of individuals to consider as Second Parent / Mother (also male individuals are possible)

best1.from.cohort

(OLD!- use selection.m.cohorts) Groups of individuals to consider as First Parent / Father (also female individuals are possible)

best2.from.cohort

(OLD! - use selection.f.cohorts) Groups of individuals to consider as Second Parent / Mother (also male individuals are possible)

add.class.cohorts

Migration levels of all cohorts selected for reproduction are automatically added to class.m/class.f (default: TRUE)

store.comp.times

If TRUE store computation times in \$info\$comp.times (default: TRUE)

store.comp.times.bve

If TRUE store computation times of breeding value estimation in \$info\$comp.times.bve (default: TRUE)

store.comp.times.generation

If TRUE store computation times of mating simulations in \$info\$comp.times.generation (default: TRUE)

import.position.calculation

Function to calculate recombination point into adjacent/following SNP

BGLR. save Method to use in BGLR (default: "RKHS" - alt: NON currently)

BGLR.save.random

Add random number to store location of internal BGLR computations (only needed when simulating a lot in parallel!)

ogc If TRUE use optimal genetic contribution theory to perform selection (This

requires the use of the R-package optiSel)

ogc.target Target of OGC (default: "min.sKin" - minimize inbreeding; alt: "max.BV" /

"min.BV" - maximize genetic gain; both under constrains selected below)

ogc.uniform This corresponds to the uniform constrain in optiSel

ogc.ub This corresponds to the ub constrain in optiSel ogc.lb This corresponds to the lb constrain in optiSel

ogc.ub.sKin

This corresponds to the ub.sKin constrain in optiSel
ogc.lb.BV

This corresponds to the lb.BV constrain in optiSel
ogc.ub.BV

This corresponds to the ub.BV constrain in optiSel

ogc.eq.BV This corresponds to the eq.BV constrain in optiSel

ogc.ub.sKin.increase

This corresponds to the upper bound (current sKin + ogc.ub.sKin.increase) as ub.sKin in optiSel

ogc.lb.BV.increase

This corresponds to the lower bound (current BV + ogc.lb.BV.increase) as lb.BV in optiSel

emmreml.bve If TRUE use REML estimator from R-package EMMREML in breeding value

estimation

rrblup.bve If TRUE use REML estimator from R-package rrBLUP in breeding value esti-

mation

sommer.bve If TRUE use REML estimator from R-package sommer in breeding value estimation

sommer.multi.bve

Set TRUE to use a mulit-trait model in the R-package sommer for BVE

nr.edits Number of edits to perform per individual

gene.editing.offspring

If TRUE perform gene editing on newly generated individuals

gene.editing.best

If TRUE perform gene editing on selected individuals

gene.editing.offspring.sex

Which sex to perform editing on (Default c(TRUE, TRUE), mw)

gene.editing.best.sex

Which sex to perform editing on (Default c(TRUE,TRUE), mw)

gwas.u If TRUE estimate u via GWAS (relevant for gene editing)

approx.residuals

If FALSE calculate the variance for each marker separatly instead of using a set

variance (doesnt change order - only p-values)

sequenceZ Split genomic matric into parts (relevent if high memory usage)

maxZ Number of SNPs to consider in each part of sequenceZ

maxZtotal Number of matrix entries to consider jointly (maxZ = maxZtotal/number of an-

imals)

delete.sex Remove all individuals from these sex from generation delete.individuals (de-

fault: 1:2; note:delete individuals=NULL)

gwas.group.standard

If TRUE standardize phenotypes by group mean

y.gwas.used What y value to use in GWAS study (Default: "pheno", alt: "bv", "bve")

gen.architecture.m

Genetic architecture for male animal (default: 0 - no transformation)

gen.architecture.f

Genetic architecture for female animal (default: gen.architecture.m - no transformation)

add.architecture

List with two vectors containing (A: length of chromosomes, B: position in cM

of SNPs)

ncore Cores used for parallel computing in compute.snps

ncore.generation

Number of cores to use in parallel generation

Z. integer If TRUE save Z as a integer in parallel computing

store.effect.freq

If TRUE store the allele frequency of effect markers per generation

backend Chose the used backend (default: "doParallel", alt: "doMPI")

randomSeed Set random seed of the process

randomSeed.generation

Set random seed for parallel generation process

Rprof Store computation times of each function

miraculix If TRUE use miraculix to perform computations (ideally already generate popu-

lation in creating.diploid with this; default: automatic detection from population

list)

miraculix.cores

Number of cores used in miraculix applications (default: 1)

miraculix.mult If TRUE use miraculix for matrix multiplications even if miraculix is not used

for storage

miraculix.chol Set to FALSE to deactive miraculix based Cholesky-decomposition (default:

TRUE)

best.selection.ratio.m

Ratio of the frequency of the selection of the best best animal and the worst best

animal (default=1)

best.selection.ratio.f

Ratio of the frequency of the selection of the best best animal and the worst best

animal (default=1)

best.selection.criteria.m

Criteria to calculate this ratio (default: "bv", alt: "bve", "pheno")

best.selection.criteria.f

Criteria to calculate this ratio (default: "bv", alt: "bve", "pheno")

best.selection.manual.ratio.m

vector containing probability to draw from for every individual (e.g. c(0.1,0.2,0.7))

best.selection.manual.ratio.f

vector containing probability to draw from for every individual (e.g. c(0.1,0.2,0.7))

best.selection.manual.reorder

Set to FALSE to not use the order from best to worst selected individual but

plain order based on database-order

bve.class Consider only animals of those class classes in breeding value estimation (de-

fault: NULL - use all)

parallel.generation

Set TRUE to active parallel computing in animal generation

name.cohort Name of the newly added cohort

display.progress

Set FALSE to not display progress bars. Setting verbose to FALSE will auto-

matically deactive progress bars

combine Copy existing individuals (e.g. to merge individuals from different groups in a

joined cohort). Individuals to use are used as the first parent

repeat.mating Generate multiple mating from the same dam/sire combination (first column:

number of offspring; second column: probability)

repeat.mating.copy

Generate multiple copies from a copy action (combine / copy.individuals.m/f)

(first column: number of offspring; second column: probability)

repeat.mating.fixed

Vector containing number of times each mating is repeated. This will overwrite sampling from repeat.mating / repeat.mating.copy (default: NULL)

repeat.mating.overwrite

Set to FALSE to not use the current repeat.mating / repeat.mating.copy input as the new standard values (default: TRUE)

time.point Time point at which the new individuals are generated

creating.type Technique to generate new individuals (usage in web-based application) multiple.observation

Set TRUE to allow for more than one phenotype observation per individual (this will decrease environmental variance!)

new.bv.observation

(OLD! - use phenotyping) Quick acces to phenotyping for (all: "all", non-phenotyped: "non_obs", non-phenotyped male: "non_obs_m", non-phenotyped female: "non_obs_f")

new.bv.observation.gen

(OLD! use phenotyping.gen) Vector of generation from which to generate additional phenotypes

new.bv.observation.cohorts

(OLD! use phenotyping.cohorts)Vector of cohorts from which to generate additional phenotype

new.bv.observation.database

(OLD! use phenotyping.database) Matrix of groups from which to generate additional phenotypes

phenotyping Quick acces to phenotyping for (all: "all", non-phenotyped: "non_obs", non-phenotyped male: "non_obs_m", non-phenotyped female: "non_obs_f")

phenotyping.gen

Vector of generation from which to generate additional phenotypes

phenotyping.cohorts

Vector of cohorts from which to generate additional phenotype

phenotyping.database

Matrix of groups from which to generate additional phenotypes

bve.gen Generations of individuals to consider in breeding value estimation (default: NULL)

bve.cohorts Cohorts of individuals to consider in breeding value estimation (default: NULL) bve.database Groups of individuals to consider in breeding value estimation (default: NULL)

sigma.e.gen Generations to consider when estimating sigma.e when using hertability

sigma.e.cohorts

Cohorts to consider when estimating sigma.e when using hertability

sigma.e.database

Groups to consider when estimating sigma.e when using hertability

sigma.g.gen Generations to consider when estimating sigma.g

sigma.g.cohorts

Cohorts to consider when estimating sigma.g

sigma.g.database

Groups to consider when estimating sigma.g

gwas.gen Generations to consider in GWAS analysis

gwas.cohorts Cohorts to consider in GWAS analysis

gwas.database Groups to consider in GWAS analysis

bve.insert.gen Generations of individuals to compute breeding values for (default: all groups

in bye.database)

bve.insert.cohorts

Cohorts of individuals to compute breeding values for (default: all groups in bye.database)

bve.insert.database

Groups of individuals to compute breeding values for (default: all groups in bye.database)

reduced.selection.panel.m

Use only a subset of individuals of the potential selected ones ("Split in user-interface")

reduced.selection.panel.f

Use only a subset of individuals of the potential selected ones ("Split in user-interface")

breeding.all.combination

Set to TRUE to automatically perform each mating combination possible exactly ones

depth.pedigree Depth of the pedigree in generations (default: 7)

depth.pedigree.ogc

Depth of the pedigree in generations (default: 7)

copy.individual.keep.bve

Set to FALSE to not keep estimated breeding value in case of use of copy.individuals

copy.individual.keep.pheno

Set to FALSE to not keep estimated breeding values in case of use of copy.individuals

bve.avoid.duplicates

If set to FALSE multiple generatations of the same individual can be used in the bye (only possible by using copy.individual to generate individuals)

report.accuracy

Report the accuracy of the breeding value estimation

share.genotyped

Share of individuals newly generated individuals that are genotyped

singlestep.active

Set TRUE to use single step in breeding value estimation (only implemented for vanRaden- G matrix and without use sequenceZ) (Legarra 2014)

remove.non.genotyped

Set to FALSE to manually include non-genotyped individuals in genetic BVE, single-step will deactive this as well

added.genotyped

Share of individuals that is additionally genotyped (only for copy.individuals)

Set to FALSE to derive inverse of A in rrBLUP fast.uhat offspring.bve.parents.gen Generations to consider to derive phenotype from offspring phenotypes offspring.bve.parents.database Groups to consider to derive phenotype from offspring phenotypes offspring.bve.parents.cohorts Cohorts to consider to derive phenotype from offspring phenotypes offspring.bve.offspring.gen Active generations for import of offspring phenotypes offspring.bve.offspring.database Active groups for import of offspring phenotypes offspring.bve.offspring.cohorts Active cohorts for import of offspring phenotypes culling.gen Generations to consider to culling culling.database Groups to consider to culling culling.cohort Cohort to consider to culling culling.time Age of the individuals at culling culling.name Name of the culling action (user-interface stuff) culling.bv1 Reference Breeding value culling. share1 Probability of death for individuals with bv1 Alternative breeding value (linear extended for other bvs) culling.bv2 culling. share 2 Probability of death for individuals with bv2 Genomic index (default:0 - no genomic impact, use: "lastindex" to use the last culling.index selection index applied in selection) culling.single Set to FALSE to not apply the culling module on all individuals of the cohort culling.all.copy Set to FALSE to not kill copies of the same individual in the culling module calculate.reliability Set TRUE to calculate a reliability when performing Direct-Mixed-Model BVE selection.m.gen Generations available for selection of paternal parent selection.f.gen Generations available for selection of maternal parent selection.m.database Groups available for selection of paternal parent selection.f.database Groups available for selection of maternal parent selection.m.cohorts Cohorts available for selection of paternal parent selection.f.cohorts

