# Package 'PedGFLMM'

March 26, 2019

Type Package

<b>Title</b> Gene-Based Association Testing of Dichotomous Traits with Generalized Linear Mixed Models for Family Data
Version 1.0.0
<b>Description</b> Implements family-based additive generalized linear mixed models (GLMM) and generalized functional linear mixed models (GFLMM) for gene-based association testing of dichotomous traits.
<pre>URL https://github.com/DanielEWeeks/PedGFLMM</pre>
<pre>BugReports https://github.com/DanielEWeeks/PedGFLMM/issues</pre>
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R topics documented:
cov

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cov	COV	

## **Description**

Example covariate data frame, available via data(exampleData).

## Usage

COV

## **Format**

An object of class data. frame with 228 rows and 4 columns.

## **Details**

A data frame containing the covariate information. The first two columns are required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data frame.

## See Also

Ped, geno, exampleData, snpPos

**DOPedGFLMM** 

PedGFLMM\_beta\_smooth\_only call back function

# Description

First, ignore call backs that have less than two polymorphic markers. Second, convert the genotypesraw() patterns of 0x10001, 0x10002 (or 0x20001), 0x20002, 0 from the genotype matrix to the numbers 0, 1, 2, 0 for each marker. (Reverse, the order if allele "1" has the minor allele frequency.) Next, prepend the pedigree and person columns of the family data to this modified genotype matrix. Finally, invoke PedGFLMM with the family data and genotype matrix to compute the PedGFLMM\_beta\_smooth\_only statistics. Save the p-values for each statistic in the *envir\$PedGFLMM\_results* data frame.

## Usage

```
DOPedGFLMM(markers_arg, range_arg, envir = ENV)
```

exampleData 3

#### **Arguments**

markers\_arg a data.frame with the following 5 observations:

locus\_link is the ordinal ranking of this marker among all loci

**locus\_link\_fill** is the position of corresponding genotype data in the *unified\_genotype\_table* 

**MarkerName** is the text name of the marker **chromosome** is the integer chromosome number **position** is the integer base pair position of marker

range\_arg one row of a ranges\_arg. The latter is a data frame of at least three integer

columns. The columns indicate a range: a chromosome number, a start base

pair value, and an end base pair value.

envir 'environment' containing SQLite database and other globals

## Value

None

#### Note

This function computes the PedGFLMM\_beta\_smooth\_only statistics and appends the output to the data frame, <code>envir\$PedGFLMM\_results</code>. It will print out the lines as they are generated if <code>envir\$verbose</code> is TRUE. The data frame <code>envir\$PedGFLMM\_results</code> is initialized by <code>init\_PedGFLMM</code>, and is appended to each time <code>DOPedGFLMM</code> is run.

#### See Also

```
init_PedGFLMM
```

## **Examples**

```
db = system.file("exdata", "seqsimmGFLMM.db", package="PedGFLMM")
ENV = init_PedGFLMM(db)
ENV$verbose = TRUE
Mega2R::applyFnToRanges(DOPedGFLMM, ENV$refRanges[50:60,], ENV$refIndices)
# Not run
Mega2R::applyFnToGenes(DOPedGFLMM, genes_arg = c("CEP104"))
```

exampleData

Example data for the PedGLMM package

# Description

Example data for the PedGLMM package

## Usage

```
data(exampleData)
```

4 geno

## **Format**

Four data frames: Ped, geno, cov, snpPos

#### Value

None

## See Also

```
Ped, geno, cov, snpPos
```

## **Examples**

```
data(exampleData)
dim(Ped)
head(Ped)
dim(geno)
head(geno[,1:10])
dim(cov)
head(cov)
dim(snpPos)
head(snpPos)
```

geno

geno

# Description

Example genotype data frame, available via data(exampleData).

## Usage

geno

# **Format**

An object of class data. frame with 228 rows and 278 columns.

## **Details**

A data frame containing the genotype information. This is a matrix with genotypes for subjects (rows) at each variant position (columns). The first two columns are required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data frame. The genotypes are coded as 0, 1, 2 for autosomal markers (typically a count of the number of the minor alleles).

