Package 'GxEprsDummy'

May 29, 2023

Title What the Package Does (One Line, Title Case)

Version 0.0.0.9000

Index

Description What the package does (one paragraph).
License `use_mit_license()`, `use_gpl3_license()` or friends to pick a license
Encoding UTF-8
Roxygen list(markdown = TRUE)
RoxygenNote 7.2.3
Depends R (>= 2.10)
LazyData true
R topics documented:
Bcov_discovery
Bcov_target
Bphe_discovery
Bphe_target
DummyData
GWAS_binary
GWEIS_binary
PRS_binary
Qcov_discovery
Qcov_target
Qphe_discovery

9

13

2 Bcov_target

Bcov_discovery	Covariate data file of the discovery dataset when the outcome is binary This contains covariate information of the individuals in the discovery dataset following confounders
	unusci jonoving conjounucis

Description

Covariate data file of the discovery dataset when the outcome is binary This contains covariate information of the individuals in the discovery dataset following confounders

Usage

Bcov_discovery

Format

A dataframe with 7916 rows and 18 columns

Family ID \item Individual ID \item standardized covariate \item square of the standardized covariate \item confounder 1 \item confounder 2 \item confounder 3 \item confounder 4 \item confounder 5 \item confounder 6 \item confounder 7 \item confounder 8 \item confounder 9 \item confounder 10 \item confounder 11 \item confounder 12 \item confounder 13 \item confounder 14

Bcov_target	Covariate data file of the target dataset when the outcome is binary This contains covariate information of the individuals in the target dataset following confounders
	,

Description

Covariate data file of the target dataset when the outcome is binary This contains covariate information of the individuals in the target dataset following confounders

Usage

Bcov_target

Format

A dataframe with 1939 rows and 18 columns

Family ID \item Individual ID \item standardized covariate \item square of the standardized covariate \item confounder 1 \item confounder 2 \item confounder 3 \item confounder 4 \item confounder 5 \item confounder 6 \item confounder 7 \item confounder 8 \item confounder 9 \item confounder 10 \item confounder 11 \item confounder 12 \item confounder 13 \item confounder 14

Bphe_discovery 3

Bphe_discovery	Phenotype data file of the discovery dataset when the outcome is binary This contains phenotype information of the individuals in the discovery dataset
	covery dataset

Description

Phenotype data file of the discovery dataset when the outcome is binary This contains phenotype information of the individuals in the discovery dataset

Usage

```
Bphe_discovery
```

Format

A dataframe with 7916 rows and 3 columns

Family ID \item Individual ID \item phenotype (1=controls, 2=cases)

Bphe_target Phenotype data file of the target dataset when the outcome is This contains phenotype information of the individuals in the dataset	
---	--

Description

Phenotype data file of the target dataset when the outcome is binary This contains phenotype information of the individuals in the target dataset

Usage

```
Bphe_target
```

Format

A dataframe with 1939 rows and 3 columns

Family ID \item Individual ID \item phenotype (0=controls, 1=cases)

GWAS_binary

, and the second	DummyData	PLINK binary data files This contains DummyData.fam, DummyData.bim and DummyData.bed files, all in one
--	-----------	--

Description

PLINK binary data files This contains DummyData.fam, DummyData.bim and DummyData.bed files, all in one

Usage

DummyData

Format

The data files follow the general PLINK format

GWAS_binary	GWAS_binary function This function performs GWAS using plink2 and outputs the GWAS summary statistics file with additive SNP effects named B_trd.sum

Description

GWAS_binary function This function performs GWAS using plink2 and outputs the GWAS summary statistics file with additive SNP effects named $B_{\rm trd.sum}$

Usage

```
GWAS_binary(
  plink_path,
  b_file,
  pheno_file,
  covar_file,
  n_confounders,
  thread,
  summary_output = NULL
)
```

Arguments

plink_path	Path to the PLINK executable application
b_file	Prefix of the binary files, where all .fam, .bed and .bim files have a common prefix
pheno_file	Name (with file extension) of the phenotype file containing family ID, individual ID and phenotype of the discovery dataset as columns, without heading
covar_file	Name (with file extension) of the covariate file containing family ID, individual ID, standardized covariate, square of standardized covariate, and/or confounders of the discovery dataset as columns, without heading

GWEIS_binary 5

```
n_confounders
```

Number of confounding variables in the discovery dataset

thread Number of threads used

Value

This function will perform GWAS and output

```
B_trd.sum
```

GWEIS_binary

GWEIS_binary function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics files with additive SNP effects named B_add.sum and interaction SNP effects named B_gxe.sum

