Package 'STAARpipelineSummary'

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```
Annotate_Single_Variants
```

Functionally annotate a list of variants

Description

The Annotate_Single_Variants function takes in a list of variants to functionally annotate the input variants

Usage

```
Annotate_Single_Variants(
   agds_dir,
   single_variants_list,
   QC_label = "annotation/filter",
   Annotation_dir = "annotation/info/FunctionalAnnotation",
   Annotation_name_catalog,
   Annotation_name
)
```

Arguments

```
agds_dir
                  file directory of annotated GDS (aGDS) files for all chromosomes (1-22).
single_variants_list
                  a data frame containing the information of variants to be functionally anno-
                  tated. The data frame must include 4 columns with the following names: "CHR"
                  (chromosome number), "POS" (position), "REF" (reference allele), and "ALT"
                  (alternative allele).
QC_label
                  channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").
Annotation_dir channel name of the annotations in the aGDS file
                  (default = "annotation/info/FunctionalAnnotation").
Annotation_name_catalog
                  a data frame containing the annotation names and the corresponding channel
                  names in the aGDS file.
Annotation_name
                  a vector of qualitative/quantitative annotation names user wants to extract.
```

Value

a data frame containing the basic information (chromosome, position, reference allele and alternative allele) and annotation scores for the input variants.

Dynamic_Window_Results_Summary

Summarize the results of dynamic window analysis generated by STAARpipeline package and perform conditional analysis for (unconditionally) significant genetic regions by adjusting for a given list of known variants

Description

The Dynamic_Window_Results_Summary function takes in the results of dynamic window analysis generated by STAARpipeline package, the object from fitting the null model, and the set of known variants to be adjusted for in conditional analysis to summarize the dynamic window analysis results and analyze the conditional association between a quantitative/dichotomous phenotype and the rare variants in the unconditional significant genetic regions.

Usage

