Package ‘hierGWAS’

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Title Asessing statistical significance in predictive GWA studies

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Author Laura Buzdugan

Maintainer Laura Buzdugan <buzdugan@stat.math.ethz.ch>

Description Testing individual SNPs, as well as arbitrarily large groups of SNPs in GWA studies, using a joint model of all SNPs. The method controls the FWER, and provides an automatic, data-driven refinement of the SNP clusters to smaller groups or single markers.

Depends R (>= 3.2.0)

License GPL-3

LazyData true

Imports fastcluster, glmnet, fmsb

Suggests BiocGenerics, RUnit, MASS

biocViews SNP, LinkageDisequilibrium, Clustering

 'iterative.DFS.R’ ’test.hierarchy.R’ ’return.r2.R’
 'compute.r2.R’

NeedsCompilation no

R topics documented:

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cluster.snp

Hierarchical Clustering of SNP Data

Description

Clusters SNPs hierachically.

Usage

cluster.snp(x = NULL, d = NULL, method = "average", SNP_index = NULL)

Arguments

x
  The SNP data matrix of size \texttt{nobs} \times \texttt{nvar}. Default value is \texttt{NULL}

d
  \texttt{NULL} or a dissimilarity matrix. See the 'Details' section.

method
  The agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC). See \texttt{hclust} for details.

SNP_index
  \texttt{NULL} or the index vector of SNPs to be clustered. See the 'Details' section.

Details

The SNPs are clustered using \texttt{hclust}, which performs a hierarchical cluster analysis using a set of dissimilarities for the \texttt{nvar} objects being clustered. There are 3 possible scenarios.

If \(d = \text{NULL}\), \(x\) is used to compute the dissimilarity matrix. The dissimilarity measure between two SNPs is \(1 - \text{LD}\) (Linkage Disequilibrium), where \(\text{LD}\) is defined as the square of the Pearson correlation coefficient. If \(\text{SNP\textunderscore index} = \text{NULL}\), all \texttt{nvar} SNPs will be clustered; otherwise only the SNPs with indices specified by \(\text{SNP\textunderscore index}\) will be considered.

If the user wishes to use a different dissimilarity measure, \(d\) needs to be provided. \(d\) must be either a square matrix of size \(\texttt{nvar} \times \texttt{nvar}\), or an object of class \texttt{dist}. If \(d\) is provided, \(x\) and \(\text{SNP\textunderscore index}\) will be ignored.

Value

An object of class \texttt{dendrogram} which describes the tree produced by the clustering algorithm \texttt{hclust}.

Examples

library(MASS)
x <- mvrnorm(60,mu = rep(0,60), Sigma = diag(60))
clust.1 <- cluster.snp(x = x, method = "average")
SNP_index <- seq(1,10)
clust.2 <- cluster.snp(x = x, method = "average", SNP_index = SNP_index)
d <- dist(x)
clust.3 <- cluster.snp(d = d, method = "single")
compute.r2

R2 computation

Description
Calculates the R2 of a cluster of SNPs.

Usage
compute.r2(x, y, res.multisplit, covar = NULL, SNP_index = NULL)

Arguments
- **x**: The input matrix, of dimension \(nobs \times nvar\). Each row represents a subject, each column a SNP.
- **y**: The response vector. It can be continuous or discrete.
- **res.multisplit**: The output of `multisplit`.
- **covar**: NULL or the matrix of covariates one wishes to control for, of size \(nobs \times ncovar\).
- **SNP_index**: NULL or the index vector of the cluster of SNPs whose R2 will be computed. See the 'Details' section.

Details
The R2 of a cluster of SNPs is computed on the second half-samples. The cluster members, are intersected with the SNPs selected by the lasso, and the R2 of this model is calculated. Thus, we obtain B R2 values. Finally, the mean of these values is taken. If the value of SNP_index is NULL, the R2 of the full model with all the SNPs will be computed.

Value
The R2 value of the SNP cluster

References

Examples
library(MASS)
x <- mvrnorm(60, mu = rep(0,60), Sigma = diag(60))
beta <- rep(0,60)
betac[5,9,3] <- 1
y <- x %*% beta + rnorm(60)
SNP_index <- c(5,9,3)
res.multisplit <- multisplit(x, y)
r2 <- compute.r2(x, y, res.multisplit, SNP_index = SNP_index)
hierGWAS  
Asessing statistical significance in predictive GWA studies

Description

Testing individual SNPs, as well as arbitrarily large groups of SNPs in GWA studies, using a joint model of all SNPs. The method controls the FWER, and provides an automatic, data-driven refinement of the SNP clusters to smaller groups or single markers.

Details

hierGWAS is a package designed to assess statistical significance in GWA studies, using a hierarchical approach.

