# Package 'CohortMethod'

December 8, 2014

Type Package

Title What the package does (short line)

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| simulateCohortData  | <br> |  |  |  |  |  |  | <br> |  |  |  |  |  |
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| stratifyByPs        |      |  |  |  |  |  |  |      |  |  |  |  |  |
| rimByPs             |      |  |  |  |  |  |  | <br> |  |  |  |  |  |
| trimByPsToEquipoise |      |  |  |  |  |  |  | <br> |  |  |  |  |  |

computeCovariateBalance

Compute covariate balance before and after matching and trimming

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# **Description**

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For every covariate, prevalence in treatment and comparator groups before and after matching/trimming are computed.

# Usage

```
computeCovariateBalance(restrictedCohorts, cohortData,
  outcomeConceptId = NULL)
```

#### **Arguments**

restrictedCohorts

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

 ${\tt cohortData} \qquad {\tt An~object~of~type~cohortData}~ as~ {\tt generated~using~getDbCohortData}.\\ {\tt outcomeConceptId}$ 

The concept ID of the outcome. Persons marked for removal for the outcome will be removed when computing the balance before matching/trimming.

#### **Details**

The restrictedCohorts 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.

| computePsAuc Compute the area under the ROC curve |  |
|---|--|
|---|--|

# **Description**

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

#### Usage

```
computePsAuc(data)
```

#### **Arguments**

data

A data frame with at least the two columns described below

#### **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 (real) 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>
```

 ${\tt createCohortDataSimulationProfile}$ 

Create simulation profile

# **Description**

createCohortDataSimulationProfile creates a profile based on the provided cohortData object, which can be used to generate simulated data that has similar characteristics.

#### Usage

```
createCohortDataSimulationProfile(cohortData)
```

# **Arguments**

cohortData

An object of type cohortData as generated using getDbCohortData.

# **Details**

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

# Value

 $An\ object\ of\ type\ cohort {\tt DataSimulationProfile}.$ 

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createPs

Create propensity scores

#### **Description**

createPs creates propensity scores using a regularized logistic regression.

#### Usage

```
createPs(cohortData, outcomeConceptId = NULL, prior = createPrior("laplace",
  useCrossValidation = TRUE), control = createControl(lowerLimit = 0.01,
  upperLimit = 10, fold = 5, noiseLevel = "quiet"))
```

#### **Arguments**

cohortData An object of type cohortData as generated using getDbCohortData. outcomeConceptId

The concept ID of the outcome. Persons marked for removal for the outcome

will be removed prior to creating the propensity score model.

prior The prior used to fit the model. See ?createPrior in the Cyclops package for

details.

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

perparameters of the prior (if applicable). See ?createControl in the Cyclops

package for details.

# **Details**

createPs creates propensity scores using a regularized logistic regression.

#### **Examples**

#todo

fitOutcomeModel

Create an outcome model, and compute the relative risk

#### **Description**

fitOutcomeModel creates an outcome model, and computes the relative risk

```
fitOutcomeModel(outcomeConceptId, cohortData, strata = NULL,
  riskWindowStart = 0, riskWindowEnd = 9999, addExposureDaysToEnd = FALSE,
  useCovariates = TRUE, fitModel = TRUE, modelType = "cox",
  prior = createPrior("laplace", useCrossValidation = TRUE),
  control = createControl(lowerLimit = 0.01, upperLimit = 10, fold = 5,
  noiseLevel = "quiet"))
```

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#### **Arguments**

riskWindowEnd

cohortData An object of type cohortData as generated using getDbCohortData.

strata A data frame specifying the (matched and/or trimmed) subpopulation to be used

in the study, as well as their strata (for conditional models). This data frame should have at least a RowId, and a StratumId when including stratification.

The maximum length (in days) of the risk window.

useCovariates Whether to use the covariate matrix in the cohortData in the outcome model. fitModel If false, the model will not be fit, and only summary statistics are available.

modelType The type of model to be fitted. See details for options.

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

#### **Details**

The model type can be one of these:

lr Logistic regression

clr Conditional logistic regression

cox Cox regression (stratified or not, depending on whether stata is specified)

pr Poisson regression

cpr Conditional Poisson regression

#### Value

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

# **Examples**

#todo

getDbCohortData Get the cohort data from the server

# Description

Todo: add description

```
getDbCohortData(connectionDetails, cdmSchema = "CDM4_SIM",
    resultsSchema = "scratch", targetDrugConceptId = 755695,
    comparatorDrugConceptId = 739138, indicationConceptIds = 439926,
    washoutWindow = 183, indicationLookbackWindow = 183,
    studyStartDate = "", studyEndDate = "", exclusionConceptIds = c(4027133,
    4032243, 4146536, 2002282, 2213572, 2005890, 43534760, 21601019),
    outcomeConceptIds = 194133, outcomeConditionTypeConceptIds = c(38000215,
    38000216, 38000217, 38000218, 38000183, 38000232), maxOutcomeCount = 1,
    exposureSchema = "CDM4_SIM", exposureTable = "DRUG_ERA",
    outcomeSchema = "CDM4_SIM", outcomeTable = "CONDITION_OCCURRENCE",
```

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```
useCovariateDemographics = TRUE, useCovariateConditionOccurrence = TRUE,
useCovariateConditionEra = FALSE, useCovariateConditionGroup = FALSE,
useCovariateDrugExposure = FALSE, useCovariateDrugEra = FALSE,
useCovariateDrugGroup = FALSE, useCovariateProcedureOccurrence = FALSE,
useCovariateProcedureGroup = FALSE, useCovariateObservation = FALSE,
useCovariateConceptCounts = FALSE, useCovariateRiskScores = FALSE,
useCovariateInteractionYear = FALSE, useCovariateInteractionMonth = FALSE,
excludedCovariateConceptIds = c(4027133, 4032243, 4146536, 2002282, 2213572,
2005890, 43534760, 21601019), deleteCovariateSmallCount = 100)
```

