Package 'gwasurvivr'

July 29, 2025

Type Package

Title gwasurvivr: an R package for genome wide survival analysis

Version 1.27.0

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Description gwasurvivr is a package to perform survival analysis using Cox proportional hazard models on imputed genetic data.

VignetteBuilder knitr

Imports GWASTools, survival, VariantAnnotation, parallel, matrixStats, SummarizedExperiment, stats, utils, SNPRelate

Suggests BiocStyle, knitr, rmarkdown

Depends R (>= 3.4.0)

License Artistic-2.0

URL https://github.com/suchestoncampbelllab/gwasurvivr

biocViews GenomeWideAssociation, Survival, Regression, Genetics, SNP, GeneticVariability, Pharmacogenomics, BiomedicalInformatics

RoxygenNote 6.1.0 Encoding UTF-8 git_url https://git.bioconductor.org/packages/gwasurvivr git_branch devel git_last_commit_ceda955 git_last_commit_date 2025-04-15 Repository Bioconductor 3.22

Date/Publication 2025-07-29

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gdsCoxSurv

Description

Performs survival analysis using Cox proportional hazard models on imputed genetic data from GDS files.

Usage

```
gdsCoxSurv(gdsfile, covariate.file, id.column, sample.ids = NULL,
    time.to.event, event, covariates, inter.term = NULL,
    print.covs = "only", out.file, chunk.size = 5000,
    maf.filter = 0.05, flip.dosage = TRUE, verbose = TRUE,
    clusterObj = NULL)
```

| gdsfile | path to .gds file. Location of the .gds file should also contain .snp.rdata and .scan.rdata files. |
|----------------|--|
| covariate.file | data.frame comprising phenotype information, all covariates to be added in the model must be numeric. |
| id.column | character giving the name of the ID column in covariate.file. |
| sample.ids | character vector of sample IDs to keep in survival analysis |
| time.to.event | character of column name in covariate.file that represents the time interval of interest in the analysis |
| event | character of column name in covariate.file that represents the event of interest to be included in the analysis |
| covariates | character vector with exact names of columns in covariate.file to include in anal- ysis |
| inter.term | character string giving the column name of the covariate that will be added to the interaction term with SNP (e.g. term*SNP). See details. |
| print.covs | character string of either "only", "all" or "some", defining which covariate statistics should be printed to the output. See details. |
| out.file | character of output file name (do not include extension) |
| chunk.size | integer number of variants to process per thread |
| maf.filter | numeric to filter minor allele frequency (i.e. choosing 0.05 means filtering MAF>0.05). User can set this to NULL if no filtering is preffered. Default is 0.05. |
| flip.dosage | $logical \ to \ flip \ which \ allele \ the \ dosage \ was \ calculated \ on, \ default \ flip \ dosage=TRUE$ |
| verbose | logical for messages that describe which part of the analysis is currently being run |
| cluster0bj | A cluster object that can be used with the parApply function. See details. |

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Details

Testing for SNP-covariate interactions: User can define the column name of the covariate that will be included in the interaction term. For example, for given covariates a and b, where c is defined as the inter.term the model will be: a + b + c + SNP + c + SNP.

Printing results of other covariates: print.covs argument controls the number of covariates will be printed as output. The function is set to only by default and will only print the SNP or if an interaction term is given, the results of the interaction term (e.g. SNP*covariate). Whereas, all will print results (coef, se.coef, p.value etc) of all covariates included in the model. some is only applicable if an interaction term is given and will print the results for SNP, covariate tested for interaction and the interaction term. User should be mindful about using the all option, as it will likely slow down the analysis and will increase the output file size.

User defined parallelization: This function uses parApply from parallel package to fit models to SNPs in parallel. User is not required to set any options for the parallelization. However, advanced users who wish to optimize it, can provide a cluster object generated by makeCluster family of functions that suits their need and platform.

