Package 'TreatmentPatterns'

April 4, 2025

```
Type Package
Title Analyzes Real-World Treatment Patterns of a Study Population of Interest
Version 3.0.2
Maintainer Maarten van Kessel <m.l.vankessel@erasmusmc.nl>
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/TreatmentPatterns, https:
     //darwin-eu-dev.github.io/TreatmentPatterns/
BugReports https://github.com/darwin-eu/TreatmentPatterns/issues
Language en-US
Depends R (>= 4.2)
Imports checkmate,
     dplyr,
     stringr,
     utils,
     stats,
     Andromeda,
     tidyr,
     R6,
     sunburstR,
     networkD3,
     htmlwidgets,
     ggplot2,
     dbplyr,
     jsonlite
Suggests knitr,
     rmarkdown,
     tibble,
     testthat (>= 3.0.0),
     usethis,
     Eunomia,
     CDMConnector,
     DatabaseConnector (>= 6.0.0),
     SqlRender,
     CohortGenerator,
```

2 Contents

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Collate 'CDMInterface.R' 'TreatmentPatterns-package.R' 'TreatmentPatternsResults.R' 'attrition.R' 'computePathways.R' 'constructPathways.R' 'createSankeyDiagram.R' 'createSunburstPlot.R' 'executeTreatmentPatterns.R' 'export.R' 'getResultsDataModelSpecification.R' 'ggSunburst.R' 'plotEventDuration.R'
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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,
  analysisId = 1,
  description = ""
  tempEmulationSchema = NULL,
  includeTreatments = "startDate",
  indexDateOffset = 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 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.
cdm
                  (CDMConnector::cdm_from_con(): NULL)
                  Optional; Ignores \ connection Details, \ cdm Schema, \ and \ result Schema.
connectionDetails
                  (DatabaseConnector::createConnectionDetails(): NULL)
                  Optional; In congruence with cdmSchema and resultSchema. Ignores cdm.
```

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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.

analysisId (character(1)) Identifier for the TreatmentPatterns analysis.

description (character(1)) Description of the analysis.

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.

indexDateOffset

(integer(1): 0)

Offset the index date of the Target cohort.

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 combina-

tion treatment

 $\verb|minPostCombinationDuration||$

(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').

maxPathLength (integer(1): 5)

Maximum number of steps included in treatment pathway

Value

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

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```
ableToRun <- all(</pre>
  require("CirceR", character.only = TRUE, quietly = TRUE),
  require("CDMConnector", character.only = TRUE, quietly = TRUE),
  require("TreatmentPatterns", character.only = TRUE, quietly = TRUE),
  require("dplyr", character.only = TRUE, quietly = TRUE)
if (ableToRun) {
  library(TreatmentPatterns)
  library(CDMConnector)
  library(dplyr)
  withr::local_envvar(
    R_USER_CACHE_DIR = tempfile(),
    EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
  )
  tryCatch({
    if (Sys.getenv("skip_eunomia_download_test") != "TRUE") {
      CDMConnector::downloadEunomiaData(overwrite = TRUE)
  }, error = function(e) NA)
  con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())</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
  Andromeda::close(outputEnv)
  DBI::dbDisconnect(con, shutdown = TRUE)
```

createSankeyDiagram

```
createSankeyDiagram
createSankeyDiagram
```

Description

Create sankey diagram.

Usage

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

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)
```

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createSunburstPlot

create Sunburst Plot

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.
```

Value

(htmlwidget)

Examples

 ${\tt executeTreatmentPatterns}$

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.

