Package ‘clinDataReview’

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Type Package

Title Clinical Data Review Tool

Version 1.6.1

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Description Creation of interactive tables, listings and figures (‘TLFs’) and associated report for exploratory analysis of data in a clinical trial, e.g. for clinical oversight activities.
Interactive figures include sunburst, treemap, scatterplot, line plot and barplot of counts data.
Interactive tables include table of summary statistics (as counts of adverse events, enrollment table) and listings.
Possibility to compare data (summary table or listing) across two data batches/sets.
A clinical data review report is created via study-specific configuration files and template ‘R Markdown’ reports contained in the package.

Imports bookdown, clinUtils (>= 0.1.0), crosstalk, data.table, ggplot2, grid, haven, htmltools, htmlwidgets, knitr, jsonlite, jsonvalidate, methods, plotly, plyr, rmarkdown, stats, stringr, utils, tools, yaml, xml2, xfun, base64enc

Suggests countrycode, inTextSummaryTable (>= 3.1.0), patientProfilesVis (>= 0.12.0), testthat, DT, scales, grDevices

SystemRequirements pandoc (to create a clinical data review report)

URL https://github.com/openanalytics/clinDataReview

BugReports https://github.com/openanalytics/clinDataReview/issues

License MIT + file LICENSE

VignetteBuilder knitr

RoxygenNote 7.3.1

NeedsCompilation no

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addDateOfReportRun  

Add date of report running

Description

Add the today’s date of when the report runs to the info of the metadata.

Usage

addDateOfReportRun(summaryInfo)

Arguments

summaryInfo  [matrix, see output from getMetadata.]

Value

A matrix, same as input summaryInfo with an extra row with the date of today.

addFacetPanel  

Add facet-panel to single plotly plot.

Description

Add facet-panel to single plotly plot.
Usage

addFacetPanel(
  pl, panelLab, panelWidth = 20, fontSize = 15, side = c("top", "right")
)

Arguments

  pl  plotly object.
  panelLab  text to be shown in the facet panel
  panelWidth  thickness of the panel in pixels.
  fontSize  fontsize of facetText
  side  the side of the plot to show the panel (currently only right panels are implemented.)

Details

  plot title clipping.
  In case side = 'top', the plot title (eg. layout(title = "title")) will clip with the top panel.
  Resolve this with the following configurations: (once all the subplots have already been combined)
  layout(# place the title at absolute top of the page title = list(text = "title", yref = "container", y = 1)) # If font size = 15 roughly equal to 20 pixels. margin = list(t = panelWidth + heightTitleTextInPixels)

Value

  plotly object with the facet panel added.

Author(s)

  lennart tuijnder

addLayerToScatterPlot  Helper function to add layer to scatter plot

Description

  Helper function to add layer to scatter plot
addReferenceLinesClinDataPlot

Usage

addLayerToScatterPlot(
  gg,
  aesVar,
  pars,
  generalPars,
  layerFunction,
  useHandlers = FALSE
)

Arguments

gg ggplot object
aesVar layers specific aesthetics list of layer specific aesthetics
pars list of parameters specific to the layer aes
generalPars overall, not layer specific parameters can be overwritten by pars
layerFunction function to use for adding the layer e.g. geom_line
useHandlers if TRUE we use handlers to repress the expected warning: Hover is set via the 'text' aesthetic in ggplot, we need to pass this aesthetic to have it available in plotly even though it is not used by geom_point. Defaults to FALSE

Value

ggplot object

Author(s)

Adriaan Blommaert Laure Cougnaud

addReferenceLinesClinDataPlot

Description

Add reference (horizontal/vertical/diagonal) lines to a clinical data plot

Usage

addReferenceLinesClinDataPlot(
  gg,
  data,
  xVar,
  yVar,
Arguments

- **gg** (ggplot object.)
- **data** (Data frame with data.)
- **xVar** (String with column of data containing x-variable.)
- **yVar** (String with column of data containing y-variable.)
- **xLim, yLim** (Numeric vector of length 2 with limits for the x/y axes.)
- **refLinePars** (optional) Nested list, with parameters for each reference line(s). Each sublist contains:
  - aesthetic value(s) or variable(s) for the lines (in this case column names of data) for reference lines. The line position is controlled by the aesthetics supported in `geom_vline`, `geom_hline` and `geom_abline`.
  - 'label': (optional) Logical specifying if the line should be annotated (FALSE to not annotate the line) or string with annotation label. By default, the value of the position of the horizontal/vertical line or the equation of the diagonal line is displayed.

- **facetPars** (List with facetting parameters, passed to the facetting function. Variables should be specified as character or formula. For 'wrap' facetting (facetType is 'wrap'), if the layout is not specified via nrow/ncol, 2 columns are used by default.)

Value

- Updated ggplot object.

Author(s)

Laure Cougnaud

---

**addSelectBtn**  
*Add selection box(es) to a plotly plot.*

Description

Add selection box(es) to a plotly plot.
Usage

addSelectBtn(
  data,
  pl,
  selectVars = NULL,
  selectLab = NULL,
  labelVars = NULL,
  id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
  keyVar
)

Arguments

data  
  SharedData object used for the plot.

pl  
  plotly object.

selectVars  
  (optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).

selectLab  
  (Named) character vector with label for selectVars.

labelVars  
  Named character vector containing variable labels.

id  
  String with general id for the plot:
  • 'id' is used as group for the SharedData
  • 'button:[id]' is used as button ID if table is TRUE

  If not specified, a random id, as 'plotClinData[X]' is used.

keyVar  
  String with unique key variable, identifying unique group for which the link between the table and the plot should be done.

Value

if selectVars is specified: a browsable object combining the select buttons and the plotly object. Otherwise, the input plotly object.

addWatermark

Add a watermark to a plotly object.

Description

Add a watermark to a plotly object.

Usage

addWatermark(pl, watermark = NULL)
**annotateData**

**Arguments**

- `pl` : plotly object.
- `watermark` : (optional) String with path to a file containing a watermark.

**Value**

plotly object with a watermark (if specified)

**See Also**

Other watermark helpers: `getWatermark()`

**Examples**

```r
watermark <- tempfile(pattern = "watermark", fileext = ".png")
getWatermark(file = watermark)
addWatermark(pl = plotly::plot_ly(), watermark = watermark)
```

---

**Annotate a dataset.**

**Description**

Standard annotation variables are available via the parameter `annotType`. Custom dataset/variables of interest are specified via the `annotDataset/annotVar` parameters.

**Usage**

```r
annotateData(
  data,
  dataPath = ".",
  annotations,
  subjectVar = "USUBJID",
  verbose = FALSE,
  labelVars = NULL,
  labelData = "data"
)
```

**Arguments**

- `data` : Data.frame with input data to annotate.
- `dataPath` : String with path to the data.
- `annotations` : Annotations (or list of those) either as a:
  - string with standard annotation type, among:
    - demographics: standard variables from the demographics data (DM or ADSL) are extracted
- exposed_subjects: a logical variable: `EXFL` is added to data, identifying exposed subjects, i.e. subjects included in the exposure dataset (EX/ADEX) dataset and with non empty and non missing start date (‘EXSTDTC’, ‘STDY’ or ‘ASTDY’)

- functional_groups_lab: a character variable: ‘LBFCTGRP’ is added to data based on standard naming of the parameter code (‘PARAMCD’ or ‘LBTESTCD’ variable)

* list of custom annotation, with:
  - (optional) annotation dataset, either:
    * ‘dataset’: String with name of the annotation dataset, e.g. ‘ex’ to import data from the file: `[dataset].sas7bdat` in dataPath
    * ‘data’: Data.frame with annotation dataset
      The input data is used if ‘data’ and ‘dataset’ are not specified.
  - ‘vars’: Either:
    * Character vector with variables of interest from annotation dataset. If not specified, all variables of the dataset are considered.
    * String with new variable name computed from varFct
  - ‘varFct’: (optional) Either:
    * function of data or string containing such function (e.g. ‘func(data) ...’)
    * string containing manipulations from column names of data (e.g. ‘col1 + col2’)
      used to create a new variable specified in vars.
  - ‘filters’: (optional) Filters for the annotation dataset, see filters parameter of filterData.
      The annotation dataset is first filtered, before being combined to the input data, such as only the records retained in the annotation dataset will be annotated in the output data. Other records will have missing values in the annotated variables.
  - ‘varLabel’: (optional) label for new variable in case varFct is specified.
  - ‘varsBy’: (optional) Character vector with variables used to merge input data and the annotation dataset. If not specified:
    * if an external dataset (dataset/data) is specified: subjectVar is used
    * otherwise: annotation dataset and input data are merged by rows IDs

subjectVar String with subject ID variable, ‘USUBJID’ by default.
verbose Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.
labelVars Named character vector containing variable labels of data. This will be updated with the labels of the extra annotation variables (in attr(output, ‘labelVars’)).
labelData (optional) String with label for input data, that will be included in progress messages.
**Value**

Annotated data. If labelVars is specified, the output contains an extra attribute: 'labelVars' containing updated labelVars (accessible via: in attr(output, 'labelVars')).

**Examples**

```r
library(clinUtils)

data(dataADaMCDISCP01)

dataLB <- dataADaMCDISCP01$ADLBC
dataDM <- dataADaMCDISCP01$ADSL
dataAE <- dataADaMCDISCP01$ADAE

labelVars <- attr(dataADaMCDISCP01, "labelVars")

# standard annotations:
# path to dataset should be specified via: 'pathData'
## Not run:
annotateData(dataLB, annotations = "demographics", pathData = ...)
## End(Not run)

# add all variables in annotation data (if not already available)
head(annotateData(dataLB, annotations = list(data = dataDM)), 1)

# only variables of interest
head(annotateData(dataLB, annotations = list(data = dataDM, vars = c("ARM", "ETHNIC"))), 1)

# filter annotation dataset
dataAnnotated <- annotateData(dataLB,
  annotations = list(
    data = dataDM,
    vars = c("ARM", "ETHNIC"),
    filters = list(var = "ARM", value = "Placebo")
  )
)

head(subset(dataAnnotated, ARM == "Placebo"), 1)
head(subset(dataAnnotated, is.na(ARM)), 1)

# worst-case scenario: add a new variable based on filtering condition
dataAE$AESEV <- factor(dataAE$AESEV, levels = c('MILD', 'MODERATE', "SEVERE"))
dataAEWC <- annotateData(
  data = dataAE,
  annotations = list(
    vars = "WORSTINT",
    # create new variable: 'WORSTINT'
    # with TRUE if maximum toxicity grade per subject/test
    # (if multiple, they are all retained)
    filters = list(
      var = "AESEV",
      # max will take latest level in a factor
```
# (so 'MODERATE' if 'MILD'/'MODERATE' are available)
valueFct = function(x) x[which.max(as.numeric(x))],
varsBy = c("USUBJID", "AEDECOD"),
keepNA = FALSE,
varNew = "WORSTINT",
labelNew = "worst-case"
)
labelVars = labelVars,
verbose = TRUE
)
attr(dataAEWC, "labelVars")["WORSTINT"]

# add a new variable based on a combination of variables:
dataLB <- annotateData(dataLB,
annotations = list(vars = "HILORATIO", varFct = "A1HI / A1LO")
)

# add a new variable based on extraction of a existing variable
# Note: slash should be doubled when the function is specified as text
dataLB <- annotateData(dataLB,
annotations = list(vars = "PERIOD", varFct = "sub(\.\* Week (.+)' Week \\1', AVISIT)"")
)

# multiple annotations:
dataAnnotated <- annotateData(dataLB,
annotations = list(
list(data = dataDM, vars = c("ARM", "ETHNIC")),
list(data = dataAE, vars = c("AESEV"))
)
)
head(dataAnnotated, 1)

---

**barplotClinData**  
*Barplot visualization of clinical data.*

**Description**

Barplot visualization of clinical data.

**Usage**

```r
barplotClinData(
  data,
  xVar,
  yVar,
  xLab = getLabelVar(xVar, labelVars = labelVars),
  xLabVar = NULL,
  yLab = getLabelVar(yVar, labelVars = labelVars),
  yLabVar = NULL,
)```
colorVar = NULL,
colorLab = getLabelVar(colorVar, labelVars = labelVars),
colorPalette = NULL,
barmode = "group",
title = paste(c(paste(yLab, "vs", xLab), titleExtra), collapse = "<br>")
titleExtra = NULL,
caption = NULL,
subtitle = NULL,
labelVars = NULL,
width = NULL,
height = NULL,
hoverVars,
hoverLab,
textVar = NULL,
pathVar = NULL,
pathLab = getLabelVar(pathVar, labelVars = labelVars),
table = FALSE,
tableVars,
tableLab,
tableButton = TRUE,
tablePars = list(),
id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
selectVars = NULL,
selectLab = getLabelVar(selectVars, labelVars = labelVars),
watermark = NULL,
verbose = FALSE
)

Arguments

data       Data.frame with data.
xVar       String with column of data containing x-variable.
yVar       String with column of data containing y-variable.
xLab       String with label for xVar.
xLabVar    (optional) Character vector with column(s) of data containing variable(s) to display in the label of the x-axis.
yLab       String with label for yVar.
yLabVar    (optional) Character vector with column(s) of data containing variable(s) to display in the label of the y-axis.
colorVar   (optional) String with color variable.
colorLab   String with label for colorVar.
colorPalette (optional) Named character vector with color palette. If not specified, the viridis color palette is used.
            See clinColors.
barmode    String with type of barplot, either: 'group' or 'stack' (see parameter in layout).
title      String with title for the plot.
<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><code>titleExtra</code></td>
<td>String with extra title for the plot (appended after title).</td>
</tr>
<tr>
<td><code>caption</code></td>
<td>String with caption. The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.</td>
</tr>
<tr>
<td><code>subtitle</code></td>
<td>String with subtitle. The subtitle is included at the top left of the plot, below the title.</td>
</tr>
<tr>
<td><code>labelVars</code></td>
<td>Named character vector containing variable labels.</td>
</tr>
<tr>
<td><code>width</code></td>
<td>Numeric, width of the plot in pixels, 800 by default.</td>
</tr>
<tr>
<td><code>height</code></td>
<td>Numeric, height of the plot in pixels, 500 by default.</td>
</tr>
<tr>
<td><code>hoverVars</code></td>
<td>Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.</td>
</tr>
<tr>
<td><code>hoverLab</code></td>
<td>Named character vector with labels for <code>hoverVars</code>.</td>
</tr>
<tr>
<td><code>textVar</code></td>
<td>(optional) String with a text variable, that will be displayed outside of each bar.</td>
</tr>
<tr>
<td><code>pathVar</code></td>
<td>String with variable of <code>data</code> containing hyperlinks with path to the subject-specific report, formatted as: &lt;a href=&quot;./path-to-report&quot;&gt;label&lt;/a&gt;</td>
</tr>
<tr>
<td><code>pathLab</code></td>
<td>String with label for <code>pathVar</code>, included in the collapsible row in the table.</td>
</tr>
<tr>
<td><code>table</code></td>
<td>Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)</td>
</tr>
<tr>
<td><code>tableVars</code></td>
<td>Character vector with variables to be included in the table.</td>
</tr>
<tr>
<td><code>tableLab</code></td>
<td>Named character vector with labels for each <code>tableVars</code>.</td>
</tr>
<tr>
<td><code>tableButton</code></td>
<td>Logical, if TRUE (by default) the table is included within an HTML button.</td>
</tr>
<tr>
<td><code>tablePars</code></td>
<td>List with parameters passed to the <code>getClinDT</code> function.</td>
</tr>
<tr>
<td><code>id</code></td>
<td>String with general id for the plot:</td>
</tr>
<tr>
<td></td>
<td>• <code>id</code> is used as group for the <code>SharedData</code></td>
</tr>
<tr>
<td></td>
<td>• <code>button:[id]</code> is used as button ID if <code>table</code> is TRUE</td>
</tr>
<tr>
<td><code>selectVars</code></td>
<td>(optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).</td>
</tr>
<tr>
<td><code>selectLab</code></td>
<td>(Named) character vector with label for <code>selectVars</code>.</td>
</tr>
<tr>
<td><code>watermark</code></td>
<td>(optional) String with path to a file containing a watermark.</td>
</tr>
<tr>
<td><code>verbose</code></td>
<td>Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.</td>
</tr>
</tbody>
</table>
**Value**

Either:

- if a table is requested: a clinDataReview object, a.k.a a list with the 'plot' (**plotly** object) and 'table' (**datatable** object)
- otherwise: a **plotly** object

**Author(s)**

Laure Cougnaud

**See Also**

Other visualizations of summary statistics for clinical data: **boxplotClinData()**, **errorbarClinData()**, **plotCountClinData()**, **sunburstClinData()**, **treemapClinData()**

**Examples**

```r
library(clinUtils)

data(dataADaMCDISCP01)
labelVars <- attr(dataADaMCDISCP01, "labelVars")

dataAE <- dataADaMCDISCP01$ADAE
dataDM <- dataADaMCDISCP01$ADSL

## example of basic barplot:

# treemap takes as input table with counts

if (requireNamespace("inTextSummaryTable", quietly = TRUE)) {

  # total counts: Safety Analysis Set (patients with start date for the first treatment)
dataTotal <- subset(dataDM, RFSTDTC != "")

  # compute adverse event table
tableAE <- inTextSummaryTable::computeSummaryStatisticsTable(
data = dataAE,
rowVar = c("AEBODSYS", "AEDECOD"),
rowOrder = "total",
dataTotal = dataTotal,
labelVars = labelVars,
stats = inTextSummaryTable::getStats("count")
)

dataPlot <- subset(tableAE, AEDECOD != "Total")
dataPlot$n <- as.numeric(dataPlot$n)

# create plot
```
barplotClinData(
  data = dataPlot,
  xVar = "AEDECOD",
  yVar = "n", yLab = "Number of patients with adverse events",
  labelVars = labelVars
)

# add number on top of the bars
barplotClinData(
  data = dataPlot,
  xVar = "AEDECOD",
  yVar = "n", yLab = "Number of patients with adverse events",
  textVar = "n",
  labelVars = labelVars
)

# add a selection box
if(interactive()){
  barplotClinData(
    data = dataPlot,
    xVar = "AEDECOD",
    yVar = "n", yLab = "Number of patients with adverse events",
    labelVars = labelVars,
    selectVars = "AEBODSYS"
  )
}

## Not run:
# display percentage of events per severity

```r

# display percentage of events per severity
tableAEBySeverity <- inTextSummaryTable::computeSummaryStatisticsTable(
  data = dataAE,
  rowVar = c("AEDECOD", "AESEV"),
  dataTotal = dataTotal,
  labelVars = labelVars,
  statsPerc = "statm",
  stats = inTextSummaryTable::getStats("%m"),
  dataTotalPerc = dataAE,
  rowVarTotalPerc = "AEDECOD"
)

barplotClinData(
  data = tableAEBySeverity,
  xVar = "AEDECOD", xLab = "Adverse event term",
  yVar = "statPercm", yLab = "Percentage of adverse events",
  labelVars = labelVars,
  colorVar = "AESEV", barmode = "stack",
  hoverVar = c("AEDECOD", "AESEV", "statN", "statm", "statPercm"),
  hoverLab = c(labelVars["AEDECOD"],
               labelVars["AESEV"],
               statN = "Number of patients",
               statm = "Number of events",
               statPercm = "Percentage of events")
```

---

**barplotClinData**
## Description

Boxplot interactive plot.

## Usage

```r
boxplotClinData(
  data,
  xVar,
  yVar,
  xLab = getLabelVar(xVar, labelVars = labelVars),
  xLabVar = NULL,
  yLab = getLabelVar(yVar, labelVars = labelVars),
  yLabVar = NULL,
  colorVar = NULL,
  colorLab = getLabelVar(colorVar, labelVars = labelVars),
  colorPalette = NULL,
  facetVar = NULL,
  facetLab = getLabelVar(facetVar, labelVars = labelVars),
  ncol = 1L,
  title = paste(c(paste(yLab, "vs", xLab), titleExtra), collapse = "<br>")
)```

### Arguments

- `data`: data frame containing the variables.
- `xVar`: variable for the x-axis.
- `yVar`: variable for the y-axis.
- `xLab`: label for the x-axis.
- `xLabVar`: variable for the x-axis label.
- `yLab`: label for the y-axis.
- `yLabVar`: variable for the y-axis label.
- `colorVar`: variable for the color.
- `colorLab`: label for the color.
- `colorPalette`: color palette.
- `facetVar`: variable for faceting.
- `facetLab`: label for faceting.
- `ncol`: number of columns for facetting.
- `title`: title of the plot.
- `titleExtra`: additional title.
- `subtitle`: subtitle of the plot.
- `caption`: caption.
- `labelVars`: label variables.
- `width`: width of the plot.
- `height`: height of the plot.
- `hoverVars`: variables for hover.
- `hoverLab`: labels for hover.
- `pathVar`: variable for path.
- `pathLab`: label for path.
- `idVar`: variable for id.
- `idLab`: label for id.
- `table`: table option.
Arguments

data Data.frame with data.
xVar String with column of data containing x-variable.
yVar String with column of data containing y-variable.
xLab String with label for xVar.
xLabVar (optional) Character vector with column(s) of data containing variable(s) to display in the label of the x-axis.
yLab String with label for xVar.
yLabVar (optional) Character vector with column(s) of data containing variable(s) to display in the label of the y-axis.
colorVar (optional) String with color variable.
colorLab String with label for colorVar.
colorPalette (optional) Named character vector with color palette. If not specified, the viridis color palette is used. See clinColors.
facetVar (optional) String with facet variable.
facetLab String with label for facetVar.
col single-length integer denoting the number of columns for the facetting.
title String with title for the plot.
titleExtra String with extra title for the plot (appended after title).
subtitle String with subtitle.
The subtitle is included at the top left of the plot, below the title.
caption String with caption.
The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.
labelVars Named character vector containing variable labels.
width Numeric, width of the plot in pixels, 800 by default.
height Numeric, height of the plot in pixels, 500 by default.
hoverVars Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.
hoverLab Named character vector with labels for hoverVars.
### boxplotClinData

**pathVar**
String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

```html
<a href="/path-to-report">label</a>
```

If multiple, they should be separated by ``,`.

