# Package 'TreatmentPatterns'

January 26, 2024

```
Type Package
Title Analyzes Real-World Treatment Patterns of a Study Population of Interest
Version 2.6.4
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Description Computes treatment patterns within a given cohort using the Observational
     Medical Outcomes Partnership (OMOP) common data model (CDM). As described
     in Markus, Verhamme, Kors, and Rijnbeek (2022) <doi:10.1016/j.cmpb.2022.107081>.
URL https://github.com/darwin-eu-dev/TreatmentPatterns
BugReports https://github.com/darwin-eu-dev/TreatmentPatterns/issues
Language en-US
Depends R (>= 4.2)
Imports checkmate,
     dplyr,
     stringr,
     stringi,
     utils,
     rjson,
     googleVis,
     stats,
     Andromeda,
     tidyr,
     R6,
     sunburstR,
     networkD3,
     htmlwidgets,
     shiny,
     shinydashboard
Suggests knitr,
     rmarkdown,
     tibble.
     testthat (>= 3.0.0),
     usethis,
     Eunomia,
     CDMConnector,
     DatabaseConnector (>= 6.0.0),
     SqlRender,
```

2 CharacterizationPlots

```
CohortGenerator,
     webshot2,
     CirceR,
     duckdb,
     DBI,
     withr,
     plotly,
     ggplot2,
     Capr
License Apache License 2.0
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LazyData true
RoxygenNote 7.2.3
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Additional_repositories https://ohdsi.github.io/drat
Roxygen list(markdown = TRUE)
```

# **R** topics documented:

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 $Characterization Plots \quad \textit{Characterization Plots}$ 

# Description

Class to handle the characterization plots.

## Super class

TreatmentPatterns::Module -> CharacterizationPlots

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#### Methods

```
Public methods:
```

• CharacterizationPlots\$uiMenu()

```
• CharacterizationPlots$uiBody()
  • CharacterizationPlots$server()
  • CharacterizationPlots$clone()
Method uiMenu(): Method to include a menuItem to link to the body.
 Usage:
 CharacterizationPlots$uiMenu(
   label = "Characteristics",
   tag = "characteristics"
 )
 Arguments:
 label (character(1))
     Label to show for the menuItem.
 tag (character(1))
     Tag to use internally in input.
 Returns: (menuItem)
Method uiBody(): Method to include a tabItem to include the body.
 Usage:
 CharacterizationPlots$uiBody()
 Returns: (tabItem)
Method server(): Method to handle the back-end.
 CharacterizationPlots$server(input, output, session, inputHandler)
 Arguments:
 input (input)
     Input from the server function.
 output (output)
     Output from the server function.
 session (session)
     Session from the server function.
 inputHandler (inputHandler)
     InputHandler class.
 Returns: (NULL)
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 CharacterizationPlots$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

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computePathways

computePathways

#### **Description**

Compute treatment patterns according to the specified parameters within specified cohorts.

#### Usage