Value

Saves two text files directly to disk: .coxph extension containing CoxPH survival analysis results. .snps_removed extension containing SNPs that were removed due to low variance or user-defined thresholds.

```
gdsfile <- system.file(package="gwasurvivr",</pre>
                        "extdata",
                        "gds_example.gds")
covariate.file <- system.file(package="gwasurvivr",</pre>
                                "extdata",
                                "simulated_pheno.txt")
covariate.file <- read.table(covariate.file,</pre>
                              sep=" ",
                               header=TRUE,
                              stringsAsFactors = FALSE)
covariate.file$SexFemale <- ifelse(covariate.file$sex=="female", 1L, 0L)</pre>
sample.ids <- covariate.file[covariate.file$group=="experimental",]$ID_2</pre>
gdsCoxSurv(gdsfile=gdsfile,
           covariate.file=covariate.file,
           id.column="ID_2",
           sample.ids=sample.ids,
           time.to.event="time",
           event="event",
           covariates=c("age", "SexFemale", "DrugTxYes"),
           inter.term=NULL,
           print.covs="only",
           out.file="impute_example",
           chunk.size=50,
           maf.filter=0.005,
           flip.dosage=TRUE,
           verbose=TRUE,
           clusterObj=NULL)
```

impute2CoxSurv

Description

Performs survival analysis using Cox proportional hazard models on imputed genetic data from IMPUTE2 output

Usage

```
impute2CoxSurv(impute.file, sample.file, chr, covariate.file, id.column,
  sample.ids = NULL, time.to.event, event, covariates,
  inter.term = NULL, print.covs = "only", out.file,
  chunk.size = 10000, maf.filter = 0.05, exclude.snps = NULL,
  flip.dosage = TRUE, verbose = TRUE, clusterObj = NULL,
  keepGDS = FALSE)
```

| <pre>impute.file</pre> | character of IMPUTE2 file |
|------------------------|--|
| sample.file | character of sample file affiliated with IMPUTE2 file |
| chr | numeric denoting chromosome number |
| covariate.file | data.frame comprising phenotype information, all covariates to be added in the model must be numeric. |
| id.column | character giving the name of the ID column in covariate.file. |
| sample.ids | character vector of sample IDs to keep in survival analysis |
| time.to.event | character of column name in covariate.file that represents the time interval of interest in the analysis |
| event | character of column name in covariate.file that represents the event of interest to be included in the analysis |
| covariates | character vector with exact names of columns in covariate.file to include in anal- ysis |
| inter.term | character string giving the column name of the covariate that will be added to the interaction term with SNP (e.g. term*SNP). See details. |
| print.covs | character string of either "only", "all" or "some", defining which covariate statistics should be printed to the output. See details. |
| out.file | character of output file name (do not include extension) |
| chunk.size | integer number of variants to process per thread |
| maf.filter | numeric to filter minor allele frequency (i.e. choosing 0.05 means filtering MAF>0.05). User can set this to NULL if no filtering is preffered. Default is 0.05. |
| exclude.snps | a character vector listing the rsIDs of SNPs that will be excluded from analyses |
| flip.dosage | logical to flip which allele the dosage was calculated on, default flip.dosage=TRUE |
| verbose | logical for messages that describe which part of the analysis is currently being run |
| cluster0bj | A cluster object that can be used with the parApply function. See details. |
| keepGDS | logical to keep GDS files (compressed IMPUTE2 files) after the analysis. Defaults to FALSE. |

Details

Testing for SNP-covariate interactions: User can define the column name of the covariate that will be included in the interaction term. For example, for given covariates a and b, where c is defined as the inter.term the model will be: $\sim a + b + c + SNP + c + SNP$.

Printing results of other covariates: print.covs argument controls the number of covariates will be printed as output. The function is set to only by default and will only print the SNP or if an interaction term is given, the results of the interaction term (e.g. SNP*covariate). Whereas, all will print results (coef, se.coef, p.value etc) of all covariates included in the model. some is only applicable if an interaction term is given and will print the results for SNP, covariate tested for interaction and the interaction term. User should be mindful about using the all option, as it will likely slow down the analysis and will increase the output file size.

User defined parallelization: This function uses parApply from parallel package to fit models to SNPs in parallel. User is not required to set any options for the parallelization. However, advanced users who wish to optimize it, can provide a cluster object generated by makeCluster family of functions that suits their need and platform.

