

Package: famSKATRC (via r-universe)

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Type Package

Title Family Sequence Kernel Association Test for Rare and Common Variants

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Description FamSKAT-RC is a family-based association kernel test for both rare and common variants. This test is general and several special cases are known as other methods: famSKAT, which only focuses on rare variants in family-based data, SKAT, which focuses on rare variants in population-based data (unrelated individuals), and SKAT-RC, which focuses on both rare and common variants in population-based data. When one applies famSKAT-RC and sets the value of phi to 1, famSKAT-RC becomes famSKAT. When one applies famSKAT-RC and set the value of phi to 1 and the kinship matrix to the identity matrix, famSKAT-RC becomes SKAT. When one applies famSKAT-RC and set the kinship matrix (fullkins) to the identity matrix (and phi is not equal to 1), famSKAT-RC becomes SKAT-RC. We also include a small sample synthetic pedigree to demonstrate the method with. For more details see Saad M and Wijsman EM (2014) <doi:10.1002/gepi.21844>.

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famSKATRC	<i>Family Sequence Kernel Association Test for Rare and Common Variants</i>
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Description

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References

Saad M and Wijsman EM (2014) Combining family- and population-based imputation data for association analysis of rare and common variants in large pedigrees. <https://www.ncbi.nlm.nih.gov/pmc/articles/PMC4190076/>

famSKAT_RC	<i>famSKAT_RC</i>
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Description

FamSKAT-RC is a family-based association kernel test for both rare and common variants.

Usage

```
famSKAT_RC (PHENO, genotypes, id, fullkins, covariates=NULL, sqrtweights_c,
            sqrtweights_r, binomialimpute=FALSE, acc=NULL, maf, phi)
```

Arguments

PHENO	The vector of the phenotype values. The missing values must be represented by NA.
genotypes	The genotype matrix. Its size should be $N * P$, where N is the number of individuals and P is the number of SNPs.
id	The vector of individual IDs to be included in the test. These IDs are present in the kinship matrix.
fullkins	The full kinship matrix that contains all individuals.
covariates	A matrix of possible covariates.
sqrtweights_c	The weight function to be assigned for common variants. An example is: <code>function(maf) ifelse(maf>0, dbeta(maf,0.5,0.5), 0)</code> .
sqrtweights_r	The weight function to be assigned for rare variants. An example is: <code>function(maf) ifelse(maf>0, dbeta(maf,1,25), 0)</code> .
binomialimpute	If TRUE, then impute missing genotypes using a binomial distribution (e.g. <code>rbinom(10, 2, MAF)</code> , if 10 genotypes are missing).
acc	The accuracy used in the Davies approximation. For example, <code>acc= 1e-06</code> .
maf	A MAF threshold used to define rare and common variants (e.g. <code>maf=0.01</code>).
phi	This parameter indicates the contribution portion of rare variants. For example, a value of <code>phi=0.5</code> means that the contribution of rare and common variants to the test is equal and a value of <code>phi=1</code> means that only rare variants contribute to the test. You can vary the phi values as you wish and you can also provide a grid of different values (e.g. <code>phi = c(0,0.2,0.5,0.9)</code>). In this case, four p-values will be obtained, one for each value of phi.

Details

FamSKAT-RC is a family-based association kernel test for both rare and common variants. This test is general and several special cases are known as other methods: famSKAT, which only focuses on rare variants in family-based data, SKAT, which focuses on rare variants in population-based data (unrelated individuals), and SKAT-RC, which focuses on both rare and common variants in

population-based data. When one applies famSKAT-RC and sets the value of phi to 1, famSKAT-RC becomes famSKAT. When one applies famSKAT-RC and set the value of phi to 1 and the kinship matrix to the identity matrix, famSKAT-RC becomes SKAT. When one applies famSKAT-RC and set the kinship matrix (fullkins) to the identity matrix (and phi is not equal to 1), famSKAT-RC becomes SKAT-RC.