```
computePathways(
  cohorts,
  cohortTableName,
  cdm = NULL,
  connectionDetails = NULL,
  cdmSchema = NULL,
  resultSchema = NULL,
  tempEmulationSchema = NULL,
  includeTreatments = "startDate",
  periodPriorToIndex = 0,
  minEraDuration = 0,
  splitEventCohorts = NULL,
  splitTime = NULL,
  eraCollapseSize = 30,
  combinationWindow = 30,
  minPostCombinationDuration = 30,
  filterTreatments = "First",
  maxPathLength = 5
```

## **Arguments**

cohorts (data.frame())

Data frame containing the following columns and data types:

**cohortId** numeric(1) Cohort ID's of the cohorts to be used in the cohort table. **cohortName** character(1) Cohort names of the cohorts to be used in the cohort table.

**type** character(1) ["target", "event', "exit" ] Cohort type, describing if the cohort is a target, event, or exit cohort

cohortTableName

(character(1))Cohort table name.

cdm (CDMConnector::cdm\_from\_con(): NULL)

Optional; Ignores connectionDetails, cdmSchema, and resultSchema.

connectionDetails

(DatabaseConnector::createConnectionDetails(): NULL)

Optional; In congruence with  ${\tt cdmSchema}$  and  ${\tt resultSchema}$ . Ignores  ${\tt cdm}$ .

cdmSchema (character(1): NULL)

Optional; In congruence with connectionDetails and resultSchema. Ignores

cdm.

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resultSchema (character(1): NULL) Optional; In congruence with connectionDetails and cdmSchema. Ignores tempEmulationSchema Schema used to emulate temp tables includeTreatments(character(1): "startDate") "startDate" Include treatments after the target cohort start date and onwards. "endDate" Include treatments before target cohort end date and before. periodPriorToIndex (integer(1): 0) Number of days prior to the index date of the target cohort | that event cohorts are allowed to start minEraDuration (integer(1): 0) Minimum time an event era should last to be included in analysis splitEventCohorts (character(n): "") Specify event cohort to split in acute (< X days) and therapy (>= X days) splitTime (integer(1): 30) Specify number of days (X) at which each of the split event cohorts should be split in acute and therapy eraCollapseSize (integer(1): 30) Window of time between which two eras of the same event cohort are collapsed into one era combinationWindow (integer(1): 30) Window of time two event cohorts need to overlap to be considered a combination treatment minPostCombinationDuration (integer(1): 30) Minimum time an event era before or after a generated combination treatment should last to be included in analysis filterTreatments (character(1): "First" ["first", "Changes", "all"]) Select first occurrence of ('First'); changes between ('Changes'); or all event cohorts ('All'). (integer(1): 5) maxPathLength Maximum number of steps included in treatment pathway

#### Value

(Andromeda::andromeda()) andromeda object containing non-sharable patient level data outcomes.

```
library(TreatmentPatterns)
library(CDMConnector)
library(dplyr)
```

createSankeyDiagram

```
if (require("CirceR", character.only = TRUE, quietly = TRUE)) {
  withr::local_envvar(
    EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
  downloadEunomiaData(overwrite = TRUE)
  con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())</pre>
  cdm <- cdmFromCon(con, cdmSchema = "main", writeSchema = "main")</pre>
  cohortSet <- readCohortSet(</pre>
    path = system.file(package = "TreatmentPatterns", "exampleCohorts")
  )
  cdm <- generateCohortSet(</pre>
   cdm = cdm,
    cohortSet = cohortSet,
    name = "cohort_table"
  cohorts <- cohortSet %>%
    \mbox{\tt\#} Remove 'cohort' and 'json' columns
    select(-"cohort", -"json") %>%
  mutate(type = c("event", "event", "event", "event", "event", "event", "event", "event", "target")) %>%
    rename(
      cohortId = "cohort_definition_id",
      cohortName = "cohort_name",
    ) %>%
    select("cohortId", "cohortName", "type")
  outputEnv <- computePathways(</pre>
    cohorts = cohorts,
    cohortTableName = "cohort_table",
    cdm = cdm
  )
  Andromeda::close(outputEnv)
 DBI::dbDisconnect(con, shutdown = TRUE)
}
```

createSankeyDiagram

#### **Description**

Create sankey diagram.

#### Usage

```
createSankeyDiagram(
  treatmentPathways,
  groupCombinations = FALSE,
  colors = NULL,
```

createSankeyDiagram2

```
)
```

# **Arguments**

```
treatmentPathways

(data.frame())
The contents of the treatmentPathways.csv-file as a data.frame().

groupCombinations
(logical(1): FALSE)

TRUE Group all combination treatments in category "Combination".
FALSE Do not group combination treatments.

colors
(character(n)) Vector of hex color codes.
...
Paramaters for sankeyNetwork.
```

## Value

```
(htmlwidget)
```

## **Examples**

createSankeyDiagram2 createSankeyDiagram2

# **Description**

DEPRECATED Create sankey diagram.

# Usage

```
createSankeyDiagram2(
  treatmentPathways,
  groupCombinations = FALSE,
  colors = NULL,
  ...
)
```

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```
Arguments
```

```
treatmentPathways

(data.frame())
The contents of the treatmentPathways.csv-file as a data.frame().

groupCombinations
(logical(1): FALSE)

TRUE Group all combination treatments in category "Combination".
FALSE Do not group combination treatments.

colors
(character(n)) Vector of hex color codes.
...
Paramaters for sankeyNetwork.
```

