Package: martini (via r-universe)

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Type Package

Title GWAS Incorporating Networks

Version 1.25.0

Description martini deals with the low power inherent to GWAS studies by using prior knowledge represented as a network. SNPs are the vertices of the network, and the edges represent biological relationships between them (genomic adjacency, belonging to the same gene, physical interaction between protein products). The network is scanned using SConES, which looks for groups of SNPs maximally associated with the phenotype, that form a close subnetwork.

License GPL-3

LazyData TRUE

Imports igraph (>= 1.0.1), Matrix, memoise (>= 2.0.0), methods (>= 3.3.2), Rcpp (>= 0.12.8), snpStats (>= 1.20.0), stats, utils,

Suggests biomaRt (>= 2.34.1), circlize (>= 0.4.11), STRINGdb (>= 2.2.0), httr (>= 1.2.1), IRanges (>= 2.8.2), S4Vectors (>= 0.12.2), knitr, testthat, readr, rmarkdown

Depends R (>= 4.0)

LinkingTo Rcpp, RcppEigen (>= 0.3.3.5.0)

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URL https://github.com/hclimente/martini

BugReports https://github.com/hclimente/martini/issues

RemoteUrl https://bioc.r-universe.dev **RemoteUrl** https://github.com/bioc/martini

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get_GI_network

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get_GI_network

Get gene-interaction network.

Description

Creates a network of SNPs where each SNP is connected as in the GM network and, in addition, to all the other SNPs pertaining to any interactor of the gene it is mapped to. Corresponds to the gene-interaction (GI) network described by Azencott et al.

```
get_GI_network(
   gwas,
   organism = 9606,
   snpMapping = snp2ensembl(gwas, organism),
   ppi = get_gxg("biogrid", organism, flush),
   col_ppi = c("gene1", "gene2"),
   col_genes = c("snp", "gene"),
   flush = FALSE
)
```

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Arguments

gwas	A SnpMatrix object with the GWAS information.
organism	Tax ID of the studied organism. The default is 9606 (human).
snpMapping	A data.frame informing how SNPs map to genes. It contains minimum two columns: SNP id and a gene it maps to. Each row corresponds to one gene-SNP mapping. Unless column names are specified using col_genes, involved columns must be named 'snp' and 'gene'.
ppi	A data.frame describing protein-protein interactions with at least two colums. Gene ids must be the contained in snpMapping. Unless column names are specified using col_ppi, involved columns must be named gene1 and gene2.
col_ppi	Optional, length-2 character vector with the names of the two columns involving the protein-protein interactions.
col_genes	Optional, length-2 character vector with the names of the two columns involving the SNP-gene mapping. The first element is the column of the SNP, and the second is the column of the gene.
flush	Remove cached results? Boolean value.

Value

An igraph network of the GI network of the SNPs.

References

Azencott, C. A., Grimm, D., Sugiyama, M., Kawahara, Y., & Borgwardt, K. M. (2013). Efficient network-guided multi-locus association mapping with graph cuts. Bioinformatics, 29(13), 171-179. https://doi.org/10.1093/bioinformatics/btt238

Examples

```
get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)

get_GM_network

Get gene membership network.
```

Description

Creates a network of SNPs where each SNP is connected as in the GS network and, in addition, to all the other SNPs pertaining to the same gene. Corresponds to the gene membership (GM) network described by Azencott et al.

```
get_GM_network(
   gwas,
   organism = 9606,
   snpMapping = snp2ensembl(gwas, organism),
   col_genes = c("snp", "gene")
)
```

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Arguments

gwas A SnpMatrix object with the GWAS information.

organism Tax ID of the studied organism. The default is 9606 (human).

snpMapping A data frame informing how SNPs map to genes. It contains minimum two

columns: SNP id and a gene it maps to. Each row corresponds to one gene-SNP mapping. Unless column names are specified using col_genes, involved

columns must be named 'snp' and 'gene'.

col_genes Optional, length-2 character vector with the names of the two columns involving

the SNP-gene mapping. The first element is the column of the SNP, and the

second is the column of the gene.

