## Package 'GWAS.BAYES'

March 30, 2021

## Type Package

Title GWAS for Selfing Species
Version 1.0.0
Description This package is built to perform GWAS analysis for selfing species. The research related to this package was supported in part by National Science Foundation Award 1853549.
License GPL-2 I GPL-3
Encoding UTF-8
LazyData true
biocViews AssayDomain, SNP
Depends R (>= 4.0), Rcpp (>= 1.0.3), RcppEigen ( $>=0.3 .3 .7 .0$ ), GA ( $>=$
3.2), caret ( $>=6.0-86$ ), ggplot2 ( $>=3.3 .0$ ), doParallel ( $>=$
1.0 .15 ), memoise ( $>=1.1 .0$ ), reshape2 $(>=1.4 .4)$, Matrix ( $>=$ 1.2-18)

LinkingTo RcppEigen ( $>=$ 0.3.3.7.0), Rcpp ( $>=1.0 .3$ )
Suggests BiocStyle, knitr, rmarkdown, formatR, rrBLUP, qqman
VignetteBuilder knitr
git_url https://git.bioconductor.org/packages/GWAS.BAYES
git_branch RELEASE_3_12
git_last_commit 919eb22
git_last_commit_date 2020-10-27
Date/Publication 2021-03-29
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## Description

Aggregate SNPs and Y by Species

## Usage

aggregate_SNPs(SNPs,Y, na.rm)

## Arguments

| SNPs | Standardized SNP data set where the values of each column are either 0 or 1 |
| :--- | :--- |
| $Y$ | The phenotype response of interest |
| na.rm | Logical value where TRUE removes NA's in response vector as well correspond- <br> ing rows in SNP matrix. |

## Value

SNPs reduced SNP dataset
Y

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
SNPs <- standardize(SNPs = SNPs,method = "major-minor",number_cores = 1)
aggregate_SNPs(SNPs = SNPs, Y = Y)
```

```
cor_plot cor_plot(SNPs,significant,info = FALSE)
```


## Description

A function that creates correlation heatmaps for given significant SNPs from a SNP dataframe.

## Usage

```
cor_plot(SNPs,significant,info = FALSE)
```


## Arguments

SNPs A standardized SNP matrix where the columns take on the values of 0 or 1 .
significant
A vector of 0's or 1's that contain which significant SNP's to look at.
info Default is FALSE. If information such as the chromosome and the position is known, then that can be entered here and will return a correlation heatmap with the labels chromosome - position. The format for this entry is the 2 row dataframe or matrix, where the first row is the chromosome and the second row is the position.

## Value

A correlation heatmap with the axis labels either SNP1 ... or chromosome - position

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
fullPreprocess <- preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01,number_cores = 1)
SNPs <- fullPreprocess$SNPs
Y <- fullPreprocess$Y
fullPreprocess$SNPs_Dropped
principal_comp <- pca_function(SNPs = SNPs,number_components = 1,plot_it = FALSE)
Significant_SNPs <- preselection(Y = Y, SNPs = SNPs,number_cores = 1, principal_components = principal_comp,fr
cor_plot(SNPs = SNPs,significant = Significant_SNPs$Significant,info = FALSE)
```

eigenMapMatMult2 eigenMapMatMult2

## Description

Matrix multiplication in $\mathrm{C}++$ between two matrices.

## Usage

eigenMapMatMult2(A, B)

## Arguments

A
First Matrix
B
Second Matrix

## Value

Returns the matrix multiplication of $A * B$

```
eigenMapMatMult3 eigenMapMatMult3
```


## Description

Matrix multiplication between 3 matrices $(\mathrm{A} * \mathrm{~B} * \mathrm{C})$

## Usage

eigenMapMatMult3(A, B, C)

## Arguments

| A | First Matrix |
| :--- | :--- |
| B | Second Matrix |
| C | Third Matrix |

## Value

The result of $A * B * C$

```
ga_modelselection_nopc
ga_modelselection_nopс
```


## Description

Performs GA model selection to identify the best model when no principal components are involved

## Usage

ga_modelselection_nopc(Y,X,significant,number_cores,maxiterations,runs_til_stop,kinship = FALSE)

## Arguments

Y The phenotype response on the reduced scale (aggregating phenotype by ecotype/taxa), this should be a matrix with 1 column.
$X \quad$ The SNP matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
significant A vector of 0 and 1's where the 1's indicate what SNP's were found to be significant in the preselection function.
number_cores Number of cores to be passed on to the genetic algorithm to increase computational speed.
maxiterations This is the maximum number of iterations the Genetic Search algorithm will run.
runs_til_stop This is the numebr of consectutive iterations where the BIC is not improved before the genetic algorithm is stopped.
kinship Default is set at FALSE. If kinship model is desired, input a kinship matrix and this will search models with the kinship component.

## Details

This function will print out lines correpsonding to the convergence of the genetic search algorithm.

## Value

A named matrix where the names corespond to the significant SNP's. This will usually out a matrix with a singular row, where the values of this row is 0 or 1 where 1 indicates significance in the final model and 0 indicates non significance in the final model. Sometimes this will output a matrix with mulitple columns. This is because there is a SNP or multiple SNPs that when added to the model create rank deficiency issues in the model. Naturally rank deficient columns are forced out but the genetic algoritm is not smart enough to sort these

```
ga_modelselection_nopc_new
    ga_modelselection_nopc_new
```


## Description

Performs GA model selection to identify the best model when no principal components are involved. Internal function for modelselection_new().

