Package 'CohortMethod'

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```
Type Package

Title New-user cohort method with large scale propensity and outcome models

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```

Description CohortMethod is an R package for performing new-user cohort studies in an observational database in the OMOP Common Data Model. It extracts the necessary data from a database in OMOP Common Data Model format, and uses a large set of covariates for both the propensity and outcome model, including for example all drugs, diagnoses, procedures, as well as age, comorbidity indexes, etc. Large scale regularized regression is used to fit the propensity and outcome models. Functions are included for trimming, stratifying and matching on propensity scores, as well as diagnostic functions, such as propensity score distribution plots and plots showing covariate balance before and after matching and/or trimming. Supported outcome models are (conditional) logistic regression, (conditional) Poisson regression, and (stratified) Cox regression.

```
License Apache License 2.0
VignetteBuilder knitr
Depends R (>= 3.2.2),
      DatabaseConnector (>= 1.3.0),
      Cyclops (>= 1.2.0),
      FeatureExtraction (>= 1.0.0)
Imports bit,
      methods,
      ggplot2,
      gridExtra,
      grid,
      ff.
      ffbase (>= 0.12.3),
      plyr,
      Rcpp (>= 0.11.2),
      RJDBC,
      SqlRender (>= 1.1.1),
```

survival,

2 R topics documented:

stringi, OhdsiRTools (>= 1.1.2)
Suggests testthat,
pROC,
gnm,
knitr,
rmarkdown,
EmpiricalCalibration
LinkingTo Rcpp
NeedsCompilation yes
RoxygenNote 6.0.1

R topics documented:

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checkCmInstallation

Check is CohortMethod and its dependencies are correctly installed

Description

Check is CohortMethod and its dependencies are correctly installed

Usage

 ${\tt checkCmInstallation(connectionDetails)}$

Arguments

connectionDetails

An R object of type

 ${\tt connectionDetails}\ created\ using\ the\ function\ create{\tt ConnectionDetails}\ in\ the\ {\tt DatabaseConnector}\ package.$

Details

This function checks whether CohortMethod and its dependencies are correctly installed. This will check the database connectivity, large scale regresion engine (Cyclops), and large data object handling (ff).

CohortMethod

CohortMethod

Description

CohortMethod

 ${\it cohort} {\it MethodDataSimulationProfile} \\ A {\it simulation profile}$

Description

A simulation profile

Usage

data(cohortMethodDataSimulationProfile)

computeCovariateBalance

Compute covariate balance before and after matching and trimming

Description

For every covariate, prevalence in treatment and comparator groups before and after matching/trimming are computed. When variable ratio matching was used the balance score will be corrected according the method described in Austin et al (2008).

Usage

computeCovariateBalance(population, cohortMethodData)

Arguments

population A data frame containing the people that are remaining after matching and/or trimming.

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

Details

The population data frame should have at least the following columns:

rowId (integer) A unique identifier for each row (e.g. the person ID)
treatment (integer) Column indicating whether the person is in the treated (1) or comparator (0)
group

Value

Returns a date frame describing the covariate balance before and after matching/trimming.

References

Austin, P.C. (2008) Assessing balance in measured baseline covariates when using many-to-one matching on the propensity-score. Pharmacoepidemiology and Drug Safety, 17: 1218-1225.

computeMdrr 5

(computeMdrr	Compute the minimum detectable relative risk
	, and the second	1

Description

Compute the minimum detectable relative risk

Usage

```
computeMdrr(population, alpha = 0.05, power = 0.8, twoSided = TRUE,
  modelType = "cox")
```

Arguments

population A data frame describing the study population as created using the createStudyPopulation

function. This should at least have these columns: subjectId, treatment, out-

comeCount, timeAtRisk.

alpha Type I error.

power 1 - beta, where beta is the type II error.

twoSided Consider a two-sided test?

modelType The type of outcome model that will be used. Possible values are "logistic",

"poisson", or "cox". Currently only "cox" is supported.

Details

Compute the minimum detectable relative risk (MDRR) for a given study population, using the actual observed sample size and number of outcomes. Currently, only computations for Cox models are implemented. For Cox model, the computations by Schoenfeld (1983) is used.

Value

A data frame with the MDRR and some counts.

References

Schoenfeld DA (1983) Sample-size formula for the proportional-hazards regression model, Biometrics, 39(3), 499-503

computePsAuc

Compute the area under the ROC curve

Description

computePsAuc computes the area under the ROC curve of the propensity score

Usage

```
computePsAuc(data, confidenceIntervals = FALSE)
```

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Arguments

data A data frame with at least the two columns described below confidenceIntervals

Compute 95 percent confidence intervals (computationally expensive for large data sets)

Details

The data frame should have a least the following two columns:

treatment (integer) Column indicating whether the person is in the treated (1) or comparator

(0) group

propensityScore (numeric) Propensity score

Value

A data frame holding the AUC and its 95 percent confidence interval

Examples

```
treatment <- rep(0:1, each = 100)
propensityScore <- c(rnorm(100, mean = 0.4, sd = 0.25), rnorm(100, mean = 0.6, sd = 0.25))
data <- data.frame(treatment = treatment, propensityScore = propensityScore)
data <- data[data$propensityScore > 0 & data$propensityScore < 1, ]
computePsAuc(data)</pre>
```

constructEras

Build eras

Description

Constructs eras (continuous periods of exposure or disease).

Usage

```
constructEras(connectionDetails, sourceDatabaseSchema,
  sourceTable = "drug_exposure",
  targetDatabaseSchema = sourceDatabaseSchema, targetTable = "drug_era",
  createTargetTable = FALSE, cdmDatabaseSchema = sourceDatabaseSchema,
  gracePeriod = 30, rollUp = TRUE, rollUpConceptClassId = "Ingredient",
  rollUpVocabularyId = "RxNorm", cdmVersion = "5")
```

Arguments

connectionDetails

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

sourceDatabaseSchema

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

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sourceTable The name of the source table. targetDatabaseSchema

The name of the database schema that contains the target table. Requires write permissions to this database. On SQL Server, this should specify both the database and the schema, so for example 'cdm_instance.dbo'.

targetTable The name of the target table.

createTargetTable

Should the target table be created? If not, the data is inserted in an existing table.

cdmDatabaseSchema

Only needed when rolling up concepts to ancestors: The name of the database schema that contains the vocabulary files. Requires read permissions to this database. On SQL Server, this should specify both the database and the schema, so for example 'cdm instance.dbo'.

gracePeriod The number of days allowed between periods for them to still be considered part

of the same era.

rollUp Should concepts be rolled up to their ancestors?

 ${\tt rollUpConceptClassId}$

The identifier of the concept class to which concepts should be rolled up.

rollUpVocabularyId

The identifier of the vocabulary to which concepts should be rolled up.

cdmVersion The verion of the CDM that is being used.

Details

This function creates eras from source data. For example, one could use this function to create drug eras based on drug exposures. The function allows drugs to be rolled up to ingredients, and prescriptions to the same ingredient that overlap in time are merged into a single ingredient. Note that stockpiling is not assumed to take place (ie. overlap is discarded), but a grace period can be specified allowing for a small gap between prescriptions when merging. The user can specify the source and target table. These tables are assumed to have the same structure as the cohort table in the Common Data Model (CDM), except when the table names are 'drug_exposure' or 'condition_occurrence' for the source table, or 'drug_era' or 'condition_era' for the target table, in which case the tables are assumed to have the structure defined for those tables in the CDM. If both the source and target table specify a field for type_concept_id, the era construction will partition by the type_concept_id, in other words periods with different type_concept_ids will be treated independently.