```
Dynamic_Window_Results_Summary(
  agds_dir,
  jobs_num,
  input_path,
  output_path,
  dynamic_window_results_name,
  obj_nullmodel,
  known_loci = NULL,
 method_cond = c("optimal", "naive"),
  QC_label = "annotation/filter",
  geno_missing_imputation = c("mean", "minor"),
  variant_type = c("SNV", "Indel", "variant"),
  Annotation_dir = "annotation/info/FunctionalAnnotation",
  Annotation_name_catalog,
 Use_annotation_weights = FALSE,
  Annotation_name = NULL,
  alpha = 0.05
```

Arguments

agds_dir a vector containing file directory of annotated GDS (aGDS) files for all chromosomes (1-22).

jobs_num a data frame containing the number of jobs for association analysis. The data frame must include a column with the name "scang_num" input_path file directory of the input dynamic window analysis results.

output_path file directory of the output summary results.

dynamic_window_results_name file names of the input dynamic window analysis results.

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel function in the GENESIS package and transformed using the genesis2staar_nullmodel

function in the STAARpipeline package.

known_loci a data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

method_cond a character value indicating the method for conditional analysis. optimal refers

> to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

channel name of the QC label in the GDS/aGDS file (default = "annotation/filter"). QC_label

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default =

"mean").

type of variant included in the conditional analysis. Choice includes "SNV", variant_type

"Indel", or "variant" (default = "SNV").

Annotation_dir channel name of the annotations in the aGDS file

(default = "annotation/info/FunctionalAnnotation").

Annotation_name_catalog

a data frame containing the name and the corresponding channel name in the

aGDS file.

Use_annotation_weights

use annotations as weights or not (default = FALSE).

Annotation_name

a vector of annotation names used in SCANG-STAAR (default = NULL).

alpha threshod to control the genome-wise (family-wise) error rate (default = 0.05).

Value

The function returns the following analysis results:

SCANG_S_res_uncond_cond.Rdata and SCANG_S_res_uncond_cond.csv: A matrix that summarized the unconditional and conditional results of the significant regions (GWER<alpha) detected by the SCANG-STAAR-S procedure (conditional results available if known_loci is not a NULL), including chromosome ("chr"), start position ("start_pos"), end position ("end_pos"), number of variants ("SNV_nos"), family-wise/genome-wide error rate (GWER), unconditional STAAR-S pvalue ("STAAR_S"), conditional STAAR-S p-value ("STAAR_S_cond"), conditional ACAT-V pvalue ("ACAT_V_cond"), conditional Burden p-value ("Burden_cond"), conditional SKAT p-value ("SKAT_cond"), and conditional STAAR-O p-value ("STAAR_O_cond").

SCANG_B_res_uncond_cond.Rdata and SCANG_B_res_uncond_cond.csv: A matrix that summarized the unconditional and conditional results of the significant regions detected by the SCANG-STAAR-B procedure (conditional results available if known_loci is not a NULL). Details see SCANG-STAAR-S.

SCANG_O_res_uncond_cond.Rdata and SCANG_O_res_uncond_cond.csv: A matrix that summarized the unconditional and conditional results of the significant regions detected by the SCANG-STAAR-O procedure (conditional results available if known_loci is not a NULL). Details see SCANG-STAAR-S.

results_dynamic_window.Rdata: A Rdata file that summarized the significant regions detected by SCANG-STAAR procedure.

SCANG_S_top1.Rdata and SCANG_S_top1.csv: A matrix that summarized the top 1 unconditional region detected by SCANG-STAAR-S, including the STAAR-S p-value ("STAAR_S"), chromosome ("chr"), start position ("start_pos"), end position ("end_pos"), family-wise/genome-wide error rate (GWER) and the number of variants ("SNV_nos").

SCANG_B_top1.Rdata and SCANG_B_top1.csv: A matrix that summarized the top 1 unconditional region detected by SCANG-STAAR-B. Details see SCANG-STAAR-B.

SCANG_0_top1.Rdata and SCANG_0_top1.csv: A matrix that summarized the top 1 unconditional region detected by SCANG-STAAR-O. Details see SCANG-STAAR-O.

SCANG_S_res.Rdata and SCANG_S_res.csv: A matrix that summarized the significant regions (GWER<alpha) detected by SCANG-STAAR-S, including the negative log transformation of STAAR-S p-value ("-logp"), chromosome ("chr"), start position ("start_pos"), end position ("end_pos"), family-wise/genome-wide error rate (GWER) and the number of variants ("SNV_num").

SCANG_B_res.Rdata and SCANG_B_res.csv: A matrix that summarized the significant regions detected by SCANG-STAAR-B. Details see SCANG-STAAR-S.

SCANG_O_res.Rdata and SCANG_O_res.csv: A matrix that summarized the significant regions detected by SCANG-STAAR-O. Details see SCANG-STAAR-S.

SCANG_S_res_cond.Rdata and SCANG_S_res_cond.csv: A matrix that summarized the conditional p-values of the significant regions (GWER<alpha) detected by SCANG-STAAR-S, including chromosome ("chr"), start position ("Start Loc"), end position ("End Loc"), the number of variants ("#SNV"), annotation-weighted ACAT-V, Burden and SKAT conditional p-values, and STAAR conditional p-values of the regions with GWER smaller than the threshold alpha (available if known_loci is not a NULL).

SCANG_B_res_cond.Rdata and SCANG_B_res_cond.csv: A matrix that summarized the conditional p-values of the significant regions (GWER<alpha) detected by SCANG-STAAR-B (available if known_loci is not a NULL), Details see SCANG-STAAR-S.

SCANG_0_res_cond.Rdata and SCANG_0_res_cond.csv: A matrix that summarized the conditional p-values of the significant regions (GWER<alpha) detected by SCANG-STAAR-O (available if known_loci is not a NULL), Details see SCANG-STAAR-S.

Gene_Centric_Coding_Info

Functionally annotate rare variants in a coding mask

Description

The Gene_Centric_Coding_Info function takes in a coding mask of a gene to functionally annotate the rare variants in the mask.

Usage

```
Gene_Centric_Coding_Info(
  category = c("plof", "plof_ds", "missense", "disruptive_missense", "synonymous"),
  chr,
  genofile,
  obj_nullmodel,
  gene_name,
  known_loci = NULL,
  rare_maf_cutoff = 0.01,
  method_cond = c("optimal", "naive"),
  QC_label = "annotation/filter",
  variant_type = c("SNV", "Indel", "variant"),
  geno_missing_imputation = c("mean", "minor"),
  Annotation_dir = "annotation/info/FunctionalAnnotation",
```

