Package 'Tendril'

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Type Package

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Title Compute and Display Tendril Plots

Description Compute the coordinates to produce a tendril plot. In the tendril plot, each tendril (branch) represents a type of events, and the direction of the tendril is dictated by on which treatment arm the event is occurring. If an event is occurring on the first of the two specified treatment arms, the tendril bends in a clockwise direction. If an event is occurring on the second of the treatment arms, the tendril bends in an anti-clockwise direction. Ref: Karpefors, M and Weatherall, J., "The Tendril Plot - a novel visual summary of the incidence, significance and temporal aspects of adverse events in clinical trials" - JAMIA 2018; 25(8): 1069-1073 <doi:10.1093/jamia/ocy016>. URL https://github.com/Karpefors/Tendril BugReports https://github.com/Karpefors/Tendril/issues **Encoding UTF-8** LazyData true **Depends** R (>= 3.5) Imports ggplot2, dplyr, plyr, reshape2, magrittr, scales, graphics, grDevices, plotly RoxygenNote 6.1.1 Suggests testthat, knitr, rmarkdown, devtools VignetteBuilder knitr NeedsCompilation no **Author** Martin Karpefors [aut], Stefano Borini [ctb, cre], Mark Edmondson-Jones [ctb], Hielke Bijlsma [ctb] Maintainer Stefano Borini <stefano.borini@astrazeneca.com>

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plot.Tendril

Plot Tendril

Description

Function to plot Tendril results

Usage

```
## S3 method for class 'Tendril'
plot(x, term = NULL, coloring = "Terms",
  interactive = FALSE, ...)
```

Arguments

x An object of class tendril, as made by Tendril()

term The term to extract.

coloring Name of column used for coloring tendrils (only basic plots). Available col-

oring choices are: "Terms" - One tendril one color; "p" - Prop.test; "p.adj" - P-values adjusted using the False discovery rate method within each tendril; "fish" - Fisher.test; "rdiff" - Risk difference; "RR" - Risk Ratio; "OR" - Odds Ratio; "FDR.tot" - P-values adjusted using the False discovery rate method for all tendrils; and "TermsCount" - Total number of events for that specific type of

event

interactive Specifies if the plot must be interactive or not. If interactive == TRUE, plotly

will be used to render the plot. Otherwise, (default) the plot will be rendered as

a static image using ggplot2.

... unused

Details

If saving the results of the function to a variable, this will be of class tendril and will contain the data passed to the plot function and the plot itself

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Examples

```
# generate data using Tendril()
data <- Tendril(mydata = TendrilData,</pre>
rotations = Rotations,
AEfregThreshold = 9,
Tag = "Comment",
Treatments = c("placebo", "active"),
Unique.Subject.Identifier = "subjid",
Terms = "ae",
Treat = "treatment",
StartDay = "day",
SubjList = SubjList,
SubjList.subject = "subjid",
SubjList.treatment = "treatment"
#Do plot
res <- plot(data, coloring = "Terms")</pre>
res <- plot(data, coloring = "p.adj")</pre>
#To re-do the plot after the first call:
print(res)
```

plot.TendrilPerm

Plot TendrilPerm

Description

Function to plot TendrilPerm results

Usage

```
## S3 method for class 'TendrilPerm'
plot(x, coloring = "p.adj", percentile = FALSE,
    ...)
```

Arguments

An object of class TendrilPerm, as made by TendrilPerm()

Name of column used for coloring tendrils. Available coloring choices are:

"Terms" - One tendril one color; "p" - Prop.test; "p.adj" - P-values adjusted using the False discovery rate method within each tendril; "fish" - Fisher.test; "rdiff" - Risk difference; "RR" - Risk Ratio; "OR" - Odds Ratio; "FDR.tot" - P-values adjusted using the False discovery rate method for all tendrils; and "TermsCount" - Total number of events for that specific type of event percentile

Specifies if the plot must show percentile values. Default is FALSE.

unused

plot_timeseries

Examples

```
# generate data using Tendril()
data <- Tendril(mydata = TendrilData,</pre>
rotations = Rotations,
AEfreqThreshold = 9,
Tag = "Comment",
Treatments = c("placebo", "active"),
Unique.Subject.Identifier = "subjid",
Terms = "ae",
Treat = "treatment",
StartDay = "day",
SubjList = SubjList,
SubjList.subject = "subjid",
SubjList.treatment = "treatment"
tendril_perm <- TendrilPerm(</pre>
  tendril = data,
  PermTerm="AE40",
  n.perm = 200,
  perm.from.day = 1)
#Do plot
res <- plot(tendril_perm)</pre>
res <- plot(tendril_perm, percentile = TRUE)</pre>
#To re-do the plot after the first call:
print(res)
```

plot_timeseries

Plot time series

Description

Plot time series of net events on a second treatment arm

Usage

```
plot_timeseries(tendril, term = NULL)
```

Arguments

tendril An object of class Tendril, as made by Tendril()

term A character vector describing the value or values of Term to select; defaults to

NULL which corresponds to all values

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Examples

```
# generate data using Tendril()
data <- Tendril(</pre>
  mydata = TendrilData,
  rotations = Rotations,
  AEfreqThreshold = 9,
  Tag = "Comment",
  Treatments = c("placebo", "active"),
  Unique.Subject.Identifier = "subjid",
  Terms = "ae",
  Treat = "treatment",
  StartDay = "day",
  SubjList = SubjList,
  SubjList.subject = "subjid",
  SubjList.treatment = "treatment"
)
# do plot
plot_timeseries(data, term="AE33")
plot_timeseries(data, term=c("AE33","AE40"))
plot_timeseries(data, term=NULL)
```

Rotations

Example of Rotations in package Tendril.

