Package 'TreatmentPatterns'

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```
Type Package
Title A package to analyse real-world treatment patterns of a study population of interest.
Version 0.2.0
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Description This package gives insight in treatment patterns, the following tasks are per-
     formed sequentially:
     1. Target/event cohorts of interest are created
     2. Characterization of study/target population
     3. Treatment pathways are constructed
     4. Output is generated (sunburst plots and more)
     5. Shiny application is available to visualize the results
     Go to extras -¿ CodeToRun.R to execute package.
Depends R (i = 3.6.1),
     data.table,
     dplyr
Imports Andromeda,
     DatabaseConnector,
     SqlRender,
     stringr,
     stringi,
     readr,
     rjson,
     reshape2,
     googleVis,
     RJSONIO
Suggests FeatureExtraction,
     OhdsiSharing,
     ROhdsiWebApi,
     knitr,
     rmarkdown
License Apache License 2.0
Encoding UTF-8
\mathbf{LazyData} true
RoxygenNote 7.1.1
VignetteBuilder knitr
```

2 cohortCharacterization

R topics documented:

cohortCharacterization	 4
constructPathways	 ٠
createCohorts	 0
executeTreatmentPatterns	
generateResults	 (
importCohorts	 7

cohortCharacterization

This function will perform baseline characterization for the the target cohorts.

Description

This function will perform baseline characterization for the target cohorts.

Usage

```
cohortCharacterization(
  connection,
  connectionDetails,
  cdmDatabaseSchema,
  cohortDatabaseSchema,
  cohortTable,
  outputFolder,
  databaseId,
  targetCohortIds,
  minCellCount
)
```

Arguments

connection Connection to database server.

connectionDetails

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

cdmDatabaseSchema

Schema name where your patient-level data resides if OMOP-CDM = TRUE. Note that for SQL Server, this should include both the database and schema name, for example 'cdm_data.dbo'.

cohortDatabaseSchema

outputFolder

Schema name where intermediate data can be stored. You will need to have write priviliges in this schema. Note that for SQL Server, this should include both the database and schema name, for example 'cdm_data.dbo'.

cohortTable The name of the table that will be created in the work database schema. This table will hold the target and event cohorts used in this study.

Name of local folder to place results; make sure to use forward slashes

(/).

databaseId Unique identifier for database (can be the same as databaseName).

constructPathways 3

targetCohortIds

IDs to refer to target cohorts.

minCellCount Minimum number of persons with a specific treatment pathway for the

pathway to be included in analysis.

constructPathways

This function will construct treatment pathways.

Description

This function will construct treatment pathways.

Usage

```
constructPathways(
  all_data,
  study_settings,
  databaseName,
  studyName,
  outputFolder
)
```

Arguments

all_data Datatable with all target and event cohorts.

study_settings Object that contains all study settings inputted by the user.

databaseName Name of the database that will appear in the results.

studyName Name for the study corresponding to the current settings.

outputFolder Name of local folder to place results; make sure to use forward slashes

(/).

createCohorts

This function will create the target and event cohorts following the definitions included in this package if OMOP-CDM = TRUE.

Description

This function will create the target and event cohorts following the definitions included in this package if OMOP-CDM = TRUE.

4 createCohorts

Usage

```
createCohorts(
  connection,
  connectionDetails,
  cdmDatabaseSchema,
  cohortDatabaseSchema,
  cohortTable,
  outputFolder,
  loadCohorts = FALSE,
  baseUrl,
  generateCohorts = TRUE,
  minCellCount,
  flowChart = TRUE
)
```

Arguments

connection Connection to database server.

connectionDetails

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

cdmDatabaseSchema

Schema name where your patient-level data resides if OMOP-CDM = TRUE. Note that for SQL Server, this should include both the database and schema name, for example 'cdm_data.dbo'.

cohortDatabaseSchema

Schema name where intermediate data can be stored. You will need to have write priviliges in this schema. Note that for SQL Server, this should include both the database and schema name, for example 'cdm_data.dbo'.

cohortTable The name of the table that will be created in the work database schema.

This table will hold the target and event cohorts used in this study.

outputFolder Name of local folder to place results; make sure to use forward slashes

(/).

loadCohorts Setting to load cohorts from ATLAS.

baseUrl The base URL for the WebApi instance, for example: "http://server.org:80/WebAPI"

Note, there is no trailing $^{\prime}/^{\prime}.$ If trailing $^{\prime}/^{\prime}$ is used, you may receive an

error.

 ${\tt generateCohorts}$

Setting to extract specified target/event cohorts from database.

minCellCount Minimum number of persons with a specific treatment pathway for the

pathway to be included in analysis.

flowChart Setting to return numbers for flowchart with inclusion/exclusion criteria.

executeTreatmentPatterns

This is the main function which runs all different parts of the treatment pathways analysis.

Description

This is the main function which runs all different parts of the treatment pathways analysis.

Usage