## See Also

```
Ped, exampleData, cov, snpPos
```

init\_PedGFLMM 5

init_PedGFLMM	load	Mega2	SQLite	database	and	perform	initialization	for
	PedG	FLMM u	sage					

## **Description**

This populates the  ${\bf R}$  data frames from the specified  ${\bf Mega2}$  SQLite database.

# Usage

```
init_PedGFLMM(db = NULL, verbose = FALSE, traitname = "default")
```

## **Arguments**

	db	specifies the path of a Mega2 SQLite database containing study data.
--	----	--

verbose TRUE indicates that diagnostic printouts should be enabled. This value is saved

in the returned environment.

traitname Name of the affection status trait to use to set the case/control values; by default,

"default"

## Value

"environment" containing data frames from an SQLite database and some computed values.

## Note

*init\_PedGFLMM* sets up the schaidPed and pedPer data frames that are used later in the *DOPedGFLMM* calculation. In addition, it initializes a matrix to aid in translating a genotype allele matrix to a genotype count matrix.

It also initializes the results data frame *envir\$PedGFLMM\_results* to zero rows.

# See Also

```
DOPedGFLMM, Mega2PedGFLMM
```

```
db = system.file("exdata", "seqsimmGFLMM.db", package="PedGFLMM")
ENV = init_PedGFLMM(db, traitname = "default")
ls(ENV)
```

6 Mega2PedGFLMM

Mega2PedGFLMM	Execute the PedGFLMM_beta_smooth_only function on a transcript
	ranges

# Description

This example function illustates how to use functions from the Mega2R R package to iterate over defined gene ranges, computing the PedGFLMM\_beta\_smooth\_only statistics for each gene that contains more than two polymorphic markers.

Execute the PedGFLMM\_beta\_smooth\_only function on the first gs default gene transcript ranges (gs = 1:100). Update the envir\$PedGFLMM\_results data frame with the results.

# Usage

```
Mega2PedGFLMM(gs = 1:100, genes = NULL, envir = ENV)
```

# **Arguments**

gs	a subrange of the default transcript ranges over which to calculate the $DOPe-dGFLMM$ function.
genes	a list of genes over which to calculate the <i>DOPedGFLMM</i> function. The value, "*", means use all the transcripts in the selected Bioconductor database. If genes is NULL, the gs range of the internal <i>refRanges</i> will be used.
envir	'environment' containing SQLite database and other globals

## Value

None the data frame with the PedGFLMM\_beta\_smooth\_only results is stored in the environment and named *PedGFLMM\_results*, viz. envir\$PedGFLMM\_results

## See Also

```
init_PedGFLMM
```

```
db = system.file("exdata", "seqsimmGFLMM.db", package="PedGFLMM")
ENV = init_PedGFLMM(db)
ENV$verbose = TRUE
Mega2PedGFLMM(gs = 50:60)
```

Ped 7

Ped Ped

## **Description**

Example pedigree data frame, available via data(exampleData)

## Usage

Ped

#### **Format**

An object of class data. frame with 228 rows and 7 columns.

#### **Details**

A data frame containing the pedigree information with the following columns:

**ID** Person ID

ped pedigree ID, character or numeric allowed.

person person ID, a unique ID within each pedigree, numeric or character allowed.

father father ID, NA if no father.

mother mother ID, NA if no mother.

sex sex, coded as 1 for male, 2 for female.

**trait** trait phenotype, either case-control status coded as 1 for affected and 0 for unaffected. Subjects with missing (NA) will be removed from the analysis.

## See Also

exampleData, geno, cov, snpPos

PedGFLMM

PedGFLMM package

# Description

This package implements family-based additive generalized linear mixed models (GLMM) and generalized functional linear mixed models (GFLMM) for gene-based association testing of dichotomous traits (Jiang et al, 2019).

# Author(s)

Yingda Jiang, Chi-Yang Chiu, Daniel E. Weeks, Ruzong Fan

PedGFLMM\_beta\_smooth\_only

PedGFLMM\_beta\_smooth\_only

## **Description**

Computes the PedGFLMM statistics under the beta smooth only model.

## Usage

```
PedGFLMM_beta_smooth_only(ped, geno, covariate = NULL, pos, order, beta_basis,
  base = "bspline", Wald = FALSE)
```

#### **Arguments**

ped A data frame containing the pedigree information with the following columns:

**ID** Person ID

**ped** pedigree ID, character or numeric allowed.

person person ID, a unique ID within each pedigree, numeric or character al-

lowed.