Value

Saves two text files directly to disk: .coxph extension containing CoxPH survival analysis results. .snps_removed extension containing SNPs that were removed due to low variance or user-defined thresholds.

```
impute.file <- system.file(package="gwasurvivr",</pre>
                             "extdata",
                            "impute_example.impute2.gz")
sample.file <- system.file(package="gwasurvivr",</pre>
                             "extdata",
                            "impute_example.impute2_sample")
covariate.file <- system.file(package="gwasurvivr",</pre>
                                "extdata",
                                "simulated_pheno.txt")
covariate.file <- read.table(covariate.file,</pre>
                               sep=" ".
                               header=TRUE,
                               stringsAsFactors = FALSE)
covariate.file$SexFemale <- ifelse(covariate.file$sex=="female", 1L, 0L)</pre>
sample.ids <- covariate.file[covariate.file$group=="experimental",]$ID_2</pre>
impute2CoxSurv(impute.file=impute.file,
               sample.file=sample.file,
               chr=14,
               covariate.file=covariate.file,
               id.column="ID_2",
               sample.ids=sample.ids,
               time.to.event="time",
               event="event",
               covariates=c("age", "SexFemale", "DrugTxYes"),
               inter.term=NULL,
               print.covs="only",
               out.file="impute_example",
               chunk.size=50,
               maf.filter=0.005,
               exclude.snps=NULL,
```

flip.dosage=TRUE, verbose=TRUE, clusterObj=NULL, keepGDS=FALSE)

michiganCoxSurv

Fit Cox survival to all variants in a .vcf.gz file from Michigan imputation server

Description

Performs survival analysis using Cox proportional hazard models on imputed genetic data stored in compressed VCF files

Usage

```
michiganCoxSurv(vcf.file, covariate.file, id.column, sample.ids = NULL,
    time.to.event, event, covariates, inter.term = NULL,
    print.covs = "only", out.file, maf.filter = 0.05, r2.filter = NULL,
    chunk.size = 5000, verbose = TRUE, clusterObj = NULL)
```

| vcf.file | character(1) path to VCF file. |
|----------------|---|
| covariate.file | matrix(1) comprising phenotype (time, event) and additional covariate data. |
| id.column | character(1) providing exact match to sample ID column from covariate.file |
| sample.ids | character vector with sample ids to include in analysis |
| time.to.event | character(1) string that matches time column name in pheno.file |
| event | character(1) string that matches event column name in pheno.file |
| covariates | character vector with matching column names in pheno.file of covariates of in- terest |
| inter.term | character(1) string giving the column name of the covariate that will be added to the interaction term with SNP (e.g. term*SNP). See details. |
| print.covs | character(1) string of either "only", "all" or "some", defining which covariate statistics should be printed to the output. See details. |
| out.file | character(1) string with output name |
| maf.filter | integer(1) filter out minor allele frequency below threshold (i.e. 0.005 will filter MAF > 0.005) |
| r2.filter | integer(1) of imputation quality score filter (i.e. 0.7 will filter r2 > 0.7) |
| chunk.size | integer(1) number of variants to process per thread |
| verbose | logical(1) for messages that describe which part of the analysis is currently being run |
| cluster0bj | A cluster object that can be used with the parApply function. See details. |

Details

Testing for SNP-covariate interactions: User can define the column name of the covariate that will be included in the interaction term. For example, for given covariates a and b, where c is defined as the inter.term the model will be: a + b + c + SNP + c + SNP.

Printing results of other covariates: print.covs argument controls the number of covariates will be printed as output. The function is set to only by default and will only print the SNP or if an interaction term is given, the results of the interaction term (e.g. SNP*covariate). Whereas, all will print results (coef, se.coef, p.value etc) of all covariates included in the model. some is only applicable if an interaction term is given and will print the results for SNP, covariate tested for interaction and the interaction term. User should be mindful about using the all option, as it will likely slow down the analysis and will increase the output file size.

User defined parallelization: This function uses parApply from parallel package to fit models to SNPs in parallel. User is not required to set any options for the parallelization. However, advanced users who wish to optimize it, can provide a cluster object generated by makeCluster family of functions that suits their need and platform.

