Package 'vartools'

August 21, 2015

Type Package

| Title Variant Association Tools R-package | |
|---|----|
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| Depends survival | |
| Description Computes statistics of rare vatiants using CMC, KBAC, VT and other association methods | |
| License GPL | |
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vartools-package

Variant Association tools

Description

vartools is an R-package for the analysis of rare variants and currently has the following methods: CMC, KBAC, RVE, VT, RareCover, RBT, C-alpha.

Installation

git clone https://github.com/izhbannikov/vartools.git R CMD INSTALL vartools

Details

Package: vartools
Type: Package
Version: 1.0

Date: 2014-12-10 License: GPL (>= 2)

Project URL: https://github.com/izhbannikov/vartools

Author(s)

Ilya Y. Zhbannikov

Maintainer: Ilya Y. Zhbannikov | i.zhbannikov@mail.ru

```
#CMC test
# Load the package
library(vartools)
pgdata <- as.matrix(read.table(system.file("extdata", "phengen.dat", package="vartools"), as.is=T, skip = 1))</pre>
cmc.pvalue <- cmc(table=pgdata)</pre>
print(cmc.pvalue)
#KBAC test
?kbac
alpha <- 0.05
num.permutation <- 3000
quiet <- 1
alternative <- 1
maf.upper.bound <- 0.05</pre>
casectrl.dat <- read.table(system.file("extdata", "phengen.dat", package="vartools"), skip = 1)</pre>
kbac.pvalue <- kbac(table=casectrl.dat, alpha, num.permutation, quiet, maf.upper.bound, alternative)</pre>
print(kbac.pvalue)
```

asum 3

| asum aSum statistical test |
|----------------------------|
|----------------------------|

Description

aSum, Adaptive Sum Test (Han and Pan, 2010) is a statistical test which utilizes the difference in direction of effects (protective or deleterious) of rare variants within the same genetic region analyzed by a rare variant association test.

Usage

```
asum(table, perm=100)
```

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'.

perm positive integer that defines the number of permutations. In permutation test, the

distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrangements of the labels on

the observed data points.

Details

• • • •

Value

A list with the following elements:

asum.stat aSum statistic

perm.pval permuted p-value; only when permutations is used

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Fang Han and Wei Pan (2010) A Data-Adaptive Sum Test for Disease Association with Multiple Common or Rare Variants. Human Heredity doi:10.1159/000288704. http://www.karger.com/doi/10.1159/000288704

```
# Load the package
library(vartools)
?asum

casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)
asum.stat <- asum(casectrl.dat)
asum.stat</pre>
```

4 calpha

| calpha | C(alpha) statistical test | |
|--------|---------------------------|--|
| Сатрпа | C(aipna) siansucai iesi | |

Description

C(alpha) (Neale at al., 2011) is a statistical test, developed for disease traits, in order to test for the hypothesis of rare variants disease association. This test has an assumption that rare variants, which were observed in cases and controls and are a mix of various types of variants: phenotypically deleterious, protective and neutral variants.

Usage

```
calpha(table, permutations=NULL)
```

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'.

permutations positive integer that defines the number of permutations. In permutation test, the

distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrangements of the labels on

the observed data points.

Details

....

Value

A list with the following elements:

calpha.stat C(alpha) statistic

asym.pval asymptotic p-value, distributed as Chi-square, with parameter degrees of free-

dom df=1

perm.pval permuted p-value; only when perm is used

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Benjamin M. Neale, Manuel A. Rivas, Benjamin F. Voight, David Altshuler, Bernie Devlin, Marju Orho-Melander, Sekar Kathiresan, Shaun M. Purcell, Kathryn Roeder and Mark J. Daly (2011), Testing for an Unusual Distribution of Rare Variants. PLoS Genetics doi:10.1371/journal.pgen.1001322. http://dx.plos.org/10.1371/journal.pgen.1001322

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Examples

```
# Load the package
library(vartools)
?calpha

casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)
calpha.stat <- calpha(casectrl.dat)
calpha.stat</pre>
```

cmc

CMC statistic implementation

Description

This program implements the CMC statistic in [Liu and Leal 2008]. It carries out case-control association testing for rare variants for whole exome association studies. Briefly, consider a gene of length n which harbors m rare variants. Genotype on the m variant sites & the disease status (case/control) are known for each individual. The program takes as input the m-site genotype and disease status (case/control) data files, and computes a p-value indicating the significance of association. In order to speed up permutation testing we use an "adaptive" approach to obtain p-values.

