Package ‘MultiPhen’

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MultiPhen-package ............................................ 2
mPhen ......................................................... 3
mPhen.assoc .................................................. 5
mPhen.cca ................................................... 6
mPhen.assoc .................................................. 7
mPhen.plotCorrelation ..................................... 8
mPhen.preparePheno ........................................ 8
mPhen.readGenotypes ...................................... 9
mPhen.readPhenoFiles .................................... 10
mPhen.sampleCovar ....................................... 11
MultiPhen-package

Description

MultiPhen is the package containing the function mPhen, which performs association testing between genetic variants (SNPs; CNVs to be added soon) and multiple phenotypes. The primary purpose is for modelling and testing multiple phenotypes jointly by performing an ordinal regression where SNPs are treated as the outcome and multiple phenotypes are predictors; this can have large increases in statistical power to detect genotype-phenotype associations over the univariate approach. However, mPhen can also be used to perform standard univariate linear regression (SNP as predictor) and univariate ordinal regression (SNP as outcome) on the phenotypes under study. mPhen can be applied to genotyped or imputed data. From version 0.4 the option “multiPhenTest” is now called “JointModel”, and its default is now “TRUE”

Details

Package: MultiPhen
Type: Package
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License: GPL-2
LazyLoad: yes

Author(s)

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References

O’Reilly et al. 2012. MultiPhen: Joint model of multiple phenotypes can increase discovery in GWAS. http://dx.plos.org/10.1371/journal.pone.0034861
**mPhen**  

_A function for the genetic association testing of multiple phenotypes_

---

**Description**

mPhen performs association testing between genetic variants (SNPs; CNVs) and multiple phenotypes. The primary purpose is for modelling and testing multiple phenotypes jointly by performing an ordinal regression where SNPs are treated as the outcome and multiple phenotypes are predictors; this can have large increases in statistical power to detect genotype-phenotype associations over the univariate approach (method described in O’Reilly et al. 2012, see below). However, mPhen can also be used to perform standard univariate linear regression (SNP as predictor) and univariate ordinal regression (SNP as outcome) on the phenotypes under study. mPhen can be applied to directly genotyped or imputed data.

**Usage**

```r
mPhen(genoData, phenoData, phenotypes = "all",
       covariates = NULL, resids = NULL, strats = NULL,
       opts = mPhen.options(c("regression","pheno.input")))
```

**Arguments**

- **genoData**
  Either a matrix (for directly measured genotypes) or 3 dimensional array (for imputed genotypes). The first dimension (rows) corresponds to individuals, and row.names are individual IDs. The second dimension corresponds to SNPS, with col.names equal to the snp identifiers. For directly measured genotypes, the value in each cell is a numeric genotype (i.e. AA = 0, AB=1, BB = 2). For imputed genotypes, the 3rd dimension corresponds to genotypes with dimnames(genoData)[[3]] equal to a numeric vector corresponding to genotype values. The values in these cells are the probability of each genotype multiplied by 1000. For copy number genotypes the numeric values correspond to numbers of copies. An example provided by ‘snps’ and ‘snps.imputed’

- **phenoData**
  Matrix containing phenotype data, where each row corresponds to an individual and row.names are individual IDs. Each column contains data on a certain phenotype across the sample of individuals (can be quantitative, case/control or ordinal. Must be numeric); the column header provides the phenotype name. An example is provided by ‘pheno’.

- **phenotypes**
  Vector of phenotype names, to be tested. If value is 'all' then all phenotypes are included after removing covariates and residuals.

- **covariates**
  Vector of phenotypes, from phenoData, to be considered as covariates to be controlled for in the regression (Default is no covariates).

- **resids**
  Vector of residuals, from phenoData, alternative way to adjust for covariates, which pre-calculates offset terms to use in the per SNP regression (Default is no residuals).

- **strats**
  Stratification vector (i.e. cases/controls, exposed/not exposed, male/female etc), from phenoData (Default is no stratification).
opts A list of options, which is obtained from mPhen.options(c("regression","pheno.input")). To get more information about these options, type mPhen.options(c("regression","pheno.input"), descr = TRUE).

Value

Returns a list, with two items. The first item (Results) is a 4 dimensional matrix, with dimensions [strata, snps, phenotypes, result_type], where result_type includes beta, pvalue and Nobs. The second item is a vector of minor allele frequencies.