Value

Saves two text files directly to disk: .coxph extension containing CoxPH survival analysis results. .snps_removed extension containing SNPs that were removed due to low variance or user-defined thresholds.

```
vcf.file <- system.file(package="gwasurvivr",</pre>
                          "extdata",
                      "michigan.chr14.dose.vcf.gz")
pheno.fl <- system.file(package="gwasurvivr",</pre>
                          "extdata",
                      "simulated_pheno.txt")
pheno.file <- read.table(pheno.fl,</pre>
                           sep=" ".
                          header=TRUE,
                          stringsAsFactors = FALSE)
pheno.file$SexFemale <- ifelse(pheno.file$sex=="female", 1L, 0L)</pre>
sample.ids <- pheno.file[pheno.file$group=="experimental",]$ID_2</pre>
michiganCoxSurv(vcf.file=vcf.file,
               covariate.file=pheno.file,
               id.column="ID_2",
               sample.ids=sample.ids,
               time.to.event="time",
               event="event",
               covariates=c("age", "SexFemale", "DrugTxYes"),
               inter.term=NULL,
               print.covs="only",
               out.file="michigan_example",
               r2.filter=0.3,
               maf.filter=0.005,
               chunk.size=50,
               verbose=TRUE,
               clusterObj=NULL)
```

```
plinkCoxSurv
```

Description

Performs survival analysis using Cox proportional hazard models on directly typed data in PLINK format

Usage

```
plinkCoxSurv(bed.file, covariate.file, id.column, sample.ids = NULL,
    time.to.event, event, covariates, inter.term = NULL,
    print.covs = "only", out.file, chunk.size = 10000,
    maf.filter = 0.005, flip.dosage = TRUE, verbose = TRUE,
    clusterObj = NULL)
```

| bed.file | character of name of plink files without extension |
|----------------|--|
| covariate.file | data.frame comprising phenotype information, all covariates to be added in the model must be numeric. |
| id.column | character giving the name of the ID column in covariate.file. |
| sample.ids | character vector of sample IDs to keep in survival analysis |
| time.to.event | character of column name in covariate.file that represents the time interval of interest in the analysis |
| event | character of column name in covariate.file that represents the event of interest to be included in the analysis |
| covariates | character vector with exact names of columns in covariate.file to include in anal- ysis |
| inter.term | character string giving the column name of the covariate that will be added to the interaction term with SNP (e.g. term*SNP). See details. |
| print.covs | character string of either "only", "all" or "some", defining which covariate statistics should be printed to the output. See details. |
| out.file | character of output file name (do not include extension) |
| chunk.size | integer number of variants to process per thread |
| maf.filter | numeric to filter minor allele frequency (i.e. choosing 0.05 means filtering MAF>0.05). User can set this to NULL if no filtering is preffered. Default is 0.05. |
| flip.dosage | $logical \ to \ flip \ which \ allele \ the \ dosage \ was \ calculated \ on, \ default \ flip \ dosage=TRUE$ |
| verbose | logical for messages that describe which part of the analysis is currently being run |
| cluster0bj | A cluster object that can be used with the parApply function. See details. |

plinkCoxSurv

Details

Testing for SNP-covariate interactions: User can define the column name of the covariate that will be included in the interaction term. For example, for given covariates a and b, where c is defined as the inter.term the model will be: a + b + c + SNP + c + SNP.

Printing results of other covariates: print.covs argument controls the number of covariates will be printed as output. The function is set to only by default and will only print the SNP or if an interaction term is given, the results of the interaction term (e.g. SNP*covariate). Whereas, all will print results (coef, se.coef, p.value etc) of all covariates included in the model. some is only applicable if an interaction term is given and will print the results for SNP, covariate tested for interaction and the interaction term. User should be mindful about using the all option, as it will likely slow down the analysis and will increase the output file size.

User defined parallelization: This function uses parApply from parallel package to fit models to SNPs in parallel. User is not required to set any options for the parallelization. However, advanced users who wish to optimize it, can provide a cluster object generated by makeCluster family of functions that suits their need and platform.

Value

Saves two text files directly to disk: .coxph extension containing CoxPH survival analysis results. .snps_removed extension containing SNPs that were removed due to low variance or user-defined thresholds.

```
bed.file <- system.file(package="gwasurvivr",</pre>
                        "extdata",
                        "plink_example.bed")
covariate.file <- system.file(package="gwasurvivr",</pre>
                                "extdata",
                                "simulated_pheno.txt")
covariate.file <- read.table(covariate.file,</pre>
                               sep=" ",
                               header=TRUE,
                               stringsAsFactors = FALSE)
covariate.file$SexFemale <- ifelse(covariate.file$sex=="female", 1L, 0L)</pre>
sample.ids <- covariate.file[covariate.file$group=="experimental",]$ID_2</pre>
plinkCoxSurv(bed.file=bed.file,
             covariate.file=covariate.file,
             id.column="ID_2",
             sample.ids=sample.ids,
             time.to.event="time",
             event="event",
             covariates=c("age", "SexFemale", "DrugTxYes"),
             inter.term=NULL,
             print.covs="only",
             out.file="impute_example",
             chunk.size=50,
             maf.filter=0.005,
             flip.dosage=TRUE,
             verbose=TRUE,
             clusterObj=NULL)
```

sangerCoxSurv

Description

Performs survival analysis using Cox proportional hazard models on imputed genetic data stored in compressed VCF files.