Examples

Constructing drug eras in CDM v5:

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createCmAnalysis

Create a CohortMethod analysis specification

Description

Create a CohortMethod analysis specification

Usage

```
createCmAnalysis(analysisId = 1, description = "", targetType = NULL,
   comparatorType = NULL, getDbCohortMethodDataArgs, createStudyPopArgs,
   createPs = FALSE, createPsArgs = NULL, trimByPs = FALSE,
   trimByPsArgs = NULL, trimByPsToEquipoise = FALSE,
   trimByPsToEquipoiseArgs = NULL, matchOnPs = FALSE, matchOnPsArgs = NULL,
   matchOnPsAndCovariates = FALSE, matchOnPsAndCovariatesArgs = NULL,
   stratifyByPs = FALSE, stratifyByPsArgs = NULL,
   stratifyByPsAndCovariates = FALSE, stratifyByPsAndCovariatesArgs = NULL,
   computeCovariateBalance = FALSE, fitOutcomeModel = FALSE,
   fitOutcomeModelArgs = NULL)
```

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.

targetType If more than one target is provided for each drugComparatorOutcome, this field

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

comparatorType If more than one comparator is provided for each drugComparatorOutcome, this

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

 ${\tt getDbCohortMethodDataArgs}$

An object representing the arguments to be used when calling the getDbCohortMethodData

function.

createStudyPopArgs

An object representing the arguments to be used when calling the createStudyPopulation

function.

createPs Should the createPs function be used in this analysis?

createPsArgs An object representing the arguments to be used when calling the createPs

function.

trimByPs Should the trimByPs function be used in this analysis?

trimByPsArgs An object representing the arguments to be used when calling the trimByPs

function.

trimByPsToEquipoise

Should the trimByPsToEquipoise function be used in this analysis?

trimByPsToEquipoiseArgs

An object representing the arguments to be used when calling the trimByPsToEquipoise

function.

matchOnPs Should the matchOnPs function be used in this analysis?

matchOnPsArgs An object representing the arguments to be used when calling the matchOnPs

function.

matchOnPsAndCovariates

Should the matchOnPsAndCovariates function be used in this analysis?

matchOnPsAndCovariatesArgs

An object representing the arguments to be used when calling the matchOnPsAndCovariates

function.

stratifyByPs Should the stratifyByPs function be used in this analysis?

stratifyByPsArgs

An object representing the arguments to be used when calling the stratifyByPs

function.

stratifyByPsAndCovariates

Should the stratifyByPsAndCovariates function be used in this analysis?

stratifyByPsAndCovariatesArgs

 $An object \ representing \ the \ arguments \ to \ be \ used \ when \ calling \ the \ stratify ByPs And Covariates$

function.

computeCovariateBalance

Should the computeCovariateBalance function be used in this analysis?

fitOutcomeModel

Should the fitOutcomeModel function be used in this analysis?

fitOutcomeModelArgs

An object representing the arguments to be used when calling the ${\tt fitOutcomeModel}$

function.

Details

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

 ${\tt createCohortMethodDataSimulationProfile}$

Create simulation profile

Description

createCohortMethodDataSimulationProfile creates a profile based on the provided cohort-MethodData object, which can be used to generate simulated data that has similar characteristics.

Usage

 $create {\tt CohortMethodDataSimulationProfile} (cohort{\tt MethodData})$

10 createCreatePsArgs

Arguments

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

Details

The output of this function is an object that can be used by the simulateCohortMethodData function to generate a cohortMethodData object.

Value

An object of type cohortDataSimulationProfile.

createCreatePsArgs

Create a parameter object for the function createPs

Description

Create a parameter object for the function createPs

Usage

```
createCreatePsArgs(excludeCovariateIds = c(), includeCovariateIds = c(),
  errorOnHighCorrelation = TRUE, stopOnError = TRUE,
  prior = createPrior("laplace", exclude = c(0), useCrossValidation = TRUE),
  control = createControl(noiseLevel = "silent", cvType = "auto", tolerance =
  2e-07, cvRepetitions = 10, startingVariance = 0.01))
```

Arguments

 ${\tt excludeCovariateIds}$

Exclude these covariates from the propensity model.

includeCovariateIds

Include only these covariates in the propensity model.

errorOnHighCorrelation

If true, the function will test each covariate for correlation withthe treatment assignment. If any covariate has an unusually highcorrelation (either positive or

negative), this will throw anderror.

stopOnError If an error occurrs, should the function stop? Else, the two cohortswill be as-

sumed to be perfectly separable.

prior The prior used to fit the model. SeecreatePrior for details.

control The control object used to control the cross-validation used todetermine the hy-

perparameters of the prior (if applicable). SeecreateControl for details.

Details

Create an object defining the parameter values.

createCreateStudyPopulationArgs

Create a parameter object for the function createStudyPopulation

Description

Create a parameter object for the function createStudyPopulation

Usage

```
createCreateStudyPopulationArgs(firstExposureOnly = FALSE,
  restrictToCommonPeriod = FALSE, washoutPeriod = 0,
  removeDuplicateSubjects = FALSE, removeSubjectsWithPriorOutcome = TRUE,
  priorOutcomeLookback = 99999, minDaysAtRisk = 1, riskWindowStart = 0,
  addExposureDaysToStart = FALSE, riskWindowEnd = 0,
  addExposureDaysToEnd = TRUE)
```

Arguments

firstExposureOnly

Should only the first exposure per subject be included? Notethat this is typically done in thecreateStudyPopulation function,

restrictToCommonPeriod

Restrict the analysis to the period when both treatments are observed?

washoutPeriod The minimum required continuous observation time prior toindex date for a person to be included in the cohort.

removeDuplicateSubjects

Remove subjects that are in both the treated and comparatorcohort?

removeSubjectsWithPriorOutcome

Remove subjects that have the outcome prior to the riskwindow start?

priorOutcomeLookback

How many days should we look back when identifying prioroutcomes?

minDaysAtRisk The minimum required number of days at risk.

riskWindowStart

The start of the risk window (in days) relative to the indexdate (+ days of exposure if theaddExposureDaysToStart parameter is specified).

 ${\it add} {\it Exposure Days To Start}$

Add the length of exposure the start of the risk window?

riskWindowEnd The end of the risk window (in days) relative to the indexdata (+ days of exposure if the addExposureDaysToEndparameter is specified).

addExposureDaysToEnd

Add the length of exposure the risk window?

Details

Create an object defining the parameter values.

createDrugComparatorOutcomes

Create drug-comparator-outcomes combinations.

Description

Create drug-comparator-outcomes combinations.

Usage

```
createDrugComparatorOutcomes(targetId, comparatorId, outcomeIds,
  excludedCovariateConceptIds = c(), includedCovariateConceptIds = c())
```

Arguments

targetId A concept ID indentifying the target drug in the exposure table. If multiple

strategies for picking the target 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 #' targetType parameter in the createCmAnalysis

function.

comparator Id A concept ID indentifying the comparator drug in the exposure table. If multi-

ple strategies for picking the comparator 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 #' comparatorType parameter in the

 ${\tt createCmAnalysis}\ function.$

outcomeIds A vector of concept IDs indentifying the outcome(s) in the outcome table.

excludedCovariateConceptIds

A list of concept IDs that cannot be used to construct covariates. This argument is to be used only for exclusion concepts that are specific to the drug-comparator

combination.

includedCovariateConceptIds

A list of concept IDs that must be used to construct covariates. This argument is to be used only for inclusion concepts that are specific to the drug-comparator combination.