```
Annotation_name_catalog,
Annotation_name
)
```

Arguments

category the coding functional category of rare variants to be functionally annotated.

Choices include plof, plof_ds, missense, disruptive_missense, synonymous

(default = plof).

chr chromosome.

genofile an object of opened annotated GDS (aGDS) file.

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel

function in the GENESIS package and transformed using the genesis2staar_nullmodel

function in the STAARpipeline package.

gene_name name of the gene to be annotated.

known_loci the data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

rare_maf_cutoff

the cutoff of maximum minor allele frequency in defining rare variants (default

= 0.01).

method_cond a character value indicating the method for conditional analysis. optimal refers

to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

QC_label channel name of the QC label in the GDS/aGDS file.

variant_type type of variant included in the conditional analysis. Choices include "SNV",

"Indel", or "variant" (default = "SNV").

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default =

"mean").

Annotation_dir channel name of the annotations in the aGDS file

(default = "annotation/info/FunctionalAnnotation").

Annotation_name_catalog

a data frame containing the name and the corresponding channel name in the

aGDS file.

Annotation_name

a vector of qualitative/quantitative annotation names user wants to extract.

Value

a data frame containing the basic information (chromosome, position, reference allele and alternative allele), unconditional and conditional the score test p-values, and annotation scores for the rare variants of the input coding mask.

```
Gene_Centric_Coding_Results_Summary
```

Summarize gene-centric coding analysis results generated by STAARpipeline package and perform conditional analysis for (unconditionally) significant coding masks by adjusting for a given list of known variants

Description

The Gene_Centric_Coding_Results_Summary function takes in the objects of gene-centric coding analysis results generated by STAARpipeline package, the object from fitting the null model, and the set of known variants to be adjusted for in conditional analysis to summarize the gene-centric coding analysis results and analyze the conditional association between a quantitative/dichotomous phenotype and the rare variants in the unconditional significant coding masks.

Usage

```
Gene_Centric_Coding_Results_Summary(
  agds_dir,
  gene_centric_coding_jobs_num,
  input_path,
  output_path,
  gene_centric_results_name,
  obj_nullmodel,
  known_loci = NULL,
  method_cond = c("optimal", "naive"),
  QC_label = "annotation/filter",
  geno_missing_imputation = c("mean", "minor"),
  variant_type = c("SNV", "Indel", "variant"),
  Annotation_dir = "annotation/info/FunctionalAnnotation",
  Annotation_name_catalog,
  Use_annotation_weights = FALSE,
  Annotation_name = NULL,
  alpha = 2.5e-06,
  manhattan_plot = FALSE,
  QQ_plot = FALSE
)
```

Arguments

```
agds_dir file directory of annotated GDS (aGDS) files for all chromosomes (1-22)

gene_centric_coding_jobs_num

the number of gene-centric coding analysis results generated by STAARpipeline
package.

input_path the directory of gene-centric coding analysis results that generated by STAARpipeline
package.

output_path the directory for the output files.

