

Package ‘psychomics’

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Title Graphical Interface for Alternative Splicing Quantification,
Analysis and Visualisation

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Description Interactive R package with a Shiny-based graphical interface for the quantification, analysis and visualisation of alternative splicing data from The Cancer Genome Atlas (TCGA) or the Genotype-Tissue Expression (GTEx) project. This tool interactively performs survival, principal components and differential splicing analyses with direct incorporation of clinical features (such as tumour stage or survival) associated with TCGA and GTEx samples.

Depends R (>= 3.3), shiny (>= 1.0.0), shinyBS

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LazyData true

RoxygenNote 6.0.1

Imports AnnotationHub, colourpicker, data.table, digest, dplyr, DT (>= 0.2), fastmatch, ggplot2, grDevices, highcharter (>= 0.5.0), htmltools, httr, jsonlite, miscTools, plyr, R.utils, shinyjs, stringr, stats, survival, Sushi, tools, utils, XML, methods

Suggests testthat, knitr, parallel, devtools, rmarkdown, gplots, covr, car

VignetteBuilder knitr

Collate 'analysis.R' 'analysis_diffSplicing.R'
'analysis_diffSplicing_event.R' 'analysis_diffSplicing_table.R'
'analysis_information.R' 'analysis_pca.R' 'analysis_survival.R'
'analysis_template.R' 'utils.R' 'globalAccess.R' 'app.R'
'data.R' 'formats.R' 'data_firebrowse.R' 'data_gtex.R'
'data_inclusionLevels.R' 'data_local.R' 'events_suppa.R'
'events_vastTools.R' 'events_miso.R' 'events_mats.R' 'events.R'
'formats_firebrowseGeneExpression.R'
'formats_firebrowseJunctionReads.R'
'formats_firebrowseMergeClinical.R' 'formats_gtexClinical.R'
'formats_gtexJunctionReads.R' 'formats_gtexSampleInfo.R'
'groups.R' 'help.R'

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Survival, BiomedicalInformatics, Transcriptomics,
Visualization, MultipleComparison

URL <https://github.com/nuno-agostinho/psychomics>

BugReports <https://github.com/nuno-agostinho/psychomics/issues>

NeedsCompilation no

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| | |
|-------------|---|
| addTCGAdata | <i>Creates a UI set with options to add data from TCGA/Firebrowse</i> |
|-------------|---|

Description

Creates a UI set with options to add data from TCGA/Firebrowse

Usage

```
addTCGAdata(ns)
```

Arguments

| | |
|----|--------------------|
| ns | Namespace function |
|----|--------------------|

Value

A UI set that can be added to a UI definition

| | |
|----------------|--------------------------------------|
| analysesServer | <i>Server logic for the analyses</i> |
|----------------|--------------------------------------|

Description

Server logic for the analyses

Usage

```
analysesServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|---|
| analysesUI | <i>User interface for the data analyses</i> |
|------------|---|

Description

User interface for the data analyses

Usage

```
analysesUI(id, tab)
```

Arguments

| | |
|-----|-----------------------------------|
| id | Character: identifier |
| tab | Function to process HTML elements |

Value

HTML element as character

| | |
|-----------|------------------------|
| appServer | <i>Server function</i> |
|-----------|------------------------|

Description

Instructions to build the Shiny app.

Usage

```
appServer(input, output, session)
```

Arguments

| | |
|---------|----------------|
| input | Input object |
| output | Output object |
| session | Session object |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-------|---|
| appUI | <i>The user interface (ui) controls the layout and appearance of the app All the CSS modifications are in the file "shiny/www/styles.css"</i> |
|-------|---|

Description

The user interface (ui) controls the layout and appearance of the app All the CSS modifications are in the file "shiny/www/styles.css"

Usage

```
appUI()
```

Value

HTML elements

| | |
|-----------|---|
| articleUI | <i>Return the interface to display an article</i> |
|-----------|---|

Description

Return the interface to display an article

Usage

```
articleUI(article)
```

Arguments

| | |
|---------|----------------|
| article | PubMed article |
|---------|----------------|

Value

HTML to render an article's interface

| | |
|------------|---|
| basicStats | <i>Basic statistics performed on data</i> |
|------------|---|

Description

Variance and median of each group. If data has 2 groups, also calculates the delta variance and delta median.

Usage