Usage

```
cmc(table, method = "fisher")
```

Arguments

table

a numeric matrix with first column having disease status '0' or '1' and the rest columns codes the locus genotype as '0', '1', and '2'. DO NOT allow for miss-

ing data.

method

statistical method, Fisher test used by default

Details

....

Value

pvalue

the p-value of test.

Author(s)

Ilya Y. Zhbannikov | i.zhbannikov@mail.ru

References

Liu DJ, Leal SM, 2008 Methods for Detecting Associations with Rare Variants for Common Diseases: Application to Analysis of Sequence Data. The American Journal of Human Genetics, 83, DOI10.1016/j.ajhg.2008.06.024

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Examples

```
# Load the package
library(vartools)
?cmc

pgdata <- as.matrix(read.table(system.file("extdata","phengen.dat",package="vartools"), as.is=T, skip = 1))
cmc.pvalue <- cmc(table=pgdata)
print(cmc.pvalue)</pre>
```

coxKM

SNP-set kernel association test for right-censored survival outcomes.

Description

Tests for association between a set of common SNPS and a right-censored survival outcome. Warnings: (1) coxKM is meant for common genetic variants, (2) for very small p-values, it is necessary to increase no. of perturbations.

Usage

```
coxKM(Z=NULL, U, Delta, X=NULL, gamma=NULL, kernel="linear", weights=NULL,
npert=10^4, npert.check=TRUE, npert.upper=10^8, npert.threshold=50,
impute.method = "fixed", is_check_genotype=TRUE,
is_dosage=FALSE, missing_cutoff=0.15, SetID=NULL)
```

Arguments

| X | is a nxR matrix of relevant covariates with each row as a different individual and each column as a separate covariate measurement. If no additional covariates are present, X can be left unspecified or left as NULL. Note that each column of X has to be a numerical variable, non-numerical variables have to be recoded appropriately before analysis. X should not include an intercept. |
|--------|--|
| Z | is a nxS numeric genotype matrix with each row as a different individual and each column as a separate snp. Each genotype should be coded as 0, 1, 2, and 9 (or NA) for AA, Aa, aa, and missing, where A is a major allele and a is a minor allele. Missing genotypes will be imputed by the simple Hardy-Weinberg equilibrium (HWE) based imputation. If kernel matrix is supplied, Z is ignored and not used in testing. |
| U | is a nx1 vector containing the observed times. Note: $U=min(C,T)$ where $C=$ censoring time, $T=$ survival time |
| Delta | is a nx1 vector containing the status/event indicator. |
| gamma | Unless $X = NULL$, gamma has to be supplied. gamma is the vector of coefficients from the null cox model corresponding to X . gamma \leftarrow coxph(Surv(U,Delta) \sim X)\$coef |
| kernel | Type of kernel. kernel can be an nxn kernel matrix OR one of these six options: "linear.weighted", "linear", "IBS", "IBS.weighted", "quadratic" or "2wayIX". If an nxn kernel matrix is supplied, Z is ignored and is not used in testing. |

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weights is a vector of length S of prespecified weights for the weighted kernels. Weights

in coxKM are defined the same way as in SKAT. The kernel matrix of the

weighted linear kernel is K=ZWWZ'.

is the number of perturbations used to calculate p-value (default =10000), npert npert

> should be at least 1000. Note that how small the p-value can be is limited by the number of perturbations. If npert.check = FALSE, the smallest possible p-value is 0.5/npert. If npert.check = TRUE, the smallest possible p-value is 0.5/(ceiling(npert.upper/10^4)*10^4). For very small p-values, to obtain accurate p-values, it is necessary to increase the number of perturbations. See

npert.check.