The report(s) will be:
- compressed to a zip file and downloaded if the user clicks on the `p` (a.k.a `profile`) key when hovering on a point of the plot
- included in a collapsible row, and clickable with hyperlinks in the table

**pathLab**
String with label for `pathVar`, included in the collapsible row in the table.

**idVar**
String with variable containing subject ID.

**idLab**
String with label for `idVar`.

**table**
Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)

**tableVars**
Character vector with variables to be included in the table.

**tableLab**
Named character vector with labels for each `tableVars`.

**tableButton**
Logical, if TRUE (by default) the table is included within an HTML button.

**tablePars**
List with parameters passed to the `getClinDT` function.

**id**
String with general id for the plot:
- `id` is used as group for the `SharedData`
- `button:[id]` is used as button ID if `table` is TRUE

If not specified, a random id, as `plotClinData[X]` is used.

**watermark**
(optional) String with path to a file containing a watermark.

**verbose**
Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.

### Value
Either:
- if a table is requested: a `clinDataReview` object, a.k.a a list with the `plot` (`plotly` object) and `table` (`datatable` object)
- otherwise: a `plotly` object

### Author(s)
Lennart Tuijnder

### See Also
Other visualizations of summary statistics for clinical data: `barplotClinData()`, `errorbarClinData()`, `plotCountClinData()`, `sunburstClinData()`, `treemapClinData()`
Examples

```r
library(clinUtils)

data(dataADaMCDISCP01)
labelVars <- attr(dataADaMCDISCP01, "labelVars")

## example of basic boxplot:

data <- subset(dataADaMCDISCP01$ADVS, PARAMCD == "DIABP" & ANL01FL == "Y" & AVISIT %in% c("Baseline", "Week 2", "Week 4", "Week 6", "Week 8")
)

## example of basic boxplot:
# With color var and facet:
boxplotClinData(
data = data,
xVar = "AVISIT",
yVar = "AVAL",
colorVar = "TRTA",
facetVar = "ATPT",
title = "Diastolic Blood Pressure distribution by actual visit and analysis timepoint",
yLab = "Actual value of the Diastolic Blood Pressure parameter (mmHg)",
labelVars = labelVars
)

# Control number of facet columns:
boxplotClinData(
data = data,
xVar = "AVISIT",
yVar = "AVAL",
colorVar = "TRTA",
facetVar = "ATPT",
ncol = 2,
title = "Diastolic Blood Pressure distribution by actual visit and analysis timepoint",
yLab = "Actual value of the Diastolic Blood Pressure parameter (mmHg)",
labelVars = labelVars
)

## Not run:

# Facet or color is optional:
boxplotClinData(
data = data,
xVar = "AVISIT",
yVar = "AVAL",
colorVar = "TRTA"
)

boxplotClinData(
data = data,
...
buildBook

xVar = "AVISIT",
yVar = "AVAL",
facetVar = "ATPT"

# add caption & subtitle
boxplotClinData(
data = data,
xVar = "AVISIT",
yVar = "AVAL",
facetVar = "ATPT", ncol = 2,
colorVar = "TRTA",
title = "Diastolic Blood Pressure distribution",
subtitle = "By actual visit and analysis timepoint",
yLab = "Actual value of the Diastolic Blood Pressure parameter (mmHg)",
caption = "Summary statistics are computed internally.",
labelVars = labelVars
)

## End(Not run)

---

**buildBook**  
*Build the book*

**Description**

Build the book

**Usage**

`buildBook(htmlFiles, verbose = TRUE)`

**Arguments**

- `htmlFiles` character vector with path to HTML files
- `verbose` Logical, if TRUE (FALSE by default) progress messages are printed during the report execution.

**Value**

String with path to the front page of the report.

**Author(s)**

Laure Cougnaud
checkAvailabilityMetadata

*Check availability of arguments in list*

**Description**

Check availability of arguments in list

**Usage**

```r
checkAvailabilityMetadata(paramsList, subListName)
```

**Arguments**

- **paramsList**: A named list.
- **subListName**: String indicating which of the sublist names to check for existance.

**Value**

The content of the sublist. If not available, returns "Not Available".

checkChapterParallel

*Check if a chapter is run internally in parallel or not.*

**Description**

This is identified via the 'parallel' parameter from the config file. If this parameter is not available in the config file (or the parameters are imported with an error), the chapter is considered to not be run in parallel.

**Usage**

```r
checkChapterParallel(
  configFile,
  configDir = file.path(inputDir, "config"),
  inputDir = "."
)
```

**Arguments**

- **configFile**: String with filename of the config file of interest in YAML format.
- **configDir**: String with directory with config files, by default a 'config' folder in `inputDir`. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general general 'config.yml'.
- **inputDir**: String with input directory, working directory by default.
checkConfigFile

Value

Logical, if TRUE, the chapter is run in parallel (FALSE if not available).

Description

Check a configuration file (in _YAML_ format) based on a requirement file in JSON Schema format.

Usage

checkConfigFile(configFile, configSpecFile, configDir = "./config")

Arguments

- **configFile**: path to the config file
- **configSpecFile**: String with path to the file containing requirements in JSON Schema format.
- **configDir**: String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general 'config.yml'.

Value

No returned value, an error message is printed in the console if the configuration file doesn’t comply to the specified specifications.

Author(s)

Laure Cougnaud

checkReportTitles

Description

Check uniqueness of report titles across the config files. If not unique titles are provided, an error is returned.
checkTemplatesName

Usage

checkReportTitles(
  configFiles,
  configDir = file.path(inputDir, "config"),
  inputDir = "."
)

Arguments

  configFiles  Character vector with config file names
  configDir    String with directory with config files, ('config' by default)
  inputDir     String with input directory, working directory by default.

Value

A named vector with the report titles and the corresponding config file

Author(s)

Michela Pasetto

See Also

Other clinical data reporting: forceParams(), getMdHeader(), getParamsFromConfig(), gitbook_clinDataReview_report(), html_clinDataReview_report(), knitPrintClinDataReview(), postProcessReport(), render_clinDataReviewReport

---

checkTemplatesName  Checks of config files template.

Description

Check if the templates specified in the input config files don’t originate from multiple sources (e.g. custom and R package via the parameter templatePackage). If so, the corresponding config files are not considered.

Usage

checkTemplatesName(
  configFiles,
  configDir = file.path(inputDir, "config"),
  inputDir = "."
)
checkValueType

Arguments

configFiles Character vector with name or path of the config file(s).
configDir String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general general 'config.yml'.
inputDir String with input directory, working directory by default.

Value

Updated configFiles

Author(s)

Laure Cougnaud

checkValueType (data, vars, valueVar, valueType = "total", labelVars = NULL)

Description

If this parameter is set to 'total' and the sum of the counts of the the children nodes is not bigger than the parent node, an empty plot is created. In this case, this function set this parameter to: 'relative'.

Usage

checkValueType(data, vars, valueVar, valueType = "total", labelVars = NULL)

Arguments

data Data.frame with data.
vars Character vector with variables of data containing the groups. If multiple, they should be specified in hierarchical order (from parent to child node).
valueVar String with numeric variable of data containing the value to display.
valueType String with type of values in valueVar (branchvalues of the plot_ly function), among others: 'total' (default, only if sum(child) <= to parent) or 'relative'.
labelVars Named character vector containing variable labels.

Value

If the condition is fullfilled: updated valueType and warning; otherwise input valueType.
Author(s)

Laure Cougnaud

---

**clinDataReview-common-args**

*Common arguments for the functions of the clinDataReview package*

---

**Description**

Common arguments for the functions of the clinDataReview package

**Arguments**

- **data**  
  Data frame with data.

- **verbose**  
  Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.

- **gg**  
  ggplot object.

- **pl**  
  plotly object.

- **xVar**  
  String with column of data containing x-variable.

- **yVar**  
  String with column of data containing y-variable.

- **xLab**  
  String with label for xVar.

- **xLabVar**  
  (optional) Character vector with column(s) of data containing variable(s) to display in the label of the x-axis.

- **yLab**  
  String with label for yVar.

- **yLabVar**  
  (optional) Character vector with column(s) of data containing variable(s) to display in the label of the y-axis.

- **xLim, yLim**  
  Numeric vector of length 2 with limits for the x/y axes.

- **idVar**  
  String with variable containing subject ID.

- **idLab**  
  String with label for idVar.

- **width**  
  Numeric, width of the plot in pixels, 800 by default.

- **height**  
  Numeric, height of the plot in pixels, 500 by default.

- **facetPars**  
  List with facetting parameters, passed to the facetting function. Variables should be specified as character or formula. For 'wrap' facetting (FacetType is 'wrap'), if the layout is not specified via nrow/ncol, 2 columns are used by default.

- **lineVars**  
  List with parameters for the reference lines.

- **hoverVars**  
  Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.

- **hoverLab**  
  Named character vector with labels for hoverVars.
**clinDataReview-common-args**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>pathExpand</td>
<td>Logical, should the variable in pathExpand be included in a collapsible row or as hyperlink in the table? Should be TRUE for if multiple paths are included for each idVar, FALSE otherwise (by default).</td>
</tr>
<tr>
<td>table</td>
<td>Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. The plot and table are linked when included in a Rmarkdown document: when clicking on an plot element, only the corresponding records are retained in the associated table; when some records are selected in the table, they are highlighted in the associated table.</td>
</tr>
<tr>
<td>refLinePars</td>
<td>(optional) Nested list, with parameters for each reference line(s). Each sublist (a.k.a reference line) contains:</td>
</tr>
<tr>
<td></td>
<td>• aesthetic value(s) or variable(s) for the lines (in this case column names of data) for reference lines. The line position is controlled by the aesthetics supported in <code>geom_vline</code>, <code>geom_hline</code> and <code>geom_abline</code>.</td>
</tr>
<tr>
<td></td>
<td>• 'label': (optional) Logical specifying if the line should be annotated (FALSE to not annotate the line) or string with annotation label. By default, the value of the position of the horizontal/vertical line or the equation of the diagonal line is displayed.</td>
</tr>
<tr>
<td>labelVars</td>
<td>Named character vector containing variable labels.</td>
</tr>
<tr>
<td>id</td>
<td>String with general id for the plot:</td>
</tr>
<tr>
<td></td>
<td>• 'id' is used as group for the <code>SharedData</code></td>
</tr>
<tr>
<td></td>
<td>• 'button:[id]' is used as button ID if <code>table</code> is TRUE</td>
</tr>
<tr>
<td></td>
<td>If not specified, a random id, as 'plotClinData[X]' is used.</td>
</tr>
<tr>
<td>title</td>
<td>String with title for the plot.</td>
</tr>
<tr>
<td>titleExtra</td>
<td>String with extra title for the plot (appended after <code>title</code>).</td>
</tr>
<tr>
<td>caption</td>
<td>String with caption.</td>
</tr>
<tr>
<td></td>
<td>The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.</td>
</tr>
<tr>
<td>subtitle</td>
<td>String with subtitle.</td>
</tr>
<tr>
<td></td>
<td>The subtitle is included at the top left of the plot, below the title.</td>
</tr>
<tr>
<td>colorVar</td>
<td>(optional) String with color variable.</td>
</tr>
<tr>
<td>colorLab</td>
<td>String with label for <code>colorVar</code>.</td>
</tr>
<tr>
<td>colorPalette</td>
<td>(optional) Named character vector with color palette. If not specified, the viridis color palette is used. See <code>clinColors</code>.</td>
</tr>
<tr>
<td>selectVars</td>
<td>(optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).</td>
</tr>
<tr>
<td>selectLab</td>
<td>(Named) character vector with label for <code>selectVars</code>.</td>
</tr>
<tr>
<td>keyVar</td>
<td>String with unique key variable, identifying unique group for which the link between the table and the plot should be done.</td>
</tr>
<tr>
<td>watermark</td>
<td>(optional) String with path to a file containing a watermark.</td>
</tr>
</tbody>
</table>

**Value**

No return value, used for the documentation of the functions of the package.
**Common parameters for the clinical data reporting function**

### Description

Common parameters for the clinical data reporting function

### Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>indexPath</strong></td>
<td>String with path to the index file, by default 'index.Rmd' in inputDir.</td>
</tr>
<tr>
<td><strong>configDir</strong></td>
<td>String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general general 'config.yml'.</td>
</tr>
<tr>
<td><strong>configFile</strong></td>
<td>String with filename of the config file of interest in YAML format.</td>
</tr>
<tr>
<td><strong>inputDir</strong></td>
<td>String with input directory, working directory by default.</td>
</tr>
<tr>
<td><strong>outputDir</strong></td>
<td>String with output directory, ('report' by default).</td>
</tr>
<tr>
<td><strong>intermediateDir</strong></td>
<td>String with intermediate directory ('interim' by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.</td>
</tr>
<tr>
<td><strong>extraDirs</strong></td>
<td>Character vector with extra directories required by the report, directory with external images. By default, the directories: 'figures', 'tables' and mentioned in the 'patientProfilePath' parameter of the general config file are included. All these folders should be available in inputDir.</td>
</tr>
<tr>
<td><strong>mdFile</strong></td>
<td>String with path of the Markdown file</td>
</tr>
<tr>
<td><strong>logFile</strong></td>
<td>(optional) String with path to a log file, where output (also error/messages/warnings) should be stored. If specified, the entire output is re-directed to this file.</td>
</tr>
<tr>
<td><strong>nCores</strong></td>
<td>Integer containing the number of cores used to render the report (1 by default). If more than 1, two steps of the report creation are run in parallel across chapters:</td>
</tr>
<tr>
<td><strong>verbose</strong></td>
<td>Logical, if TRUE (FALSE by default) progress messages are printed during the report execution.</td>
</tr>
</tbody>
</table>

### Value

No return value, used for the documentation of the clinical data reporting functions of the package.
Common arguments for the plotting functions summary statistics of the clinDataReview package

**Arguments**

- **vars**
  Character vector with variables of data containing the groups. If multiple, they should be specified in hierarchical order (from parent to child node).

- **varsLab**
  Named character vector with labels for `vars`.

- **valueVar**
  String with numeric variable of data containing the value to display.

- **valueLab**
  String with label for the `valueVar` variable.

- **valueType**
  String with type of values in `valueVar` (`branchvalues` of the `plot_ly` function), among others: 'total' (default, only if sum(child) <= to parent) or 'relative'.

- **pathVar**
  String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

  `<a href="./path-to-report">label</a>`

  If multiple, they should be separated by ': , '.

  The report(s) will be:
  - compressed to a zip file and downloaded if the user clicks on the 'p' (a.k.a 'profile') key when hovering on a point of the plot
  - included in a collapsible row, and clickable with hyperlinks in the table

- **pathLab**
  String with label for `pathVar`, included in the collapsible row in the table.

- **table**
  Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)

**Value**

No return value, used for the documentation of the plotting functions of summary statistics of the package.
Description
Template reports with standard visualizations/tables available in the package are described here.

Details
For each template, required parameters are indicated in **bold**.

Value
No return value, used for the documentation of the Rmarkdown template reports contained in the package.

Parameter type
Please note that the type mentioned below corresponds to the type in the config file (in YAML/JSON format).
The mapping to R data type is as followed:

- **string**: character vector of length 1
- **integer**: integer vector of length 1
- **array**: vector/list without names
- **object**: list with names

Clinical data template for a visualization of count data: **countsVisualizationTemplate**
This report compute counts of variable(s) of interest (with the inTextSummaryTable package) and visualize them with a treemap and/or sunburst.
The following parameters are available:

- **template**: string set to: 'countsVisualizationTemplate.Rmd', name of the template report
- **templatePackage**: string set to: 'clinDataReview', package from which the template should be extracted
- **reportTitle**: string, header title
- **reportTitleLevel**: (optional) integer, header level, 1 by default (1: 'chapter', '2': 'section', '3': subsection, ...)
- **parallel**: (optional) boolean, does this chapter use parallel execution? If the entire report is run in parallel, this ensures that this specific chapter is created outside of the report parallelization.
- **split_by**: (optional) integer of length: minimum 0, maximum 7 or string among: 'none', 'chapter', 'section', split the chapter at the specified level: 1 (for 'chapter'), 2 (for 'section') until 7. This overwrites the 'split_by' parameter defined in the output format of the report.
• `dataFileName`: string or array, name of the data file(s) of interest. If multiple files are specified, the data are combined by rows ("row bind"), with a column: "DATASET" containing the name of the file (in upper case and without extension) the data originate from.

• `dataProcessing`: (optional) array, data processing parameters, passed to `processData`

• `dataTotalFileName`: (optional) string, filename of the total dataset

• `dataTotalProcessing`: (optional) array, data processing parameters for 'dataTotalFileName', passed to `processData`

• `countVar`: string, variable of data to count on

• `parentVar`: (optional) string or array, parent variable(s) of the counting variable, used for grouping

• `colorVar`: (optional) string or object, numeric variable(s) to consider for coloring, named by count/parent variable if different for each variable

• `colorRange`: (optional) array of number(s) of length: minimum 2, maximum 2, range of the color variable for the visualization

• `loopingVar`: (optional) array or string, data variable(s) to loop over. Each group of the variable(s) is displayed in a separated section of the report.

• `loopingNMax`: (optional) integer, maximum number of elements of `loopingVar` to include in the report

• `loopingTotal`: (optional) boolean, should the total also be computed by `loopingVar` (TRUE by default)?

• `typePlot`: (optional) array or string of string(s) among: 'sunburst', 'treemap'

• `startup`: (optional) array or string, R commands that should be run at the start of the report

Clinical data template to create a report division: `divisionTemplate`

This report includes a division, i.e. extra chapter, section of subsection in the report. The following parameters are available:

• `template`: string set to: 'divisionTemplate.Rmd', name of the template report

• `templatePackage`: string set to: 'clinDataReview', package from which the template should be extracted

• `reportTitle`: string, header title

• `reportTitleLevel`: (optional) integer, header level, 1 by default (1: 'chapter', '2': 'section', '3': 'subsection', ...)

• `parallel`: (optional) boolean, does this chapter use parallel execution? If the entire report is run in parallel, this ensures that this specific chapter is created outside of the report parallelization.

• `split_by`: (optional) integer of length: minimum 0, maximum 7 or string among: 'none', 'chapter', 'section', split the chapter at the specified level: 1 (for 'chapter'), 2 (for 'section') until 7. This overwrites the 'split_by' parameter defined in the output format of the report.

• `content`: (optional) string, any content that should be included after the title
Clinical data template to create a listing: listingTemplate

This report displays a listing of the variables and data of interest, displayed in an interactive table. This table can contain comparison with a previous batch ('comparisonTable' parameters).

The following parameters are available:

- **template**: string set to: 'listingTemplate.Rmd', name of the template report
- **templatePackage**: string set to: 'clinDataReview', package from which the template should be extracted
- **reportTitle**: string, header title
- **reportTitleLevel**: (optional) integer, header level, 1 by default (1: 'chapter', 2: 'section', 3: 'subsection', ...)
- **parallel**: (optional) boolean, does this chapter use parallel execution? If the entire report is run in parallel, this ensures that this specific chapter is created outside of the report parallelization.
- **split_by**: (optional) integer of length: minimum 0, maximum 7 or string among: 'none', 'chapter', 'section', split the chapter at the specified level: 1 (for 'chapter'), 2 (for 'section') until 7. This overwrites the 'split_by' parameter defined in the output format of the report.
- **dataFileName**: string or array, name of the data file(s) of interest. If multiple files are specified, the data are combined by rows ('row bind'), with a column: 'DATASET' containing the name of the file (in upper case and without extension) the data originate from.
- **dataProcessing**: (optional) array, data processing parameters, passed to processData
- **dataTotalFileName**: (optional) string, filename of the total dataset
- **dataTotalProcessing**: (optional) array, data processing parameters for 'dataTotalFileName', passed to processData
- **tableParams**: (optional) object, parameters to create the table, passed to tableClinData
- **comparisonTableType**: (optional) string among: 'none', 'newData-diff-interactive', 'table-comparison-interactive', output type of the comparison table
- **comparisonTableParams**: (optional) object, parameters for the comparison table, passed to compareTables
- **loopingVar**: (optional) array or string, data variable(s) to loop over. Each group of the variable(s) is displayed in a separated section of the report.
- **loopingNMax**: (optional) integer, maximum number of elements of loopingVar to include in the report
- **listingDocx**: (optional) boolean, export listing to Word

Clinical data template for the creation of patient profiles: patientProfilesTemplate

This report creates the specified patient profiles (with the patientProfilesVis package) by subject, and export them to a specified directory.

The following parameters are available:

- **template**: string set to: 'patientProfilesTemplate.Rmd', name of the template report
- **templatePackage**: string set to: 'clinDataReview', package from which the template should be extracted
• reportTitle: string, header title
• reportTitleLevel: (optional) integer, header level, 1 by default (1: 'chapter', 2: 'section', 3: subsection, ...)
• parallel: (optional) boolean, does this chapter use parallel execution? If the entire report is run in parallel, this ensures that this specific chapter is created outside of the report parallelization.
• split_by: (optional) integer of length: minimum 0, maximum 7 or string among: 'none', 'chapter', 'section', split the chapter at the specified level: 1 (for 'chapter'), 2 (for 'section') until 7. This overwrites the 'split_by' parameter defined in the output format of the report.
• createPatientProfiles: (optional) boolean, Should the patient profiles be created or only loaded from a previous execution?
• patientProfilesGeneralParams: (optional) object, set of parameters used for all patient profiles modules. These parameters are passed to all subjectProfile[]Plot functions.
• patientProfilesParams: array of object(s)
The following parameters are available:
  – typePlot: string among: 'text', 'line', 'interval', 'event', plot type, used to get the appropriate plot module function:
    * 'text': subjectProfileTextPlot
    * 'line': subjectProfileLinePlot
    * 'interval': subjectProfileIntervalPlot
    * 'event': subjectProfileEventPlot
  – dataFileName: string, name of the data file of interest
  – dataProcessing: (optional) array, data processing parameters, passed to processData
  – plotParams: object, parameters for the plotting function. Parameters depending on the dataset of interest can be specified as:
    [parameterName]: !r-lazy [dataI]
    , parameters for each patient profile module
• patientProfilesCreateReportParams: (optional) object, parameters for the creation of the patient profile report(s), passed to createSubjectProfileReport
• tableParams: (optional) object, parameters specifying a table containing data of interest and links to created patient profiles
• startup: (optional) array or string, R commands that should be run at the start of the report

Clinical data generic template for visualization: plotTemplate

This report visualizes input data with a function of the clinical data review package. The data can be compared to the data of a previous batch, in the table associated to the plot ('comparisonTable' parameters). Summary statistics can be computed optionally and included in the plot (see 'tableParams' parameter).
The following parameters are available:

• template: string set to: 'plotTemplate.Rmd', name of the template report
• templatePackage: string set to: 'clinDataReview', package from which the template should be extracted
Clinical data template for visualization of summarized data : summaryPlotTemplate

This report summarizes the data of interest (with the inTextSummaryTable package) and visualize it with any clinical data review plot function.

The following parameters are available:

- template: string set to: 'summaryPlotTemplate.Rmd', name of the template report
- templatePackage: string set to: 'clinDataReview', package from which the template should be extracted
- reportTitle: string, header title
• reportTitleLevel: (optional) integer, header level, 1 by default (1: 'chapter', 2: 'section', 3: 'subsection', ...)
• parallel: (optional) boolean, does this chapter use parallel execution? If the entire report is run in parallel, this ensures that this specific chapter is created outside of the report parallelization.
• split_by: (optional) integer of length: minimum 0, maximum 7 or string among: 'none', 'chapter', 'section', split the chapter at the specified level: 1 (for 'chapter'), 2 (for 'section') until 7. This overwrites the 'split_by' parameter defined in the output format of the report.
• dataFileName: string or array, name of the data file(s) of interest. If multiple files are specified, the data are combined by rows ('row bind'), with a column: 'DATASET' containing the name of the file (in upper case and without extension) the data originate from.
• dataProcessing: (optional) array, data processing parameters, passed to processData
• dataTotalFileName: (optional) string, filename of the total dataset
• dataTotalProcessing: (optional) array, data processing parameters for 'dataTotalFileName', passed to processData
• tableParams: object, parameters to summarize the data in a table, passed to computeSummaryStatisticsTable
• tableProcessing: (optional) array, data processing parameters for the summary table, passed to processData
• plotFunction: string among: 'timeProfileIntervalPlot', 'scatterplotClinData', 'sunburstClinData', 'treeMapClinData', 'barplotClinData', 'errorbarClinData', plotting function to visualize summary data
• plotParams: object, parameters for the plotting function. Parameters depending on the dataset of interest can be specified as:
  [parameterName]: !r-lazy [summaryTableI]
• loopingVar: (optional) array or string, data variable(s) to loop over. Each group of the variable(s) is displayed in a separated section of the report.
• loopingNMax: (optional) integer, maximum number of elements of loopingVar to include in the report
• loopingTotal: (optional) boolean, should the total also be computed by loopingVar (TRUE by default)?
• startup: (optional) array or string, R commands that should be run at the start of the report

Clinical data template for a summary table of the data: summaryTableTemplate

This report summarizes the data of interest (with the inTextSummaryTable package). This table is displayed with an interactive table in the report, and exported to a docx file. This table can be compared to a summary table of a previous batch ('comparisonTable' parameters).

The following parameters are available:
• template: string set to: 'summaryTableTemplate.Rmd', name of the template report
• templatePackage: string set to: 'clinDataReview', package from which the template should be extracted
• reportTitle: string, header title
• reportTitleLevel: (optional) integer, header level, 1 by default (1: 'chapter', '2': 'section', '3': subsection, ...)

• parallel: (optional) boolean, does this chapter use parallel execution? If the entire report is run in parallel, this ensures that this specific chapter is created outside of the report parallelization.

• split_by: (optional) integer of length: minimum 0, maximum 7 or string among: 'none', 'chapter', 'section', split the chapter at the specified level: 1 (for 'chapter'), 2 (for 'section') until 7. This overwrites the 'split_by' parameter defined in the output format of the report.

• dataFileName: string or array, name of the data file(s) of interest. If multiple files are specified, the data are combined by rows ('row bind'), with a column: 'DATASET' containing the name of the file (in upper case and without extension) the data originate from.

• dataProcessing: (optional) array, data processing parameters, passed to processData

• dataTotalFileName: (optional) string, filename of the total dataset

• dataTotalProcessing: (optional) array, data processing parameters for 'dataTotalFileName', passed to processData

• tableParams: object, parameters to summarize the data in a table, passed to computeSummaryStatisticsTable. Parameters depending on the dataset of interest can be specified as: [parameterName]: !r-lazy [dataI].

• tableParamsDocx: (optional) object, parameters to format the table to the docx format, passed to exportSummaryStatisticsTable

• tableParamsDT: (optional) object, parameters to format the table to the DT interactive table included in the report, passed to exportSummaryStatisticsTable

• comparisonTableType: (optional) string among: 'none', 'table-comparison-interactive', 'newData-diff', 'table-combine', output type of the comparison table:
  - 'table-comparison-interactive': the summary statistics are computed for the old and new batch separately, and the statistics are compared
  - 'newData-diff': the differences at a record level between the new and old batch are first identified, and then summarized
  - 'table-combine': the summary tables are computed for the old and new batch and displayed next to each other in a combined table

• comparisonTableParams: (optional) object, parameters for the comparison table, passed to compareTables. By default, statistics variables are compared across row and column elements.

• loopingVar: (optional) array or string, data variable(s) to loop over. Each group of the variable(s) is displayed in a separated section of the report.

• loopingNMax: (optional) integer, maximum number of elements of loopingVar to include in the report

• loopingTotal: (optional) boolean, should the total also be computed by loopingVar (TRUE by default)?
collapseHtmlContent  
*Function to create collapsible HTML content*

**Description**
Please note that the button is of class: `hideshow`, defined in the `input.hideshow.js` js file included in the package.

**Usage**
collapseHtmlContent(input, title = "Click to show or hide")

**Arguments**
- **input**: Object to be collapse, e.g. datatable.
- **title**: String with button title.

**Value**
- **tag object**

**Author(s)**
Laure Cougnaud

------------------------

combineButtonsAndPlot  
*Combine select box(es) and the plot*

**Description**
Combine select box(es) and the plot

**Usage**
combineButtonsAndPlot(x)

**Arguments**
- **x**: Object of class clinDataReview

**Value**
- **x object**:
  - with the plot element containing a combination of the buttons and the plot
  - without the buttons element
**convertMdToHtml**  
*Convert the Md file for a specific chapter to html*

**Description**

Convert the Md file for a specific chapter to html

**Usage**