Description

Example of Rotations in package Tendril.

Usage

Rotations

Format

A numeric vector with 1000 number 3s:

SubjList

Example of SubjList in package Tendril.

Description

Example of SubjList in package Tendril.

Usage

SubjList

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Format

A data frame with 500 rows and 2 variables:

subjid: The subject IDs

treatment: The two treatments causing the tendrils to bend left or right

Tendril *Tendril*

Description

Function to calculate coordinates and statistical measures used to create a tendril plot

Usage

```
Tendril(mydata, rotations, AEfreqThreshold = 50, Tag = "Comment",
   Treatments = c("Active", "Placebo"),
   Unique.Subject.Identifier = "Unique.Subject.Identifier",
   Terms = "Dictionary.Derived.Term", Treat = "Actual.Treatment...DB",
   StartDay = "Analysis.Start.Relative.Day", SubjList = NULL,
   SubjList.subject = NULL, SubjList.treatment = NULL,
   SubjList.dropoutday = NULL, compensate_imbalance_groups = FALSE,
   filter_double_events = FALSE, suppress_warnings = FALSE)
```

Arguments

mydata A dataframe containing the data for the tendril calculations

rotations a vector of same length as mydata containing the rotation factors for all the

events

AEfreqThreshold

The minimum frequency threshold of events to be included in the analysis. De-

fault is 50

Tag A tag or comment associated with the analysis

Treatments The names of the two treatments to be included in the tendril. The first treatment

bends to the right and second treatment bends to the left. Must be a vector of

two elements and the two elements must be found in the Treatment column

Unique.Subject.Identifier

The name of the column containing the unique patients IDs

Terms The name of the column containing the name of the tendrils (e.g. adverse event

terms)

Treat The name of the column containing the name of the treatments

StartDay The name column containing the days of the events
SubjList A dataframe containing subject IDs and treatments

SubjList.subject

The name of the columns in SubjList containing the subjects IDs

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SubjList.treatment

The name of the columns in SubjList containing the treatments

SubjList.dropoutday

The name of the column in SubjList containing the dropoutday

compensate_imbalance_groups

Boolean Whether the rotation factors have been compensated for imbalance in the groups

filter_double_events

Boolean whether to filter out events duplicated in subject id and adverse effect suppress_warnings

Boolean whether to suppress warnings from chi squared approximation may be incorrect

Details

The function accepts a dataframe with at least 4 columns named as the arguments Unique.Subject.Identifier, Terms, Treat and StartDay.

Two treatments must be given as arguments, and at least one of the two treatments must be found in the Treatment column

The function returns an object of class tendril. The object contains the coordinates for the tendril plot and the arguments of the tendril function

The result of the function can be plotted with plot()

The result can be saved to file using write.table() with argument row.names = FALSE

Value

The function return an object of class tendril. The object contains the original dataset added with the tendril coordinates, all the function arguments and a dataframe with the results from statistical analysis

- data\$data: Dataframe of orginal data, coordinates and stat results
- data\$Unique.Subject.Identifier : Column containing subject IDs
- data\$Terms : Column containing the name of the tendrils
- data\$Treat : Column containing the name of the treatments
- data\$StartDay: Column containing the days of the events
- data\$Treatments: The names of the treatments causing the tendrils to bend
- data\$AEfreqThreshold: The frequency threshold for the events to be included in the analysis
- data\$Tag: A tag or comment associated with the analysis
- data\$n.tot : A dataframe with the total number of events for each treatment. Used in the statistical calculations
- data\$SubjList : A dataframe containing subject IDs and treatments
- data\$SubjList.subject: The name of the columns in SubjList containing the subjects IDs
- data\$SubjList.treatment: The name of the columns in SubjList containing the treatments
- data\$SubjList.dropoutday: The name of the column in SubjList containing the dropoutday

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- data\$rotation_vector : Rotation vector used to generate the tendril
- data\$compensate_imbalance_groups : Boolean Whether the rotation factors have been compensated for imbalance in the groups

Examples

```
data <- Tendril(mydata = TendrilData,
rotations = Rotations,
AEfreqThreshold=9,
Tag = "Comment",
Treatments = c("placebo", "active"),
Unique.Subject.Identifier = "subjid",
Terms = "ae",
Treat = "treatment",
StartDay = "day",
SubjList = SubjList,
SubjList.subject = "subjid",
SubjList.treatment = "treatment")
plot(data)</pre>
```

Tendril.perm.res

Example object Tendril as generated by Tendril() and Tendril.perm().