Note

The user should remember that the genotype data file is always a matrix of at least a column, hence if taking a subset of 1 SNP in the non-imputed genotype data matrix, the option drop = FALSE should be used (see the example below).

Author(s)

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References

O’Reilly et al. 2012. MultiPhen: Joint model of multiple phenotypes can increase discovery in GWAS. http://dx.plos.org/10.1371/journal.pone.0034861

Examples

data(snps); data(snps.imputed); data(pheno)
opts = mPhen.options(c("regression","pheno.input"))
res = mPhen(snps, pheno, phenotypes = "all",
  covariates = c("testPheno3", "testPheno4"), opts = opts)
# performs a MultiPhen analysis, with snp as outcome,
# with testPheno3 and testPheno4 as predictors using ordinal regression
res = mPhen(snps, pheno, phenotypes = c("testPheno1", "testPheno2"),
  covariates = c("testPheno3", "testPheno4"), resids = "testPheno5", opts = opts)
# the same as above, with the fifth phenotype as residual
res = mPhen(snps[,2, drop = FALSE], pheno, phenotypes = c("testPheno1", "testPheno2"),
  covariates = "testPheno3", opts = opts)
# please note the use of drop = FALSE if analysing only one SNP
res = mPhen(snps.imputed, pheno, phenotypes = c("testPheno1", "testPheno2"),
  covariates = "testPheno3", opts = opts)
# for imputed data
A function for the genetic association testing of multiple phenotypes

Description

This function is called by mPhen. If you are doing association on multiple batches of genotype data, it is more efficient to use this function, and to pre-prepare a 'phenoObject' object once and then use this function.

Usage

mPhen.assoc(genoData, phenoObject, opts = mPhen.options("regression"), subinds = 1:(dim(genoData)[1]))

Arguments

- **phenoObject**: A phenotype object prepared by mPhen.preparePheno
- **genoData**: This can be obtained from mPhen.readGenoConnection(...)$genoData. It is either a matrix (for directly measured genotypes) or a 3-dimensional array (for imputed genotypes). The first dimension (rows) corresponds to individuals, and row.names are individual IDs. The second dimension corresponds to SNPS, with col.names equal to the snp identifiers. For directly measured genotypes, the value in each cell is a numeric genotype (i.e., AA = 0, AB = 1, BB = 2). For imputed genotypes, the 3rd dimension corresponds to genotypes (with dimnames(genoData)[[3]]) equal to a numeric vector corresponding to genotype values. The values in these cells are the probability of each genotype multiplied by 1000. For copy number genotypes the numeric values correspond to numbers of copies. An example provided by 'snps'.
- **opts**: A list of options, which is obtained from mPhen.options("regression"). To get more information about these options, type mPhen.options("regression", descr=TRUE)
- **subinds**: This indicates the indices of individuals to include in the analysis. It is possible to have repeat indices (i.e., for bootstrap)

Value

Returns a list, with two items. The first item (Results) is a Results is a 4-dimensional matrix, with dimensions [strata, snps, phenotypes, result_type], where result_type includes beta, pvalue and Nobs. The second item is a vector of minor allele frequencies.
**mPhen.cca**  

_Sparse canonical correlation analysis_

**Description**

Carries out a sparse canonical correlation analysis

**Usage**

```r
mPhen.cca(genoData, phenoObject, opts = mPhen.options("regression"),
subinds = 1:(dim(genoData)[1]),
vs.G = opts$mPhen.variable.selection,
   vs.P = opts$mPhen.variable.selection
)
```

**Arguments**

- **genoData**
  A 2 dimensional array. The first dimension (rows) corresponds to individuals, and row.names are individual IDs. The second dimension corresponds to SNPs, with col.names equal to the snp identifiers. The entries are either genotypes, or expected genotypes.

- **phenoObject**
  A phenotype object prepared by mPhen.preparePheno.

- **opts**
  A list of options, which is obtained from mPhen.options("regression"). To get more information about these options, type mPhen.options("regression",descr=TRUE).

