Package 'Deconfounder'

November 16, 2020

Title Deconfounder **Version** 0.0.0.9000

Description MvDeconfounder is a causal inference model for estimating the treatment effects of medications with eletronic health records. We aim to assess the treatment effects of multiple medications on measurements using the MvDeconfounder. MvDeconfounder identifies the causal medications that have either direct effect or adverse effect on each clinical measurement. The inputs to MvDeconfounder are medication records and pre-treatment and post-treatment measurement values. The MvDeconfounder fits a probabilistic factor model (e.g., poisson factoriztion or deep exponential family) to the medication records to construct substitute confounders and adjusts for substitute confounders in the outcome model for assessing the causal effects of medications..

License Apache License 2.0 **RoxygenNote** 7.0.0

R topics documented:

creat	eCohorts	Gei	ner	at	e c	col	hoi	rt													
ndex																					7
	fitDeconfounder generateData preprocessingData .																 				4
	createCohorts												 				 				1

Description

Generate cohort

Usage

```
createCohorts(
  connection,
  cdmDatabaseSchema,
  oracleTempSchema = NULL,
  vocabularyDatabaseSchema = cdmDatabaseSchema,
  cohortDatabaseSchema,
```

2 createCohorts

```
targetCohortTable,
  createTargetCohortTable = TRUE,
  conditionConceptIds,
  measurementConceptId,
  observationWindowBefore,
  observationWindowAfter,
  drugWindow,
  targetCohortId
)
```

Arguments

connection

Connection to the database. An object of class connect as created by the connect function in the DatabaseConnector package.

cdmDatabaseSchema

A schema where OMOP CDM data are stored.

oracleTempSchema

A schema that can be used to create temp tables in when using Oracle.

vocabularyDatabaseSchema

A schema where vocabulary is stored

cohortDatabaseSchema

A schema where the cohort is stored

targetCohortTable

A string corresponds to the name of cohort table

 ${\tt createTargetCohortTable}$

A boolean that indicates whether the targetCohortTable will be created. Default TRUE.

 ${\tt conditionConceptIds}$

A list of condition concept IDs that correspond to the disease of interest.

 ${\tt measurementConceptId}$

A numeric of measurement concept ID

 $observation {\tt WindowBefore}$

An integer indicates the number of pre-treatment days used to look for pre-treatment measurement.

observationWindowAfter

An integer indicates the number of post-treatment days used to look for post-treatment measurement.

drugWindow

An integer indicates the number of post-treatment days during which drug exposure are also considered. Default is 0

targetCohortId An integer of the cohort ID.

Details

This function generates cohort for the medical deconfounder (single outcome)

Value

Returns a string containing the rendered SQL.

fitDeconfounder 3

 $\verb|fitDeconfounder||\\$

fit the deconfounder to estimate average treatment effect

Description

fit the deconfounder to estimate average treatment effect

Usage

```
fitDeconfounder(
  data_dir,
  save_dir,
  factor_model,
  learning_rate = 1e-04,
  max\_steps = 1e+05,
  latent_dim = 1,
  layer_dim = c(30, 10),
  batch_size = 1024,
  num\_samples = 1,
  holdout_portion = 0.5,
  print_steps = 50,
  tolerance = 3,
  num_confounder_samples = 30,
  CV = 5,
  outcome_type = "linear"
```

Arguments

tolerance

data_dir	String: the directory where cohort data are stored							
save_dir	String: the directory where results will be stored							
factor_model	String: the type of probabilistic factor model to fit. Choices are: PMF or DEF.							
learning_rate	Float: The learning rate for the probabilistic factor model.							
max_steps	Integer: the maximum steps to run the probabilistic factor model.							
latent_dim	Integer: the number of latent dimensions in PMF.							
layer_dim	List: a list of length 2. The number of latent dimensions in each layer of the 2-layer DEF.							
batch_size	Integer: the number of datapoints to use in each training step of the probabilistic model.							
num_samples	Integer: number of samples from variational distribution used in updating variational parameters.							
holdout_portion								
	Float: A value between 0 and 1. The proportion of data heldout for predictive model checking in checking the probabilistic model.							
print_steps	Integer: Print the results during training.							

Integer: The termination criteria for training the probabilistic model.

4 generateData

```
num_confounder_samples
```

Integer: number of samples of substitute confounder from the posterior, input

for the outcome model for estimating ATE.

CV Integer: Fold of cross validation in the outcome model.

outcome_type String: The type of outcome. Choices are: linear

generateData

Generate data

Description

Generate data

Usage

```
generateData(
  connection,
  cdmDatabaseSchema,
  oracleTempSchema = NULL,
  vocabularyDatabaseSchema = cdmDatabaseSchema,
  cohortDatabaseSchema,
  targetCohortTable,
  drugExposureTable,
  measurementTable,
  conditionConceptIds,
  measurementConceptId,
  observationWindowBefore,
  observationWindowAfter,
  drugWindow,
  createTargetCohortTable = T,
  createTargetCohort = T,
  extractFeature = T,
  targetCohortId = NULL,
  dataFolder,
  drugFilename,
  measFilename
)
```

Arguments

connection

Connection to the database. An object of class connect as created by the connect function in the DatabaseConnector package.

 ${\tt cdmDatabaseSchema}$

A schema where OMOP CDM data are stored.

 ${\tt oracleTempSchema}$

A schema that can be used to create temp tables in when using Oracle.

vocabularyDatabaseSchema

A schema where vocabulary is stored

cohortDatabaseSchema

A schema where the cohort is stored

preprocessingData 5

targetCohortTable

A string corresponds to the name of the cohort table

drugExposureTable

A string corresponds to the name of the drug exposure table

measurementTable

A string corresponds to the name of the measurement table

condition Concept Ids

A list of condition concept IDs that correspond to the disease of interest.

measurementConceptId

A numeric of measurement concept ID

observationWindowBefore

An integer indicates the number of pre-treatment days used to look for pre-treatment measurement.

observationWindowAfter

An integer indicates the number of post-treatment days used to look for post-

treatment measurement.

drugWindow An integer indicates the number of post-treatment days during which drug ex-

posure are also considered. Default is 0

 ${\tt createTargetCohortTable}$

A boolean that indicates whether the targetCohortTable will be created. Default TRUE.

createTargetCohort

A boolean that indicates whether to create the cohort. Default TRUE.

extractFeature A boolean that indicates whether to extract features for the cohort. Default

TRUE.

targetCohortId An integer of the cohort ID.

dataFolder A string indicates where output will be stored.

drugFilename A string for the name of the table to store drug exposure.

Matring for the name of the table to store measurement.

Details

This function generates cohort and extracts data of the cohort.

Value

Returns a string containing the rendered SQL.

preprocessingData Preprocess data

Description

Preprocess data

Usage

preprocessingData(dataFolder, measFilename, drugFilename, drugWindow)

6 preprocessingData

Arguments

dataFolder A string indicates where output will be stored.

measFilename A string for the name of the table to store measurement.

drugFilename A string for the name of the table to store drug exposure.

drugWindow An integer indicates the number of post-treatment days during which drug ex-

posure are also considered. Default is 0

Index

```
createCohorts, 1
fitDeconfounder, 3
generateData, 4
preprocessingData, 5
```