```r
convertMdToHtml(
  mdFile,
  configFile = NULL,
  indexPath = "index.Rmd",
  intermediateDir = "/interim",
  outputDir = "/report",
  setTitle = TRUE,
  verbose = TRUE,
  ...
)
```

**Arguments**

- **mdFile**: String with path of the Markdown file
- **configFile**: String with filename of the config file of interest in YAML format.
- **indexPath**: String with path to the index file, by default 'index.Rmd' in inputDir.
- **intermediateDir**: String with intermediate directory ('interim' by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.
- **outputDir**: String with output directory, ('report' by default).
- **setTitle**: Logical (TRUE by default), should the title be set to the document? If so, the pandoc metadata option: 'pagetitle' is set to: base file name of mdFile.
- **verbose**: Logical, if TRUE (FALSE by default) progress messages are printed during the report execution.
- **...**: Arguments passed to `renderFile`

**Value**

No returned value, the files in the intermediateDir are converted to HTML

**Author(s)**

Laure Cougnaud
**convertReportToAspx**  
*Convert report format from .html to .aspx*

**Description**

Report files generated as output of `render_clinDataReviewReport` are converted from .html to .aspx format by changing extensions and cross-links of all files within the directory `reportDir`. This allows for deployment on SharePoint.

**Usage**

```
convertReportToAspx(reportDir = ".")
```

**Arguments**

- `reportDir`  
  String for the path to the directory where the clinical data reports are stored, defaults to current directory

**Value**

- no return value, files in directory are modified

---

**countNLines**  
*Count number of lines in a vector*

**Description**

Count number of lines in a vector

**Usage**

```
countNLines(x)
```

**Arguments**

- `x`  
  Character vector.

**Value**

- Integer vector of length `x` with number of lines

**Author(s)**

Laure Cougnaud

**Examples**

```
clinDataReview:::countNLines(x = c("A\nB", "blabla", "This\nis\na\nsentence."))
```
createClinDataReviewReportSkeleton

*Create the skeleton of a report*

**Description**

Creates the skeleton of a report to start running the analyses.

**Usage**

```r
createClinDataReviewReportSkeleton(dir = ".")
```

**Arguments**

- `dir`  
  String with the path of the directory where the skeleton should be created. The current working directory is used as default.

**Details**

This function is meant to get familiar with the use of the package and the necessary files to create a report.

It will create a ready-to-use report with example data from the clinUtils package. After getting use to the file structure, the user can substitute the example data with custom data sets and add specific configuration files.

**Value**

The files to run a report are written in the specified directory. To run the report, the user can call the `render_clinDataReviewReport`.

createExampleMetadata

*Create an example metadata file*

**Description**

Create an example of metadata file for the createClinDataReviewReportSkeleton.

**Usage**

```r
createExampleMetadata(dir)
```

**Arguments**

- `dir`  
  String, path to the directory.

**Value**

Nothing, the example metadata file is created in the specified directory.
createMainConfigSkeleton

*Create the config file for the skeleton*

**Description**

This function creates the main config file for the `createClinDataReviewReportSkeleton` with the directory where the data are stored.

**Usage**

```r
createMainConfigSkeleton(dir, dirData)
```

**Arguments**

- **dir** String, path to the directory.
- **dirData** String, path to the directory of the data.

**Value**

No return value, a file `_config.yml` is created in the specified directory.

createOutputYaml

*Create a output YAML file*

**Description**

This file containing the contents of the output field of the YAML header of a Markdown file. It can be passed to the `output_yaml` parameter of the `render` function.

**Usage**

```r
createOutputYaml(indexPath, outputDir)
```

**Arguments**

- **indexPath** String with path to the index file, by default 'index.Rmd' in `inputDir`.
- **outputDir** String with output directory, ('report' by default).

**Value**

String with file to the `_output.yml` file in a temporary folder.
createPatientProfileVar

Create link to patient profile

Description

Create a link to a patient profile directory (where the patient profile files are saved) by adding an extra column with the link in the data. The path to the patient profile is built as: \[patientProfilePath\]/subjectProfile-[subjectID].pdf, where '/' are replaced with '.' in the subject identifier (subjectVar).

Usage

\[
\text{createPatientProfileVar}(\text{data, patientProfilePath, subjectVar = "USUBJID", checkExist = TRUE})
\]

Arguments

- **data**: a data.frame
- **patientProfilePath**: string indicating the directory where the patient profiles are stored.
- **subjectVar**: string indicating which column in the data represents the unique subject identifier, "USUBJID" by default.
- **checkExist**: Logical, if TRUE (by default) the patientProfilePath is checked for existence, and an error is returned if this directory doesn’t exist.

Value

A data.frame with two extra columns: patientProfilePath and patientProfileLink with the path to the patient profile and an hyperlink to it, respectively.

Author(s)

Michela Pasetto

Examples

# Typical CDISC dataset contains universal subject ID (USUBJID)
data <- data.frame(USUBJID = c("subj1", "subj2", "subj3"))
dataWithPatientProfileVar <- createPatientProfileVar(
data = data,
patientProfilePath = "pathProfiles",
checkExist = FALSE
)
# path and HTML link are included in the output dataset
head(dataWithPatientProfileVar[, c("USUBJID", "patientProfilePath", "patientProfileLink")])
createRedirectPage

Create a redirect page

Description
Create an html page that redirects to the "1-introduction.html" page of the clinical data report available in a directory. See output from `render_clinDataReviewReport`.

Usage
```r
createRedirectPage(redirectPage = "report.html", dir = "report_dependencies")
```

Arguments
- `redirectPage`: String with the path of the html file that redirects to the "1-introduction.html" page of the report.
- `dir`: String for the path where the "1-introduction.html" is stored.

Value
The html file is created.

createTemplateDoc

Create documentation for clinical data template reports available in the 'template' folder of the package.

Description
If a JSON schema file available, the information relative to the template is extracted from this file with the function `JSONSchToRd`.

Usage
```r
createTemplateDoc(
  templatePath = system.file("template", package = "clinDataReview")
)
```

Arguments
- `templatePath`: String with path where the template Rmd reports and associated JSON schema files are stored, by default path of the installed version of the package. This parameter is only for expert use of the package.

Value
Character vector with Rd code containing description for all template documents.
Author(s)
Laure Cougnaud

References
JSON schema specification

errorbarClinData
Interactive plot of confidence interval/error interval of clinical data.

Description
This plot is designed to display summary statistics of a continuous variable with (confidence) intervals.
The intervals are either displayed:
• vertically if yErrorVar is specified
• horizontally if xErrorVar is specified

Error bars can visualized by group, via the color variable parameter.
Different symbols are set for each central point of the error bar via the shape variable parameter.

Usage
errorbarClinData(
data,
xVar,
xLab = getLabelVar(xVar, labelVars = labelVars),
xLabVar = NULL,
yVar,
yLab = getLabelVar(yVar, labelVars = labelVars),
yLabVar = NULL,
yErrorVar = NULL,
yErrorLab = getLabelVar(yErrorVar, labelVars = labelVars),
xErrorVar = NULL,
xErrorLab = getLabelVar(xErrorVar, labelVars = labelVars),
xLabVars = NULL,
xAxisLab = paste(c(xLab, xErrorLab), collapse = " and "),
yAxisLab = paste(c(yLab, yErrorLab), collapse = " and "),
colorVar = NULL,
colorLab = getLabelVar(colorVar, labelVars = labelVars),
colorPalette = NULL,
shapeVar = NULL,
shapeLab = getLabelVar(shapeVar, labelVars = labelVars),
shapePalette = NULL,
size = 6,
title = paste(c(paste(yAxisLab, " vs ", xAxisLab), titleExtra), collapse = "<br>")},
titleExtra = NULL,
subtitle = NULL,
caption = NULL,
labelVars = NULL,
mode = "markers",
legendPosition = "bottom",
width = NULL,
height = NULL,
pathVar = NULL,
pathLab = getLabelVar(pathVar, labelVars = labelVars),
hoverVars,
hoverLab,
id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
selectVars = NULL,
selectLab = getLabelVar(selectVars, labelVars = labelVars),
table = FALSE,
tableVars,
tableLab,
tableButton = TRUE,
tablePars = list(),
watermark = NULL,
verbose = FALSE
)

Arguments

data  Data.frame with data.
xVar  String with column of data containing x-variable.
xLab  String with label for xVar.
xLabVar (optional) Character vector with column(s) of data containing variable(s) to display in the label of the x-axis.
yVar  String with column of data containing y-variable.
yLab  String with label for xVar.
yLabVar (optional) Character vector with column(s) of data containing variable(s) to display in the label of the y-axis.
xErrorVar, yErrorVar  String with variable of data containing the width of the interval (from the center of the interval) for horizontal or vertical intervals.
xErrorLab, yErrorLab  String with labels for xErrorVar/yErrorVar variables.
xLabVars (vertical error bars) Character vector with variable(s) to be displayed as the labels of the ticks in the x-axis.
  By default, xVar is displayed.
  In case the variable(s) contain different elements by xVar, they are combined and displayed below each other.
xAxisLab, yAxisLab  Label for the x/y-axis.
colorVar (optional) String with color variable.
colorLab String with label for colorVar.
colorPalette (optional) Named character vector with color palette. If not specified, the viridis color palette is used. See clinColors.
shapeVar (optional) String with shape variable.
shapeLab String with label for shapeVar.
shapePalette (optional) Named character vector with shape palette, clinShapes by default.
size Integer with size of markers in pixels, 6 by default.
title String with title for the plot.
titleExtra String with extra title for the plot (appended after title).
subtitle String with subtitle.
The subtitle is included at the top left of the plot, below the title.
caption String with caption.
The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.
labelVars Named character vector containing variable labels.
mode String with the mode of the plot, 'markers' by default, so only data points are displayed.
This can also be set to 'lines' to include a line connecting the center of the error bars instead; or 'lines+markers' to include both a marker and a line.
See mode attribute for plotly scatter.
legendPosition String with position of the legend, among: 'top'/'left'/'bottom'/'right', 'bottom' by default.
width Numeric, width of the plot in pixels, 800 by default.
height Numeric, height of the plot in pixels, 500 by default.
pathVar String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

$$<a href="./path-to-report">label</a>$$

If multiple, they should be separated by ' , '.
The report(s) will be:

- compressed to a zip file and downloaded if the user clicks on the 'p' (a.k.a 'profile') key when hovering on a point of the plot
- included in a collapsible row, and clickable with hyperlinks in the table

pathLab String with label for pathVar, included in the collapsible row in the table.
hoverVars Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.
hoverLab Named character vector with labels for hoverVars.
id String with general id for the plot:
### errorbarClinData

- 'id' is used as group for the **SharedData**
- 'button:[id]' is used as button ID if table is TRUE

If not specified, a random id, as 'plotClinData[X]' is used.

**selectVars** (optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).

**selectLab** (Named) character vector with label for selectVars.

**table** Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)

**tableVars** Character vector with variables to be included in the table.

**tableLab** Named character vector with labels for each tableVars.

**tableButton** Logical, if TRUE (by default) the table is included within an HTML button.

**tablePars** List with parameters passed to the getClinDT function.

**watermark** (optional) String with path to a file containing a watermark.

**verbose** Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.

**Value**

Either:

- if a table is requested: a clinDataReview object, a.k.a a list with the 'plot' (**plotly** object) and 'table' (**datatable** object)
- otherwise: a **plotly** object

**Author(s)**

Laure Cougnaud

**See Also**

Other visualizations of summary statistics for clinical data: barplotClinData(), boxplotClinData(), plotCountClinData(), sunburstClinData(), treemapClinData()

**Examples**

```r
library(clinUtils)

data(dataADaMCDISCP01)
labelVars <- attr(dataADaMCDISCP01, "labelVars")