Description

The object contains the following 15 items:

Usage

```
Tendril.perm.res
```

Format

A list of 15 elements:

Tendril.perm.res\$data: Dataframe of orginal data and coordinates

Tendril.perm.res\$Terms: Column containing the name of the tendrils

Tendril.perm.res\$Unique.Subject.Identifier : Columns containing subject IDs

Tendril.perm.res\$Treat: Column containing the name of the treatments

Tendril.perm.res\$StartDay: Column containing the start days of the events

Tendril.perm.res\$Treatments: The names of the treatments causing the tendrils to bend

Tendril.perm.res\$AEfreqThreshold: The frequency threshold for the events to be included in the analysis

Tendril.perm.res\$Tag: A tag or comment associated with the analysis

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Tendril.perm.res\$SubjList : A dataframe containing subject IDs and treatments

Tendril.perm.res\$SubjList.subject: The name of the column in SubjList containing the subjects IDs

Tendril.perm.res\$SubjList.treatment: The name of the columns in SubjList containing the treatments

Tendril.perm.res\$n.tot: A data frame of 2 variables for number of subjects in each treatment

Tendril.perm.res\$PermTerm: Name of the tendril used for the permutations **Tendril.perm.res\$perm.data**: A dataframe containing the permutation results **Tendril.perm.res\$tendril.pi**: A list of permutation percentile calculation results

Tendril.res

Example object Tendril as generated by Tendril().

Description

The object contains the following 12 items:

Usage

Tendril.res

Format

A list of 12 elements:

Tendril.res\$data: Dataframe of orginal data and coordinates **Tendril.res\$Terms**: Column containing the name of the tendrils

Tendril.res\$Unique.Subject.Identifier: Columns containing subject IDs

Tendril.res\$Treat: Column containing the name of the treatments **Tendril.res\$StartDay**: Column containing the start days of the events

Tendril.res\$Treatments: The names of the treatments causing the tendrils to bend

Tendril.res\$AEfreqThreshold: The frequency threshold for the events to be included in the analysis

Tendril.res\$Tag : A tag or comment associated with the analysis

Tendril.res\$SubjList : A dataframe containing subject IDs and treatments

Tendril.res\$SubjList.subject: The name of the column in SubjList containing the subjects IDs **Tendril.res\$SubjList.treatment**: The name of the columns in SubjList containing the treatments **Tendril.res\$SubjList.dropoutday**: The name of the column in SubjList containing the dropout-

day

Tendril.res\$rotation_vector: The rotation vector used for the tendril analysis

Tendril.res\$compensate_imbalance: Boolean indicating whether the imbalance in the groups is compensated

Tendril.res\$n.tot: A data frame of 2 variables for number of subjects in each treatment

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| TendrilData | Example dataframe in package Tendril. | |
|-------------|---------------------------------------|--|
| | | |

Description

A dataset containing the minimal 4 columns required to produe a Tendril plot

Usage

TendrilData

Format

A data frame with 1000 rows and 4 variables:

```
subjid The subject IDs
```

treatment The two treatments causing the tendrils to bend right or left, respectively

ae The names of the tendrils

day The day of the event relative to the treatment start date

| TendrilPerm | Tendril permutations |
|-------------|----------------------|
| | |

Description

Function to compute the permutations of one specified tendril, starting from a specific day. Permutations are simulated under the null hypothesis. Thus, on average, there will be an equal number of events on each treatment arm.

Usage

```
TendrilPerm(tendril, PermTerm, n.perm = 100, perm.from.day = 1,
    pi.low = 0.1, pi.high = 0.9)
```

Arguments

tendril an object of class tendril as produced by Tendril()

PermTerm the name of the type of event (tendril) to calculate permutations on

n.perm the number of permutations. Default 100

perm. from. day the starting day for the permutation calculations. Default 1

pi.low percentile low value. Default 0.1 pi.high percentile high value. Default 0.9 TendrilPerm 11

Details

Make permutation analysis to a specific type of event, as specified in PermTerm.

Value

The function return an object of class TendrilPerm containing all the input data and a dataframe of permutation results. Use:

data\$perm.data and data\$tendril.pi and data\$tendril

to access the permutations, percentiles dataframes, and tendril data respectively

Examples

```
# Create tendril
tendril <- Tendril(mydata = TendrilData,</pre>
  rotations = Rotations,
  AEfregThreshold = 9,
  Tag = "Comment",
  Treatments = c("placebo", "active"),
  Unique.Subject.Identifier = "subjid",
  Terms = "ae",
  Treat = "treatment",
  StartDay = "day",
  SubjList = SubjList,
  SubjList.subject = "subjid",
  SubjList.treatment = "treatment"
)
# Compute permutations
perm.data <- TendrilPerm(tendril = tendril,</pre>
 PermTerm="AE40",
  n.perm = 200,
  perm.from.day = 1)
# Plot results
plot(perm.data)
plot(perm.data, percentile = TRUE)
```

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