

Package ‘bmass’

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Type Package

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Title Bayesian Multivariate Analysis of Summary Statistics

Description Multivariate tool for analyzing genome-wide association study results in the form of univariate summary statistics. The goal of 'bmass' is to comprehensively test all possible multivariate models given the phenotypes and datasets provided. Multivariate models are determined by assigning each phenotype to being either Unassociated (U), Directly associated (D) or Indirectly associated (I) with the genetic variant of interest. Test results for each model are presented in the form of Bayes factors, thereby allowing direct comparisons between models. The underlying framework implemented here is based on the modeling developed in ``A Unified Framework for Association Analysis with Multiple Related Phenotypes'', M. Stephens (2013) <[doi:10.1371/journal.pone.0065245](https://doi.org/10.1371/journal.pone.0065245)>.

License GPL (>= 3)

URL <https://github.com/mturchin20/bmass>

BugReports <https://github.com/mturchin20/bmass/issues>

Depends R (>= 3.3.0)

Imports utils, stats

Suggests testthat, knitr, rmarkdown

LazyData true

NeedsCompilation no

RoxygenNote 6.1.1

VignetteBuilder knitr

Author Michael Turchin [aut, cre],
Matthew Stephens [aut],
Peter Carbonetto [ctb]

Maintainer Michael Turchin <mturchin20@uchicago.edu>
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bmass	<i>Bayesian multivariate analysis of summary statistics (bmass)</i>
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Description

Run bmass on a set of phenotypes that each have univariate GWAS statistics on the same set of SNPs

Usage

```
bmass(DataSources, GWASsnps = NULL,  
      SNPMarginalUnivariateThreshold = 1e-06,  
      SNPMarginalMultivariateThreshold = 1e-06, GWASThreshFlag = TRUE,  
      GWASThreshValue = 5e-08, NminThreshold = 0,  
      PrintMergedData = FALSE, PrintProgress = FALSE, ...)
```

Arguments

- | | |
|--------------------------------|---|
| DataSources | A string indicating the variable names of the input datafiles and phenotypes. No default value. |
| GWASsnps | A data.table containing rows of SNPs that were univariate genome-wide significant in the phenotypes being used for analysis; GWASsnps input file should have two columns, one for chromosome and another for basepair position (with column headers of Chr and BP). No default value. |
| SNPMarginalUnivariateThreshold | A numerical value indicating the univariate p-value threshold to use when collecting marginally significant SNPs for final bmass analysis. Default is 1e-6. |

SNPMarginalMultivariateThreshold	A numerical value indicating the basic multivariate p-value threshold to use when collecting marginally significant SNPs for final bmass analysis. Default is $1e-6$.
GWASThreshFlag	A logical TRUE/FALSE flag that indicates whether to threshold input GWASsnps list by a univariate GWAS p-value or not (eg the input GWASsnps list contains variants that are significant from discovery + replication data, but the input summary statistics are just from the discovery cohort). Default is TRUE.
GWASThreshValue	A numerical value indicating the univariate p-value threshold to use in conjunction with the GWASThreshFlag. Default is $5e-8$.
NminThreshold	A numerical value that indicates a sample size threshold to use where SNPs below which are removed. Default is 0.
PrintMergedData	A logical TRUE/FALSE flag that indicates whether the intermediary 'merged datafile' should be included in the final bmass output; this file combines all the phenotypes for every SNP provided just prior to thresholding for marginally significant SNPs. Default is FALSE.
PrintProgress	A logical TRUE/FALSE flag that indicates whether progress statements should be printed to stderr during the course of running bmass or not. Default is FALSE.
...	Additional optional arguments.

Value

A list containing model, SNP, and posterior information for both the previously significant univariate SNPs (PreviousSNPs) and the newly significant multivariate SNPs (NewSNPs). For a full breakdown of the bmass output list structure, please see the associated vignettes.

