Package 'GeneticMediation'

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Type Package

Title An R Package for Causal Mediation Analysis of ROSMAP Data
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https://tydarnell.github.io/GeneticMediation
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Description Provides methods for cleaning and processing data from the ROSMAP study and preparing it for causal mediation analysis
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R topics documented:
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check_data

Check Data

Description

Checks if the data is in the data folder

Usage

check_data()

chr_numeric

Chromosome Numeric

Description

Convert chromosome names from character to numeric. Useful when sorting a dataframe by chromosome number.

Usage

chr_numeric(Chr)

Arguments

Chr

a character column or vector of chromosome names

clean_data 3

clean_data

Clean Data

Description

Clean snp.info, peak.info data and save as data/chipseq.RData

Usage

```
clean_data(snp.path, peak.path)
```

Arguments

snp.path path to SNP information dataframe peak.path path to Peak annotation dataframe

combine_data_spc

Combine Data SNP-Peak-Clinical

Description

SNP-Peak-Clinical data: combine projid, SNPs, PCs, membership, age, gender, peaks, and outcome data for a chromosome

Usage

```
combine_data_spc(chr)
```

Arguments

chr

chromosome name

combine_data_spg

Combine Data SNP-Peak-Gene

Description

SNP-Peak-Gene Data: combine projid, SNPs, PCs, membership, age, gender, peaks, gene and outcome data for a chromosome.

Usage

```
combine_data_spg(chr)
```

Arguments

chr

chromosome name

make_folder

last_to_first

Last to First

Description

Make the last column the first column in a dataframe

Usage

```
last_to_first(df)
```

Arguments

df

dataframe

lu

Length Unique

Description

Get the length of unique values in a vector

Usage

lu(x)

Arguments

Х

vector

make_folder

Make Folder

Description

Make a folder only if the folder does not already exist

Usage

```
make_folder(path)
```

Arguments

path

folder path

match_all 5

match_all Match	SNPs Peaks All Chromosomes
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Description

Match SNPs and Peaks in all chromosomes and return dataframe of matches

Usage

```
match_all(snp.info, peak.info, chrs)
```

Arguments

snp.info SNP information peak.info Peak information

chrs character vector of chromosome names

Description

Match SNPs and Peaks in a chromosome

Usage

```
match_snp_peak(snp.info, peak.info)
```

Arguments

snp.info SNP information for chr peak.info Peak information chr

med.res_spc SNP-Peak-Clinical Mediation Data Prep

Description

Prepare a chromosome of SNP-Peak-Clinical data for mediation analysis and save as "data/spc.res/res_chr.RData"

Usage

```
med.res_spc(chr, all_matches)
```

Arguments

chr chromosome name

all_matches dataframe of SNP-Peak matches for all chromosomes

6 med_all_spc

med.res_spg SNP-Peak-Gene Mediation Data Prep

Description

Prepare a chromosome SNP-Peak-Gene data for mediation analysis data and save as "data/spg.res/res_chr.RData"

Usage

```
med.res_spg(chr, med.data, matches, mediator.path)
```

Arguments

chr chromosome name
med.data gene mediation data

matches SNP-Peak-Gene matches dataframe

med_all_spc SNP-Peak-Clinical Mediation Table

Description

Create a table of SNP-Peak-Clinical mediation results for all chromosomes

Usage

```
med_all_spc(chrs, covar, simulations)
```

Arguments

chrs character vector of chromosome names

covar covariates string, each covariate separated by +

simulations number of simulations to run

med_chr_spc 7

med_chr_spc SNP-Peak-Clinical Mediation Table Chromosome	med_chr_spc	SNP-Peak-Clinical Mediation Table Chromosome
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Description

Create SNP-Peak-Clinical mediation table for SNP-Peak matches in a chromosome

Usage

```
med_chr_spc(matches, med.res.data, covar, simulations)
```

Arguments

matches SNP-Peak matches dataframe for a chromosome

med.res.data mediation data

covar covariates string, separate covariates with +

simulations number of simulations to run

med_chr_spg SNP-Peak-Gene Mediation Table Chromosome

Description

Create a SNP-Peak-Gene mediation table for a chromosome

Usage

```
med_chr_spg(chr, gene_matches, covar, simulations)
```

Arguments

chr chromosome name

gene_matches list of peak-SNP matches for each gene

covar covariates string, separate covariates with +

simulations number of simulations to run

8 med_table_spc

Description

Mediation summary for one SNP-Peak-Gene match

Usage

```
med_summary_spg(med.res.data, match_row, simulations = 1000)
```

Arguments

med.res.data gene mediation data

match_row row in match dataframe: col1 SNP, col2 Peak, col3 Gene

simulations number of simulations, 1000 by default

med_table_spc SNP-Peak-Clinical Mediation Table

Description

Create SNP-Peak-Clinical mediation table for a SNP-Peak match

Usage

```
med_table_spc(med.res.data, match_row, covar, simulations)
```

Arguments

med.res.data mediation data

match_row row in match dataframe, should have 2 columns

covar covariates character variable, separate covariates with +

simulations number of simulations to run

med_table_spg 9

med_table_spg

SNP-Peak-Gene Mediation Table

Description

Create a mediation table for one SNP-Peak-Gene match

Usage

```
med_table_spg(med.res.data, match_row, covar, simulations)
```

Arguments

med.res.data gene mediation data

match_row row in match dataframe: col1 SNP, col2 Peak, col3 Gene covar covariates character variable, separate covariates with +

simulations number of simulations to run

merge_csv

Merge CSV

Description

merge multiple csv files into a single dataframe

Usage

```
merge_csv(path)
```

Arguments

path to folder containing csv files

 ${\tt setup_folders}$

Setup Folders

Description

Creates data, data/mediator, results folders

Usage

```
setup_folders()
```

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snp_peak_bychr

SNPs and Peaks by Chromosome

Description

save snp.info, peak.info for each chromosome as "data/chipseq/chipseq_chr.RData"

Usage

```
snp_peak_bychr(chrs, snps, peaks)
```

Arguments

chrs Character vector of chromosomes names

snps SNP information dataframe peaks Peak information dataframe

sum_zero

Sum Zero

Description

Get names of columns that have a sum of zero

Usage

```
sum\_zero(df)
```

Arguments

df

dataframe

transpose_readcount

Transpose Readcount

Description

Read in and transpose ChIP-seq Readcount dataframe and match project id

Usage

```
transpose_readcount(readcount.path)
```

Arguments

readcount.path ChIP-seq readcount dataframe file path

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