# Package 'MORST'

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Type Package

Title Minimax Optimal Ridge-type Set Test

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<b>Description</b> This package implemented MORST, a fast and powerful test that is designed to be power robust to the strength of signals. While mainly motivated from the linear regression setting, MORST is a generic test and has several versions. This package implemented the score version of MORST in GLMs and a specific function of MORST tailored to genetic association studies. At the core of MORST, it is the choice of the ridge-type parameter tau_c, which is implemented in this package. With tau_c, one can easily develop other versions of MORST or adapt MROST to other applications.
<b>Depends</b> R (>= $3.2.0$ )
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R topics documented:
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Get\_Q\_pval

Get the p-value of a quadratic test

## **Description**

The survival function (i.e., p-value) of the weighted sum of i.i.d. Chi-squared variables with df=1 (i.e., a quadratic test). A hybrid of davies's method, saddle point method, and liu's method is used.

# Usage

```
Get_Q_pval(Q, w)
```

#### **Arguments**

Q The value of the quadratic test statistic. It must be positive.

w a numeric vector of positive weights. See details.

#### **Details**

Compute P[T > Q], where  $T = \sum_{i=1}^k w_i X_i$ ,  $X_i$ 's are i.i.d. Chi-squared variables with df=1, and  $w_i$ 's are the elements of w.

A hybrid of several methods is used to acheive both accuracy and computation effeciency. Specifically, when the p-value > 1e-10, the davies' method is used; when 1e-10 < p-value < 1e-15, the saddle point method is used; when p-value < 1e-15, liu's method is used.

### Value

The p-value of the quadratic test.

# Author(s)

Yaowu Liu

## References

Davies, R. B. (1980). Algorithm AS 155: The distribution of a linear combination of X2 random variables. *Journal of the Royal Statistical Society. Series C (Applied Statistics)*, 29(3):323-333.

Kuonen, D. (1999). Miscellanea. saddlepoint approximations for distributions of quadratic forms in normal variables. *Biometrika*, 86(4):929-935.

Liu, H., Tang, Y., and Zhang, H. H. (2009). A new chi-square approximation to the distribution of non-negative definite quadratic forms in non-central normal variables. *Computational Statistics & Data Analysis*,53(4):853-856.

## **Examples**

```
Get_Q_pval(50, seq(1,10,1))
Get_Q_pval(200, seq(1,10,1))
Get_Q_pval(1000, seq(1,10,1))
```

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MORST_glm	The score version of MORST for Generalized Linear Models.
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## **Description**

Calculate the score version MORST p-value for GLM. For genetic association study, please use the function SetBasedTests.

# Usage

```
MORST_glm(
   X,
   obj,
   alpha = 0.05,
   weights = NULL,
   tau.type = "approx",
   target_power = 0.5,
   n.points = 50
)
```

## **Arguments**

```
a numeric matrix or dgCMatrix of predictors.

an output from Null_model_glm.

alpha the alpha parameter in MORST. It is suggested to be the significance level.

weights a numeric vector of nonnegative weights. If NULL, the equal weight is used.

tau.type either "minimax" or "approx". See tau_c for details.

target_power a value that is used when tau.type == "approx". See tau_c for details.

n.points number of grid points used when tau.type == "minimax". See tau_c for details.
```

#### Value

The p-value of MORST.

## Author(s)

Yaowu Liu

#### References

Liu, Y., Li, Z., and Lin, X. (2020+) A Minimax Optimal Ridge-Type Set Test for Global Hypothesis with Applications in Whole Genome Sequencing Association Studies. *Journal of the American Statistical Association*. Accepted.

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#### **Examples**

```
X<-matrix(rnorm(20000),ncol=20); Z=matrix(rnorm(nrow(X)*4),ncol=4)

### linear regression for continuous outcome
Y<-rnorm(nrow(X));obj<-Null_model_glm(Y,Z,family="gaussian")

MORST_glm(X,obj)

### Logistic regression for binary outcome
Y<-rbinom(nrow(X),1,0.4);obj<-Null_model_glm(Y,Z,family="binomial")

MORST_glm(X,obj,alpha = 1e-04)

### Binomial outcome
Y<-rbinom(nrow(X),5,0.4);Y<-cbind(Y,5-Y);obj<-Null_model_glm(Y,Z,family="binomial")

MORST_glm(X,obj,weights=runif(ncol(X)))

### Poisson outcome
Y<-rpois(nrow(X),5);obj<-Null_model_glm(Y,Z,family="poisson")

MORST_glm(X,obj,alpha = 1e-04,weights=runif(ncol(X)),tau.type = "minimax")</pre>
```

Null\_model\_glm

Fit the GLM null model for MORST

#### **Description**

fit a GLM null model

# Usage

```
Null_model_glm(Y, Z, family = "gaussian", include_intercept = TRUE)
```

## **Arguments**

Y a numeric vector of outcomes.

Z a numeric matrix of covariates that need to be adjusted.

include\_intercept

logical. If TRUE, the intercept will be included in the null model.

famlily a character. Should be "gaussian", "binomial" or "poisson". For each famliy, the

cannonical link is used.

# Value

This function returns an object that has model parameters and residuals of the NULL model of no association between outcomes Y and predictors X after adjusting for covariates Z. After obtaining it, please use MORST\_glm or SetBasedTests to conduct the association test.

