## Package 'GxEprs'

May 29, 2024

Version 1.2
<b>Description</b> A novel PRS model is introduced to enhance the prediction accuracy by utilising GxE ef-
fects. This package performs Genome Wide Association Studies (GWAS) and Genome Wide En
vironment Interaction Studies (GWEIS) using a discovery dataset. The package has the abil-

ity to obtain polygenic risk scores (PRSs) for a target sample. Finally it predicts the risk values of each individual in the target sample. Users have the choice of using existing models (Li et al., 2015) <doi:10.1093/annonc/mdu565>, (Pan-

dis et al., 2013) <doi:10.1093/ejo/cjt054>, (Pey-

Title Genotype-by-Environment Interaction in Polygenic Score Models

rot et al., 2018) <a href="https://doi:10.1016/j.biopsych.2017.09.009">doi:10.1038/s41467-022-32407-9</a>, as well as newly proposed models for genomic risk prediction (refer to the URL for more details).

URL https://github.com/DoviniJ/GxEprs

License GPL (>=3)

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LazyData true

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## 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 800 rows and 18 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Standardized covariate

Column 4 Square of the standardized covariate

Column 5 Confounder 1

Column 6 Confounder 2

Column 7 Confounder 3

Column 8 Confounder 4

Column 9 Confounder 5

Column 10 Confounder 6

Column 11 Confounder 7

Column 12 Confounder 8

Column 13 Confounder 9

Column 14 Confounder 10

Column 15 Confounder 11

Column 16 Confounder 12

Column 17 Confounder 13

Column 18 Confounder 14

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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 200 rows and 18 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Standardized covariate

Column 4 Square of the standardized covariate

Column 5 Confounder 1

Column 6 Confounder 2

Column 7 Confounder 3

Column 8 Confounder 4

Column 9 Confounder 5

Column 10 Confounder 6

Column 11 Confounder 7

Column 12 Confounder 8

Column 13 Confounder 9

Column 14 Confounder 10

Column 15 Confounder 11

Column 16 Confounder 12

Column 17 Confounder 13

Column 18 Confounder 14

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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.
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## 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 800 rows and 3 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Phenotype (1=controls, 2=cases)

Bphe_target	Phenotype data file of the target dataset when the outcome is binary.
	This contains phenotype information of the individuals in the target
	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 200 rows and 3 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Phenotype (0=controls, 1=cases)

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DummyData.bim

PLINK .bim file

## Description

PLINK .bim file

## Usage

DummyData.bim

### **Format**

This follows PLINK general format

Column 1 Chromosome ID

Column 2 SNP ID

Column 3 Position of centimorgans

Column 4 Base-pair coordinate

Column 5 Minor Allele

Column 6 Reference Allele

DummyData.fam

PLINK .fam file

## Description

PLINK .fam file

## Usage

DummyData.fam

#### **Format**

This follows PLINK general format

Column 1 Family ID

Column 2 Individual ID

Column 3 Father's ID

Column 4 Mother's ID

Column 5 Sex

Column 6 Phenotype value

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DummyData.map

PLINK .map file

## Description

PLINK .map file

## Usage

DummyData.map

#### **Format**

This follows PLINK general format

DummyData.ped

PLINK .ped file

## Description

PLINK .ped file

## Usage

DummyData.ped

#### **Format**

This follows PLINK general format

GWAS\_binary

GWAS\_binary function This function performs GWAS using plink2 and outputs the GWAS summary statistics with additive SNP effects. Users may save the output in a user-specified file (see example).

## **Description**

GWAS\_binary function This function performs GWAS using plink2 and outputs the GWAS summary statistics with additive SNP effects. Users may save the output in a user-specified file (see example).