## Usage

ga_modelselection_nopc_new( $\mathrm{Y}, \mathrm{X}$, regions,regionsnames, significant, number_cores, maxiterations,runs

## Arguments

$Y \quad$ The phenotype response on the reduced scale (aggregating phenotype by ecotype/taxa), this should be a matrix with 1 column.
$X \quad$ The SNP matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
significant A vector of 0 and 1's where the 1's indicate what SNP's were found to be significant in the preselection function.
regions A matrix where each column represents a principal component for each region.
regionsnames A named list which highlights which SNPs fall into which region.
number_cores Number of cores to be passed on to the genetic algorithm to increase computational speed.
maxiterations This is the maximum number of iterations the Genetic Search algorithm will run.
runs_til_stop This is the number of consecutive iterations where the BIC is not improved before the genetic algorithm is stopped.
kinship The kinship matrix associated with the SNPs in this analysis.

## Details

This function will print out lines corresponding to the convergence of the genetic search algorithm.

## Value

A named matrix where the names correspond to the significant SNP's. This will usually out a matrix with a singular row, where the values of this row is 0 or 1 where 1 indicates significance in the final model and 0 indicates non significance in the final model. Sometimes this will output a matrix with multiple columns. This is because there is a SNP or multiple SNPs that when added to the model create rank deficiency issues in the model. Naturally rank deficient columns are forced out but the genetic algorithm is not smart enough to sort these.

```
ga_modelselection_pcs ga_modelselection_pcs
```


## Description

Performs GA model selection to identify the best model when principal components are involved

## Usage

ga_modelselection_pcs(Y,X, significant, number_cores, principal_components,maxiterations,runs_til_

## Arguments

Y The phenotype response on the reduced scale (aggregating phenotype by ecotype/taxa), this should be a matrix with 1 column.
$X \quad$ The SNP matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
significant A vector of 0 and 1's where the 1's indicate what SNP's were found to be significant in the preselection function.
number_cores Number of cores to be passed on to the genetic algorithm to increase computational speed.
principal_components
The principal component matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
maxiterations This is the maximum number of iterations the Genetic Search algorithm will run.
runs_til_stop This is the numebr of consectutive iterations where the BIC is not improved before the genetic algorithm is stopped.
kinship Default is set at FALSE. If kinship model is desired, input a kinship matrix and this will search models with the kinship component.

## Details

This function will print out lines correpsonding to the convergence of the genetic search algorithm.

## Value

A named matrix where the names corespond to the significant SNP's. This will usually out a matrix with a singular row, where the values of this row is 0 or 1 where 1 indicates significance in the final model and 0 indicates non significance in the final model. Sometimes this will output a matrix with mulitple columns. This is because there is a SNP or multiple SNPs that when added to the model create rank deficiency issues in the model. Naturally rank deficient columns are forced out but the genetic algoritm is not smart enough to sort these.

```
ga_modelselection_pcs_new
    ga_modelselection_pcs_new
```


## Description

Performs GA model selection to identify the best model when principal components are involved

## Usage

ga_modelselection_pcs_new(Y,X,regions,regionsnames, significant, number_cores, principal_components

## Arguments

Y The phenotype response on the reduced scale (aggregating phenotype by ecotype/taxa), this should be a matrix with 1 column.
$X \quad$ The SNP matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
significant A vector of 0 and 1's where the 1's indicate what SNP's were found to be significant in the preselection function.
regions A matrix where each column represents a principal component for each region.
regionsnames A named list which highlights which SNPs fall into which region.
number_cores Number of cores to be passed on to the genetic algorithm to increase computational speed.
principal_components
The principal component matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
maxiterations This is the maximum number of iterations the Genetic Search algorithm will run.
runs_til_stop This is the numebr of consectutive iterations where the BIC is not improved before the genetic algorithm is stopped.
kinship Default is set at FALSE. If kinship model is desired, input a kinship matrix and this will search models with the kinship component.

## Details

This function will print out lines correpsonding to the convergence of the genetic search algorithm.

## Value

A named matrix where the names corespond to the significant SNP's. This will usually out a matrix with a singular row, where the values of this row is 0 or 1 where 1 indicates significance in the final model and 0 indicates non significance in the final model. Sometimes this will output a matrix with mulitple columns. This is because there is a SNP or multiple SNPs that when added to the model create rank deficiency issues in the model. Naturally rank deficient columns are forced out but the genetic algoritm is not smart enough to sort these.

```
level_function level_function
```


## Description

Removes all SNPs that only have one level in it

## Usage

level_function(SNPs,MAF = 0.01)

## Arguments

SNPs The standardized SNP data where the columns take on the values of 0 or 1
MAF The minor allele frequency at which to drop SNPs. Default is set to 0.01 , meaning if the minor allele occurs less than 1 percent of the time in a given SNP, that given SNP will be dropped from the dataset.