TRUE/FALSE (default=TRUE). If npert.check=TRUE, coxKM first uses npert npert.check

> perturbations to obtain an initial p-value and checks to see if the initial p-value <= npert.threshold/npert. If the initial p-value <= npert.threshold/npert, then npert.upper perturbations is used to obtain a more accurate p-value. Setting npert.check=TRUE allows a larger number of perturbations to be used to obtain more accurate p-values only when it is necessary. For very small p-values, it

may be necessary to further increase npert.upper.

npert.upper default=10^8. Used only if npert.check=TRUE. See npert.check.

npert.threshold

default=50. Used only if npert.check=TRUE. See npert.check.

impute.method a method to impute missing genotypes (default= "fixed"). "random" imputes

> missing genotypes by generating binomial(2,p) random variables (p is the MAF), and "fixed" imputes missing genotypes by assigning the mean genotype value (2p). If you use "random", you will have different p-values for different runs because imputed values are randomly assigned. Can use set.seed() to replicate

results.

is_check_genotype

a logical value indicating whether to check the validity of the genotype matrix Z (default= TRUE). If you use non-SNP type data and want to run coxKM, please set it to FALSE. If you use SNP data or imputed data, please set it to TRUE. If is_check_genotype=FALSE, missing values in Z have to be coded only as NA

since 9 will not be treated as a missing value.

is_dosage a logical value indicating whether the matrix Z is a dosage matrix (default=

> FALSE). If is_dosage=TRUE, "is_check_genotype" and "impute.method" will be ignored and coxKM will check the genotype matrix and set impute.method="fixed".

Note that coxKM will also treat 9 as missing in Z.

missing_cutoff a cutoff of the missing rates of SNPs (default=0.15). Any SNPs with missing

rates higher than cutoff will be excluded from the analysis.

SetID SetID.

Details

If kernel is not a matrix and Z is supplied, and either is check genotype=TRUE OR is dosage=TRUE, coxKM will check the Z matrix for missing values (missing values must be coded either as NA or 9) and apply imputation. If you are using coxKM for non-SNP/dosage data, set is_check_genotype=FALSE and is_dosage=FALSE, in which case missing values must be coded as NA (9 is not considered a missing value).

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Value

p.value the p-value of coxKM based on resampling. Note that if the p-value takes on the smallest possible value based on the number of perturbations, it may be necessary to increase npert and npert.upper. See npert.check.

Q the unscaled score test statistic of coxKM.

 $\mbox{n.marker.test} \quad \mbox{no. of SNPs used for testing,} <= \!\! S.$

n.indiv n = no. of samples

df the estimated degrees of freedom of the test statistic (for reference only, not used

in association testing).

Author(s)

Xinyi (Cindy) Lin, Qian Zhou

References

Lin X, Cai T, Wu M, Zhou Q, Liu G, Christiani D and Lin X. 2011. Survival Kernel Machine SNP-set Analysis for Genome-wide Association Studies. Genetic Epidemiology 35:620-31. doi: 10.1002/gepi.20610