## Usage

GWAS\_binary(plink\_path, b\_file, Bphe\_discovery, Bcov\_discovery, thread = 20)

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#### **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

Bphe\_discovery Name (with file extension) of the phenotype file containing family ID, individual

ID and phenotype of the discovery dataset as columns, without heading

Bcov\_discovery 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

thread Number of threads used

#### Value

This function will perform GWAS and output

B\_out.trd.sum GWAS summary statistics with additive SNP effects

#### **Examples**

```
## Not run:
x <- GWAS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery,</pre>
thread = 20)
sink("B_out.trd.sum") #to create a file in the working directory
write.table(x[c("ID", "A1", "BETA")], sep = " ",
row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
head(x) #to obtain the head of GWAS summary statistics of additive SNP effects
x$CHROM #to extract the chromosome number
x$POS #to extract the base pair position
x$ID #to extract the SNP ID
x$REF #to extract the reference allele
x$ALT #to extract the alternate allele
x$A1 #to extract the minor allele
x$OBS_CT #to extract the number of allele observations
x$BETA #to extract the SNP effects
x$SE #to extract the standard errors of the SNP effects
x$Z_STAT #to extract the test statistics
x$P #to extract the p values
## End(Not run)
```

 ${\tt GWAS\_quantitative}$ 

GWAS\_quantitative function This function performs GWAS using plink2 and outputs the GWAS summary statistics with additive SNP effects. Users may save the output in a user-specified file (see example).

#### **Description**

GWAS\_quantitative function This function performs GWAS using plink2 and outputs the GWAS summary statistics with additive SNP effects. Users may save the output in a user-specified file (see example).

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#### Usage

```
GWAS_quantitative(
  plink_path,
  b_file,
  Qphe_discovery,
  Qcov_discovery,
  thread = 20
)
```

#### **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

Qphe\_discovery Name (with file extension) of the phenotype file containing family ID, individual

ID and phenotype of the discovery dataset as columns, without heading

Qcov\_discovery 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

thread Number of threads used

#### Value

This function will perform GWAS and output

Q\_out.trd.sum GWAS summary statistics with additive SNP effects

```
## Not run:
x <- GWAS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery,</pre>
thread = 20)
sink("Q_out.trd.sum") #to create a file in the working directory
write.table(x[c("ID", "A1", "BETA")], sep = " ",
row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
head(x) #to obtain the head of GWAS summary statistics of additive SNP effects
x$CHROM #to extract the chromosome number
x$POS #to extract the base pair position
x$ID #to extract the SNP ID
x$REF #to extract the reference allele
x$ALT #to extract the alternate allele
x$A1 #to extract the minor allele
x$OBS_CT #to extract the number of allele observations
x$BETA #to extract the SNP effects
x$SE #to extract the standard errors of the SNP effects
x$T_STAT #to extract the test statistics
x$P #to extract the p values
## End(Not run)
```

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GWEIS_binary	GWEIS_binary function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics with additive SNP effects and interaction SNP effects. Users may save the outputs in separate user-specified files (see examples).

## **Description**

GWEIS\_binary function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics with additive SNP effects and interaction SNP effects. Users may save the outputs in separate user-specified files (see examples).

#### Usage

```
GWEIS_binary(plink_path, b_file, Bphe_discovery, Bcov_discovery, thread = 20)
```

## **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
Bphe_discovery	Phenotype file containing family ID, individual ID and phenotype of the discovery dataset as columns, without heading
Bcov_discovery	Covariate file containing family ID, individual ID, standardized covariate, square of standardized covariate, and/or confounders of the discovery dataset as columns, without heading
thread	Number of threads used

#### Value

This function will perform GWEIS and output

B\_out.sum GWEIS summary statistics with additive and interaction SNP effects

```
## Not run:
x <- GWEIS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery,</pre>
thread = 20)
sink("B_out.add.sum") #to create a file in the working directory
write.table(x[c("ID", "A1", "ADD_BETA")], sep = " ",
row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
sink("B_out.gxe.sum") #to create a file in the working directory
write.table(x[c("ID", "A1", "INTERACTION_BETA")], sep = " ",
row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
head(x) #to extract the head of all columns in GWEIS summary
#statistics of additive and interaction SNP effects
x$CHROM #to extract the chromosome number
x$POS #to extract the base pair position
xID #to extract the SNP ID
```