Value

An igraph network of the GM network of the SNPs.

References

Azencott, C. A., Grimm, D., Sugiyama, M., Kawahara, Y., & Borgwardt, K. M. (2013). Efficient network-guided multi-locus association mapping with graph cuts. Bioinformatics, 29(13), 171-179. https://doi.org/10.1093/bioinformatics/btt238

Examples

```
get_GM_network(minigwas, snpMapping = minisnpMapping)
```

get_GS_network

Get genomic sequence network

Description

Creates a network of SNPs where each SNP is connected to its adjacent SNPs in the genome sequence. Corresponds to the genomic sequence (GS) network described by Azencott et al.

Usage

```
get_GS_network(gwas)
```

Arguments

gwas

A SnpMatrix object with the GWAS information.

Value

An igraph network of the GS network of the SNPs.

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References

Azencott, C. A., Grimm, D., Sugiyama, M., Kawahara, Y., & Borgwardt, K. M. (2013). Efficient network-guided multi-locus association mapping with graph cuts. Bioinformatics, 29(13), 171-179. https://doi.org/10.1093/bioinformatics/btt238

Examples

```
get_GS_network(minigwas)
```

gwas2bed

Converts a MAP data.frame to a BED data.frame

Description

Takes a map file and:

- column 1: Used as the chromosome column in the BED file.
- column 4: Used as start and end in the BED data.frame (as we work with SNPs).

Usage

```
gwas2bed(gwas)
```

Arguments

gwas

A SnpMatrix object with the GWAS information.

Value

A BED data.frame.

ldweight_edges

Include linkage disequilibrium information in the network.

Description

Include linkage disequilibrium information in the SNP network. The weight of the edges will be lower the higher the linkage is.

```
ldweight_edges(net, ld, method = "inverse")
```

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Arguments

net A SNP network.

ld A dsCMatrix or dgCMatrix containing linkage disequilibrium measures, like

the output of 1d.

method How to incorporate linkage-disequilibrium values into the network.

Value

An copy of net where the edges weighted according to linkage disequilibrium.

Examples

```
ld <- snpStats::ld(minigwas[['genotypes']], depth = 2, stats = "R.squared")
# don't weight edges for which LD cannot be calculated
ld[is.na(ld)] <- 0
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
ldGi <- ldweight_edges(gi, ld)</pre>
```

maxflow

Maxflow algorithm

Description

Run the maxflow algorithm.

Usage

```
maxflow(A, As, At)
```

Arguments

A A sparse matrix with the connectivity.

As A vector containing the edges to the source.

At A vector containing the edges to the sink.

Value

A list with vector indicating if the feature was selected and the objective score.

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mincut_c

Min-cut algorithm

Description

Run the mincut algorithm.

Usage

```
mincut_c(c, eta, lambda, W)
```

Arguments

c A vector with the association of each SNP with the phenotype.

eta A numeric with the value of the eta parameter.

lambda A numeric with the value of the eta parameter.

W A sparse matrix with the connectivity.

Value

A list with vector indicating if the feature was selected and the objective score.

minigwas

Description of the minigwas dataset.

Description

Small GWAS example.

Format

A list with 3 items:

genotypes Genotype and phenotype information.

fam Simulated network.

map Result of runing find_cones with gwas and net.

Examples

```
data(minigwas)

# access different elements
minigwas[["genotypes"]]
minigwas[["map"]]
minigwas[["fam"]]
```

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minippi

PPIs for the minigwas dataset.

Description

data.frame describing pairs of proteins that interact for minigwas.

Examples

```
data(minippi)
```

head(minippi)

minisnpMapping

Genes for the minigwas dataset.

Description

data.frame that maps SNPs from minigwas to their gene.

Examples

```
data(minisnpMapping)
head(minisnpMapping)
```

plot_ideogram

Ideogram of SConES results.

Description

Create a circular ideogram of the a network results using the circlize package (Gu et al., 2014).