Cohorts available for selection of maternal parent

selection.m.miesenberger

Use Weighted selection index according to Miesenberger 1997 for paternal selection

selection.f.miesenberger

Use Weighted selection index according to Miesenberger 1997 for maternal selection

selection.miesenberger.reliability.est

If available reliability estimated are used. If not use default: "estimated" (SD BVE / SD Pheno), alt: "heritability", "derived" (cor(BVE,BV)^2) as replacement

miesenberger.trafo

Ignore all eigenvalues below this threshold and apply dimension reduction (default: 0 - use all)

multiple.bve.weights.m

Weighting between traits when using "add" (default: 1)

multiple.bve.weights.f

Weighting between traits when using "add" (default: same as multiple.bve.weights.m)

multiple.bve.scale.m

Default: "bv_sd"; Set to "pheno_sd" when using gains per phenotypic SD, "unit" when using gains per unit, "bve" when using estimated breeding values

multiple.bve.scale.f

Default: "bv_sd"; Set to "pheno_sd" when using gains per phenotypic SD, "unit" when using gains per unit, "bve" when using estimated breeding values

verbose Set to FALSE to not display any prints

bve.parent.mean

Set to TRUE to use the average parental performance as the breeding value estimate

bve.grandparent.mean

Set to TRUE to use the average grandparental performance as the breeding value estimate

bve.mean.between

Select if you want to use the "bve", "bv", "pheno" or "bvepheno" to form the mean (default: "bvepheno" - if available bve, else pheno)

bve.direct.est If TRUE predict BVEs in direct estimation according to vanRaden 2008 method 2 (default: TRUE)

bve.pseudo If set to TRUE the breeding value estimation will be simulated with resulting accuracy bve.pseudo.accuracy (default: 1)

bve.pseudo.accuracy

The accuracy to be obtained in the "pseudo" - breeding value estimation

miraculix.destroyA

If FALSE A will not be destroyed in the process of inversion (less computing / more memory)

mas.bve If TRUE use marker assisted selection in the breeding value estimation
mas.markers Vector containing markers to be used in marker assisted selection

mas.number If no markers are provided this nr of markers is selected (if single marker QTL are present highest effect markers are prioritized)

mas.effects Effects assigned to the MAS markers (Default: estimated via lm()) threshold.selection

Minimum value in the selection index selected individuals have to have

threshold.sign Pick all individuals above (">") the threshold. Alt: ("<", "=", "<=", ">=") input.phenotype

Select what to use in BVE (default: own phenotype ("own"), offspring phenotype ("off"), their average ("mean") or a weighted average ("weighted"))

bve.ignore.traits

Vector of traits to ignore in the breeding value estimation (default: NULL, use: "zero" to not consider traits with 0 index weight in multiple.bve.weights.m/.w)

bv.ignore.traits

Vector of traits to ignore in the calculation of the genomic value (default: NULL; Only recommended for high number of traits and experienced users!)

genotyped.database

Groups to generate genotype data (that can be used in a BVE)

genotyped.gen Generations to generate genotype data (that can be used in a BVE) genotyped.cohorts

Cohorts to generate genotype data (that can be used in a BVE)

genotyped.share

Share of individuals in genotyped.gen/database/cohort to generate genotype data from (default: 1)

genotyped.array

Genotyping array used

sex.s Specify which newly added individuals are male (1) or female (2)

bve.imputation Set to FALSE to not perform imputation up to the highest marker density of genotyping data that is available

bve.imputation.errorrate

Share of errors in the imputation procedure (default: 0)

share.phenotyped

Share of the individuals to phenotype

avoid.mating.fullsib

Set to TRUE to not generate offspring of full siblings

avoid.mating.halfsib

Set to TRUE to not generate offspring from half or full siblings

max.mating.pair

Set to the maximum number of matings between two individuals (default: Inf)

bve.per.sample.sigma.e

Set to FALSE to deactivate the use of a heritablity based on the number of observations generated per sample

bve.solve Provide solver to be used in BVE (default: "exact" solution via inversion, alt: "pcg", function with inputs A, b and output y_hat)

Value

Population-list

breeding.intern 27

Examples

```
population <- creating.diploid(nsnp=1000, nindi=100)
population <- breeding.diploid(population, breeding.size=100, selection.size=c(25,25))</pre>
```

breeding.intern

Internal function to simulate one meiosis

Description

Internal function to simulate one meiosis

Usage

```
breeding.intern(
  info.parent,
  parent,
  population,
 mutation.rate = 10^-5,
  remutation.rate = 10^-5,
  recombination.rate = 1,
  recom.f.indicator = NULL,
  duplication.rate = 0,
  duplication.length = 0.01,
  duplication.recombination = 1,
  delete.same.origin = FALSE,
  gene.editing = FALSE,
  nr.edits = 0,
  gen.architecture = 0,
  decodeOriginsU = MoBPS::decodeOriginsR
)
```

Arguments

```
info.parent position of the parent in the dataset

parent list of information regarding the parent

population Population list

mutation.rate Mutation rate in each marker (default: 10^-5)

remutation.rate

Remutation rate in each marker (default: 10^-5)

recombination.rate

Average number of recombination per 1 length unit (default: 1M)

recom.f.indicator

Use step function for recombination map (transform snp.positions if possible instead)
```

28 bv.development

```
duplication.rate
                  Share of recombination points with a duplication (default: 0 - DEACTIVATED)
duplication.length
                  Average length of a duplication (Exponentially distributed)
duplication.recombination
                  Average number of recombinations per 1 length uit of duplication (default: 1)
delete.same.origin
                 If TRUE delete recombination points when genetic origin of adjacent segments
                 is the same
gene.editing
                 If TRUE perform gene editing on newly generated individual
                  Number of edits to perform per individual
nr.edits
gen.architecture
                  Used underlying genetic architecture (genome length in M)
decodeOriginsU Used function for the decoding of genetic origins [[5]]/[[6]]
```

Value

Inherited parent gamete

Examples

bv.development

Development of genetic/breeding value

Description

Function to plot genetic/breeding values for multiple generation/cohorts

Usage

```
bv.development(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  confidence = c(1, 2, 3),
  development = c(1, 2, 3),
  quantile = 0.95,
  bvrow = "all",
  ignore.zero = TRUE,
  json = FALSE,
  display.time.point = FALSE,
```

bv.development 29

```
display.creating.type = FALSE,
  display.cohort.name = FALSE,
  display.sex = FALSE,
  equal.spacing = FALSE,
  time_reorder = FALSE,
  display.line = TRUE,
  ylim = NULL,
  fix_mfrow = FALSE
)
```

Arguments

population	population list				
database	Groups of individuals to consider for the export				
gen	Quick-insert for database (vector of all generations to export)				
cohorts	Quick-insert for database (vector of names of cohorts to export)				
confidence	Draw confidence intervals for (1- bv, 2- bve, 3- pheno; default: c(1,2,3))				
development	Include development of (1- bv, 2- bve, 3- pheno; default: c(1,2,3))				
quantile	Quantile of the confidence interval to draw (default: 0.05)				
bvrow	Which traits to display (for multiple traits separate plots (par(mfrow)))				
ignore.zero	Cohorts with only 0 individuals are not displayed (default: TRUE)				
json	If TRUE extract which cohorts to plot according to the json-file used in json.simulation				
display.time.pd	pint				
	Set TRUE to use time point of generated to sort groups				
display.creating.type					
	Set TRUE to show Breedingtype used in generation (web-interface)				
display.cohort.name					
	Set TRUE to display the name of the cohort in the x-axis				
display.sex	Set TRUE to display the creating.type (Shape of Points - web-based-application)				
equal.spacing	Equal distance between groups (independent of time.point)				
time_reorder	Set TRUE to order cohorts according to the time point of generation				
display.line	Set FALSE to not display the line connecting cohorts				
ylim	Set this to fix the y-axis of the plot				
fix_mfrow	Set TRUE to not use mfrow - use for custom plots				

Value

Genomic values of selected gen/database/cohort

Examples

```
data(ex_pop)
bv.development(ex_pop, gen=1:5)
```

30 bv.development.box

bv.development.box

Development of genetic/breeding value using a boxplot

Description

Function to plot genetic/breeding values for multiple generation/cohorts using box plots

Usage

```
bv.development.box(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  bvrow = "all",
   json = FALSE,
  display = "bv",
  display.selection = FALSE,
  display.reproduction = FALSE,
  ylim = NULL,
  fix_mfrow = FALSE
)
```

Arguments

population population list database Groups of individuals to consider for the export gen Quick-insert for database (vector of all generations to export) Quick-insert for database (vector of names of cohorts to export) cohorts bvrow Which traits to display (for multiple traits separte plots (par(mfrow))) If TRUE extract which cohorts to plot according to the json-file used in json.simulation json Choose between "bv", "pheno", "bve" (default: "bv") display display.selection Display lines between generated cohorts via selection (webinterface) display.reproduction Display lines between generated cohorts via reproduction (webinterface) Set this to fix the y-axis of the plot ylim fix_mfrow Set TRUE to not use mfrow - use for custom plots

Value

Genomic values of selected gen/database/cohort

bv.standardization 31

Examples

```
data(ex_pop)
bv.development.box(ex_pop, gen=1:5)
```

bv.standardization

BV standardization

Description

Function to get mean and genetic variance of a trait to a fixed value

Usage

```
bv.standardization(
  population,
  mean.target = 100,
  var.target = 10,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  adapt.bve = FALSE,
  adapt.pheno = FALSE,
  verbose = FALSE
```

Arguments

population	Population list
mean.target	Target mean
var.target	Target variance
gen	Quick-insert for database (vector of all generations to export)
database	Groups of individuals to consider for the export
cohorts	Quick-insert for database (vector of names of cohorts to export)
adapt.bve	Modify previous breeding value estimations by scaling (default: FALSE)
adapt.pheno	Modify previous phenotypes by scaling (default: FALSE)
verbose	Set to TRUE to display prints

Value

Population-list with scaled QTL-effects

Examples

```
population <- creating.diploid(nsnp=1000, nindi=100, n.additive=100)
population <- bv.standardization(population, mean.target=200, var.target=5)</pre>
```

32 calculate.by

calculate.bv

Calculate breeding values

Description

Internal function to calculate the breeding value of a given individual

Usage

```
calculate.bv(
  population,
  gen,
  sex,
  nr,
  activ_bv,
  import.position.calculation = NULL,
  decodeOriginsU = decodeOriginsR,
  store.effect.freq = FALSE,
  bit.storing = FALSE,
  nbits = 30,
  output_compressed = FALSE,
  bv.ignore.traits = NULL
)
```

Arguments

population	Population list
gen	Generation of the individual of interest
sex	Sex of the individual of interest
nr	Number of the individual of interest
activ_bv import.position	traits to consider
	Function to calculate recombination point into adjacent/following SNP
decodeOriginsU	Used function for the decoding of genetic origins [[5]]/[[6]]
store.effect.fr	req
	If TRUE store the allele frequency of effect markers per generation
bit.storing	Set to TRUE if the MoBPS (not-miraculix! bit-storing is used)
nbits	Bits available in MoBPS-bit-storing
output_compress	sed
	Set to TRUE to get a miraculix-compressed genotype/haplotype
bv.ignore.trait	ts
	Vector of traits to ignore in the calculation of the genomic value (default: NULL; Only recommended for high number of traits and experienced users!)

cattle_chip 33

Value

[1] true genomic value [[2]] allele frequency at QTL markers

Examples

```
data(ex_pop)
calculate.bv(ex_pop, gen=1, sex=1, nr=1, activ_bv = 1)
```

cattle_chip

Cattle chip

Description

Genome for cattle according to Ma et al.

Usage

```
cattle_chip
```

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

Ma et al 2015

check.parents

Relatedness check between two individuals

Description

Internal function to check the relatedness between two individuals

Usage

```
check.parents(population, info.father, info.mother, max.rel = 2)
```

Arguments

population	Population list
info.father	position of the first parent in the dataset
info.mother	position of the second parent in the dataset
max.rel	maximal allowed relationship (default: 2, alt: 1 no full-sibs, 0 no half-sibs)

34 clean.up

Value

logical with TRUE if relatedness does not exceed max.rel / FALSE otherwise.

Examples

```
\label{lem:data} \begin{array}{ll} \mbox{data(ex\_pop)} \\ \mbox{check.parents(ex\_pop, info.father=c(4,1,1,1), info.mother=c(4,2,1,1))} \end{array}
```

chicken_chip

chicken chip

Description

Genome for chicken according to Groenen et al.