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```
x$REF #to extract the reference allele
x$ALT #to extract the alternate allele
x$A1 #to extract the minor allele
x$OBS_CT #to extract the number of allele observations
x$ADD_BETA #to extract the additive SNP effects
x$ADD_SE #to extract the standard errors of the
#additive SNP effects
x$ADD_Z_STAT #to extract the test statistics of additive
#SNP effects
x$ADD_P #to extract the p values of additive SNP effects
x$INTERACTION_BETA #to extract the interaction SNP effects
x$INTERACTION_SE #to extract the standard errors of the
#interaction SNP effects
x$INTERACTION_Z_STAT #to extract the test statistics of
#interaction SNP effects
x$INTERACTION_P #to extract the p values of interaction
#SNP effects
## End(Not run)
```

GWEIS\_quantitative

GWEIS\_quantitative function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics with additive SNP effects and interaction SNP effects separately. It is recommended to save the outputs in separate user-specified files (see examples).

### **Description**

GWEIS\_quantitative function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics with additive SNP effects and interaction SNP effects separately. It is recommended to save the outputs in separate user-specified files (see examples).

#### Usage

```
GWEIS_quantitative(
  plink_path,
  b_file,
  Qphe_discovery,
  Qcov_discovery,
  thread = 20
)
```

#### **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
	Phenotype file containing family ID, individual ID and phenotype of the discovery dataset as columns, without heading
-	Covariate file containing family ID, individual ID, standardized covariate, square of standardized covariate, and/or confounders of the discovery dataset as columns, without heading
thread	Number of threads used

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#### Value

This function will perform GWEIS and output

Q\_out.sum GWEIS summary statistics with additive and interaction SNP effects

#### **Examples**

```
## Not run:
x <- GWEIS_quantitative (plink_path, DummyData, Qphe_discovery, Qcov_discovery,
thread = 20)
sink("Q_out.add.sum") #to create a file in the working directory
write.table(x[c("ID", "A1", "ADD_BETA")], sep = " ",
row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
sink("0_out.gxe.sum") #to create a file in the working directory
write.table(x[c("ID", "A1", "INTERACTION_BETA")], sep = " ",
row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
head(x) #to extract the head of all columns in GWEIS summary
#statistics of additive and interaction SNP effects
x$CHROM #to extract the chromosome number
x$POS #to extract the base pair position
x$ID #to extract the SNP ID
x$REF #to extract the reference allele
x$ALT #to extract the alternate allele
x$A1 #to extract the minor allele
x$OBS_CT #to extract the number of allele observations
x$ADD_BETA #to extract the additive SNP effects
x$ADD_SE #to extract the standard errors of the
#additive SNP effects
x$ADD_T_STAT #to extract the test statistics of additive
#SNP effects
x$ADD_P #to extract the p values of additive SNP effects
x$INTERACTION BETA #to extract the interaction SNP effects
x$INTERACTION SE #to extract the standard errors of the
#interaction SNP effects
x$INTERACTION_T_STAT #to extract the test statistics of
#interaction SNP effects
x$INTERACTION_P #to extract the p values of interaction
#SNP effects
## End(Not run)
```

PRS\_binary

PRS\_binary function This function uses plink2 and outputs Polygenic Risk Scores (PRSs) of all the individuals, using pre-generated GWAS and/or GWEIS summary statistics. Note that the input used in this function can be generated by using GWAS\_binary and/or GWEIS\_binary functions. Users may save the output in a user-specified file (see examples).

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#### **Description**

PRS\_binary function This function uses plink2 and outputs Polygenic Risk Scores (PRSs) of all the individuals, using pre-generated GWAS and/or GWEIS summary statistics. Note that the input used in this function can be generated by using GWAS\_binary and/or GWEIS\_binary functions. Users may save the output in a user-specified file (see examples).

#### Usage