Details

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

createFitOutcomeModelArgs

Create a parameter object for the function fitOutcomeModel

Description

Create a parameter object for the function fitOutcomeModel

Usage

```
createFitOutcomeModelArgs(modelType = "logistic", stratified = TRUE,
  useCovariates = TRUE, excludeCovariateIds = c(),
  includeCovariateIds = c(), prior = createPrior("laplace",
  useCrossValidation = TRUE), control = createControl(cvType = "auto",
  startingVariance = 0.01, tolerance = 2e-07, cvRepetitions = 10, noiseLevel =
  "quiet"))
```

Arguments

modelType The type of outcome model that will be used. Possible values are "logistic",

"poisson", or "cox".

stratified Should the regression be conditioned on the strata defined in the population ob-

ject (e.g. by matching or stratifying on propensityscores)?

useCovariates Whether to use the covariate matrix in the cohortMethodDataobject in the out-

come model.

excludeCovariateIds

Exclude these covariates from the outcome model.

includeCovariateIds

Include only these covariates in the outcome model.

prior The prior used to fit the model. See createPriorfor details.

control The control object used to control the cross-validation used todetermine the hy-

perparameters of the prior (if applicable). SeecreateControl for details.

Details

Create an object defining the parameter values.

 ${\tt createGetDbCohortMethodDataArgs}$

Create a parameter object for the function getDbCohortMethodData

Description

Create a parameter object for the function getDbCohortMethodData

Usage

```
createGetDbCohortMethodDataArgs(studyStartDate = "", studyEndDate = "",
  excludeDrugsFromCovariates = TRUE, firstExposureOnly = FALSE,
  removeDuplicateSubjects = FALSE, restrictToCommonPeriod = FALSE,
  washoutPeriod = 0, covariateSettings)
```

Arguments

studyStartDate A calendar date specifying the minimum date that a cohort indexdate can appear. Date format is 'yyyymmdd'.

studyEndDate

A calendar date specifying the maximum date that a cohort indexdate can appear. Date format is 'yyyymmdd'. Important: the studyend data is also used to truncate risk windows, meaning nooutcomes beyond the study end date will be considered.

excludeDrugsFromCovariates

Should the target and comparator drugs (and their descendantconcepts) be excluded from the covariates? Note that this willwork if the drugs are actualy drug concept IDs (and not cohortIDs).

firstExposureOnly

Should only the first exposure per subject be included? Notethat this is typically done in the createStudyPopulationfunction, but can already be done here for efficiency reasons.

removeDuplicateSubjects

Remove subjects that are in both the treated and comparatorcohort? Note that this is typically done in thecreateStudyPopulation function, but can already be donehere for efficiency reasons.

restrictToCommonPeriod

Restrict the analysis to the period when both treatments are observed?

washoutPeriod

The mininum required continuous observation time prior to indexdate for a person to be included in the cohort. Note that thisis typically done in the createStudyPopulation function,but can already be done here for efficiency reasons.

covariateSettings

An object of type covariateSettings as created using thecreateCovariateSettings function in theFeatureExtraction package.

Details

Create an object defining the parameter values.

createMatchOnPsAndCovariatesArgs

Create a parameter object for the function matchOnPsAndCovariates

Description

Create a parameter object for the function matchOnPsAndCovariates

Usage

```
createMatchOnPsAndCovariatesArgs(caliper = 0.25,
  caliperScale = "standardized", maxRatio = 1, covariateIds)
```

Arguments

caliper The caliper for matching. A caliper is the distance which is acceptable for any

match. Observations which are outside of the caliper are dropped. A caliper of $\boldsymbol{0}$

means no caliper is used.

caliperScale

The scale on which the caliper is defined. Two scales are supported: caliper Scale = 'propensity score' or caliper Scale = 'standardized'. On the standardized scale, the caliper is interpreted in standard deviations of the propensity scoredistribution.

createMatchOnPsArgs 15

maxRatio The maximum number of persons int the comparator arm to be matched to each-

person in the treatment arm. A maxRatio of 0 means no maximum: all compara-

tors will be assigned to a treated person.

covariateIds One or more covariate IDs in the cohortMethodData object on whichsubjects

should be also matched.

Details

Create an object defining the parameter values.

createMatchOnPsArgs

Create a parameter object for the function matchOnPs

Description

Create a parameter object for the function matchOnPs

Usage

```
createMatchOnPsArgs(caliper = 0.25, caliperScale = "standardized",
    maxRatio = 1, stratificationColumns = c())
```

Arguments

caliper The caliper for matching. A caliper is the distance which isacceptable for any

match. Observations which are outside of the caliper are dropped. A caliper of $\boldsymbol{0}$

means no caliper is used.

caliperScale The scale on which the caliper is defined. Two scales are supported:caliperScale

= 'propensity score' or caliperScale = 'standardized'. On the standardized scale, the caliper isinterpreted in standard deviations of the propensity scoredistribu-

tion.

maxRatio The maximum number of persons int the comparator arm to be matched toeach

person in the treatment arm. A maxRatio of 0 means no maximum:all compara-

tors will be assigned to a treated person.

stratificationColumns

Names or numbers of one or more columns in the data data.frameon which subjects should be stratified prior to matching. No personswill be matched with persons outside of the strata identified by thevalues in these columns.

Details

Create an object defining the parameter values.

16 createPs

createPs	Create propensity scores
Ci Catci 5	Credite propertient, scores

Description

createPs creates propensity scores using a regularized logistic regression.

Usage

```
createPs(cohortMethodData, population, excludeCovariateIds = c(),
  includeCovariateIds = c(), errorOnHighCorrelation = TRUE,
  stopOnError = TRUE, prior = createPrior("laplace", exclude = c(0),
 useCrossValidation = TRUE), control = createControl(noiseLevel = "silent",
 cvType = "auto", tolerance = 2e-07, cvRepetitions = 10, startingVariance =
 0.01))
```

Arguments

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

population

A data frame describing the population. This should at least have a 'rowld' column corresponding to the rowId column in the cohortMethodData covariates object and a 'treatment' column. If population is not specified, the full popula-

tion in the cohortMethodData will be used.

excludeCovariateIds

Exclude these covariates from the propensity model.

includeCovariateIds

Include only these covariates in the propensity model.

errorOnHighCorrelation

If true, the function will test each covariate for correlation with the treatment assignment. If any covariate has an unusually high correlation (either positive

or negative), this will throw and error.

If an error occurrs, should the function stop? Else, the two cohorts will be stopOnError

assumed to be perfectly separable.

prior The prior used to fit the model. See createPrior for details.

control The control object used to control the cross-validation used to determine the

hyperparameters of the prior (if applicable). See createControl for details.

Details

createPs creates propensity scores using a regularized logistic regression.