Description

GWEIS_binary function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics files with additive SNP effects named $B_{add.sum}$ and interaction SNP effects named $B_{add.sum}$

Usage

```
GWEIS_binary(
  plink_path,
  b_file,
  pheno_file,
  covar_file,
  n_confounders,
  thread,
  summary_output1 = NULL,
  summary_output2 = NULL)
```

Arguments

plink_path	Path to the PLINK executable application
b_file	Prefix of the binary files, where all .fam, .bed and .bim files have a common prefix
pheno_file	Phenotype file containing family ID, individual ID and phenotype of the discovery dataset as columns, without heading
covar_file	Covariate file containing family ID, individual ID, standardized covariate, square of standardized covariate, and/or confounders of the discovery dataset as columns, without heading
n_confounder	s

Number of confounding variables in the discovery dataset

thread Number of threads used

6 PRS_binary

Value

This function will perform GWEIS and output

```
B_add.sum B_gxe.sum
```

PRS_binary

PRS_binary function This function uses plink2 and outputs PRSs of each individual in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Description

PRS_binary function This function uses plink2 and outputs PRSs of each individual in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Usage

```
PRS_binary(
  plink_path,
  b_file,
  summary_input1 = "B_trd.sum",
  summary_input2 = "B_add.sum",
  summary_input3 = "B_gxe.sum",
  summary_output1 = NULL,
  summary_output2 = NULL,
  summary_output3 = NULL
)
```

Arguments

plink_path Path to the PLINK executable application
b_file Prefix of the binary files, where all .fam, .bed and .bim files have a common prefix

Value

This function will output

```
B_trd.sscore
B_add.sscore
B_gxe.sscore
```

Qcov_discovery 7

Qcov_discovery	Covariate data file of the discovery dataset when the outcome is quantitative This contains covariate information of the individuals in the
	discovery dataset following confounders

Description

Covariate data file of the discovery dataset when the outcome is quantitative This contains covariate information of the individuals in the discovery dataset following confounders

Usage

Qcov_discovery

Format

A dataframe with 6426 rows and 18 columns

Family ID \item Individual ID \item standardized covariate \item square of the standardized covariate \item confounder 1 \item confounder 2 \item confounder 3 \item confounder 4 \item confounder 5 \item confounder 6 \item confounder 7 \item confounder 8 \item confounder 9 \item confounder 10 \item confounder 11 \item confounder 12 \item confounder 13 \item confounder 14

Qcov_target	Covariate data file of the target dataset when the outcome is quantita- tive This contains covariate information of the individuals in the target
	dataset following confounders

Description

Covariate data file of the target dataset when the outcome is quantitative This contains covariate information of the individuals in the target dataset following confounders

Usage

Qcov_target

Format

A dataframe with 1579 rows and 18 columns

Family ID \item Individual ID \item standardized covariate \item square of the standardized covariate \item confounder 1 \item confounder 2 \item confounder 3 \item confounder 4 \item confounder 5 \item confounder 6 \item confounder 7 \item confounder 8 \item confounder 9 \item confounder 10 \item confounder 11 \item confounder 12 \item confounder 13 \item confounder 14

Qphe_target

Qphe_discovery Phenotype data file of the discovery dataset when the or titative This contains phenotype information of the ind discovery dataset	*
---	---

Description

Phenotype data file of the discovery dataset when the outcome is quantitative This contains phenotype information of the individuals in the discovery dataset

Usage

```
Qphe_discovery
```

Format

A dataframe with 6426 rows and 3 columns Family ID \item Individual ID \item phenotype

Qphe_target	Phenotype data file of the target dataset when the outcome is quantitative This contains phenotype information of the individuals in the target dataset
	8

Description

Phenotype data file of the target dataset when the outcome is quantitative This contains phenotype information of the individuals in the target dataset

Usage

```
Qphe_target
```

Format

A dataframe with 1579 rows and 3 columns
Family ID \item Individual ID \item phenotype

```
results_permuted_binary
```

results_permuted_binary function This function uses plink2 and outputs PRSs of each individual in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Description

results_permuted_binary function This function uses plink2 and outputs PRSs of each individual in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Usage

```
results_permuted_binary(
   Bphe_target,
   Bcov_target,
   n_confounders,
   input_score1 = "B_trd.sscore",
   input_score2 = "B_add.sscore",
   input_score3 = "B_gxe.sscore"
)
```