Cai T, Tonini G and Lin X. 2011. Kernel machine approach to testing the significance of multiple genetic markers for risk prediction. Biometrics, 67:975-86. doi:10.1111/j.1541-0420.2010.01544.x

```
\verb|data| (examples npset, example covariates, example phenotype 1, example phenotype 2, example phenotype 3)|
Z <- as.matrix(examplesnpset)</pre>
X <- as.matrix(examplecovariates)</pre>
phenotype1 <- examplephenotype1</pre>
phenotype2 <- examplephenotype2</pre>
phenotype3 <- examplephenotype3</pre>
set.seed(1)
#-----
# coxKM without covariates
#-----
coxKM(Z=Z, U=phenotype1$time, Delta=phenotype1$event, kernel="IBS")
coxKM(Z=Z, U=phenotype1$time, Delta=phenotype1$event, kernel="linear")
coxKM(Z=Z, U=phenotype3$time, Delta=phenotype3$event, kernel="IBS")
coxKM(Z=Z, U=phenotype3$time, Delta=phenotype3$event, kernel="linear")
# coxKM with covariates
#-----
Gamma <- coxph(Surv(phenotype2$time, phenotype2$event)~X)$coef</pre>
coxKM(Z=Z, U=phenotype2$time, Delta=phenotype2$event, X=X, gamma=Gamma, kernel="IBS")
coxKM(Z=Z, U=phenotype2$time, Delta=phenotype2$event, X=X, gamma=Gamma, kernel="linear")
```

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examplecovariates

Example covariates dataset for coxKM.

Description

Example covariates dataset for coxKM.

Format

examplecovariates contains:

a numeric matrix of 2000 individuals and 2 covariates. Each row represents a different individual. coxKM.examplecovariates is identical to X in SKAT.example.

Author(s)

Xinyi (Cindy) Lin

examplephenotype1

Example phenotype for coxKM.

Description

Example phenotype for coxKM.

Format

examplephenotype1 contains:

a numeric matrix with two columns, the first column is the event time, the second column is the status indicator. Each row represents a different individual.

Author(s)

Xinyi (Cindy) Lin

examplephenotype2

Example phenotype for coxKM.

Description

Example phenotype for coxKM.

Format

examplephenotype2 contains:

a numeric matrix with two columns, the first column is the event time, the second column is the status indicator. Each row represents a different individual.

Author(s)

Xinyi (Cindy) Lin

10 kbac

Description

Example phenotype for coxKM.

Format

examplephenotype3 contains:

a numeric matrix with two columns, the first column is the event time, the second column is the status indicator. Each row represents a different individual.

Author(s)

Xinyi (Cindy) Lin

| Example SNP-set for coxKM. |
|----------------------------|
|----------------------------|

Description

Example SNP-set for coxKM.

Format

examplesnpset contains:

a numeric genotype matrix of 2000 individuals and 11 SNPs. Each row represents a different individual, and each column represents a different SNP marker. coxKM.examplesnpset is subset of Z in SKAT.example.

Author(s)

Xinyi (Cindy) Lin

| kbac | KBAC statistic implementation | |
|------|-------------------------------|--|
| | | |

Description

This program implements the KBAC statistic in [Liu and Leal 2010]. It carries out case-control association testing for rare variants for whole exome association studies. Briefly, consider a gene of length n which harbors m rare variants. Genotype on the m variant sites & the disease status (case/control) are known for each individual. The program takes as input the m-site genotype and disease status (case/control) data files, and computes a p-value indicating the significance of association. In order to speed up permutation testing we use an "adaptive" approach to obtain p-values.

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Usage

kbac(table, alpha = NULL, num.permutation = NULL, quiet = T, maf.upper.bound = 1.0, alternative = 1

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'. DO NOT allow for miss-

ing data.

alpha size of test, or the significant level. This feature will be useful in adaptive p-

value calculation. If you do not want to use adaptive p-value, set alpha = 999 (or

any number greater than 1.0).

num.permutation

number of permutations for p-value calculation. In permutation test, the distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrangements of the labels on the

observed data points.

quiet when quiet = 0 the screen output would contain a summary of the KBAC test;

otherwise only the p-value will be printed on screen.

maf.upper.bound

The upper bound of the MAF to be included in analysis. MAF is calculated based on observed sample. This can be arbitary although it is usually defined as

0.01 for analysis of rare variants.

alternative Set alternative = 1 for test of deleterious variants, = 2 for test of both deleterious

and protective variants. Please note that this is different from the "one/two-

sided" definition in the KBAC paper.

Details

....

Value

pvalue the p-value of test.

alternative can have 0 or 1, depending on hyphotesis.

kbac.stat test statistics.