Examples

```
data(cohortMethodDataSimulationProfile)
cohortMethodData <- simulateCohortMethodData(cohortMethodDataSimulationProfile, n = 1000)</pre>
ps <- createPs(cohortMethodData)</pre>
```

 ${\tt createStratifyByPsAndCovariatesArgs}$

Create a parameter object for the function stratifyByPsAndCovariates

Description

Create a parameter object for the function stratifyByPsAndCovariates

Usage

createStratifyByPsAndCovariatesArgs(numberOfStrata = 5, covariateIds)

Arguments

numberOfStrata Into how many strata should the propensity score be divided? Theboundaries of the strata are automatically defined to contain equalnumbers of treated persons.

 ${\tt covariateIds} \qquad {\tt One \ or \ more \ covariate \ IDs \ in \ the \ cohortMethodData \ object \ on \ which subjects}$

should also be stratified.

Details

Create an object defining the parameter values.

createStratifyByPsArgs

Create a parameter object for the function stratifyByPs

Description

Create a parameter object for the function stratifyByPs

Usage

```
createStratifyByPsArgs(numberOfStrata = 5, stratificationColumns = c())
```

Arguments

numberOfStrata How many strata? The boundaries of the strata are automatically defined to contain equal numbers of treated persons.

stratificationColumns

Names of one or more columns in the data data.frame on which subjects should also be stratified in addition to stratification on propensity score.

Details

Create an object defining the parameter values.

createStudyPopulation Create a study population

Description

Create a study population

Usage

```
createStudyPopulation(cohortMethodData, population = NULL, outcomeId,
  firstExposureOnly = FALSE, restrictToCommonPeriod = FALSE,
  washoutPeriod = 0, removeDuplicateSubjects = FALSE,
  removeSubjectsWithPriorOutcome = TRUE, priorOutcomeLookback = 99999,
  minDaysAtRisk = 1, riskWindowStart = 0, addExposureDaysToStart = FALSE,
  riskWindowEnd = 0, addExposureDaysToEnd = TRUE)
```

Arguments

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

population If specified, this population will be used as the starting point instead of the co-

horts in the cohortMethodData object.

outcomeId The ID of the outcome. If not specified, no outcome-specific transformations

will be performed.

firstExposureOnly

Should only the first exposure per subject be included? Note that this is typically done in the createStudyPopulation function,

restrictToCommonPeriod

Restrict the analysis to the period when both treatments are observed?

washoutPeriod The minimum required continuous observation time prior to index date for a

person to be included in the cohort.

removeDuplicateSubjects

Remove subjects that are in both the treated and comparator cohort?

 ${\tt remove Subjects With Prior Outcome}$

Remove subjects that have the outcome prior to the risk window start?

 $\verb"priorOutcomeLookback"$

How many days should we look back when identifying prior outcomes?

minDaysAtRisk The minimum required number of days at risk.

riskWindowStart

The start of the risk window (in days) relative to the index date (+ days of exposure if the addExposureDaysToStart parameter is specified).

 ${\it add} {\it Exposure Days To Start}$

Add the length of exposure the start of the risk window?

riskWindowEnd The end of the risk window (in days) relative to the index data (+ days of exposure if the addExposureDaysToEnd parameter is specified).

addExposureDaysToEnd

Add the length of exposure the risk window?

createTrimByPsArgs 19

Details

Create a study population by enforcing certain inclusion and exclusion criteria, defining a risk window, and determining which outcomes fall inside the risk window.

Value

A data frame specifying the study population. This data frame will have the following columns:

rowId A unique identifier for an exposure

subjectId The person ID of the subject

cohortStartdate The index date

outcomeCount The number of outcomes observed during the risk window

timeAtRisk The number of days in the risk window

survivalTime The number of days until either the outcome or the end of the risk window

createTrimByPsArgs

Create a parameter object for the function trimByPs

Description

Create a parameter object for the function trimByPs

Usage

```
createTrimByPsArgs(trimFraction = 0.05)
```

Arguments

trimFraction

This fraction will be removed from each treatment group. In the treatmentgroup, persons with the highest propensity scores will be removed, in the comparator group person with the lowest scores will be removed.

Details

Create an object defining the parameter values.

20 drawAttritionDiagram

```
createTrimByPsToEquipoiseArgs
```

Create a parameter object for the function trimByPsToEquipoise

Description

Create a parameter object for the function trimByPsToEquipoise

Usage

```
createTrimByPsToEquipoiseArgs(bounds = c(0.25, 0.75))
```

Arguments

bounds

The upper and lower bound on the preference score for keeping persons

Details

Create an object defining the parameter values.

Description

drawAttritionDiagram draws the attition diagram, showing how many people were excluded from the study population, and for what reasons.

Usage

```
drawAttritionDiagram(object, treatmentLabel = "Treated",
  comparatorLabel = "Comparator", fileName = NULL)
```

Arguments

object

Either an object of type cohortMethodData, a population object generated by functions like createStudyPopulation, or an object of type outcomeModel.

treatmentLabel A label to us for the treated cohort.

comparatorLabel

A label to us for the comparator cohort.

fileName

Name of the file where the plot should be saved, for example 'plot.png'. See the

function ggsave in the ggplot2 package for supported file formats.

Value

A ggplot object. Use the ggsave function to save to file in a different format.

fitOutcomeModel 21

fitOutcomeModel Create an outcome model, and compute the relative risk
--

Description

fitOutcomeModel creates an outcome model, and computes the relative risk

Usage

```
fitOutcomeModel(population, cohortMethodData, modelType = "logistic",
    stratified = TRUE, useCovariates = TRUE, excludeCovariateIds = c(),
    includeCovariateIds = c(), prior = createPrior("laplace",
    useCrossValidation = TRUE), control = createControl(cvType = "auto",
    startingVariance = 0.01, tolerance = 2e-07, cvRepetitions = 10, noiseLevel =
    "quiet"))
```

Arguments

population A population object generated by createStudyPopulation, potentially filtered

by other functions.

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

modelType The type of outcome model that will be used. Possible values are "logistic",

"poisson", or "cox".

stratified Should the regression be conditioned on the strata defined in the population

object (e.g. by matching or stratifying on propensity scores)?

useCovariates Whether to use the covariate matrix in the cohortMethodData object in the

outcome model.

excludeCovariateIds

Exclude these covariates from the outcome model.

includeCovariateIds

Include only these covariates in the outcome model.

prior The prior used to fit the model. See createPrior for details.

control The control object used to control the cross-validation used to determine the

hyperparameters of the prior (if applicable). See createControl for details.

Value

An object of class outcomeModel. Generic function summary, coef, and confint are available.

 ${\tt getAttritionTable}$

Get the attrition table for a population

Description

Get the attrition table for a population

Usage

```
getAttritionTable(object)
```

Arguments

object

Either an object of type cohortMethodData, a population object generated by functions like createStudyPopulation, or an object of type outcomeModel.

Value

A data frame specifying the number of people and exposures in the population after specific steps of filtering.

getDbCohortMethodData Get the cohort data from the server

Description

This function executes a large set of SQL statements against the database in OMOP CDM format to extract the data needed to perform the analysis.