gene_centric_results_name
file name of gene-centric coding analysis results generated by STAARpipeline
package.
```

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel

function in the GENESIS package and transformed using the genesis2staar_nullmodel

function in the STAARpipeline package.

known_loci the data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

method_cond a character value indicating the method for conditional analysis. optimal refers

to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

QC_label channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default =

"mean").

variant_type type of variant included in the analysis. Choices include "SNV", "Indel", or

"variant" (default = "SNV").

Annotation_dir channel name of the annotations in the aGDS file

(default = "annotation/info/FunctionalAnnotation").

Annotation_name_catalog

a data frame containing the name and the corresponding channel name in the

aGDS file.

Use_annotation_weights

use annotations as weights or not (default = FALSE).

Annotation_name

a vector of annotation names used in STAAR (default = NULL).

alpha p-value threshold of significant results (default=2.5E-06).

manhattan_plot output manhattan plot or not (default = FALSE).

QQ_plot output Q-Q plot or not (default = FALSE).

Value

The function returns the following analysis results:

coding_sig.csv: a matrix that summarizes the unconditional significant coding masks detected by STAAR-O (STAAR-O pvalue smaller than the threshold alpha), including gene name ("Gene name"), chromosome ("chr"), coding functional category ("Category"), number of variants ("#SNV"), and unconditional p-values of set-based tests SKAT ("SKAT(1,25)"), Burden ("Burden(1,1)"), ACAT-V ("ACAT-V(1,25)") and STAAR-O ("STAAR-O").

coding_sig_cond.csv: a matrix that summarized the conditional analysis results of unconditional significant coding masks detected by STAAR-O (available if known_loci is not a NULL), including gene name ("Gene name"), chromosome ("chr"), coding functional category ("Category"), number of variants ("#SNV"), and conditional p-values of set-based tests SKAT ("SKAT(1,25)"), Burden ("Burden(1,1)"), ACAT-V ("ACAT-V(1,25)") and STAAR-O ("STAAR-O").

results_plof_genome.Rdata: a matrix contains the STAAR p-values (including STAAR-O) of the coding mask defined by the putative loss of function variants (plof) for all protein-coding genes across the genome.

plof_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant plof masks.

plof_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant plof masks (available if known loci is not a NULL).

results_plof_ds_genome.Rdata: a matrix contains the STAAR p-values (including STAAR-O) of the coding mask defined by the putative loss of function variants and disruptive missense variants (plof_ds) for all protein-coding genes across the genome.

plof_ds_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant plof ds masks.

plof_ds_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant plof_ds masks (available if known_loci is not a NULL).

results_disruptive_missense_genome.Rdata: a matrix contains the STAAR p-values (including STAAR-O) of the coding mask defined by the disruptive missense variants (disruptive_missense) for all protein-coding genes across the genome.

disruptive_missense_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant disruptive_missense masks.

disruptive_missense_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant disruptive_missense masks (available if known_loci is not a NULL).

results_missense_genome.Rdata: a matrix contains the STAAR p-values (including STAAR-O) of the coding mask defined by the missense variants (missense) for all protein-coding genes across the genome.

missense_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant missense masks.

missense_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant missense masks (available if known_loci is not a NULL).

results_synonymous_genome.Rdata: a matrix contains the STAAR p-values (including STAAR-O) of the coding mask defined by the synonymous variants (synonymous) for all protein-coding genes across the genome.

synonymous_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant synonymous masks.

synonymous_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant synonymous masks (available if known_loci is not a NULL).

Gene_Centric_Noncoding_Info

Functionally annotate rare variants in a noncoding mask

Description

The Gene_Centric_Noncoding_Info function takes in a noncoding mask of a gene to functionally annotate the rare variants in the mask.

Usage

```
Gene_Centric_Noncoding_Info(
  category = c("downstream", "upstream", "UTR", "promoter_CAGE", "promoter_DHS",
        "enhancer_CAGE", "enhancer_DHS", "ncRNA"),
    chr,
```

```
genofile,
obj_nullmodel,
gene_name,
known_loci = NULL,
rare_maf_cutoff = 0.01,
method_cond = c("optimal", "naive"),
QC_label = "annotation/filter",
variant_type = c("SNV", "Indel", "variant"),
geno_missing_imputation = c("mean", "minor"),
Annotation_dir = "annotation/info/FunctionalAnnotation",
Annotation_name_catalog,
Annotation_name
```

Arguments

category the noncoding functional category to be functionally annotated. Choices include

downstream, upstream, UTR, promoter_CAGE, promoter_DHS, enhancer_CAGE,

enhancer_DHS, ncRNA (default = downstream).

chr chromosome.

genofile an object of opened annotated GDS (aGDS) file.

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel

 $function\ in\ the\ {\tt GENESIS}\ package\ and\ transformed\ using\ the\ {\tt genesis2staar_nullmodel}$

function in the STAARpipeline package.

gene_name name of the gene to be annotated.

known_loci the data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

rare_maf_cutoff

the cutoff of maximum minor allele frequency in defining rare variants (default

= 0.01).

method_cond a character value indicating the method for conditional analysis. optimal refers

to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

QC_label channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").

variant_type type of variant included in the conditional analysis. Choices include "SNV",

"Indel", or "variant" (default = "SNV").

 ${\tt geno_missing_imputation}$

method of handling missing genotypes. Either "mean" or "minor" (default = "mean").

Annotation_dir channel name of the annotations in the aGDS file

(default = "annotation/info/FunctionalAnnotation").

Annotation_name_catalog

a data frame containing the name and the corresponding channel name in the aGDS file.

Annotation_name

a vector of qualitative/quantitative annotation names user wants to extract.

Value

a data frame containing the basic information (chromosome, position, reference allele and alternative allele), unconditional and conditional the score test p-values, and annotation scores for the rare variants of the input noncoding mask.