## Value

SNPs The SNP matrix where columns that were either all 1's or all 0's are removed
SNPs_Dropped A true/false vector with length ncol(SNPs), where the TRUE's indicate that the column was not dropped and the FALSE's indicate that the column was dropped

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
SNPs <- standardize(SNPs = SNPs,method = "major-minor",number_cores = 1)
list1 <- aggregate_SNPs(SNPs = SNPs, Y = Y)
SNPs <- list1[[1]]
Y <- list1[[2]]
level_function(SNPs, MAF = .01)
```

log_profile_likelihood_REML
log_profile_likelihood_REML

## Description

The log likelihood that needs to be optimized for the full kinship model.

## Usage

log_profile_likelihood_REML(x,t,y,d)

## Arguments

X
The reduced design matrix with principal components, intercept and SNP of interest
$t \quad$ Tau value for the random effect term
$y \quad$ The reduced matrix for the response value of interest
d The spectral decomposition diagonal matrix of eigen values

## Value

This returns the REML value

```
optim_llik_RE_BIC optim_llik_RE_BIC
```


## Description

Calculates the BIC in the full kinship model scenario

## Usage

optim_llik_RE_BIC(x,y,d)

## Arguments

x
$y \quad$ The reduced response matrix.
d The spectral decomposition diagonal matrix of eigen values

## Value

Returns the BIC for the model with this SNP

```
optim_llik_RE_p optim_llik_RE_p
```


## Description

This will calculate the p -value and perform the optimization in the full kinship model case

## Usage

optim_llik_RE_p(x,y,d)

## Arguments

$x \quad$ Reduced design matrix with principal components, intercept and SNP of interest
$y \quad$ The reduced matrix of the response value of interest
d The spectral decomposition diagonal matrix of eigen values

## Value

Returns a p-value for the specified data

```
optim_llik_SLR_BIC optim_llik_SLR_BIC
```


## Description

Calculates the BIC in the SLR scenario

## Usage

optim_llik_SLR_BIC(x,y)

## Arguments

$x \quad$ The reduced design matrix that includes intercept, SNP, and principal components
y
The reduced response matrix

## Value

The BIC for the given data

```
optim_llik_SLR_p optim_llik_SLR_p
```


## Description

This calculates the p -value in the simple linear regression scenario

## Usage

optim_llik_SLR_p(x,y)

## Arguments

$x \quad$ Design Matrix for a single SNP.
y Phenotype Response

```
pca_function pca_function
```


## Description

Create Principal Components from Standardized Set of SNPs.

## Usage

pca_function(SNPs,number_components,plot_it)

## Arguments

SNPs The SNP matrix where the columns consist of 0 and 1's.
number_components
The number of principal components desired, if you don't know put a random value and use plot_it = TRUE.
plot_it A TRUE FALSE logical equality, if TRUE will plot the percent variation explained by the components, if FALSE will not create a plot. In both scenarios this function will return a matrix of principal components.

## Details

This will work with both the full SNP matrix and the reduced SNP matrix. If you use the full SNP matrix you will have to reduce it yourself and this is at a higher computational burden. If you use the reduced SNP matrix you will get the same values as if you aggregated the principal components from the full SNP matrix, but this will be much faster.

## Value

| Plot | A plot of the percent variation explained by the components when plot_it $=$ |
| :--- | :--- |
| TRUE |  |
| Matrix | A matrix that the number of columns is the number of principal components and |
| the number of rows is the same number of rows as the inputted data matrix. |  |

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
fullPreprocess <- preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01,number_cores = 1)
SNPs <- fullPreprocess$SNPs
Y <- fullPreprocess$Y
fullPreprocess$SNPs_Dropped
pca_function(SNPs = SNPs,number_components = 3,plot_it = TRUE)
```

```
postGWAS postGWAS
```


## Description

Performs GA model selection to identify the best model

## Usage

postGWAS(Y, SNPs, significant, number_cores, principal_components, maxiterations, runs_til_stop,kinshi

## Arguments

Y The phenotype response on the reduced scale (aggregating phenotype by ecotype/taxa), this should be a matrix with 1 column.

SNPs The SNP matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
significant A vector of 0 and 1's where the 1's indicate what SNP's were found to be significant in the preselection function.
number_cores Number of cores to be passed on to the genetic algorithm to increase computational speed.
principal_components
The principal component matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
maxiterations This is the maximum number of iterations the Genetic Search algorithm will run.
runs_til_stop This is the number of consecutive iterations where the BIC is not improved before the genetic algorithm is stopped.
kinship Default is set at FALSE. If kinship model is desired, input a kinship matrix and this will search models with the kinship component.
info Default is set at FALSE. An information matrix where the first row is the chromosomes and the second row is the position information

## Details

This function will print out lines corresponding to the convergence of the genetic search algorithm.

## Value

A named matrix where the names correspond to the significant SNP's. This will usually out a matrix with a singular row, where the values of this row is 0 or 1 where 1 indicates significance in the final model and 0 indicates non significance in the final model. Sometimes this will output a matrix with multiple columns. This is because there is a SNP or multiple SNPs that when added to the model create rank deficiency issues in the model. Naturally rank deficient columns are forced out but the genetic algorithm is not smart enough to sort these.