```
executeTreatmentPatterns(
 OMOP\_CDM = TRUE,
  connection = NULL.
  connectionDetails = NULL,
  cdmDatabaseSchema,
  cohortDatabaseSchema = cdmDatabaseSchema,
  cohortTable = "cohort",
  outputFolder,
  databaseId = "Unknown";
  databaseName = "Unknown",
  baseUrl = "...",
  cohortLocation = "inst/Settings/input_cohorts.csv",
  runCreateCohorts = TRUE,
  runCohortCharacterization = FALSE,
  runConstructPathways = TRUE,
  runGenerateResults = TRUE,
  study_settings = study_settings
)
```

Arguments

 $\begin{array}{c} {\rm connection} & {\rm Connection} \ {\rm to} \ {\rm database} \ {\rm server}. \\ {\rm connectionDetails} \\ \end{array}$

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

cdmDatabaseSchema

Schema name where your patient-level data resides if OMOP-CDM = TRUE. Note that for SQL Server, this should include both the database and schema name, for example 'cdm_data.dbo'.

cohortDatabaseSchema

Schema name where intermediate data can be stored. You will need to have write priviliges in this schema. Note that for SQL Server, this should include both the database and schema name, for example 'cdm_data.dbo'.

cohortTable The name of the table that will be created in the work database schema.

This table will hold the target and event cohorts used in this study.

outputFolder Name of local folder to place results; make sure to use forward slashes

(/).

databaseId Unique identifier for database (can be the same as databaseName).

databaseName Name of the database that will appear in the results.

6 generateResults

baseUrl The base URL for the WebApi instance, for example: "http://server.org:80/WebAPI"

Note, there is no trailing '/'. If trailing '/' is used, you may receive an

error.

cohortLocation Location where cohorts are saved if OMOP-CDM = FALSE.

runCreateCohorts

Setting to run part of the analysis where cohorts are created.

runCohortCharacterization

Setting to run part of the analysis where characterization of target cohorts

is done.

runConstructPathways

Setting to run part of the analysis where treatment pathways are con-

structed.

runGenerateResults

Setting to run part of the analysis where final result files and plots are

generated.

study_settings Object that contains all study settings inputted by the user.

OMOP-CDM Format of 'Observational Medical Outcomes Partnership Common Data

Model' = TRUE or 'Other' = FALSE

generateResults

This function will generate all result files and plots.

Description

This function will generate all result files and plots.

Usage

generateResults(study_settings, databaseName, studyName, outputFolder)

Arguments

study_settings Object that contains all study settings inputted by the user.

databaseName Name of the database that will appear in the results.

 ${\tt studyName} \qquad \qquad {\tt Name} \ \ {\tt for} \ \ {\tt the} \ \ {\tt study} \ \ {\tt corresponding} \ \ {\tt to} \ \ {\tt the} \ \ {\tt current} \ \ {\tt settings}.$

outputFolder Name of local folder to place results; make sure to use forward slashes

(/).

importCohorts 7

importCohorts	This function will import the target and event cohorts following the path included in this package if OMOP-CDM = FALSE.
	the pain included in this package if $OMOP-ODM = FALSE$.

${\bf Description}$

This function will import the target and event cohorts following the path included in this package if OMOP-CDM = FALSE.

Usage

importCohorts(cohortLocation, outputFolder)

Arguments

```
\label{eq:cohortLocation} \begin{array}{ll} \text{Location where cohorts are saved if OMOP-CDM} = \text{FALSE}. \\ \text{outputFolder} & \text{Name of local folder to place results; make sure to use forward slashes} \\ & (/). \end{array}
```