Value

The test statistic p-value.

temp_pvalues Some Description

Examples

```
library(kinship2)
sample.ped.geno <- process_data()
KIN = kinship(sample.ped.geno$IID, sample.ped.geno$FA, sample.ped.geno$M0)
IID = sample.ped.geno$IID
wuweights_r <- function(maf) ifelse(maf>0, dbeta(maf,1,25), 0)
wuweights_c <- function(maf) ifelse(maf>0, dbeta(maf,0.5,0.5), 0)
P_VALUES <- famSKAT_RC(PHENO=sample.ped.geno[, "Phenotype"], genotypes=as.matrix(
  sample.ped.geno[, 7:ncol(sample.ped.geno)]), binomialimpute=TRUE,
  id=IID, fullkins=KIN, maf=0.05, sqrtweights_c=wuweights_c,
  sqrtweights_r=wuweights_r, phi = c(0,0.2,0.5,0.9))
print(P_VALUES)
```

process_data

Process Data

Description

Processes the raw data included to produce data identical to the (also included) sample.ped.geno data set. This can be used for processing your own pedigrees with SNP data.

Usage

```
process_data(Data = read.table(system.file("extdata", "data",
  package = "famSKATRC"), header = TRUE))
```

Arguments

Data A string, the path to the location of the data file you are processing, formatted as the included example, which can be loaded with: `read.table(system.file("extdata", "data", package = "famSKATRC"), header = TRUE)` and can be found in your filesystem at: `system.file("extdata", "data", package = "famSKATRC")`

Value

Returns the data frame with completed preprocessing changes for famSKATRC. Mainly reworking IDs so there are not duplicates.

See Also[famSKATRC](#)**Examples**

```

sample.ped.geno <- process_data()
## The function is currently defined as
function(Data = read.table(system.file("extdata", "data",
                                     package = "famSKATRC"), header = TRUE))
{
  Data[, "IID"] = paste(Data[, "FID"] , Data[, "IID"] ,sep=".")
  Data[Data[,"FA"]!=0 , "FA"] = paste(Data[Data[,"FA"]!=0 , "FID"], Data[Data[,"FA"]!=0,
                                     "FA"] ,sep=".")
  Data[Data[,"FA"]!=0 , "MO"] = paste(Data[Data[,"FA"]!=0 , "FID"], Data[Data[,"FA"]!=0,
                                     "MO"] ,sep=".")
  return(Data)
}

```

sample.ped.geno

*Sample Pedigree Genotype Data***Description**

A sample pedigree file with SNP data, already processed by the `process_data()` function. The raw data is also included in the package.

Usage

```
data("sample.ped.geno")
```

Format

A data frame with 20 observations on the following 36 variables.

FID a numeric vector

IID a character vector

FA a character vector

MO a character vector

SEX a numeric vector

Phenotype a numeric vector

rs1 a numeric vector

rs2 a numeric vector

rs3 a numeric vector

rs4 a numeric vector

rs5 a numeric vector

rs6 a numeric vector
rs7 a numeric vector
rs8 a numeric vector
rs9 a numeric vector
rs10 a numeric vector
rs11 a numeric vector
rs12 a numeric vector
rs13 a numeric vector
rs14 a numeric vector
rs15 a numeric vector
rs16 a numeric vector
rs17 a numeric vector
rs18 a numeric vector
rs19 a numeric vector
rs20 a numeric vector
rs21 a numeric vector
rs22 a numeric vector
rs23 a numeric vector
rs24 a numeric vector
rs25 a numeric vector
rs26 a numeric vector
rs27 a numeric vector
rs28 a numeric vector
rs29 a numeric vector
rs30 a numeric vector

Details

A pedigree file with 20 individuals from two families. The first column is a family ID, the second an individual ID, the 3rd gives their father, the 4th their mother, and the 5th their sex. The 6th column gives their phenotype and columns 7 through 36 give their genotype, 30 SNP loci.

Source

This is synthetically generated data.

Examples

```
data(sample.ped.geno)
library(kinship2)
sample.ped.geno$FA[sample.ped.geno$FA == "0"] <- NA
sample.ped.geno$MO[sample.ped.geno$MO == "0"] <- NA
sample.ped.geno$Phenotype[sample.ped.geno$Phenotype >= 0] <- 1
sample.ped.geno$Phenotype[sample.ped.geno$Phenotype < 0] <- 0
ped.list <- pedigree(id = sample.ped.geno$IID, dadid = sample.ped.geno$FA,
                    momid = sample.ped.geno$MO, sex = sample.ped.geno$SEX,
                    famid = sample.ped.geno$FID,
                    affected = sample.ped.geno$Phenotype)

ped1 <- ped.list[1]
ped2 <- ped.list[2]
plot(ped1)
plot(ped2)
```

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