#### Value

(htmlwidget)

## **Examples**

createSunburstPlot

createSunburstPlot

# Description

New sunburstPlot function

# Usage

```
createSunburstPlot(treatmentPathways, groupCombinations = FALSE, ...)
```

#### **Arguments**

```
treatmentPathways

(data.frame())

The contents of the treatmentPathways.csv-file as a data.frame().

groupCombinations

(logical(1): FALSE)

TRUE Group all combination treatments in category "Combination".

FALSE Do not group combination treatments.

... Paramaters for sunburst.
```

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#### Value

```
(htmlwidget)
```

#### **Examples**

createSunburstPlot2

createSunburstPlot2

## **Description**

DEPRECATED New sunburstPlot function

# Usage

```
createSunburstPlot2(treatmentPathways, groupCombinations = FALSE, ...)
```

# **Arguments**

```
treatmentPathways

(data.frame())

The contents of the treatmentPathways.csv-file as a data.frame().

groupCombinations

(logical(1): FALSE)

TRUE Group all combination treatments in category "Combination".

FALSE Do not group combination treatments.

Paramaters for sunburst.
```

# Value

```
(htmlwidget)
```

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```
age = rep("all", 5),
index_year = rep("all", 5)
)
createSunburstPlot2(treatmentPatwhays)
```

executeTreatmentPatterns

executeTreatmentPatterns

# Description

Compute treatment patterns according to the specified parameters within specified cohorts. For more customization, or investigation of patient level outcomes, you can run computePathways and export separately.

## Usage

```
executeTreatmentPatterns(
  cohorts,
  cohortTableName,
  outputPath,
  cdm = NULL,
  connectionDetails = NULL,
  cdmSchema = NULL,
  resultSchema = NULL,
  tempEmulationSchema = NULL,
  minEraDuration = 0,
  eraCollapseSize = 30,
  combinationWindow = 30,
  minCellCount = 5
)
```

## **Arguments**

```
(data.frame())
cohorts
                  Data frame containing the following columns and data types:
                  cohortId numeric(1) Cohort ID's of the cohorts to be used in the cohort table.
                  cohortName character(1) Cohort names of the cohorts to be used in the co-
                      hort table.
                  type character(1) ["target", "event', "exit" ] Cohort type, describing if the
                      cohort is a target, event, or exit cohort
cohortTableName
                  (character(1))
                  Cohort table name.
outputPath
                  (character(1))
                  (CDMConnector::cdm_from_con(): NULL)
cdm
                  Optional; Ignores connectionDetails, cdmSchema, and resultSchema.
```

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```
connectionDetails
                 (DatabaseConnector::createConnectionDetails(): NULL)
                 Optional; In congruence with cdmSchema and resultSchema. Ignores cdm.
cdmSchema
                 (character(1): NULL)
                 Optional; In congruence with connectionDetails and resultSchema. Ignores
                 cdm.
resultSchema
                 (character(1): NULL)
                 Optional; In congruence with connectionDetails and cdmSchema. Ignores
                 cdm.
tempEmulationSchema
                 (character(1)) Schema to emulate temp tables.
minEraDuration (integer(1): 0)
                 Minimum time an event era should last to be included in analysis
eraCollapseSize
                 (integer(1): 30)
                 Window of time between which two eras of the same event cohort are collapsed
                 into one era
combinationWindow
                 (integer(1): 30)
                 Window of time two event cohorts need to overlap to be considered a combina-
                 tion treatment
minCellCount
                 (integer(1): 5)
                 Minimum count required per pathway. Censors data below x as <x. This mini-
                 mum value will carry over to the sankey diagram and sunburst plot.
```

#### Value

```
(invisible(NULL))
```

```
library(TreatmentPatterns)
library(CDMConnector)
library(dplyr)
if (require("CirceR", character.only = TRUE, quietly = TRUE)) {
    withr::local_envvar(
        EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
    )

    downloadEunomiaData(overwrite = TRUE)

    con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())
    cdm <- cdmFromCon(con, cdmSchema = "main", writeSchema = "main")

    cohortSet <- readCohortSet(
        path = system.file(package = "TreatmentPatterns", "exampleCohorts")
    )

    cdm <- generateCohortSet(
    cdm = cdm,
    cohortSet = cohortSet,
    name = "cohort_table"
    )</pre>
```