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Usage

```
executeTreatmentPatterns(
      cohorts,
      cohortTableName,
      cdm = NULL,
      connectionDetails = NULL,
      cdmSchema = NULL,
      resultSchema = NULL,
      tempEmulationSchema = NULL,
      minEraDuration = 0,
      eraCollapseSize = 30,
      combinationWindow = 30,
      minCellCount = 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 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.
    cdm
                     (CDMConnector::cdm_from_con(): NULL)
                     Optional; Ignores connectionDetails, cdmSchema, and resultSchema.
    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
    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
```

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```
minCellCount (integer(1): 5)
```

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

Value

TreatmentPatternsResults

```
ableToRun <- all(</pre>
  require("CirceR", character.only = TRUE, quietly = TRUE),
  require("CDMConnector", character.only = TRUE, quietly = TRUE),
  require("TreatmentPatterns", character.only = TRUE, quietly = TRUE),
  require("dplyr", character.only = TRUE, quietly = TRUE)
)
if (require("CirceR", character.only = TRUE, quietly = TRUE)) {
  library(TreatmentPatterns)
  library(CDMConnector)
  library(dplyr)
  withr::local_envvar(
    R_USER_CACHE_DIR = tempfile(),
    EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
  )
  tryCatch({
    if (Sys.getenv("skip_eunomia_download_test") != "TRUE") {
      CDMConnector::downloadEunomiaData(overwrite = TRUE)
    }
  },
  error = function(e) NA)
  con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())</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 %>%
    # 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")
```

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```
executeTreatmentPatterns(
  cohorts = cohorts,
  cohortTableName = "cohort_table",
  cdm = cdm
)

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 = NULL,
  ageWindow = 10,
  minCellCount = 5,
  censorType = "minCellCount",
  archiveName = NULL,
  nonePaths = FALSE,
  stratify = FALSE
)
```

Arguments

andromeda

(character: NULL) Output path where to write output files to. When set to NULL outputPath no files will be written, and only the results object is returned. 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 minimum value will carry over to the sankey diagram and sunburst plot. 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

(Andromeda::andromeda()) Andromeda object.

archiveName (character(1): NULL)

If not NULL adds the exported files to a ZIP-file with the specified archive name.

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nonePaths (logical(1)) Should None paths be included? This will fetch all persons included in the target cohort and assign them a "None" pathway. Significantly impacts performance.

stratify (logical(1)) Should pathways be stratified? This will perform pairwise stratification between age, sex, and index year. Significantly impacts performance.

Value

TreatmentPatternsResults object

```
ableToRun <- all(</pre>
  require("CirceR", character.only = TRUE, quietly = TRUE),
  require("CDMConnector", character.only = TRUE, quietly = TRUE),
  require("TreatmentPatterns", character.only = TRUE, quietly = TRUE),
  require("dplyr", character.only = TRUE, quietly = TRUE)
if (ableToRun) {
  library(TreatmentPatterns)
  library(CDMConnector)
  library(dplyr)
  withr::local_envvar(
    R_USER_CACHE_DIR = tempfile(),
    EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
  )
  tryCatch({
    if (Sys.getenv("skip_eunomia_download_test") != "TRUE") {
      CDMConnector::downloadEunomiaData(overwrite = TRUE)
    }
  }, error = function(e) NA)
  con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())</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 %>%
    # 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",
    ) %>%
```

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```
select("cohortId", "cohortName", "type")

outputEnv <- computePathways(
   cohorts = cohorts,
   cohortTableName = "cohort_table",
   cdm = cdm
)

results <- export(
   andromeda = outputEnv
)

Andromeda::close(outputEnv)

DBI::dbDisconnect(con, shutdown = TRUE)
}</pre>
```

 $get Results Data Model Specifications \\ get Results Data Model Specifications$

Description

Gets the results data model specifications of TreatmentPatterns.

Usage

```
getResultsDataModelSpecifications()
```

Value

data.frame

Examples

```
{
getResultsDataModelSpecifications()
}
```

ggSunburst

ggSunburst

Description

ggSunburst

Usage

```
ggSunburst(treatmentPathways, groupCombinations = FALSE, unit = "percent")
```

plotEventDuration 13

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.

unit (character(1)) Either "count" or "percent", to scale the plot to.
```

