Package 'CaseCrossover'

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CaseCrossoverData-class

2 CaseCrossoverData-class

	createExposureOutcomervestingConort	- 4
	createGetDbCaseCrossoverDataArgs	4
	createGetExposureStatusArgs	5
	createMatchingCriteria	6
	createSelectSubjectsToIncludeArgs	7
	fitCaseCrossoverModel	
	getAttritionTable	8
	getDbCaseCrossoverData	9
	getExposureStatus	11
	isCaseCrossoverData	12
	loadCaseCrossoverData	12
	loadCcrAnalysisList	13
	loadExposureOutcomeNestingCohortList	13
	runCcrAnalyses	14
	saveCaseCrossoverData	15
	saveCcrAnalysisList	16
	saveExposureOutcomeNestingCohortList	16
	selectSubjectsToInclude	17
	summarizeCcrAnalyses	18
Index		19

CaseCrossoverData-class

Case Data

Description

CaseCrossoverData is an S4 class that inherits from Andromeda. It contains information on cases.

A CaseCrossoverData object is typically created using getDbCaseCrossoverData, can only be saved using saveCaseCrossoverData, and loaded using loadCaseCrossoverData.

Usage

```
## S4 method for signature 'CaseCrossoverData'
show(object)
## S4 method for signature 'CaseCrossoverData'
summary(object)
```

Arguments

object An object of class 'CaseCrossoverData'.

See Also

isCaseCrossoverData

createCcrAnalysis 3

createCcrAnalysis

Create a case-crossover analysis specification

Description

Create a case-crossover analysis specification

Usage

```
createCcrAnalysis(
  analysisId = 1,
  description = "",
  exposureType = NULL,
  outcomeType = NULL,
  nestingCohortType = NULL,
  getDbCaseCrossoverDataArgs,
  selectSubjectsToIncludeArgs,
  getExposureStatusArgs
)
```

Arguments

analysisId An integer that will be used later to refer to this specific set of analysis choices.

description A short description of the analysis.

exposureType If more than one exposure is provided for each exposureOutcomeNestingCo-

hort, this field should be used to select the specific exposure to use in this anal-

ysis.

outcomeType If more than one outcome is provided for each exposureOutcomeNestingCohort,

this field should be used to select the specific outcome to use in this analysis.

nestingCohortType

If more than one nesting cohort is provided for each exposureOutcomeNesting-Cohort, this field should be used to select the specific nesting cohort to use in

this analysis.

getDbCaseCrossoverDataArgs

An object representing the arguments to be used when calling the createGetDbCaseCrossoverDataA function.

selectSubjectsToIncludeArgs

An object representing the arguments to be used when calling the selectSubjectsToInclude function.

getExposureStatusArgs

An object representing the arguments to be used when calling the getExposureStatus function.

Details

Create a set of analysis choices, to be used with the runCcrAnalyses function.

 ${\tt createExposureOutcomeNestingCohort}$

Create exposure-outcome-nesting-cohort combinations.

Description

Create exposure-outcome-nesting-cohort combinations.

Usage

```
createExposureOutcomeNestingCohort(
  exposureId,
  outcomeId,
  nestingCohortId = NULL
)
```

Arguments

exposureId

A concept ID identifying the target drug in the exposure table. If multiple strategies for picking the exposure will be tested in the analysis, a named list of numbers can be provided instead. In the analysis, the name of the number to be used can be specified using the exposureType parameter in the createCcrAnalysis function.

outcomeId

A concept ID identifying the outcome in the outcome table. If multiple strategies for picking the outcome will be tested in the analysis, a named list of numbers can be provided instead. In the analysis, the name of the number to be used can be specified using the outcomeType parameter in the createCcrAnalysis function.

nestingCohortId

A concept ID identifying the nesting cohort in the nesting cohort table. If multiple strategies for picking the nesting cohort will be tested in the analysis, a named list of numbers can be provided instead. In the analysis, the name of the number to be used can be specified using the nestingCohortType parameter in the createCcrAnalysis function.