- **vs.G**
  If true performs variable selection on genotypes. Is equal to opts$mPhen.variable_selection by default

- **vs.P**
  If true performs variable selection on phenotypes. Is equal to opts$mPhen.variable_selection by default

- **subinds**
  This indicates the indices of individuals to include in the analysis. It is possible to have repeat indices (i.e. for bootstrap)

**Value**

A list with following elements

- **betasp**
  Phenotype weights

- **betasg**
  Genotype weights

- **resultsGeno**
  Single (combined) phenotype analysis results against all genotypes

- **resultsPheno**
  Multiple phenotype analysis results against single (combined) genotype
**mPhen.defineOptions**

*Defines and modifies options.*

**Description**

This reads options from the command-line, if provided. Also it can replace references to system variables in option values, such as "*" or \$WORK, with fully qualified values. Also translates coordinates which use Gb, Mb, Kb to integer values. All defined options of the form mPhen.xxx are examined and modified.

**Usage**

mPhen.defineOptions(file = NULL, getOptionsFromCommandLine = TRUE)

**Arguments**

- **file** specified if the options are in a script file and not set manually
- **getOptionsFromCommandLine**
  
  If running from a script using command Rscript, then this will read in command line options, such as \`--mPhen.logp=FALSE\`

**Value**

None

---

**mPhen.options**

*Retrieves default mPhen options, and descriptions.*

**Description**

This command is used to get options which can be modified to control the behaviour of MultiPhen commands. It provides a list of options which are relevant to a particular command. For example, mPhen.assoc() has its behaviour controlled by options in mPhen.options("regression"). In order to get a list of all options, you can type mPhen.options(descr=TRUE).

**Usage**

mPhen.options(type=c("regression", "plot", "geno.input", "pheno.input","meta.analysis","misc"), descr = FALSE)

**Arguments**

- **type**
  
  A value which can take any of the following values: "regression", "plot", "geno.input", "pheno.input", "meta.analysis", "misc"

- **descr**
  
  If set to TRUE, then returns descriptions of all the options. If FALSE, then returns the values of all the options.
**mPhen.plotCorrelation**  
Plots correlation between phenotype values conditional on genotype

**Description**

Plots the correlation between phenotype values, with different genotypes coloured differently. Note that this will plot \((\text{dim(pheno\_to\_plot)[2]} - 1) \times \text{dim(geno)[2]}\) plots

**Usage**

```r
mPhen.plotCorrelation(pheno\_to\_plot, geno, title = \"\", cex = 0.25, cols = c(1, 2, 3))
```

**Arguments**

- **pheno\_to\_plot**  
  the phenotype to plot
- **geno**  
  Genotypes to use for stratifying samples
- **title**  
  Title of plots
- **cex**  
  Scaling of points
- **cols**  
  The colours to be used in the plot

**Value**

None

---

**mPhen.preparePheno**  
Prepare phenotype data for analysis

**Description**

This harmonises the phenotype data with genotype data, and also extracts the relevant columns from a larger phenotype matrix, and also pre-calculates stratification indices and residuals. This is called by mPhen function, but quicker to do this just once for batched genotype data.

**Usage**

```r
mPhen.preparePheno(phenoData,  
  pcs = NULL, indiv = if (is.null(pcs)) rownames(phenoData) else rownames(pcs),  
  opts = mPhen.options("regression"))
```
Arguments

phenoData  This is typically the output of mPhen.readPhenoFiles(...) or mPhen.simulate(...). It is a list containing the elements phenoData$pheno and phenoData$limit. The element phenoData$pheno is a matrix containing phenotype data, where each row corresponds to an individual and row.names are individual IDs. Each column contains data on a certain phenotype across the sample of individuals (can be quantitative, case/control or ordinal. Must be numeric); the column header provides the phenotype name. An example is provided by 'pheno'. The element phenoData$limit is a list which specifies which of the phenotypes to use as covariates, variables to associate, etc, in the following way:

limit$phenotypes - vector of phenotypes to be tested. If set to 'all' then all phenotypes are used. limit$covariates - vector of phenotypes to be considered as covariates to be controlled for in the regression, limit$resids - vector of phenotypes to be considered as residuals, which is an alternative way to adjust for covariates, which pre-calculates offset terms to use in the per SNP regression limit$strats - stratification vector (i.e. cases/controls, exposed/not exposed, male/female etc). limit$excls - Exclusion vector, ie. names of phenotypes which should be used as exclusion criteria respectively. Rows will be excluded if the value in any of exclusion columns is NA or 1