```
Gene_Centric_Noncoding_Results_Summary

Summarize gene-centric noncoding analysis results generated by

STAARpipeline package
```

Description

The Gene_Centric_Noncoding_Results_Summary function takes in the objects of gene-centric noncoding analysis results generated by STAARpipeline package, the object from fitting the null model, and the set of known variants to be adjusted for in conditional analysis to summarize the gene-centric noncoding analysis results and analyze the conditional association between a quantitative/dichotomous phenotype and the rare variants in the unconditional significant noncoding masks.

Usage

```
Gene_Centric_Noncoding_Results_Summary(
  agds_dir,
  gene_centric_noncoding_jobs_num,
  input_path,
 output_path,
  gene_centric_results_name,
  ncRNA_jobs_num,
 ncRNA_input_path,
 ncRNA_output_path,
  ncRNA_results_name,
  obj_nullmodel,
  known_loci = NULL,
 method_cond = c("optimal", "naive"),
 QC_label = "annotation/filter",
  geno_missing_imputation = c("mean", "minor"),
  variant_type = c("SNV", "Indel", "variant"),
  Annotation_dir = "annotation/info/FunctionalAnnotation",
  Annotation_name_catalog,
  Use_annotation_weights = FALSE,
  Annotation_name = NULL,
 alpha = 2.5e-06,
 manhattan_plot = FALSE,
  QQ_plot = FALSE
)
```

Arguments