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```
cohorts <- cohortSet %>%
    # Remove 'cohort' and 'json' columns
    select(-"cohort", -"json") %>%
mutate(type = c("event", "event", "event", "event", "exit", "event", "event", "target")) %>%
    rename(
        cohortId = "cohort_definition_id",
        cohortName = "cohort_name",
        ) %>%
        select("cohortId", "cohortName", "type")

executeTreatmentPatterns(
        cohorts = cohorts,
        cohortTableName = "cohort_table",
        cdm = cdm,
        outputPath = tempdir()
)

DBI::dbDisconnect(con, shutdown = TRUE)
}
```

export

export

## **Description**

Export andromeda generated by computePathways object to sharable csv-files and/or a zip archive.

## Usage

```
export(
  andromeda,
  outputPath,
  ageWindow = 10,
  minCellCount = 5,
  censorType = "minCellCount",
  archiveName = NULL
)
```

#### **Arguments**

```
andromeda (Andromeda::andromeda()) Andromeda object.

outputPath (character(1))

ageWindow (integer(n): 10)

Number of years to bin age groups into. It may also be a vector of integers. I.e. c(0, 18, 150) which will results in age group 0-18 which includes subjects < 19. And age group 18-150 which includes subjects > 18.

minCellCount (integer(1): 5)

Minimum count required per pathway. Censors data below x as <x. This mini-
```

Minimum count required per pathway. Censors data below x as <x. This minimum value will carry over to the sankey diagram and sunburst plot.

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```
censorType (character(1))

"minCellCount" Censors pathways <minCellCount to minCellCount.

"remove" Censors pathways <minCellCount by removing them completely.

"mean" Censors pathways <minCellCount to the mean of all frequencies below minCellCount

archiveName (character(1): NULL)

If not NULL adds the exported files to a ZIP-file with the specified archive name.</pre>
```

#### Value

```
(invisible(NULL))
```

```
library(TreatmentPatterns)
library(CDMConnector)
library(dplyr)
if (require("CirceR", character.only = TRUE, quietly = TRUE)) {
  withr::local_envvar(
    EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
  downloadEunomiaData(overwrite = TRUE)
  con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomia_dir())</pre>
  cdm <- cdmFromCon(con, cdmSchema = "main", writeSchema = "main")</pre>
  cohortSet <- readCohortSet(</pre>
    path = system.file(package = "TreatmentPatterns", "exampleCohorts")
  cdm <- generateCohortSet(</pre>
    cdm = cdm,
    cohortSet = cohortSet,
    name = "cohort_table"
  )
  cohorts <- cohortSet %>%
    \mbox{\tt\#} Remove 'cohort' and 'json' columns
    select(-"cohort", -"json") %>%
  mutate(type = c("event", "event", "event", "event", "event", "event", "event", "target")) %>%
    rename(
      cohortId = "cohort_definition_id",
      cohortName = "cohort_name",
    select("cohortId", "cohortName", "type")
  outputEnv <- computePathways(</pre>
    cohorts = cohorts,
    cohortTableName = "cohort_table",
    cdm = cdm
```

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```
export(
   andromeda = outputEnv,
   outputPath = tempdir()
)

Andromeda::close(outputEnv)
   DBI::dbDisconnect(con, shutdown = TRUE)
}
```

InputHandler

InputHandler

# Description

Class to handle input from the user. Supports direct paths or input fields through setDataPath().

# Super class

```
TreatmentPatterns::Module -> InputHandler
```

## **Active bindings**

```
reactiveValues (reactiveValues) reactiveValues class created by reactiveValues.
```

#### Methods

#### **Public methods:**

- InputHandler\$uiMenu()
- InputHandler\$uiBody()
- InputHandler\$server()
- InputHandler\$uiDatabaseSelector()
- InputHandler\$setDataPath()
- InputHandler\$clone()

**Method** uiMenu(): Method to include a menuItem to link to the body.