**father** father ID, NA if no father.

**mother** mother ID, NA if no mother.

sex sex, coded as 1 for male, 2 for female.

**trait** trait phenotype, case-control status coded as 1 for affected and 0 for unaffected. Subjects with missing (NA) will be removed from the analysis.

geno A data frame containing the genotype information. This is a matrix with geno-

types for subjects (rows) at each variant position (columns). The first two columns are required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data.frame. The genotypes are coded as 0, 1, 2 for subjects to the property of the purple of the principal leads.

autosomal markers (typically a count of the number of the minor alleles).

covariate A data frame containing the covariate information. The first two columns are

required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data frame. This is optional and the default "covariate

= NULL" is for the case when the covariate matrix is not provided.

pos Position of the markers in base pairs.

order The order used to generate the B-spline basis.

beta\_basis The number of basis functions used to estimate the genetic effect function.

base Can be either 'bspline' or 'fspline'.

Wald If Wald is set to true, return the Wald p-value in addition to the LRT p-value

(Default: Wald = FALSE).

# Value

A list containing the following components:

LRT The p-value based on a likelihood ratio test

Wald The p-value based on a Wald test, returned if 'Wald' is TRUE

#### References

Chiu CY, Yuan F, Zhang BS, Yuan A, Li X, Fang HB, Lange K, Weeks DE, Wilson AF, Bailey-Wilson JE, Lakhal-Chaieb ML, Cook RJ, McMahon FJ, Amos CI, Xiong MM, and Fan RZ (2019) Pedigree-based linear mixed models for association analysis of quantitative traits with next-generation sequencing data. Genetic Epidemiology 43(2):189-206.

Fan RZ, Wang YF, Mills JL, Wilson AF, Bailey-Wilson JE, and Xiong MM (2013) Functional linear models for association analysis of quantitative traits. Genetic Epidemiology 37 (7):726-742.

Fan RZ, Wang YF, Mills JL, Carter TC, Lobach I, Wilson AF, Bailey-Wilson JE, Weeks DE, and Xiong MM (2014) Generalized functional linear models for case-control association studies. Genetic Epidemiology 38 (7):622-637.

Jiang YD, Chiu CY, Yan Q, Chen W, Gorin MB, Conley YP, Lakhal-Chaieb ML, Cook RJ, Amos CI, Wilson AF, Bailey-Wilson JE, McMahon FJ, Vazquez AI, Yuan A, Zhong XG, Xiong MM, Weeks DE, and Fan RZ (2019) Gene-based association testing of dichotomous traits with generalized linear mixed models for family data.

Schaid DJ, McDonnell SK, Sinnwell JP, and Thibodeau SN (2013) Multiple genetic variant association testing by collapsing and kernel methods with pedigree or population structured data. Genetic Epidemiology 37:409-418.

#### See Also

PedGLMM\_additive\_effect\_model, PedGFLMM\_fixed\_model, exampleData

```
data(exampleData)
betabasis_Bsp = 10
betabasis_Fsp = 11
order = 4
bsmooth_bsp=PedGFLMM_beta_smooth_only(ped = Ped, geno = as.matrix(geno),
  pos = snpPos$pos, order = order, beta_basis=betabasis_Bsp, covariate = as.matrix(cov),
   base = "bspline")
bsmooth_bsp
bsmooth_fsp=PedGFLMM_beta_smooth_only(ped = Ped, geno = as.matrix(geno),
  pos = snpPos$pos, order = order, beta_basis=betabasis_Fsp, covariate = as.matrix(cov),
   base = "fspline")
bsmooth_fsp
bsmooth_bsp_no_cov=PedGFLMM_beta_smooth_only(ped = Ped, geno = as.matrix(geno),
   pos = snpPos$pos, order = order, beta_basis=betabasis_Bsp, covariate = NULL,
   base = "bspline")
bsmooth_bsp_no_cov
bsmooth_fsp_no_cov=PedGFLMM_beta_smooth_only(ped = Ped, geno = as.matrix(geno),
   pos = snpPos$pos, order = order, beta_basis=betabasis_Fsp, covariate = NULL,
   base = "fspline")
bsmooth_fsp_no_cov
```

#### **Description**

Computes the PedGFLMM statistics under a fixed model.