There are 4 functions provided: cluster.snp, multisplit, test.hierarchy and compute.r2. cluster.snp performs the hierarchical clustering of the SNPs, while multisplit runs multiple penalized regressions on repeated random subsamples. These 2 functions need to be executed before test.hierarchy, which does the hierarchical testing, though the order in which the 2 functions are executed does not matter. test.hierarchy provides the final output of the method: a list of SNP groups or individual SNPs, along with their corresponding p-values. Finally, compute.r2 computes the explained variance of an arbitrary group of SNPs, of any size. This group can encompass all SNPs, SNPs belonging to a certain chromosome, or an individual SNP.

Author(s)

Laura Buzdugan laura.buzdugan@stat.math.ethz.ch

References


multisplit  
Variable Selection on Random Sample Splits.

Description

Performs repeated variable selection via the lasso on random sample splits.

Usage

multisplit(x, y, covar = NULL, B = 50)

Arguments

x  The SNP data matrix, of size nobs x nvar. Each row represents a subject, each column a SNP.
y  The response vector. It can be continuous or discrete.
covar  NULL or the matrix of covariates one wishes to control for, of size nobs x ncovar.
B  The number of random splits. Default value is 50.
simGWAS

Details

The samples are divided into two random splits of approximately equal size. The first subsample is used for variable selection, which is implemented using glmnet. The first \([\text{nobs}/6]\) variables which enter the lasso path are selected. The procedure is repeated \(B\) times.

If one or more covariates are specified, these will be added unpenalized to the regression.

Value

A data frame with 2 components. A matrix of size \(B \times \lceil \text{nobs}/2 \rceil\) containing the second subsample of each split, and a matrix of size \(B \times \lceil \text{nobs}/6 \rceil\) containing the selected variables in each split.

References


Examples

library(MASS)
\[
x <- mvrnorm(60, mu = rep(0, 200), Sigma = diag(200))
\]
\[
beta <- rep(1, 200)
\]
\[
beta[c(5, 9, 3)] <- 3
\]
\[
y <- x %*% beta + rnorm(60)
\]
\[
res.multisplit <- multisplit(x, y)
\]

simGWAS

Simulated GWAS data

Description

This data set was simulated using PLINK. Please refer to the vignette for more details.

Usage

simGWAS

Format

The dataset contains the following components:

- **SNP.1**: The first SNP, of dimension 500 \( \times \) 1. Each row represents a subject.
- **SNP.1000**: The last SNP, of dimension 500 \( \times \) 1. Each row represents a subject.
- **y**: The response vector. It can be continuous or discrete.
- **sex**: The first covariate, representing the sex of the subjects: 0 for men and 1 for women.
- **age**: The second covariate, representing the age of the subjects.

Value

data.frame
test.hierarchy

Description

Performs hierarchical testing of SNPs.

Usage

test.hierarchy(x, y, dendr, res.multisplit, covar = NULL, SNP_index = NULL, alpha = 0.05, global.test = TRUE, verbose = TRUE)

Arguments

x
The input matrix, of dimension nobs \times nvar. Each row represents a subject, each column a SNP.

y
The response vector. It can be continuous or discrete.

dendr
The cluster tree obtained by hierarchically clustering the SNPs using cluster.snp.

res.multisplit
The output of multisplit.

covar
NULL or the matrix of covariates one wishes to control for, of size nobs \times ncovar.

SNP_index
NULL or the index vector of SNP to be tested. See the 'Details' section.

alpha
The significance level at which the FWER is controlled. Default value is 0.05.

global.test
Specifies whether the global null hypothesis should be tested. Default value is TRUE. See the 'Details' section.

verbose
Report information on progress. Default value is TRUE

Details

The testing is performed on the cluster tree given by dendr. If the SNP data matrix was divided (e.g. by chromosome), and clustered separately, the user must provide the argument SNP_index, to specify which part of the data is being tested.

Testing starts at the highest level, which includes all variables specified by SNP_index, and moves down the cluster tree. It stops when a cluster’s null hypothesis cannot be rejected anymore. The smallest, still significant clusters will be returned.

By default the parameter global.test = TRUE, which means that first the global null hypothesis is tested. If the data is divided (e.g. by chromosome), and clustered separately, this parameter can be set to FALSE once the global null has been rejected. This helps save time.

Value

A list of significant SNP groups with the following components:

SNP_index
The indeces of the SNPs in the group

pval
The p-value of the SNP group
References

Buzdugan, L. et al. (2015), Assessing statistical significance in predictive genome-wide association studies

Examples

```r
library(MASS)
x <- mvrnorm(60, mu = rep(0, 60), Sigma = diag(60))
beta <- rep(0, 60)
beta[c(5, 9, 3)] <- 1
y <- x %*% beta + rnorm(60)
dendr <- cluster.snp(x = x, method = "average")
res.multisplit <- multisplit(x, y)
sign.clusters <- test.hierarchy(x, y, dendr, res.multisplit)
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