Details

Create a set of hypotheses of interest, to be used with the runCcrAnalyses function.

createGetDbCaseCrossoverDataArgs

Create a parameter object for the function getDbCaseCrossoverData

Description

Create a parameter object for the function getDbCaseCrossoverData

Usage

```
createGetDbCaseCrossoverDataArgs(
  useNestingCohort = FALSE,
  useObservationEndAsNestingEndDate = TRUE,
  getVisits = FALSE,
  studyStartDate = ""
  studyEndDate = "",
  getTimeControlData = FALSE,
  maxNestingCohortSize = 1e+07,
  maxCasesPerOutcome = 5e+05
)
```

Arguments

useNestingCohort

Should the study be nested in a cohort (e.g. people with a specific indication)? If not, the study will be nested in the general population.

use Observation End As Nesting End Date

When using a nesting cohort, should the observation period end date be used instead of the cohort end date?

getVisits

Get data on visits? This is needed when performing a time- case-control study and matching on visit date is requested later on.

studyStartDate A calendar date specifying the minimum date where data is used. Date format is 'yyyymmdd'.

A calendar date specifying the maximum date where data is used. Date format studyEndDate is 'yyyymmdd'.

getTimeControlData

Should data for time controls be fetched? (needed for case-time-control analyses).

maxNestingCohortSize

If the nesting cohort is larger than this number it will be sampled to this size. maxCohortSize = 0 indicates no maximum size. (needed for case-time-control analyses).

 ${\tt maxCasesPerOutcome}$

If there are more than this number of cases for a single outcome cases will be sampled to this size. maxCasesPerOutcome = 0 indicates no maximum size.

Details

Create an object defining the parameter values.

createGetExposureStatusArgs

Create a parameter object for the function getExposureStatus

Description

Create a parameter object for the function getExposureStatus

Usage

```
createGetExposureStatusArgs(
  firstExposureOnly = FALSE,
  riskWindowStart = -30,
  riskWindowEnd = 0,
  controlWindowOffsets = c(-60)
)
```

Arguments

firstExposureOnly

Should only the first exposure per subject be included?

riskWindowStart

The start of the risk window (in days) relative to the index date. This number should be non-positive.

 ${\tt riskWindowEnd}$

The end of the risk window (in days) relative to the index date. This number should be non-positive.

controlWindowOffsets

Offsets in days of the control windows relative to the case window.

Details

Create an object defining the parameter values.

createMatchingCriteria

Create matching criteria

Description

Create matching criteria

Usage

```
createMatchingCriteria(
  controlsPerCase = 1,
  matchOnAge = TRUE,
  ageCaliper = 2,
  matchOnGender = TRUE,
  matchOnProvider = FALSE,
  matchOnCareSite = FALSE,
  matchOnVisitDate = FALSE,
  visitDateCaliper = 30,
  matchOnTimeInCohort = FALSE,
  daysInCohortCaliper = 30
)
```

Arguments

controlsPerCase

Maximum number of controls to select per case.

matchOnAge Match on age?

ageCaliper Maximum difference (in years) in age when matching on age.

matchOnGender Match on gender?

matchOnProvider

Match on provider (as specified in the person table)?

matchOnCareSite

Match on care site (as specified in the person table)?

matchOnVisitDate

Should the index date of the control be changed to the nearest visit date?

visitDateCaliper

Maximum difference (in days) between the index date and the visit date when matching on visit date.

matchOnTimeInCohort

Match on time in nesting cohort? When not using nesting, this is interpreted as time observed prior to index.

daysInCohortCaliper

Maximum difference (in days) in time in cohort.

createSelectSubjectsToIncludeArgs

Create a parameter object for the function selectSubjectsToInclude

Description

Create a parameter object for the function selectSubjectsToInclude

Usage

```
createSelectSubjectsToIncludeArgs(
  firstOutcomeOnly = TRUE,
  washoutPeriod = 180,
  matchingCriteria = NULL,
  minAge = NULL,
  maxAge = NULL
)
```

Arguments

firstOutcomeOnly

Use the first outcome per person?

washoutPeriod

Minimum required numbers of days of observation for inclusion as either case or control.

matchingCriteria

If provided, a case-time-control analysis will be performed and controls will be matched based on these criteria.

