

Package ‘PAST’

October 18, 2022

Type Package

Title Pathway Association Study Tool (PAST)

Version 1.12.0

Description PAST takes GWAS output and assigns SNPs to genes, uses those genes to find pathways associated with the genes, and plots pathways based on significance. Implements methods for reading GWAS input data, finding genes associated with SNPs, calculating enrichment score and significance of pathways, and plotting pathways.

License GPL (>=3) + file LICENSE

Encoding UTF-8

Depends R (>= 4.0)

Imports stats, utils, dplyr, rlang, iterators, parallel, foreach,
doParallel, qvalue, rtracklayer, ggplot2, GenomicRanges,
S4Vectors

Suggests knitr, rmarkdown

VignetteBuilder knitr

RoxygenNote 7.1.0

URL <https://github.com/IGBB/past>

BugReports <https://github.com/IGBB/past/issues>

biocViews Pathways, GeneSetEnrichment

git_url <https://git.bioconductor.org/packages/PAST>

git_branch RELEASE_3_15

git_last_commit 4659dad

git_last_commit_date 2022-04-26

Date/Publication 2022-10-18

Author Thrash Adam [cre, aut],
DeOrnellis Mason [aut]

Maintainer Thrash Adam <thrash@igbb.msstate.edu>

R topics documented:

assign_chunk	2
assign_SNPs_to_genes	2
determine_linkage	3
find_pathway_significance	4
findRepresentativeSNP	5
findRepresentativeSNP_gene_pairing	5
load_GWAS_data	6
load_LD	7
plot_pathways	7

Index	9
--------------	----------

assign_chunk	<i>Assign SNPs in a chunk to genes</i>
--------------	--

Description

Assign SNPs in a chunk to genes

Usage

```
assign_chunk(gff, chunk, window)
```

Arguments

gff	The GFF data for the chromosome being parsed
chunk	The dataframe containing SNP data
window	The search window around the SNPs

Value

tagSNPs labeled with gene names

assign_SNPs_to_genes	<i>Assign SNPs to genes</i>
----------------------	-----------------------------

Description

Assign SNPs to genes

Usage

```
assign_SNPs_to_genes(
  gwas_data,
  LD,
  gff_file,
  filter_type,
  window,
  r_squared_cutoff,
  num_cores
)
```

Arguments

gwas_data	Merged association and effects data from merge_data()
LD	Linkage disequilibrium data from parse_LD()
gff_file	The path to a GFF file
window	The search window for genes around the SNP
r_squared_cutoff	The R^2 value used to determine SNP significance
num_cores	The number of cores to use in parallelizing PAST

Value

A dataframe of genes from the SNP data

Examples

```
example("load_GWAS_data")
example("load_LD")
demo_genes_file = system.file("extdata", "genes.gff",
  package = "PAST", mustWork = TRUE)
filter_type = c("gene")
genes <- assign_SNPs_to_genes(gwas_data, LD, demo_genes_file, filter_type, 1000, 0.8, 2)
```

determine_linkage	<i>Determine Linkage</i>
-------------------	--------------------------

Description

Determine Linkage

Usage

```
determine_linkage(chunk, r_squared_cutoff)
```

Arguments

`chunk` A chunk of data to be processed
`r_squared_cutoff` The R² value to check against

Value

Either the first unlinked SNP or a set of linked SNPs

find_pathway_significance
Find Pathway Significance

Description

Find Pathway Significance

Usage

```
find_pathway_significance(  
  genes,  
  pathways_file,  
  gene_number_cutoff = 5,  
  mode,  
  sample_size = 1000,  
  num_cores  
)
```

Arguments

`genes` Genes from `assign_SNPs_to_genes()`
`pathways_file` A file containing the pathway IDs, their names, and the genes in the pathway
`gene_number_cutoff` A cut-off for the minimum number of genes in a pathway
`mode` increasing/decreasing
`sample_size` How many times to sample the effects data during random sampling
`num_cores` The number of cores to use in parallelizing PAST