## Summary plot with vertical error bars

dataVSDIABP <- subset(dataADaMCDISCP01$ADVS,
  PARAMCD == "DIABP" & ANL01FL == "Y" &
  AVISIT %in% c("Baseline", "Week 2", "Week 4", "Week 6", "Week 8")
```
# compute summary statistics by visit
if (requireNamespace("inTextSummaryTable", quietly = TRUE)) {

  summaryTableVSDIABP <- inTextSummaryTable::computeSummaryStatisticsTable(
    data = dataVSDIABP,
    rowVar = c("AVISIT", "ATPT"),
    var = "AVAL",
    stats = inTextSummaryTable::getStats(c("n", "Mean", "SE")),
    labelVars = labelVars
  )
  dataPlot <- subset(summaryTableVSDIABP, !isTotal)

  errorbarClinData(
    data = dataPlot,
    xVar = "AVISIT",
    colorVar = "ATPT",
    yVar = "statMean", yErrorVar = "statSE",
    yLab = "Mean", yErrorLab = "Standard Error",
    # include lines connecting the error bars
    mode = "markers+lines",
    labelVars = labelVars
  )

  # add number of subjects in labels
  dataPlot$nSubj <- with(dataPlot, paste0("N=", n))
  errorbarClinData(
    data = dataPlot,
    xVar = "AVISIT",
    xLabVars = c("AVISIT", "nSubj"),
    colorVar = "ATPT",
    yVar = "statMean", yLab = "Mean",
    yErrorVar = "statSE", yErrorLab = "Standard error",
    mode = "markers+lines",
    title = paste("Diastolic Blood Pressure summary profile by actual visit",
                  "and analysis timepoint"),
    labelVars = labelVars
  )

  ## Add a selection box
  if(interactive()){
    summaryTable <- inTextSummaryTable::computeSummaryStatisticsTable(
      data = subset(dataADaMCDISCP01$ADVS,
                    ANL01FL == "Y" &
                   AVISIT %in% c("Baseline", "Week 2", "Week 4", "Week 6", "Week 8")),
      rowVar = c("PARAM", "AVISIT", "ATPT"),
      var = "AVAL",
      stats = inTextSummaryTable::getStats(c("Mean", "SE")),
      labelVars = labelVars
    )
  }
}
dataPlot <- subset(summaryTable, !isTotal)

errorbarClinData(
  data = dataPlot,
  xVar = "AVISIT",
  colorVar = "ATPT",
  yVar = "statMean", yLab = "Mean",
  yErrorVar = "statSE", yErrorLab = "Standard error",
  mode = "markers+lines",
  title = paste("Lab parameters summary profile by actual visit",
                "and analysis timepoint"),
  labelVars = labelVars,
  selectVars = "PARAM"
)

## Summary plot with horizontal error bars
# Data of interest: ratio from baseline at week 16
dataLBW8 <- subset(dataADaMCDISCP01$ADLBC, grepl("Week 8", AVISIT))
# compute ratio from baseline
dataLBW8$R2BASE <- with(dataLBW8, AVAL/BASE)
dataLBW8 <- subset(dataLBW8, !is.na(R2BASE))
# Order actual treatments
dataLBW8$TRTA <- with(dataLBW8, reorder(TRTA, TRTAN))

# compute summary statistics of the ratio per baseline per parameter
summaryTableLBW8 <- inTextSummaryTable::computeSummaryStatisticsTable(
  data = dataLBW8,
  var = "R2BASE",
  rowVar = "PARAM",
  colVar = "TRTA",
  stats = inTextSummaryTable::getStats(x = dataLBW8$R2BASE, type = c("n", "Median", "SD"))
)
dataPlot <- subset(summaryTableLBW8, !isTotal)
# extract direction of ratio
dataPlot$dir <- factor(
  ifelse(dataPlot$statMedian >= 1, "Increase", "Decrease"),
  levels = c("Decrease", "Increase")
)
# compute relative ratio (percentage above 1)
dataPlot$statMedianRelative <- with(dataPlot, 
  ifelse(statMedian < 1, 1/statMedian, statMedian)
)
# order based on mean relative ratio across treatment arms
params <- names(sort(with(dataPlot, tapply(statMedianRelative, PARAM, mean))))
dataPlot$PARAM <- factor(dataPlot$PARAM, levels = params)
errorbarClinData(
  data = dataPlot,
  xVar = "statMedianRelative", xErrorVar = "statSD",
  xLab = "Median", xErrorLab = "Standard deviation",
  xAxisLab = "Relative ratio from baseline (Median +- SD)",
  yVar = "PARAM",
  colorVar = "TRTA",
)
exportSessionInfoToMd

Combine all session informations across all clinical data reports and export them into a dedicated Markdown document

Description

Combine all session informations across all clinical data reports and export them into a dedicated Markdown document

Usage

exportSessionInfoToMd(
  sessionInfos,
  intermediateDir = "interim",
  logFile = NULL,
  ...
)

Arguments

sessionInfos  List with sessionInfo objects
intermediateDir String with intermediate directory ('interim' by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.
logFile (optional) String with path to a log file, where output (also error/messages/warnings) should be stored. If specified, the entire output is re-directed to this file.
... Any parameters passed to renderFile, for expert use only.

Value

String with path to Markdown file containing the session information, NULL if no session information(s) are provided.

Author(s)

Laure Cougnaud
filterData

Filter a dataset based on specified filters.

Description
A dataset can be filtered:

• on a specific value of interest
• on a function of a variable (valueFct parameter), e.g. maximum of the variable
• to retain only non missing values of a variable (keepNA set to FALSE)
• by groups (varsBy parameter)

Note that by default, missing values in the filtering variable are retained (which differs from the default behaviour in R). To filter missing records, please set the keepNA parameter to FALSE.

Usage
filterData(
  data,
  filters,
  keepNA = TRUE,
  returnAll = FALSE,
  verbose = FALSE,
  labelVars = NULL,
  labelData = "data"
)

Arguments
data
  Data.frame with data.
filters
  Unique filter or list of filters.

Each filter is a list containing:

• 'var': String with variable from data to filter on.
• 'value': (optional) Character vector with values from var to consider/keep.
• 'valueFct': (optional) Function (or string with this function) to be applied on var to extract value to consider.
  For example, valueFct = max will extract the records with the maximum value of the variable.
• 'op': (optional) String with operator used to retain records from value. If not specified, the inclusion operator: '%in%' is considered, so records with var in value are retained.
• 'rev': (optional) Logical, if TRUE (FALSE by default), filtering condition based on value/valueFct is reversed.
• 'keepNA': (optional) Logical, if TRUE (by default), missing values in var are retained.
  If not specified, keepNA general parameter is used.
filterData

- 'varsBy': (optional) Character vector with variables in data containing groups to filter by.
- 'postFct': (optional) Function (or string with this function) with post-processing applied on the results of the filtering criteria (TRUE/FALSE for each record). This function should return TRUE/FALSE (for each record or for all considered records).
  For example, `postFct = any, varsBy = "group"` retains all groups which contain at least one record that fulfills the criteria.
- 'varNew': (optional) String with name of a new variable containing the results of the filtering criteria (as TRUE/FALSE).
- 'labelNew': (optional) String with label for the varNew variable.

If a list of filters is specified, the different filters are independently executed on the entire dataset to identify the records to retain for each filtering condition.

The resulting selections are combined with a Logic operator ('&' by default, i.e. 'AND' condition). A custom logic operator can be specified between the lists describing the filter, for example:

```
list(list(var = "SEX", value = "F"), ",", list(var = "COUNTRY", value = "DEU"))
```

keepNA Logical, if TRUE (by default) missing values in var are retained. If set to FALSE, missing values are ignored for all filters. The specification within filters prevails on this parameter.

returnAll Logical:
- if FALSE (by default): the data for only the filtered records is returned.
- if TRUE: the full data is returned. Records are flagged based on the filters condition, in a new column: varNew (if specified), or 'keep' otherwise; containing TRUE if the record fulfill all conditions, FALSE otherwise.

verbose Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.

Value

If returnAll
- is FALSE: data filtered with the specified filters
- is TRUE: data with the additional column: keep or varNew (if specified), containing TRUE for records which fulfill the specified condition(s) and FALSE otherwise.

The output contains the additional attribute: msg which contains a message describing the filtered records.

Author(s)

Laure Cougnaud
Examples

```r
library(clinUtils)

data(dataADaMCDISCP01)
labelVars <- attr(dataADaMCDISCP01, "labelVars")

dataDM <- dataADaMCDISCP01$ADSL

## single filter

# filter with inclusion criteria:
filterData(
data = dataDM,
filters = list(var = "SEX", value = "M"),
# optional
labelVars = labelVars, verbose = TRUE
)

# filter with non-inclusion criteria
filterData(
data = dataDM,
filters = list(var = "SEX", value = "M", rev = TRUE),
# optional
labelVars = labelVars, verbose = TRUE
)

# filter based on inequality operator
filterData(
data = dataDM,
filters = list(var = "AGE", value = 75, op = "<="),
# optional
labelVars = labelVars, verbose = TRUE
)

# missing values are retained by default!
dataDMNA <- dataDM
dataDMNA[1:2, "AGE"] <- NA
filterData(
data = dataDMNA,
filters = list(var = "AGE", value = 75, op = "<="),
# optional
labelVars = labelVars, verbose = TRUE
)

# filter missing values on variable
filterData(
data = dataDMNA,
filters = list(var = "AGE", value = 75, op = "<="), keepNA = FALSE,
# optional
labelVars = labelVars, verbose = TRUE
)
```
# retain only missing values
filterData(
data = dataDMNA,
filters = list(var = "AGE", value = NA, keepNA = TRUE),
# optional
labelVars = labelVars, verbose = TRUE
)

# filter missing values
filterData(
data = dataDMNA,
filters = list(var = "AGE", keepNA = FALSE),
# optional
labelVars = labelVars, verbose = TRUE
)

## multiple filters
# by default the records fulfilling all conditions are retained ('AND')
filterData(
data = dataDM,
filters = list(
list(var = "AGE", value = 75, op = "<="),
list(var = "SEX", value = "M")
),
# optional
labelVars = labelVars, verbose = TRUE
)

# custom operator:
filterData(
data = dataDM,
filters = list(
list(var = "AGE", value = 75, op = "<="),
"|",
list(var = "SEX", value = "M")
),
# optional
labelVars = labelVars, verbose = TRUE
)

# filter by group
# only retain adverse event records with worst-case severity

dataAE <- dataADaMCDISCP01$AE
dataAE$AESEV <- factor(dataAE$AESEV, levels = c("MILD", "MODERATE", "SEVERE"))
dataAE$AESEVN <- as.numeric(dataAE$AESEV)
nrow(dataAE)
dataAEWorst <- filterData(
data = dataAE,
filters = list(
var = "AESEVN",
)
filterDataSingle

Filter data for a single filter

Description

Filter data for a single filter

Usage

filterDataSingle(data, filters,
Arguments

data Data.frame with data.
filters Unique filter or list of filters.
keepNA Logical, if TRUE (by default) missing values in var are retained. If set to FALSE, missing values are ignored for all filters. The specification within filters prevails on this parameter.
returnAll Logical:
  • if FALSE (by default): the data for only the filtered records is returned.
  • if TRUE: the full data is returned. Records are flagged based on the filters condition, in a new column: varNew (if specified), or 'keep' otherwise; containing TRUE if the record fulfill all conditions, FALSE otherwise
labelVars Named character vector containing variable labels.
labelData (optional) String with label for input data, that will be included in progress messages.

Value

Updated data with attributes:

  • 'labelVars': input labelVars with any new variables if labelNew is specified.
  • 'msg': message describing the filtering process
  • 'warn': warning describing the filtering process

Author(s)

Laure Cougnaud

Description

This function is only useful if some parameters should be lazy-evaluated in the report. These parameters should have the class: r-lazy. A typical use case is a parameter that consists of a R expression depending on objects created in a template report (typically data). Parameters are searched in the environment in which this function is called from.
formatDataForPlotClinData

Usage

forceParams(params)

Arguments

params List of parameters as obtained via the `getParamsFromConfig` function.

Value

Input parameter list, with object(s) of class `r-lazy` evaluated.

Author(s)

Laure Cougnaud

See Also

`getParamsFromConfig`

Other clinical data reporting: `checkReportTitles()`, `getMdHeader()`, `getParamsFromConfig()`, `gitbook_clinDataReview_report()`, `html_clinDataReview_report()`, `knitPrintClinDataReview()`, `postProcessReport()`, `render_clinDataReviewReport()

Examples

data <- mtcars
params <- list(label = "Cars dataset", nrow = structure("nrow(data)", class = "r-lazy"))
str(params)
str(forceParams(params))

formatDataForPlotClinData

Format data for interactive plot for clinical data

Description

Format data for interactive plot for clinical data

Usage

formatDataForPlotClinData(
  data,
  hoverVars = NULL,
  hoverLab = getLabelVar(hoverVars, labelVars = labelVars),
  hoverByVar = NULL,
  keyVar = NULL,
  id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
  labelVars = NULL
)

**Arguments**

- **data**: Data.frame with data.
- **hoverVars**: Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.
- **hoverLab**: Named character vector with labels for hoverVars.
- **hoverByVar**: Character vector with variables identifying unique elements in the plot, usually x, y, facet variables. These variables are used to identify records with the same position in the plot, their information are combined in the hover.
- **keyVar**: String with unique key variable, identifying unique group for which the link between the table and the plot should be done.
- **id**: String with general id for the plot:
  - 'id' is used as group for the `SharedData`
  - 'button:[id]' is used as button ID if table is TRUE
  If not specified, a random id, as 'plotClinData[X]' is used.
- **labelVars**: Named character vector containing variable labels.

**Value**

A `SharedData` object containing the data, with an extra column: 'hover' with the combined info from hoverVars, and the key defined as keyVar and group as id.

**Author(s)**

Laure Cougnaud

---

**formatHoverText**

Format hover text for use in plotly interactive plots. The labels are wrapped to multiple lines if exceed the width of the plotly hover box, e.g. in case labels for points with same x/y coordinates overlap, and corresponding labels are truncated.

**Description**

Format hover text for use in plotly interactive plots. The labels are wrapped to multiple lines if exceed the width of the plotly hover box, e.g. in case labels for points with same x/y coordinates overlap, and corresponding labels are truncated.

**Usage**

formatHoverText(x, label, width = 50)
Arguments

- **x**: Vector with hover text information.
- **label**: Label for the variable
- **width**: Integer, number of characters at which the hover text should be cut at to multiple lines.

Value

String with formatted hover label.

Author(s)

Laure Cougnaud

---

**formatPathDateInfoMetadata**

*Format the info on paths from metadata*

Description

Format the info on paths from metadata

Usage

`formatPathDateInfoMetadata(summaryInfo, namesInfo)`

Arguments

- **summaryInfo**: matrix, see output from `getMetadata`.
- **namesInfo**: Named vector to rename the final output.

Value

A kable object, to be printed.
formatPlotlyClinData  
Format interactive plot, with possibility to download patient profiles on a click event.

Description

Format interactive plot, with possibility to download patient profiles on a click event.

Usage

formatPlotlyClinData(
  pl,  
data,  
idVar = "USUBJID",  
pathVar = NULL,  
pathDownload = TRUE,  
idFromDataPlot = FALSE,  
idVarPlot = "key",  
labelVarPlot = NULL,  
highlightOn = "plotly_click",  
highlightOff = "plotly_doubleclick",  
id = paste0("plotClinData", sample.int(n = 1000, size = 1)),  
selectVars = NULL,  
selectLab = getLabelVar(selectVars, labelVars = labelVars),  
keyVar = NULL,  
keyHighlightBox = FALSE,  
labelVars = NULL,  
verbose = FALSE  
)

Arguments

pl  
plotly object.

data  
Data.frame with data.

idVar  
String with variable of data containing plot element.

pathVar  
String with variable of data containing path to a subject-specific report (e.g. patient profiles).

pathDownload  
Logical, if TRUE (by default) the subject-specific report(s) are downloaded in a zip compressed file. If FALSE (only available if unique report per idVarPlot), each report is opened in a new window.

idFromDataPlot  
Logical, if TRUE (by default) idVarPlot is extracted from the data of the plot output object (e.g. if this plot was created from ggplotly), otherwise directly from the plot object (if the plot was created from plot_ly directly).

idVarPlot  
String with variable in the plotly output containing IDs.
formatToHierarchicalData

LabelVarPlot String with plotly variable used to extract label to build the file name of the zip compressed file containing patient report. If not specified, the label are extracted based on the idVarPlot of the selected plot element.

highlightOn String with event to turn on the selection (on parameter of highlight), 'plotly_click' by default.

highlightOff String with event to turn off the selection (off parameter of highlight), 'plotly_doubleclick' by default.

id String with general id for the plot:
  - 'id' is used as group for the SharedData
  - 'button:[id]' is used as button ID if table is TRUE
If not specified, a random id, as 'plotClinData[X]' is used.

selectVars (optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).

selectLab (Named) character vector with label for selectVars.

keyVar String with unique key variable, identifying unique group for which the link between the table and the plot should be done.

keyHighlightBox Logical, if TRUE (FALSE by default) a selectize box is included to highlight selected element(s) of the key variable.

labelVars Named character vector containing variable labels.

verbose Logical, if TRUE report progress messages during execution (included in the browser 'Console').

Value

Updated plotly object.

Author(s)

Laure Cougnaud

Format data to a hierarchical data, in the format as required by the plotly sunburst and treemap.

Description

Note that new variables are created for each variable of interest (the variables are not overwritten) to avoid issues with cases where the value in the child and parent variables are the same.

Usage

formatToHierarchicalData(data, vars, valueVar)
Arguments

data Data.frame with data.
vars Character vector with variables of data containing the groups. If multiple, they should be specified in hierarchical order (from parent to child node).
valueVar String with numeric variable of data containing the value to display.

Value

Updated data.frame with vars in hierarchical format, with extra attributes (in 'metadat'):

- 'varID': String with column of output containing ID of specific element.
  This is a combination from the specified vars, or 'Overall' for the grand total.
- 'varParent': String with column of output containing ID of the parent element
- 'varLabel': String with column of output containing the label to display.
  This is usually the name of the child element.

Author(s)

Laure Cougnaud

getAxisLab Get axis label

Description

Get axis label

Usage

getAxisLab(axisVar, axisLab, labVar = NULL, data, labelVars)

Arguments

axisVar String with column of data containing variable for the axis.
axisLab (optional) String with label for axisVar
labVar String with column of data containing variable(s) whose content should be displayed in the axis (with its labels).
data Data.frame with data.
lableVars Named character vector containing variable labels.

Value

String with label for the axis.
getAxisLabs

*Set different variables for the x-axis labels*

**Description**
Set different variables for the x-axis labels

**Usage**
getAxisLabs(data, var, labVars)

**Arguments**
data Data.frame with data.
var String with variable displayed in the axis.
labVars Character vector with variable(s) to be displayed as the labels of the ticks in the axis.

**Value**
Named character vector. The names are the position in the x-axis, the values are the new labels.

**Author(s)**
Laure Cougnaud

getAxisLimPlot

*Get axis limits for a ggplot plot from the input dataset.*

**Description**
Get axis limits for a ggplot plot from the input dataset.

**Usage**
getAxisLimPlot(
  data,
  xVar,
  yVar,
  xLim = NULL,
  yLim = NULL,
  facetPars = NULL,
  refLineData = NULL
)


getDataReferenceLines

Arguments

- **data**: Data.frame with data.
- **xVar**: String with column of data containing x-variable.
- **yVar**: String with column of data containing y-variable.
- **xLim, yLim**: Numeric vector of length 2 with limits for the x/y axes.
- **facetPars**: List with facetting parameters, passed to the facetting function. Variables should be specified as character or formula. For 'wrap' facetting (facetType is 'wrap'), if the layout is not specified via nrow/ncol, 2 columns are used by default.
- **refLineData**: Data used for the reference lines, as output of the `getDataReferenceLines` function.

Value

Data.frame with limits of the:

- x-axis: 'xmin'/xmax'
- y-axis: 'ymin'/ymax'

for each element of the facetting variable (if any).

---

**getDescription**

*Extract data for the reference lines*

**Description**

This function especially extracts the data if an aesthetic variable is specified in the reference line parameters.

**Usage**

`getDataReferenceLines(refLinePars, data, facetPars = NULL)`

**Arguments**

- **refLinePars** (optional): Nested list, with parameters for each reference line(s). Each sublist (a.k.a reference line) contains:
  - aesthetic value(s) or variable(s) for the lines (in this case column names of data) for reference lines. The line position is controlled by the aesthetics supported in `geom_vline`, `geom_hline` and `geom_abline`.
  - 'label': (optional) Logical specifying if the line should be annotated (FALSE to not annotate the line) or string with annotation label. By default, the value of the position of the horizontal/vertical line or the equation of the diagonal line is displayed.
- **data**: Data.frame with data.
- **facetPars**: List with facetting parameters, passed to the facetting function. Variables should be specified as character or formula. For 'wrap' facetting (facetType is 'wrap'), if the layout is not specified via nrow/ncol, 2 columns are used by default.
**getDimGgplot**

*Get plot dimensions*

**Value**

List of data for the lines

**Author(s)**

Laure Cougnaud

---

**getExtraDirs**

*Get extra directory(ies) required for the clinical data review report*

**Description**

By default, the 'figures', 'tables' and patient profiles folders (patientProfilePath parameter in the general config file, if specified) in the input directory are considered.

**Usage**