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
fullPreprocess <- preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01, number_cores = 1)
SNPs <- fullPreprocess$SNPs
Y <- fullPreprocess$Y
fullPreprocess$SNPs_Dropped
principal_comp <- pca_function(SNPs = SNPs,number_components = 1,plot_it = FALSE)
Significant_SNPs <- preselection(Y = Y, SNPs = SNPs,number_cores = 1, principal_components = principal_comp,fr
postGWAS(Y = Y, SNPs = SNPs,number_cores = 1, significant = Significant_SNPs$Significant,principal_components =
```

postGWAS_Haplotype postGWAS_Haplotype

## Description

Performs GA model selection to identify the best model

## Usage

postGWAS_Haplotype(Y,SNPs,info,size $=10$, significant, number_cores,principal_components, maxiterat

## Arguments

Y The phenotype response on the reduced scale (aggregating phenotype by ecotype/taxa), this should be a matrix with 1 column.

SNPs The SNP matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
info A dataframe where the first row is the chromosome info for the SNP's and the second is the location of each SNP represented by its base pair.
size The number of kilobase pairs to search for regions.
significant A vector of 0 and 1's where the 1's indicate what SNP's were found to be significant in the preselection function.
number_cores Number of cores to be passed on to the genetic algorithm to increase computational speed.
principal_components
The principal component matrix on the reduced scale (aggregating phenotype by ecotype/taxa).
maxiterations This is the maximum number of iterations the Genetic Search algorithm will run.
runs_til_stop This is the number of consecutive iterations where the BIC is not improved before the genetic algorithm is stopped.
kinship Default is set at FALSE. If kinship model is desired, input a kinship matrix and this will search models with the kinship component.

## Details

This function will print out lines corresponding to the convergence of the genetic search algorithm.

## Value

A named matrix where the names correspond to the significant SNP's. This will usually out a matrix with a singular row, where the values of this row is 0 or 1 where 1 indicates significance in the final model and 0 indicates non significance in the final model. Sometimes this will output a matrix with multiple columns. This is because there is a SNP or multiple SNPs that when added to the model create rank deficiency issues in the model. Naturally rank deficient columns are forced out but the genetic algorithm is not smart enough to sort these.

## Examples

```
data("RealDataSNPs_Y")
Y <- RealDataSNPs_Y$Phenotype
SNPs <- subset(RealDataSNPs_Y,select = -c(Phenotype))
fullPreprocess <- preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01,number_cores = 1,na.rm = FALSE)
SNPs <- fullPreprocess$SNPs
Y <- fullPreprocess$Y
data("RealDataInfo")
RealDataInfo <- RealDataInfo[,-fullPreprocess$SNPs_Dropped]
data("RealDataKinship")
kinship <- as.matrix(RealDataKinship)
Significant_SNPs <- preselection(Y = log(Y), SNPs = SNPs,number_cores = 1, principal_components = FALSE,frequer
#postGWAS_Haplotype(Y = log(Y),SNPs = SNPs,info = RealDataInfo,size = 10,number_cores = 1, significant = Signif
```

```
preprocess_SNPs preprocess_SNPs
```


## Description

This functions takes raw SNP data and the associated phenotype response and returns a SNP dataset and phenotype response variable that can be used in the preselection function.

## Usage

preprocess_SNPs(SNPs,Y,MAF $=0.01$, number_cores, na.rm)

## Arguments

SNPs SNP data where each column is a SNP and the SNP column takes on the values A, C, T, or G.
Y The phenotype response of interest. Should be a numeric vector.
MAF The minor allele frequency at which to drop SNPs. Default is set to 0.01 , meaning if the minor allele occurs less than 1 percent of the time in a given SNP, that given SNP will be dropped from the dataset.
number_cores The number of cores one would wish to parallelize over.
na.rm If there is NA's in the vector Y, set na.rm = TRUE and the Y values that are NA will be removed as well as the corresponding rows of the SNP matrix.

## Value

| SNPs | A new SNP matrix. The matrix will be formatted so the minor allele's are coded <br> as 0's and the major allele's are coded as 1's. This matrix will have columns <br> dropped that have minor allele frequency less than the specified value. It will <br> also aggregate over replications, so SNP's and the vector Y will be aggregated <br> according to replications in the SNP matrix. |
| :--- | :--- |
| Y | The new aggregated response vector Y. If you did not have any replications then <br> this vector will be the exact same as the one entered. |
| SNPs_Dropped $\quad$This will tell you which SNPs were dropped if the had minor allele frequency <br> less than the specified value, it will be in the form of column index number. If <br> no SNPs were dropped this will be the character string "None". |  |

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01,number_cores = 1,na.rm = FALSE)
```

```
preselection preselection
```


## Description

Finds significant SNP's

## Usage

preselection(Y, SNPs, number_cores, principal_components, frequentist, controlrate, threshold, nullprob

## Arguments

Y The reduced matrix of response values
SNPs The reduced SNP matrix where the columns are either 1's or 0's.
number_cores The number of cores on which you would like to parallize this procedure principal_components

The reduced matrix of the principal components.
frequentist A logical value to see whether one would like to use a frequentist multiple comparison test or Bayesian False Discovery based on BIC's. The value of this affects whether values of the next parameters are needed.
controlrate Only used when frequentist $=$ TRUE. This is for which multiple comparison method you would like to use. Examples are "bonferroni" and "BH". See p.adjust for a full list of methods.
threshold The value at which type 1 error rate is held at. .05 in most common literature. Used when frequentist is TRUE or FALSE
nullprob Used when frequentist = FALSE, the probability that is assigned to the null hypothesis.

| alterprob | Used when frequentist = FALSE, the probability that is assigned to the alternate <br> hypothesis. |
| :--- | :--- |
| kinship | The kinship matrix if a model with a kinship component is desired. If not set <br> kinship = FALSE. |
| info | An information matrix where the first row is the chromosome information and <br> the second row in the position information for the SNP's. |

## Value <br> Frequentist Matrix

The matrix of results when Frequentist $=$ TRUE. The results are formatted as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The $\mathrm{P}_{-}$values column will be the p -values that were calculated for each SNP.

## Bayesian Matrix

The matrix of results when Frequentist $=$ FALSE. The results are formatted as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The ApprPosteriorProbs column will be the Approximate Posterior Probabilities that were calculated for each SNP.