Other Examples

```
bmass(c("HDL", "LDL", "TG", "TC"), GWASsnps, NminThreshold = 50000) bmass(c("HDL", "LDL", "TG", "TC"),
GWASsnps, GWASThreshValue = 1e-8, NminThreshold = 50000, PrintProgress = TRUE) bmass(c("HDL",
"LDL", "TG", "TC"), GWASsnps, GWASThreshFlag = FALSE, SNPMarginalUnivariateThreshold
= 1e-4, SNPMarginalMultivariateThreshold = 1e-4, PrintMergedData = TRUE) bmassOutput
<- bmass(c("HDL", "LDL", "TG", "TC"), GWASsnps, NminThreshold = 50000)
```

Examples

```
Phenotypes <- c("bmass_SimulatedData1", "bmass_SimulatedData2")
bmassOutput <- bmass(Phenotypes, bmass_SimulatedSigSNPs)
summary(bmassOutput)
bmassOutput$NewSNPs$SNPs
```

bmass_SimulatedData1 *bmass Simulated Dataset 1*

Description

A manually created sample dataset for use in Roxygen2 documents and vignettes.

Format

A data frame with 11 rows and 9 variables:

Chr chromosome

BP basepair position

Marker rsID# or other identifier

MAF Minor Allele Frequency

A1 reference allele

A2 alternative allele

Direction direction of association effect size, + or -

pValue p-Value of GWAS association

N sample size

Source

Manually created

bmass_SimulatedData2 *bmass Simulated Dataset 2*

Description

A manually created sample dataset for use in Roxygen2 documents and vignettes.

Format

A data frame with 11 rows and 9 variables:

Chr chromosome

BP basepair position

Marker rsID# or other identifier

MAF Minor Allele Frequency

A1 reference allele

A2 alternative allele

Direction direction of association effect size, + or -

pValue p-Value of GWAS association

N sample size

Source

Manually created

bmass_SimulatedSigSNPs

bmass Simulated GWAS SNPs

Description

A manually created list of GWAS significant SNPs to be used in conjunction with 'bmass_SimulatedData1' and 'bmass_SimulatedData2'.

Format

A data frame with 2 rows and 2 variables:

Chr chromosome

BP basepair position

Source

Manually created

bmass_TestData1

bmass Test Dataset 1

Description

A manually created sample dataset for use in unit tests.

Format

A data frame with 11 rows and 9 variables:

Chr chromosome

BP basepair position

Marker rsID# or other identifier

MAF Minor Allele Frequency

A1 reference allele

A2 alternative allele

Direction direction of association effect size, + or -

pValue p-Value of GWAS association

N sample size

Source

Manually created

bmass_TestData2	<i>bmass Test Dataset 2</i>
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Description

A manually created sample dataset for use in unit tests.

Format

A data frame with 11 rows and 9 variables:

Chr chromosome

BP basepair position

Marker rsID# or other identifier

MAF Minor Allele Frequency

A1 reference allele

A2 alternative allele

Direction direction of association effect size, + or -

pValue p-Value of GWAS association

N sample size

Source

Manually created

bmass_TestSigSNPs	<i>bmass Test GWAS SNPs</i>
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Description

A manually created list of GWAS significant SNPs to be used in conjunction with 'bmass_TestData1' and 'bmass_TestData2'.

Format

A data frame with 2 rows and 2 variables:

Chr chromosome

BP basepair position

Source

Manually created

GetMarginalPosteriors *Get Marginal {U,D,I} Posteriors*

Description

Get marginal posteriors for how much every individual phenotype belongs to categories {U,D,I} across each SNP

Usage

```
GetMarginalPosteriors(DataSources, ListSNPs, Models, LogFile)
```

Arguments

DataSources	A string indicating the variable names of the input datafiles and phenotypes.
ListSNPs	A list produced from running bmass containing the SNPs of interest to get marginal posteriors for.
Models	A matrix describing the models being explored (default output from running bmass).
LogFile	A matrix of string outputs for function logging purposes (default output from running bmass).

Value

A list containing three matrices of SNPs x Phenotypes marginal posteriors for each category {U,D,I}; this list is appended to the input ListSNPs as a new object, Marginals (the full returned object is a list containing the input ListSNPs and the input LogFile).