```r
getExtraDirs(inputDir = ".", configDir = file.path(inputDir, "config"))
```
Arguments

inputDir  String with input directory, working directory by default.
configDir String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general general 'config.yml'.

Value

Character vector with extra directories required by the report

Author(s)

Laure Cougnaud

---

getFacetVars  Get faceting variables from facet parameters.

Description

Get faceting variables from facet parameters.

Usage

getFacetVars(facetPars = list())

Arguments

facetPars List with faceting parameters, passed to the faceting function. Variables should be specified as character or formula. For 'wrap' faceting (facetType is 'wrap'), if the layout is not specified via nrow/ncol, 2 columns are used by default.

Value

Character vector with faceting variable

Author(s)

Laure Cougnaud
getFctCode

Get function code

Description
Get function code

Usage
getFctCode(fct)

Arguments
fct a R function

Value
String with function code

getFctTypeReferenceLines

Get the names of the ggplot function to use for the reference lines

Description
Get the names of the ggplot function to use for the reference lines

Usage
getFctTypeReferenceLines(refLinePars)

Arguments
refLinePars (optional) Nested list, with parameters for each reference line(s). Each sublist (a.k.a reference line) contains:
  • aesthetic value(s) or variable(s) for the lines (in this case column names of data) for reference lines. The line position is controlled by the aesthetics supported in geom_vline, geom_hline and geom_abline.
  • 'label': (optional) Logical specifying if the line should be annotated (FALSE to not annotate the line) or string with annotation label. By default, the value of the position of the horizontal/vertical line or the equation of the diagonal line is displayed.

Value
List of type of each reference lines, among: 'vline', 'hline' and 'abline'.
**getHTMLToc**

**Description**
Get HTML toc

**Usage**
getHTMLToc(toc)

**Arguments**
toc data.frame with TOC info

**Value**
Character vector with HTML toc

---

**getHeightLab**

**Get height of labels: title, subtitle or caption**

**Description**
Get height of labels: title, subtitle or caption

**Usage**
getHeightLab(lab)

**Arguments**
lab String with label.

**Value**
Integer with height in pixels for this element.

---

**Author(s)**
Laure Cougnaud

---

**Author(s)**
Laure Cougnaud
getIndexHTMLTitle

Get index of the line containing the HTML title in a vector of HTML strings

Description

Get index of the line containing the HTML title in a vector of HTML strings

Usage

getIndexHTMLTitle(x)

Arguments

x Character vector with HTML

Value

Integer vector with index of the title in the vector x

getInterimResFile

Get interim res file

Description

Get interim res file

Usage

getInterimResFile(intermediateDir = "./interim", mdFile)

Arguments

intermediateDir String with intermediate directory ("interim" by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.

mdFile String with path of the Markdown file

Value

String with path to the file with intermediate results.
getJitterVar

Add jitter to the variable of the plot, based on the different groups of a grouping variable

Description

Add jitter to the variable of the plot, based on the different groups of a grouping variable

Usage

getJitterVar(data, var, byVar)

Arguments

data Data.frame with data.
var String with variable to add a jitter to.
byVar String with variable containing the groups to jitter by.

Value

Numeric vector of length nrow(data) containing the jittered variable.

Author(s)

Laure Cougnaud

getJsDepClinDataReview

Get Javascript custom scripts required for specific clinical data functionalities.

Description

Get Javascript custom scripts required for specific clinical data functionalities.

Usage

getJsDepClinDataReview(
  type = c("collapsibleButton", "patientProfiles"),
  dep = NULL
)
getMdFromConfig

Arguments

- **type** (optional) Character vector with type of dependencies, either: 'collapsibleButton' or 'patientProfiles'.
- **dep** (optional) Character vector with names of Javascript dependencies By default, all dependencies are included.

Value

List of htmlDependency. To include this dependency in a report e.g. generated with rmarkdown, these can be passed to the: extra_dependencies parameter of the output_format specific function, e.g.: rmarkdown::render(...,output_format = rmarkdown::html_document(extra_dependencies = dep))

Author(s)

Laure Cougnaud

---

**getMdFromConfig**  
*Get path of the HTML file corresponding to a specific config file.*

Description

The name of the Markdown file is based on:

- for the general config.yml file: the basename of the specified indexPath
- for other config file (each sub-report): the name of the config file, after removal of the 'config-' part.

Usage

```
getMdFromConfig(
  configFiles,
  indexPath = "index.Rmd",
  intermediateDir = "./interim"
)
```

Arguments

- **configFiles** Character vector with name or path of the config file(s).
- **indexPath** String with path to the index file, by default 'index.Rmd' in inputDir.
- **intermediateDir** String with intermediate directory ('interim' by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.
Value
String with path to the HTML file

Author(s)
Laure Cougnaud

---

getMdHeader
Get Markdown header, for creation in Rmarkdown.

Description
The depth is extracted:

1. from the settings if a specified depth is provided in the `rmd_file_depth` for the current knitted file
2. level parameter otherwise

Usage
getMdHeader(title, level = 1)

Arguments
- **title**: String with header title.
- **level**: Integer of length 1 with header depth/level, 1 by default

Value
String with Markdown header, to be included in R within cat.

See Also
Other clinical data reporting: checkReportTitles(), forceParams(), getParamsFromConfig(),
gitbook_clinDataReview_report(), html_clinDataReview_report(), knitPrintClinDataReview(),
postProcessReport(), render_clinDataReviewReport()
Description

Read the metadata file from a yaml format. This function checks for existance of the metadata file and its content. In particular, within the yaml file matches the following strings:

- path: Path to the data. More than one path is allowed.
- dateTime: Date and time, usually of the SDTM data creation. When printing the metadata in Rmd document, there is the possibility to add the date and time of the report generation. See `knit_print.clinDataReviewMetadata`.
- datasetInfo: General information about the data sets.

Usage

```r
getMetadata(filePath, namesInfo)
```

Arguments

- **filePath**
  String of path to file. Currently only one file path is supported. If more than one paths are provided, a warning will be printed and the first path will be used.

- **namesInfo**
  Named vector to rename the final output when printed in Rmd. The renaming happens only if the metadata info are printed in Rmd and not in the console.

Details

Note that the input names do not necessarily have to match the exact names. For instance, the user can also write "dataTimeMySDTMData", and the function will parse for existance of the string "dataTime".

Value

A list of:

- **summaryInfo** Information extracted from the inputs path, and dateTime.
- **datasetInfo** Information extracted from datasetInfo.

Examples

```r
# Create temporary yaml file
tmpdir <- tempdir()
library(yaml)

tmpYamlFile <- tempfile(  
  pattern = "file", tmpdir = tempdir(), fileext = ".yml"
)
listArgs <- list(
  path = "myPath",
  dateTime = "2023-04-15 09:00:00",
  datasetInfo = c(  
    title = "My Dataset",
    description = "This is a description of my dataset."
  )
)}
```
getParamsFromConfig

Get parameters from a config file

description

Please note that the information from this config file and the general config file: config.yml are considered.
In case parameters are defined both in the general and specific config files, the parameter from the general config file is ignored.

Usage

going getParamsFromConfig(
    configFile,
    configDir = file.path(inputDir, "config"),
    inputDir = "."
)

Arguments

- configFile: String with filename of the config file of interest in YAML format.
- configDir: String with directory with config files, by default a `config` folder in inputDir. It should contain a general `config.yml` file and dedicated `config-[X].yml` for each chapter. The order of each chapter is specified in the `config` slot in the general general `config.yml`.
- inputDir: String with input directory, working directory by default.
Value

List with parameters from the specified configFile and the general config file: config.yml.
There are two specific handlers:

- parameters tagged with `'[param] !r [value]'` are evaluated in R, and their evaluated value is returned
- parameters tagged with `'[param] !r-lazy [value]'` are imported as character, and need to be further process with `forceParams` inside the report.

Parameters with YAML type 'r-lazy' are imported as character, with this additional class.

Author(s)

Laure Cougnaud

See Also

`forceParams`


---

getParFctReferenceLines

*Get parameter of function used for reference lines*

Description

Get parameter of function used for reference lines

Usage

```r
getParFctReferenceLines(type)
```

Arguments

- `type` : string with line type, either: 'hline', 'abline' or 'vline'.

Value

Character vector with parameter names of the functions

Author(s)

Laure Cougnaud
getPathHyperlink

Get path ('href') property from hyperlink(s).

Description

Get path ('href') property from hyperlink(s).

Usage

gPathHyperlink(x)

Arguments

x
Character vector with hyperlink(s). If multiple, the hyperlinks should be sepa-
rated by: ',', '.

Value

Character vector of length x containing only the hyperlinks.

Author(s)

Laure Cougnaud

getPathTemplate

Get path of template clinical data report

Description

Get path of template clinical data report

Usage

gPathTemplate(file, package = "clinDataReview")

Arguments

file
String with name of the template Rmd document

package
String, which package the template should be extracted from, by default the
clinDataReview package.

Value

String with path to the template in the installed clinDataReview package
getPlotTableVars

Author(s)
Laure Cougnaud

Examples

```r
## Not run:
pathDivisionTemplate <- getPathTemplate("divisionTemplate.Rmd") # get path template in the package
file.copy(from = pathDivisionTemplate, to = ".") # copy to current directory
rmarkdown::render(pathDivisionTemplate) # run file

## End(Not run)
```

getPlotTableVars Extract variables displayed in the attached table, for each available plotting function of the clinDataReview package.

Description

This function is used in each plotting function of the package to extract the variable(s) displayed in the table associated to the plot and their associated labels. This can also be used in the template reports, e.g. to extract reference variable(s) for the comparison table functionality in the plot template report. The following framework is used:

- if variables to be displayed in the table (tableVars) are not specified:
  - all variables displayed in the plot are selected, based on the plot arguments.
  - For example: the variables displayed in the x and y axis and for coloring are extracted for the `scatterplotClinData` plotting function.
  - Label for these variable(s) are extracted from the associated parameter (e.g. `xLab` for `xVar` and so on) or the general parameter for the variable labels (`labelVars`) if not specified.

- if variables to be displayed in the table (tableVars) are specified:
  - these variable(s) are returned.
  - The associated label(s) are extracted from the associated parameter (tableLab) or the general parameter for the variable labels (labelVars) if not specified.

For the functions: `plotCountClinData`, `treemapClinData`, `sunburstClinData`: value to represent are included in the table and colored with a bar.

Usage

```r
getPlotTableVars(plotFunction, plotArgs)
```

Arguments

- **plotFunction**:
  - String with name of the plotting function, be available in the clinDataReview package.
- **plotArgs**:
  - List with parameters passed to the plotting function.
getPositionAndMargins

Value

Character vector with variable to include in the table, with extra attributes (passed to `tableClinData`):

- ‘tableLab’: Named character vector with labels for the table variables
- ‘tablePars’: extra table parameters, only included if specified as input or specified internally.

labels and the table parameters.

Author(s)

Laure Cougnaud

getPositionAndMargins  Get margins and positions of specific elements for a clinical data plot

Description

The elements are positioned as following:

- on top of the plot
  1. title
  2. subtitle
  3. legend, if positioned on top of the plot
  4. facet title
- at the bottom of the plot
  1. label for the x-axis
  2. legend, if positioned on the bottom of the plot
  3. caption

Margins are computed based on the presence of these elements. Only one line is counted for the legend, as plotly will extend the margin if necessary for the legend.

Usage

```r
getPositionAndMargins(
  title = NULL,
  subtitle = NULL,
  xLab = NULL,
  caption = NULL,
  facet = FALSE,
  includeLegend = TRUE,
  legendPosition = "right"
)
```
**getSizePlot**

**Arguments**

- **title** String with title for the plot.
- **subtitle** String with subtitle. The subtitle is included at the top left of the plot, below the title.
- **xLab** String with label for xVar.
- **caption** String with caption. The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.
- **facet** Logical, if TRUE the plot contains facets.
- **includeLegend** Logical, if TRUE (by default) a legend is available in the plot.
- **legendPosition** String with position of the legend, among: 'top'/'left'/'bottom'/'right', 'right' by default.

**Value**

List with:

- 'margin': List with bottom ('t') and top ('t') margins in pixels
- 'position': List with position of the following plot elements:
  - on top of the plot: subtitle and legend (if positioned at the top). The position is defined as the distance in pixels from the top of the plotting area to the bottom of the element (yanchor = 'bottom')
  - at the bottom of the plot: caption, xLab and legend (if positioned at the bottom). The position is defined as the distance in pixels from the bottom of the plotting area to the top of the element (yanchor = 'top')
  - Especially, the legend should be positioned with anchor 'top' such as the margins are automatically expanded if the legend contains multiple rows.

**Author(s)**

Laure Cougnaud

---

**getDescription**

Get dimensions for a clinical data plot

**Description**

This function set sensitive defaults dimensions for a plot in the package. This includes:

- setting a default width for a figure to fit in a standard clinical data review report
- increasing the figure height:
  - for faceted plot, ensuring that each facet is relatively squared
  - if a caption, subtitle, title, title for the x-axis are specified
  - if a legend is set at the bottom or the top of the plot
- increasing the figure width if a legend is set at the left or the right of the plot
Usage

getSizePlot(
  width = NULL,
  height = NULL,
  gg = NULL,
  nrow = 1L,
  ncol = 1L,
  title = NULL,
  subtitle = NULL,
  caption = NULL,
  xLab = NULL,
  facet = FALSE,
  includeLegend = TRUE,
  legendPosition = "right",
  y = NULL
)

Arguments

width Numeric, width of the plot in pixels, 800 by default.
height Numeric, height of the plot in pixels, 500 by default.
gg ggplot
nrow single-length integer specifying the number of facet rows in the plot. (default = 1) Overwritten if gg is specified.
ncol single-length integer specifying the number of facet columns in the plot. (default = 1) Overwritten if gg is specified.
title String with title for the plot.
subtitle String with subtitle.
The subtitle is included at the top left of the plot, below the title.
caption String with caption.
The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.
xLab String with label for xVar.
facet Logical, if TRUE the plot contains facets.
includeLegend Logical, if TRUE (by default) a legend is available in the plot.
legendPosition String with position of the legend, among: 'top'/left'/bottom'/right', 'right' by default.
y Character vector or factor with elements in the y-axis, or list of such vectors. If a list is provided, the maximum height obtained across the different list elements is used.

Value

Numeric vector with width ('width') and height ('height') of the plot in pixels.
getTocNumbering

**Author(s)**
Laure Cougnaud

---

---

**getTocNumbering**  
*Get TOC numbering*

**Description**
Get TOC numbering

**Usage**

getTocNumbering(levels)

**Arguments**

- **levels**
  vector with levels of the section, in the order as available in the book.

**Value**
Character vector with section numbers

**Author(s)**
Laure Cougnaud

---

---

**getWatermark**  
*Get a watermark, to be included in a clinical visualization*

**Description**
Get a watermark, to be included in a clinical visualization

**Usage**

getWatermark(
  label = "EXPLORATORY",
  color = "lightgrey",
  fontsize = 20,
  file = NULL,
  width = 800,
  height = 500
)

Arguments

- **label**: String with label for the watermark, 'EXPLORATORY' by default.
- **color**: String with color for the watermark, 'lightgrey' by default.
- **fontsize**: Integer with fontsize for the watermark, 20 by default.
- **file**: (optional) String with path to a file where the watermark will be exported to.
- **width**: Numeric, width of the plot in pixels, 800 by default.
- **height**: Numeric, height of the plot in pixels, 500 by default.

Value

The file with the watermark (invisibly).

Author(s)

Laure Cougnaud, Kirsten van Hoorde

See Also

Other watermark helpers: addWatermark()

Examples

```r
# export a watermark to a file
watermark <- tempfile(pattern = "watermark", fileext = ".png")
getWatermark(file = watermark)
```

Description

This function is only meant to set sensitive defaults for gitbook. `gitbook` can be used instead.

Usage

```r
gitbook_clinDataReview_report(
  logo = NULL,
  logoHeight = "60px",
  split_by = "section+number",
  config = list(sharing = NULL, toc = list(collapse = "section")),
  extra_dependencies = NULL,
  css = NULL,
  ...
)
```
Arguments

- **logo**: String, path to the logo. No logo is printed by default.
- **logoHeight**: String, indicating the logo height; 60px height by default.
- **split_by**: String, how the reports should be split, (see help of the `gitbook` function)
- **config**: List with config parameters, by default: no sharing and collapsed by section. (see help of the `gitbook` function)
- **extra_dependencies**: NULL by default
- **css**: String, path to the css.
- **...**: Extra parameters passed to the `gitbook` function.

Value

R Markdown output format to pass to `render_book`.

Author(s)

Laure Cougnaud

See Also


---

**html_clinDataReview_report**

*Clinical data format for rmarkdown report.*

Description

This function only kept for back-compatibility, `html_document` can be used instead.

Usage

`html_clinDataReview_report(extra_dependencies = NULL, ...)`

Arguments

- **extra_dependencies**: NULL by default.
- **...**: Extra parameters passed to the `html_document` function.

Value

R Markdown output format to pass to `render`. 
Author(s)

Laure Cougnaud

See Also

Other clinical data reporting: `checkReportTitles()`, `forceParams()`, `getMdHeader()`, `getParamsFromConfig()`, `gitbook_clinDataReview_report()`, `knitPrintClinDataReview()`, `postProcessReport()`, `render_clinDataReviewReport()`

---

JSONSchToRd

Get R Documentation from a JSON schema.

Description

Note: this function doesn’t support the full JSON schema specification, currently only the functionalities required by the templates of the package are implemented.

Usage

```r
JSONSchToRd(JSONSch, title = NULL)
```

Arguments

- **JSONSch**: List with JSON schema, as returned by `fromJSON`.
- **title**: (optional) String with title. This will combined with the JSON schema 'title' tag if this is specified. is not available.

Value

Character vector with R documentation for the specified JSON schema.

Supported JSON schema tags

- 'title' is used as Rd section header
- 'description' is included in the text
- parameters are extracted from the following 'properties' tag:
  - 'type': object type
  - 'doc': documentation for the parameter (custom JSON schema tag). This can contain any Roxygen tags, e.g.: `\link[package]{function}`.
  - 'pattern' (optional): required value for the parameter
  - 'items' (optional): JSON schema for the different elements of an 'object'
  - 'minItems'/'maxItems' (optional): minimum/maximum number of elements in an 'array'
  - 'enum' (optional): set of possible values
  - 'const' (optional): fixed value for the parameter (a.k.a 'constant')

If a parameter is required, it should be listed in the 'required' tag of the schema (outside of the 'properties' tag).
In clinical data reporting, it is often necessary to include output from clinical data, or list of such outputs in a Rmarkdown report, with an appropriate title. The `knitPrintClinDataReview` function facilitates this process.

### Description
Include output from clinical data, or list of such outputs in a Rmarkdown report, with an appropriate title.

### Usage
```r
knitPrintClinDataReview(list, sep = ".", level = 1)
```

### Arguments
- **list**: Named list of clinical data plots, the names are used for the section header. If several section header should be created, either:
  - a list of level 1 named by the different group elements, separated by `sep`, e.g. `list('group1.param1' = ..., 'group1.param2' = ...)`. Such list is e.g. created with `dplyr`.
  - a nested list, named with the different groups, e.g. created with `lapply`.
- **sep**: String with separator used to distinguish different levels in the labels of the list. e.g. `"."` by default.
- **level**: Integer with base level for section, 1 by default.

### Value
No returned value, the plots are included in the report. If an element in the list are empty (NULL), these elements (and the associated sections) are not included in the report.

### Author(s)
Laure Cougnaud

### See Also
Other clinical data reporting: `checkReportTitles()`, `forceParams()`, `getMdHeader()`, `getParamsFromConfig()`, `gitbook_clinDataReview_report()`, `html_clinDataReview_report()`, `postProcessReport()`, `render_clinDataReviewReport()`
knit_print.clinDataReview

Print clinDataReviewTable object in a knitted document (e.g. Rmarkdown document).

Description

Print clinDataReviewTable object in a knitted document (e.g. Rmarkdown document).

Usage

```r
## S3 method for class 'clinDataReview'
knit_print(x, ...)
```

Arguments

- `x`: Object of class clinDataReview
- `...`: Extra parameters for compatibility with `knit_print`, not used currently.

Author(s)

Laure Cougnaud

---

knit_print.clinDataReviewMetadata

Print metadata file in the clinical data report

Description

This function receives the metadata information from `getMetadata` and prints them in a format for an Rmd report. In general, any list could be called as long as it is composed by two elements:

- `summaryInfo`: an R object.
- `datasetInfo`: a data.frame or a matrix.

The first (`summaryInfo`) is printed as `kable` object and the second (`datasetInfo`) is printed as hide/show html button with the function `collapseHtmlContent`.

Usage