Usage

```
chicken_chip
```

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

Groenen et al 2009

clean.up

Clean-up recombination points

Description

Function to remove recombination points + origins with no influence on markers

Usage

```
clean.up(population, gen = "all", database = NULL, cohorts = NULL)
```

Arguments

population Population list

gen Generations to clean up (default: "current")

database Groups of individuals to consider

cohorts Quick-insert for database (vector of names of cohorts to export)

codeOriginsR 35

Value

Population-list with deleted irrelevant recombination points

Examples

```
data(ex_pop)
ex_pop <- clean.up(ex_pop)</pre>
```

codeOriginsR

Origins-coding(R)

Description

R-Version of the internal bitwise-coding of origins

Usage

```
codeOriginsR(M)
```

Arguments

Μ

Origins matrix

Value

Bit-wise coded origins

Examples

```
codeOriginsR(cbind(1,1,1,1))
```

 ${\tt combine.traits}$

Combine traits

Description

Function to combine traits in the BVE

Usage

```
combine.traits(
  population,
  combine.traits = NULL,
  combine.name = NULL,
  remove.combine = NULL,
  remove.all = FALSE
)
```

36 compute.costs

Arguments

```
population Population list

combine.traits Vector containing the traits (numbers) to combine into a joined trait

combine.name Name of the combined trait

remove.combine Remove a selected previously generated combined trait

set TRUE to remove all previously generated combined traits
```

Value

Population-list

Examples

```
population <- creating.diploid(nsnp=100, nindi=100, n.additive = c(50,50))
population <- combine.traits(population, combine.traits=1:2)
population <- breeding.diploid(population, bve=TRUE, phenotyping.gen=1, heritability=0.3)</pre>
```

compute.costs

Compute costs of a breeding program

Description

Function to derive the costs of a breeding program / population-list

Usage

```
compute.costs(
  population,
  phenotyping.costs = 10,
  genotyping.costs = 100,
  fix.costs = 0,
  fix.costs.annual = 0,
  profit.per.bv = 1,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  interest.rate = 1,
  base.gen = 1
)
```

Arguments

```
population population-list phenotyping.costs
```

Costs for the generation of a phenotype

compute.costs.cohorts 37

interest.rate Applied yearly interest rate

base generation (application of interest rate)

Value

Cost-table for selected gen/database/cohorts of a population-list

Examples

```
data(ex_pop)
compute.costs(ex_pop, gen=1:5)
```

Description

Function to derive the costs of a breeding program / population-list by cohorts

```
compute.costs.cohorts(
  population,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  json = TRUE,
  phenotyping.costs = NULL,
  genotyping.costs = 0,
  housing.costs = NULL,
  fix.costs = 0,
  fix.costs.annual = 0,
  profit.per.bv = 1,
  interest.rate = 1,
  verbose = TRUE
)
```

38 compute.snps

Arguments

population population-list

gen Quick-insert for database (vector of all generations to consider)

database Groups of individuals to consider

cohorts Quick-insert for database (vector of names of cohorts to consider)

json If TRUE extract which cohorts to plot according to the json-file used in json.simulation

phenotyping.costs

Costs for the generation of a phenotype

genotyping.costs

Costs for the geneation of a genotype

housing.costs Costs for housing

fix.costs one time occuring fixed costs

fix.costs.annual

annually occuring fixed costs

profit.per.bv profit generated by bv per animal

interest.rate Applied yearly interest rate

verbose Set to FALSE to not display any prints

Value

Cost-table for selected gen/database/cohorts of a population-list

Examples

```
data(ex_pop)
compute.costs.cohorts(ex_pop, gen=1:5, genotyping.costs=25, json=FALSE)
```

compute.snps

Compute genotype/haplotype

Description

Internal function for the computation of genotypes & haplotypes

```
compute.snps(
  population,
  gen,
  sex,
  nr,
  faster = TRUE,
  import.position.calculation = NULL,
  from_p = 1,
```

compute.snps_single 39

```
to_p = Inf,
  decodeOriginsU = decodeOriginsR,
  bit.storing = FALSE,
  nbits = 30,
  output_compressed = FALSE
)
```

Arguments

population	Population list	
gen	Generation of the individual to compute	
sex	Gender of the individual to compute	
nr	Number of the individual to compute	
faster	If FALSE use slower version to compute markers between recombination points	
import.position	calculation Function to calculate recombination point into adjacent/following SNP	
from_p	First SNP to consider	
to_p	Last SNP to consider	
decodeOriginsU	Used function for the decoding of genetic origins [[5]]/[[6]]	
bit.storing	Set to TRUE if the MoBPS (not-miraculix! bit-storing is used)	
nbits	Bits available in MoBPS-bit-storing	
output_compressed		
	Set to TRUE to get a miraculix-compressed genotype/haplotype	

Value

haplotypes for the selected individual

Examples

```
data(ex_pop)
compute.snps(ex_pop, gen=1, sex=1, nr=1)
```

compute.snps_single

Compute genotype/haplotype in gene editing application

Description

Internal function for the computation of genotypes & haplotypes in gene editing application

Usage

```
compute.snps_single(
  population,
  current.recombi,
  current.mut,
  current.ursprung,
  faster = TRUE,
  import.position.calculation = NULL,
  decodeOriginsU = decodeOriginsR
)
```

Arguments

```
population Population list
current.recombi
vector of currently activ recombination points

current.mut vector of currently activ mutations
current.ursprung
vector of currently activ origins

faster If FALSE use slower version to compute markers between recombination points import.position.calculation
Function to calculate recombination point into adjacent/following SNP

decodeOriginsU Used function for the decoding of genetic origins [[5]]/[[6]]
```

Value

haplotypes for the selected individual

creating.diploid

Generation of the starting population

Description

Generation of the starting population

```
creating.diploid(
  dataset = NULL,
  vcf = NULL,
  chr.nr = NULL,
  bp = NULL,
  snp.name = NULL,
  hom0 = NULL,
  hom1 = NULL,
```

```
bpcm.conversion = 0,
nsnp = 0,
nindi = 0,
freq = "beta",
population = NULL,
sex.s = "fixed",
add.chromosome = FALSE,
generation = 1,
class = 0L,
sex.quota = 0.5,
chromosome.length = NULL,
length.before = 5,
length.behind = 5,
real.bv.add = NULL,
real.bv.mult = NULL,
real.bv.dice = NULL,
snps.equidistant = NULL,
change.order = FALSE,
bv.total = 0,
polygenic.variance = 100,
bve.mult.factor = NULL,
bve.poly.factor = NULL,
base.bv = NULL,
add.chromosome.ends = TRUE,
new.phenotype.correlation = NULL,
new.residual.correlation = NULL,
new.breeding.correlation = NULL,
add.architecture = NULL,
snp.position = NULL,
position.scaling = FALSE,
bit.storing = FALSE,
nbits = 30,
randomSeed = NULL,
miraculix = TRUE,
miraculix.dataset = TRUE,
n.additive = 0,
n.equal.additive = 0,
n.dominant = 0,
n.equal.dominant = 0,
n.qualitative = 0,
n.quantitative = 0,
dominant.only.positive = FALSE,
var.additive.l = NULL,
var.dominant.1 = NULL,
var.qualitative.l = NULL,
var.quantitative.l = NULL,
effect.size.equal.add = 1,
effect.size.equal.dom = 1,
```

```
exclude.snps = NULL,
  replace.real.bv = FALSE,
  shuffle.traits = NULL,
  shuffle.cor = NULL,
  skip.rest = FALSE,
  enter.bv = TRUE,
  name.cohort = NULL,
  template.chip = NULL,
  beta.shape1 = 1,
  beta.shape2 = 1,
  time.point = 0,
  creating.type = 0,
  trait.name = NULL,
  share.genotyped = 1,
  genotyped.s = NULL,
  map = NULL,
  remove.invalid.qtl = TRUE,
  verbose = TRUE,
  bv.standard = FALSE,
 mean.target = NULL,
  var.target = NULL,
  is.maternal = NULL,
  is.paternal = NULL,
  vcf.maxsnp = Inf,
  internal = FALSE
)
```

Arguments

dataset SN	√P dataset, ι	ıse "random"	, "allhetero" "all0	" when generating a	a dataset via
------------	---------------	--------------	---------------------	---------------------	---------------

nsnp,nindi

vcf Path to a vcf-file used as input genotypes (correct haplotype phase is assumed!)

chr.nr Vector containing the assosiated chromosome for each marker (default: all on

the same)

bp Vector containing the physical position (bp) for each marker (default: 1,2,3...)

snp.name Vector containing the name of each marker (default ChrXSNPY - XY chosen

accordingly)

hom0 Vector containing the first allelic variant in each marker (default: 0)
hom1 Vector containing the second allelic variant in each marker (default: 1)

bpcm.conversion

Convert physical position (bp) into a cM position (default: 0 - not done)

nsnp number of markers to generate in a random dataset

nindi number of inidividuals to generate in a random dataset

freq frequency of allele 1 when randomly generating a dataset

population Population list

Specify which newly added individuals are male (1) or female (2) sex.s If TRUE add an additional chromosome to the dataset add.chromosome generation Generation of the newly added individuals (default: 1) class Migration level of the newly added individuals sex.quota Share of newly added female individuals (deterministic if sex.s="fixed", alt: sex.s="random") chromosome.length Length of the newly added chromosome (default: 5) Length before the first SNP of the dataset (default: 5) length.before length.behind Length after the last SNP of the dataset (default: 5) real.bv.add Single Marker effects real.bv.mult Two Marker effects real.bv.dice Multi-marker effects snps.equidistant Use equidistant markers (computationally faster!; default: TRUE) change.order If TRUE sort markers according to given marker positions bv.total Number of traits (If more than traits via real.bv.X use traits with no directly underlying QTL) polygenic.variance Genetic variance of traits with no underlying QTL bve.mult.factor Multiplicate trait value times this bve.poly.factor Potency trait value over this base.bv Average genetic value of a trait add.chromosome.ends Add chromosome ends as recombination points new.phenotype.correlation (OLD! - use new.residual.correlation) Correlation of the simulated environmental variance new.residual.correlation Correlation of the simulated environmental variance new.breeding.correlation Correlation of the simulated genetic variance (child share! heritage is not influadd.architecture Add genetic architecture (marker positions) snp.position Location of each marker on the genetic map position.scaling Manual scaling of snp.position Set to TRUE if the MoBPS (not-miraculix! bit-storing is used) bit.storing

Bits available in MoBPS-bit-storing

nbits

Set random seed of the process randomSeed miraculix If TRUE use miraculix package for data storage, computations and dataset generation miraculix.dataset Set FALSE to deactive miraculix package for dataset generation n.additive Number of additive QTL with effect size drawn from a gaussian distribution n.equal.additive Number of additive QTL with equal effect size (effect.size) n.dominant Number of dominant QTL with effect size drawn from a gaussian distribution n.equal.dominant Number of n.equal.dominant QTL with equal effect size n. qualitative Number of qualitative epistatic QTL n. quantitative Number of quantitative epistatic QTL dominant.only.positive Set to TRUE to always asign the heterozygous variant with the higher of the two homozygous effects (e.g. hybrid breeding); default: FALSE var.additive.l Variance of additive QTL var.dominant.l Variance of dominante OTL var.qualitative.l Variance of qualitative epistatic QTL var.quantitative.l Variance of quantitative epistatic QTL effect.size.equal.add Effect size of the QTLs in n.equal.additive effect.size.equal.dom Effect size of the QTLs in n.equal.dominant exclude.snps Marker were no QTL are simulated on replace.real.bv If TRUE delete the simulated traits added before shuffle.traits Combine different traits into a joined trait shuffle.cor Target Correlation between shuffeled traits skip.rest Internal variable needed when adding multipe chromosomes jointly enter.bv Internal parameter name.cohort Name of the newly added cohort template.chip Import genetic map and chip from a species ("cattle", "chicken", "pig") First parameter of the beta distribution for simulating allele frequencies beta.shape1

Second parameter of the beta distribution for simulating allele frequencies

Technique to generate new individuals (usage in web-based application)

Time point at which the new individuals are generated

Name of the trait generated

beta.shape2

time.point

creating.type
trait.name

share.genotyped

Share of individuals genotyped in the founders

genotyped.s Specify with newly added individuals are genotyped (1) or not (0)

map map-file that contains up to 5 colums (Chromsome, SNP-id, M-position, Bp-

position, allele freq - Everything not provides it set to NA). A map can be im-

ported via MoBPSmaps::ensembl.map()

remove.invalid.qtl

Set to FALSE to deactive the automatic removal of QTLs on markers that do not

exist

verbose Set to FALSE to not display any prints

bv.standard Set TRUE to standardize trait mean and variance via bv.standardization() - au-

tomatically set to TRUE when mean/var.target are used

mean.target Target mean
var.target Target variance

is.maternal Vector coding if a trait is caused by a maternal effect (Default: all FALSE)

is.paternal Vector coding if a trait is caused by a paternal effect (Default: all FALSE)

vcf.maxsnp Maximum number of SNPs to include in the genotype file (default: Inf)

internal Dont touch!