```
basicStats(psi, groups)
```

Arguments

| | |
|--------|---|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |

Value

HTML elements

| | |
|----------------|----------------------------------|
| browserHistory | <i>Enable history navigation</i> |
|----------------|----------------------------------|

Description

Navigate app according to the location given by the navigation bar. Code and logic adapted from <https://github.com/daattali/advanced-shiny/blob/master/navigate-history>

Usage

```
browserHistory(navId, input, session)
```

Arguments

| | |
|---------|---|
| navId | Character: identifier of the navigation bar |
| input | Input object |
| session | Session object |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|----------|---|
| bsModal2 | <i>Modified version of shinyBS::bsModal</i> |
|----------|---|

Description

bsModal is used within the UI to create a modal window. This allows to use the footer.

Usage

```
bsModal2(id, title, trigger, ..., size = NULL, footer = NULL,
  style = NULL)
```

Arguments

| | |
|---------|--|
| id | A unique identifier for the modal window |
| title | The title to appear at the top of the modal |
| trigger | The id of a button or link that will open the modal. |
| ... | UI elements to include within the modal |
| size | Character: Modal size ("small", "default" or "large") |
| footer | UI set: List of elements to include in the footer |
| style | Character: message style can be "warning", "error", "info" or NULL |

Value

HTML element to create a modified modal

calculateInclusionLevels

Calculate inclusion levels using alternative splicing event annotation and junction quantification for many samples

Description

Calculate inclusion levels using alternative splicing event annotation and junction quantification for many samples

Usage

```
calculateInclusionLevels(eventType, junctionQuant, annotation, minReads = 10)
```

Arguments

| | |
|---------------|--|
| eventType | Character: type of the alternative event to calculate |
| junctionQuant | Data.frame: junction quantification with samples as columns and junctions as rows |
| annotation | Data.frame: alternative splicing annotation related to event type |
| minReads | Integer: minimum of total reads required to consider the quantification as valid (10 by default) |

Value

Matrix with inclusion levels

checkFileFormat

Checks the format of a file

Description

Checks the format of a file

Usage

```
checkFileFormat(format, head, filename)
```

Arguments

| | |
|----------|---------------------------------------|
| format | Environment: format of the file |
| head | Data.frame: head of the file to check |
| filename | Character: name of the file |

Details

The name of the file may also be required to be considered of a certain format.

Value

TRUE if the file is of the given format; otherwise, returns FALSE

| | |
|-----------------|---|
| checkFirebrowse | <i>Return an user interface depending on the status of the Firebrowse API</i> |
|-----------------|---|

Description

If the API is working, it'll be loaded. Else, a message will appear warning the user that the API is down and that will let check again if the API is back online.

Usage

```
checkFirebrowse(ns)
```

Arguments

| | |
|----|--------------------|
| ns | Namespace function |
|----|--------------------|

Value

HTML elements

| | |
|----------------|--|
| checkIntegrity | <i>Compute the 32-byte MD5 hashes of one or more files and check with given md5 file</i> |
|----------------|--|

Description

Compute the 32-byte MD5 hashes of one or more files and check with given md5 file

Usage

```
checkIntegrity(filesToCheck, md5file)
```

Arguments

| | |
|--------------|--|
| filesToCheck | Character: files to calculate and match MD5 hashes |
| md5file | Character: file containing correct MD5 hashes |

Value

Logical vector showing TRUE for files with matching md5sums and FALSE for files with non-matching md5sums

checkSurvivalInput *Prepare survival terms in case of valid input*

Description

Prepare survival terms in case of valid input

Usage

```
checkSurvivalInput(session, input, coxph = FALSE)
```

Arguments

| | |
|---------|--|
| session | Shiny session |
| input | Shiny input |
| coxph | Boolean: prepare data for Cox models? FALSE by default |

Value

NULL (this function is used to modify the Shiny session's state)

closeProgress *Close the progress even if there's an error*

Description

Close the progress even if there's an error

Usage

```
closeProgress(message = NULL, global = sharedData)
```

Arguments

| | |
|---------|--|
| message | Character: message to show in progress bar |
| global | Global Shiny variable where all data is stored |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|---------------|--|
| createDataTab | <i>Render a specific data tab (including data table and related interface)</i> |
|---------------|--|

Description

Render a specific data tab (including data table and related interface)

Usage