```
Usage:
InputHandler$uiMenu(label = "File upload", tag = "fileUpload")
Arguments:
label (character(1))
    Label to show for the menuItem.
tag (character(1))
    Tag to use internally in input.
Returns: (menuItem)
```

**Method** uiBody(): Method to include a tabItem to include the body.

```
Usage:
 InputHandler$uiBody()
 Returns: (tabItem)
Method server(): Method to handle the back-end.
 Usage:
 InputHandler$server(input, output, session)
 Arguments:
 input (input)
     Input from the server function.
 output (output)
     Output from the server function.
 session (session)
     Session from the server function.
 Returns: (NULL)
Method uiDatabaseSelector(): Method to include a uiOutput to select between multiple
uploaded files.
 Usage:
 InputHandler$uiDatabaseSelector()
 Returns: (uiOutput)
Method setDataPath(): Method to dictate where the data is coming from, either from the
input through the shiny application, or from a specified path. When one is provided, the other is
ignored.
 Usage:
 InputHandler$setDataPath(tag = "uploadField", input = NULL, path = NULL)
 Arguments:
 tag (character(1))
     Tag to use internally in input.
 input (input)
     Input from the server function of the shiny app.
 path (character(1))
     Path to a zip-file containing TreatmentPatterns output files.
 Returns: (invisible(self))
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 InputHandler$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

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InteractivePlots

InteractivePlots

# Description

Class to handle the interactive plots of TreatmentPatterns (Sunburst plot & Sankey diagram)

## Super class

```
TreatmentPatterns::Module -> InteractivePlots
```

#### Methods

```
Public methods:
```

```
• InteractivePlots$uiMenu()
```

- InteractivePlots\$uiBody()
- InteractivePlots\$server()
- InteractivePlots\$clone()

```
Method uiMenu(): Method to include a menuItem to link to the body.
```

```
Usage:
InteractivePlots$uiMenu(label = "Plots", tag = "plots")
Arguments:
label (character(1))
    Label to show for the menuItem.
tag (character(1))
    Tag to use internally in input.
Returns: (menuItem)
```

**Method** uiBody(): Method to include a tabItem to include the body.

```
Interact
```

InteractivePlots\$uiBody()

Returns: (tabItem)

**Method** server(): Method to handle the back-end.

```
Usage:
```

```
InteractivePlots$server(input, output, session, inputHandler)
```

## Arguments:

```
input (input)
```

Input from the server function.

output (output)

Output from the server function.

session (session)

Session from the server function.

inputHandler (inputHandler)

InputHandler class.

Returns: (NULL)

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```
Method clone(): The objects of this class are cloneable with this method.
```

```
Usage:
InteractivePlots%clone
```

InteractivePlots\$clone(deep = FALSE)

Arguments:

deep Whether to make a deep clone.

 $launch Results Explorer \ \ launch Results Explorer$ 

## **Description**

Launches the ResultExplorer shinyApp.

## Usage

```
launchResultsExplorer()
```

#### Value

(shinyApp)

# **Examples**

```
if (interactive()) {
  launchResultsExplorer()
}
```

Module

Module

## **Description**

Module super class

# **Active bindings**

namespace Namespace of the module.

# Methods

#### **Public methods:**

- Module\$new()
- Module\$validate()
- Module\$uiMenu()
- Module\$uiBody()
- Module\$server()
- Module\$clone()

Method new(): Initializer method

Module Module

```
Usage:
 Module$new(namespace)
 Arguments:
 namespace (character(1))
 Returns: (invisible(self))
Method validate(): Validator method
 Usage:
 Module$validate()
 Returns: (invisible(self))
Method uiMenu(): Method to include a menuItem to link to the body.
 Usage:
 Module$uiMenu(label, tag)
 Arguments:
 label (character(1))
     Label to show for the menuItem.
 tag (character(1))
     Tag to use internally in input.
 Returns: (menuItem)
Method uiBody(): Method to include a tabItem to include the body.
 Usage:
 Module$uiBody()
 Returns: (tabItem)
Method server(): Method to handle the back-end.
 Usage:
 Module$server(input, output, session)
 Arguments:
 input (input)
     Input from the server function.
 output (output)
     Output from the server function.
 session (session)
     Session from the server function.
 Returns: (NULL)
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 Module$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

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