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
fullPreprocess <- preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01,number_cores = 1)
SNPs <- fullPreprocess$SNPs
Y <- fullPreprocess$Y
fullPreprocess$SNPs_Dropped
principal_comp <- pca_function(SNPs = SNPs,number_components = 1,plot_it = FALSE)
preselection(Y = Y, SNPs = SNPs, number_cores = 1, principal_components = principal_comp,frequentist = TRUE,con
```

```
preselection_nopc preselection_nopc
```


## Description

Finds significant SNP's when no principal components are present.

## Usage

preselection_nopc ( $\mathrm{Y}, \mathrm{X}$, number_cores, frequentist, controlrate, threshold, nullprob, alterprob,kinship

## Arguments

$Y \quad$ The reduced matrix of response values
X The reduced SNP matrix where th columns are either 1's or 0's.
number_cores The number of cores on which you would like to parallize this procedure

| frequentist | A logical value to see whether one would like to use a frequentist multiple com- <br> parison test or Bayesian False Discovery based on BIC's. The value of this <br> affects whether values of the next parameters are needed. |
| :--- | :--- |
| controlrate | Only used when frequentist = TRUE. This is for which multiple comparison <br> method you would like to use. Examples are "bonferroni" and "BH". See <br> p.adjust for a full list of methods. <br> The value at which type 1 error rate is held at. . 05 in most common literature. <br> Used when frequentist is TRUE or FALSE |
| threshold | Used when frequentist = FALSE, the probability that is assigned to the null <br> hypothesis. |
| nullprob | Used when frequentist = FALSE, the probability that is assigned to the alternate <br> hypothesis. |
| kinship | The kinship matrix if a model with a kinship component is desired. If not set <br> kinship = FALSE. |

## Value <br> Frequentist Matrix

The matrix of results when Frequentist $=$ TRUE. The results are formated as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The $\mathrm{P}_{-}$values column will be the p -values that were calculated for each SNP.
Bayesian Matrix
The matrix of results when Frequentist $=$ FALSE. The results are formated as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The ApprPosteriorProbs column will be the Approximate Posterior Probabilities that were calculated for each SNP.

```
preselection_pc preselection_pc
```


## Description

Finds significant SNP's when principal components are present

## Usage

preselection_pc( $\mathrm{Y}, \mathrm{X}$, number_cores, principal_components, frequentist, controlrate, threshold, nullprob

## Arguments

$Y \quad$ The reduced matrix of response values
$\mathrm{X} \quad$ The reduced SNP matrix where th columns are either 1's or 0's.
number_cores The number of cores on which you would like to parallize this procedure principal_components

The reduced matrix of the principal components.
frequentist A logical value to see whether one would like to use a frequentist multiple comparison test or Bayesian False Discovery based on BIC's. The value of this affects whether values of the next parameters are needed.

```
    controlrate Only used when frequentist = TRUE. This is for which multiple comparison
        method you would like to use. Examples are "bonferroni" and "BH". See
        p.adjust for a full list of methods.
    threshold The value at which type 1 error rate is held at. . 05 in most common literature.
        Used when frequentist is TRUE or FALSE
    nullprob Used when frequentist \(=\) FALSE, the probability that is assigned to the null
        hypothesis.
    alterprob Used when frequentist = FALSE, the probability that is assigned to the alternate
        hypothesis.
    kinship The kinship matrix if a model with a kinship component is desired. If not set
        kinship = FALSE.
```


## Value

Frequentist Matrix
The matrix of results when Frequentist $=$ TRUE. The results are formated as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The $P_{-}$values column will be the $p$-values that were calculated for each SNP.
Bayesian Matrix
The matrix of results when Frequentist $=$ FALSE. The results are formated as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The ApprPosteriorProbs column will be the Approximate Posterior Probabilities that were calculated for each SNP.

## Pval_function Pval_function

## Description

Performs multiple comparison corrections on p-values and returns significant SNP's. This is used internally for the preselection function.

## Usage

Pval_function(p_vals,n, thresh, control)

## Arguments

p_vals A vector of p-values calculated by the preselection function.
$\mathrm{n} \quad$ The original number of SNPs
thresh The type 1 error rate
control The multiple comparison correction one would like to apply.

## Value

The results are formated as a data.frame with the column Significant being 1 or 0 depending on if the SNP was significant ( 1 for significant). The P_values column will be the p-values that were calculated for each SNP.

## RealDataInfo RealDataInfo

## Description

A information matrix, where the first row is the chromosome information and the second row is the position information.

## Usage

data("RealDataInfo")

## Format

A data frame with 2 observations on the following 1500 variables.
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V1111 a numeric vector
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V1121 a numeric vector
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V1123 a numeric vector
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V1125 a numeric vector
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V1128 a numeric vector
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V1198 a numeric vector
V1199 a numeric vector
V1200 a numeric vector
V1201 a numeric vector
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V1497 a numeric vector
V1498 a numeric vector
V1499 a numeric vector
V1500 a numeric vector
V1501 a numeric vector
V1502 a numeric vector
V1503 a numeric vector
V1504 a numeric vector
V1505 a numeric vector
V1506 a numeric vector

## Examples

data(RealDataInfo)

RealDataKinship RealDataKinship

## Description

A kinship matrix for a section of the vignette for GWAS.BAYES.

## Usage

data("RealDataKinship")

## Format

A data frame with 328 observations on the following 328 variables.
V1 a numeric vector
V2 a numeric vector
V3 a numeric vector
V4 a numeric vector
V5 a numeric vector
V6 a numeric vector
V7 a numeric vector
V8 a numeric vector
V9 a numeric vector
V10 a numeric vector
V11 a numeric vector
V12 a numeric vector
V13 a numeric vector
V14 a numeric vector
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V136 a numeric vector
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V141 a numeric vector
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V211 a numeric vector
V212 a numeric vector
V213 a numeric vector
V222
V22 a a numeric vector
V22 a numeric vector
V22 a numeric vector
V22 a numeric vector
V215 a numeric vector
V216 a numeric vector
V217 a numeric vector
V218 a numeric vector
V219 a numeric vector
V22 a numeric vector
Vector

V229 a numeric vector
V230 a numeric vector
V231 a numeric vector
V232 a numeric vector
V233 a numeric vector
V234 a numeric vector
V235 a numeric vector
V236 a numeric vector
V237 a numeric vector
V238 a numeric vector
V239 a numeric vector
V240 a numeric vector
V241 a numeric vector
V242 a numeric vector
V243 a numeric vector
V244 a numeric vector
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V292 a numeric vector
V293 a numeric vector
V294 a numeric vector
V305 a numeric vector
V306 a numeric vector
V30 a numeric vector
V30 a numeric vector
V296 a numeric vector
V2 numeric vector
V297 a numeric vector
V298 a numeric vector
V299 a numeric vector
V300 a numeric vector
Vector

V309 a numeric vector
V310 a numeric vector
V311 a numeric vector
V312 a numeric vector
V313 a numeric vector
V314 a numeric vector
V315 a numeric vector
V316 a numeric vector
V317 a numeric vector
V318 a numeric vector
V319 a numeric vector
V320 a numeric vector
V321 a numeric vector
V322 a numeric vector
V323 a numeric vector
V324 a numeric vector
V325 a numeric vector
V326 a numeric vector
V327 a numeric vector
V328 a numeric vector

## Examples

data(RealDataKinship)