Value

```
(gg, ggplot)
```

Examples

 ${\tt plotEventDuration}$

plotEventDuration

Description

plotEventDuration

Usage

```
plotEventDuration(
  eventDurations,
  minCellCount = 0,
  treatmentGroups = "both",
  eventLines = NULL,
  includeOverall = TRUE
)
```

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Arguments

```
eventDurations (data.frame) Contents of summaryEventDuration.csv file.

minCellCount (numeric(1): 0) Min Cell Count per event group.

treatmentGroups

(character(1): "both") "group": Only mono-, and combination-events. "individual":

Only individual (combination) events. "both": Both mono-, and combination-events, and individual (combination) events.

eventLines (numeric(n): NULL) Event lines to include, i.e. c(1, 2, 3) includes first (1), second (2), and third (3) lines of events. NULL will include all eventLines.

includeOverall (logical(1): TRUE) TRUE: Include an overall column with the eventLines.

FALSE: Exclude the overall column.
```

Value

ggplot

```
ableToRun <- all(</pre>
  require("CirceR", character.only = TRUE, quietly = TRUE),
  require("CDMConnector", character.only = TRUE, quietly = TRUE),
  require("TreatmentPatterns", character.only = TRUE, quietly = TRUE),
  require("dplyr", character.only = TRUE, quietly = TRUE)
if (ableToRun) {
  withr::local_envvar(
    R_USER_CACHE_DIR = tempfile(),
    EUNOMIA_DATA_FOLDER = Sys.getenv("EUNOMIA_DATA_FOLDER", unset = tempfile())
  )
  tryCatch({
    if (Sys.getenv("skip_eunomia_download_test") != "TRUE") {
      CDMConnector::downloadEunomiaData(overwrite = TRUE)
    }
  }, error = function(e) NA)
  con <- DBI::dbConnect(duckdb::duckdb(), dbdir = eunomiaDir())</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 %>%
    # Remove 'cohort' and 'json' columns
    select(-"cohort", -"json") %>%
  mutate(type = c("event", "event", "event", "event", "event", "event", "event", "event", "target")) %>%
```

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```
rename(
    cohortId = "cohort_definition_id",
    cohortName = "cohort_name",
  ) %>%
  select("cohortId", "cohortName", "type")
outputEnv <- computePathways(</pre>
  cohorts = cohorts,
  cohortTableName = "cohort_table",
  cdm = cdm
results <- export(outputEnv)</pre>
plotEventDuration(
  eventDurations = results$summary_event_duration,
  minCellCount = 5,
  treatmentGroups = "group",
  eventLines = 1:4,
  includeOverall = FALSE
Andromeda::close(outputEnv)
DBI::dbDisconnect(con, shutdown = TRUE)
```

TreatmentPatternsResults

TreatmentPatternsResults Class

Description

Houses the results of a TreatmentPatterns analysis. Each field corresponds to a file. Plotting methods are provided.

Active bindings

```
attrition (data.frame)
metadata (data.frame)
treatment_pathways (data.frame)
summary_event_duration (data.frame)
counts_age (data.frame)
counts_sex (data.frame)
counts_year (data.frame)
cdm_source_info (data.frame)
analyses (data.frame)
arguments (list)
```

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Methods

Public methods:

- TreatmentPatternsResults\$new()
- TreatmentPatternsResults\$saveAsZip()
- TreatmentPatternsResults\$saveAsCsv()
- TreatmentPatternsResults\$uploadResultsToDb()
- TreatmentPatternsResults\$load()
- TreatmentPatternsResults\$plotSunburst()
- TreatmentPatternsResults\$plotSankey()
- TreatmentPatternsResults\$plotEventDuration()
- TreatmentPatternsResults\$clone()

