# Package 'GxEprsDummy'

July 6, 2023

Title What the Package Does (One Line, Title Case)	
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<b>Description</b> What the package does (one paragraph).	
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2 Bcov\_discovery

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

# 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

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

Bcov\_target 3

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

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

4 Bphe\_target

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

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 1939 rows and 3 columns

Column 1 Family ID

Column 2 Individual ID

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

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DummyData	PLINK binary data files This contains DummyData.fam, Dummy-
	Data.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 function This function performs GWAS using plink2 and outputs the GWAS summary statistics file with additive SNP effects named  $B_{\rm out.trd.sum}$ 

#### **Description**

GWAS\_binary function This function performs GWAS using plink2 and outputs the GWAS summary statistics file with additive SNP effects named B\_out.trd.sum

# Usage

```
GWAS_binary(
  plink_path,
  b_file,
  Bphe_discovery,
  Bcov_discovery,
  thread = 20,
  summary_output = "B_out.trd.sum")
```

#### **Arguments**

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

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```
thread Number of threads used summary_output
```

Name of the SNP effects of the GWAS summary statistics file specified by the user

#### Value

This function will perform GWAS and output

```
B_out.trd.sum
```

GWAS summary statistics file with additive SNP effects

#### **Examples**

```
## Not run:
x <- GWAS_binary(plink_path, DummyData, Bphe_discovery, Bcov_discovery,
                 thread = 20, summary_output = "B_out.trd.sum")
head(x) #to read the head of all columns in B_out.trd.sum file
x$V1 #to extract the chromosome number (CHROM)
x$V2 #to extract the base pair position (POS)
x$V3 #to extract the SNP ID (ID)
x$V4 #to extract the reference allele (REF)
x$V5 #to extract the alternate allele (ALT)
x$V6 #to extract the minor allele (A1)
x$V7 #to extract whether firth regression is used (FIRTH?)
x$V8 #to extract the type of test performed (TEST)
x$V9 #to extract the nmber of allele observations (OBS_CT)
x$V10 #to extract the odds ration of the SNP effect (OR)
x$V11 #to extract the standard error of log odds (LOG(OR)_SE)
x$V12 $$to extract the test statistic (Z_STAT)
x$V13 #to extract the p value (P)
x$V14 #to extract the error code (ERRCODE)
## End(Not run)
```

GWAS\_quantitative GWAS\_quantitative function This function performs GWAS using plink2 and outputs the GWAS summary statistics file with additive SNP effects named Q\_out.trd.sum

# Description

GWAS\_quantitative function This function performs GWAS using plink2 and outputs the GWAS summary statistics file with additive SNP effects named Q\_out.trd.sum

## Usage

```
GWAS_quantitative(
  plink_path,
  b_file,
  Qphe_discovery,
  Qcov_discovery,
  thread = 20,
  summary_output = "Q_out.trd.sum"
)
```

GWEIS\_binary 7

#### **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

summary\_output

Name of the SNP effects of the GWAS summary statistics file specified by the user

#### Value

This function will perform GWAS and output

Q\_out.trd.sum

GWAS summary statistics file with additive SNP effects

#### **Examples**

```
x <- GWAS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery,
                       thread = 20, summary_output = "Q_out.trd.sum")
head(x) #to read the head of all columns in Q_out.trd.sum file
x$V1 #to extract the chromosome number (CHROM)
x$V2 $\#to extract the base pair position (POS)
x$V3 $\#to extract the SNP ID (ID)
x$V4 #to extract the reference allele (REF)
x$V5 #to extract the alternate allele (ALT)
x$V6 #to extract the minor allele (A1)
x$V7 $$#to extract the type of test performed (TEST)
x$V8 #to extract the nmber of allele observations (OBS_CT)
x$V9 $\#$to extract the SNP effect (BETA)
x$V10 #to extract the standard error of each SNP effect (SE)
x$V11 #to extract the test statistic (T_STAT)
x$V12 #to extract the p value (P)
x$V13 $$to extract the error code (ERRCODE)
## End(Not run)
```

GWEIS\_binary

GWEIS\_binary function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics files with additive SNP effects named B\_out.add.sum and interaction SNP effects named B\_out.gxe.sum

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#### **Description**

GWEIS\_binary function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics files with additive SNP effects named B\_out.add.sum and interaction SNP effects named B\_out.gxe.sum

#### Usage

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

#### **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

 $\begin{array}{cc} \text{thread} & \text{Number of threads used} \\ \text{summary\_output} \end{array}$ 

Name (prefix) of the additive or interaction SNP effects of the GWEIS summary statistics file specified by the user

### Value

This function will perform GWEIS and output