Author(s)

Gao Wang | wangow@gmail.com

References

Liu DJ, Leal SM, 2010 A Novel Adaptive Method for the Analysis of Next-Generation Sequencing Data to Detect Complex Trait Associations with Rare Variants Due to Gene Main Effects and Interactions. PLoS Genet 6(10): e1001156. doi:10.1371/journal.pgen.1001156

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Examples

```
# Load the package
library(vartools)
?kbac
# Set parameters and use the kbac() function to obtain p-value
alpha <- 0.05
num.permutation <- 3000
quiet <- 1
alternative <- 1
maf.upper.bound <- 0.05
casectrl.dat <- read.table(system.file("extdata", "phengen.dat", package="vartools"), skip = 1)</pre>
res <- kbac(table=casectrl.dat, alpha, num.permutation, quiet, maf.upper.bound, alternative)</pre>
print(res)
# To evaluate test at small alpha we need huge number of permutations. Adaptive approach is thus necessary.
casectrl.dat <- read.table(system.file("extdata", "phengen.dat", package="vartools"), skip = 1)</pre>
res <- kbac(table=casectrl.dat, 0.00001, 1000000, quiet, maf.upper.bound, alternative)
print(res)
# Not using adaptive p-value calculation; will take longer time
casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)</pre>
kbac.pvalue <- kbac(table=casectrl.dat, 9, 1000000, quiet, maf.upper.bound, alternative)</pre>
print(kbac.pvalue)
```

rarecover

RareCover statistical test

Description

Testing for rare variants with RareCover algorithm. This algorithm is similar to CMC, meaning that it follows its collapsing strategy, but uses greedy algorithm to find an optimized combination of variants in a loci for which its association signal is strongest.

Usage

```
rarecover(table, maf=0.05, dif=0.5, perm=250)
```

Arguments

| table | a numeric matrix with first column having disease status '0' or '1' and the rest columns codes the locus genotype as '0', '1', and '2'. |
|-------|--|
| maf | numeric value indicating the minor allele frequency threshold for rare variants |
| dif | numeric value between 0 and 1 as a threshold for the decision criterion in the RareCover algorithm |
| perm | positive integer that defines the number of permutations. In permutation test, the distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrangements of the labels on the observed data points. |

Details

...

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Value

A list with the following elements:

rc.stat RareCover statistic
perm.pval permuted p-value
set set of selected variants

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Bhatia G, Bansal V, Harismendy O, Schork NJ, Topol EJ, Frazer K, Bafna V (2010) A Covering Method for Detecting Genetic Associations between Rare Variants and Common Phenotypes. *PLoS Computational Biology*, **6(10)**: e1000954

Examples

```
# Load the package
library(vartools)
?rarecover

casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)
rarecover.stat <- rarecover(casectrl.dat)
rarecover.stat</pre>
```

rbt

Replication-Based statistical test

Description

Implementation of the replication base test (RBT) proposed by Ionita-Laza et al., 2011. Replication means that it computes evidences to reject each of the two hypothesis: (1) Deleterious rare variants are enriched in cases; (2) Protective rare variants are enriched in controls.

Usage

```
rbt(table, perm=150)
```

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'.

perm positive integer that defines the number of permutations, 150 by default. In per-

mutation test, the distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrange-

ments of the labels on the observed data points.

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Details

....

Value

A list with the following elements:

rbt.stat RBT statistic
perm.pval permuted p-value

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Ionita-Laza I, Buxbaum JD, Laird NM, Lange C (2011) A New Testing Strategy to Identify Rare Variants with Either risk or Protective Effects on Disease. *PLoS Genetics*, **7(2)**: e1001289

Examples

```
# Load the package
library(vartools)
?rbt

casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)
rbt.stat <- rbt(casectrl.dat)
rbt.stat</pre>
```

skat

SKAT statistical test

Description

SKAT (Wu at al., 2010, 2011) is a statistical test, developed for disease traits, in order to test for the hypothesis of rare variants disease association. This is a score-based variance-component test.