8 getAttritionTable

minAge Minimum age at which patient time will be included in the analysis. Note that

information prior to the min age is still used to determine exposure status after the minimum age (e.g. when a prescription was started just prior to reaching the minimum age). Also, outcomes occurring before the minimum age is reached will be considered as prior outcomes when using first outcomes only. Age should be specified in years, but non-integer values are allowed. If not

specified, no age restriction will be applied.

maxAge Maximum age at which patient time will be included in the analysis. Age should

be specified in years, but non-integer values are allowed. If not specified, no age

restriction will be applied.

Details

Create an object defining the parameter values.

fitCaseCrossoverModel Fit case-crossover model

Description

Fit case-crossover model

Usage

fitCaseCrossoverModel(exposureStatus)

Arguments

exposureStatus A data frame as generated using the getExposureStatus function.

Details

Fits a conditional logistic regression on the case-crossover data.

getAttritionTable Get the attrition table for a set of subjects

Description

Get the attrition table for a set of subjects

Usage

getAttritionTable(subjects)

Arguments

subjects A data frame of subjects as generated by the function selectSubjectsToInclude.

Value

A data frame specifying the number of cases and events after various steps of filtering.

getDbCaseCrossoverData

Load case-crossover data from the database

Description

Load all data about the cases from the database.

Usage

```
getDbCaseCrossoverData(
  connectionDetails,
  cdmDatabaseSchema,
 oracleTempSchema = cdmDatabaseSchema,
  outcomeDatabaseSchema = cdmDatabaseSchema,
  outcomeTable = "condition_era",
  outcomeIds = c(),
 useNestingCohort = FALSE,
 nestingCohortDatabaseSchema = cdmDatabaseSchema,
 nestingCohortTable = "cohort",
  nestingCohortId = NULL,
  useObservationEndAsNestingEndDate = TRUE,
  getVisits = FALSE,
  exposureDatabaseSchema = cdmDatabaseSchema,
  exposureTable = "drug_era",
  exposureIds = c(),
  studyStartDate = ""
  studyEndDate = "",
  getTimeControlData = FALSE,
 maxNestingCohortSize = 1e+07,
 maxCasesPerOutcome = 5e+05
)
```

Arguments

connectionDetails

An R object of type ConnectionDetails created using the function createConnectionDetails in the DatabaseConnector package.

 ${\tt cdmDatabaseSchema}$

The name of the database schema that contains the OMOP CDM instance. Requires read permissions to this database. On SQL Server, this should specify both the database and the schema, so for example 'cdm_instance.dbo'.

oracleTempSchema

A schema where temp tables can be created in Oracle.

 ${\tt outcomeDatabaseSchema}$

The name of the database schema that is the location where the data used to define the outcome cohorts is available. If outcomeTable = CONDITION_ERA, outcomeDatabaseSchema is not used. Requires read permissions to this database.

outcomeTable

The tablename that contains the outcome cohorts. If outcome Table is not CON-DITION_OCCURRENCE or CONDITION_ERA, then expectation is outcome Table

has format of COHORT table: COHORT DEFINITION ID, SUBJECT ID, COHORT START DATE, COHORT END DATE.

outcomeIds

A list of ids used to define outcomes. If outcomeTable = CONDITION_OCCURRENCE, the list is a set of ancestor CONCEPT_IDs, and all occurrences of all descendant concepts will be selected. If outcome Table <> CONDITION OCCURRENCE, the list contains records found in COHORT_DEFINITION_ID field.

useNestingCohort

Should the study be nested in a cohort (e.g. people with a specific indication)? If not, the study will be nested in the general population.

nestingCohortDatabaseSchema

The name of the database schema that is the location where the nesting cohort is defined.

nestingCohortTable

Name of the table holding the nesting cohort. This table should have the same structure as the cohort table.

nestingCohortId

A cohort definition ID identifying the records in the nestingCohortTable to use as nesting cohort.

use Observation End As Nesting End Date

When using a nesting cohort, should the observation period end date be used instead of the cohort end date?

getVisits

Get data on visits? This is needed when performing a time- case-control study and matching on visit date is requested later on.

exposureDatabaseSchema

The name of the database schema that is the location where the exposure data used to define the exposure cohorts is available. If exposure Table = DRUG_ERA, exposureDatabaseSchema is not used but assumed to be cdmSchema. Requires read permissions to this database.