```
B_out.add.sum
```

GWEIS summary statistics file with additive SNP effects

B\_out.gxe.sum

GWEIS summary statistics file with interaction SNP effects

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```
x[[1]] $V6 #to extract the minor allele (A1)
x[[1]]$V7 #to extract whether firth regression is used (FIRTH?)
x[[1]] $V8 #to extract the type of test performed (TEST)
x[[1]] $V9 #to extract the nmber of allele observations (OBS_CT)
x[[1]] $V10 #to extract the odds ration of the SNP effect (OR)
x[[1]] $V11 #to extract the standard error of log odds (LOG(OR)_SE)
x[[1]]$V12 #to extract the test statistic (Z_STAT)
x[[1]]$V13 #to extract the p value (P)
x[[1]]$V14 #to extract the error code (ERRCODE)
head(x[[2]]) #to extract the head of all columns in B_out.gxe.sum file
x[[2]] $V1 #to extract the chromosome number (CHROM)
x[[2]]$V2 #to extract the base pair position (POS)
x[[2]]$V3 #to extract the SNP ID (ID)
x[[2]]$V4 #to extract the reference allele (REF)
x[[2]] $V5 #to extract the alternate allele (ALT)
x[[2]] $V6 #to extract the minor allele (A1)
x[[2]] $V7 #to extract whether firth regression is used (FIRTH?)
x[[2]] $V8 #to extract the type of test performed (TEST)
x[[2]] $V9 #to extract the nmber of allele observations (OBS_CT)
x[[2]] $V10 #to extract the odds ration of the SNP effect (OR)
x[[2]] $V11 #to extract the standard error of log odds (LOG(OR)_SE)
x[[2]]$V12 #to extract the test statistic (Z_STAT)
x[[2]]$V13 #to extract the p value (P)
x[[2]]$V14 #to extract the error code (ERRCODE)
## End(Not run)
```

GWEIS\_quantitative GWEIS\_quantitative function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics files with additive SNP effects named Q\_out.add.sum and interaction SNP effects named Q\_out.gxe.sum

#### **Description**

GWEIS\_quantitative function This function performs GWEIS using plink2 and outputs the GWEIS summary statistics files with additive SNP effects named Q\_out.add.sum and interaction SNP effects named Q\_out.gxe.sum

#### Usage

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

#### **Arguments**

plink\_path Path to the PLINK executable application

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b\_file Prefix of the binary files, where all .fam, .bed and .bim files have a common prefix

Qphe\_discovery

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

Qcov\_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

 $\begin{array}{cc} \text{thread} & \text{Number of threads used} \\ \text{summary\_output} \end{array}$ 

Name (prefix) of the additive or interaction SNP effects of the GWEIS summary statistics file specified by the user

#### Value

This function will perform GWEIS and output

Q\_out.add.sum

GWEIS summary statistics file with additive SNP effects

Q\_out.gxe.sum

GWEIS summary statistics file with interaction SNP effects

```
## Not run:
x <- GWEIS_quantitative(plink_path, DummyData, Qphe_discovery, Qcov_discovery,
                        thread = 20, summary_output = "Q_out")
head(x[[1]]) #to read the head of all columns in Q_out.add.sum file
x[[1]]$V1 #to extract the chromosome number (CHROM)
x[[1]]$V2 #to extract the base pair position (POS)
x[[1]]$V3 #to extract the SNP ID (ID)
x[[1]]$V4 #to extract the reference allele (REF)
x[[1]] $V5 #to extract the alternate allele (ALT)
x[[1]] $V6 #to extract the minor allele (A1)
x[[1]] $V7 #to extract the type of test performed (TEST)
x[[1]] $V8 #to extract the nmber of allele observations (OBS_CT)
x[[1]] $V9 #to extract the SNP effect (BETA)
x[[1]]$V10 #to extract the standard error of each SNP effect (SE)
x[[1]]$V11 #to extract the test statistic (T_STAT)
x[[1]]$V12 #to extract the p value (P)
x[[1]]$V13 #to extract the error code (ERRCODE)
head(x[[2]]) #to read the head of all columns in Q_out.gxe.sum file
x[[2]] $V1 #to extract the chromosome number (CHROM)
x[[2]] $V2 #to extract the base pair position (POS)
x[[2]]$V3 #to extract the SNP ID (ID)
x[[2]] $V4 #to extract the reference allele (REF)
x[[2]$V5 $$to extract the alternate allele (ALT)
x[[2]] $V6 #to extract the minor allele (A1)
x[[2]]$V7 #to extract the type of test performed (TEST)
x[[2]] $V8 #to extract the nmber of allele observations (OBS_CT)
x[[2]]$V9 #to extract the SNP effect (BETA)
x[[2]] $V10 #to extract the standard error of each SNP effect (SE)
x[[2]] $V11 #to extract the test statistic (T_STAT)
x[[2]]$V12 #to extract the p value (P)
```