## Author(s)

Yaowu Liu

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#### **Examples**

```
X<-matrix(rnorm(20000),ncol=20); Z=matrix(rnorm(nrow(X)*4),ncol=4)
### linear regression for continuous outcome
Y<-rnorm(nrow(X));obj<-Null_model_glm(Y,Z,family="gaussian")
### Logistic regression for binary outcome
Y<-rbinom(nrow(X),1,0.4);obj<-Null_model_glm(Y,Z,family="binomial")
### Binomial outcome
Y<-rbinom(nrow(X),5,0.4);Y<-cbind(Y,5-Y);obj<-Null_model_glm(Y,Z,family="binomial")
### Poisson outcome
Y<-rpois(nrow(X),5);obj<-Null_model_glm(Y,Z,family="poisson")</pre>
```

SetBasedTests

Set based tests for testing the association between a set of genetic variants and a phenotype

#### **Description**

Calculate the Burden, SKAT, ACAT-V and MORST p-values under one or multiple sets of weights in GLMs or GLMMs.

## Usage

```
SetBasedTests(
   G,
   obj,
   alpha = 1e-06,
   weights.beta = matrix(c(1, 25, 1, 1), nrow = 2),
   weights = NULL,
   tau.type = "approx",
   target_power = 0.5,
   n.points = 50,
   mac.thresh = 10
)
```

## **Arguments**

a numeric matrix or dgCMatrix with each row as a different individual and each column as a separate gene/snp. Each genotype should be coded as 0, 1, 2.

obj an output from Null\_model\_glm or the function glmmkin from the GMMAT pack-

age. See details.

alpha the alpha parameter in MORST. It is suggested to be the significance level.

weights.beta a numeric vector/matrix of parameters for the beta weights for the weighted

kernels. If it is a matrix, each column corresponds to one set of the beta-weights parameters. If you want to use your own weights, please use the "weights"

parameter. It will be ignored if "weights" parameter is not null.

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weights a numeric vector/matrix of weights for the SNPs. If it is a matrix, each column

corresponds to one set of weights. When it is NULL, the beta weight with the

"weights.beta" parameter is used.

tau.type either "minimax" or "approx". See tau\_c for details.

target\_power a value that is used when tau.type == "approx". See tau\_c for details.

n.points number of grid points used when tau.type == "minimax". See tau\_c for details.

mac.thresh a threshold of the minor allele count (MAC) that is used in the ACAT-V test.

SNPs with MAC less than this thrshold will be frist aggregated by the Burden

test in ACAT-V.

#### **Details**

If you want to fit a GLM, please use the Null\_model\_glm function to obtain the null model *obj*. If you have a kinship matrix/GRM and would like to fit a GLMM, please use the function glmmkin from the the GMMAT package. Both dense and sparse kinship/GRM can be used.

The ACAT-V p-value might be slightly different from the result from the ACAT\_V function in the ACAT package. This is because the variant-level p-values are calculated using slighly different methods

While the *alpha* parameter is suggested to be the significance level, practically there is no need to set *alpha* less than 1e-08. In most situations, the MORST p-values would only have negligible difference for values of *alpha* less than 1e-06. A super small *alpha* could slow down the computation and might cause some numerical issue. Therefore, the default value for *alpha* is 1e-06.

#### Value

The p-values of Burden, SKAT, ACAT-V and MORST under under one or multiple choices of weights.

#### Author(s)

Yaowu Liu

## **Examples**

```
library(Matrix)
data(Geno)
G<-Geno[,1:100] # Geno is a dgCMatrix of genotypes
Y<-rnorm(nrow(G)); Z<-matrix(rnorm(nrow(G)*4),ncol=4)
obj<-Null_model_glm(Y,Z,family="gaussian")
SetBasedTests(G,obj)</pre>
```

tau\_c

The choice of tau in MORST

## **Description**

Calculate the parameter tau in MORST based on either the minimax criterion or an approximation of the minimax solution

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## Usage

```
tau_c(eg.values, alpha, tau.type = "approx", target_power = 0.5, n.points = 50)
```

#### **Arguments**

eg.values a numeric vector of non-negative eigenvalues.

alpha the alpha parameter in MORST. It is suggested to be the significance level.

tau.type either "minimax" or "approx". If tau.type == "minimax", tau is calculated based

on the minimax criterion; if tau.type == "approx", tau is an approximation to

the minimax tau. Default value is "approx".

target\_power a value that is used when tau.type == "approx". See details.

n.points number of grid points used when tau.type == "minimax". Should be at least 20

to have reasonable accuracy.

#### **Details**

The approximation method is substantially faster than the minimax method and will be introduced soon.

#### Value

the parameter tau in MORST

#### Author(s)

Yaowu Liu

# References

Liu, Y., Li, Z., and Lin, X. (2020+) A Minimax Optimal Ridge-Type Set Test for Global Hypothesis with Applications in Whole Genome Sequencing Association Studies. *Journal of the American Statistical Association*. Accepted.

## **Examples**

```
tau_c(seq(1,100,1),1e-04)

tau_c(seq(1,100,1),1e-04,tau.type = "minimax")

tau_c(c(2,rep(0.2,20)),0.05,tau.type = "minimax",n.points = 200)
```

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