Usage

```
getDbCohortMethodData(connectionDetails, cdmDatabaseSchema,
  oracleTempSchema = cdmDatabaseSchema, targetId, comparatorId, outcomeIds,
  studyStartDate = "", studyEndDate = "",
  exposureDatabaseSchema = cdmDatabaseSchema, exposureTable = "drug_era",
  outcomeDatabaseSchema = cdmDatabaseSchema,
  outcomeTable = "condition_occurrence", cdmVersion = "5",
  excludeDrugsFromCovariates = TRUE, firstExposureOnly = FALSE,
  removeDuplicateSubjects = FALSE, restrictToCommonPeriod = FALSE,
  washoutPeriod = 0, covariateSettings)
```

Arguments

connectionDetails

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

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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.

A unique identifier to define the target cohort. If exposureTable = DRUG_ERA, targetId is a CONCEPT_ID and all descendant concepts within that CONCEPT_ID will be used to define the cohort. If exposureTable <> DRUG_ERA, targetId is

used to select the cohort_concept_id in the cohort-like table.

comparatorId A unique identifier to define the comparator cohort. If exposureTable = DRUG_ERA,

comparatorId is a CONCEPT_ID and all descendant concepts within that CON-CEPT_ID will be used to define the cohort. If exposureTable <> DRUG_ERA, comparatorId is used to select the cohort_concept_id in the cohort-like table.

outcomeIds A list of cohort definition ids used to define outcomes.

studyStartDate A calendar date specifying the minimum date that a cohort index date can ap-

pear. Date format is 'yyyymmdd'.

studyEndDate A calendar date specifying the maximum date that a cohort index date can appear. Date format is 'yyyymmdd'. Important: the study end data is also used to

truncate risk windows, meaning no outcomes beyond the study end date will be

considered.

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_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 <> CONDI-

TION_OCCURRENCE, then expectation is outcomeTable has format of CO-HORT table: COHORT_DEFINITION_ID, SUBJECT_ID, COHORT_START_DATE,

COHORT_END_DATE.

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

 ${\tt excludeDrugsFromCovariates}$

Should the target and comparator drugs (and their descendant concepts) be excluded from the covariates? Note that this will work if the drugs are actualy drug concept IDs (and not cohort IDs).

firstExposureOnly

Should only the first exposure per subject be included? Note that this is typically done in the createStudyPopulation function, but can already be done here for efficiency reasons.

removeDuplicateSubjects

Remove subjects that are in both the treated and comparator cohort? Note that this is typically done in the createStudyPopulation function, but can already be done here for efficiency reasons.

restrictToCommonPeriod

Restrict the analysis to the period when both treatments are observed?

washoutPeriod

The minimum required continuous observation time prior to index date for a person to be included in the cohort. Note that this is typically done in the createStudyPopulation function, but can already be done here for efficiency reasons.

covariateSettings

An object of type covariateSettings as created using the createCovariateSettings function in the FeatureExtraction package.

Details

Based on the arguments, the treatment and comparator cohorts are retrieved, as well as outcomes occurring in exposed subjects. The treatment and comparator cohorts can be identified using the drug_era table, or through user-defined cohorts in a cohort table either inside the CDM instance or in a separate schema. Similarly, outcomes are identified using the condition_era table or through user-defined cohorts in a cohort table either inside the CDM instance or in a separate schema. Covariates are automatically extracted from the appropriate tables within the CDM. Important: The target and comparator drug must not be included in the covariates, including any descendant concepts. If the targetId and comparatorId arguments represent real concept IDs, you can set the excludeDrugsFromCovariates argument to TRUE and automatically the drugs and their descendants will be excluded from the covariates. However, if the targetId and comparatorId arguments do not represent concept IDs, you will need to manually add the drugs and descendants to the excludedCovariateConceptIds of the covariateSettings argument.

Value

Returns an object of type cohortMethodData, containing information on the cohorts, their outcomes, and baseline covariates. Information about multiple outcomes can be captured at once for efficiency reasons. This object is a list with the following components:

outcomes A data frame listing the outcomes per person, including the time to event, and the outcome id. Outcomes are not yet filtered based on risk window, since this is done at a later stage.

cohorts A data frame listing the persons in each cohort, listing their exposure status as well as the time to the end of the observation period and time to the end of the cohort (usually the end of the exposure era).

covariates An ffdf object listing the baseline covariates per person in the two cohorts. This is done using a sparse representation: covariates with a value of 0 are omitted to save space.

covariateRef An ffdf object describing the covariates that have been extracted.

metaData A list of objects with information on how the cohortMethodData object was constructed.

The generic print() and summary() functions have been implemented for this object.

getFollowUpDistribution

Get the distribution of follow-up time

Description

Get the distribution of follow-up time

Usage

```
getFollowUpDistribution(population, quantiles = c(0, 0.25, 0.5, 0.75, 1))
```

Arguments

population A data frame describing the study population as created using the createStudyPopulation

function. This should at least have these columns: treatment, timeAtRisk.

quantiles The quantiles of the population to compute minimum follow-up time for.

Details

Get the distribution of follow-up time as quantiles. Follow-up time is defined as time-at-risk, so not censored at the outcome.

Value

A data frame with per treatment group at each quantile the amount of follow-up time available.

getOutcomeModel Get the outcome model

Description

getOutcomeModel shows the full outcome model, so showing the betas of all variables included in the outcome model, not just the treatment variable.

Usage

getOutcomeModel(outcomeModel, cohortMethodData)

Arguments

outcomeModel An object of type outcomeModel as generated using he createOutcomeMode function.

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

Details

Shows the coefficients and names of the covariates with non-zero coefficients.

26 grepCovariateNames

Examples

todo

getPsModel

Get the propensity model

Description

getPsModel shows the propensity score model

Usage

```
getPsModel(propensityScore, cohortMethodData)
```

Arguments

propensityScore

The propensity scores as generated using the createPs function.

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

Details

Shows the coefficients and names of the covariates with non-zero coefficients.

Examples

todo

grepCovariateNames

Extract covariate names

Description

Extracts covariate names using a regular-expression.

Usage

```
grepCovariateNames(pattern, object)
```

Arguments

pattern A regular expression with which to name covariate names object An R object of type cohortMethodData or covariateData.

Details

This function extracts covariate names that match a regular-expression for a cohortMethodData or covariateData object.

insertDbPopulation 27

Value

Returns a data.frame containing information about covariates that match a regular expression. This data.frame has the following columns:

covariateId Numerical identifier for use in model fitting using these covariates

covariateName Text identifier analysisId Analysis identifier

conceptId OMOP common data model concept identifier, or 0

insertDbPopulation

Insert a population into a database

Description

Insert a population into a database

Usage

```
insertDbPopulation(population, cohortIds = c(1, 0), connectionDetails, cohortDatabaseSchema, cohortTable = "cohort", createTable = FALSE, dropTableIfExists = TRUE, cdmVersion = "5")
```

Arguments

population Either an object of type cohortMethodData or a population object generated by

functions like createStudyPopulation.

cohortIds The IDs to be used for the treated and comparator cohort, respectively.

connectionDetails

An R object of type

 ${\tt connectionDetails}\ created\ using\ the\ function\ {\tt createConnectionDetails}\ in$

the DatabaseConnector package.

cohortDatabaseSchema

The name of the database schema where the data will be written. Requires write permissions to this database. On SQL Server, this should specify both the

database and the schema, so for example 'cdm_instance.dbo'.

cohortTable The name of the table in the database schema where the data will be written.

createTable Should a new table be created? If not, the data will be inserted into an existing

table.

dropTableIfExists

If createTable = TRUE and the table already exists it will be overwritten.