```
PRS_binary(plink_path, b_file, summary_input)
```

#### **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

summary\_input Pre-generated GWAS and/or GWEIS summary statistics

#### Value

This function will output

prs.sscore PRSs for each individual

```
## Not run:
a <- GWAS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery)</pre>
trd <- a[c("ID", "A1", "BETA")]</pre>
b <- GWEIS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery)</pre>
add <- b[c("ID", "A1", "ADD_BETA")]
gxe <- b[c("ID", "A1", "INTERACTION_BETA")]</pre>
x <- PRS_binary(plink_path, DummyData, summary_input = trd)</pre>
sink("B_trd.sscore") #to create a file in the working directory
write.table(x, sep = " ", row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
head(x) #to read the head of all columns in the output
x$FID #to extract the family ID's of full dataset
x$IID #to extract the individual ID's of full dataset
x$PRS #to extract the polygenic risk scores of full dataset
y <- PRS_binary(plink_path, DummyData, summary_input = add)</pre>
sink("B_add.sscore") #to create a file in the working directory
write.table(y, sep = " ", row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
z <- PRS_binary(plink_path, DummyData, summary_input = gxe)</pre>
sink("B_gxe.sscore") #to create a file in the working directory
write.table(z, sep = " ", row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
## End(Not run)
```

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PRS	quantitative
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PRS\_quantitative function This function uses plink2 and outputs Polygenic Risk Scores (PRSs) of all the individuals, using pre-generated GWAS and/or GWEIS summary statistics. Note that the input used in this function can be generated by using GWAS\_quantitative and/or GWEIS\_quantitative functions. Users may save the output in a user-specified file (see examples).

#### **Description**

PRS\_quantitative function This function uses plink2 and outputs Polygenic Risk Scores (PRSs) of all the individuals, using pre-generated GWAS and/or GWEIS summary statistics. Note that the input used in this function can be generated by using GWAS\_quantitative and/or GWEIS\_quantitative functions. Users may save the output in a user-specified file (see examples).

#### Usage

```
PRS_quantitative(plink_path, b_file, summary_input)
```

## 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

summary\_input Pre-generated GWAS and/or GWEIS summary statistics

#### Value

This function will output

prs.sscore PRSs for each individual

```
## Not run:
a <- GWAS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery)</pre>
trd <- a[c("ID", "A1", "BETA")]</pre>
b <- GWEIS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery)</pre>
add <- b[c("ID", "A1", "ADD_BETA")]
gxe <- b[c("ID", "A1", "INTERACTION_BETA")]</pre>
x <- PRS_quantitative(plink_path, DummyData, summary_input = trd)</pre>
sink("Q_trd.sscore") #to create a file in the working directory
write.table(x, sep = " ", row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
head(x) #to read the head of all columns in the output
x$FID #to extract the family ID's of full dataset
x$IID #to extract the individual ID's of full dataset
x$PRS #to extract the polygenic risk scores of full dataset
y <- PRS_quantitative(plink_path, DummyData, summary_input = add)</pre>
sink("Q_add.sscore") #to create a file in the working directory
write.table(y, sep = " ", row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
z <- PRS_quantitative(plink_path, DummyData, summary_input = gxe)</pre>
```

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```
sink("Q_gxe.sscore") #to create a file in the working directory
write.table(z, sep = " ", row.names = FALSE, quote = FALSE) #to write the output
sink() #to save the output
## End(Not run)
```

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 800 rows and 18 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Standardized covariate

Column 4 Square of the standardized covariate

Column 5 Confounder 1

Column 6 Confounder 2

Column 7 Confounder 3

Column 8 Confounder 4

Column 9 Confounder 5

Column 10 Confounder 6

Column 11 Confounder 7

Column 12 Confounder 8

Column 13 Confounder 9

Column 14 Confounder 10

Column 15 Confounder 11

Column 16 Confounder 12

Column 17 Confounder 13

Column 18 Confounder 14

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Qcov_target	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.
	target adiaset 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 200 rows and 18 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Standardized covariate