```
agds_dir a data farme containing directory of GDS/aGDS files. gene_centric_noncoding_jobs_num
```

the number of results for gene-centric noncoding analysis of protein-coding genes generated by STAARpipeline package.

input_path the directory of gene-centric noncoding analysis results for protein-coding genes

that generated by STAARpipeline package.

output_path the directory for the output files of the summary of gene-centric noncoding anal-

ysis results for protein-coding genes.

gene_centric_results_name

the file name of gene-centric noncoding analysis results for protein-coding genes generated by STAARpipeline package.

ncRNA_jobs_num the number of results for gene-centric noncoding analysis of ncRNA genes gen-

erated by STAARpipeline package..

 $ncRNA_input_path$

the directory of gene-centric noncoding analysis results for ncRNA genes that generated by STAARpipeline package.

ncRNA_output_path

the directory for the output files of the summary of gene-centric noncoding analysis results for ncRNA genes.

ncRNA_results_name

file name of gene-centric noncoding analysis results for ncRNA genes that generated by STAARpipeline package.

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel function in the GENESIS package and transformed using the genesis2staar_nullmodel

function in the STAARpipeline package.

known_loci the data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

method_cond a character value indicating the method for conditional analysis. optimal refers

to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

QC_label channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default =

"mean").

variant_type type of variant included in the analysis. Choices include "SNV", "Indel", or

"variant" (default = "SNV").

Annotation_dir channel name of the annotations in the aGDS file

(default = "annotation/info/Functional Annotation").

Annotation_name_catalog

a data frame containing the name and the corresponding channel name in the

aGDS file.

 ${\tt Use_annotation_weights}$

use annotations as weights or not (default = FALSE).

Annotation_name

a vector of annotation names used in STAAR (default = NULL).

alpha p-value threshold of significant results (default=2.5E-06).

manhattan_plot output manhattan plot or not (default = FALSE).

QQ_plot output Q-Q plot or not (default = FALSE).

Value

The function returns the following analysis results:

noncoding_sig.csv: a matrix that summarized the unconditional significant region detected by STAAR-O (STAAR-O pvalue smaller than the threshold alpha), including gene name ("Gene name"), chromosome ("chr"), coding functional category ("Category"), number of variants ("#SNV"), and the unconditional STAAR p-values (including STAAR-O).

noncoding_sig_cond.csv: a matrix that summarized the conditional analysis results of the unconditional significant region detected by STAAR-O (available if known_loci is not a NULL), including gene name ("Gene name"), chromosome ("chr"), coding functional category ("Category"), number of variants ("#SNV"), and the conditional STAAR p-values (including STAAR-O).

results_UTR_genome: a matrix contains the STAAR p-values (including STAAR-O) of the non-coding masks defined by UTR variants (UTR) for all protein-coding genes across the genome.

UTR_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant UTR masks.

UTR_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant UTR masks (available if known_loci is not a NULL).

results_upstream_genome: a matrix contains the STAAR p-values (including STAAR-O) of the noncoding masks defined by upstream variants (upstream) for all protein-coding genes across the genome.

upstream_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant upstream masks.

upstream_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant upstream masks (available if known_loci is not a NULL).

results_downstream_genome: a matrix contains the STAAR p-values (including STAAR-O) of the noncoding masks defined by downstream variants (downstream) for all protein-coding genes across the genome.

downstream_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant downstream masks.

downstream_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant downstream masks (available if known loci is not a NULL).

results_promoter_CAGE_genome: a matrix contains the STAAR p-values (including STAAR-O) of the noncoding masks defined by variants overlaid with CAGE sites in the promoter (promoter_CAGE) for all protein-coding genes across the genome.

promoter_CAGE_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant promoter CAGE masks.

promoter_CAGE_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant promoter_CAGE masks (available if known_loci is not a NULL).

results_promoter_DHS_genome: a matrix contains the STAAR p-values (including STAAR-O) of the noncoding masks defined by variants overlaid with DHS sites in the promoter (promoter_DHS) for all protein-coding genes across the genome.

promoter_DHS_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant promoter_DHS masks.

promoter_DHS_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant promoter_DHS masks (available if known_loci is not a NULL).

results_enhancer_CAGE_genome: a matrix contains the STAAR p-values (including STAAR-O) of the noncoding masks defined by variants overlaid with CAGE sites in the enhancer (enhancer_CAGE) for all protein-coding genes across the genome.

enhancer_CAGE_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant enhancer_CAGE masks.

enhancer_CAGE_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant enhancer_CAGE masks (available if known_loci is not a NULL).

results_enhancer_DHS_genome: a matrix contains the STAAR p-values (including STAAR-O) of the noncoding masks defined by variants overlaid with DHS sites in the enhancer (enhancer_DHS) for all protein-coding genes across the genome.

enhancer_DHS_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant enhancer_DHS masks.

enhancer_DHS_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant enhancer_DHS masks (available if known_loci is not a NULL).

results_ncRNA_genome: a matrix contains the STAAR p-values (including STAAR-O) of the non-coding masks defined by exonic and splicing ncRNA variants (ncRNA) for all ncRNA genes across the genome.

ncRNA_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the unconditional significant ncRNA masks.

ncRNA_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the unconditional significant ncRNA masks (available if known_loci is not a NULL).

```
\label{lem:continuous} Individual\_Analysis\_Results\_Summary \\ Summarize individual-variant analysis results generated by \\ {\tt STAARpipeline} \ package
```

Description

The Individual_Analysis_Results_Summary function takes in the objects of individual analysis results generated by STAARpipeline package, the object from fitting the null model, and the set of known variants to be adjusted for in conditional analysis to summarize the individual analysis results and analyze the conditional association between a quantitative/dichotomous phenotype and the unconditional significant single variants.

Usage

```
Individual_Analysis_Results_Summary(
   agds_dir,
   jobs_num,
   input_path,
   output_path,
   individual_results_name,
   obj_nullmodel,
   known_loci = NULL,
   method_cond = c("optimal", "naive"),
   QC_label = "annotation/filter",
```