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

Details

Inserts a population table into a database. The table in the database will have the same structure as the 'cohort' table in the Common Data Model.

28 loadCohortMethodData

loadCmAnalysisList

Load a list of cmAnalysis from file

Description

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

Usage

```
loadCmAnalysisList(file)
```

Arguments

file

The name of the file

Value

A list of objects of type cmAnalysis.

loadCohortMethodData

Load the cohort data from a folder

Description

loadCohortMethodData loads an object of type cohortMethodData from a folder in the file system.

Usage

```
loadCohortMethodData(file, readOnly = TRUE)
```

Arguments

file

The name of the folder containing the data.

readOnly

If true, the data is opened read only.

Details

The data will be written to a set of files in the folder specified by the user.

Value

An object of class cohortMethodData.

Examples

todo

loadDrugComparatorOutcomesList

Load a list of drugComparatorOutcomes from file

Description

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

Usage

loadDrugComparatorOutcomesList(file)

Arguments

file The name of the file

Value

A list of objects of type drugComparatorOutcome.

matchOnPs

Match persons by propensity score

Description

matchOnPs uses the provided propensity scores to match treated to comparator persons.

Usage

```
matchOnPs(population, caliper = 0.25, caliperScale = "standardized",
    maxRatio = 1, stratificationColumns = c())
```

Arguments

population A data frame with the three columns described below.

caliper The caliper for matching. A caliper is the distance which is acceptable for any

match. Observations which are outside of the caliper are dropped. A caliper of

0 means no caliper is used.

caliperScale The scale on which the caliper is defined. Two scales are supported:

caliperScale = 'propensity score' or caliperScale = 'standardized'. On the standardized scale, the caliper is interpreted in standard deviations of the

propensity score distribution.

maxRatio The maximum number of persons int the comparator arm to be matched to each

person in the treatment arm. A maxRatio of 0 means no maximum: all com-

parators will be assigned to a treated person.

stratificationColumns

Names or numbers of one or more columns in the data data.frame on which subjects should be stratified prior to matching. No persons will be matched with persons outside of the strata identified by the values in these columns.

Details

The data frame should have at least the following three columns:

```
rowId (numeric) A unique identifier for each row (e.g. the person ID)
treatment (integer) Column indicating whether the person is in the treated (1) or comparator (0) group
propensityScore (numeric) Propensity score
```

This function implements the greedy variable-ratio matching algorithm described in Rassen et al (2012).

Value

Returns a date frame with the same columns as the input data plus one extra column: stratumId. Any rows that could not be matched are removed

References

Rassen JA, Shelat AA, Myers J, Glynn RJ, Rothman KJ, Schneeweiss S. (2012) One-to-many propensity score matching in cohort studies, Pharmacoepidemiology and Drug Safety, May, 21 Suppl 2:69-80.

Examples

matchOnPsAndCovariates

Match by propensity score as well as other covariates

Description

matchOnPsAndCovariates uses the provided propensity scores and a set of covariates to match treated to comparator persons.

Usage

```
matchOnPsAndCovariates(population, caliper = 0.25,
    caliperScale = "standardized", maxRatio = 1, cohortMethodData,
    covariateIds)
```

matchOnPsAndCovariates 31

Arguments

population A data frame with the three columns described below.

caliper The caliper for matching. A caliper is the distance which is acceptable for any

match. Observations which are outside of the caliper are dropped. A caliper of

0 means no caliper is used.

caliperScale The scale on which the caliper is defined. Two scales are supported:

caliperScale = 'propensity score' or caliperScale = 'standardized'. On the standardized scale, the caliper is interpreted in standard deviations of the

propensity score distribution.

maxRatio The maximum number of persons int the comparator arm to be matched to each

person in the treatment arm. A maxRatio of 0 means no maximum: all com-

parators will be assigned to a treated person.

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

covariateIds One or more covariate IDs in the cohortMethodData object on which subjects

should be also matched.

Details

The data frame should have at least the following three columns:

rowId (numeric) A unique identifier for each row (e.g. the person ID)

treatment (integer) Column indicating whether the person is in the treated (1) or comparator

(0) group

propensityScore (numeric) Propensity score

This function implements the greedy variable-ratio matching algorithm described in Rassen et al (2012).

Value

Returns a date frame with the same columns as the input data plus one extra column: stratumId. Any rows that could not be matched are removed

References

Rassen JA, Shelat AA, Myers J, Glynn RJ, Rothman KJ, Schneeweiss S. (2012) One-to-many propensity score matching in cohort studies, Pharmacoepidemiology and Drug Safety, May, 21 Suppl 2:69-80.

Examples

todo

```
plotCovariateBalanceOfTopVariables
```

Plot variables with largest imbalance

Description

Create a plot showing those variables having the largest imbalance before matching, and those variables having the largest imbalance after matching. Requires running computeCovariateBalance first.

Usage

```
plotCovariateBalanceOfTopVariables(balance, n = 20, maxNameWidth = 100,
  fileName = NULL, beforeLabel = "before matching",
  afterLabel = "after matching")
```

Arguments

balance A data frame created by the computeCovariateBalance funcion.

n Count of variates to plot.

maxNameWidth Covariate names longer than this number of characters are truncated to create a

nicer plot.

fileName Name of the file where the plot should be saved, for example 'plot.png'. See the

function ggsave in the ggplot2 package for supported file formats.

beforeLabel Label for identifying data before matching / stratification / trimming.

Label for identifying data after matching / stratification / trimming.

Value

A ggplot object. Use the ggsave function to save to file in a different format.

```
\verb|plotCovariateBalanceScatterPlot|
```

Create a scatterplot of the covariate balance

Description

Create a scatterplot of the covariate balance, showing all variables with balance before and after matching on the x and y axis respectively. Requires running computeCovariateBalance first.

Usage

```
plotCovariateBalanceScatterPlot(balance, absolute = TRUE, threshold = 0,
  fileName = NULL, beforeLabel = "Before matching",
  afterLabel = "After matching")
```

Arguments

balance A data frame created by the computeCovariateBalance funcion.

absolute Should the absolute value of the difference be used?

threshold Show a threshold value for after matching standardized difference.

fileName Name of the file where the plot should be saved, for example 'plot.png'. See the

function ggsave in the ggplot2 package for supported file formats.

beforeLabel Label for the x-axis. afterLabel Label for the y-axis.

Value

A ggplot object. Use the ggsave function to save to file in a different format.

plotFollowUpDistribution

Plot the distribution of follow-up time

Description

Plot the distribution of follow-up time

Usage

```
plotFollowUpDistribution(population, targetLabel = "Target",
  comparatorLabel = "Comparator", yScale = "percent", logYScale = FALSE,
  dataCutoff = 0.95, title = "Follow-up distribution", fileName = NULL)
```

Arguments

population A data frame describing the study population as created using the createStudyPopulation

function. This should at least have these columns: treatment, timeAtRisk.

targetLabel A label to us for the target cohort.

comparatorLabel

A label to us for the comparator cohort.

yScale Should be either 'percent' or 'count'.
logYScale Should the Y axis be on the log scale?

dataCutoff Fraction of the data (number censored) after which the graph will not be shown.

title The main title of the plot.

fileName Name of the file where the plot should be saved, for example 'plot.png'. See the

function ggsave in the ggplot2 package for supported file formats.

Details

Plot the distribution of follow-up time, stratified by treatment group. Follow-up time is defined as time-at-risk, so not censored at the outcome.

