Package ‘MACAU2’

April 8, 2017

Type Package

Title MACAU 2.0: Efficient Mixed Model Analysis of Count Data

Version 1.10

Date 2017-03-31

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Description MACAU2 is an R package designed for generalized linear mixed model inference in genomic sequencing studies.

Depends R (>= 3.0.1), spam, gtools, Matrix, INLA

License GPL-3

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Imports Rcpp (>= 0.11.6), foreach, doParallel, parallel

LinkingTo Rcpp, RcppArmadillo, foreach, doParallel, parallel

NeedsCompilation yes

R topics documented:

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MACAU2-package  Efficient Mixed Model Analysis of Count Data in Large-Scale Genomic Sequencing Studies

Description

MACAU2 is an R package for efficient differential analysis of large-scale RNA sequencing (RNAseq) data and Bisulfite sequencing (BSseq) data in the presence of individual relatedness and population structure. MACAU2 first fits a GLMM with adjusted covariates, predictor of interest and random effects to account for population structure and individual relatedness, and then performs wald tests for each gene in RNAseq or site for BSseq.

Details

Package: MACAU2
Type: Package
Version: 1.10
Date: 2017-03-31
License: GPL-3

Author(s)

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References


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ExampleBSseq

BSseq example dataset

Description

A simulated example dataset of BSseq for MACAU2.

Format

Contains the following objects:

- `mcount`: a matrix containing the read counts for 100 sites.
- `predictor`: a vector (size=200) of observations on a continuous variable.
- `totalcount`: a matrix containing the total read counts for 100 sites.

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ExampleRNAseq

RNAseq example dataset

Description

A simulated example dataset of RNAseq for MACAU2.

Format

Contains the following objects:

- `count`: a matrix containing the read counts for 100 genes.
- `predictor`: a vector (size=200) of observations on a continuous variable.
- `totalcount`: a vector (size=200) with one read depth number for each individual.
macau2

*Fit generalized linear mixed model with known kinship matrices through penalized-quasi likelihood.*

**Description**

Fit a generalized linear mixed model with a random intercept. The covariance matrix of the random intercept is proportional to a known kinship matrix.

**Usage**

```r
macau2(RawCountDataSet, Phenotypes, Covariates=NULL, RelatednessMatrix=NULL, LibSize=NULL, fit.model="PMM", fit.method = "AI.REML", fit.maxiter=500, fit.tol=1e-5, numCore=1, filtering=TRUE, verbose=FALSE, ...)
```

**Arguments**

- **RawCountDataSet**
  a data frame containing the read count.
- **Phenotypes**
  a vector containing the predictor of interest.
- **Covariates**
  a data frame containing the covariates subject to adjustment (Default = NULL).
- **RelatednessMatrix**
  a known relationship matrix (e.g. kinship matrix in genetic studies). When supplied with a matrix, this matrix should be a positive semi-definite matrix with dimensions equal to the sample size in count data, and the order of subjects in this matrix should also match the order of subjects in count data. Currently there is no ID checking feature implemented, and it is the user’s responsibility to match the orders.
- **LibSize**
  a data frame containing the total read count. For posission mixed model, it will be calculated automatically if users do not provide. For binomial mixed model, it is required.
- **fit.model**
  a description of the error distribution and link function to be used in the model. Either "PMM" for posission model, or "BMM" for binomial model (default="PMM").
- **fit.method**
  method of fitting the generalized linear mixed model, currently only "REML" version is available.
- **fit.maxiter**
  a positive integer specifying the maximum number of iterations when fitting the generalized linear mixed model (default = 500).
- **fit.tol**
  a positive number specifying tolerance, the difference threshold for parameter estimates below which iterations should be stopped (default = 1e-5).
- **numCore**
  a positive integer specifying the number of cores for parallel computing (default = 1).
- **filtering**
  a logical switch for RNAseq data. By default, for each gene, at least two individuals should have read counts greater than 5. Otherwise, the gene is filtered (default = TRUE).
verbose a logical switch for printing detailed information (parameter estimates in each iteration) for testing and debugging purpose (default = FALSE).

... additional arguments that could be passed to glm.

Details

Generalized linear mixed models (GLMM) are fitted using the penalized quasi-likelihood (PQL) method proposed by Breslow and Clayton (1993). Statistical inference in GLMM is notoriously difficult because of an intractable high-dimensional integral in the likelihood (Chen, 2016 and Lea, 2015), and by default we use the Average Information REML algorithm (Gilmour, Thompson and Cullis, 1995; Yang et al., 2011) to fit the model. An eigen-decomposition is performed in each outer iteration and the estimate of the variance component parameter $\tau$ is obtained by maximizing the profiled log restricted likelihood. When the Average Information REML algorithm fails to converge, a warning message is given and the algorithm is default to INLA approaches (Rue, 2009).

Value

- **numIDV**: number of individuals with data being analyzed
- **beta**: the fixed effect parameter estimate for the predictor of interest.
- **se_beta**: the standard deviation of fixed effect.
- **pvalue**: P value for the fixed effect, based on the wald test.
- **h2**: heritability of the transformed rate.
- **sigma2**: total variance component.
- **converged**: a logical indicator for convergence.

Author(s)

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References


Examples

```r
data(ExampleBSseq)
attach(ExampleBSseq)
model_DNA=macau2(RawCountDataSet=mcount, Phenotypes=predictor, RelatednessMatrix=relatednessmatrix, LibSize=totalcount, fit.model="BMM",numCore=10)
head(model_DNA)
detach(ExampleBSseq)

data(ExampleRNAseq)
attach(ExampleRNAseq)
model_RNA=macau2(RawCountDataSet=count, Phenotypes=predictor, RelatednessMatrix=relatednessmatrix, fit.model="PMM",numCore=10)
head(model_RNA)
detach(ExampleRNAseq)
```
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