```
RealDataSNPs_Y RealDataSNPs_Y
```


## Description

A dataset associated with the Vignette for GWAS.BAYES.

## Usage

data("RealDataSNPs_Y")

## Format

A data frame with 328 observations on the following 1501 variables.
Phenotype a numeric vector
SNP1 a character vector
SNP2 a character vector
SNP3 a character vector
SNP4 a character vector

SNP5 a character vector
SNP6 a character vector
SNP7 a character vector
SNP8 a character vector
SNP9 a character vector
SNP10 a character vector
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## Examples

```
data(RealDataSNPs_Y)
```

```
resids_diag resids_diag
```


## Description

Tests residuals to see if they are normal. This looks at the model with all significant SNPs from the preselection phase.

## Usage

resids_diag(Y,SNPs,significant,kinship = FALSE, principal_components = FALSE, plot_it = TRUE)

## Arguments

| Y | The phenotype response of interest |
| :--- | :--- |
| SNPs | Standardized SNP data set where the values of each column are either 0 or 1 |
| significant | A vector of 0's and 1's where the 1's indicate a significant SNP. This is returned <br> in the output of the preselection function. |
| kinship | A kinship matrix, can be calculated from the rrBLUP package. |
| principal_components |  |
| A matrix or vector of the principal components one would like to include in the |  |
| analysis. |  |

## Value

value 1 The output of a Shapiro-Wilk test for the residuals. If the p-value is above .05 , there is no evidence that the residuals are not normal. If the p-value is below . 05 there is evidence that the residuals are not normal, and some transformation is suggested.
value 2 A histogram of the residuals when plot_it = TRUE

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
fullPreprocess <- preprocess_SNPs(SNPs = SNPs,Y = Y,MAF = 0.01, number_cores = 1)
SNPs <- fullPreprocess$SNPs
Y <- fullPreprocess$Y
fullPreprocess$SNPs_Dropped
principal_comp <- pca_function(SNPs = SNPs,number_components = 1,plot_it = FALSE)
Significant_SNPs <- preselection(Y = Y, SNPs = SNPs,number_cores = 1, principal_components = principal_comp,fr
resids_diag(Y = Y,SNPs = SNPs,significant = Significant_SNPs$Significant,kinship = FALSE,principal_components
```

```
SNP_data_function_nopcp
```

    SNP_data_function_nopср
    
## Description

This is used internally in the preselection function to sort the SNPs into datasets

## Usage

SNP_data_function_nopcp(x, int)

## Arguments

x
The SNP of interest.
int The intercept.

## Value

Returns a dataframe combining all three entries using cbind.

```
SNP_data_function_pcp SNP_data_function_pcp
```


## Description

This is used internally in the preselection function to sort the SNPs into datasets

## Usage

SNP_data_function_pcp(x, pcp,int)

## Arguments

$x \quad$ The SNP of interest.
pcp The principal components.
int The intercept.

## Value

Returns a dataframe combining all three entries using cbind.

```
standardize standardize
```


## Description

Standardize the SNPs to the $0-1$ scale

## Usage

standardize(SNPs,method=c("major-minor", "alphabetical"), number_cores)

## Arguments

SNPs The SNP dataset with columns of the values A, C, T, and G
method The method in which to standarize. If "major-minor" is selected then the major allele gets the value 1 and the minor allele gets the value 0 . If "alphabetical" is selected the first allele alphabetically gets a value of 0 and the second allele alphabetically gets the value of 1 .
number_cores The number of cores on which to parallize over.

## Value

Returns a matrix of SNPs with the same dimension as th input, the columns are now defined as 0 and 1's.