```r
## S3 method for class 'clinDataReviewMetadata'
knit_print(x, options = list(), ...)
```
Arguments

- **xLab**
  - String with label for xVar.

- **yLab**
  - String with label for xVar.

- **title**
  - String with title for the plot.

- **caption**
  - String with caption.
  - The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.

Value

Nothing. The tables are ready to be printed in Rmd.

html code to include metadata in a report

---

**layoutClinData**

*Set layout for a clinical data plot.*

Description

Set layout for a clinical data plot.

Usage

```
layoutClinData(
  xLab = NULL,
  yLab = NULL,
  title = NULL,
  caption = NULL,
  subtitle = NULL,
  includeLegend = FALSE,
  legendPosition = "right",
  facet = FALSE,
  nrow = 1L,
  ncol = 1L,
  width,
  height,
  watermark = NULL,
  ...
)
```

Arguments

- **xLab**
  - String with label for xVar.

- **yLab**
  - String with label for xVar.

- **title**
  - String with title for the plot.

- **caption**
  - String with caption.
  - The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.
subtitle
String with subtitle.
The subtitle is included at the top left of the plot, below the title.

includeLegend
Logical, if TRUE (by default) a legend is available in the plot.

legendPosition
String with position of the legend, among: ‘top’/’left’/’bottom’/’right’, ’right’ by default.

facet
Logical (FALSE by default), does the plot contains facets?

nrow
single-length integer specifying the number of facet rows in the plot. (default = 1) Overwritten if gg is specified.

ncol
single-length integer specifying the number of facet columns in the plot. (default = 1) Overwritten if gg is specified.

width
Numeric, width of the plot in pixels, 800 by default.

height
Numeric, height of the plot in pixels, 500 by default.

watermark
(optional) String with path to a file containing a watermark.

...
Any parameters for the layout function. This should contain at least the plot object.

Value
The updated plotly object

Author(s)
Laure Cougnaud

merge.sessionInfo  Merge multiple session information

Description
Merge multiple session information

Usage
## S3 method for class 'sessionInfo'
merge(...)

Arguments
...
objects of type sessionInfo

Value
sessionInfo with combined information

Author(s)
Laure Cougnaud
moveSkeletonFiles  Move skeleton files from the package to a directory

Description
This function moves the files used to create the skeleton from the package to a specified directory.

Usage
moveSkeletonFiles(dir)

Arguments

dir  String, path to the directory.

Value
Nothing, the files are available in the specified directory.

moveXpt  Move data sets from clinUtils

Description
Move SDTM data sets available in clinUtils into a specified local directory.

Usage
moveXpt(dir)

Arguments

dir  String, path to the directory.

Value
Nothing, the data are saved in the dedicated location.
**plotCountClinData**  
*Interactive plot of 'count' data*

**Description**

Note: the table and plot are not (yet) linked.

**Usage**

```r
plotCountClinData(
  data,
  vars,
  varsLab = getLabelVar(vars, labelVars = labelVars),
  valueVar,
  valueLab = getLabelVar(valueVar, labelVars = labelVars),
  colorVar = NULL,
  colorLab = getLabelVar(valueVar, labelVars = labelVars),
  colorPalette = getOption("clinDataReview.colors"),
  colorRange = NULL,
  valueType = "total",
  title = paste(c(paste(valueLab, "by", paste(varsLab, collapse = " and ")), titleExtra),
    collapse = "<br>"),
  titleExtra = NULL,
  subtitle = NULL,
  caption = NULL,
  labelVars = NULL,
  width = NULL,
  height = NULL,
  pathVar = NULL,
  pathLab = getLabelVar(pathVar, labelVars = labelVars),
  hoverVars = c(vars, valueVar, colorVar),
  hoverLab = getLabelVar(hoverVars, labelVars = labelVars),
  table = FALSE,
  tableVars,
  tableLab,
  tableButton = TRUE,
  tablePars = list(),
  id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
  verbose = FALSE,
  typePlot = c("sunburst", "treemap"),
  watermark = NULL
)
```

**Arguments**

- **data**  
  Data.frame with data.
vars Character vector with variables of data containing the groups. If multiple, they should be specified in hierarchical order (from parent to child node).

varsLab Named character vector with labels for vars.

valueVar String with numeric variable of data containing the value to display.

valueLab String with label for the valueVar variable.

colorVar (optional) String with coloring variable (NULL by default). By default, the treemap is colored based by section.

colorLab String with label for colorVar.

colorPalette (optional) Named character vector with color palette. If not specified, the viridis color palette is used. See clinColors.

colorRange (optional) Numeric vector of length 2 with range for the color variable, in case it is a numeric variable.

valueType String with type of values in valueVar (branchvalues of the plot_ly function), among others: 'total' (default, only if sum(child) <= to parent) or 'relative'.

title String with title for the plot.

titleExtra String with extra title for the plot (appended after title).

subtitle String with subtitle.
The subtitle is included at the top left of the plot, below the title.

caption String with caption.
The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.

labelVars Named character vector containing variable labels.

width Numeric, width of the plot in pixels, 800 by default.

height Numeric, height of the plot in pixels, 500 by default.

pathVar String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

\(<a\ href="./path-to-report">label</a>\)

If multiple, they should be separated by ' , '.
The report(s) will be:

- compressed to a zip file and downloaded if the user clicks on the 'p' (a.k.a 'profile') key when hovering on a point of the plot
- included in a collapsible row, and clickable with hyperlinks in the table

pathLab String with label for pathVar, included in the collapsible row in the table.

hoverVars Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.

hoverLab Named character vector with labels for hoverVars.
postProcessReport  

Convert clinical data Markdown files to HTML

### Description

Convert clinical data Markdown files to HTML.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>table</td>
<td>Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)</td>
</tr>
<tr>
<td>tableVars</td>
<td>Character vector with variables to be included in the table.</td>
</tr>
<tr>
<td>tableLab</td>
<td>Named character vector with labels for each tableVars.</td>
</tr>
<tr>
<td>tableButton</td>
<td>Logical, if TRUE (by default) the table is included within an HTML button.</td>
</tr>
<tr>
<td>tablePars</td>
<td>List with parameters passed to the getClinDT function.</td>
</tr>
<tr>
<td>id</td>
<td>String with general id for the plot:</td>
</tr>
<tr>
<td></td>
<td>• 'id' is used as group for the SharedData</td>
</tr>
<tr>
<td></td>
<td>• 'button:[id]' is used as button ID if table is TRUE</td>
</tr>
<tr>
<td></td>
<td>If not specified, a random id, as 'plotClinData[X]' is used.</td>
</tr>
<tr>
<td>verbose</td>
<td>Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.</td>
</tr>
<tr>
<td>typePlot</td>
<td>String with plot type, 'treemap' or 'sunburst'.</td>
</tr>
<tr>
<td>watermark</td>
<td>(optional) String with path to a file containing a watermark.</td>
</tr>
</tbody>
</table>

### Value

Either:

- if a table is requested: a clinDataReview object, a.k.a a list with the 'plot' (plotly object) and 'table' (datatable object)
- otherwise: a plotly object

### Author(s)

Laure Cougnaud

### See Also

Other visualizations of summary statistics for clinical data: barplotClinData(), boxplotClinData(), errorbarClinData(), sunburstClinData(), treemapClinData()
Usage

postProcessReport(
  inputDir = ".",
  configDir = file.path(inputDir, "config"),
  indexPath = file.path(inputDir, "index.Rmd"),
  extraDirs = getExtraDirs(inputDir = inputDir, configDir = configDir),
  outputDir = ".\report",
  intermediateDir = ".\interim",
  mdFiles = NULL,
  nCores = 1,
  logFile = NULL,
  verbose = TRUE,
  ...
)

Arguments

inputDir String with input directory, working directory by default.
configDir String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general 'config.yml'.
indexPath String with path to the index file, by default 'index.Rmd' in inputDir.
extraDirs Character vector with extra directories required by the report, directory with external images. By default, the directories: 'figures', 'tables' and mentioned in the 'patientProfilePath' parameter of the general config file are included. All these folders should be available in inputDir.
outputDir String with output directory, ('report' by default).
intermediateDir String with intermediate directory ('interim' by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.
mdFiles (optional) Path to the Markdown files that should be converted. If specified, the specified config files in configDir are ignored.
nCores Integer containing the number of cores used to render the report (1 by default). If more than 1, two steps of the report creation are run in parallel across chapters:
  • the rendering of the Rmarkdown file to Markdown
  • the conversion from Markdown to HTML
logFile (optional) String with path to a log file, where output (also error/messages/warnings) should be stored. If specified, the entire output is re-directed to this file.
verbose Logical, if TRUE (FALSE by default) progress messages are printed during the report execution.
... Any parameters passed to render, for expert use only.
Value

String with path to the front page of the report.

Author(s)

Laure Cougnaud

See Also

Other clinical data reporting: `checkReportTitles()`, `forceParams()`, `getMdHeader()`, `getParamsFromConfig()`, `gitbook_clinDataReview_report()`, `html_clinDataReview_report()`, `knitPrintClinDataReview()`, `render_clinDataReviewReport()`

---

**print.clinDataReview**

*Print a clinDataReview object in the console*

Description

Print a clinDataReview object in the console

Usage

```r
## S3 method for class 'clinDataReview'
print(x, ...)
```

Arguments

- `x` Object of class clinDataReview
- `...` Extra parameters for compatibility with `print`, not used currently.

Value

No returned value, the object is printed into the console.
### processData

**Process a dataset.**

**Description**

This function is intended to automate all data processing steps for use in the 'clinDataReview' reports using config files.

**Usage**

```
processData(data, processing, labelVars = NULL, ...)
```

**Arguments**

- `data`  
  Data.frame with data.

- `processing`  
  List with processing steps for the data. Each element in the list should be a named list containing the parameters for the specific processing function. The name specifies the processing step, among:
  - 'annotate' for `annotateData` (annotations parameter)
  - 'filter' for `filterData` (filters parameter)
  - 'transform' for `transformData` (transformations parameter)

  Multiple steps of each kind can be specified after each other (e.g. 1: filter, 2: transform, 3: filter, ...).

  If a filter step is specified as a list of multiple filters, the filters are run independently of each other on the entire dataset (see the documentation of `filterData`).

  If filters should be run sequentially, i.e. filter from step 2 should be applied on the filtered dataset from step 1, **separated filtering steps** should be specified, e.g.

  ```
  list(filter = list(var = "ANL01FL", value = "Y"), filter = list(var = "PARAM", value = "QTCF"))
  ```

- `labelVars`  
  Named character vector containing variable labels.

- `...`  
  Any parameters passed to all processing functions (if this parameter is available). If specified, these parameters shouldn’t be specified also in `processing`.

**Value**

Data.frame with processed data, with extra attribute: `labelVars`.

**Author(s)**

Laure Cougnaud
Examples

```r
library(clinUtils)

data(dataADaMCDISCP01)

dataLB <- dataADaMCDISCP01$ADLBC

# filter and annotate data
processData(
  data = dataLB,
  processing = list(
    list(filter = list(var = "ANL01FL", value = "Y")),
    list(annotate = list(vars = "ANRIND", varFct = 'factor(ANRIND, levels = c("L", "N", "H"))'))
  )
)

## multiple filtering steps:
# If these are specified in the same 'filter' step condition, these are considered independently,
# and the selected records combined with an 'AND' operator.
# Example: consider only records:
# - with analysis flag AND
# - from subject with high/low measurement (for all records) for each parameter
processData(
  data = dataLB,
  processing = list(
    list(filter = list(
      list(var = "ANL01FL", value = "Y"),
      list(var = "ANRIND", value = c("L", "H"),
           postFct = any, varsBy = c("USUBJID", "PARAM"))
    ))
  )
)

# a custom operator to combine the selected records can be specified
# Example: consider only records:
# - with analysis flag OR
# - from subject with high/low measurement (for all records) for each parameter
processData(
  data = dataLB,
  processing = list(
    list(filter = list(
      list(var = "ANL01FL", value = "Y"),
      "|",
      list(var = "ANRIND", value = c("L", "H"),
           postFct = any, varsBy = c("USUBJID", "PARAM"))
    ))
  )
)
```
# If the filtering conditions are specified in different filtering steps, these are
# considered sequentially.
# Example:
# 1) consider only analysis records and
# 2) from these records, consider only subject with high/low measurement for
# each parameter
processData(
    data = dataLB,
    processing = list(
        list(filter = list(var = "ANL01FL", value = "Y")),
        list(filter = list(var = "ANRIND", value = c("L", "H")),
            postFct = any, varsBy = c("USUBJID", "PARAM"))
    )
)
# Note for this particular

renamePathDateInfoMetadata

*Rename variable names of metadata info*

**Description**

Rename variable names referring to the paths and the date.

**Usage**

`renamePathDateInfoMetadata(summaryInfo, namesInfo)`

**Arguments**

- `summaryInfo` A matrix, see output from `getMetadata`.
- `namesInfo` Named vector to rename the final output.

**Value**

A matrix, same as input `summaryInfo` with renamed variable names.
renderChapter

Render one chapter of a clinical report, based on a configuration file

Description

Render one chapter of a clinical report, based on a configuration file

Usage

renderChapter(
  configFile,
  configGeneralParams = getParamsFromConfig(configFile = "config.yml", configDir =
    configDir, inputDir = inputDir),
  configDir = file.path(inputDir, "config"),
  indexPath = file.path(inputDir, "index.Rmd"),
  inputDir = ".",
  intermediateDir = "./interim",
  logFile = NULL,
  verbose = TRUE,
  ...
)

Arguments

configFile  String with filename of the config file of interest in YAML format.
configGeneralParams
  List with parameters from the general config file
configDir  String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general general 'config.yml'.
indexPath  String with path to the index file, by default 'index.Rmd' in inputDir.
inputDir  String with input directory, working directory by default.
intermediateDir  String with intermediate directory ('interim' by default), where markdown files and rds file specifying Js libraries (with knit_meta) for each sub report are stored.
logFile  (optional) String with path to a log file, where output (also error/messages/warnings) should be stored. If specified, the entire output is re-directed to this file.
verbose  Logical, if TRUE (FALSE by default) progress messages are printed during the report execution.
...  options passed to renderFile
renderFile

Value

No output file, the Markdown report for the chapter and the knit_meta object is available in the intermediateDir directory.
If the input parameters are not correctly extracted, NULL is returned.

renderFile Render a rmarkdown file, possibly in a new R session

Description

This has the possibility to save output in a log file, and saving also session information.

Usage

renderFile(input, encoding = "UTF-8", params = NULL, logFile = NULL, ...)

Arguments

input Input file to be rendered.
encoding String with encoding, 'UTF-8' by default.
params List with input parameters for this document.
These parameters should be accessed in the Rmd document via params$....
These parameters will be saved to a RDS file and imported during the report rendering.
logFile (optional) String with path to a log file, where output (also error/messages/warnings) should be stored. If specified, the entire output is re-directed to this file.
... Any extra parameters passed to render, for expert use only.

Details

Note: this function is inspired from xfun::Rscript_call

Value

Output of the function with additional attribute: 'sessionInfo' containing the details of the session information. If the report fails, an error message is returned.

Author(s)

Laure Cougnaud
render_clinDataReviewReport

Render a clinical data review report.

Description

Render a clinical data review report.

Usage

render_clinDataReviewReport(
  configFiles = NULL,
  configDir = file.path(inputDir, "config"),
  logFile = NULL,
  indexPath = file.path(inputDir, "index.Rmd"),
  inputDir = ".",
  outputDir = ".\report",
  intermediateDir = ".\interim",
  extraDirs = getExtraDirs(inputDir = inputDir, configDir = configDir),
  quiet = FALSE,
  verbose = TRUE,
  nCores = 1
)

Arguments

configFiles (optional) Character vector with specific config files to be converted from Rmarkdown to Markdown. If
  • not specified (by default): all config files specified in the general 'config.yml' will be run (Rmd -> md)
  • specified (expert use only): only the specified files will be run (Rmd -> md). Other config files mentioned in the general 'config.yml' file won't be rerun, so the associated 'md' file should be already available in the intermediateDir folder.

configDir String with directory with config files, by default a 'config' folder in inputDir. It should contain a general 'config.yml' file and dedicated 'config-[X].yml' for each chapter. The order of each chapter is specified in the 'config' slot in the general 'config.yml'.

logFile (optional) String with path to a log file, where output (also error/messages/warnings) should be stored. If specified, the entire output is re-directed to this file.

indexPath String with path to the index file, by default 'index.Rmd' in inputDir.

inputDir String with input directory, working directory by default.

outputDir String with output directory, ('report' by default).
**render_clinDataReviewReport**

- **intermediateDir**
  String with intermediate directory (‘interim’ by default), where markdown files
  and rds file specifying Js libraries (with knit_meta) for each sub report are
  stored.

- **extraDirs**
  Character vector with extra directories required by the report, directory with
  external images. By default, the directories: ‘figures’, ‘tables’ and mentioned
  in the ‘patientProfilePath’ parameter of the general config file are included. All
  these folders should be available in inputDir.

- **quiet**
  Logical, if TRUE (FALSE by default) messages during the execution of each
  report are not displayed in the console (see render).

- **verbose**
  Logical, if TRUE (FALSE by default) progress messages are printed during the
  report execution.

- **nCores**
  Integer containing the number of cores used to render the report (1 by default). If
  more than 1, two steps of the report creation are run in parallel across chapters:
  - the rendering of the Rmarkdown file to Markdown
  - the conversion from Markdown to HTML

**Value**

String with path to the front page of the report.

**Process**

This function is based on the render_book function, with the extra functionalities:

- specification of chapter-specific input parameters, specified in YAML configuration files
- (optional) creation of each chapter in parallel if nCores > 1. In that case, all chapters are run
  in parallel, excepted the chapter(s) run internally in parallel (config file with parallel set to
  ‘TRUE’).
- (optional) split of each chapter into html file specific for each chapter, by specifying the
  split_by parameter in the chapter-specific config file

This consists of:

1. importing the general config file (‘config’.yml) to identify each report of interest (‘config’ tag)
2. for each report of interest:
   - loading the report specific parameters from the associated ‘config’ file (see the getParamsFromConfig
     function)
   - if the template should be extracted from a specified package (templatePackage tag), this
     template is copied to the current directory. Please note that if a file with same name is
     available in the working directory, this file will be overwritten.
   - running the report (‘template’ tag) with the associated parameters in a new R session for
     reproducibility, to obtain the associated Markdown file.
     This step is parallelized across the different config files, if the nCores parameter is spec-     ified.
3. checking if the associated Markdown and rds file (list of Js dependencies) are available in
   intermediateDir
4. split each chapter into separated Markdown documents, based on the split_by parameter (specified at the report or config level)

5. conversion of each Markdown document to an HTML document. This step is parallelized across the different Markdown documents, if the nCores parameter is specified.

6. build the book:
   (a) creation of a common TOC for the book
   (b) inclusion of the TOC in each Markdown file
   (c) update of the section number in each chapter
   (d) inclusion of the section number in each HTML file name

If the execution of a specific report fails with error, a warning message is triggered. A report containing only the specified title is created, to ensure output consistency (especially html file numbering) in case the report succeeds.

Available template report

see ?‘clinDataReview-templates‘ for a list of clinical data template report available in the package.

Extension to chapter-specific split

The bookdown ‘split_by’ parameter is extended, to support:

- chapter-specific split, specified in the configuration file of the specific chapter, via the split_by parameter
- specification as a number (if specified within a config file), e.g. ’0’ for no split, 1’ for chapter, ’2’ for section, ’3’ for subsection, ...
- split at section level higher than 2 (until 7) (if specified within a config file)

Author(s)

Laure Cougnaud

See Also

Other clinical data reporting: checkReportTitles(), forceParams(), getMdHeader(), getParamsFromConfig(), gitbook_clinDataReview_report(), html_clinDataReview_report(), knitPrintClinDataReview(), postProcessReport()
scatterplotClinData  

Scatterplot of variables of interest for clinical data visualization.

Description

The parameters for this visualization are based on ggplot2 (aesthetic, scale, ...), parameter specification, unlike the other visualizations of the package.

Usage

scatterplotClinData(
  data,
  xVar,
  yVar,
  xLab = getLabelVar(var = xVar, labelVars = labelVars),
  xLabVar = NULL,
  yLab = getLabelVar(var = yVar, labelVars = labelVars),
  yLabVar = NULL,
  aesPointVar = list(),
  pointPars = list(),
  aesLineVar = list(),
  linePars = list(),
  lineInclude = length(aesLineVar) > 0,
  aesSmoothVar = list(),
  smoothPars = list(),
  smoothInclude = length(c(aesSmoothVar, smoothPars)) > 0,
  aesLab,
  xTrans = "identity",
  yTrans = "identity",
  xPars = list(),
  yPars = list(),
  xLimVars = NULL,
  yLim = NULL,
  xLim = NULL,
  yLimExpandData = TRUE,
  xLimExpandData = TRUE,
  title = paste(c(paste(yLab, "vs", xLab), titleExtra), collapse = "<br>")
  titleExtra = NULL,
  caption = NULL,
  subtitle = NULL,
  facetPars = list(),
  facetType = c("wrap", "grid"),
  scalePars = list(),
  themePars = list(legend.position = "bottom"),
  refLinePars = NULL,
  labelVars = NULL,
  width = NULL,
height = NULL,
hoverVars,
hoverLab,
idVar = "USUBJID",
idLab = getLabelVar(idVar, labelVars = labelVars),
idHighlightBox = FALSE,
pathVar = NULL,
pathExpand = FALSE,
id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
selectVars = NULL,
selectLab = getLabelVar(selectVars, labelVars = labelVars),
table = FALSE,
tableVars,
tableLab,
tableButton = TRUE,
tablePars = list(),
watermark = NULL,
verbose = FALSE
)

Arguments

data Data.frame with input data.
xVar String with column of data containing x-variable.
yVar String with column of data containing y-variable.
xLab String with label for xVar.
xLabVar (optional) Character vector with column(s) of data containing variable(s) to display in the label of the x-axis.
yLab String with label for xVar.
yLabVar (optional) Character vector with column(s) of data containing variable(s) to display in the label of the y-axis.
aesPointVar List with specification of aesthetic variable(s), for the point, passed to the mapping parameter of geom_point, e.g. list(color = "TRTP"). Please note by default symbols with fill and color are used. Color is used for the outside of the points, fill for the inside and the hover. Usually, you might want to specify both filling and coloring.
pointPars List with parameters other than aesthetic variables to pass to geom_point, defaults to empty list.
aesLineVar List with specification of aesthetic variable(s), for the line, passed to the mapping parameter of geom_line, e.g. list(group = "USUBJID").
linePars List with parameters other than aesthetic variables to pass to geom_line, defaults to empty list.
lineInclude Logical, if TRUE (by default if aesLineVar is specified) include a scatterplot.
aesSmoothVar List with specification of aesthetic variable(s), for the smoothing layer, passed to the mapping parameter of geom_smooth defaults to empty list.
scatterplotClinData

smoothPars List with parameters other than aesthetic variables to pass to `geom_smooth`, defaults to empty list. Note this parameter overwrites other parameters set by `aesSmoothVar`.

smoothInclude Logical, if TRUE (by default if one of `aesSmoothVar` or `smoothPars` is non-empty)

aesLab Named character vector with labels for each aesthetic variable.

xTrans, yTrans Transformation for the x/y- variables, passed to the `trans` parameter of `scale_x_continuous/ scale_y_continuous`.

xPars, yPars List with extra parameters for x/y axis, passed to the `scale_x_continuous/ scale_y_continuous` functions, besides `trans` and `limits`.

xLabVars Character vector with variable(s) to be displayed as the labels of the ticks in the x-axis.

By default, xVar is displayed.

If specified, this overwrites any labels specified via `xPars`.

In case the variable(s) contain different elements by `xVar` or between facets, they are combined and displayed below each other.

xLim, yLim Numeric vector of length 2 with limits for the x/y axes.

xLimExpandData, yLimExpandData Logical (TRUE by default), should the limits specified via `xLim/yLim` be expanded to include any data points outside of these limits? Please note that the same limits are set for all facets.

title String with title for the plot.

titleExtra String with extra title for the plot (appended after `title`).

caption String with caption.

The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.

subtitle String with subtitle.

The subtitle is included at the top left of the plot, below the title.

facetPars List with facetting parameters, passed to the facetting function.

facetType String with facetting type, either:

- 'wrap': `facet_wrap`
- 'grid': `facet_grid`

scalePars List with parameters to customize scales. Each sublist contains a set of parameters passed to the `scale_discrete_manual` function.

If palette(s) are not specified, default palettes are used (see `getColorPalette, getShapePalette, getLinetypePalette`)

themePars List with general theme parameters (see `theme`).

refLinePars (optional) Nested list, with parameters for each reference line(s). Each sublist (a.k.a reference line) contains:

- aesthetic value(s) or variable(s) for the lines (in this case column names of data) for reference lines. The line position is controlled by the aesthetics supported in `geom_vline, geom_hline` and `geom_abline`. 


• 'label': (optional) Logical specifying if the line should be annotated (FALSE to not annotate the line) or string with annotation label. By default, the value of the position of the horizontal/vertical line or the equation of the diagonal line is displayed.

labelVars Named character vector containing variable labels.
width Numeric, width of the plot in pixels, 800 by default.
height Numeric, height of the plot in pixels, 500 by default.
hoverVars Character vector with variables to be displayed in the hover, by default xVar, yVar and any aesthetic variables.
hoverLab Named character vector with labels for hoverVars.
idVar String with variable containing subject ID.
idLab String with label for idVar.
idHighlightBox Logical, if TRUE (FALSE by default) a selectize box is included to highlight selected element(s) of the ID variable (idVar).
pathVar String with variable of data containing path to a subject-specific report. The report info should be unique for each element of idVar. The report will be:
  • opened in a different window in the browser if the user clicks on the 'p' (a.k.a 'profile') key when hovering on a point of the plot
  • opened in the browser via hyperlink in the table
pathExpand Logical, if FALSE (by default) the path to subject-report is included in an hyperlink in the table, otherwise a collapsed row is created. This should be set to TRUE only if multiple paths are included for each row in pathVar (e.g. in case of summary table).
idd String with general id for the plot:
  • 'id' is used as group for the SharedData
  • 'button:[id]' is used as button ID if table is TRUE
If not specified, a random id, as 'plotClinData[X]' is used.
selectVars (optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).
selectLab (Named) character vector with label for selectVars.
table Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. The plot and table are linked when included in a Rmarkdown document: when clicking on an plot element, only the corresponding records are retained in the associated table; when some records are selected in the table, they are highlighted in the associated table.
tableVars Character vector with variables to be included in the table.
tableLab Named character vector with labels for each tableVars.
tableButton Logical, if TRUE (by default) the table is included within an HTML button.
tablePars List with parameters passed to the getClinDT function.
watermark (optional) String with path to a file containing a watermark.
verbose Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.
Value

Either:

- if a table is requested: a `clinDataReview` object, a.k.a a list with the 'plot' (`plotly` object) and 'table' (`datatable` object)
- otherwise: a `plotly` object

Author(s)

Laure Cougnaud

See Also

Other Clinical data visualization of individual profiles: `timeProfileIntervalPlot()`

Examples