Value

A ggplot object. Use the ggsave function to save to file in a different format.

34 plotKaplanMeier

Plot the Kaplan-Meier curve	plotKaplanMeier

Description

plotKaplanMeier creates the Kaplain-Meier survival plot. Based (partially) on recommendations in Pocock et al (2002).

Usage

```
plotKaplanMeier(population, censorMarks = FALSE, confidenceIntervals = TRUE,
  includeZero = FALSE, dataTable = TRUE, dataCutoff = 0.9,
  treatmentLabel = "Treated", comparatorLabel = "Comparator", title,
  fileName = NULL)
```

Arguments

population	A population object	t generated by createStudy	yPopulation, potentially filtered
------------	---------------------	----------------------------	-----------------------------------

by other functions.

censorMarks Whether or not to include censor marks in the plot.

confidenceIntervals

Plot 95 percent confidence intervals? Default is TRUE, as recommended by

Pocock et al.

includeZero Should the y axis include zero, or only go down to the lowest observed survival?

The default is FALSE, as recommended by Pocock et al.

dataTable Should the numbers at risk be shown in a table? Default is TRUE, as recom-

mended by Pocock et al.

dataCutoff Fraction of the data (number censored) after which the graph will not be shown.

The default is 90 percent as recommended by Pocock et al.

treatmentLabel A label to us for the treated cohort.

comparatorLabel

A label to us for the comparator cohort.

title The main title of the plot.

fileName Name of the file where the plot should be saved, for example 'plot.png'. See the

function ggsave in the ggplot2 package for supported file formats.

Value

A ggplot object. Use the ggsave function to save to file in a different format.

References

Pocock SJ, Clayton TC, Altman DG. (2002) Survival plots of time-to-event outcomes in clinical trials: good practice and pitfalls, Lancet, 359:1686-89.

plotPs 35

plotPs	Plot the propensity score distribution

Description

plotPs shows the propensity (or preference) score distribution

Usage

```
plotPs(data, unfilteredData = NULL, scale = "preference",
  type = "density", binWidth = 0.05, treatmentLabel = "Treated",
  comparatorLabel = "Comparator", fileName = NULL)
```

Arguments

data A data frame with at least the two columns described below unfilteredData To be used when computing preference scores on data from which subjects have already been removed, e.g. through trimming and/or matching. This data frame should have the same structure as data. scale The scale of the graph. Two scales are supported: scale = 'propensity' or scale = 'preference'. The preference score scale is defined by Walker et al (2013).Type of plot. Two possible values: type = 'density' or type = 'histogram' type binWidth For histograms, the width of the bins treatmentLabel A label to us for the treated cohort. comparator LabelA label to us for the comparator cohort. Name of the file where the plot should be saved, for example 'plot.png'. See the fileName

Details

The data frame should have a least the following two columns:

treatment	(integer)	Column indicating whether the person is in the treated (1) or comparat	
		(0) group	
propensityScore	(numeric)	Propensity score	

function ggsave in the ggplot2 package for supported file formats.

Value

A ggplot object. Use the ggsave function to save to file in a different format.

References

Walker AM, Patrick AR, Lauer MS, Hornbrook MC, Marin MG, Platt R, Roger VL, Stang P, and Schneeweiss S. (2013) A tool for assessing the feasibility of comparative effectiveness research, Comparative Effective Research, 3, 11-20

36 runCmAnalyses

Examples

```
treatment <- rep(0:1, each = 100)
propensityScore <- c(rnorm(100, mean = 0.4, sd = 0.25), rnorm(100, mean = 0.6, sd = 0.25))
data <- data.frame(treatment = treatment, propensityScore = propensityScore)
data <- data[data$propensityScore > 0 & data$propensityScore < 1, ]
plotPs(data)</pre>
```

runCmAnalyses

Run a list of analyses

Description

Run a list of analyses

Usage

```
runCmAnalyses(connectionDetails, cdmDatabaseSchema,
  oracleTempSchema = cdmDatabaseSchema,
  exposureDatabaseSchema = cdmDatabaseSchema, exposureTable = "drug_era",
  outcomeDatabaseSchema = cdmDatabaseSchema,
  outcomeTable = "condition_occurrence", cdmVersion = 4,
  outputFolder = "./CohortMethodOutput", cmAnalysisList,
  drugComparatorOutcomesList, refitPsForEveryOutcome = FALSE,
  getDbCohortMethodDataThreads = 1, createPsThreads = 1, psCvThreads = 1,
  createStudyPopThreads = 1, trimMatchStratifyThreads = 1,
  computeCovarBalThreads = 1, fitOutcomeModelThreads = 1,
  outcomeCvThreads = 1, outcomeIdsOfInterest)
```

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.

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.

runCmAnalyses 37

outcomeDatabaseSchema

outcomeTable

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.

read permissions to any dataou

The tablename that contains the outcome cohorts. If outcome Table <> CONDI-

TION_OCCURRENCE, then expectation is outcomeTable has format of CO-HORT 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.

 $\verb|cmAnalysisList| A list of objects of type \verb|cmAnalysis| as created using the \verb|createCmAnalysis| \\$

function.

drugComparatorOutcomesList

A list of objects of type drugComparatorOutcomes as created using the createDrugComparatorOutcomes function.

refitPsForEveryOutcome

Should the propensity model be fitted for every outcome (i.e. after people who already had the outcome are removed)? If false, a single propensity model will be fitted, and people who had the outcome previously will be removed afterwards.

getDbCohortMethodDataThreads

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

createPsThreads

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

psCvThreads The number of parallel threads to use for the cross-validation when estimating

the hyperparameter for the propensity model. Note that the total number of CV

threads at one time could be 'createPsThreads * psCvThreads'.

createStudyPopThreads

The number of parallel threads to use for creating the study population.

trimMatchStratifyThreads

The number of parallel threads to use for trimming, matching and stratifying.

computeCovarBalThreads

The number of parallel threads to use for computing the covariate balance.

fitOutcomeModelThreads

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

outcomeCvThreads

The number of parallel threads to use for the cross-validation when estimating the hyperparameter for the outcome model. Note that the total number of CV threads at one time could be 'fitOutcomeModelThreads * outcomeCvThreads'.

outcomeIdsOfInterest

If provided, creation of non-essential files will be skipped for all other outcome IDs. This could be helpful to speed up analyses with many controls.

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

38 saveCmAnalysisList

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.

Value

studyPopFile

A data frame with the following columns:

analysisId The unique identifier for a set of analysis choices.

targetId The ID of the target drug.
comparatorId The ID of the comparator group.

 $\begin{array}{ll} \text{excludedCovariateConceptIds} & \text{The ID(s) of concepts that cannot be used to construct covariates.} \\ \text{includedCovariateConceptIds} & \text{The ID(s) of concepts that should be used to construct covariates.} \\ \end{array}$

outcomeId The ID of the outcome cohortMethodDataFolder The ID of the outcome.

sharedPsFile The name of the file containing the propensity scores of the shared

propensity model. This model is used to create the outcome-specific

propensity scores by removing people with prior outcomes. The name of the file containing the study population (prior

and trimming, matching, or stratification on the PS.

psFile The name of file containing the propensity scores for a specific

outcomes (ie after people with prior outcomes have been removed).

strataFile The name of the file containing the identifiers of the population

after any trimming, matching or stratifying, including their strata.

covariateBalanceFile The name of the file containing the covariate balance (ie. the

 $output\ of\ the\ compute {\tt CovariateBalance}\ function.$

outcomeModelFile The name of the file containing the outcome model.

saveCmAnalysisList Save a list of cmAnalysis to file

Description

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

Usage

```
saveCmAnalysisList(cmAnalysisList, file)
```

Arguments

cmAnalysisList The cmAnalysis list to be written to file

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

saveCohortMethodData 39

saveCohortMethodData Save the cohort data to folder

Description

saveCohortMethodData saves an object of type cohortMethodData to folder.