Column 4 Square of the standardized covariate

Column 5 Confounder 1

Column 6 Confounder 2

Column 7 Confounder 3

Column 8 Confounder 4

Column 9 Confounder 5

Column 10 Confounder 6

Column 11 Confounder 7

Column 12 Confounder 8

Column 13 Confounder 9

Column 14 Confounder 10

Column 15 Confounder 11

Column 16 Confounder 12

Column 17 Confounder 13

Column 18 Confounder 14

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Qphe_discovery	Phenotype data file of the discovery dataset when the outcome is quantitative. This contains phenotype information of the individuals in the 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 800 rows and 3 columns

Column 1 Family IDColumn 2 Individual IDColumn 3 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.

### **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 200 rows and 3 columns

Column 1 Family IDColumn 2 Individual IDColumn 3 Phenotype

summary\_permuted\_binary

summary\_permuted\_binary function This function outputs the p value of permuted model in the target dataset, using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_quantitative function. It is recommended to run this function, if you choose to fit 'PRS\_gxe x E' interaction component (i.e. novel proposed model, Model 5) when generating risk scores. If the 'PRS\_gxe x E' term is significant in Model 5, and insignificant in Model 5\* (permuted p value), consider that the 'PRS\_gxe x E' interaction component is actually insignificant (always give priority to the p value obtained from the permuted model).

#### **Description**

summary\_permuted\_binary function This function outputs the p value of permuted model in the target dataset, using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_quantitative function. It is recommended to run this function, if you choose to fit 'PRS\_gxe x E' interaction component (i.e. novel proposed model, Model 5) when generating risk scores. If the 'PRS\_gxe x E' term is significant in Model 5, and insignificant in Model 5\* (permuted p value), consider that the 'PRS\_gxe x E' interaction component is actually insignificant (always give priority to the p value obtained from the permuted model).

#### Usage

```
summary_permuted_binary(
   Bphe_target,
   Bcov_target,
   iterations = 1000,
   add_score,
   gxe_score
)
```

## 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
iterations	Number of iterations used in permutation
add_score	PRSs generated using additive SNP effects of GWEIS summary statistics
gxe_score	PRSs generated using interaction SNP effects of GWEIS summary statistics

## Value

This function will output

B\_permuted\_p the p value of the permuted model

#### **Examples**

```
## Not run:
a <- GWEIS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery)
add <- a[c("ID", "A1", "ADD_OR")]
gxe <- a[c("ID", "A1", "INTERACTION_OR")]
p <- PRS_binary(plink_path, DummyData, summary_input = add)
q <- PRS_binary(plink_path, DummyData, summary_input = gxe)
x <- summary_permuted_binary(Bphe_target, Bcov_target, iterations = 1000, add_score = p, gxe_score = q)
x
## End(Not run)</pre>
```

summary\_permuted\_quantitative

summary\_permuted\_quantitative function This function outputs the p value of permuted model in the target dataset, using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_quantitative functions. It is recommended to run this function, if you choose to fit 'PRS\_gxe x E' interaction component (i.e. novel proposed model, Model 4) when generating risk scores. If the 'PRS\_gxe x E' term is significant in Model 4, and insignificant in Model 4\* (permuted p value), consider that the 'PRS\_gxe x E' interaction component is actually insignificant (always give priority to the p value obtained from the permuted model).

### **Description**

summary\_permuted\_quantitative function This function outputs the p value of permuted model in the target dataset, using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_quantitative functions. It is recommended to run this function, if you choose to fit 'PRS\_gxe x E' interaction component (i.e. novel proposed model, Model 4) when generating risk scores. If the 'PRS\_gxe x E' term is significant in Model 4, and insignificant in Model 4\* (permuted p value), consider that the 'PRS\_gxe x E' interaction component is actually insignificant (always give priority to the p value obtained from the permuted model).