Alternatively, phenoData can simply be a matrix containing phenotype data, in which case, the default value of limit used is limit = list(phenotypes="all")

pcs  This is the genotype pcs which should be used in the analysis. If specified it should be in the same sample order as 'inidv'. The user still needs to specify in the phenoData$limit$covariate or phenoData$limit$resids the names of the PCs they wish to include (i.e. covariate = c("PC1","PC2")). These would typically be obtained from mPhen.readGenotypes.

indiv  individuals to be used. If unspecified the function defaults to using all individuals

opts  A list of options, which is obtained from mPhen.options("regression"). To get more information about these options, type mPhen.options("regression",descr=TRUE)

Value

A list object, which can then be used in mPhen.assoc.

mPhen.readGenotypes  Open, and read from a read connection to a genotype file

Description

Opens a read connection to a list of files which have a VCF-like format. Can read a .gz file. Also supports plink bed format. Also supports cnvPipe format. Also supports a .zip file format used by cnvHap.
Usage

mPhen.readGenotypes(genoConnection, indiv = NULL, opts = mPhen.options("geno.input"))

Arguments

genoConnection: A list of paths to genotype files. Each file might represent a different cohort with overlapping SNP set. These files can be .vcf, .vcf.gz, and .zip files. They can also be plink bed files. In the case that bed files are used, the root name of the file should be given (i.e. hapmap2 instead of hapmap2.bed).

Alternatively, this can also be the object returned by mPhen.readGenotypes(...). This means that the connection only has to be established once, and can be read from multiple times in batches.

indiv: A list of individual ids. If provided the results will be in this order. If not provided the results will be in the order given by the genotype file

opts: A list of options, which is obtained from mPhen.options("geno.input"). To get more information about these options, type mPhen.options("geno.input", descr = TRUE)

Value

Returns a list object which can be used by subsequent calls to readGenotypes(....). This list also includes a value genoData, which contains the genotypes which have been read. This list includes a connection to the underlying file (conn), a list of sample ids, and a flag indicating whether the file is zipped. It also includes 'pcs', which is a matrix of genotype pcs. These are only calculated if opts$mPhen.numGenoPCs>0, and are only calculated once all batches have been read (but will include all genotypes which have been read previously using the same connection).

mPhen.readPhenoFiles Read and merge phenotype files

Description

This helper function merges multiple phenotype files into a single phenotype matrix, and applies missing value and exclusion criteria

Usage

mPhen.readPhenoFiles(phenofiles,
   limitFile = getOption("mPhen.limitFile","./limit.txt"),
   excludeFile = getOption("mPhen.excludeFile","./exclude.txt"),
   opts = mPhen.options("pheno.input"),
   indiv = NULL)
mPhen.sampleCovar

Arguments

- **phenoFiles**: A list of paths to phenotype files (can be more than 1)
- **excludeFile**: A path to a file which lists ids to exclude from further analysis, or alternatively is a two column file, with the first column of ids and a second column of numerical values which are used in conjunction with opts$quantileThresh
- **limitFile**: As an alternative to specifying covariates, resids,strats and excl in mPhen.preparePheno(..), you can also specify this information via a limitfile, which is tab delimited file in which the first column specifies the type of variable to set (pheno,covar,resid,strat,excl), the second column specifies the phenotype name, and the third column optionally specifies a transformation Different lines can then be used for different values. The transformation syntax includes 'quantile' and 'factor', and also 'thresh_x_y' in which values less than x are coded 0 and greater than y are coded 1; and also 'toptail_x_y' where values less than x percentile are coded 0 and greater than y percentile are coded 1.
- **opts**: A list of options, which is obtained from mPhen.options("pheno.input"). To get more information about these options, type mPhen.options("pheno.input",descr=TRUE)
- **indiv**: A list of individual ids. If provided the phenotype matrix will be re-ordered to match

Value

An object consisting of a single merged phenotype matrix, and also a 'limit' object which specifies phenotypes to include in analyses. The limit is a list with the following entries: phenotypes - vector of phenotypes to be tested. If set to 'all' then all phenotypes are used. covariates - vector of phenotypes to be considered as covariates to be controlled for in the regression, resids - vector of phenotypes to be considered as residuals, which is an alternative way to adjust for covariates, which pre-calculates offset terms to use in the per SNP regression strats - statification vector (i.e. cases/controls, exposed/not exposed, male/female etc).

