Package 'SelfControlledCohort'

November 23, 2015

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Type Package	
Title Population-level estimation method that estimates incidence rate comparison of exposed/unexposed time within an exposed cohort	
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createExposureOutcome Create exposure-outcome combinations.

Description

Create exposure-outcome combinations.

Usage

createExposureOutcome(exposureId, outcomeId)

Arguments

exposureId

A concept ID indentifying the drug of interest 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 createSccAnalysis function.

outcomeId

A concept ID indentifying the outcome of interest 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 createSccAnalysis function.

Details

Create a hypothesis of interest, to be used with the runSccAnalyses function.

create Run Self Controlled Cohort Args

Create a parameter object for the function runSelfControlledCohort

Description

Create a parameter object for the function runSelfControlledCohort

Usage

```
createRunSelfControlledCohortArgs(firstOccurrenceDrugOnly = TRUE,
  firstOccurrenceConditionOnly = TRUE,
  outcomeConditionTypeConceptIds = c(38000247), genderConceptIds = c(8507,
  8532), minAge = "", maxAge = "", studyStartDate = "",
  studyEndDate = "", stratifyByGender = FALSE, stratifyByAge = FALSE,
  stratifyByYear = FALSE, useLengthOfExposureExposed = TRUE,
  timeAtRiskExposedStart = 1, surveillanceExposed = 30,
  useLengthOfExposureUnexposed = TRUE, timeAtRiskUnexposedStart = -1,
  surveillanceUnexposed = -30, hasFullTimeAtRisk = FALSE,
  washoutWindow = 0, followupWindow = 0, shrinkage = 1e-04)
```

Arguments

firstOccurrenceDrugOnly

If TRUE, only use first occurrence of each drug concept idfor each person

firstOccurrenceConditionOnly

If TRUE, only use first occurrence of each condition conceptid for each person.

outcomeConditionTypeConceptIds

A list of TYPE_CONCEPT_ID values that will restrict condition occurrences.

Only applicable if outcomeTable =CONDITION_OCCURRENCE.

genderConceptIds

of gender_concept_id, generally use MALE (8507) and FEMALE(8532).

minAge Integer for minimum allowable age.
maxAge Integer for maximum allowable age.

studyStartDate Date for minimum allowable data for index exposure.

studyEndDate Date for maximum allowable data for index exposure.

stratifyByGender

If TRUE, analysis will be calculated overall, and stratified cross all gender

groups.

stratifyByAge If TRUE, analysis will be calculated overall, and stratified across all age groups

(using AGE_GROUP table below).

stratifyByYear If TRUE, analysis will be calculated overall, and stratified across all years of the

index dates.

useLengthOfExposureExposed

If TRUE, use the duration from drugEraStart -> drugEraEnd aspart of timeAtRisk.

timeAtRiskExposedStart

Integer of days to add to drugEraStart for start of timeAtRisk (0 to include index $\,$

date, 1 to start the dayafter).

surveillanceExposed

Additional window to add to end of exposure period (ifuseLengthOfExposure-

Exposed = TRUE, then add to exposure enddate, else add to exposure start date).

useLengthOfExposureUnexposed

If TRUE, use the duration from exposure start -> exposureend as part of timeAtRisk

looking back before exposurestart.

timeAtRiskUnexposedStart

Integer of days to add to exposure start for start of timeAtRisk (0 to include index

date, -1 to start the daybefore).

surveillanceUnexposed

Additional window to add to end of exposure period (ifuseLengthOfExposure-

Unexposed = TRUE, then add to exposureend date, else add to exposure start

date).

hasFullTimeAtRisk

If TRUE, restrict to people who have full time-at-riskexposed and unexposed.

washoutWindow Integer to define required time observed before exposurestart.

followupWindow Integer to define required time observed after exposurestart.

shrinkage used in IRR calculations, required >0 to deal with0 case counts, but

larger number means more shrinkage.

Details

Create an object defining the parameter values.

createSccAnalysis

Create a SelfControlledCohort analysis specification

Description

Create a SelfControlledCohort analysis specification

Usage

```
createSccAnalysis(analysisId = 1, description = "", exposureType = NULL,
  outcomeType = NULL, runSelfControlledCohortArgs)
```

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.

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

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

runSelfControlledCohortArgs

 $An object \ representing \ the \ arguments \ to \ be \ used \ when \ calling \ the \ runSelfControlledCohort$

function.

