Package 'RVMMAT'

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Title Retrospective Varying Coefficient Mixed Model Association Test

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Author Gang Xu [aut, cre], Amei Amei [aut], Zuoheng Wang [aut]				
Maintainer Gang Xu <xug3@unlv.nevada.edu></xug3@unlv.nevada.edu>				
Description We propose a GLMM-based retrospective varying coefficient mixed model association test, RVMMAT, to detect genetic variant that affects the developmental patterns of a trait.				
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VignetteBuilder knitr				
R topics documented:				
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glmm_est

NULL model estimation for GMMAT and RGMMAT

Description

This function estimate the parameters and residuals for the NULL generalized linear mixed effect model

Usage

```
glmm_est(
   y.long,
   time,
   y.cov,
   phe.model = phe.model,
   maxiter = 50,
   tol = 10^(-6)
)
```

Arguments

y.long	Long-formatted phenotype vector
time	Time covarites matched with phenotype vector
y.cov	Covariate matrix denoting the covariate variables measured at each time
phe.model	String, the phenotype model, two optional values: 'logistic', 'Gaussian'
maxiter	Numeric, the maximum count for the iterative estimation in when using mixture correlation structure
tol	Numeric, tolerance for the iterative estimation in when using mixture correlation structure

Value

This function returns a list object with model parameters and residuals of the NULL generalized linear mixed effect model

rvmmat_est	NULL model estimation
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Description

This function estimate the parameters and residuals for the NULL model in RVMMAT

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Usage

```
rvmmat_est(
  y.long,
  time,
  y.cov,
  phe.model = phe.model,
  maxiter = 50,
  tol = 10^(-6),
  Hd = 2,
  VCcorrection = FALSE
)
```

Arguments

y.lo	ng	Long-formatted phenotype vector
time		Time covarites matched with phenotype vector
y.co	V	Covariate matrix denoting the covariate variables measured at each time
phe.	model	String, the phenotype model, two optional values: 'logistic', 'Gaussian'
maxi	ter	Numeric, the maximum count for the iterative estimation in when using mixture correlation structure
tol		Numeric, tolerance for the iterative estimation in when using mixture correlation structure
Hd		Numeric, degree of derivative of smooth function which lead to smooth spline of order 2Hd
VCco	rrection	Logical variable, indicating whether apply correction for variance components and natural parameters

Value

This function returns a list object with model parameters and residuals of the NULL model

rvmmat_simu Simulation for RVMMAT test

Description

This function use pre-defined parameters to make the simulation data for the RVMMAT test (including type I and power test)

Usage

```
rvmmat_simu(
    n.sample = 1000,
    n.time = 5,
    par = list(),
    genetic.effect = 0.6,
    time_cov = TRUE,
    snp.count = 100,
    intercept = TRUE,
```

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```
disease.para = list(),
onlypower = FALSE,
phe.model = "logistic",
oversampling = "random"
)
```

Arguments

n. sample Numeric, sample size, number of individuals

n.time Numeric, number of measurements for each individual

par List, the parameters for the phenotype traits, including covariates and individual

specific time dependent random effects

genetic.effect Numeric, genetic effect size (gamma) in genetic effect function

time_cov Logical variable, indicating whether time effect is included in phenotypic traits

snp.count Numeric, number of SNPs

intercept Logical variable, indicating whether intercept is used in phenotypic traits disease.para List, the parameters for disease allele and its effect size for power simulation onlypower Logical variable, indicating whether include disease SNPs in the generated SNPs

phe.model String, the phenotype model, two optional values: 'logistic', 'liability'

oversampling String, the ascertainment scheme, three optional value: 'random', 'baseline',

'sum

Value

A list object is returned to be used as object for RVMMAT test

rvmmat_test Calculate prospective and retrospective P-values for RVMMAT test

Description

This function tests a SNPs for a given SNP set for a given estimated null model.

Usage

```
rvmmat_test(rvmmat.est, G, impute.method = "fixed", GRM = NULL)
```

Arguments

rvmmat.est The output of function "rvmmat_est()"

G The genotype matrix, an m*q matrix where m is the number of subjects and q is

the total number genetic variants.

impute.method choose the iputation method when there is missing genotype. Optional options

are: 'random', 'fixed' or 'bestguess'.

GRM takes m-by-m genetic correlation matrix or kinship matrix.

Value

This function returns a dataframe. The row name is the SNP ID, the first column is the prospective pvalue and the second colum is the restrospective pvalue.

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