Examples

```
Phenotypes <- c("bmass_SimulatedData1", "bmass_SimulatedData2")
bmassOutput <- bmass(Phenotypes, bmass_SimulatedSigSNPs)
bmassOutput[c("PreviousSNPs", "LogFile")] <-
  GetMarginalPosteriors(Phenotypes, bmassOutput$PreviousSNPs,
    bmassOutput$Models, bmassOutput$LogFile)
bmassOutput$PreviousSNPs$Marginals
```

GetModelPriorMatrix *Get Model Prior Matrix*

Description

Creates a matrix containing the model descriptions and their associated priors.

Usage

```
GetModelPriorMatrix(DataSources, Models, ModelPriors, LogFile,
  SigmaAlphas = c(0.005, 0.0075, 0.01, 0.015, 0.02, 0.03, 0.04, 0.05,
    0.06, 0.07, 0.08, 0.09, 0.1, 0.15))
```

Arguments

DataSources	A string indicating the variable names of the input datafiles and phenotypes.
Models	A matrix describing the models being explored (default output from running bmass).
ModelPriors	A vector containing the priors on each model across each tranche of sigma alpha (default output from running bmass ; length is number of models times number of sigma alphas).
LogFile	A matrix of string outputs for function logging purposes (default output from running bmass).
SigmaAlphas	A vector containing the different values traversed for this 'effect size controlling' hyperparameter (see "Prior on Sigma_Alpha" in Stephens 2013 PLoS ONE, https://doi.org/10.1371/journal.pone.0065245).

Value

A matrix containing the original description of each model sort by prior, each model's trained prior, the cummulative prior distribution, and the model's original order position.

Examples

```
Phenotypes <- c("bmass_SimulatedData1", "bmass_SimulatedData2")
bmassOutput <- bmass(Phenotypes,bmass_SimulatedSigSNPs)
bmassOutput[c("ModelPriorMatrix", "LogFile")] <-
  GetModelPriorMatrix(Phenotypes, bmassOutput$Models,
    bmassOutput$ModelPriors, bmassOutput$LogFile)
head(bmassOutput$ModelPriorMatrix)
```

GetTopModelsPerSNPViaPosteriors
Get Top Multivariate Models

Description

Get a summary of the top models per SNP across all multivariate $\{\mathbf{U}, \mathbf{D}, \mathbf{I}\}$ combinations based on posterior probabilities.

Usage

```
GetTopModelsPerSNPViaPosteriors(DataSources, ListSNPs, ModelPriorMatrix,
                                LogFile)
```

Arguments

DataSources	A string indicating the variable names of the input datafiles and phenotypes.
ListSNPs	A list produced from running bmass containing the SNPs of interest to get marginal posteriors for.
ModelPriorMatrix	A matrix detailing the models being explored and their associated priors (obtained by running GetModelPriorMatrix)
LogFile	A matrix of string outputs for function logging purposes (default output from running bmass).

Value

A matrix containing each model that was a SNP's top model at least once, along with related information; this matrix is appended to the input ListSNPs as a new object, TopModels (the full returned object is a list containing the input ListSNPs and the input LogFile).

Examples

```
Phenotypes <- c("bmass_SimulatedData1", "bmass_SimulatedData2")
bmassOutput <- bmass(Phenotypes, bmass_SimulatedSigSNPs)
bmassOutput[c("ModelPriorMatrix", "LogFile")] <-
  GetModelPriorMatrix(Phenotypes, bmassOutput$Models,
                      bmassOutput$ModelPriors, bmassOutput$LogFile)
bmassOutput[c("PreviousSNPs", "LogFile")] <-
  GetTopModelsPerSNPViaPosteriors(Phenotypes,
                                  bmassOutput$PreviousSNPs, bmassOutput$ModelPriorMatrix, bmassOutput$LogFile)
head(bmassOutput$PreviousSNPs$TopModels)
```

GlobalLipids2013.GWASsnps

GlobalLipids2013 GWAS SNPs

Description

A list of the univariate GWAS significant SNPs from the GlobalLipids2013 dataset to be used in the second introductory bmass vignette.

Format

A data frame with 157 rows and 2 variables:

Chr chromosome

BP basepair position

Source

Supplementary Tables 2 and 3 from <https://doi.org/10.1038/ng.2797>.

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