Details

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

loadExposureOutcomeList

Load a list of exposureOutcome from file

Description

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

Usage

loadExposureOutcomeList(file)

Arguments

file

The name of the file

Value

A list of objects of type exposureOutcome.

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loadsccAnalysisList

Load a list of sccAnalysis from file

Description

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

Usage

```
loadsccAnalysisList(file)
```

Arguments

file

The name of the file

Value

A list of objects of type sccAnalysis.

runSccAnalyses

Run a list of analyses

Description

Run a list of analyses

Usage

```
runSccAnalyses(connectionDetails, cdmDatabaseSchema,
  oracleTempSchema = cdmDatabaseSchema,
  exposureDatabaseSchema = cdmDatabaseSchema, exposureTable = "drug_era",
  outcomeDatabaseSchema = cdmDatabaseSchema,
  outcomeTable = "condition_occurrence", cdmVersion = 4,
  outputFolder = "./SelfControlledCohortOutput", sccAnalysisList,
  exposureOutcomeList, analysisTreads = 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

For Oracle only: the name of the database schema where you want all temporary tables to be managed. Requires create/insert permissions to this database.

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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 by assumed to be cdmSchema. Requires read permissions to this database.

exposureTable

The tablename that contains the exposure cohorts. If exposureTable <> DRUG_ERA, then expectation is exposureTable 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 exposureTable = CONDITION_ERA, exposureDatabaseSchema is not used by assumed to be cdmSchema. Requires read permissions to this database.

outcomeTable

The tablename that contains the outcome cohorts. If outcomeTable <> CONDITION_OCCURRENCE, then expectation is outcomeTable has format of COHORT table: COHORT_DEFINITION_ID, SUBJECT_ID, COHORT_START_DATE, COHORT_END_DATE.

cdmVersion

Define the OMOP CDM version used: currently support "4" and "5".

outputFolder

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

sccAnalysisList

A list of objects of type sccAnalysis as created using the createSccAnalysis function.

exposureOutcomeList

A list of objects of type exposureOutcome as created using the ${\tt createExposureOutcome}$ function.

analysisTreads The number of parallel threads to use to execute the analyses.

Details

Run a list of analyses for the drug-comparator-outcomes of interest. This function will run all specified analyses against all hypotheses of interest, meaning that the total number of outcome models is 'length(cmAnalysisList) * length(drugComparatorOutcomesList)' (if all analyses specify an outcome model should be fitted). 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 outcome model is fitted, then this function will extract the data and fit the propensity model only once, and re-use this in all the analysis.

 $\verb"runSelfControlledCohort"$

Run self-controlled cohort

Description

runSelfControlledCohort generates population-level estimation by comparing exposed and unexposed time among exposed cohort.

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Usage