Value

Population-list

Examples

```
population <- creating.diploid(nsnp=1000, nindi=100)</pre>
```

 ${\tt creating.phenotypic.transform}$

Create a phenotypic transformation

Description

Function to perform create a transformation of phenotypes

```
creating.phenotypic.transform(
  population,
  phenotypic.transform.function = NULL,
  trait = 1
)
```

46 creating.trait

Arguments

population

```
phenotypic.transform.function
Phenotypic transformation to apply

trait
Trait for which a transformation is to be applied data(ex_pop) trafo <- function(x) return(x^2) ex_pop <- creating.phenotypic.transform(ex_pop, phenotypic.transform.function=traf
```

Value

Population-list with a new phenotypic transformation function

Population list

creating.trait

Generation of genomic traits

Description

Generation of the trait in a starting population

```
creating.trait(
  population,
  real.bv.add = NULL,
  real.bv.mult = NULL,
  real.bv.dice = NULL,
  bv.total = 0,
  polygenic.variance = 100,
  bve.mult.factor = NULL,
  bve.poly.factor = NULL,
  base.bv = NULL,
  new.phenotype.correlation = NULL,
  new.residual.correlation = NULL,
  new.breeding.correlation = NULL,
  n.additive = 0,
  n.equal.additive = 0,
  n.dominant = 0,
  n.equal.dominant = 0,
  n.qualitative = 0,
  n.quantitative = 0,
  dominant.only.positive = FALSE,
  var.additive.l = NULL,
  var.dominant.1 = NULL,
  var.qualitative.l = NULL,
  var.quantitative.1 = NULL,
  effect.size.equal.add = 1,
  effect.size.equal.dom = 1,
```

47 creating.trait

```
exclude.snps = NULL,
      randomSeed = NULL,
      shuffle.traits = NULL,
      shuffle.cor = NULL,
      replace.traits = FALSE,
      trait.name = NULL,
      remove.invalid.qtl = TRUE,
      bv.standard = FALSE,
      mean.target = NULL,
      var.target = NULL,
      verbose = TRUE,
      is.maternal = NULL,
      is.paternal = NULL
    )
Arguments
                     Population list
    population
    real.bv.add
                     Single Marker effects
    real.bv.mult
                     Two Marker effects
    real.bv.dice
                     Multi-marker effects
    bv.total
                     Number of traits (If more than traits via real.bv.X use traits with no directly
                     underlying QTL)
    polygenic.variance
                      Genetic variance of traits with no underlying QTL
    bve.mult.factor
                     Multiplicate trait value times this
    bve.poly.factor
                     Potency trait value over this
    base.bv
                     Average genetic value of a trait
    new.phenotype.correlation
                     (OLD! - use new.residual.correlation) Correlation of the simulated environmental
                     variance
    new.residual.correlation
                     Correlation of the simulated environmental variance
    new.breeding.correlation
                     Correlation of the simulated genetic variance (child share! heritage is not influ-
                     enced!
    n.additive
                     Number of additive QTL with effect size drawn from a gaussian distribution
    n.equal.additive
                     Number of additive QTL with equal effect size (effect.size)
    n.dominant
                     Number of dominant QTL with effect size drawn from a gaussian distribution
    n.equal.dominant
                     Number of n.equal.dominant QTL with equal effect size
                     Number of qualitative epistatic QTL
```

n.qualitative

48 creating.trait

```
n. quantitative Number of quantitative epistatic QTL
dominant.only.positive
                  Set to TRUE to always asign the heterozygous variant with the higher of the two
                  homozygous effects (e.g. hybrid breeding); default: FALSE
var.additive.l Variance of additive QTL
var.dominant.l Variance of dominante QTL
var.qualitative.l
                  Variance of qualitative epistatic QTL
var.quantitative.l
                  Variance of quantitative epistatic QTL
effect.size.equal.add
                  Effect size of the QTLs in n.equal.additive
effect.size.equal.dom
                  Effect size of the QTLs in n.equal.dominant
exclude.snps
                  Marker were no QTL are simulated on
randomSeed
                  Set random seed of the process
shuffle.traits Combine different traits into a joined trait
shuffle.cor
                  Target Correlation between shuffeled traits
replace.traits If TRUE delete the simulated traits added before
trait.name
                  Name of the trait generated
remove.invalid.qtl
                  Set to FALSE to deactive the automatic removal of QTLs on markers that do not
                  exist
bv.standard
                  Set TRUE to standardize trait mean and variance via by.standardization()
mean.target
                  Target mean
var.target
                  Target variance
verbose
                  Set to FALSE to not display any prints
                  Vector coding if a trait is caused by a maternal effect (Default: all FALSE)
is.maternal
is.paternal
                  Vector coding if a trait is caused by a paternal effect (Default: all FALSE)
```

Value

Population-list with one or more additional new traits

```
population <- creating.diploid(nsnp=1000, nindi=100)
population <- creating.trait(population, n.additive=100)</pre>
```

decodeOriginsR 49

decodeOriginsR

Origins-Decoding(R)

Description

R-Version of the internal bitwise-decoding of origins

Usage

```
decodeOriginsR(P, row)
```

Arguments

P coded origins vector row row to decode

Value

de-coded origins

Examples

decodeOriginsR(0L)

demiraculix

Remove miraculix-coding for genotypes

Description

Internal function to decode all genotypes to non-miraculix objects

Usage

```
demiraculix(population)
```

Arguments

population

Population list

Value

Population list

```
# This is only relevant with the package miraculix is installed and used
population <- creating.diploid(nsnp=100, nindi=50)
population <- demiraculix(population)</pre>
```

50 derive.loop.elements

Description

Internal function to derive the position of all individuals to consider for BVE/GWAS

Usage

```
derive.loop.elements(
  population,
  bve.database,
  bve.class,
  bve.avoid.duplicates,
  store.adding = FALSE,
  store.which.adding = FALSE,
  list.of.copys = FALSE
)
```

Arguments

population	Population list	
bve.database	Groups of individuals to consider in breeding value estimation	
bve.class	Consider only animals of those class classes in breeding value estimation (default: NULL - use all)	
bve.avoid.duplicates		
	If set to FALSE multiple generatations of the same individual can be used in the bye (only possible by using copy.individual to generate individuals)	
store.adding	Internal parameter to derive number of added individuals per database entry (only relevant internally for GWAS)	
store.which.adding		
	Internal parameter to derive which individuals are copy entries	
list.of.copys	Internal parameter to derive further information on the copies individuals	

Value

Matrix of individuals in the entered database

```
data(ex_pop)
derive.loop.elements(ex_pop, bve.database=get.database(ex_pop, gen=2),
bve.class=NULL, bve.avoid.duplicates=TRUE)
```

diag.mobps 51

diag.mobps

Add a genotyping array

Description

Function to add a genotyping array for the population

Usage

```
diag.mobps(elements)
```

Arguments

elements

vector with entries to put on the diagonal of a matrix

Value

Diagonal matrix

Examples

diag.mobps(5)

edges.fromto

Detection of parental/child nodes

Description

Internal function to extract parental/child node of an edge

Usage

```
edges.fromto(edges)
```

Arguments

edges

Edges of the json-file generated via the web-interface

Value

Matrix of Parent/Child-nodes for the considered edges

52 edit_animal

edit_animal

Internal gene editing function

Description

Internal function to perform gene editing

Usage

```
edit_animal(
  population,
  gen,
  sex,
  nr,
  nr.edits,
  decodeOriginsU = decodeOriginsR,
  bit.storing = FALSE,
  nbits = 30
)
```

Arguments

population	Population list
gen	Generation of the individual to edit
sex	Gender of the individual to edit
nr	Number of the individual to edit
nr.edits	Number of edits to perform
decodeOriginsU	Used function for the decoding of genetic origins [[5]]/[[6]]
bit.storing	Set to TRUE if the MoBPS (not-miraculix! bit-storing is used)
nbits	Bits available in MoBPS-bit-storing

Value

animal after genome editing

effect.estimate.add 53

effect.estimate.add Estimation of marker effects

Description

Function to estimate marker effects

Usage

```
effect.estimate.add(geno, pheno, map = NULL, scaling = TRUE)
```

Arguments

geno genotype dataset (marker x individuals)
pheno phenotype dataset (each phenotype in a row)

map genomic map

scaling Set FALSE to not perform variance scaling

Value

Empirical kinship matrix (IBD-based since Founders)

Examples

```
data(ex_pop)
pheno <- get.pheno(ex_pop, gen=1:5)
geno <- get.geno(ex_pop, gen=1:5)
map <- get.map(ex_pop, use.snp.nr=TRUE)
real.bv.add <- effect.estimate.add(geno, pheno, map)</pre>
```

effective.size

Estimate effective population size

Description

Internal function to estimate the effective population size

Usage

```
effective.size(ld, dist, n)
```

Arguments

ld ld between markers

dist distance between markers in Morgan

n Population size

54 ex_json

Value

Estimated effective population size

epi

Martini-Test function

Description

Internal function to perform martini test

Usage

$$epi(y, Z, G = NULL)$$

Arguments

y y

Z genomic information matrix

G kinship matrix

Value

Estimated breeding values

 ex_json

ex_json

Description

Exemplary json-data

Usage

 ex_json

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

Web-interface

ex_pop 55

Description

Exemplary population-list

Usage

ex_pop

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

MoBPS

find.chromo

Position detection (chromosome)

Description

Internal function for the detection on which chromosome each marker is

Usage

```
find.chromo(position, length.total)
```

Arguments

position position in the genome

length.total Length of each chromosome

Value

Chromosome the marker is part of

56 founder.simulation

find.snpbefore

Position detection (SNPs)

Description

Internal function for the detection on which position each marker is

Usage

```
find.snpbefore(position, snp.position)
```

Arguments

position Position on the genome snp.position Position of the SNPs on the genome

Value

SNP-position of the target position

founder.simulation

Founder simulation

Description

Function to generate founder genotypes

```
founder.simulation(
 nindi = 100,
  sex.quota = 0.5,
 nsnp = 0,
 n.gen = 100,
  nfinal = NULL,
  sex.quota.final = NULL,
 big.output = FALSE,
 plot = TRUE,
 display.progress = TRUE,
 depth.pedigree = 7,
 dataset = NULL,
  vcf = NULL,
  chr.nr = NULL,
  bp = NULL,
  snp.name = NULL,
```

founder.simulation 57

```
hom0 = NULL,
  hom1 = NULL,
  bpcm.conversion = 0,
  freq = "beta",
  sex.s = "fixed"
  chromosome.length = NULL,
  length.before = 5,
  length.behind = 5,
  snps.equidistant = NULL,
  change.order = FALSE,
  snp.position = NULL,
  position.scaling = FALSE,
  bit.storing = FALSE,
  nbits = 30,
  randomSeed = NULL,
  miraculix = TRUE,
 miraculix.dataset = TRUE,
  template.chip = NULL,
  beta.shape1 = 1,
  beta.shape2 = 1,
 map = NULL,
  verbose = TRUE,
  vcf.maxsnp = Inf
)
```

Arguments

nindi number of inidividuals to generate in a random dataset

sex.quota Share of newly added female individuals (deterministic if sex.s="fixed", alt:

sex.s="random")

nsnp number of markers to generate in a random dataset

n.gen Number of generations to simulate (default: 100)

nfinal Number of final individuals to include (default: nindi)

sex.quota.final

Share of female individuals in the final generation

big.output Set to TRUE to export map, population list and pedigree relationship

plot Set to FALSE to not generate LD-decay plot and allele frequency spectrum

display.progress

Set FALSE to not display progress bars. Setting verbose to FALSE will auto-

matically deactive progress bars

depth.pedigree Depth of the pedigree in generations (default: 7)

dataset SNP dataset, use "random", "allhetero" "all0" when generating a dataset via

nsnp,nindi

Path to a vcf-file used as input genotypes (correct haplotype phase is assumed!)

chr.nr Vector containing the assosiated chromosome for each marker (default: all on

the same)

58 founder.simulation

bp Vector containing the physical position (bp) for each marker (default: 1,2,3...)