#### Usage

```
summary_permuted_quantitative(
    Qphe_target,
    Qcov_target,
    iterations = 1000,
    add_score,
    gxe_score
)
```

#### **Arguments**

Qphe\_target

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

Qcov_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	
iterations	Number of iterations used in permutation	
add_score	PRSs generated using additive SNP effects of GWEIS summary statistics	
gxe_score	PRSs generated using interaction SNP effects of GWEIS summary statistics	

#### Value

This function will output

Q\_permuted\_p the p value of the permuted model

## **Examples**

```
## Not run:
a <- GWEIS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery)
add <- a[c("ID", "A1", "ADD_BETA")]
gxe <- a[c("ID", "A1", "INTERACTION_BETA")]
p <- PRS_quantitative(plink_path, DummyData, summary_input = add)
q <- PRS_quantitative(plink_path, DummyData, summary_input = gxe)
x <- summary_permuted_quantitative(Qphe_target, Qcov_target, iterations = 1000, add_score = p, gxe_score = q)
x
## End(Not run)</pre>
```

summary\_regular\_binary

summary\_regular\_binary function This function outputs the summary of regular model and final risk score values of each individual in the target dataset using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_binary function.

#### **Description**

summary\_regular\_binary function This function outputs the summary of regular model and final risk score values of each individual in the target dataset using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_binary function.

## Usage

```
summary_regular_binary(
   Bphe_target,
   Bcov_target,
   add_score = NULL,
   gxe_score = NULL,
   Model
)
```

#### **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

add\_score PRSs generated using additive SNP effects of GWAS/GWEIS summary statis-

tics

gxe\_score PRSs generated using interaction SNP effects of GWEIS summary statistics

Model Specify the model number (0:  $y = PRS_{trd} + E + confounders$ , 1:  $y = PRS_{trd}$ 

+ E + PRS\_trd x E + confounders, 2: y = PRS\_add + E + PRS\_add x E + confounders, 3: y = PRS\_add + E + PRS\_gxe x E + confounders, 4: y = PRS\_add + E + PRS\_gxe + PRS\_gxe x E + confounders, 5: y = PRS\_add + E + E^2 + PRS\_gxe + PRS\_gxe x E + confounders, where y is the outcome variable, E is the covariate of interest, PRS\_trd and PRS\_add are the polygenic risk scores computed using additive SNP effects of GWAS and GWEIS summary statistics respectively, and PRS\_gxe is the polygenic risk scores computed using GxE

interaction SNP effects of GWEIS summary statistics.)

#### Value

This function will output

Bsummary the summary of the fitted model

Individual\_risk\_values

the estimated risk values of individuals in the target sample

```
## Not run:
a <- GWAS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery)</pre>
trd <- a[c("ID", "A1", "OR")]</pre>
b <- GWEIS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery)</pre>
add <- b[c("ID", "A1", "ADD_OR")]
gxe <- b[c("ID", "A1", "INTERACTION_OR")]</pre>
p <- PRS_binary(plink_path, DummyData, summary_input = trd)</pre>
q <- PRS_binary(plink_path, DummyData, summary_input = add)</pre>
r <- PRS_binary(plink_path, DummyData, summary_input = gxe)</pre>
summary_regular_binary(Bphe_target, Bcov_target,
                              add_score = p,
                              Model = 0)
summary_regular_binary(Bphe_target, Bcov_target,
                              add_score = p,
                              Model = 1)
summary_regular_binary(Bphe_target, Bcov_target,
                              add_score = q,
                              Model = 2)
\verb|summary_regular_binary| Bphe\_target, Bcov\_target, \\
                              add_score = q,
                              gxe\_score = r,
                              Model = 3)
summary_regular_binary(Bphe_target, Bcov_target,
                              add_score = q,
```

```
gxe\_score = r,
                            Model = 4)
x <- summary_regular_binary(Bphe_target, Bcov_target,</pre>
                            add_score = q,
                             gxe\_score = r,
                            Model = 5)
sink("Bsummary.txt") #to create a file in the working directory
print(x$summary) #to write the output
sink() #to save the output
sink("Individual_risk_values.txt") #to create a file in the working directory
write.table(x$risk.values, sep = " ", row.names = FALSE, col.names = FALSE,
quote = FALSE) #to write the output
sink() #to save the output
x$summary #to obtain the model summary output
x$risk.values #to obtain the predicted risk values of target individuals
## End(Not run)
```