```r
library(clinUtils)

data(dataADaMCDISCP01)
labelVars <- attr(dataADaMCDISCP01, "labelVars")

dataLB <- dataADaMCDISCP01$ADLBC
dataOM <- dataADaMCDISCP01$ADSL
dataLB <- annotateData(dataLB, annotations = list(data = dataOM))
# subset of the data for the example
dataLB <- subset(dataLB, VISIT %in% c(“SCREENING 1”, “WEEK 2”, “WEEK 8”))

## time profile

dataPlot <- subset(dataLB, PARAMCD == "ALT")

# with relative day
scatterplotClinData(
 data = dataPlot,
 xVar = "ADY",
 yVar = "LBSTRESN",
 aesPointVar = list(color = "TRTP", fill = "TRTP"),
 aesLineVar = list(group = "USUBJID", color = "TRTP"),
 labelVars = labelVars
)

# with actual visit
dataPlot$AVISIT <- with(dataPlot, reorder(trimws(AVISIT), AVISITN))
scatterplotClinData(
 data = dataPlot,
 xVar = "AVISIT",
 yVar = "LBSTRESN",
 aesPointVar = list(color = "TRTP", fill = "TRTP"),
 aesLineVar = list(group = "USUBJID", color = "TRTP"),
 labelVars = labelVars
)
```
## Not run:

# add number of subjects below each visit

if (requireNamespace("inTextSummaryTable", quietly = TRUE)) {

# compute number of subjects by visit
summaryTable <- inTextSummaryTable::computeSummaryStatisticsTable(
  dataPlot,
  rowVar = "AVISIT",
  stats = "n"
)

# add it in the data
dataPlot <- merge(dataPlot, summaryTable[, c("AVISIT", "n")], all.x = TRUE)
dataPlot$n <- paste0("N=", dataPlot$n)

scatterplotClinData(
  data = dataPlot,
  xVar = "AVISIT", xLabVars = c("AVISIT", "n"),
  yVar = "LBSTRESN",
  aesPointVar = list(color = "TRTP", fill = "TRTP"),
  aesLineVar = list(group = "USUBJID", color = "TRTP"),
  labelVars = labelVars
)
}

## End(Not run)

## pairwise comparison plot of two parameters of interest:

# format data long -> wide format (one column per lab param)
dataPlot <- subset(dataLB, PARAMCD %in% c("ALT", "AST"))
dataPlot <- stats::aggregate(
  LBSTRESN ~ USUBJID + VISIT + VISITNUM + PARAMCD,
  data = dataPlot,
  FUN = mean
)
dataPlotWide <- stats::reshape(
  data = dataPlot,
  timevar = "PARAMCD", idvar = c("USUBJID", "VISIT", "VISITNUM"),
  direction = "wide"
)
colnames(dataPlotWide) <- sub("^LBSTRESN.", "", colnames(dataPlotWide))

# scatterplot per visit
scatterplotClinData(
  data = dataPlotWide,
  xVar = "ALT", yVar = "AST",
  aesPointVar = list(color = "USUBJID", fill = "USUBJID"),
  aesLineVar = list(group = "USUBJID", color = "USUBJID"),
  labelVars = labelVars
)
themePars = list(legend.position = "none"),
facetPars = list(facets = "VISIT"),
labelVars = labelVars,
subtitle = "Visualization is split by visit",
caption = "Points are colored by subject ID"
)

## Not run:

# scatterplot with all visits, link subjects
xLab <- getLabelParamcd(paramcd = "ALT", data = dataLB,
paramcdVar = "PARAMCD", paramVar = "PARAM")
yLab <- getLabelParamcd(paramcd = "AST", data = dataLB,
paramcdVar = "PARAMCD", paramVar = "PARAM")
scatterplotClinData(
  data = dataPlotWide,
  xVar = "ALT", yVar = "AST",
xLab = xLab,
yLab = yLab,
aesPointVar = list(color = "VISIT", fill = "VISIT"),
aesLineVar = list(group = "USUBJID"),
labelVars = labelVars
)

# scatterplot of different visits versus baseline

# add baseline as extra column:
dataPlot <- subset(dataLB, PARAMCD == "ALT")
dataPlotBL <- subset(dataPlot, VISIT == "SCREENING 1")
dataPlotBL <- dataPlotBL[with(dataPlotBL, order(USUBJID, -ADY)), ]
dataPlotBL <- dataPlotBL[!duplicated(dataPlotBL$USUBJID), ]
dataPlot$LBSTRESNBL <- dataPlot[match(dataPlot$USUBJID, dataPlotBL$USUBJID), "LBSTRESN"]

# sort visits:
dataPlot$VISIT <- with(dataPlot, reorder(VISIT, VISITNUM))

xLab <- paste(labelVars["LBSTRESN"], "for last screening visit")
yLab <- paste(labelVars["LBSTRESN"], "at visit X")
paramLab <- getLabelParamcd(paramcd = "ALT", data = dataLB,
paramcdVar = "PARAMCD", paramVar = "PARAM")
scatterplotClinData(
  data = dataPlot,
  xVar = "LBSTRESNBL", xLab = xLab,
yVar = "LBSTRESN", yLab = yLab,
aesPointVar = list(color = "USUBJID", fill = "USUBJID"),
aesLineVar = list(group = "USUBJID", color = "USUBJID"),
hoverVars = c("USUBJID", "VISIT", "ADY", "LBSTRESN"),
labelVars = labelVars,
facetPars = list(facets = "VISIT"),
themePars = list(legend.position = "none"),
title = paste("Comparison of actual value of ",
paramLab,
"at each visit versus baseline")
refLinePars = list(
  list(slope = 1, intercept = 0, linetype = 1, color = "black",
       label = FALSE),
  list(yintercept = "A1LO", linetype = 2, color = "blue"),
  list(yintercept = "A1HI", linetype = 2, color = "purple",
       label = "Reference Range Upper Limit")
)

## scatterplot with smoothing layer

data <- data.frame(
  subj = c(rep('subj1', 20), rep('subj2', 20)),
  time = rep(1:20, 2),
  response = c(1:20, 50:31) + runif(min = -3, max = +3, 40),
  treat = rep(c('trA', 'trB'), 20),
  stringsAsFactors = FALSE
)

# smoothing per subject
smoothPlot <- scatterplotClinData(
  data = data,
  xVar = "time", yVar = "response",
  aesPointVar = list(color = "treat"),
  aesLineVar = list(group = 'subj'),
  linePars = list(linetype='dotted'),
  aesSmoothVar = list(color='subj', group='subj'),
  smoothPars = list(alpha=0.5, size=0.3, se=TRUE, color = 'black')
)
smoothPlot

# plot smoothing over subjects
smoothPlot <- scatterplotClinData(
  data = data,
  xVar = "time", yVar = "response",
  aesPointVar = list(color = "treat"),
  aesLineVar = list(group = 'subj'),
  linePars = list(linetype='dotted'),
  aesSmoothVar = list(),
  smoothPars = list(alpha=0.5, size=0.3, se=TRUE, color = 'black')
)
smoothPlot

## End(Not run)

# add a selection box
if(interactive()){
  dataPlot <- subset(dataLB, PARAMCD == "ALT")
  dataPlot$TRTA <- with(dataPlot, reorder(TRTA, TRTAN))
setFacetLayoutWrap

scatterplotClinData(
data = dataPlot,
xVar = "ADY",
yVar = "LBSTRESN",
aesPointVar = list(fill = "TRTA", color = "TRTA"),
aesLineVar = list(group = "USUBJID", color = "TRTA"),
selectVars = "TRTA",
labelVars = labelVars
)
)

setFacetLayoutWrap

Set facetting layout for 'wrap' faceting.

Description

By default, the number of columns is 2.

Usage

setFacetLayoutWrap(data, facetPars = list())

Arguments

data Data.frame with data.

facetPars List with facetting parameters, passed to the facetting function. Variables should be specified as character or formula. For 'wrap' faceting (FacetType is 'wrap'), if the layout is not specified via nrow/ncol, 2 columns are used by default.

Value

Updated facetPars.

Author(s)

Laure Cougnaud
setPaletteStaticScatterplotClinData

_Get standard palette for the staticScatterplotClinData function._

Description

Get standard palette for the staticScatterplotClinData function.

Usage

```r
setPaletteStaticScatterplotClinData(data, var, aes, scalePars, geomAes, ...)
```

Arguments

- **data**  
  Data.frame with data for the plot.

- **var**  
  Character vector with variable(s) to consider. If multiple, currently only the first one is considered.

- **aes**  
  String with aesthetic, either: 'color', 'shape' or 'linetype'.

- **scalePars**  
  List with parameters to customize scales. Each sublist should contains a set of parameters passed to the `scale_discrete_manual` function.  
  If palette(s) are not specified, default palettes are used (see `getColorPalette`, `getShapePalette`, `getLinetypePalette`)

- **geomAes**  
  List with aesthetic for each geom.

- **...**  
  Any extra parameters than `x` and `n` for the default palette fcts.

Value

List with: `scalePars` and `geomAes`, each of those potentially updated with default palette(s).

Author(s)

Laure Cougnaud

---

_splitChapter

_Split a chapter based on the 'split_by' parameter._

Description

Split a chapter based on the 'split_by' parameter.
splitChapter

Usage

```r
splitChapter(
  configFile = NULL,
  configDir = './config',
  mdFile = NULL,
  indexPath = 'index.Rmd',
  intermediateDir = './interim',
  outputDir = './report',
  verbose = TRUE
)
```

Arguments

- `configFile` String with filename of the config file of interest in YAML format.
- `configDir` String with directory with config files, by default a `config` folder in `inputDir`. It should contain a general `config.yml` file and dedicated `config-[X].yml` for each chapter. The order of each chapter is specified in the `config` slot in the general `config.yml`.
- `mdFile` (optional) Path to the Markdown file containing the chapter. If not specified, the Markdown file corresponding to the specified `configFile` parameter is used.
- `indexPath` String with path to the index file, by default `index.Rmd` in `inputDir`.
- `intermediateDir` String with intermediate directory (`interim` by default), where markdown files and `rds` file specifying Js libraries (with `knit_meta`) for each sub report are stored.
- `outputDir` String with output directory, (`report` by default).
- `verbose` Logical, if TRUE (FALSE by default) progress messages are printed during the report execution.

Value

No return value, the Markdown files are split as specified.

Extension to chapter-specific split

The bookdown `split_by` parameter is extended, to support:

- chapter-specific split, specified in the configuration file of the specific chapter, via the `split_by` parameter
- specification as a number (if specified within a config file), e.g. '0' for no split, '1' for chapter, '2' for section, '3' for subsection, ...
- split at section level higher than 2 (until 7) (if specified within a config file)

Author(s)

Laure Cougnaud
staticScatterplotClinData

Scatterplot of variables of interest for clinical data visualization

Description

Scatterplot of variables of interest for clinical data visualization

Usage