snp.name Vector containing the name of each marker (default ChrXSNPY - XY chosen

accordingly)

hom0 Vector containing the first allelic variant in each marker (default: 0)
hom1 Vector containing the second allelic variant in each marker (default: 1)

bpcm.conversion

Convert physical position (bp) into a cM position (default: 0 - not done)

freq frequency of allele 1 when randomly generating a dataset

sex.s Specify which newly added individuals are male (1) or female (2)

chromosome.length

Length of the newly added chromosome (default: 5)

length.before Length before the first SNP of the dataset (default: 5)
length.behind Length after the last SNP of the dataset (default: 5)

snps.equidistant

Use equidistant markers (computationally faster!; default: TRUE)

change.order If TRUE sort markers according to given marker positions

snp.position Location of each marker on the genetic map

position.scaling

Manual scaling of snp.position

bit.storing Set to TRUE if the MoBPS (not-miraculix! bit-storing is used)

nbits Bits available in MoBPS-bit-storing

randomSeed Set random seed of the process

miraculix If TRUE use miraculix package for data storage, computations and dataset gen-

eration

miraculix.dataset

Set FALSE to deactive miraculix package for dataset generation

template.chip Import genetic map and chip from a species ("cattle", "chicken", "pig")
beta.shape1 First parameter of the beta distribution for simulating allele frequencies
beta.shape2 Second parameter of the beta distribution for simulating allele frequencies

map map-file that contains up to 5 colums (Chromsome, SNP-id, M-position, Bp-

position, allele freq - Everything not provides it set to NA). A map can be im-

ported via MoBPSmaps::ensembl.map()

verbose Set to FALSE to not display any prints

vcf.maxsnp Maximum number of SNPs to include in the genotype file (default: Inf)

Examples

population <- founder.simulation(nindi=100, nsnp=1000, n.gen=5)</pre>

generation.individual 59

generation.individual Function to generate a new individual

Description

Function to generate a new individual

Usage

```
generation.individual(
  indexb,
  population,
  info_father_list,
  info_mother_list,
  copy.individual,
 mutation.rate,
  remutation.rate,
  recombination.rate,
  recom.f.indicator,
  duplication.rate,
  duplication.length,
  duplication.recombination,
  delete.same.origin,
  gene.editing,
  nr.edits,
  gen.architecture.m,
  gen.architecture.f,
  decodeOriginsU,
  current.gen,
  save.recombination.history,
  new.bv.child,
  dh.mating,
  share.genotyped,
  added.genotyped,
  genotyped.array,
  dh.sex,
  n.observation
)
```

Arguments

60 generation.individual

copy.individual

windows parallel internal test

mutation.rate windows parallel internal test

remutation.rate

windows parallel internal test

recombination.rate

windows parallel internal test

recom.f.indicator

windows parallel internal test

duplication.rate

windows parallel internal test

duplication.length

windows parallel internal test

duplication.recombination

windows parallel internal test

delete.same.origin

windows parallel internal test

gene.editing windows parallel internal test nr.edits windows parallel internal test

gen.architecture.m

windows parallel internal test

gen.architecture.f

windows parallel internal test

decodeOriginsU windows parallel internal test

windows parallel internal test current.gen

save.recombination.history

windows parallel internal test

new.bv.child windows parallel internal test

windows parallel internal test share.genotyped

dh.mating

windows parallel internal test

added.genotyped

windows parallel internal test

genotyped.array

windows parallel internal test

dh.sex windows parallel internal test windows parallel internal test n.observation

Value

Offspring individual

get.admixture 61

Description

Function to generate admixture plots

Usage

```
get.admixture(
  population,
  geno = NULL,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  d = NULL,
  verbose = TRUE,
  plot = TRUE,
  sort = FALSE,
  sort.cutoff = 0.01
)
```

Arguments

population	Population list
geno	Manually provided genotype dataset to use instead of gen/database/cohorts
gen	Quick-insert for database (vector of all generations to consider)
database	Groups of individuals to consider
cohorts	Quick-insert for database (vector of names of cohorts to consider)
d	dimensions to consider in admixture plot (default: automatically estimate a reasonable number)
verbose	Set to FALSE to not display any prints
plot	Set to FALSE to not generate an admixture plot
sort	Set to TRUE to sort individuals according to contributes from the first dimension
sort.cutoff	Skip individuals with contributions under this threshold (and use next dimension instead) data(ex_pop) get.admixture(ex_pop, gen=4:6, d=2, sort=TRUE)

Value

Matrix with admixture proportion

62 get.age.point

Description

Function to devide age point for each individual (Same as time.point unless copy.individual is used for aging)

Usage

```
get.age.point(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Time point selected gen/database/cohorts-individuals are born

```
data(ex_pop)
get.age.point(ex_pop, gen=2)
```

get.bv 63

get.bv

Export underlying true breeding values

Description

Function to export underlying true breeding values

Usage

```
get.bv(population, database = NULL, gen = NULL, cohorts = NULL, use.id = FALSE)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

Value

Genomic value of in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.bv(ex_pop, gen=2)
```

get.bve

Export estimated breeding values

Description

Function to export estimated breeding values

```
get.bve(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

64 get.class

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Estimated breeding value of in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.bve(ex_pop, gen=2)
```

get.class

Derive class

Description

Function to devide the class for each individual

Usage

```
get.class(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Class of in gen/database/cohorts selected individuals

get.cohorts 65

Examples

```
data(ex_pop)
get.class(ex_pop, gen=2)
```

get.cohorts

Export Cohort-names

Description

Function to export cohort names for the population list

Usage

```
get.cohorts(population, extended = FALSE)
```

Arguments

population Population list extended extended cohorts

Value

List of all cohorts in the population-list

Examples

```
data(ex_pop)
get.cohorts(ex_pop)
```

get.creating.type

Derive creating type

Description

Function to devide creating type for each individual

```
get.creating.type(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

get.cullingtime

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

Value

Creating type of in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.creating.type(ex_pop, gen=2)
```

get.cullingtime

Derive time of culling

Description

Function to devide the time of culling for all individuals

Usage

```
get.cullingtime(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

Value

Time of death of in gen/database/cohorts selected individuals

get.database 67

Examples

```
data(ex_pop)
get.cullingtime(ex_pop, gen=2)
```

get.database

gen/database/cohorts conversion

Description

Function to derive a database based on gen/database/cohorts

Usage

```
get.database(
  population,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  avoid.merging = FALSE
)
```

Arguments

population Population list

gen Quick-insert for database (vector of all generations to export)

database Groups of individuals to consider for the export

cohorts Quick-insert for database (vector of names of cohorts to export)

avoid.merging Set to TRUE to avoid different cohorts to be merged in a joint group when

possible

Value

Combine gen/database/cohorts to a joined database

```
data(ex_pop)
get.database(ex_pop, gen=2)
```

68 get.dendrogram

get.death.point

Derive death point

Description

Function to devide the time of death for each individual (NA for individuals that are still alive))

Usage

```
get.death.point(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Time of death of in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.death.point(ex_pop, gen=2)
```

get.dendrogram

Dendrogram

Description

Function calculate a dendogram

69

Usage

```
get.dendrogram(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  method = NULL,
  individual.names = NULL)
```

Arguments

population Population list

path provide a path if the dendrogram would be saved as a png-file

database Groups of individuals to consider

gen Quick-insert for database (vector of all generations to consider)

cohorts Quick-insert for database (vector of names of cohorts to consider)

method Method used to calculate genetic distances (default: "Nei", alt: "Rogers", "Pre-

vosti", "Modified Rogers"

individual.names

Names of the individuals in the database ((default are MoBPS internal names

based on position))

Value

Dendrogram plot for genotypes

Examples

```
data(ex_pop)
get.dendrogram(ex_pop, gen=2)
```

```
\verb"get.dendrogram.heatmap"
```

Dendrogram Heatmap

Description

Function calculate a dendogram

Usage

```
get.dendrogram.heatmap(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
 method = NULL,
  individual.names = NULL,
  traits = NULL,
  type = "pheno"
)
```

Arguments

	population	Population list	
	path	provide a path if the dendrogram would be saved as a png-file	
	database	Groups of individuals to consider	
	gen	Quick-insert for database (vector of all generations to consider)	
	cohorts	Quick-insert for database (vector of names of cohorts to consider)	
	method	Method used to calculate genetic distances (default: "Nei", alt: "Rogers", "Prevosti", "Modified Rogers"	
individual.names			
		Names of the individuals in the database ((default are MoBPS internal names based on position)) $ \\$	
	traits	Traits to include in the dendrogram (default: all traits)	
	type	Which traits values to consider (default: "pheno", alt: "bv", "bve")	

Value

Dendrogram plot of genotypes vs phenotypes

```
population <- creating.diploid(nsnp=1000, nindi=40, n.additive = c(100,100,100),</pre>
          shuffle.cor = matrix(c(1,0.8,0.2,0.8,1,0.2,0.2,0.2,1), ncol=3), shuffle.traits = 1:3)
population <- breeding.diploid(population, phenotyping = "all", heritability = 0.5)
get.dendrogram.heatmap(population, gen=1, type="pheno")
```

get.dendrogram.trait 71

```
get.dendrogram.trait Dendrogram
```

Description

Function calculate a dendogram for the traits

Usage

```
get.dendrogram.trait(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  traits = NULL,
  type = "pheno"
)
```

Arguments

population	Population list
path	provide a path if the dendrogram would be saved as a png-file
database	Groups of individuals to consider
gen	Quick-insert for database (vector of all generations to consider)
cohorts	Quick-insert for database (vector of names of cohorts to consider)
traits	Traits to include in the dendrogram (default: all traits)
type	Which traits values to consider (default: "pheno", alt: "bv", "bve")

Value

Dendrogram plot for traits

```
population <- creating.diploid(nsnp=1000, nindi=100, n.additive = c(100,100,100), shuffle.cor = matrix(c(1,0.8,0.2,0.8,1,0.2,0.2,0.2,1), ncol=3), shuffle.traits = 1:3) population <- breeding.diploid(population, phenotyping = "all", heritability = 0.5) get.dendrogram.trait(population, gen=1, type="pheno")
```

72 get.distance

get.distance

Calculate Nei distance between two or more population

Description

Function to calculate Nei's distance between two or more population

Usage

```
get.distance(
  population,
  type = "nei",
  marker = "all",
  per.marker = FALSE,
  gen1 = NULL,
  database1 = NULL,
  cohorts1 = NULL,
  gen2 = NULL,
  database2 = NULL,
  cohorts2 = NULL,
  database.list = NULL,
  gen.list = NULL,
  cohorts.list = NULL
```

Arguments

population	population list
type	Chose type of distance to compute (default: Neis standard genetic distance "nei"). Alt: Reynolds distance ("reynold"), Cavalli-Sforza ("cavalli"), Neis distance ("nei_distance"), Neis minimum distance ("nei_minimum")
marker	Vector with SNPs to consider (Default: "all" - use of all markers)
per.marker	Set to TRUE to return per marker statistics on genetic distances
gen1	Quick-insert for database (vector of all generations to consider)
database1	First Groups of individuals to consider
cohorts1	Quick-insert for database (vector of names of cohorts to consider)
gen2	Quick-insert for database (vector of all generations to consider)
database2	Second Groups of individuals to consider
cohorts2	Quick-insert for database (vector of names of cohorts to consider)
database.list	List of databases to consider (use when working with more than 2 populations)
gen.list	Quick-insert for database (vector of all generations to consider)
cohorts.list	Quick-insert for database (vector of names of cohorts to consider)

get.effect.freq 73

Value

Population list

Examples

```
data(ex_pop)
get.distance(ex_pop, database1 = cbind(1,1), database2 = cbind(1,2))
```

get.effect.freq

Compute marker frequency in QTL-markers

Description

Function to compute marker frequency in QTL-markers

Usage

```
get.effect.freq(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  sort = FALSE
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

sort Set to FALSE to not sort markers according to position on the genome

Value

Matrix with allele frequencies in the QTLs

```
data(ex_pop)
get.effect.freq(ex_pop, gen=1)
```

74 get.geno

```
get.effective.size
```

Estimate effective population size

Description

Function to estimate the effective population size

Usage

```
get.effective.size(population, gen = NULL, database = NULL, cohorts = NULL)
```

Arguments

population Population list

gen Quick-insert for database (vector of all generations to export)

database Groups of individuals to consider for the export

cohorts Quick-insert for database (vector of names of cohorts to export)