# Arguments

connectionDetails

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

cdmSchema
resultsSchema
targetDrugConceptId

comparator Drug Concept Id

indication Concept Ids

washoutWindow
indicationLookbackWindow

studyStartDate
studyEndDate
exclusionConceptIds

 $\verb"outcomeConceptIds"$ 

 $outcome {\tt ConditionTypeConceptIds}$ 

maxOutcomeCount

exposureSchema

exposureTable

outcomeSchema

outcomeTable

use Covariate Demographics

use Covariate Condition Occurrence

useCovariateConditionEra

useCovariateConditionGroup

useCovariateDrugExposure

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```
useCovariateDrugEra
useCovariateDrugGroup
useCovariateProcedureOccurrence
useCovariateProcedureGroup
useCovariateObservation
useCovariateConceptCounts
useCovariateRiskScores
useCovariateInteractionYear
useCovariateInteractionMonth
excludedCovariateConceptIds
deleteCovariateSmallCount
```

# **Details**

Todo: add details

# Value

Returns an object of type cohortData, containing information on the cohorts, their outcomes, and baseline covariates.

getOutcomeModel Get the outcome model

# **Description**

getFullOutcomeModel 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, cohortData)

# Arguments

outcomeModel An object of type outcomeModel as generated using he createOutcomeMode

function.

cohortData An object of type cohortData as generated using getDbCohortData.

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#### **Details**

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

#### **Examples**

#todo

getPsModel

Get the propensity model

# Description

getPsModel shows the propensity score model

# Usage

```
getPsModel(propensityScore, cohortData)
```

# **Arguments**

propensityScore

The propensity scores as generated using the createPs function.

cohortData

An object of type cohortData as generated using getDbCohortData.

#### **Details**

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

# **Examples**

#todo

loadCohortData

Load the cohort data from a folder

# **Description**

loadCohortData loads an object of type cohortData from a folder in the file system.

# Usage

loadCohortData(file)

# Arguments

file

The name of the folder containing the data.

# **Details**

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

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#### Value

An object of class cohortData.

#### **Examples**

#todo

matchOnPs

Match persons by propensity score

#### **Description**

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

#### Usage

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

#### **Arguments**

data 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

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.

stratification Columns

Names 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 (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 (real) 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**

```
rowId = 1:5 treatment = c(1,0,1,0,1) propensityScore = c(0,0.1,0.3,0.4,1) age_group =c(1,1,1,1,1) #everyone in the same age group, so will not influence the matching data <- data.frame(rowId = rowId, treatment = treatment, propensityScore = propensityScore, age_group = age_group <- matchOnPs(data, caliper = 0, maxRatio = 1, stratificationColumns = "age_group")
```

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, fileName = NULL)
```

#### **Arguments**

balance A data frame created by the computeCovariateBalance funcion.

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.

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.

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#### Usage

```
plotCovariateBalanceScatterPlot(balance, fileName = NULL)
```

# **Arguments**

balance A data frame created by the computeCovariateBalance funcion.

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.

plotKaplanMeier Plot the Kaplan-Meier curve

#### **Description**

plotKaplanMeier creates the Kaplain-Meier survival plot

# Usage

```
plotKaplanMeier(outcomeModel, censorMarks = FALSE, legend = FALSE,
  labelsInGraph = TRUE, fileName = NULL)
```

# **Arguments**

outcomeModel An object of type outcomeModel as generated using he fitOutcomeModel func-

tion.

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

legend Whether or not to include a legend in the plot.

labelsInGraph If true, the labels identifying the two curves will be added to the graph.

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.

# **Examples**

#todo

plotPs Plot the propensity score distribution

# Description

plotPs shows the propensity (or preference) score distribution

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

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#### **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 Type of plot. Two possible values: type = "density" or type = "histogram"

binWidth For histograms, the width of the bins

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**

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 (real) Propensity score

#### 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**

```
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>
```

saveCohortData

Save the cohort data to folder

# **Description**

 ${\tt saveCohortData}\ saves\ an\ object\ of\ type\ cohortData\ to\ folder.$ 

# Usage

```
saveCohortData(cohortData, file)
```

#### **Arguments**

cohortData An object of type cohortData as generated using getDbCohortData.

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

exist.

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#### **Details**

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

#### **Examples**

#todo

simulateCohortData

Generate simulated data

# **Description**

 $\verb|simulateCohortData| creates a cohortData| object with simulated data.$ 

# Usage

simulateCohortData(cohortDataSimulationProfile, n = 10000)

# **Arguments**

cohortDataSimulationProfile

An object of type cohortDataSimulationProfile as generated using the createCohortDataSimul 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 cohortData.

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.

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

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#### **Arguments**

data 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.

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 (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 (real) 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>
```

trimByPs

Trim persons by propensity score

# Description

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

# Usage

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

#### **Arguments**

data 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:

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| rowld     | (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 (real) 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 persons in clinical equipoise

# **Description**

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

# Usage

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

# **Arguments**

data 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 (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 (real) 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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