Value

Rugplots data

Examples

```
example("assign_SNPs_to_genes")
demo_pathways_file = system.file("extdata", "pathways.txt.xz",
  package = "PAST", mustWork = TRUE)
rugplots_data <- find_pathway_significance(genes, demo_pathways_file, 5,
  "increasing", 1000, 2)
```

findRepresentativeSNP

Find representative SNP for a chunk of SNPs

Description

Find representative SNP for a chunk of SNPs

Usage

```
findRepresentativeSNP(chunk, r_squared_cutoff)
```

Arguments

chunk	A chunk of data to parse
r_squared_cutoff	The R ² value to check against when counting SNPs

Value

A single SNP representing the whole chunk

findRepresentativeSNP_gene_pairing

Find the SNP-gene assignment that represents SNPs assigned to a gene

Description

Find the SNP-gene assignment that represents SNPs assigned to a gene

Usage

```
findRepresentativeSNP_gene_pairing(chunk)
```

Arguments

chunk	A chunk of gene assignments
-------	-----------------------------

Value

A single SNP-gene assignment representing all SNPs assigned to the same gene to a gene

load_GWAS_data	<i>Load GWAS data</i>
----------------	-----------------------

Description

Load GWAS data

Usage

```
load_GWAS_data(
  association_file,
  effects_file,
  association_columns = c("Trait", "Marker", "Locus", "Site", "p", "marker_R2"),
  effects_columns = c("Trait", "Marker", "Locus", "Site", "Effect")
)
```

Arguments

association_file	The association file
effects_file	The effects file
association_columns	The names of the columns in your association data for Trait, Marker, Chromosome, Site, F, p, and marker_Rsquared
effects_columns	The names of the columns in your effects data for Trait, Marker, Chromosome, Site, and effect

Value

The association data and the effects data merged into a dataframe with one row for each SNP

Examples

```
demo_association_file = system.file("extdata", "association.txt.xz",
  package = "PAST", mustWork = TRUE)
demo_effects_file = system.file("extdata", "effects.txt.xz",
  package = "PAST", mustWork = TRUE)
gwas_data <- load_GWAS_data(demo_association_file, demo_effects_file)
```

load_LD

Load Linkage Disequilibrium

Description

Load Linkage Disequilibrium

Usage

```
load_LD(  
  LD_file,  
  LD_columns = c("Locus1", "Position1", "Site1", "Position2", "Site2", "Dist_bp",  
  "R.2")  
)
```

Arguments

LD_file	The file containing linkage disequilibrium data
LD_columns	The names of the columns in your linkage disequilibrium data for the chromosome of the first SNP, the position of the first SNP, the site of the first SNP, the chromosome of the second SNP, the position of the second SNP, the site of the second SNP, the distance between the two SNPs, and the R.2

Value

The linkage disequilibrium data in a list containing dataframes for each chromosome.

Examples

```
demo_LD_file = system.file("extdata", "LD.txt.xz",  
  package = "PAST", mustWork = TRUE)  
LD <- load_LD(demo_LD_file)
```

plot_pathways

Plot Rugplots for Selected Pathways

Description

Plot Rugplots for Selected Pathways

Usage

```
plot_pathways(  
  rugplots_data,  
  filter_type,  
  filter_parameter,  
  mode,  
  output_directory  
)
```

Arguments

rugplots_data The data to be plotted (returned from `find_pathway_significance()`)
filter_type The parameter to be used for filtering
filter_parameter The cut-off value of the filtering parameter
mode The mode used to create the data (increasing/decreasing)
output_directory An existing directory to save results in

Value

Does not return a value

Examples

```
example("find_pathway_significance")  
plot_pathways(rugplots_data, "pvalue", "0.03", "decreasing", tempdir())
```

Index

assign_chunk, 2
assign_SNPs_to_genes, 2

determine_linkage, 3

find_pathway_significance, 4
findRepresentativeSNP, 5
findRepresentativeSNP_gene_pairing,
 5

load_GWAS_data, 6
load_LD, 7

plot_pathways, 7