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```
x[[2]]$V13 #to extract the error code (ERRCODE)
## End(Not run)
```

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\_out.trd.sum, B\_out.add.sum or B\_out.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\_out.trd.sum, B\_out.add.sum or B\_out.gxe.sum

#### Usage

```
PRS_binary(
  plink_path,
  b_file,
  summary_input = "B_out.trd.sum",
  summary_output = "B_trd"
)
```

#### **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

Name of the summary statistics file specified by the user

summary\_output

Name of the PRS file generated using provided summary statistics file specified

by the user

#### Value

This function will output

B\_trd.sscore PRSs for each target individual using GWAS additive effects

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```
x$V5 #to extract the polygenic risk scores (SCORE1_AVG)
## End(Not run)
```

PRS\_quantitative

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 Q\_out.trd.sum, Q\_out.add.sum or Q\_out.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 Q\_out.trd.sum, Q\_out.add.sum or Q\_out.gxe.sum

#### Usage

```
PRS_quantitative(
  plink_path,
  b_file,
  summary_input = "Q_out.trd.sum",
  summary_output = "Q_trd"
)
```

#### **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

Name of the summary statistics file specified by the user

summary\_output

Name of the PRS file generated using provided summary statistics file specified by the user

# Value

This function will output

Q\_trd.sscore PRSs for each target individual using GWAS additive effects

Qcov\_discovery 13

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

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

Qcov\_target

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

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

Qphe\_discovery 15

Qphe_discovery	Phenotype data file of the discovery dataset when the outcome is quan-
	titative 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 6426 rows and 3 columns

Column 1 Family IDColumn 2 Individual ID

Column 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 1579 rows and 3 columns

Column 1 Family ID

Column 2 Individual ID

Column 3 Phenotype

```
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,
   iterations = 1000,
   add_score = "B_add.sscore",
   gxe_score = "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
iterations	Number of iterations used in permutation
add_score	The .sscore file generated using additive SNP effects of GWEIS summary statistics
gxe_score	The .sscore file generated using interaction SNP effects of GWEIS summary statistics

# Value

This function will output

```
B_permuted_p.txt
```

the p value of the permuted model

```
## Not run:
x <- summary_permuted_binary(Bphe_target, Bcov_target)
x
## End(Not run)</pre>
```

```
summary_permuted_quantitative
```

summary\_permuted\_quantitative 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 Q\_trd.sum, Q\_add.sum and Q\_gxe.sum

# Description

summary\_permuted\_quantitative 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 Q\_trd.sum, Q\_add.sum and Q\_gxe.sum

# Usage

```
summary_permuted_quantitative(
    Qphe_target,
    Qcov_target,
    iterations = 1000,
    add_score = "Q_add.sscore",
    gxe_score = "Q_gxe.sscore")
```

#### **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	The .sscore file generated using additive SNP effects of GWEIS summary statistics
gxe_score	The .sscore file generated using interaction SNP effects of GWEIS summary statistics

# Value

This function will output

```
Q_permuted_p.txt
```

the p value of the permuted model

```
## Not run:
x <- summary_permuted_quantitative(Qphe_target, Qcov_target)
x
## End(Not run)</pre>
```

```
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,
   trd_score = "B_trd.sscore",
   add_score = "B_add.sscore",
   gxe_score = "B_gxe.sscore",
   Model,
   summary_output = "Bsummary.txt",
   risk_output = "Individual_risk_values.txt")
```

# 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
trd_score	The .sscore file generated using additive SNP effects of GWAS summary statistics $% \left( 1\right) =\left( 1\right) \left( 1\right) $
add_score	The .sscore file generated using additive SNP effects of GWEIS summary statistics $% \left( 1\right) =\left( 1\right) \left( 1\right)$
gxe_score	The .sscore file generated using interaction SNP effects of GWEIS summary statistics $% \left( 1\right) =\left( 1\right) \left( $
Model	Specify the model number (1: $y = PRS\_trd + E + PRS\_trd \times E + confounders$ , 2: $y = PRS\_add + E + PRS\_add \times E + confounders$ , 3: $y = PRS\_add + E + PRS\_gxe \times E + confounders$ , 4: $y = PRS\_add + E + PRS\_gxe + PRS\_gxe \times E + confounders$ , 5: $y = PRS\_add + E + E^2 + PRS\_gxe + PRS\_gxe \times E + confounders$ , where $y$ is the outcome variable, $E$ is the covariate of interest, $E$ and $E$ and $E$ are the polygenic risk scores computed using additive SNP effects of GWAS and GWEIS summary statistics respectively, and $E$ are the polygenic risk scores computed using $E$ interaction SNP effects of GWEIS summary statistics.)
summary_output	

Name of the model summary file specified by the user

risk\_output Name of the file containing risk scores of the target individuals specified by the

#### Value

This function will output