#### **Usage**

```
PedGFLMM_fixed_model(ped, geno, covariate = NULL, pos, order, beta_basis,
  geno_basis, base = "bspline", Wald = FALSE)
```

## **Arguments**

ped A data frame containing the pedigree information with the following columns:

**ID** Person ID

**ped** pedigree ID, character or numeric allowed.

person person ID, a unique ID within each pedigree, numeric or character al-

lowed.

father father ID, NA if no father.

mother mother ID, NA if no mother.

sex sex, coded as 1 for male, 2 for female.

**trait** trait phenotype, case-control status coded as 1 for affected and 0 for unaffected. Subjects with missing (NA) will be removed from the analysis.

geno A data frame containing the genotype information. This is a matrix with geno-

types for subjects (rows) at each variant position (columns). The first two columns are required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data.frame. The genotypes are coded as 0, 1, 2 for

autosomal markers (typically a count of the number of the minor alleles).

covariate A data frame containing the covariate information. The first two columns are

required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data frame. This is optional and the default "covariate

= NULL" is for the case when the covariate matrix is not provided.

pos Position of the markers in base pairs.

order The order used to generate the B-spline basis.

beta\_basis The number of basis functions used to estimate the genetic effect function.

geno\_basis The number of basis functions used to estimate the genetic variant functions.

base Can be either 'bspline' or 'fspline'.

Wald If Wald is set to true, return the Wald p-value in addition to the LRT p-value

(Default: Wald = FALSE).

### Value

A list containing the following components:

LRT The p-value based on a likelihood ratio test

Wald The p-value based on a Wald test, returned if 'Wald' is TRUE

#### References

Chiu CY, Yuan F, Zhang BS, Yuan A, Li X, Fang HB, Lange K, Weeks DE, Wilson AF, Bailey-Wilson JE, Lakhal-Chaieb ML, Cook RJ, McMahon FJ, Amos CI, Xiong MM, and Fan RZ (2019) Pedigree-based linear mixed models for association analysis of quantitative traits with next-generation sequencing data. Genetic Epidemiology 43(2):189-206.

Fan RZ, Wang YF, Mills JL, Wilson AF, Bailey-Wilson JE, and Xiong MM (2013) Functional linear models for association analysis of quantitative traits. Genetic Epidemiology 37 (7):726-742.

Fan RZ, Wang YF, Mills JL, Carter TC, Lobach I, Wilson AF, Bailey-Wilson JE, Weeks DE, and Xiong MM (2014) Generalized functional linear models for case-control association studies. Genetic Epidemiology 38 (7):622-637.

Jiang YD, Chiu CY, Yan Q, Chen W, Gorin MB, Conley YP, Lakhal-Chaieb ML, Cook RJ, Amos CI, Wilson AF, Bailey-Wilson JE, McMahon FJ, Vazquez AI, Yuan A, Zhong XG, Xiong MM, Weeks DE, and Fan RZ (2019) Gene-based association testing of dichotomous traits with generalized linear mixed models for family data.

Schaid DJ, McDonnell SK, Sinnwell JP, and Thibodeau SN (2013) Multiple genetic variant association testing by collapsing and kernel methods with pedigree or population structured data. Genetic Epidemiology 37:409-418.

#### See Also

PedGFLMM\_beta\_smooth\_only, PedGLMM\_additive\_effect\_model, exampleData

```
data(exampleData)
betabasis_Bsp = 10
genobasis_Bsp = 10
betabasis_Fsp = 11
genobasis_Fsp = 11
order = 4
fixed_bsp=PedGFLMM_fixed_model(ped = Ped, geno = as.matrix(geno), pos = snpPos$pos,
    order = order, beta_basis=betabasis_Bsp, geno_basis = genobasis_Bsp,
    covariate = as.matrix(cov), base = "bspline")
fixed_bsp
fixed_fsp=PedGFLMM_fixed_model(ped = Ped, geno = as.matrix(geno), pos = snpPos$pos,
    order = order, beta_basis=betabasis_Fsp, geno_basis = genobasis_Fsp,
    covariate = as.matrix(cov), base = "fspline")
fixed_fsp
fixed_bsp_no_cov=PedGFLMM_fixed_model(ped = Ped, geno = as.matrix(geno), pos = snpPos$pos,
    order = order, beta_basis=betabasis_Bsp, geno_basis = genobasis_Bsp,
    covariate = NULL, base = "bspline")
fixed_bsp_no_cov
fixed_fsp_no_cov=PedGFLMM_fixed_model(ped = Ped, geno = as.matrix(geno), pos = snpPos$pos,
    order = order, beta_basis=betabasis_Fsp, geno_basis = genobasis_Fsp,
    covariate = NULL, base = "fspline")
fixed_fsp_no_cov
```