```
Method new(): Initializer method
```

```
Usage:
TreatmentPatternsResults$new(
  attrition = NULL,
  metadata = NULL,
  treatmentPathways = NULL,
  summaryEventDuration = NULL,
  countsAge = NULL,
  countsSex = NULL,
  countsYear = NULL,
  cdmSourceInfo = NULL,
  analyses = NULL,
  arguments = NULL,
  filePath = NULL
)
Arguments:
attrition (data.frame) attrition result.
metadata (data.frame)) metadata result.
treatmentPathways (data.frame)) treatmentPathways result.
summaryEventDuration (data.frame)) summaryEventDuration result.
countsAge (data.frame)) countsAge result.
countsSex (data.frame)) countsSex result.
countsYear (data.frame)) countsYear result.
cdmSourceInfo (data.frame) cdmSourceInfo result.
analyses (data.frame) Analyses result.
arguments (list) Named list of arguments used.
```

Method saveAsZip(): Save the results as a zip-file.

```
Usage:
TreatmentPatternsResults$saveAsZip(path, name, verbose = TRUE)
Arguments:
path (character(1)) Path to write to.
name (character(1)) File name.
verbose (logical: TRUE) Verbose messaging.
```

filePath (character) File path to either a directory or zip-file, containing the csv-files.

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```
Returns: self
Method saveAsCsv(): Save the results as csv-files.
 Usage:
 TreatmentPatternsResults$saveAsCsv(path, verbose = TRUE)
 Arguments:
 path (character(1)) Path to write to.
 verbose (logical: TRUE) Verbose messaging.
 Returns: self
Method uploadResultsToDb(): Upload results to a resultsDatabase using ResultModelManager.
 TreatmentPatternsResults$uploadResultsToDb(
   connectionDetails,
   schema,
   prefix = "tp_"
   overwrite = TRUE,
   purgeSiteDataBeforeUploading = FALSE
 )
 Arguments:
 connection Details\ (Connection Details)\ Connection Details\ object\ from\ Database Connector.
 schema (character(1)) Schema to write tables to.
 prefix (character(1): "tp_") Table prefix.
 overwrite (logical(1): TRUE) Should tables be overwritten?
 purgeSiteDataBeforeUploading (logical: FALSE) Should site data be purged before up-
     loading?
 Returns: self
Method load(): Load data from files.
 Usage:
 TreatmentPatternsResults$load(filePath)
 Arguments:
 filePath (character(1)) Path to a directory or zip-file containing the result csv-files.
 Returns: self
Method plotSunburst(): Wrapper for TreatmentPatterns::createSunburstPlot(), but
with data filtering step.
 Usage:
 TreatmentPatternsResults$plotSunburst(
   age = "all",
   sex = "all",
   indexYear = "all",
   nonePaths = FALSE,
 )
 Arguments:
```

age (character(1)) Age group.

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```
sex (character(1)) Sex group.
 indexYear (character(1)) Index year group.
 nonePaths (logical(1)) Should None paths be included?
 ... Parameters for TreatmentPatterns::createSunburstPlot()
 Returns: htmlwidget
Method plotSankey():
                        Wrapper for TreatmentPatterns::createSankeyDiagram(), but
with data filtering step.
 Usage:
 TreatmentPatternsResults$plotSankey(
   age = "all",
   sex = "all",
   indexYear = "all",
   nonePaths = FALSE,
 )
 Arguments:
 age (character(1)) Age group.
 sex (character(1)) Sex group.
 indexYear (character(1)) Index year group.
 nonePaths (logical(1)) Should None paths be included?
 ... Parameters for TreatmentPatterns::createSankeyDiagram()
 Returns: htmlwidget
Method plotEventDuration(): Wrapper for TreatmentPatterns::plotEventDuration().
 TreatmentPatternsResults$plotEventDuration(...)
 Arguments:
 ... Parameters for TreatmentPatterns::plotEventDuration()
 Returns: ggplot
Method clone(): The objects of this class are cloneable with this method.
 Usage:
 TreatmentPatternsResults$clone(deep = FALSE)
 Arguments:
 deep Whether to make a deep clone.
```

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