- **excls**: Exclusion vector, ie. names of phenotypes which should be used as exclusion criteria respectively. Rows will be excluded if the value in any of exclusion columns is NA or 1

---

mPhen.sampleCovar *Generates a covariance matrix.*

Description

This function can be use to sample covariance matrices. This is useful when simulating data to test Multiphenotype based association strategies. This function lets the user decide on the orthoganality within 'blocks' and between 'blocks' of correlated variables/

Usage

mPhen.sampleCovar(noPhenos, blockSize, orthogAll = c(0.9, 0.5), dirichletScale = 50, resample = FALSE, sd = rgamma(noPhenos, shape=10, rate = 10))
mPhen.sampleGeno

Arguments

- **noPhenos**: The number of phenotypes to simulate
- **blockSize**: The number of phenotypes per covariance block
- **orthogAll**: The orthogonality relationships between and within blocks expressed as a number on the interval \((0,1)\). A number closer to one indicates closer to orthogonality, whereas 0 indicates non-orthogonality. First number is orthogonality between blocks, second is orthogonality within blocks.
- **dirichletScale**: When sampling off diagonal elements of the cholesky decomposition, how much deviation from uniform to allow. Should be a number in interval \((0,+\text{Inf})\). Smaller value leads to greater variation
- **resample**: Whether to randomly shuffle phenotype columns after sampling.
- **sd**: Standard deviation for each phenotype

Value

Simulated covariance matrix

---

**mPhen.sampleGeno**

*Sample genotypes*

Description

...

Usage

```r
mPhen.sampleGeno(n = 100, sampSize = 100, chr="0", pos = 1:n,
snpids = paste(chr, pos, sep="_"), meanAlleleFreq=0.2, mu = 10,
samples = paste("id",1:sampSize,sep="_"), imputed = FALSE, dirichlet = 1)
```

Arguments

- **n**: Number of genotypes to sample
- **sampSize**: Number of individuals to sample
- **chr**: Name of chromosome
- **pos**: Positions of genotypes on chromosome
- **snpids**: Ids of genotypes
- **meanAlleleFreq**: The mean allele frequency to simulate
- **mu**: A weight parameter which controls how close to the meanAlleleFreq the allele frequencies are sampled, via a beta distribution. A higher number implies allele frequencies stay closer to mean
- **samples**: The sample ids
- **imputed**: Whether to simulate imputed data
- **dirichlet**: The weight of a dirichlet distribution used to simulated imputed data
**mPhen.simulate**

Simulates phenotypes according to a correlation structure.

**Description**

This function simulates phenotypes based on a pre-defined correlation structure (which can also be obtained from mPhen.sampleCovar), and a genetic effect x. The function works by sampling a phenotype from a correlation matrix in a linearly transformed space such that the genetic effect direction is only in the direction of the x-axis, then transforming back into the original space. If inverse is TRUE, then the phenotypes are sampled first with no genetic effect, then the genotype is sampled according to the effect direction.

**Usage**

```r
mPhen.simulate(x,sample_names, covar,effDir,
varexp,inverse=FALSE, geno.link="gaussian",
effDirInReverseEigenspace=FALSE, freq = 0.1)
```

**Arguments**

- `x` A vector of genotype effect. If a single SNP has an effect, this will just be genotypes at this SNP.
- `sample_names` Vector of sample names, should have same length as x.
- `covar` Covariance of phenotypes, should be an n by n matrix, where n is the number of phenotypes to simulate.
- `effDir` Direction in phenotype space in which to simulate the effect.
- `varexp` The proportion of variance of phenotype variation in the target direction explained by the genotypic effect overall.
- `inverse` If TRUE, then simulates correlated phenotypes, and then simulates genotypes from phenotypes in specified direction. Otherwise, simulates correlated phenotypes with direction of effect based on input genotypes.
- `geno.link` Only applicable if inverse = TRUE, in which case it specifies a link function for genotypes. Can be binomial, gaussian or ordinal.
- `effDirInReverseEigenspace` If TRUE, then effDir is interpreted as eigenvector weights, ordered from eigenvector with smallest eigenvalue to eigenvector with biggest eigenvector (i.e. in reverse direction). This is useful if you want to simulate directs which are in the least variable axis of variation.
- `freq` If inverse = TRUE, then this is target allele frequency.
Value