```
runSelfControlledCohort(connectionDetails, cdmDatabaseSchema, cdmVersion = 5,
  oracleTempSchema, exposureIds, outcomeId,
  exposureDatabaseSchema = cdmDatabaseSchema, exposureTable = "drug_era",
  outcomeDatabaseSchema = cdmDatabaseSchema, outcomeTable = "condition_era",
  firstOccurrenceDrugOnly = TRUE, firstOccurrenceConditionOnly = TRUE,
  outcomeConditionTypeConceptIds = c(38000247), genderConceptIds = c(8507,
  8532), minAge = "", maxAge = "", studyStartDate = "",
  studyEndDate = "", stratifyByGender = FALSE, stratifyByAge = FALSE,
  stratifyByYear = FALSE, useLengthOfExposureExposed = TRUE,
  timeAtRiskExposedStart = 1, surveillanceExposed = 30,
  useLengthOfExposureUnexposed = TRUE, timeAtRiskUnexposedStart = -1,
  surveillanceUnexposed = -30, hasFullTimeAtRisk = FALSE,
  washoutWindow = 0, followupWindow = 0, shrinkage = 1e-04)
```

Arguments

connectionDetails

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

cdmDatabaseSchema

Name of database schema that contains the OMOP CDM and vocabulary.

cdmVersion Define the OMOP CDM version used: currently support "4" and "5".

oracleTempSchema

For Oracle only: the name of the database schema where you want all temporary tables to be managed. Requires create/insert permissions to this database.

exposureIds A vector containing the drug_concept_ids or cohort_definition_ids of the exposures of interest

 ${\tt outcomeId} \qquad {\tt The\ condition_concept_id\ or\ cohort_definition_id\ of\ the\ outcome\ of\ interest\ exposure{\tt DatabaseSchema}}$

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 by assumed to be cdmSchema. Requires read permissions to this database.

exposureTable The tablename that contains the exposure cohorts. If exposureTable <> DRUG_ERA, then expectation is exposureTable has format of COHORT table: cohort_concept_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 exposureTable = CONDITION_ERA, exposureDatabaseSchema is not used by assumed to be cdmSchema. Requires read permissions to this database.

outcomeTable The tablename that contains the outcome cohorts. If outcomeTable <> CONDITION_OCCURRENCE, then expectation is outcomeTable has format of COHORT table: COHORT_DEFINITION_ID, SUBJECT_ID, COHORT_START_DATE, COHORT_END_DATE.

firstOccurrenceDrugOnly

 $\label{thm:constraint} If \ TRUE, only \ use \ first \ occurrence \ of each \ drug \ concept \ id \ for \ each \ person \\ first \ Occurrence \ Condition \ Only$

If TRUE, only use first occurrence of each condition concept id for each person.

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 $\verb"outcomeConditionTypeConceptIds"$

A list of TYPE_CONCEPT_ID values that will restrict condition occurrences. Only applicable if outcomeTable = CONDITION_OCCURRENCE.

genderConceptIds

of gender_concept_id, generally use MALE (8507) and FEMALE (8532).

minAge Integer for minimum allowable age.
maxAge Integer for maximum allowable age.

studyStartDate Date for minimum allowable data for index exposure. Date format is 'yyyymmdd'.

studyEndDate Date for maximum allowable data for index exposure. Date format is 'yyyym-

mdd'.

stratifyByGender

If TRUE, analysis will be calculated overall, and stratified across all gender

stratifyByAge If TRUE, analysis will be calculated overall, and stratified across all age groups (using AGE_GROUP table below).

stratifyByYear If TRUE, analysis will be calculated overall, and stratified across all years of the index dates.

useLengthOfExposureExposed

If TRUE, use the duration from drugEraStart -> drugEraEnd as part of timeAtRisk.

timeAtRiskExposedStart

Integer of days to add to drugEraStart for start of timeAtRisk (0 to include index date, 1 to start the day after).

surveillanceExposed

Additional window to add to end of exposure period (if useLengthOfExposure-Exposed = TRUE, then add to exposure end date, else add to exposure start date).

useLengthOfExposureUnexposed

If TRUE, use the duration from exposure start -> exposure end as part of timeAtRisk looking back before exposure start.

 ${\tt timeAtRiskUnexposedStart}$

Integer of days to add to exposure start for start of timeAtRisk (0 to include index date, -1 to start the day before).

surveillanceUnexposed

Additional window to add to end of exposure period (if useLengthOfExposure-Unexposed = TRUE, then add to exposure end date, else add to exposure start date).

hasFullTimeAtRisk

If TRUE, restrict to people who have full time-at-risk exposed and unexposed.

washoutWindow Integer to define required time observed before exposure start.

followupWindow Integer to define required time observed after exposure start.

shrinkage used in IRR calculations, required >0 to deal with 0 case counts, but

larger number means more shrinkage.

Details

Population-level estimation method that estimates incidence rate comparison of exposed/unexposed time within an exposed cohort.

Value

An object of type sccResults containing the results of the analysis.

References

Ryan PB, Schuemie MJ, Madigan D.Empirical performance of a self-controlled cohort method: lessons for developing a risk identification and analysis system. Drug Safety 36 Suppl1:S95-106, 2013

Examples

saveExposureOutcomeList

Save a list of exposureOutcome to file

Description

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

Usage

```
saveExposureOutcomeList(exposureOutcomeList, file)
```

Arguments

exposureOutcomeList

The exposureOutcome list to be written to file

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

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saveSccAnalysisList

Save a list of sccAnalysis to file

Description

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

Usage

```
saveSccAnalysisList(sccAnalysisList, file)
```

Arguments

sccAnalysisList

The sccAnalysis list to be written to file

file

The name of the file where the results will be written

 ${\tt SelfControlledCohort} \quad \textit{SelfControlledCohort}$

Description

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 ${\it summarize Analyses}$

Create a summary report of the analyses

Description

Create a summary report of the analyses

Usage

```
summarizeAnalyses(resultsReference)
```

Arguments

resultsReference

A data.frame as created by the runSccAnalyses function.

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