Usage

```
plot_ideogram(gwas, net, covars = data.frame(), genome = "hg19")
```

Arguments

gwas A SnpMatrix object with the GWAS information.

net An igraph network that connects the SNPs.

covars A data frame with the covariates. It must contain a column 'sample' containing

the sample IDs, and an additional columns for each covariate.

genome Abbreviations of the genome to use: hg19 for human (default), mm10 for mouse,

etc.

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Value

A circular ideogram, including the manhattan plot, and the interactions between the selected SNPs.

References

Gu, Z., Gu, L., Eils, R., Schlesner, M., & Brors, B. (2014). circlize Implements and enhances circular visualization in R. Bioinformatics (Oxford, England), 30(19), 2811-2. https://doi.org/10.1093/bioinformatics/btu393

scones

Find connected explanatory SNPs

Description

Finds the SNPs maximally associated with a phenotype while being connected in an underlying network.

Usage

```
scones(
  gwas,
  net,
  eta,
  lambda,
  covars = data.frame(),
  score = c("chi2", "glm", "r2"),
  family = c("binomial", "poisson", "gaussian", "gamma"),
  link = c("logit", "log", "identity", "inverse")
)
```

Arguments

gwas	A SnpMatrix object with the GWAS information.
net	An igraph network that connects the SNPs.
eta	Value of the eta parameter.
lambda	Value of the lambda parameter.
covars	A data frame with the covariates. It must contain a column 'sample' containing the sample IDs, and an additional columns for each covariate.
score	Association score to measure association between genotype and phenotype. Possible values: $chi2$ (default), glm .
family	A string defining the generalized linear model family. This should match one of "binomial", "poisson", "gaussian" or "gamma". See snp.rhs.tests for details.
link	A string defining the link function for the GLM. This should match one of "logit", "log", "identity" or "inverse". See spp.rhs.tests for details.

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Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

References

Azencott, C. A., Grimm, D., Sugiyama, M., Kawahara, Y., & Borgwardt, K. M. (2013). Efficient network-guided multi-locus association mapping with graph cuts. Bioinformatics, 29(13), 171-179. https://doi.org/10.1093/bioinformatics/btt238

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
scones(minigwas, gi, 10, 1)</pre>
```

scones.cv

Find connected explanatory SNPs.

Description

Finds the SNPs maximally associated with a phenotype while being connected in an underlying network. Select the hyperparameters by cross-validation.

```
scones.cv(
  gwas,
  net,
  covars = data.frame(),
  score = c("chi2", "glm", "r2"),
  criterion = c("stability", "bic", "aic", "aicc", "global_clustering",
      "local_clustering"),
  etas = numeric(),
  lambdas = numeric(),
  family = c("binomial", "poisson", "gaussian", "gamma"),
  link = c("logit", "log", "identity", "inverse"),
  max_prop_snp = 0.5
)
```

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Arguments

gwas	A SnpMatrix object with the GWAS information.
net	An igraph network that connects the SNPs.
covars	A data frame with the covariates. It must contain a column 'sample' containing the sample IDs, and an additional columns for each covariate.
score	Association score to measure association between genotype and phenotype. Possible values: chi2 (default), glm.
criterion	String with the function to measure the quality of a split.
etas	Numeric vector with the etas to explore in the grid search. If ommited, it's automatically created based on the association scores.
lambdas	Numeric vector with the lambdas to explore in the grid search. If ommited, it's automatically created based on the association scores.
family	A string defining the generalized linear model family. This should match one of "binomial", "poisson", "gaussian" or "gamma". See snp.rhs.tests for details.
link	A string defining the link function for the GLM. This should match one of "logit", "log", "identity" or "inverse". See snp.rhs.tests for details.
max_prop_snp	Maximum proportion of SNPs accepted in the model (between 0 and 1). Larger solutions will be discarded.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

References

Azencott, C. A., Grimm, D., Sugiyama, M., Kawahara, Y., & Borgwardt, K. M. (2013). Efficient network-guided multi-locus association mapping with graph cuts. Bioinformatics, 29(13), 171-179. https://doi.org/10.1093/bioinformatics/btt238

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
scones.cv(minigwas, gi)
scones.cv(minigwas, gi, score = "glm")</pre>
```

scones_

scones.cv_

Find connected explanatory features

Description

Finds the features maximally associated with a phenotype while being connected in an underlying network. Select the hyperparameters by cross-validation.