```
geno_missing_imputation = c("mean", "minor"),
alpha = 5e-09,
manhattan_plot = FALSE,
QQ_plot = FALSE
)
```

Arguments

agds_dir a data farme containing directory of GDS/aGDS files.

jobs_num a data frame containing the number of analysis results, including the number of

individual analysis results, the number of sliding window analysis results, and

the number of dynamic window analysis results.

input_path the directory of individual analysis results that generated by STAARpipeline

package.

output_path the directory for the output files.

individual_results_name

the file name of individual analysis results generated by STAARpipeline pack-

age.

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel

function in the GENESIS package and transformed using the genesis2staar_nullmodel

function in the STAARpipeline package.

known_loci the data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

method_cond a character value indicating the method for conditional analysis. optimal refers

to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

QC_label channel name of the QC label in the GDS/aGDS file.

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default =

"mean").

alpha p-value threshold of significant results (default=5E-09).

manhattan_plot output manhattan plot or not (default = FALSE).

QQ_plot output Q-Q plot or not (default = FALSE).

Value

The function returns the following analysis results:

results_individual_analysis_genome.Rdata: a matrix contains the score test p-value and effect size estimation of each variant across the genome.

results_individual_analysis_sig.Rdata and results_individual_analysis_sig.csv: a matrix contains the score test p-values and effect size estimations of significant results (p-value < alpha).

results_sig_cond.Rdata and results_sig_cond.csv: a matrix contains the conditional score test p-values for each significant variant (available if known_loci is not a NULL).

```
Single_Variants_List_Analysis
```

Calculate individual-variant p-values of a list of variants

Description

The Single_Variants_List_Analysis function takes in a list of variants to calculate the p-values and effect sizes of the input variants

Usage

```
Single_Variants_List_Analysis(
   agds_dir,
   single_variants_list,
   obj_nullmodel,
   QC_label = "annotation/filter",
   geno_missing_imputation = c("mean", "minor")
)
```

Arguments

agds_dir file directory of annotated GDS (aGDS) files for all chromosomes (1-22). single_variants_list

name a data frame containing the information of variants to be functionally annotated. The data frame must include 4 columns with the following names: "CHR" (chromosome number), "POS" (position), "REF" (reference allele), and "ALT" (alternative allele).

obj_nullmodel

an object from fitting the null model, which is either the output from fit_nullmodel function in the STAARpipeline package, or the output from fitNullModel function in the GENESIS package and transformed using the genesis2staar_nullmodel function in the STAARpipeline package.

QC_label channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default = "mean").

Value

a data frame containing the basic information (chromosome, position, reference allele and alternative allele) the score test p-values, and the effect sizes for the input variants.

Sliding_Window_Info Functionally annotate rare variants in a genetic region

Description

The Sliding_Window_Info function takes in the location of a genetic region to functionally annotate the rare variants in the region.

Usage

```
Sliding_Window_Info(
  chr,
  genofile,
  obj_nullmodel,
  start_loc,
  end_loc,
  known_loci = NULL,
  rare_maf_cutoff = 0.01,
  method_cond = c("optimal", "naive"),
  QC_label = "annotation/filter",
  variant_type = c("SNV", "Indel", "variant"),
  geno_missing_imputation = c("mean", "minor"),
  Annotation_dir = "annotation/info/FunctionalAnnotation",
  Annotation_name_catalog,
  Annotation_name
)
```

Arguments

chr	chromosome.	
genofile	an object of opened annotated GDS (aGDS) file.	
obj_nullmodel	an object from fitting the null model, which is either the output from fit_nullmodel function in the STAARpipeline package, or the output from fitNullModel function in the GENESIS package and transformed using the genesis2staar_nullmodel function in the STAARpipeline package.	
start_loc	starting location (position) of the genetic region to be annotated.	
end_loc	ending location (position) of the genetic region to be annotated.	
known_loci	the data frame of variants to be adjusted for in conditional analysis and should contain 4 columns in the following order: chromosome (CHR), position (POS), reference allele (REF), and alternative allele (ALT) (default = NULL).	
rare_maf_cutoff		
	the cutoff of maximum minor allele frequency in defining rare variants (default $= 0.01$).	
method_cond	a character value indicating the method for conditional analysis. optimal refers to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and taking the residuals (default = optimal).	
QC_label	channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").	

Value

a data frame containing the basic information (chromosome, position, reference allele and alternative allele), unconditional and conditional the score test p-values, and annotation scores for the input variants.