```
\label{eq:boundary} \textbf{Bsummary.txt} \ \ \textbf{the summary of the fitted model} \\ \textbf{Individual\_risk\_values.txt}
```

the estimated risk values of individuals in the target sample

# **Examples**

```
## Not run:
x <- summary_regular_binary(Bphe_target, Bcov_target,
                            add_score = "B_add.sscore",
                            gxe_score = "B_gxe.sscore",
                            Model = 5,
                            summary_output = "Bsummary.txt",
                            risk_output = "Individual_risk_values.txt")
x[[1]][[1]] #to obtain the content of the Bsummary.txt file
x[[1]][[2]] #to extract "Call" of the model summary
x[[1]][[3]] #to extract terms of the model summary
x[[1]][[4]] #to extract family information of the model summary
x[[1]][[5]] #to extract deviance information of the model summary
x[[1]][[6]] #to extract AIC information of the model summary
x[[1]][[7]] #to extract contrasts of the model summary
x[[1]][[8]] #to extract degrees of freedom (df) of residuals of
            #the model summary
x[[1]][[9]] #to extract "Null deviance" of the model summary
x[[1]][[10]] #to extract degrees of freedom (df) of null deviance
             #of the model summary
x[[1]][[11]] #to extract "iter" of the model summary
x[[1]][[12]] #to extract deviance residuals
x[[1]][[13]] #to extract regression coefficients of the model summary
x[[1]][[14]] #to extract aliesed information of the model summary
x[[1]][[15]] #to extract dispersion information of the model summary
x[[1]][[16]] #to extract degrees of freedom of the model summary
x[[1]][[17]] #to extract unscaled variance covariance matrix of all variables
x[[1]][[18]] #to extract scaled variance covariance matrix of all variables
head(x[[2]]) #to read the head of all columns in Individual_risk_values.txt file
x[[2]][,1] #to extract the column containing family ID's
x[[2]][,2] #to extract the column containing individual ID's
x[[2]][,3] #to extract the column containing predicted risk scores
## End(Not run)
```

```
summary_regular_quantitative
```

summary\_regular\_quantitative 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 Q\_trd.sum, Q\_add.sum and Q\_gxe.sum

#### **Description**

summary\_regular\_quantitative 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 Q\_trd.sum, Q\_add.sum and Q\_gxe.sum

#### Usage

```
summary_regular_quantitative(
    Qphe_target,
    Qcov_target,
    trd_score = "Q_trd.sscore",
    add_score = "Q_add.sscore",
    gxe_score = "Q_gxe.sscore",
    Model,
    summary_output = "Qsummary.txt",
    risk_output = "Individual_risk_values.txt")
```

#### **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
trd_score	The .sscore file generated using additive SNP effects of GWAS summary statistics
add_score	The .sscore file generated using additive SNP effects of GWEIS summary statistics
gxe_score	The .sscore file generated using interaction SNP effects of GWEIS summary statistics
Model	Specify the model number (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.)
summary_output	

Name of the model summary file specified by the user

Name of the file containing risk scores of the target individuals specified by the

#### Value

# This function will output

user

risk\_output

```
Qsummary.txt the summary of the fitted model
Individual_risk_values.txt
the estimated risk values of individuals in the target sample
```

```
## Not run:
x <- summary_regular_quantitative(Qphe_target, Qcov_target,
                                  add_score = "Q_add.sscore",
                                  gxe_score = "Q_gxe.sscore",
                                  Model = 4,
                                  summary_output = "Qsummary.txt",
                                  risk_output = "Individual_risk_values.txt")
x[[1]][[1]] #to obtain the content of the Qsummary.txt file
x[[1]][[2]] #to extract "Call" of the model summary
x[[1]][[3]] #to extract terms of the model summary
x[[1]][[4]] #to extract the residuals
x[[1]][[5]] #to extract regression coefficients of the model summary
x[[1]][[6]] #to extract aliesed information of the model summary
x[[1]][[7]] #to extract "sigma" (residual standard error) information
            #of the model summary
x[[1]][[8]] #to extract degrees of freedom of the model summary
x[[1]][[9]] #to extract the R squared value of the model summary
\mathbf{x}[[1]][[10]] #to extract the adjusted R squared value of the model summary
\mathbf{x}[[1]][[11]] #to extract the test statistic values of the model summary
x[[1]][[12]] #to extract unscaled variance covariance matrix of all variables
head(x[[2]]) #to read the head of all columns in Individual_risk_values.txt file
x[[2]][,1] #to extract the column containing family ID's
x[[2]][,2] #to extract the column containing individual ID's
x[[2]][,3] #to extract the column containing predicted risk scores
## End(Not run)
```

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