exposureTable

The tablename that contains the exposure cohorts. If exposure Table <> DRUG_ERA, then expectation is exposure Table has format of COHORT table: cohort_concept_id, SUBJECT_ID, COHORT_START_DATE, COHORT_END_DATE.

exposureIds

A list of identifiers to define the exposures of interest. If exposure Table = DRUG_ERA, exposureIds should be CONCEPT_ID. If exposureTable <> DRUG_ERA, exposureIds is used to select the cohort_concept_id in the cohort-like table. If no exposureIds are provided, all drugs or cohorts in the exposureTable are included as exposures.

studyStartDate A calendar date specifying the minimum date where data is used. Date format is 'yyyymmdd'.

studyEndDate

A calendar date specifying the maximum date where data is used. Date format is 'yyyymmdd'.

getTimeControlData

Should data for time controls be fetched? (needed for case-time-control analy-

maxNestingCohortSize

If the nesting cohort is larger than this number it will be sampled to this size. maxCohortSize = 0 indicates no maximum size. (needed for case-time-control analyses).

maxCasesPerOutcome

If there are more than this number of cases for a single outcome cases will be sampled to this size. maxCasesPerOutcome = 0 indicates no maximum size.

getExposureStatus 11

Value

Returns an object of type CaseCrossoverData, containing information on the cases, the nesting cohort, exposures, and optionally visits. Information about multiple outcomes can be captured at once for efficiency reasons. The generic summary() function has been implemented for this object.

getExposureStatus

Get the exposure status for cases (and controls).

Description

Get the exposure status for cases (and controls).

Usage

```
getExposureStatus(
  subjects,
  caseCrossoverData,
  exposureId,
  firstExposureOnly = FALSE,
  riskWindowStart = -30,
  riskWindowEnd = 0,
  controlWindowOffsets = c(-60)
)
```

Arguments

subjects

A data frame as generated using the selectSubjectsToInclude function.

caseCrossoverData

An object of type caseCrossoverData as generated using the getDbCaseCrossoverData function.

exposureId '

The identifier of the exposure.

firstExposureOnly

Should only the first exposure per subject be included?

riskWindowStart

The start of the risk window (in days) relative to the index date. This number should be non-positive.

riskWindowEnd

The end of the risk window (in days) relative to the index date. This number should be non-positive.

controlWindowOffsets

Offsets in days of the control windows relative to the case window.

Details

This function determines the exposure status for a give, exposure ID in various windows relative to the index date.

12 loadCaseCrossoverData

Value

A data frame with these columns:

personId The person ID
indexDate The index date

isCase Is the person a case or a control?

stratumId The ID linking cases and controls in a matched set

isCaseWindow Is this a case window (as opposed to a control window)?

exposed Was the person exposed during the window?

isCaseCrossoverData

Check whether an object is a CaseCrossoverData object

Description

Check whether an object is a CaseCrossoverData object

Usage

isCaseCrossoverData(x)

Arguments

Х

The object to check.

Value

A logical value.

loadCaseCrossoverData Load the covariate data from a folder

Description

loadCaseCrossoverData loads an object of type caseCrossoverData from a file in the file system.

Usage

loadCaseCrossoverData(file)

Arguments

file

The name of the file containing the data.

Details

The data will be read from the file specified by the user.

Value

An object of class CaseCrossoverData.

loadCcrAnalysisList 13

loadCcrAnalysisList

Load a list of ccrAnalysis from file

Description

Load a list of objects of type ccrAnalysis from file. The file is in JSON format.

Usage

loadCcrAnalysisList(file)

Arguments

file

The name of the file

Value

A list of objects of type ccrAnalysis.

 ${\tt loadExposureOutcomeNestingCohortList}$

 $Load\ a\ list\ of\ exposureOutcomeNestingCohort\ from\ file$

Description

Load a list of objects of type exposureOutcomeNestingCohort from file. The file is in JSON format.

Usage

loadExposureOutcomeNestingCohortList(file)

Arguments

file

The name of the file

Value

A list of objects of type drugComparatorOutcome.