```
Sliding\_Window\_Results\_Summary \\ Summarize \ the \ sliding \ window \ analysis \ results \ generated \ by \\ STAARpipeline \ package
```

Description

The Sliding_Window_Results_Summary function takes in the results of sliding window analysis, the object from fitting the null model, and the set of known variants to be adjusted for in conditional analysis to summarize the sliding window analysis results and analyze the conditional association between a quantitative/dichotomous phenotype and the rare variants in the unconditional significant genetic region.

Usage

```
Sliding_Window_Results_Summary(
  agds_dir,
  jobs_num,
  input_path,
  output_path,
  sliding_window_results_name,
  obj_nullmodel,
  known_loci = NULL,
 method_cond = c("optimal", "naive"),
 QC_label = "annotation/filter",
 geno_missing_imputation = c("mean", "minor"),
  variant_type = c("SNV", "Indel", "variant"),
  Annotation_dir = "annotation/info/FunctionalAnnotation",
  Annotation_name_catalog,
 Use_annotation_weights = FALSE,
  Annotation_name = NULL,
```

```
alpha = 0.05,
manhattan_plot = FALSE,
QQ_plot = FALSE
)
```

Arguments

agds_dir file directory of annotated GDS (aGDS) files for all chromosomes (1-22).

jobs_num a data frame containing the number of jobs for association analysis. The data

frame must include a column with the name "sliding_window_num"

input_path file directory of the sliding window analysis results.

output_path file output directory of the summary results.

sliding_window_results_name

the file name of the input sliding window analysis results.

obj_nullmodel an object from fitting the null model, which is either the output from fit_nullmodel

function in the STAARpipeline package, or the output from fitNullModel

function in the GENESIS package and transformed using the genesis2staar_nullmodel

function in the STAARpipeline package.

known_loci the data frame of variants to be adjusted for in conditional analysis and should

contain 4 columns in the following order: chromosome (CHR), position (POS),

reference allele (REF), and alternative allele (ALT) (default = NULL).

method_cond a character value indicating the method for conditional analysis. optimal refers

to regressing residuals from the null model on known_loci as well as all covariates used in fitting the null model (fully adjusted) and taking the residuals; naive refers to regressing residuals from the null model on known_loci and

taking the residuals (default = optimal).

QC_label channel name of the QC label in the GDS/aGDS file (default = "annotation/filter").

geno_missing_imputation

method of handling missing genotypes. Either "mean" or "minor" (default =

"mean").

variant_type variants include in the conditional analysis. Choices include "variant", "SNV",

or "Indel" (default = "SNV").

Annotation_dir channel name of the annotations in the aGDS file

(default = "annotation/info/FunctionalAnnotation").

Annotation_name_catalog

a data frame containing the name and the corresponding channel name in the

aGDS file.

Use_annotation_weights

use annotations as weights or not (default = FALSE).

Annotation_name

a vector of annotation names used in STAAR (default = NULL).

alpha threshod to control the genome-wise (family-wise) error rate (default = 0.05),

the p-value threshold is alpha/total number of sliding windows

 $manhattan_plot \quad output \ manhattan \ plot \ or \ not \ (default = FALSE).$

QQ_plot output Q-Q plot or not (default = FALSE).

Value

The function returns the following analysis results:

results_sliding_window_genome.Rdata: a matrix contains the STAAR p-values (including STAAR-O) of the sliding windows across the genome.

sliding_window_sig.Rdata and sliding_window_sig.csv: a matrix contains the unconditional STAAR p-values (including STAAR-O) of the significant sliding windows (unconditional p-value<alpha/total number of sliding windows).

sliding_window_sig_cond.Rdata and sliding_window_sig_cond.csv: a matrix contains the conditional STAAR p-values (including STAAR-O) of the significant sliding windows (available if known_loci is not a NULL).

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