Usage

```
scones.cv_(X, y, featnames, net)
```

Arguments

X n x d design matrix

y Vector of length n with the outcomes

featnames Vector of length d with the feature names net An igraph network that connects the SNPs.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

Examples

```
X <- as(minigwas[['genotypes']], 'numeric')
X <- X + matrix(rnorm(2500, sd = 0.1), nrow(X), ncol(X))
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
scones.cv_(X, minigwas[['fam']]$affected, minigwas[['map']]$snp, gi)</pre>
```

scones_

Find connected explanatory features

Description

Finds the features maximally associated with a phenotype while being connected in an underlying network.

```
scones_(X, y, featnames, net, eta, lambda)
```

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Arguments

X	n x d design matrix
У	Vector of length n with the outcomes
featnames	Vector of length d with the feature names
net	An igraph network that connects the SNPs.
eta	Value of the eta parameter.
lambda	Value of the lambda parameter.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

Examples

```
X <- as(minigwas[['genotypes']], 'numeric')
X <- X + matrix(rnorm(2500, sd = 0.1), nrow(X), ncol(X))
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
scones_(X, minigwas[['fam']]$affected, minigwas[['map']]$snp, gi, 10, 1)</pre>
```

search_cones

Find connected explanatory SNPs.

Description

Finds the SNPs maximally associated with a phenotype while being connected in an underlying network (Azencott et al., 2013).

```
search_cones(
   gwas,
   net,
   encoding = "additive",
   sigmod = FALSE,
   covars = data.frame(),
   associationScore = c("chi2", "glm"),
   modelScore = c("stability", "bic", "aic", "aicc", "global_clustering",
        "local_clustering"),
   etas = numeric(),
   lambdas = numeric()
)
```

search_cones

Arguments

gwas A SnpMatrix object with the GWAS information.

net An igraph network that connects the SNPs.

encoding SNP encoding (unused argument).

sigmod Boolean. If TRUE, use the Sigmod variant of SConES, meant to prioritize

tightly connected clusters of SNPs.

covars A data frame with the covariates. It must contain a column 'sample' containing

the sample IDs, and an additional columns for each covariate.

associationScore

Association score to measure association between genotype and phenotype.

modelScore String with the function to measure the quality of a split.

etas Numeric vector with the etas to explore in the grid search. If ommited, it's

automatically created based on the association scores.

lambdas Numeric vector with the lambdas to explore in the grid search. If ommited, it's

automatically created based on the association scores.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

References

Azencott, C. A., Grimm, D., Sugiyama, M., Kawahara, Y., & Borgwardt, K. M. (2013). Efficient network-guided multi-locus association mapping with graph cuts. Bioinformatics, 29(13), 171-179. https://doi.org/10.1093/bioinformatics/btt238

Examples

```
## Not run: gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
search_cones(minigwas, gi)
search_cones(minigwas, gi, encoding = "recessive")
search_cones(minigwas, gi, associationScore = "skat")
## End(Not run)</pre>
```

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sigmod	Find connected explanatory SNPs
G	*

Description

Finds the SNPs maximally associated with a phenotype while being connected in an underlying network.