Usage

```
skat(table, kernel="linear", weights=NULL, a=1, b=25, permutations=NULL)
```

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'.

kernel Kernel function used. By default, a linear function is used. Other available

kernels are: "wlinear", "quadratic", "IBS", "wIBS", "twowayx"

weights a set of numeric weights for genetic variants.

a a positive numeric value for the parameter a in the Beta distribution (a=1 by

default)

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b a positive numeric value for the parameter b in the Beta distribution (b=1 by

default)

permutations positive integer that defines the number of permutations. In permutation test, the

distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrangements of the labels on

the observed data points.

Details

....

Value

A list with the following elements:

skat.stat SKAT statistic

asym.pval asymptotic p-value, distributed as Chi-square, with parameter degrees of free-

dom df=1

perm.pval permuted p-value; only when permutations is used

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Wu MC, Kraft P, Epstein MP, Taylor DM, Chanock SJ, Hunter DJ, Lin X (2010) Powerful SNP-Set Analysis for Case-Control Genome-wide Association Studies. *The American Journal of Human Genetics*, **86**: 929-942

Wu MC, Lee S, Cai T, Li Y, Boehnke M, Lin X (2011) Rare-Variant Association Testing for Sequencing Data with the Sequence Kernel Association Test. *The American Journal of Human Genetics*, **89**: 82-93

Examples

```
# Load the package
library(vartools)
?skat

casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)
skat.stat <- skat(casectrl.dat)
skat.stat</pre>
```

٧t

Variable Threshold statistical test

Description

This is implementation for the Variable Threshold statistical test. It assigns different weights to rare variants base on their frequencies.

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Usage

```
vt(table, maf=0.05, perm=50)
```

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'.

maf numeric value indicating the minor allele frequency threshold for rare variants

(must be a positive number between 0 and 1, maf=0.05 by default).

perm positive integer that defines the number of permutations, perm=50 by default. In

permutation test, the distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrange-

ments of the labels on the observed data points.

Details

•••

Value

A list with the following elements:

vt.stat VT statistic

perm.pval permuted p-value

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Price AL, Kryukov GV, de Bakker PIW, Purcell SM, Staples J, Wei LJ, Sunyaev SR (2010) Pooled Association Tests for Rare Variants in Exon-Sequencing Studies. *The American Journal of Human Genetics*, **86**: 832-838

```
# Load the package
library(vartools)
?vt

casectrl.dat <- read.table(system.file("extdata","phengen2.dat",package="vartools"), skip = 1)
vt.stat <- vt(casectrl.dat)
vt.stat</pre>
```

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wss WSS statistical test

Description

WSS (Madsen and Browning, 2009) is a statistical test, developed for disease traits, in order to test for the hypothesis of rare variants disease association. WSS test introduces the method of assigning "weights" to rare variants found in a genetic region before they are collapsed.

Usage

```
wss(table, perm=100)
```

Arguments

table a numeric matrix with first column having disease status '0' or '1' and the rest

columns codes the locus genotype as '0', '1', and '2'.

perm positive integer that defines the number of permutations. In permutation test, the

distribution of the test statistic under the null hypothesis is obtained by calculating all possible values of the test statistic under rearrangements of the labels on

the observed data points.

Details

• • • •

Value

A list with the following elements:

wss.stat WSS statistic

perm.pval permuted p-value; only when permutations is used

args descriptive information with number of controls, cases, variants, and permuta-

tions

name of the statistic

References

Bo Eskerod Madsen and Sharon R. Browning (2009) A Groupwise Association Test for Rare Mutations Using a Weighted Sum Statistic. PLoS Genetics doi:10.1371/journal.pgen.1000384. http://dx.plos.org/10.1371/journal.pgen.1000384

```
# Load the package
library(vartools)
?wss

casectrl.dat <- read.table(system.file("extdata","phengen.dat",package="vartools"), skip = 1)
wss.stat <- wss(casectrl.dat)
wss.stat</pre>
```

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