summary\_regular\_quantitative

summary\_regular\_quantitative function This function outputs the summary of regular model and final risk score values of each individual in the target dataset using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_quantitative function.

## **Description**

summary\_regular\_quantitative function This function outputs the summary of regular model and final risk score values of each individual in the target dataset using pre-generated Polygenic Risk Scores (PRSs) of all the individuals. Note that the input used in this function can be generated by using PRS\_quantitative function.

#### Usage

```
summary_regular_quantitative(
    Qphe_target,
    Qcov_target,
    add_score = NULL,
    gxe_score = NULL,
    Model
)
```

### **Arguments**

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

Ocov\_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

add\_score PRSs generated using additive SNP effects of GWAS/GWEIS summary statis-

gxe\_score

PRSs generated using interaction SNP effects of GWEIS summary statistics

Model

Specify the model number (0: y = PRS\_trd + E + confounders, 1: y = PRS\_trd + E + PRS\_trd x E + confounders, 2: y = PRS\_add + E + PRS\_add x E + confounders, 3: y = PRS\_add + E + PRS\_gxe x E + confounders, 4: y = PRS\_add + E + PRS\_gxe + PRS\_gxe x E + confounders, where y is the outcome variable, E is the covariate of interest, PRS\_trd and PRS\_add are the polygenic risk scores computed using additive SNP effects of GWAS and GWEIS summary statistics respectively, and PRS\_gxe is the polygenic risk scores computed using GxE interaction SNP effects of GWEIS summary statistics.)

#### Value

This function will output

 $\label{lem:continuous} {\tt Qsummary.txt} \qquad {\tt the summary of the fitted model} \\ {\tt Individual\_risk\_values.txt}$ 

the estimated risk values of individuals in the target sample

```
## Not run:
a <- GWAS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery)</pre>
trd <- a[c("ID", "A1", "BETA")]</pre>
b <- GWEIS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery)</pre>
add <- b[c("ID", "A1", "ADD_BETA")]
gxe <- b[c("ID", "A1", "INTERACTION_BETA")]</pre>
p <- PRS_quantitative(plink_path, DummyData, summary_input = trd)</pre>
q <- PRS_quantitative(plink_path, DummyData, summary_input = add)</pre>
r <- PRS_quantitative(plink_path, DummyData, summary_input = gxe)</pre>
\verb|summary_regular_quantitative(Qphe\_target, Qcov\_target,
                              add score = p.
                              Model = 0)
summary_regular_quantitative(Qphe_target, Qcov_target,
                              add_score = p,
                              Model = 1)
summary_regular_quantitative(Qphe_target, Qcov_target,
                              add_score = q,
                              Model = 2)
summary_regular_quantitative(Qphe_target, Qcov_target,
                              add_score = q,
                              gxe\_score = r,
                              Model = 3)
x <- summary_regular_quantitative(Qphe_target, Qcov_target,</pre>
                              add_score = q,
                              gxe\_score = r,
                              Model = 4)
sink("Qsummary.txt") #to create a file in the working directory
print(x$summary) #to write the output
sink() #to save the output
sink("Individual_risk_values.txt") #to create a file in the working directory
write.table(x$risk.values, sep = " ", row.names = FALSE, col.names = FALSE,
quote = FALSE) #to write the output
sink() #to save the output
x$summary #to obtain the model summary output
x$risk.values #to obtain the predicted risk values of target individuals
```

## End(Not run)

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