Returns a list of three elements: pheno - a matrix of phenotype values, with phenotypes in columns and samples in rows; and geno - a vector genotypes, either sampled if inverse=TRUE, or the original genetic effect x; and limit - this is a default list of phenotypes to include in subsequent association analysis, as well as covariates (empty) and residuals (empty). This object can then be used by mPhen.preparePheno

Description

Writes output to files defined in mPhen.openOutputConnection, and extracts pvalues and betas for further plots.

Usage

mPhen.writeOutput(results,
output = getOption("mPhen.resultsName","resultsDir/"),geno = NULL,
towrite = list(long.txt = getOption("mPhen.writeLong",TRUE),
qc.txt = getOption("mPhen.writeQC",FALSE),
wide.txt = getOption("mPhen.writeWide",TRUE)),
toplots = list(.manh = TRUE, .qq = TRUE,.heatm = TRUE,

.fprint = !is.null(geno)),
opts = mPhen.options("plot")

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>results</td>
<td>Output of mPhen.assoc</td>
</tr>
<tr>
<td>output</td>
<td>Directory to write results, or object returned by mPhen.writeOutput(..)</td>
</tr>
<tr>
<td>towrite</td>
<td>List specifying which formats to write output - long.txt and wide.txt for standard results; qc.txt for per-sample qc output.</td>
</tr>
<tr>
<td>toplot</td>
<td>List specifying which formats to plot output - .qq for qq plot, .manh for manhattan, .heatm for pvalue heatmap, .fprint for fingerprint plot</td>
</tr>
<tr>
<td>geno</td>
<td>Genotype matrix. Note that attr(geno,&quot;closeConnection&quot;) controls whether plots are produced, as this indicates whether all batches of genotype data have been analysed</td>
</tr>
<tr>
<td>opts</td>
<td>A list of options, which is obtained from mPhen.options(&quot;plot&quot;). To get more information about these options, type mPhen.options(&quot;plot&quot;,descr=TRUE)</td>
</tr>
</tbody>
</table>

Value

Returns an outputConnection, which can be used to write further results.
A dummy phenotype dataset that provides an example of the input phenotype data used by the package

Description

A dummy dataset of 5 phenotypes measured in 150 individuals. The data has been generated to yield significant results for SNP1 and SNP2 of the snps dataset. The first two columns have been generated as alpha + beta1*snp + beta2*snp2 + error (with different alphas, betas and errors for each phenotype), the third has been generated as alpha + beta1*testPheno2 + beta2*snp3 + error, the fourth column is the results of sample of a binomial distribution correlated with testPheno3, and the final column is the 1st PC of the principal component analysis of the snps matrix.

Format

A matrix with 150 phenotype observations.

testPheno1 a numeric vector
testPheno2 a numeric vector
testPheno3 a numeric vector
testPheno4 a numeric vector
testPheno5 a numeric vector

Details

Please note the following IMPORTANT issue: the 'pheno' matrix has both column names and row names! The column names MUST be the names of the phenotypes and the row names MUST be the codes representing each individual in the pheno matrix, one individual for each row. Both row names and column names are extracted by the main function and are therefore mandatory.

Examples

data(pheno)
head(pheno)
dimnames(pheno)[[1]] # the row names
dimnames(pheno)[[2]] # the column names
read.plink

A function to read (small) binary PLINK binary files in a R session

Description

read.plink is a convenience function designed to read PLINK binary files (i.e. files that end with the suffix "\b.bed") in a R session. Please be aware that binary PLINK files are binary for a reason, i.e. to store genotype data in a compact way. Once they are imported in R they exist in R in a un "unpacked" form, and can therefore be very big. If the \b.bed file is big, or very big, the result will be that R will run out of memory and crash, or make the whole system slow or unresponsive. It is MANDATORY that in the directory containing the binary file also reside two accesory files, with the same name as the binary file but with extensions \b.fam and \b.bim, both produced by PLINK.