Arguments

Bphe_target Phenotype file containing family ID, individual ID and phenotype of the target

dataset as columns, without heading

 ${\tt Bcov_target} \quad Covariate \ file \ containing \ family \ ID, individual \ ID, standardized \ covariate, square$

of standardized covariate, and/or confounders of the target dataset as columns, without heading

without headii

Number of confounding variables in the target dataset

Value

This function will output

n_confounders

```
Individual_risk_values.txt
```

```
results_regular_binary
```

results_regular_binary function This function uses plink2 and outputs PRSs of each individual in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Description

results_regular_binary function This function uses plink2 and outputs PRSs of each individual in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Usage

```
results_regular_binary(
   Bphe_target,
   Bcov_target,
   n_confounders,
   input_score1 = "B_trd.sscore",
   input_score2 = "B_add.sscore",
   input_score3 = "B_gxe.sscore")
```

Arguments

Bphe_target Phenotype file containing family ID, individual ID and phenotype of the target dataset as columns, without heading

Bcov_target Covariate file containing family ID, individual ID, standardized covariate, square of standardized covariate, and/or confounders of the target dataset as columns, without heading

n confounders

Number of confounding variables in the target dataset

Value

This function will output

```
Individual_risk_values.txt
```

```
summary_permuted_binary
```

summary_permuted_binary function This function uses plink2 and outputs the p value of permuted model in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Description

summary_permuted_binary function This function uses plink2 and outputs the p value of permuted model in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Usage

```
summary_permuted_binary(
   Bphe_target,
   Bcov_target,
   n_confounders,
   iterations = 1000,
   input_score1 = "B_trd.sscore",
   input_score2 = "B_add.sscore",
   input_score3 = "B_gxe.sscore")
```

Arguments

Bphe_target Phenotype file containing family ID, individual ID and phenotype of the target dataset as columns, without heading

Bcov_target Covariate file containing family ID, individual ID, standardized covariate, square of standardized covariate, and/or confounders of the target dataset as columns, without heading

n_confounders

Number of confounding variables in the target dataset

Value

This function will output

```
B_permuted_p.txt
```

```
summary_regular_binary
```

summary_regular_binary function This function uses plink2 and outputs the summary of regular model in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B trd.sum, B add.sum and B gxe.sum

Description

summary_regular_binary function This function uses plink2 and outputs the summary of regular model in the target dataset, using pre-generated GWAS and GWEIS summary statistics files named B_trd.sum, B_add.sum and B_gxe.sum

Usage

```
summary_regular_binary(
   Bphe_target,
   Bcov_target,
   n_confounders,
   input_score1 = "B_trd.sscore",
   input_score2 = "B_add.sscore",
   input_score3 = "B_gxe.sscore")
```

Arguments

Bphe_target Phenotype file containing family ID, individual ID and phenotype of the target

dataset as columns, without heading

 ${\tt Bcov_target} \quad Covariate file \ containing \ family \ ID, individual \ ID, standardized \ covariate, square$

of standardized covariate, and/or confounders of the target dataset as columns,

without heading

n_confounders

Number of confounding variables in the target dataset

Value

This function will output

Bsummary.txt

Index

* datasets	summary_permuted_binary, 10
Bcov_discovery, 2	summary_regular_binary,11
Bcov_target, 2	* values,
Bphe_discovery, 3	results_permuted_binary, 9
Bphe_target, 3	results_regular_binary, 9
DummyData,4	_
Qcov_discovery,7	Bcov_discovery, 2
Qcov_target, 7	Bcov_target, 2
Qphe_discovery, 8	Bphe_discovery, 3
Qphe_target, 8	Bphe_target, 3
* disease	Diametr Data 4
results_permuted_binary,9	DummyData,4
results_regular_binary,9	GWAS_binary,4
* gwas	GWEIS_binary, 5
GWAS_binary,4	
* gwies,	PRS_binary, 6
GWEIS_binary,5	_ •
* gxe	Qcov_discovery,7
GWEIS_binary,5	Qcov_target, 7
* individual	Qphe_discovery, 8
results_permuted_binary,9	<code>Qphe_target, 8</code>
results_regular_binary,9	
* interaction,	results_permuted_binary,9
GWEIS_binary, 5	results_regular_binary,9
* profile	summary_permuted_binary, 10
PRS_binary, 6	summary_regular_binary, 11
* prs,	Sammary_regurar_bridary, 11
PRS_binary, 6	
* regression	
summary_permuted_binary, 10	
summary_regular_binary,11	
* risk,	
results_permuted_binary,9	
results_regular_binary,9	
* risk	
results_permuted_binary,9	
results_regular_binary,9	
* scores,	
results_permuted_binary,9	
results_regular_binary, 9	
* scores	
PRS_binary, 6	
* summary	