Usage

```
sigmod(
  gwas,
  net,
  eta,
  lambda,
  covars = data.frame(),
  score = c("chi2", "glm", "r2"),
  family = c("binomial", "poisson", "gaussian", "gamma"),
  link = c("logit", "log", "identity", "inverse")
)
```

Arguments

gwas	A SnpMatrix object with the GWAS information.
net	An igraph network that connects the SNPs.
eta	Value of the eta parameter.
lambda	Value of the lambda parameter.
covars	A data frame with the covariates. It must contain a column 'sample' containing the sample IDs, and an additional columns for each covariate.
score	Association score to measure association between genotype and phenotype. Possible values: chi2 (default), glm.
family	A string defining the generalized linear model family. This should match one of "binomial", "poisson", "gaussian" or "gamma". See snp.rhs.tests for details.
link	A string defining the link function for the GLM. This should match one of "logit", "log", "identity" or "inverse". See snp.rhs.tests for details.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

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References

Liu, Y., Brossard, M., Roqueiro, D., Margaritte-Jeannin, P., Sarnowski, C., Bouzigon, E., Demenais, F. (2017). SigMod: an exact and efficient method to identify a strongly interconnected disease-associated module in a gene network. Bioinformatics, 33(10), 1536–1544. https://doi.org/10.1093/bioinformatics/btx004

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
sigmod(minigwas, gi, 10, 1)</pre>
```

sigmod.cv

Find connected explanatory SNPs.

Description

Finds the SNPs maximally associated with a phenotype while being connected in an underlying network. Select the hyperparameters by cross-validation.

Usage

```
sigmod.cv(
  gwas,
  net,
  covars = data.frame(),
  score = c("chi2", "glm", "r2"),
  criterion = c("stability", "bic", "aic", "aicc", "global_clustering",
       "local_clustering"),
  etas = numeric(),
  lambdas = numeric(),
  family = c("binomial", "poisson", "gaussian", "gamma"),
  link = c("logit", "log", "identity", "inverse"),
  max_prop_snp = 0.5
)
```

Arguments

gwas	A SnpMatrix object with the GWAS information.
net	An igraph network that connects the SNPs.
covars	A data frame with the covariates. It must contain a column 'sample' containing the sample IDs, and an additional columns for each covariate.
score	Association score to measure association between genotype and phenotype. Possible values: chi2 (default), glm.
criterion	String with the function to measure the quality of a split.
etas	Numeric vector with the etas to explore in the grid search. If ommited, it's automatically created based on the association scores.

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lambdas	Numeric vector with the lambdas to explore in the grid search. If ommited, it's automatically created based on the association scores.
family	A string defining the generalized linear model family. This should match one of "binomial", "poisson", "gaussian" or "gamma". See snp.rhs.tests for details.
link	A string defining the link function for the GLM. This should match one of "logit", "log", "identity" or "inverse". See snp.rhs.tests for details.
max_prop_snp	Maximum proportion of SNPs accepted in the model (between 0 and 1). Larger solutions will be discarded.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

References

Liu, Y., Brossard, M., Roqueiro, D., Margaritte-Jeannin, P., Sarnowski, C., Bouzigon, E., Demenais, F. (2017). SigMod: an exact and efficient method to identify a strongly interconnected disease-associated module in a gene network. Bioinformatics, 33(10), 1536–1544. https://doi.org/10.1093/bioinformatics/btx004

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
sigmod.cv(minigwas, gi)
sigmod.cv(minigwas, gi, score = "glm")</pre>
```

sigmod.cv_

Find connected explanatory features

Description

Finds the features maximally associated with a phenotype while being connected in an underlying network. Select the hyperparameters by cross-validation.

Usage

```
sigmod.cv_(X, y, featnames, net)
```

Arguments

Χ	n x d design matrix
у	Vector of length n with the outcomes
featnames	Vector of length d with the feature names
net	An igraph network that connects the SNPs.

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Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

Examples

```
X <- as(minigwas[['genotypes']], 'numeric')
X <- X + matrix(rnorm(2500, sd = 0.1), nrow(X), ncol(X))
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
sigmod.cv_(X, minigwas[['fam']]$affected, minigwas[['map']]$snp, gi)</pre>
```

sigmod_

Find connected explanatory features

Description

Finds the features maximally associated with a phenotype while being connected in an underlying network.

Usage

```
sigmod_(X, y, featnames, net, eta, lambda)
```

Arguments

x n x d design matrix

y Vector of length n with the outcomes
featnames Vector of length d with the feature names
net An igraph network that connects the SNPs.

eta Value of the eta parameter.

lambda Value of the lambda parameter.