```r
staticScatterplotClinData(
  data, 
  xVar, 
  yVar, 
  xLab = getLabelVar(xVar, labelVars = labelVars), 
  yLab = getLabelVar(yVar, labelVars = labelVars), 
  aesPointVar = list(), 
  pointPars = list(), 
  aesLineVar = list(), 
  linePars = list(), 
  lineInclude = length(c(aesLineVar, linePars)) > 0, 
  aesSmoothVar = list(), 
  smoothPars = list(), 
  smoothInclude = length(c(aesSmoothVar, smoothPars)) > 0, 
  aesLab, 
  xTrans = "identity", 
  yTrans = "identity", 
  xPars = list(), 
  yPars = list(), 
  xLabVars = NULL, 
  yLim = NULL, 
  xLim = NULL, 
  yLimExpandData = TRUE, 
  xLimExpandData = TRUE, 
  title = paste(c(paste(yLab, "vs", xLab), titleExtra), collapse = "<br>"), 
  titleExtra = NULL, 
  facetPars = list(), 
  facetType = c("wrap", "grid"), 
  scalePars = list(), 
  themePars = list(legend.position = "bottom"), 
  refLinePars = NULL, 
  labelVars = NULL, 
  hoverVars = NULL, 
  geomType = c("point", "col")
)
```
Arguments

data Data.frame with input data.
xVar String with column of data containing x-variable.
yVar String with column of data containing y-variable.
xLab String with label for xVar.
yLab String with label for xVar.
aesPointVar List with specification of aesthetic variable(s), for the point, passed to the mapping parameter of `geom_point`, e.g. `list(color = "TRTP")`. Please note by default symbols with fill and color are used. Color is used for the outside of the points, fill for the inside and the hover. Usually, you might want to specify both filling and coloring.
pointPars List with parameters other than aesthetic variables to pass to `geom_point`, defaults to empty list.
aesLineVar List with specification of aesthetic variable(s), for the line, passed to the mapping parameter of `geom_line`, e.g. `list(group = "USUBJID")`.
linePars List with parameters other than aesthetic variables to pass to `geom_line`, defaults to empty list.
lineInclude Logical, if TRUE (by default if `aesLineVar` is specified) include a scatterplot.
aesSmoothVar List with specification of aesthetic variable(s), for the smoothing layer, passed to the mapping parameter of `geom_smooth` defaults to empty list.
smoothPars List with parameters other than aesthetic variables to pass to `geom_smooth`, defaults to empty list. Note this parameter overwrites other parameters set by `aesSmoothVar`.
smoothInclude Logical, if TRUE (by default if one of `aesSmoothVar` or `smoothPars` is non-empty)
aesLab Named character vector with labels for each aesthetic variable.
xTrans, yTrans Transformation for the x/y- variables, passed to the `trans` parameter of `scale_x_continuous`/`scale_y_continuous`.
xPars, yPars List with extra parameters for x/y axis, passed to the `scale_x_continuous`/`scale_y_continuous` functions, besides `trans` and limits.
xLabVars Character vector with variable(s) to be displayed as the labels of the ticks in the x-axis.
By default, xVar is displayed.
If specified, this overwrites any labels specified via `xPars`.
In case the variable(s) contain different elements by `xVar` or between facets, they are combined and displayed below each other.
xLim, yLim Numeric vector of length 2 with limits for the x/y axes.
xLimExpandData, yLimExpandData Logical (TRUE by default), should the limits specified via `xLim/yLim` be expanded to include any data points outside of these limits? Please note that the same limits are set for all facets.
title String with title for the plot.
titleExtra string with extra title for the plot (appended after title).

facetPars list with facetting parameters, passed to the facetting function.

facetType string with facetting type, either:
  • 'wrap': facet_wrap
  • 'grid': facet_grid

scalePars list with parameters to customize scales. Each sublist should contain a set of parameters passed to the scale_discrete_manual function. If palette(s) are not specified, default palettes are used (see getColorPalette, getShapePalette, getLinetypePalette).

themePars list with general theme parameters (see theme).

refLinePars (optional) Nested list, with parameters for each reference line(s). Each sublist (a.k.a a reference line) contains:
  • aesthetic value(s) or variable(s) for the lines (in this case column names of data) for reference lines. The line position is controlled by the aesthetics supported in geom_vline, geom_hline and geom_abline.
  • 'label': (optional) Logical specifying if the line should be annotated (TRUE to annotate the line) or string with annotation label. By default, the value of the position of the horizontal/vertical line or the equation of the diagonal line is displayed.

labelVars named character vector containing variable labels.

hoverVars character vector with variables to be displayed in the hover, by default xVar, yVar and any aesthetic variables.

geomType string with type of the geom used, either:
  • 'point': scatterplot with geom_point is created
  • 'col': barplot with geom_col is created

Value

ggplot object

Author(s)

Laure Cougnaud, Adriaan Blommaert

Description

Note: the table and plot are not (yet) linked.

Usage

sunburstClinData(...)
Arguments

Arguments passed on to plotCountClinData

colorVar (optional) String with coloring variable (NULL by default). By default, the treemap is colored based by section.
colorRange (optional) Numeric vector of length 2 with range for the color variable, in case it is a numeric variable.
vars Character vector with variables of data containing the groups. If multiple, they should be specified in hierarchical order (from parent to child node).
varsLab Named character vector with labels for vars.
valueVar String with numeric variable of data containing the value to display.
valueLab String with label for the valueVar variable.
valueType String with type of values in valueVar (branchvalues of the plot_ly function), among others: 'total' (default, only if sum(child) <= to parent) or 'relative'.
pathVar String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

\(<a \text{href}=\"./path-to-report\"></a\>

If multiple, they should be separated by: ', '.
The report(s) will be:

- compressed to a zip file and downloaded if the user clicks on the 'p' (a.k.a 'profile') key when hovering on a point of the plot
- included in a collapsible row, and clickable with hyperlinks in the table

pathLab String with label for pathVar, included in the collapsible row in the table.
table Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)
data Data.frame with data.
verbose Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.
width Numeric, width of the plot in pixels, 800 by default.
height Numeric, height of the plot in pixels, 500 by default.
hoverVars Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.
hoverLab Named character vector with labels for hoverVars.
labelVars Named character vector containing variable labels.
id String with general id for the plot:

- 'id' is used as group for the SharedData
- 'button:[id]' is used as button ID if table is TRUE
  If not specified, a random id, as 'plotClinData[X]' is used.
title String with title for the plot.
titleExtra String with extra title for the plot (appended after title).
caption String with caption.
   The caption is included at the bottom right of the plot. Please note that this
   might overlap with vertical or rotated x-axis labels.
subtitle String with subtitle.
   The subtitle is included at the top left of the plot, below the title.
colorLab String with label for colorVar.
colorPalette (optional) Named character vector with color palette. If not
   specified, the viridis color palette is used.
   See clinColors.
watermark (optional) String with path to a file containing a watermark.
tableButton Logical, if TRUE (by default) the table is included within an
   HTML button.
tableVars Character vector with variables to be included in the table.
tableLab Named character vector with labels for each tableVars.
tablePars List with parameters passed to the getClinDT function.

Value
   Either:
   • if a table is requested: a clinDataReview object, a.k.a a list with the 'plot' (plotly object)
     and 'table' (datatable object)
   • otherwise: a plotly object

Author(s)
   Laure Cougnaud

See Also
   Other visualizations of summary statistics for clinical data: barplotClinData(), boxplotClinData(),
   errorbarClinData(), plotCountClinData(), treemapClinData()

Examples
   library(clinUtils)
   data(dataADaMCDISP01)
   labelVars <- attr(dataADaMCDISP01, "labelVars")

   dataAE <- dataADaMCDISP01$ADAE
dataDM <- dataADaMCDISP01$ADSL

   ## example of basic sunburst:
   
   # sunburst takes as input table with counts
   if (requireNamespace("inTextSummaryTable", quietly = TRUE)) {

   # total counts: Safety Analysis Set (patients with start date for the first treatment)
dataTotal <- subset(dataDM, RFSTDTC != "")

# compute adverse event table
tableAE <- inTextSummaryTable::getSummaryStatisticsTable(
data = dataAE,
rowVar = c("AESOC", "AEDECOD"),
dataTotal = dataTotal,
rowOrder = "total",
labelVars = labelVars,
stats = inTextSummaryTable::getStats("count"),

# plotly treemap requires records (rows) for each group
rowVarTotalInclude = "AEDECOD",
outputType = "data.frame-base"
)

dataSunburst <- tableAE
dataSunburst$n <- as.numeric(dataSunburst$n)

# create plot
sunburstClinData(
data = dataSunburst,
vars = c("AESOC", "AEDECOD"),
valueVar = "n",
      valueLab = "Number of patients with adverse events"
)

## example where sum(counts) of child = counts of parent

# counts of patients per arm/site
tableDM <- inTextSummaryTable::getSummaryStatisticsTable(
data = dataDM,
rowVar = c("ARM", "SITEID"),
labelVars = labelVars,
# plotly treemap requires records (rows) for each group
rowVarTotalInclude = "SITEID",
rowTotalInclude = TRUE,
outputType = "data.frame-base"
)
tableDM$statN <- as.numeric(tableDM$statN)

# create the plot
sunburstClinData(
data = tableDM,
vars = c("ARM", "SITEID"),
valueVar = "statN", valueLab = "Counts of patients",
valueType = "total",
caption = "The sectors are colored by category.",
subtitle = "Group: treatment and site"
tableClinData

Create a 'clinical data table', associated to a plot.

Description

Interactive table is created, with the possibility to have clickable link to patient-specific report, and included within a button.

Usage

tableClinData(
  data,
  idVar = "USUBJID",
  idLab = getLabelVar(idVar, labelVars = labelVars),
  keyVar = NULL,
  keyLab = getLabelVar(keyVar, labelVars = labelVars),
  pathVar = NULL,
  pathLab = getLabelVar(pathVar, labelVars = labelVars),
  pathExpand = FALSE,
  tableVars = colnames(data),
  tableLab = getLabelVar(tableVars, labelVars = labelVars),
  tableButton = TRUE,
  tablePars = list(),
  id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
  labelVars = NULL,
  verbose = FALSE
)

Arguments

data          Data.frame with data.
idVar          String with variable containing subject ID.
idLab          String with label for idVar.
keyVar         String with unique key variable, identifying unique group for which the link between the table and the plot should be done.
keyLab         String with label for keyVar.
pathVar        String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

|<a href="./path-to-report">label</a> |
|...
|If multiple, they should be separated by ':'.
The report(s) will be:
timeProfileIntervalPlot

Visualize time intervals across subjects/parameters.

Usage

timeProfileIntervalPlot(
  data,
  paramVar,
  paramLab = getLabelVar(paramVar, labelVars = labelVars),
  paramVarSep = " - ",
  paramGroupVar = NULL,
  timeStartVar,
  ...,
  pathLab,
  pathExpand,
  tableVars
  ...,
  id
  ...,
  labelVars
  ...,
  verbose)

Value

datatable

Author(s)

Laure Cougnaud
timeProfileIntervalPlot

timeStartLab = getLabelVar(timeStartVar, labelVars = labelVars),
timeEndVar,
timeEndLab = getLabelVar(timeEndVar, labelVars = labelVars),
timeStartShapeVar = NULL,
timeStartShapeLab = getLabelVar(timeStartShapeVar, labelVars = labelVars),
timeEndShapeVar = NULL,
timeEndShapeLab = getLabelVar(timeEndShapeVar, labelVars = labelVars),
shapePalette = NULL,
colorVar = NULL,
colorLab = getLabelVar(colorVar, labelVars = labelVars),
colorPalette = NULL,
alpha = 1,
yLab = NULL,
xLab = paste(c(timeStartLab, timeEndLab), collapse = " and "),
title = NULL,
subtitle = NULL,
caption = NULL,
labelVars = NULL,
width = 800,
height = NULL,
hoverVars,
hoverLab,
idVar = "USUBJID",
idLab = getLabelVar(idVar, labelVars = labelVars),
pathVar = NULL,
pathLab = getLabelVar(pathVar, labelVars = labelVars),
id = paste0("plotClinData", sample.int(n = 1000, size = 1)),
selectVars = NULL,
selectLab = getLabelVar(selectVars, labelVars = labelVars),
table = FALSE,
tableVars,
tableLab,
tableButton = TRUE,
tablePars = list(),
watermark = NULL,
verbose = FALSE

Arguments

data Data.frame with data.

paramVar Character vector with variable of data to represent in the y-axis.

paramLab (optional) String with label for paramVar.

paramVarSep (optional) String with separator used to combined paramVar if multiple.

paramGroupVar (optional) Character vector with variable(s) to group/order the paramVar elements in the y-axis.

timeStartVar String with variable with the start of the time interval.
timeProfileIntervalPlot

timeStartLab  (optional) String with label for timeStartVar.
timeEndVar    String with variable with the end of the time interval.
timeEndLab    (optional) String with label for timeEndVar.
timeStartShapeVar (optional) String with variable used for the shape of the start of the time interval.
timeStartShapeLab  (optional) String with label for timeStartShapeVar.
timeEndShapeVar  (optional) String with variable used for the shape of the end of the time interval.
timeEndShapeLab  (optional) String with label for timeEndShapeVar.
shapePalette    (optional) Character vector with shape palette for timeStartShapeVar and timeEndShapeVar.
colorVar       (optional) String with color variable.
colorLab       String with label for colorVar.
colorPalette   (optional) Named character vector with color palette. If not specified, the viridis color palette is used. See clinColors.
alpha          (optional) Numeric with transparency, 1 by default.
xLab, yLab     (optional) String with labels for the x/y-axis.
title          String with title for the plot.
subtitle       String with subtitle. The subtitle is included at the top left of the plot, below the title.
caption        String with caption. The caption is included at the bottom right of the plot. Please note that this might overlap with vertical or rotated x-axis labels.
labelVars      Named character vector containing variable labels.
width          Numeric, width of the plot in pixels, 800 by default.
height         Numeric, height of the plot in pixels, 500 by default.
hoverVars      Character vector with variable(s) to be displayed in the hover, by default any position (and axis) and aesthetic variables displayed in the plot.
hoverLab       Named character vector with labels for hoverVars.
idVar          String with variable containing subject ID.
idLab          String with label for idVar.
pathVar        String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

        <a href="./path-to-report">label</a>

If multiple, they should be separated by ",". The report(s) will be:
• compressed to a zip file and downloaded if the user clicks on the 'p' (a.k.a 'profile') key when hovering on a point of the plot
• included in a collapsible row, and clickable with hyperlinks in the table

pathLab
String with label for pathVar, included in the collapsible row in the table.

id
String with general id for the plot:
• 'id' is used as group for the SharedData
• 'button:[id]' is used as button ID if table is TRUE

If not specified, a random id, as 'plotClinData[X]' is used.

selectVars
(optional) Character vector with variable(s) from data for which a selection box should be included. This enables to select the data displayed in the plot (and associated table).

selectLab
(Named) character vector with label for selectVars.

table
Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. The plot and table are linked when included in a Rmarkdown document: when clicking on an plot element, only the corresponding records are retained in the associated table; when some records are selected in the table, they are highlighted in the associated table.

tableVars
Character vector with variables to be included in the table.

tableLab
Named character vector with labels for each tableVars.

tableButton
Logical, if TRUE (by default) the table is included within an HTML button.

tablePars
List with parameters passed to the getClinDT function.

watermark
(optional) String with path to a file containing a watermark.

verbose
Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.

Value
Either:

• if a table is requested: a clinDataReview object, a.k.a a list with the 'plot' (plotly object) and 'table' (datatable object)
• otherwise: a plotly object

Author(s)
Laure Cougnaud

See Also
Other Clinical data visualization of individual profiles.: scatterplotClinData()
Examples

```r
library(clinUtils)

data(dataADaMCDISCP01)
labelVars <- attr(dataADaMCDISCP01, "labelVars")

dataAE <- dataADaMCDISCP01$ADAE

# basic plot
timeProfileIntervalPlot(
data = dataAE,
paramVar = "USUBJID",
# time-variables
timeStartVar = "ASTDY",
timeEndVar = "ASTDY",
# colored by severity
colorVar = "AESEV",
labelVars = labelVars
)

# add caption & subtitle
timeProfileIntervalPlot(
data = dataAE,
paramVar = "USUBJID",
timeStartVar = "ASTDY",
timeEndVar = "ASTDY",
colorVar = "AESEV",
labelVars = labelVars,
title = "Adverse events",
subtitle = "Time intervals",
caption = "Day is relative to the study baseline"
)

# add a selection box
if(interactive()){t
    timeProfileIntervalPlot(
data = dataAE,
paramVar = "USUBJID",
# time-variables
timeStartVar = "ASTDY",
timeEndVar = "ASTDY",
# colored by severity
colorVar = "AESEV",
labelVars = labelVars,
selectVars = "AEDECOD"
}
```

transformData

**Transform data.**
transformData

Description

Transform data from long to wide format. This function converts formats with the stats::reshape function.

Usage

transformData(data, transformations, verbose = FALSE, labelVars = NULL)

Arguments

data Data.frame with input data to transform.

transformations Transformations (or list of those) as a list with:

• 'type': String with type of transformation. Currently, only: 'pivot_wider' is available
• extra parameters for the transformation, for:
  – 'pivot_wider':
    * 'varsID': Character vector with variable(s) of data defining unique records in the wide format. Corresponds to the idvar parameter of the reshape function.
    * 'varPivot': String with unique variable of data containing elements to pivot in different columns in the wide format (used for column names). Corresponds to the timevar parameter of the reshape function.
    * 'varsValue': Character vector with variable(s) of data used to fill the columns in the wide format. Corresponds to the v.names parameter of the reshape function.

verbose Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.

labelVars Named character vector containing variable labels.

Value

A data.frame in wide format.

Author(s)

Laure Cougnaud
treemapClinData

Treemap interactive plot.

Description

Note: the table and plot are not (yet) linked.

Usage

treemapClinData(...)

Arguments

Arguments passed on to plotCountClinData

colorVar (optional) String with coloring variable (NULL by default). By default, the treemap is colored based by section.

colorRange (optional) Numeric vector of length 2 with range for the color variable, in case it is a numeric variable.

vars Character vector with variables of data containing the groups. If multiple, they should be specified in hierarchical order (from parent to child node).

varsLab Named character vector with labels for vars.

valueVar String with numeric variable of data containing the value to display.

valueLab String with label for the valueVar variable.

valueType String with type of values in valueVar (branchvalues of the plot_ly function), among others: ‘total’ (default, only if sum(child) <= to parent) or ‘relative’.

pathVar String with variable of data containing hyperlinks with path to the subject-specific report, formatted as:

    <a href="./path-to-report">label</a>

If multiple, they should be separated by: ‘, ’.

The report(s) will be:

• compressed to a zip file and downloaded if the user clicks on the ‘p’ (a.k.a ‘profile’) key when hovering on a point of the plot
• included in a collapsible row, and clickable with hyperlinks in the table

pathLab String with label for pathVar, included in the collapsible row in the table.

table Logical, if TRUE (FALSE by default) returns also a datatable containing the plot data. (The plot and the table are not linked.)

data Data.frame with data.

verbose Logical, if TRUE (FALSE by default) progress messages are printed in the current console. For the visualizations, progress messages during download of subject-specific report are displayed in the browser console.
width  Numeric, width of the plot in pixels, 800 by default.
height  Numeric, height of the plot in pixels, 500 by default.
hoverVars  Character vector with variable(s) to be displayed in the hover, by
default any position (and axis) and aesthetic variables displayed in the plot.
hoverLab  Named character vector with labels for hoverVars.
labelVars  Named character vector containing variable labels.
id  String with general id for the plot:
  • 'id' is used as group for the SharedData
  • 'button:[id]' is used as button ID if table is TRUE
If not specified, a random id, as 'plotClinData[X]' is used.
title  String with title for the plot.
titleExtra  String with extra title for the plot (appended after title).
caption  String with caption.
The caption is included at the bottom right of the plot. Please note that this
might overlap with vertical or rotated x-axis labels.
subtitle  String with subtitle.
The subtitle is included at the top left of the plot, below the title.
colorLab  String with label for colorVar.
colorPalette  (optional) Named character vector with color palette. If not
specified, the viridis color palette is used.
See clinColors.
watermark  (optional) String with path to a file containing a watermark.
tableButton  Logical, if TRUE (by default) the table is included within an
HTML button.
tableVars  Character vector with variables to be included in the table.
tableLab  Named character vector with labels for each tableVars.
tablePars  List with parameters passed to the getClinDT function.

Value

Either:

• if a table is requested: a clinDataReview object, a.k.a a list with the 'plot' (plotly object)
  and 'table' (datatable object)
• otherwise: a plotly object

Author(s)

Laure Cougnaud

See Also

Other visualizations of summary statistics for clinical data: barplotClinData(), boxplotClinData(),
errorbarClinData(), plotCountClinData(), sunburstClinData()
Examples

```r
library(clinUtils)

data(dataADaMCIDSCP01)
labelVars <- attr(dataADaMCIDSCP01, "labelVars")

dataDM <- dataADaMCIDSCP01$ADSL
dataAE <- dataADaMCIDSCP01$ADAE

library(plyr)

## basic treemap:
# treemap takes as input table with counts
if (requireNamespace("inTextSummaryTable", quietly = TRUE)) {

# total counts: Safety Analysis Set (patients with start date for the first treatment)
dataTotal <- subset(dataDM, RFSTDTC != "")

# compute adverse event table
tableAE <- inTextSummaryTable::getSummaryStatisticsTable(
  data = dataAE, rowVar = c("AESOC", "AEDECOD"),
  dataTotal = dataTotal, rowOrder = "total",
  labelVars = labelVars, stats = inTextSummaryTable::getStats("count"),
  rowVarTotalInclude = "AEDECOD", outputType = "data.frame-base"
)

dataPlot <- tableAE

dataPlot$n <- as.numeric(dataPlot$n)

# create plot
treemapClinData(
  data = dataPlot, vars = c("AESOC", "AEDECOD"),
  valueVar = "n",
  valueLab = "Number of patients with adverse events"
)

## treemap with coloring
# extract worst-case scenario
dataAE$AESEVN <- as.numeric(factor(dataAE$AESEV, levels = c("MILD", "MODERATE", "SEVERE")))
if(any(is.na(dataAE$AESEVN)))
```

stop("Severity should be filled for all subjects.")

dataAEWC <- ddply(dataAE, c("AESOC", "AEDECOD", "USUBJID"), function(x){
  x[which.max(x$AESEVN), ]
})
dataTotalRow <- list(AEDECOD =
  ddply(dataAEWC, c("AESOC", "USUBJID"), function(x){
    x[which.max(x$AESEVN), ]
  })
)

# compute adverse event table
tableAE <- inTextSummaryTable::getSummaryStatisticsTable(
  data = dataAEWC,
  rowVar = c("AESOC", "AEDECOD"),
  var = "AESEVN",
  dataTotal = dataTotal,
  rowOrder = "total",
  labelVars = labelVars,
  # plotly treemap requires records (rows) for each group
  rowVarTotalInclude = "AEDECOD",
  dataTotalRow = dataTotalRow,
  outputType = "data.frame-base"
)
dataPlot <- tableAE

dataPlot$statN <- as.numeric(dataPlot$statN)
dataPlot$statMean <- as.numeric(dataPlot$statMean)

# create plot
treemapClinData(
  data = dataPlot,
  vars = c("AESOC", "AEDECOD"),
  valueVar = "statN", valueLab = "Number of patients with adverse events",
  colorVar = "statMean", colorLab = "Mean severity"
)

---

**varToFm**

Get formula for a specific variable, to be used in aesthetic specification in plot_ly.

**Description**

Get formula for a specific variable, to be used in aesthetic specification in plot_ly.
**zipClinDataReview**

**Usage**

`varToFm(var)`

**Arguments**

- `var` Character vector with variable to combine. Otherwise with the `+` operator.

**Value**

`as.formula`

**Author(s)**

Laure Cougnaud

---

**zipClinDataReview** Zip the clinical data report

**Description**

Create a zip folder of clinical data reports with a redirect page. The clinical data report out of the `render_clinDataReviewReport` is copied into a new folder. A redirect html page is created to enable the user to navigate the report without needing to look into the new directory.

**Usage**

```r
zipClinDataReview(
  reportDir = "report",
  newDir = "report_dependencies",
  redirectPage = "report.html",
  zipFolder = "report.zip"
)
```

**Arguments**

- `reportDir` String for the path to the directory where the clinical data reports are stored
- `newDir` String for the path where the files from `reportDir` should be copied to.
- `redirectPage` String with the path of the html file that redirects to the "1-introduction.html" page of the report.
- `zipFolder` String with the path to the zipped folder.

**Value**

The zip folder is created in the specified location.
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