Value

Estimated effective population size

Examples

```
data(ex_pop)
get.effective.size(population=ex_pop, gen=5)
```

get.geno

Derive genotypes of selected individuals

Description

Function to devide genotypes of selected individuals

```
get.geno(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  chromosomen = "all",
  export.alleles = FALSE,
  non.genotyped.as.missing = FALSE,
  use.id = FALSE
)
```

75 get.genotyped

Arguments

population Population list database Groups of individuals to consider for the export gen Quick-insert for database (vector of all generations to export) cohorts Quick-insert for database (vector of names of cohorts to export) Beschraenkung des Genotypen auf bestimmte Chromosomen (default: 1) chromosomen export.alleles If TRUE export underlying alleles instead of just 012 non.genotyped.as.missing Set to TRUE to replace non-genotyped markers with NA

Set to TRUE to use MoBPS ids instead of Sex Nr Gen based names (default: use.id

FALSE)

Value

Genotype data for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
geno <- get.geno(ex_pop, gen=2)</pre>
```

get.genotyped

Derive genotyping status

Description

Function to if selected individuals are genotyped

Usage

```
get.genotyped(
 population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

Population list population

database Groups of individuals to consider for the export

Quick-insert for database (vector of all generations to export) gen cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

76 get.genotyped.snp

Value

Check if in gen/database/cohorts selected individuals are genotyped

Examples

```
data(ex_pop)
get.genotyped(ex_pop, gen=2)
```

get.genotyped.snp

Derive which markers are genotyped of selected individuals

Description

Function to devide which markers are genotyped for the selected individuals

Usage

```
get.genotyped.snp(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  export.alleles = FALSE,
  use.id = FALSE
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

export.alleles If TRUE export underlying alleles instead of just 012

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Binary Coded is/isnot genotyped level for in gen/database/cohorts selected individuals

```
data(ex_pop)
genotyped.snps <- get.genotyped.snp(ex_pop, gen=2)</pre>
```

get.haplo 77

get.haplo

Derive haplotypes of selected individuals

Description

Function to devide haplotypes of selected individuals

Usage

```
get.haplo(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  chromosomen = "all",
  export.alleles = FALSE,
  non.genotyped.as.missing = FALSE,
  use.id = FALSE
)
```

Arguments

population	Population list	
database	Groups of individuals to consider for the export	
gen	Quick-insert for database (vector of all generations to export)	
cohorts	Quick-insert for database (vector of names of cohorts to export)	
chromosomen	Beschraenkung der Haplotypen auf bestimmte Chromosomen (default: 1)	
export.alleles	If TRUE export underlying alleles instead of just 012	
non.genotyped.as.missing		
	Set to TRUE to replace non-genotyped markers with NA	
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)	

Value

Haplotype data for in gen/database/cohorts selected individuals

```
data(ex_pop)
haplo <- get.haplo(ex_pop, gen=2)</pre>
```

78 get.individual.loc

get.id	Derive ID on an individual
get.iu	Derive ID on an maiviauai

Description

Function to derive the internal ID given to each individual

Usage

```
get.id(population, database = NULL, gen = NULL, cohorts = NULL, use.id = FALSE)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names

Value

Individual ID for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.id(ex_pop, gen=2)
```

get.individual.loc

Export location of individuals from the population list

Description

Export location of individuals from the population list

Usage

```
get.individual.loc(population, database = NULL, gen = NULL, cohorts = NULL)
```

Arguments

population	Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

get.infos 79

Value

Storage Position for in gen/database/cohorts selected individuals (Generation/Sex/IndividualNr)

Examples

```
data(ex_pop)
get.individual.loc(ex_pop, gen=2)
```

get.infos

Extract bv/pheno/geno of selected individuals

Description

Function to extract bv/pheno/geno of selected individuals

Usage

```
get.infos(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

Value

Info list [[1]] phenotypes [[2]] genomic values [[3]] Z [[4/5/6]] additive/epistatic/dice marker effects

Examples

```
data(ex_pop)
get.infos(ex_pop, gen=2)
```

FALSE)

get.npheno

get.map

Map generation

Description

Function to derive the genomic map for a given population list

Usage

```
get.map(population, use.snp.nr = FALSE)
```

Arguments

```
population Population list
use.snp.nr Set to TRUE to display SNP number and not SNP name
```

Value

Genomic map of the population list

Examples

```
data(ex_pop)
map <- get.map(ex_pop)</pre>
```

get.npheno

Export underlying number of observations per phenotype

Description

Function to export the number of observation of each underlying phenotype

```
get.npheno(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.all.copy = FALSE,
  use.id = FALSE
)
```

get.pca 81

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.all.copy	Set to TRUE to extract phenotyping
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Phenotypes for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.pheno(ex_pop, gen=2)
```

get.pca

Principle components analysis

Description

Function to perform a principle component analysis

```
get.pca(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  coloring = "group",
  components = c(1, 2),
  plot = TRUE,
  pch = 1,
  export.color = FALSE
)
```

get.pedigree

Arguments

population Population list

path Location were to save the PCA-plot

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

coloring Coloring by "group", "sex", "plain"

components Default: c(1,2) for the first two principle components

plot Set to FALSE to not generate a plot

pch Point type in the PCA plot

export.color Set to TRUE to export the per point coloring

Value

Genotype data for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.pca(ex_pop, gen=2)
```

get.pedigree

Derive pedigree

Description

Derive pedigree for selected individuals

```
get.pedigree(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  founder.zero = TRUE,
  raw = FALSE,
  id = FALSE
)
```

get.pedigree2 83

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
founder.zero	Parents of founders are displayed as "0" (default: TRUE)
raw	Set to TRUE to not convert numbers into Sex etc.
id	Set to TRUE to extract individual IDs

Value

Pedigree-file for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.pedigree(ex_pop, gen=2)
```

get.pedigree2

Derive pedigree including grandparents

Description

Derive pedigree for selected individuals including grandparents

Usage

```
get.pedigree2(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  shares = FALSE,
  founder.zero = TRUE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
shares	Determine actual inherited shares of grandparents
founder.zero	Parents of founders are displayed as "0" (default: TRUE)

84 get.pedigree3

Value

Pedigree-file (grandparents) for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.pedigree2(ex_pop, gen=2)
```

get.pedigree3

Derive pedigree parents and grandparents

Description

Derive pedigree for selected individuals including parents/grandparents

Usage

```
get.pedigree3(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  founder.zero = TRUE
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

founder.zero Parents of founders are displayed as "0" (default: TRUE)

Value

Pedigree-file (parents + grandparents) for in gen/database/cohorts selected individuals

```
data(ex_pop)
get.pedigree3(ex_pop, gen=3)
```

get.pedmap 85

get.pedmap Generate plink-file (pedmap)	get.pedmap	Generate	plink-file	(pedmap)
---	------------	----------	------------	----------

Description

Generate a ped and map file (PLINK format) for selected groups and chromosome

Usage

```
get.pedmap(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  non.genotyped.as.missing = FALSE,
  use.id = FALSE
)
```

Arguments

population Population list

path Location to save pedmap-file

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

non.genotyped.as.missing

Set to TRUE to replaced non-genotyped entries with "./."

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names

Value

Ped and map-file for in gen/database/cohorts selected individuals

```
data(ex_pop)
file_path <- tempdir()
get.pedmap(path=file_path, ex_pop, gen=2)
file.remove(paste0(file_path, ".ped"))
file.remove(paste0(file_path, ".map"))</pre>
```

get.pheno

get.pheno

Export underlying phenotypes

Description

Function to export underlying phenotypes

Usage

```
get.pheno(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.all.copy = FALSE,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.all.copy	Set to TRUE to extract phenotyping
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Phenotypes for in gen/database/cohorts selected individuals

```
data(ex_pop)
get.pheno(ex_pop, gen=2)
```

get.pheno.off 87

get.pheno.off

Export underlying offspring phenotypes

Description

Function to export offspring phenotypes

Usage

```
get.pheno.off(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

Value

Avg. phenotype of the offspring of in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.pheno.off(ex_pop, gen=2)
```

get.pheno.off.count

Export underlying number of used offspring for offspring phenotypes

Description

Function to export number of observations used for offspring phenotypes

```
get.pheno.off.count(population, database = NULL, gen = NULL, cohorts = NULL)
```

get.phylogenetic.tree

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

Value

Number of offspring with phenotypes for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.pheno.off.count(ex_pop, gen=2)
```

```
get.phylogenetic.tree Phylogenetic Tree
```

Description

Function calculate a phylogenetic tree

Usage

```
get.phylogenetic.tree(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  method = NULL,
  individual.names = NULL,
  circular = FALSE
)
```

Arguments

population Population list

path provide a path if the dendrogram would be saved as a png-file

database Groups of individuals to consider

gen Quick-insert for database (vector of all generations to consider)

cohorts Quick-insert for database (vector of names of cohorts to consider)

method Method used to calculate genetic distances (default: "Nei", alt: "Rogers", "Pre-

vosti", "Modified Rogers"

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individual.names

Names of the individuals in the database ((default are MoBPS internal names

based on position))

circular

Set to TRUE to generate a fan/circular layout tree

Value

Dendrogram plot for traits

Examples

```
data(ex_pop)
get.phylogenetic.tree(ex_pop, gen=1, circular=TRUE)
```

get.qtl

QTL extraction

Description

Function to the position of QTLs (for snp/chr use get.qtl.effects()

Usage

```
get.qtl(population)
```

Arguments

population

Population list

Value

Vector of SNP positions

```
data(ex_pop)
positions <- get.qtl(ex_pop)</pre>
```

90 get.qtl.variance

get.qtl.effects

QTL effect extraction

Description

Function to extract QTL effect sizes

Usage

```
get.qtl.effects(population)
```

Arguments

population

Population list

Value

List with [[1]] single SNP QTLs [[2]] epistatic SNP QTLs [[3]] dice QTL

Examples

```
data(ex_pop)
effects <- get.qtl.effects(ex_pop)</pre>
```

get.qtl.variance

QTL effect variance extraction

Description

Function to extract QTL effect variance for single SNP QTLs in a given gen/database/cohort

Usage

```
get.qtl.variance(population, gen = NULL, database = NULL, cohorts = NULL)
```

Arguments

population Population list

gen Quick-insert for database (vector of all generations to consider)

database Groups of individuals to consider

cohorts Quick-insert for database (vector of names of cohorts to consider)

Value

matrix with SNP / Chr / estimated effect variance

get.recombi 91

Examples

```
data(ex_pop)
effects <- get.qtl.variance(ex_pop)</pre>
```

get.recombi

Derive genetic origins

Description

Function to derive genetic origin

Usage

```
get.recombi(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Recombination points for in gen/database/cohorts selected individuals

```
data(ex_pop)
get.recombi(ex_pop, gen=2)
```

92 get.selectionbve

get.reliabilities

Export underlying reliabilities

Description

Function to export underlying reliabilities

Usage

```
get.reliabilities(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Estimated reliability for BVE for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.reliabilities(ex_pop, gen=2)
```

get.selectionbve

Export derived breeding values based on the selection index

Description

Function to export last breeding values based on the selection index

get.selectionindex 93

Usage

```
get.selectionbve(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population	Population list
database	Groups of individuals to consider for the export
gen	Quick-insert for database (vector of all generations to export)
cohorts	Quick-insert for database (vector of names of cohorts to export)
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default: FALSE)

Value

Last applied selection index for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.selectionindex(ex_pop, gen=2)
```

get.selectionindex

Export underlying last used selection index

Description

Function to export last used selection index (mostly relevant for Miesenberger 1997 stuff)

```
get.selectionindex(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

94 get.time.point

Arguments

population	Population list
database	Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)
cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

Value

Last applied selection index for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
get.selectionindex(ex_pop, gen=2)
```

get.time.point

Derive time point

Description

Function to devide time point for each individual

Usage

```
get.time.point(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  use.id = FALSE
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

use.id Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names (default:

FALSE)

Value

Time point of generation for in gen/database/cohorts selected individuals

get.vcf 95

Examples

```
data(ex_pop)
get.time.point(ex_pop, gen=2)
```

get.vcf

Generate vcf-file

Description

Generate a vcf-file for selected groups and chromosome

Usage

```
get.vcf(
  population,
  path = NULL,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  chromosomen = "all",
  non.genotyped.as.missing = FALSE,
  use.id = FALSE
)
```

Arguments

population	Population list	
path	Location to save vcf-file	
database	base Groups of individuals to consider for the export	
gen	Quick-insert for database (vector of all generations to export)	
cohorts Quick-insert for database (vector of names of cohorts to export)		
chromosomen	Beschraenkung des Genotypen auf bestimmte Chromosomen (default: 1)	
non.genotyped.as.missing		
	Set to TRUE to replaced non-genotyped entries with "./."	
use.id	Set to TRUE to use MoBPS ids instead of Sex_Nr_Gen based names	

Value

VCF-file for in gen/database/cohorts selected individuals

96 group.diff

Examples

```
data(ex_pop)
data(ex_pop)

file_path <- tempdir()
get.vcf(path=file_path, ex_pop, gen=2)
file.remove(paste0(file_path, ".vcf"))</pre>
```

group.diff

Function to exclude individuals from a database

Description

Function to exclude individuals from a database

Usage

```
group.diff(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  remove.gen = NULL,
  remove.database = NULL,
  remove.cohorts = NULL
)
```

Arguments

population Population list

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

remove.gen Generations of individuals to remove from the database (same IDs!)

remove.database

Groups of individuals to remove from the database (same IDs!)

remove.cohorts Cohorts of individuals to remove from the database (same IDs!)