PedGLMM\_additive\_effect\_model

PedGLMM\_additive\_effect\_model

## **Description**

Computes the PedGFLMM statistics under an additive effect model

## Usage

PedGLMM\_additive\_effect\_model(ped, geno, covariate = NULL, Wald = FALSE)

## **Arguments**

ped A data frame containing the pedigree information with the following columns:

ID Person ID

**ped** pedigree ID, character or numeric allowed.

person person ID, a unique ID within each pedigree, numeric or character al-

lowed.

father father ID. NA if no father.

mother mother ID, NA if no mother.

sex sex, coded as 1 for male, 2 for female.

trait trait phenotype, case-control status coded as 1 for affected and 0 for unaf-

fected. Subjects with missing (NA) will be removed from the analysis.

geno A data frame containing the genotype information. This is a matrix with geno-

types for subjects (rows) at each variant position (columns). The first two columns are required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data.frame. The genotypes are coded as 0, 1, 2 for

autosomal markers (typically a count of the number of the minor alleles).

covariate A data frame containing the covariate information. The first two columns are

required to be named "ped" and "person", which are used to match subjects to their data in the pedigree data frame. This is optional and the default "covariate

= NULL" is for the case when the covariate matrix is not provided.

Wald If Wald is set to true, return the Wald p-value in addition to the LRT p-value

(Default: Wald = FALSE).

#### Value

A list containing the following components:

LRT The p-value based on a likelihood ratio test

Wald The p-value based on a Wald test, returned if 'Wald' is TRUE

### References

Chiu CY, Yuan F, Zhang BS, Yuan A, Li X, Fang HB, Lange K, Weeks DE, Wilson AF, Bailey-Wilson JE, Lakhal-Chaieb ML, Cook RJ, McMahon FJ, Amos CI, Xiong MM, and Fan RZ (2019) Pedigree-based linear mixed models for association analysis of quantitative traits with next-generation sequencing data. Genetic Epidemiology 43(2):189-206.

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Fan RZ, Wang YF, Mills JL, Wilson AF, Bailey-Wilson JE, and Xiong MM (2013) Functional linear models for association analysis of quantitative traits. Genetic Epidemiology 37 (7):726-742.

Fan RZ, Wang YF, Mills JL, Carter TC, Lobach I, Wilson AF, Bailey-Wilson JE, Weeks DE, and Xiong MM (2014) Generalized functional linear models for case-control association studies. Genetic Epidemiology 38 (7):622-637.

Jiang YD, Chiu CY, Yan Q, Chen W, Gorin MB, Conley YP, Lakhal-Chaieb ML, Cook RJ, Amos CI, Wilson AF, Bailey-Wilson JE, McMahon FJ, Vazquez AI, Yuan A, Zhong XG, Xiong MM, Weeks DE, and Fan RZ (2019) Gene-based association testing of dichotomous traits with generalized linear mixed models for family data.

Schaid DJ, McDonnell SK, Sinnwell JP, and Thibodeau SN (2013) Multiple genetic variant association testing by collapsing and kernel methods with pedigree or population structured data. Genetic Epidemiology 37:409-418.

#### See Also

```
PedGFLMM_beta_smooth_only, PedGFLMM_fixed_model, exampleData
```

#### **Examples**

snpPos

snpPos

## Description

Example marker position data frame, available via data(exampleData).

# Usage

snpPos

## **Format**

An object of class data. frame with 276 rows and 3 columns.

## **Details**

This data frame provides marker positions for each SNP. The first column, chr, contains the chromosome number, the second column, snp, contains the SNP name, and the third column, pos, contains the position of the SNP in base pairs.

## See Also

```
Ped, geno, cov, exampleData
```

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