14 runCcrAnalyses

runCcrAnalyses

Run a list of analyses

Description

Run a list of analyses

Usage

```
runCcrAnalyses(
  connectionDetails,
  cdmDatabaseSchema,
 oracleTempSchema = cdmDatabaseSchema,
  exposureDatabaseSchema = cdmDatabaseSchema,
  exposureTable = "drug_era",
  outcomeDatabaseSchema = cdmDatabaseSchema,
  outcomeTable = "condition_era",
 nestingCohortDatabaseSchema = cdmDatabaseSchema,
 nestingCohortTable = "condition_era",
  outputFolder = "./CcrOutput",
  ccrAnalysisList,
  exposureOutcomeNestingCohortList,
  getDbCaseCrossoverDataThreads = 1,
  selectSubjectsToIncludeThreads = 1,
 getExposureStatusThreads = 1,
  fitCaseCrossoverModelThreads = 1
```

Arguments

connectionDetails

An R object of type ConnectionDetails created using the function createConnectionDetails in the DatabaseConnector package.

cdmDatabaseSchema

The name of the database schema that contains the OMOP CDM instance. Requires read permissions to this database. On SQL Server, this should specify both the database and the schema, so for example 'cdm_instance.dbo'.

oracleTempSchema

A schema where temp tables can be created in Oracle.

exposureDatabaseSchema

The name of the database schema that is the location where the exposure data used to define the exposure cohorts is available. If exposureTable = DRUG_ERA, exposureDatabaseSchema is not used but assumed to be cdmSchema. Requires read permissions to this database.

exposureTable

The tablename that contains the exposure cohorts. If exposure Table <> drug_era, then expectation is exposure Table has format of COHORT table: cohort_definition_id, subject_id, cohort_start_date, cohort_end_date.

outcomeDatabaseSchema

The name of the database schema that is the location where the data used to define the outcome cohorts is available. If outcomeTable = CONDITION_ERA, outcomeDatabaseSchema is not used. Requires read permissions to this database.

saveCaseCrossoverData 15

outcomeTable

The tablename that contains the outcome cohorts. If outcomeTable is not CON-DITION_OCCURRENCE or CONDITION_ERA, then expectation is outcomeTable has format of COHORT table: COHORT_DEFINITION_ID, SUBJECT_ID, COHORT_START_DATE, COHORT_END_DATE.

nestingCohortDatabaseSchema

The name of the database schema that is the location where the nesting cohort is defined.

nestingCohortTable

Name of the table holding the nesting cohort. This table should have the same structure as the cohort table.

outputFolder

Name of the folder where all the outputs will written to.

ccrAnalysisList

A list of objects of type ccrAnalysis as created using the createCcrAnalysis function.

 ${\tt exposureOutcomeNestingCohortList}$

A list of objects of type exposureOutcomeNestingCohort as created using the createExposureOutcomeNestingCohort function.

getDbCaseCrossoverDataThreads

The number of parallel threads to use for building the caseControlData objects.

selectSubjectsToIncludeThreads

The number of parallel threads to use for selecting subjects

 ${\tt getExposureStatusThreads}$

The number of parallel threads to use for getting exposure status.

fit Case Crossover Model Threads

The number of parallel threads to use for fitting the models.

Details

Run a list of analyses for the exposure-outcome-nesting cohorts of interest. This function will run all specified analyses against all hypotheses of interest, meaning that the total number of outcome models is 'length(ccrAnalysisList) * length(exposureOutcomeNestingCohortList)'. When you provide several analyses it will determine whether any of the analyses have anything in common, and will take advantage of this fact. For example, if we specify several analyses that only differ in the way the control windows are specified then this function will extract the data and select the subjects only once, and re-use this in all the analysis.

saveCaseCrossoverData Save the case data to file

Description

saveCaseCrossoverData saves an object of type caseCrossoverData to file

Usage

saveCaseCrossoverData(caseCrossoverData, file)

Arguments

caseCrossoverData

An object of type CaseCrossoverData as generated using getDbCaseCrossoverData.

file

The name of the file where the data will be written. If the file exists it will be overwritten.

Details

The data will be written to the file specified by the user.

saveCcrAnalysisList

Save a list of ccrAnalysis to file

Description

Write a list of objects of type ccrAnalysis to file. The file is in JSON format.