## Examples

```
data("vignette_lm_dat")
Y <- vignette_lm_dat$Phenotype
SNPs <- vignette_lm_dat[,-1]
standardize(SNPs = SNPs,method = "major-minor",number_cores = 1)
```

```
vignette_kinship_dat vignette_kinship_dat
```


## Description

Dataset associated with the Vignette for the GWAS.BAYES package.

## Usage

```
    data("vignette_kinship_dat")
```


## Format

A data frame with 4075 observations on the following 1001 variables.
Phenotype a numeric vector
SNP1 a factor with levels G T
SNP2 a factor with levels C T
SNP3 a factor with levels C T
SNP4 a factor with levels C T
SNP5 a factor with levels A C
SNP6 a factor with levels C T
SNP7 a factor with levels C T
SNP8 a factor with levels A C
SNP9 a factor with levels A G
SNP10 a factor with levels C G
SNP11 a factor with levels A T
SNP12 a factor with levels A T
SNP13 a factor with levels C T
SNP14 a factor with levels A G
SNP15 a factor with levels A C
SNP16 a factor with levels C T
SNP17 a factor with levels C T
SNP18 a factor with levels C T
SNP19 a factor with levels A T
SNP20 a factor with levels C T
SNP21 a factor with levels A G

SNP22 a factor with levels C T
SNP23 a factor with levels C T
SNP24 a factor with levels A T
SNP25 a factor with levels C G
SNP26 a factor with levels A T
SNP27 a factor with levels C T
SNP28 a factor with levels C G
SNP29 a factor with levels G T
SNP30 a factor with levels C G
SNP31 a factor with levels G T
SNP32 a factor with levels A G
SNP33 a factor with levels A G
SNP34 a factor with levels C G
SNP35 a factor with levels A C
SNP36 a factor with levels A G
SNP37 a factor with levels A C
SNP38 a factor with levels A G
SNP39 a factor with levels A T
SNP40 a factor with levels C G
SNP41 a factor with levels A G
SNP42 a factor with levels C G
SNP43 a factor with levels G T
SNP44 a factor with levels A C
SNP45 a factor with levels C T
SNP46 a factor with levels C G
SNP47 a factor with levels G T
SNP48 a factor with levels A C
SNP49 a factor with levels G T
SNP50 a factor with levels A T
SNP51 a factor with levels A T
SNP52 a factor with levels C G
SNP53 a factor with levels G T
SNP54 a factor with levels C T
SNP55 a factor with levels C T
SNP56 a factor with levels C G
SNP57 a factor with levels C T
SNP58 a factor with levels A C
SNP59 a factor with levels A G
SNP60 a factor with levels A T
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SNP62 a factor with levels A T
SNP63 a factor with levels G T
SNP64 a factor with levels A T
SNP65 a factor with levels A T
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SNP72 a factor with levels A G
SNP73 a factor with levels C T
SNP74 a factor with levels A G
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SNP76 a factor with levels G T
SNP77 a factor with levels G T
SNP78 a factor with levels G T
SNP79 a factor with levels A T
SNP80 a factor with levels A C
SNP81 a factor with levels G T
SNP82 a factor with levels A C
SNP83 a factor with levels G T
SNP84 a factor with levels C G
SNP85 a factor with levels C G
SNP86 a factor with levels C G
SNP87 a factor with levels C T
SNP88 a factor with levels A G
SNP89 a factor with levels A T
SNP90 a factor with levels G T
SNP91 a factor with levels C T
SNP92 a factor with levels G T
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SNP748 a factor with levels A C
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SNP773 a factor with levels A G
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SNP775 a factor with levels A G
SNP776 a factor with levels A G
SNP777 a factor with levels A C
SNP778 a factor with levels C G
SNP779 a factor with levels A C
SNP780 a factor with levels A T
SNP781 a factor with levels C G

SNP782 a factor with levels A T
SNP783 a factor with levels A C
SNP784 a factor with levels A C
SNP785 a factor with levels C T
SNP786 a factor with levels C G
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SNP791 a factor with levels A T
SNP792 a factor with levels A G
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SNP794 a factor with levels A T
SNP795 a factor with levels C G
SNP796 a factor with levels G T
SNP797 a factor with levels C T
SNP798 a factor with levels A T
SNP799 a factor with levels G T
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SNP801 a factor with levels G T
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SNP803 a factor with levels A C
SNP804 a factor with levels C T
SNP805 a factor with levels A G
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SNP809 a factor with levels C T
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SNP816 a factor with levels G T
SNP817 a factor with levels G T
SNP818 a factor with levels C G
SNP819 a factor with levels A G
SNP820 a factor with levels G T
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SNP849 a factor with levels A C
SNP850 a factor with levels C G
SNP851 a factor with levels A T
SNP852 a factor with levels G T
SNP853 a factor with levels A T
SNP854 a factor with levels A C
SNP855 a factor with levels A T
SNP856 a factor with levels A T
SNP857 a factor with levels A G
SNP858 a factor with levels C G
SNP859 a factor with levels A C
SNP860 a factor with levels A T
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SNP862 a factor with levels C G
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SNP889 a factor with levels A G
SNP890 a factor with levels G T
SNP891 a factor with levels A C
SNP892 a factor with levels C G
SNP893 a factor with levels A G
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SNP895 a factor with levels A G
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SNP897 a factor with levels C G
SNP898 a factor with levels C G
SNP899 a factor with levels C T
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SNP901 a factor with levels A G