Usage

saveCohortMethodData(cohortMethodData, file)

Arguments

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

file

The name of the folder where the data will be written. The folder should not yet exist.

Details

The data will be written to a set of files in the folder specified by the user.

Examples

todo

saveDrugComparatorOutcomesList

Save a list of drugComparatorOutcome to file

Description

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

Usage

save Drug Comparator Outcomes List (drug Comparator Outcomes List, file)

Arguments

drugComparatorOutcomesList

The drugComparatorOutcomes list to be written to file

The name of the file where the results will be written

40 stratifyByPs

simulateCohortMethodData

Generate simulated data

Description

simulateCohortMethodData creates a cohortMethodData object with simulated data.

Usage

```
simulateCohortMethodData(profile, n = 10000)
```

Arguments

profile An object of type cohortMethodDataSimulationProfile as generated using

the

 ${\tt createCohortMethodDataSimulationProfile\ function}.$

n The size of the population to be generated.

Details

This function generates simulated data that is in many ways similar to the original data on which the simulation profile is based. The contains same outcome, comparator, and outcome concept IDs, and the covariates and their 1st order statistics should be comparable.

Value

An object of type cohortMethodData.

stratifyByPs

Stratify persons by propensity score

Description

stratifyByPs uses the provided propensity scores to stratify persons. Additional stratification variables for stratifications can also be used.

Usage

```
stratifyByPs(population, numberOfStrata = 5, stratificationColumns = c())
```

Arguments

population A data frame with the three columns described below

numberOfStrata How many strata? The boundaries of the strata are automatically defined to

contain equal numbers of treated persons.

 ${\it stratification Columns}$

Names of one or more columns in the data data.frame on which subjects should also be stratified in addition to stratification on propensity score.

Details

The data frame should have the following three columns:

rowId (numeric) A unique identifier for each row (e.g. the person ID)

treatment (integer) Column indicating whether the person is in the treated (1) or comparator

(0) group

propensityScore (numeric) Propensity score

Value

Returns a date frame with the same columns as the input data plus one extra column: stratumId.

Examples

```
rowId <- 1:200
treatment <- rep(0:1, each = 100)
propensityScore <- c(runif(100, min = 0, max = 1), runif(100, min = 0, max = 1))
data <- data.frame(rowId = rowId, treatment = treatment, propensityScore = propensityScore)
result <- stratifyByPs(data, 5)</pre>
```

stratifyByPsAndCovariates

Stratify persons by propensity score and other covariates

Description

 ${\sf stratifyByPsAndCovariates}$ uses the provided propensity scores and covariates to stratify persons.

Usage

```
stratifyByPsAndCovariates(population, numberOfStrata = 5, cohortMethodData,
    covariateIds)
```

Arguments

population A data frame with the three columns described below

numberOfStrata Into how many strata should the propensity score be divided? The boundaries of

the strata are automatically defined to contain equal numbers of treated persons.

cohortMethodData

An object of type cohortMethodData as generated using getDbCohortMethodData.

covariateIds One or more covariate IDs in the cohortMethodData object on which subjects

should also be stratified.

42 summarizeAnalyses

Details

The data frame should have the following three columns:

rowId (integer) A unique identifier for each row (e.g. the person ID)

treatment (integer) Column indicating whether the person is in the treated (1) or comparator

(0) group

propensityScore (numeric) Propensity score

Value

Returns a date frame with the same columns as the input population plus one extra column: stratumId.

Examples

todo

summarizeAnalyses

Create a summary report of the analyses

Description

Create a summary report of the analyses

Usage

summarizeAnalyses(referenceTable)

Arguments

referenceTable A data.frame as created by the runCmAnalyses function.

Value

A data frame with the following columns:

analysisId The unique identifier for a set of analysis choices.

targetId The ID of the target drug.
comparatorId The ID of the comparator group.

indicationConceptIds The ID(s) of indications in which to nest to study.

outcomeId The ID of the outcome.
rr The estimated effect size.

ci95lb The lower bound of the 95 percent confidence interval. ci95ub The upper bound of the 95 percent confidence interval.

treated The number of subjects in the treated group (after any trimming and matching).

The number of subjects in the comparator group (after any trimming and matching).

The number of outcomes in the treated group (after any trimming and matching).

The number of outcomes in the comparator group (after any trimming and matching).

matching).

logRr The log of the estimated relative risk.

seLogRr The standard error of the log of the estimated relative risk.

trimByPsToEquipoise

trimByPs	Trim persons by propensity score	

Description

trimByPs uses the provided propensity scores to trim subjects with extreme scores.

Usage

```
trimByPs(population, trimFraction = 0.05)
```

Arguments

population A data frame with the three columns described below

trimFraction This fraction will be removed from each treatment group. In the treatment group,

persons with the highest propensity scores will be removed, in the comparator

group person with the lowest scores will be removed.

Details

The data frame should have the following three columns:

rowId (numeric) A unique identifier for each row (e.g. the person ID)

treatment (integer) Column indicating whether the person is in the treated (1) or comparator

(0) group

propensityScore (numeric) Propensity score

Value

Returns a date frame with the same three columns as the input.

Examples

```
rowId <- 1:2000
treatment <- rep(0:1, each = 1000)
propensityScore <- c(runif(1000, min = 0, max = 1), runif(1000, min = 0, max = 1))
data <- data.frame(rowId = rowId, treatment = treatment, propensityScore = propensityScore)
result <- trimByPs(data, 0.05)</pre>
```

trimByPsToEquipoise Keep only

Keep only persons in clinical equipoise

Description

trimByPsToEquipoise uses the preference score to trim subjects that are not in clinical equipoise

Usage

```
trimByPsToEquipoise(population, bounds = c(0.25, 0.75))
```

Arguments

population A data frame with at least the three columns described below

bounds The upper and lower bound on the preference score for keeping persons

Details

The data frame should have the following three columns:

rowId (numeric) A unique identifier for each row (e.g. the person ID)

treatment (integer) Column indicating whether the person is in the treated (1) or comparator

(0) group

propensityScore (numeric) Propensity score

Value

Returns a date frame with the same three columns as the input.

References

Walker AM, Patrick AR, Lauer MS, Hornbrook MC, Marin MG, Platt R, Roger VL, Stang P, and Schneeweiss S. (2013) A tool for assessing the feasibility of comparative effectiveness research, Comparative Effective Research, 3, 11-20

Examples

```
rowId <- 1:2000
treatment <- rep(0:1, each = 1000)
propensityScore <- c(runif(1000, min = 0, max = 1), runif(1000, min = 0, max = 1))
data <- data.frame(rowId = rowId, treatment = treatment, propensityScore = propensityScore)
result <- trimByPsToEquipoise(data)</pre>
```

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