Value

Database excluding removals

```
data(ex_pop)
database <- group.diff(ex_pop, gen=1, remove.database=cbind(1,1))</pre>
```

insert.bve 97

insert.bve

Manually enter estimated breeding values

Description

Function to manually enter estimated breeding values

Usage

```
insert.bve(
  population,
  bves,
  type = "bve",
  na.override = FALSE,
  count = 1,
  count.only.increase = TRUE
)
```

Arguments

populat	Population list
bves	Matrix of breeding values to enter (one row per individual with 1 element coding individual name)
type	which time of values to input (default: "bve", alt: "bv", "pheno")
na.over	e Set to TRUE to also enter NA values (Default: FALSE - those entries will be skipped)
count	Counting for economic cost calculation (default: 1 - (one observation (for "pheno"), one genotyping (for "bve")))
count.o	. increase
	Set to FALSE to reduce the number of observation for a phenotype to "count" (default: TRUE)

Value

Population-List with newly entered estimated breeding values

```
data(ex_pop)
bv <- get.bv(ex_pop, gen=2)
new.bve <- cbind( colnames(bv), bv[,1]) ## Unrealistic but you do not get better than this!
ex_pop <- insert.bve(ex_pop, bves=new.bve)</pre>
```

98 json.simulation

json.simulation	Simulation of a breeding program based on a JSON-file from MoBP-Sweb

Description

Function to simulate a breeding program based on a JSON-file from MoBPSweb

Usage

```
json.simulation(
 file = NULL,
  log = NULL,
  total = NULL,
  fast.mode = FALSE,
 progress.bars = FALSE,
  size.scaling = NULL,
  rep.max = 1,
  verbose = TRUE,
 miraculix.cores = NULL,
 miraculix.chol = NULL,
  skip.population = FALSE,
  time.check = FALSE,
  time.max = 7200,
  export.population = FALSE,
  export.gen = NULL,
  export.timepoint = NULL,
  fixed.generation.order = NULL
)
```

Arguments

file	Path to a json-file generated by the user-interface	
log	Provide Path where to write a log-file of your simulation (or false to not write a log-file)	
total	Json-file imported via jsonlite::read_json	
fast.mode	Set to TRUE work on a small genome with few markers	
progress.bars Set to TRUE to display progress bars		
size.scaling Scale the size of nodes by this factor (especially for testing smaller examples)		
rep.max Maximum number of repeats to use in fast.mode		
verbose Set to FALSE to not display any prints		
miraculix.cores		
	Number of cores used in miraculix applications (default: 1)	
miraculix.chol	Set to FALSE to manually deactive the use of miraculix for any cholesky decompostion even though miraculix is actived	

kinship.development 99

skip.population

Set to TRUE to not execute breeding actions (only cost/time estimation will be

performed)

time.check Set to TRUE to automatically check simulation run-time before executing breed-

ing actions

time.max Maximum length of the simulation in seconds when time.check is active

export.population

Path were to export the population to (at state selected in export.gen/timepoint)

export.gen Last generation to simulate before exporting population to file

export.timepoint

Last timepoint to simulate before exporting population to file

fixed.generation.order

Vector containing the order of cohorts to generate (Advanced // Testing Param-

eter!)

Value

Population-list

Examples

```
data(ex_json)
population <- json.simulation(total=ex_json)</pre>
```

kinship.development

Devolopment of genetic/breeding value

Description

Function to plot genetic/breeding values for multiple generation/cohorts

```
kinship.development(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  json = FALSE,
  ibd.obs = 50,
  hbd.obs = 10,
  display.cohort.name = FALSE,
  display.time.point = FALSE,
  equal.spacing = FALSE,
  time_reorder = FALSE,
  display.hbd = FALSE
```

100 kinship.emp

Arguments

population	population list	
database	Groups of individuals to consider for the export	
gen	Quick-insert for database (vector of all generations to export)	
cohorts	Quick-insert for database (vector of names of cohorts to export)	
json	If TRUE extract which cohorts to plot according to the json-file used in json.simulation	
ibd.obs	Number of Individual pairs to sample for IBD estimation	
hbd.obs	Number of Individuals to sample for HBD estimation	
display.cohort	.name	
	Set TRUE to display the name of the cohort in the x-axis	
display.time.point		
	Set TRUE to use time point of generated to sort groups	
equal.spacing	Equal distance between groups (independent of time.point)	
time_reorder	Set TRUE to order cohorts according to the time point of generation	
display.hbd	Set to TRUE to also display HBD in plot	

Value

Estimated of avg. kinship/inbreeding based on IBD/HBD

Examples

```
data(ex_pop)
kinship.development(ex_pop,gen=1:5)
```

kinship.emp

Empirical kinship

Description

Function to compute empirical kinship for a set of individuals)

```
kinship.emp(
   animals = NULL,
   population = NULL,
   gen = NULL,
   database = NULL,
   cohorts = NULL,
   sym = FALSE
)
```

kinship.emp.fast 101

Arguments

animals List of animals to compute kinship for

population Population list

gen Quick-insert for database (vector of all generations to export)

database Groups of individuals to consider for the export

cohorts Quick-insert for database (vector of names of cohorts to export)

sym If True derive matrix entries below principle-diagonal

Value

Empirical kinship matrix (IBD-based since Founders)

Examples

```
data(ex_pop)
kinship <- kinship.emp(population=ex_pop, database=cbind(2,1,1,25))</pre>
```

kinship.emp.fast

Approximate empirical kinship

Description

Function to compute empirical kinship for a set of individuals (not all pairs of individuals are evaluated)

Usage

```
kinship.emp.fast(
  animals = NULL,
  population = NULL,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  sym = FALSE,
  ibd.obs = 50,
  hbd.obs = 10
)
```

Arguments

animals List of animals to compute kinship for

population Population list

gen Quick-insert for database (vector of all generations to export)

database Groups of individuals to consider for the export

102 kinship.exp

cohorts	Quick-insert for database (vector of names of cohorts to export)
sym	If True derive matrix entries below principle-diagonal
ibd.obs	Number of Individual pairs to sample for IBD estimation
hbd.obs	Number of Individuals to sample for HBD estimation

Value

Empirical kinship matrix (IBD-based since Founders) per gen/database/cohort

Examples

```
data(ex_pop)
kinship.emp.fast(population=ex_pop,gen=2)
```

kinship.exp

Derive expected kinship

Description

Function to derive expected kinship

Usage

```
kinship.exp(
  population,
  gen = NULL,
  database = NULL,
  cohorts = NULL,
  depth.pedigree = 7,
  start.kinship = NULL,
  elements = NULL,
  mult = 2,
  storage.save = 1.5,
  verbose = TRUE
)
```

Population list

Arguments

population

gen	Quick-insert for database (vector of all generations to export)
database	Groups of individuals to consider for the export
cohorts	Quick-insert for database (vector of names of cohorts to export)
depth.pedigree	Depth of the pedigree in generations
start.kinship	Relationship matrix of the individuals in the first considered generation
elements	Vector of individuals from the database to include in pedigree matrix

ld.decay 103

mult Multiplicator of kinship matrix (default: 2)

storage.save Lower numbers will lead to less memory but slightly higher computing time

(default: 1.5, min: 1)

verbose Set to FALSE to not display any prints

Value

Pedigree-based kinship matrix for in gen/database/cohort selected individuals

Examples

```
data(ex_pop)
kinship <- kinship.exp(population=ex_pop, gen=2)</pre>
```

ld.decay

Generate LD plot

Description

Generate LD pot

Usage

```
ld.decay(
  population,
  genotype.dataset = NULL,
  chromosomen = 1,
  dist = NULL,
  step = 5,
  max = 500,
  max.cases = 100,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  type = "snp",
  plot = FALSE
)
```

Arguments

population Population list genotype.dataset

Genotype dataset (default: NULL - just to save computation time when get.geno

was already run)

chromosomen Only consider a specific chromosome in calculations (default: 1)

dist Manuel input of marker distances to analyse

104 maize_chip

step Stepsize to calculate LD

max Maximum distance between markers to consider for LD-plot

max.cases Maximum number of marker pairs to consider of each distance (default: 100;

randomly sampled!)

database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

type Compute LD decay according to following distance measure between markers

(default: "snp", alt: "bp", "cM")

plot Set to FALSE to not generate an LD plot

Value

LD-decay plot for in gen/database/cohorts selected individuals

Examples

```
data(ex_pop)
ld.decay(population=ex_pop, gen=5)
```

maize_chip

maize chip

Description

Genome for maize according to Lee et al.

Usage

maize_chip

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

Lee et al 2002

miesenberger.index 105

miesenher	ger.index

Miesenberger Index

Description

Function to selection index weights according to Miesenberger 1997

Usage

```
miesenberger.index(V, G, V1 = NULL, RG = NULL, r, w, zw = NULL)
```

Arguments

V	Phenotypic covarianz matrix
G	Genomic covarianz matrix
V1	Inverted phenotypic covarianz matrix
RG	Genomic correlation matrix
r	reliability for the breeding value estimation
W	relative weighting of each trait (per genetic SD)
ZW	Estimated breeding value

Value

weights of the selection index

		-	
m 1 r	^acı	пl	1 X

Add miraculix-coding for genotypes

Description

Internal function to store genotypes bit-wise

Usage

```
miraculix(population)
```

Arguments

population

Population list

Value

Population list

106 mutation.intro

Examples

```
# This is only relevant with the package miraculix is installed and used
population <- creating.diploid(nsnp=100, nindi=50, miraculix=FALSE)
population <- miraculix(population)</pre>
```

mutation.intro

Mutation intro

Description

Function to change the base-pair in a specific loci

Usage

```
mutation.intro(population, gen, sex, individual.nr, qtl.posi, haplo.set = 1)
```

Arguments

population Population list

gen Generation of the individual to introduce a mutation in

sex Sex of the individual to introduce a mutation in

individual.nr Individual Nr. of the individual to introduce a mutation in

qtl.posi Marker number to mutate

haplo.set Select chromosome set (default: 1, alt: 2)

Value

Population-List with mutated marker for the selected individual

```
data(ex_pop)
ex_pop <- mutation.intro(ex_pop, 1,1,1, qtl.posi=100)</pre>
```

new.base.generation 107

new.base.generation Set new base generation

Description

Function to set a new base generation for the population

Usage

```
new.base.generation(
  population,
  base.gen = NULL,
  delete.previous.gen = FALSE,
  delete.breeding.totals = FALSE,
  delete.bve.data = FALSE,
  add.chromosome.ends = TRUE
)
```

Arguments

Value

Population-List with mutated marker for the selected individual

```
data(ex_pop)
ex_pop <- new.base.generation(ex_pop, base.gen=2)</pre>
```

108 OGC

OGC

Optimal genetic contribution

Description

In this function the OGC selection according to Meuwissen 1997 is performed

Usage

```
OGC(
A,
u,
Q,
cAc = NA,
single = TRUE,
verbose = FALSE,
max_male = Inf,
max_female = Inf)
```