Usage

read.plink(root, indiv = NULL, opts = mPhen.options("geno.input"))

Arguments

- \texttt{root}: filename of the dataset in PLINK binary format, WITHOUT the \b.bed extension.
- \texttt{indiv}: List of individuals, results will be in this order
- \texttt{opts}: List of options, use mPhen.options("geno.input", descr=TRUE) for more details of each option.

Details

Please note that, if the binary file is listed a "\b.mydata.bed", the filename is "\b.mydata", and the exten-sion is "\b.bed". In this case "\b.mydata" would be used as \texttt{root} value.

Value

A matrix of dimensions n by m, with n rows corresponding to the n individuals in the dataset, and m columns corresponding to the m markers. The colnames are retrieved from the \b.fam file, and (should) correspond to the markers’ names.

Note

Please do note that the concept of a “big” binary file, or a binary file that is “too big” is purely dependent on the computer on which the code is running. A computer with 512MB of RAM will stop being able to read in a whole binary file well before a 16GB RAM machine.

Author(s)

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References

The plink homepage is at: \url{https://www.cog-genomics.org/plink2}
Description

A dummy dataset of three SNPs, as a matrix of 3 column and 150 rows. The genotypes are in 0/1/2 format (0 for “AA”, 1 for “Aa” and 2 for “aa”, where A and a correspond, arbitrarily, to the two alleles). The data has been randomly generated, for instructional purposes only, and do not yield a significant association with any of the example phenotypes.

Usage

data(snps)

Format

A data frame with 150 genotype observations.

rsID1 a numeric vector
rsID2 a numeric vector
rsID3 a numeric vector

Details

The 150 genotypes in ‘snp’ correspond to the phenotype data on 150 individuals in ‘pheno’, i.e. one individual for each line. Please note the following important points: genotype data must be in matrix format, with one row for each individual and as many columns for each SNP. In the case of one single genotype the data must still conform to this format, as a matrix of as many rows as individuals and one single column for the one genotype present. A second important point is that the column names must be the rsID of the SNP for genotypes in the 0/1/2 format. Further options of genotype format (incorporating raw genotype data, and CNV genotypes) will be available and documented in future releases.

Examples

data(snps)
dim(snps)
colnames(snps)
Description

A toy dataset of three imputed SNP, for 150 individuals. For each individual, and for each SNP, the first column is the probability of a minor allele homozygote genotype (genotype “0”), the second column is the probability of an heterozygote genotype (genotype “1”) and the third and last column is the probability of a major allele homozygote (genotype “2”).

Usage

data(snps.imputed)

Format

A matrix of 150 rows and nine columns.

Examples

data(snps.imputed)
Index

*Topic GWAS
  mPhen, 3
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  MultiPhen-package, 2

*Topic association
  mPhen, 3
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  MultiPhen-package, 2

*Topic connection
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  MultiPhen-package, 2
  read.plink, 16

*Topic data imput
  read.plink, 16

*Topic datasets
  pheno, 15
  snps, 17
  snps.imputed, 18

*Topic file
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  read.plink, 16

*Topic genetics
  mPhen, 3
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  MultiPhen-package, 2
  read.plink, 16

*Topic multiple phenotypes
  mPhen, 3
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  MultiPhen-package, 2

*Topic package

19
MultiPhen-package, 2

+ Topic regression
  mPhen, 3
  mPhen.assoc, 5
  mPhen.cca, 6
  mPhen.defineOptions, 7
  mPhen.options, 7
  mPhen.plotCorrelation, 8
  mPhen.preparePheno, 8
  mPhen.readGenotypes, 9
  mPhen.readPhenoFiles, 10
  mPhen.writeOutput, 14
  MultiPhen-package, 2

mPhen, 3
mPhen.assoc, 5
mPhen.cca, 6
mPhen.defineOptions, 7
mPhen.options, 7
mPhen.plotCorrelation, 8
mPhen.preparePheno, 8
mPhen.readGenotypes, 9
mPhen.readPhenoFiles, 10
mPhen.sampleCovar, 11
mPhen.sampleGeno, 12
mPhen.simulate, 13
mPhen.writeOutput, 14
MultiPhen (MultiPhen-package), 2
MultiPhen-package, 2

pheno, 15
read.plink, 16
snps, 17
snps.imputed, 18