Usage

```
saveCcrAnalysisList(ccrAnalysisList, file)
```

Arguments

ccrAnalysisList

The ccArnalysis list to be written to file

file

The name of the file where the results will be written

 ${\tt save Exposure Out come Nesting Cohort List}$

Save a list of drugComparatorOutcome to file

Description

Write a list of objects of type exposureOutcomeNestingCohort to file. The file is in JSON format.

Usage

saveExposureOutcomeNestingCohortList(exposureOutcomeNestingCohortList, file)

Arguments

 ${\tt exposureOutcomeNestingCohortList}$

The exposureOutcomeNestingCohort list to be written to file

file The name of the file where the results will be written

selectSubjectsToInclude

Select subjects to include

Description

Select subjects to include

Usage

```
selectSubjectsToInclude(
  caseCrossoverData,
 outcomeId,
  firstOutcomeOnly = TRUE,
  washoutPeriod = 180,
 matchingCriteria = NULL,
 minAge = NULL,
 maxAge = NULL
)
```

Arguments

caseCrossoverData

An object of type caseCrossoverData as generated using the getDbCaseCrossoverData function.

outcomeId

The outcome ID of the cases for which we need to pick controls.

firstOutcomeOnly

Use the first outcome per person?

washoutPeriod

Minimum required numbers of days of observation for inclusion as either case

or control.

matchingCriteria

If provided, a case-time-control analysis will be performed and controls will be

matched based on these criteria.

minAge

Minimum age at which patient time will be included in the analysis. Note that information prior to the min age is still used to determine exposure status after the minimum age (e.g. when a prescription was started just prior to reaching the minimum age). Also, outcomes occurring before the minimum age is reached will be considered as prior outcomes when using first outcomes only. Age should be specified in years, but non-integer values are allowed. If not

specified, no age restriction will be applied.

maxAge

Maximum age at which patient time will be included in the analysis. Age should be specified in years, but non-integer values are allowed. If not specified, no age

restriction will be applied.

Details

Subject to include in the study are selected for a specific outcome, optionally filtering using a washout period, restricting to first occurrences of the outcome only, and restricting on age. If matching criteria are provided controls will be selected for each case. These controls will be used to adjust for time trends in exposure, turning the analysis into a case-time-control analysis (Suissa, 1995).

Value

A data frame with these columns:

personId The person ID
indexDate The index date

isCase Is the person a case or a control?

stratumId The ID linking cases and controls in a matched set **observationPeriodStartDate** The observation period start date

References

Suissa S (1995) The case-time-control design. Epidemiology; 6:248-253.

summarizeCcrAnalyses

Create a summary report of the analyses

Description

Create a summary report of the analyses

Usage

summarizeCcrAnalyses(outcomeReference, outputFolder)

Arguments

outcomeReference

A data.frame as created by the runCcrAnalyses function.

outputFolder Name of the folder where all the outputs have been written to.

Index

```
Andromeda, 2
CaseCrossoverData
        (CaseCrossoverData-class), 2
CaseCrossoverData-class, 2
createCcrAnalysis, 3, 4, 15
createExposureOutcomeNestingCohort, 4,
createGetDbCaseCrossoverDataArgs, 3, 4
createGetExposureStatusArgs, 5
createMatchingCriteria, 6
create Select Subjects To Include Args, \\ 7
fitCaseCrossoverModel, 8
getAttritionTable, 8
getDbCaseCrossoverData, 2, 9, 11, 16, 17
getExposureStatus, 3, 8, 11
isCaseCrossoverData, 2, 12
loadCaseCrossoverData, 2, 12
loadCcrAnalysisList, 13
loadExposureOutcomeNestingCohortList,
        13
runCcrAnalyses, 3, 4, 14, 18
saveCaseCrossoverData, 2, 15
saveCcrAnalysisList, 16
{\tt save Exposure Outcome Nesting Cohort List},
selectSubjectsToInclude, 3, 8, 11, 17
show,CaseCrossoverData-method
        (CaseCrossoverData-class), 2
summarizeCcrAnalyses, 18
summary,CaseCrossoverData-method
        (CaseCrossoverData-class), 2
```