Arguments

A	relationship matrix
u	breeding values
Q	sex indicator
cAc	target gain in inbreeding
single	If FALSE multiple individuals can be removed at the same type (this is faster but potentially inaccurate!)
verbose	Set to FALSE to not display any prints
max_male	maximum number of male with positive contributions
max_female	maximum number of females with positive contributions

Value

[1] Contributions [[2]] expected inbreeding gain

pedigree.simulation Simulation of a given pedigree

Description

Function to simulate a given pedigree

Usage

```
pedigree.simulation(
  pedigree,
  keep.ids = FALSE,
 plot = TRUE,
 dataset = NULL,
  vcf = NULL,
  chr.nr = NULL,
 bp = NULL,
  snp.name = NULL,
  hom0 = NULL,
  hom1 = NULL,
  bpcm.conversion = 0,
  nsnp = 0,
  freq = "beta",
  sex.s = "fixed",
  chromosome.length = NULL,
  length.before = 5,
  length.behind = 5,
  real.bv.add = NULL,
  real.bv.mult = NULL,
  real.bv.dice = NULL,
  snps.equidistant = NULL,
  change.order = FALSE,
  bv.total = 0,
  polygenic.variance = 100,
  bve.mult.factor = NULL,
  bve.poly.factor = NULL,
  base.bv = NULL,
  add.chromosome.ends = TRUE,
  new.phenotype.correlation = NULL,
  new.residual.correlation = NULL,
  new.breeding.correlation = NULL,
  add.architecture = NULL,
  snp.position = NULL,
  position.scaling = FALSE,
  bit.storing = FALSE,
  nbits = 30,
  randomSeed = NULL,
```

```
miraculix = TRUE,
miraculix.dataset = TRUE,
n.additive = 0,
n.dominant = 0,
n.qualitative = 0,
n.quantitative = 0,
var.additive.l = NULL,
var.dominant.l = NULL,
var.qualitative.1 = NULL,
var.quantitative.1 = NULL,
exclude.snps = NULL,
replace.real.bv = FALSE,
shuffle.traits = NULL,
shuffle.cor = NULL,
skip.rest = FALSE,
enter.bv = TRUE,
name.cohort = NULL,
template.chip = NULL,
beta.shape1 = 1,
beta.shape2 = 1,
time.point = 0,
creating.type = 0,
trait.name = NULL,
share.genotyped = 1,
genotyped.s = NULL,
map = NULL,
remove.invalid.qtl = TRUE,
verbose = TRUE,
bv.standard = FALSE,
mean.target = NULL,
var.target = NULL,
is.maternal = NULL,
is.paternal = NULL,
vcf.maxsnp = Inf
```

Arguments

)

pedigree	Pedigree-file (matrix with 3 columns (Individual ID, Father ID, Mother ID), optional forth columns with earliest generations to generate an individual)
keep.ids	Set to TRUE to keep the IDs from the pedigree-file instead of the default MoBPS ids
plot	Set to FALSE to not generate an overview of inbreeding and number of individuals over time
dataset	SNP dataset, use "random", "allhetero" "all0" when generating a dataset via nsnp,nindi
vcf	Path to a vcf-file used as input genotypes (correct haplotype phase is assumed!)

chr.nr Vector containing the assosiated chromosome for each marker (default: all on

the same)

bp Vector containing the physical position (bp) for each marker (default: 1,2,3...)

snp.name Vector containing the name of each marker (default ChrXSNPY - XY chosen

accordingly)

hom0 Vector containing the first allelic variant in each marker (default: 0)
hom1 Vector containing the second allelic variant in each marker (default: 1)

bpcm.conversion

Convert physical position (bp) into a cM position (default: 0 - not done)

nsnp number of markers to generate in a random dataset

freq frequency of allele 1 when randomly generating a dataset

sex.s Specify which newly added individuals are male (1) or female (2)

chromosome.length

Length of the newly added chromosome (default: 5)

length.before Length before the first SNP of the dataset (default: 5) length.behind Length after the last SNP of the dataset (default: 5)

real.bv.add Single Marker effects
real.bv.mult Two Marker effects
real.bv.dice Multi-marker effects

snps.equidistant

Use equidistant markers (computationally faster!; default: TRUE)

change.order If TRUE sort markers according to given marker positions

bv.total Number of traits (If more than traits via real.bv.X use traits with no directly

underlying QTL)

polygenic.variance

Genetic variance of traits with no underlying QTL

bve.mult.factor

Multiplicate trait value times this

bve.poly.factor

Potency trait value over this

base.bv Average genetic value of a trait

add.chromosome.ends

Add chromosome ends as recombination points

new.phenotype.correlation

(OLD! - use new.residual.correlation) Correlation of the simulated environmental

variance

new.residual.correlation

Correlation of the simulated environmental variance

new.breeding.correlation

Correlation of the simulated genetic variance (child share! heritage is not influ-

enced!

add.architecture

Add genetic architecture (marker positions)

snp.position Location of each marker on the genetic map

position.scaling

Manual scaling of snp.position

bit.storing Set to TRUE if the MoBPS (not-miraculix! bit-storing is used)

nbits Bits available in MoBPS-bit-storing randomSeed Set random seed of the process

miraculix If TRUE use miraculix package for data storage, computations and dataset gen-

eration

miraculix.dataset

Set FALSE to deactive miraculix package for dataset generation

n.additive Number of additive QTL n.dominant Number of dominante QTL

n.qualitative Number of qualitative epistatic QTL n.quantitative Number of quantitative epistatic QTL

var.additive.l Variance of additive QTL var.dominant.l Variance of dominante QTL

var.qualitative.l

Variance of qualitative epistatic QTL

var.quantitative.l

Variance of quantitative epistatic QTL

exclude.snps Marker were no QTL are simulated on

replace.real.bv

If TRUE delete the simulated traits added before

shuffle.traits Combine different traits into a joined trait shuffle.cor Target Correlation between shuffeled traits

skip.rest Internal variable needed when adding multipe chromosomes jointly

enter.bv Internal parameter

name.cohort Name of the newly added cohort

template.chip Import genetic map and chip from a species ("cattle", "chicken", "pig")
beta.shape1 First parameter of the beta distribution for simulating allele frequencies
beta.shape2 Second parameter of the beta distribution for simulating allele frequencies

time.point Time point at which the new individuals are generated

creating.type Technique to generate new individuals (usage in web-based application)

trait.name Name of the trait generated

share.genotyped

Share of individuals genotyped in the founders

genotyped.s Specify with newly added individuals are genotyped (1) or not (0)

map map-file that contains up to 5 colums (Chromsome, SNP-id, M-position, Bp-

position, allele freq - Everything not provides it set to NA). A map can be im-

ported via MoBPSmaps::ensembl.map()

```
remove.invalid.qtl
                  Set to FALSE to deactive the automatic removal of QTLs on markers that do not
verbose
                  Set to FALSE to not display any prints
                  Set TRUE to standardize trait mean and variance via bv.standardization() - au-
bv.standard
                  tomatically set to TRUE when mean/var.target are used
mean.target
                  Target mean
var.target
                  Target variance
                  Vector coding if a trait is caused by a maternal effect (Default: all FALSE)
is.maternal
                  Vector coding if a trait is caused by a paternal effect (Default: all FALSE)
is.paternal
vcf.maxsnp
                  Maximum number of SNPs to include in the genotype file (default: Inf)
add.chromosome If TRUE add an additional chromosome to the dataset
```

Value

Population-list

Examples

```
pedigree <- matrix(c(1,0,0,
2,0,0,
3,0,0,
4,1,2,
5,1,3,
6,1,3,
7,1,3,
8,4,6,
9,4,7), ncol=3, byrow=TRUE)
population <- pedigree.simulation(pedigree, nsnp=1000)</pre>
```

pedmap.to.phasedbeaglevcf

Internal function to perform imputing/phasing

Description

Internal function to perform imputing/phasing (path chosen for the web-based application)

Usage

```
pedmap.to.phasedbeaglevcf(
  ped_path = NULL,
  map_path = NULL,
  vcf_path = NULL,
  beagle_jar = "/home/nha/beagle.03Jul18.40b.jar",
  plink_dir = "/home/nha/Plink/plink",
```

pig_chip

```
db_dir = "/home/nha/Plink/DB/",
  verbose = TRUE
)
```

Arguments

ped_path	Directory of the ped-file
map_path	Directory of the map-file
vcf_path	Directory of the vcf-file (this will override any ped/map-file input)
beagle_jar	Directory of BEAGLE
plink_dir	Directory of Plink
db_dir	Directory to save newly generated files (ped/map will be stored in the original folder)

verbose Set to FALSE to not display any prints

Value

Phased vcf file in vcf_path

Description

Genome for pig according to Rohrer et al.

Usage

```
pig_chip
```

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

Rohrer et al 1994

plot.population 115

7 .	7
$n \cap t$	population
ртос.	population

Plot Population

Description

Basic plot of the population list

Usage

```
## S3 method for class 'population'
plot(x, type = "bve", gen = NULL, database = NULL, cohorts = NULL, ...)
```

Arguments

```
x Population-list

type Default "bve" - bv.development, alt: "kinship" - kinship.development(), "pca" -
get.pca()

gen generations to consider

database groups to consider

cohorts cohorts to consider

... remaining stuff
```

Value

Summary of the population list including number of individuals, genone length and trait overview

Examples

```
data(ex_pop)
plot(ex_pop)
```

set.class

Export estimated breeding values

Description

Function to export estimated breeding values

Usage

```
set.class(
  population,
  database = NULL,
  gen = NULL,
  cohorts = NULL,
  new.class = 0
)
```

116 set.default

Arguments

population Population list
database Groups of individuals to consider for the export

gen Quick-insert for database (vector of all generations to export)

cohorts Quick-insert for database (vector of names of cohorts to export)

new.class Class to change to (either single character or vector for each individual when

just a single group is selected)

Value

Population-List with newly entered class values

Examples

```
data(ex_pop)
population <- set.class(ex_pop, database=cbind(1,1), new.class = 2)</pre>
```

set.default

Set defaults

Description

Set default parameter values in breeding.diploid

Usage

```
set.default(
  population,
  parameter.name = NULL,
  parameter.value = NULL,
  parameter.remove = NULL,
  reset.all = FALSE
)
```

Arguments

population Population list

parameter.name Number of traits (If more than traits via real.bv.X use traits with no directly underlying QTL)

parameter.value Genetic variance of traits with no underlying QTL

parameter.remove Remove a specific previously generated parameter default

reset.all Set to TRUE to remove all prior parameter values

sheep_chip 117

Value

Population-list with one or more additional new traits

Examples

```
data(ex_pop)
population <- set.default(ex_pop, parameter.name="heritability", parameter.value=0.3)</pre>
```

sheep_chip

sheep chip

Description

Genome for sheep according to Prieur et al.

Usage

sheep_chip

Author(s)

Torsten Pook <torsten.pook@uni-goettingen.de>

Source

Prieur et al 2017

sortd

Apply sort and unique

Description

Efficient function to perform sort(unique(v))

Usage

sortd(v)

Arguments

٧

Vector

Value

numerical sorted vector without duplicates

118 summary.population

Examples

```
v <- c(1,1,4,5)
sortd(v)</pre>
```

ssGBLUP

Single Step GBLUP

Description

Function to perform single step GBLUP according to Legarra 2014

Usage

```
ssGBLUP(A11, A12, A22, G)
```

Arguments

A11	pedigree relationship matrix of non-genotyped individuals
A12	pedigree relationship matrix between non-genotyped and genotyped individuals
A22	pedigree relationship matrix of genotyped individuals
G	genomic relationship matrix of genotyped individuals

Value

Single step relationship matrix

summary.population

Summary Population

Description

Summary of the population list

Usage

```
## S3 method for class 'population'
summary(object, ...)
```

Arguments

object Population-list

. . . additional arguments affecting the summary produced

vlist 119

Value

Summary of the population list including number of individuals, genone length and trait overview

Examples

```
data(ex_pop)
summary(ex_pop)
```

vlist

Generation of a sublist

Description

Internal function to write a couple of list entries in a new list

Usage

```
vlist(list, skip = NULL, first = NULL, select = NULL)
```

Arguments

list	list you want to print details of
skip	Skip first that many list-elements

first Only display first that many list-elements

select Display only selected list-elements

Value

Selected elements of a list

Examples

```
data(ex_pop)
vlist(ex_pop$breeding[[1]], select=3:10)
```

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```