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SNP903 a factor with levels A G
SNP904 a factor with levels G T
SNP905 a factor with levels A T
SNP906 a factor with levels A T
SNP907 a factor with levels A T
SNP908 a factor with levels A C
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SNP910 a factor with levels C G
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SNP915 a factor with levels C G
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SNP922 a factor with levels A C
SNP923 a factor with levels A C
SNP924 a factor with levels A C
SNP925 a factor with levels C T
SNP926 a factor with levels G T
SNP927 a factor with levels G T
SNP928 a factor with levels A C
SNP929 a factor with levels A T
SNP930 a factor with levels C G
SNP931 a factor with levels A C
SNP932 a factor with levels A C
SNP933 a factor with levels C G
SNP934 a factor with levels A G
SNP935 a factor with levels C T
SNP936 a factor with levels A C
SNP937 a factor with levels A T
SNP938 a factor with levels C G
SNP939 a factor with levels C G
SNP940 a factor with levels A C
SNP941 a factor with levels A C

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SNP967 a factor with levels C T
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SNP988 a factor with levels G T
SNP989 a factor with levels G T
SNP990 a factor with levels C T
SNP991 a factor with levels A G
SNP992 a factor with levels G T
SNP993 a factor with levels C G
SNP994 a factor with levels A G
SNP995 a factor with levels A C
SNP996 a factor with levels C G
SNP997 a factor with levels A G
SNP998 a factor with levels A T
SNP999 a factor with levels A T
SNP1000 a factor with levels A C

## Examples

```
data(vignette_kinship_dat)
```

```
vignette_lm_dat vignette_lm_dat
```


## Description

Dataset associated with the Vignette for the GWAS.BAYES package.

## Usage

data("vignette_lm_dat")

## Format

A data frame with 4075 observations on the following 1001 variables.
Phenotype a numeric vector
SNP1 a factor with levels A G
SNP2 a factor with levels G T
SNP3 a factor with levels A G
SNP4 a factor with levels A T
SNP5 a factor with levels A T

SNP6 a factor with levels C T
SNP7 a factor with levels C G
SNP8 a factor with levels A C
SNP9 a factor with levels C G
SNP10 a factor with levels G T
SNP11 a factor with levels G T
SNP12 a factor with levels G T
SNP13 a factor with levels A G
SNP14 a factor with levels C T
SNP15 a factor with levels G T
SNP16 a factor with levels A C
SNP17 a factor with levels G T
SNP18 a factor with levels A T
SNP19 a factor with levels C T
SNP20 a factor with levels A C
SNP21 a factor with levels G T
SNP22 a factor with levels A T
SNP23 a factor with levels A C
SNP24 a factor with levels A T
SNP25 a factor with levels A C
SNP26 a factor with levels C T
SNP27 a factor with levels G T
SNP28 a factor with levels A T
SNP29 a factor with levels A T
SNP30 a factor with levels A G
SNP31 a factor with levels A T
SNP32 a factor with levels C T
SNP33 a factor with levels A G
SNP34 a factor with levels A C
SNP35 a factor with levels A G
SNP36 a factor with levels C G
SNP37 a factor with levels A T
SNP38 a factor with levels C G
SNP39 a factor with levels A G
SNP40 a factor with levels A G
SNP41 a factor with levels A C
SNP42 a factor with levels A G
SNP43 a factor with levels A T
SNP44 a factor with levels A G
SNP45 a factor with levels C G

SNP46 a factor with levels C T
SNP47 a factor with levels A T
SNP48 a factor with levels C T
SNP49 a factor with levels A G
SNP50 a factor with levels A T
SNP51 a factor with levels A C
SNP52 a factor with levels G T
SNP53 a factor with levels G T
SNP54 a factor with levels A C
SNP55 a factor with levels A C
SNP56 a factor with levels A C
SNP57 a factor with levels G T
SNP58 a factor with levels G T
SNP59 a factor with levels A G
SNP60 a factor with levels C T
SNP61 a factor with levels A C
SNP62 a factor with levels A T
SNP63 a factor with levels C T
SNP64 a factor with levels C T
SNP65 a factor with levels A T
SNP66 a factor with levels C T
SNP67 a factor with levels G T
SNP68 a factor with levels C T
SNP69 a factor with levels C T
SNP70 a factor with levels A T
SNP71 a factor with levels C G
SNP72 a factor with levels A T
SNP73 a factor with levels A C
SNP74 a factor with levels C T
SNP75 a factor with levels A G
SNP76 a factor with levels A G
SNP77 a factor with levels A T
SNP78 a factor with levels A C
SNP79 a factor with levels A C
SNP80 a factor with levels G T
SNP81 a factor with levels G T
SNP82 a factor with levels A C
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SNP90 a factor with levels G T
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SNP93 a factor with levels A G
SNP94 a factor with levels C T
SNP95 a factor with levels A G
SNP96 a factor with levels A C
SNP97 a factor with levels A C
SNP98 a factor with levels C G
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SNP100 a factor with levels A C
SNP101 a factor with levels A G
SNP102 a factor with levels A T
SNP103 a factor with levels A T
SNP104 a factor with levels G T
SNP105 a factor with levels A C
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SNP107 a factor with levels A G
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SNP179 a factor with levels C G
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SNP181 a factor with levels C T
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SNP229 a factor with levels G T
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SNP326 a factor with levels A C
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SNP516 a factor with levels A G
SNP517 a factor with levels G T
SNP518 a factor with levels A T
SNP519 a factor with levels C G
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SNP521 a factor with levels A C
SNP522 a factor with levels A C
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SNP524 a factor with levels A G
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SNP526 a factor with levels A C
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## Examples

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