Usage

```
sangerCoxSurv(vcf.file, covariate.file, id.column, sample.ids = NULL,
    time.to.event, event, covariates, inter.term = NULL,
    print.covs = "only", out.file, maf.filter = 0.05,
    info.filter = NULL, chunk.size = 5000, verbose = TRUE,
    clusterObj = NULL)
```

Arguments

| vcf.file | character(1) path to VCF file. |
|--------------------------|---|
| covariate.file | matrix(1) comprising phenotype (time, event) and additional covariate data. |
| id.column | character(1) providing exact match to sample ID column from covariate.file |
| sample.ids | character vector with sample ids to include in analysis |
| <pre>time.to.event</pre> | character(1) string that matches time column name in pheno.file |
| event | character(1) string that matches event column name in pheno.file |
| covariates | character vector with matching column names in pheno.file of covariates of in- terest |
| inter.term | character(1) string giving the column name of the covariate that will be added to the interaction term with SNP (e.g. term*SNP). See details. |
| print.covs | character(1) string of either "only", "all" or "some", defining which covariate statistics should be printed to the output. See details. |
| out.file | character(1) string with output name |
| maf.filter | integer(1) filter out minor allele frequency below threshold (i.e. 0.005 will filter $MAF > 0.005$) |
| info.filter | integer(1) of imputation quality score filter (i.e. 0.7 will filter info > 0.7) |
| chunk.size | integer(1) number of variants to process per thread |
| verbose | logical(1) for messages that describe which part of the analysis is currently being run |
| cluster0bj | A cluster object that can be used with the parApply function. See details. |

Details

Testing for SNP-covariate interactions: User can define the column name of the covariate that will be included in the interaction term. For example, for given covariates a and b, where c is defined as the inter.term the model will be: $\sim a + b + c + SNP + c + SNP$.

Printing results of other covariates: print.covs argument controls the number of covariates will be printed as output. The function is set to only by default and will only print the SNP or if an

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interaction term is given, the results of the interaction term (e.g. SNP*covariate). Whereas, all will print results (coef, se.coef, p.value etc) of all covariates included in the model. some is only applicable if an interaction term is given and will print the results for SNP, covariate tested for interaction and the interaction term. User should be mindful about using the all option, as it will likely slow down the analysis and will increase the output file size.

User defined parallelization: This function uses parApply from parallel package to fit models to SNPs in parallel. User is not required to set any options for the parallelization. However, advanced users who wish to optimize it, can provide a cluster object generated by makeCluster family of functions that suits their need and platform.

Value

Saves two text files directly to disk: .coxph extension containing CoxPH survival analysis results. .snps_removed extension containing SNPs that were removed due to low variance or user-defined thresholds.

```
vcf.file <- system.file(package="gwasurvivr",</pre>
                         "extdata"
                         "sanger.pbwt_reference_impute.vcf.gz")
pheno.fl <- system.file(package="gwasurvivr",</pre>
                          "extdata",
                      "simulated_pheno.txt")
pheno.file <- read.table(pheno.fl,</pre>
                          sep=" ",
                          header=TRUE,
                          stringsAsFactors = FALSE)
pheno.file$SexFemale <- ifelse(pheno.file$sex=="female", 1L, 0L)</pre>
sample.ids <- pheno.file[pheno.file$group=="experimental",]$ID_2</pre>
sangerCoxSurv(vcf.file=vcf.file,
               covariate.file=pheno.file,
               id.column="ID_2",
               sample.ids=sample.ids,
               time.to.event="time",
               event="event",
               covariates=c("age", "SexFemale", "DrugTxYes"),
               inter.term=NULL,
               print.covs="only",
               out.file="sanger_example",
               info.filter=0.3,
               maf.filter=0.005,
               chunk.size=50,
               verbose=TRUE,
               clusterObj=NULL)
```

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