Value

A copy of the SnpMatrix\$map data.frame, with the following additions:

- c: contains the univariate association score for every single SNP.
- selected: logical vector indicating if the SNP was selected by SConES or not.
- module: integer with the number of the module the SNP belongs to.

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Examples

```
X <- as(minigwas[['genotypes']], 'numeric')
X <- X + matrix(rnorm(2500, sd = 0.1), nrow(X), ncol(X))
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
sigmod_(X, minigwas[['fam']]$affected, minigwas[['map']]$snp, gi, 10, 1)</pre>
```

Description

Selects randomly interconnected genes as causal, then selects a proportion of them as causal.

Usage

```
simulate_causal_snps(net, ngenes = 20, pcausal = 1)
```

Arguments

net An igraph gene-interaction (GI) network that connects the SNPs.

ngenes Number of causal genes.

pcausal Number between 0 and 1, proportion of the SNPs in causal genes that are causal

themselves.

Value

A vector with the ids of the simulated causal SNPs.

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
simulate_causal_snps(gi, ngenes=2)</pre>
```

simulate_phenotype
Simulate phenotype

Description

Simulates a phenotype from a GWAS experiment and a specified set of causal SNPs. If the data is qualitative, only controls are used.

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Usage

```
simulate_phenotype(
  gwas,
  snps,
  h2,
  model = "additive",
  effectSize = rnorm(length(snps)),
  qualitative = FALSE,
  ncases,
  ncontrols,
  prevalence
)
```

Arguments

gwas A SnpMatrix object with the GWAS information.

snps Character vector with the SNP ids of the causal SNPs. Must match SNPs in

gwas[["map"]][["snp.name"]].

h2 Heritability of the phenotype (between 0 and 1).

model String specifying the genetic model under the phenotype. Accepted values: "ad-

ditive".

effectSize Numeric vector with the same length as the number of causal SNPs. It indicates

the effect size of each of the SNPs; if absent, they are sampled fron a normal

distribution.

qualitative Bool indicating if the phenotype is qualitative or not (quantitative).

ncases Integer specifying the number of cases to simulate in a qualitative phenotype.

Required if qualitative = TRUE.

ncontrols Integer specifying the number of controls to simulate in a qualitative phenotype.

Required if qualitative = TRUE.

prevalence Value between 0 and 1 specifying the population prevalence of the disease. Note

that neases cannot be greater than prevalence * number of samples. Required if

qualitative = TRUE.

Value

A copy of the GWAS experiment with the new phenotypes in gwas[["fam"]][["affected"]].

References

Inspired from GCTA simulation tool: http://cnsgenomics.com/software/gcta/Simu.html.

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
causal <- simulate_causal_snps(gi, ngenes = 2)
simulate_phenotype(minigwas, causal, h2 = 1)</pre>
```

subvert 21

subvert	Vertices with an attribute
---------	----------------------------

Description

Returns the nodes matching some condition.

Usage

```
subvert(net, attr, values, affirmative = TRUE)
```

Arguments

net An igraph network.

attr An attribute of the vertices.
values Possible values of attr

affirmative Logical. States if a condition must be its affirmation (e.g. all nodes with gene

name "X"), or its negation (all nodes not with gene name "X").

Value

The vertices with attribute equal to any of the values in values.

Examples

```
gi <- get_GI_network(minigwas, snpMapping = minisnpMapping, ppi = minippi)
martini:::subvert(gi, "gene", "A")
martini:::subvert(gi, "name", c("1A1", "1A3"))</pre>
```

wrap_Xy

Make pseudo SnpMatrix object

Description

Wrap design matrix and outcome vector into a pseudo SnpMatrix object.

Usage

```
wrap_Xy(X, y, featnames, net)
```

Arguments

X n x d design matrix

y Vector of length n with the outcomes

featnames Vector of length d with the feature names

net An igraph network that connects the SNPs.

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