```
createDataTab(index, data, name, input, output)
```

Arguments

| | |
|--------|--|
| index | Integer: index of the data to load |
| data | Data frame: data with everything to load |
| name | Character: name of the dataset |
| input | Shiny session input |
| output | Shiny session output |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-------------------------|---|
| createDensitySparklines | <i>Create density sparklines for inclusion levels</i> |
|-------------------------|---|

Description

Create density sparklines for inclusion levels

Usage

```
createDensitySparklines(data, events, delim = NULL)
```

Arguments

| | |
|--------|---|
| data | Character: HTML-formatted data series of interest |
| events | Character: event identifiers |
| delim | Character: left and right delimiters in groups that should be removed |

Value

HTML element with sparkline data (character)

createEventPlotting *Create plot for events*

Description

Create plot for events

Usage

```
createEventPlotting(df, x, y, params, highlightX, highlightY, highlightParams,
  selected, selectedParams, xlim, ylim)
```

Arguments

| | |
|-----------------|--|
| df | Data frame |
| x | Character: name of the variable used for the X axis |
| y | Character: name of the variable used for the Y axis |
| params | List of parameters to pass to geom_point related to most points |
| highlightX | Integer: region of points in X axis to highlight |
| highlightY | Integer: region of points in Y axis to highlight |
| highlightParams | List of parameters to pass to geom_point related to highlighted points |
| selected | Integer: index of rows/points to be coloured |
| selectedParams | List of parameters to pass to geom_point related to selected points |
| xlim | Numeric: limits of X axis |
| ylim | Numeric: limits of Y axis |

Value

HTML elements

createGroup *Prepare to create group according to specific details*

Description

Prepare to create group according to specific details

Usage

```
createGroup(session, input, output, id, type)
```

Arguments

| | |
|---------|--|
| session | Shiny session |
| input | Shiny input |
| output | Shiny output |
| id | Character: identifier of the group selection |
| type | Character: type of group to create |

Value

NULL (this function is used to modify the Shiny session's state)

createGroupByAttribute

Create groups with the indexes from the unique values of a given column from a dataset

Description

Create groups with the indexes from the unique values of a given column from a dataset

Usage

```
createGroupByAttribute(col, dataset)
```

Arguments

| | |
|---------|-------------------------------|
| col | Character: column name |
| dataset | Matrix or data frame: dataset |

Value

Named list with the indexes of each unique value from a given column

Examples

```
df <- data.frame(gender=c("male", "female"),
                 stage=paste("stage", c(1, 3, 1, 4, 2, 3, 2, 2)))
createGroupByAttribute(col="stage", dataset=df)
```

`createGroupByColumn` *Create groups with the indexes from the unique values of a given column from a dataset*

Description

Create groups with the indexes from the unique values of a given column from a dataset

Usage

```
createGroupByColumn(col, dataset)
```

Arguments

| | |
|----------------------|-------------------------------|
| <code>col</code> | Character: column name |
| <code>dataset</code> | Matrix or data frame: dataset |

Value

Named list with the indexes of each unique value from a given column

`createGroupById` *Create groups from a given string of rows*

Description

Create groups from a given string of rows

Usage

```
createGroupById(session, rows, dataset, identifiers)
```

Arguments

| | |
|--------------------------|--------------------------------------|
| <code>session</code> | Shiny session |
| <code>rows</code> | Character: rows separated by a comma |
| <code>dataset</code> | Matrix or data frame: dataset |
| <code>identifiers</code> | Character: available identifiers |

Value

NULL (this function is used to modify the Shiny session's state)

createGroupFromInput *Set new groups according to the user input*

Description

Set new groups according to the user input

Usage

```
createGroupFromInput(session, input, output, dataset, id, type)
```

Arguments

| | |
|---------|--|
| session | Shiny session |
| input | Shiny input |
| output | Shiny output |
| dataset | Data frame or matrix: dataset of interest |
| id | Character: identifier of the group selection |
| type | Character: type of group to create |

Value

Matrix with the group names and respective indexes

createJunctionsTemplate
Creates a template of alternative splicing junctions

Description

Creates a template of alternative splicing junctions

Usage

```
createJunctionsTemplate(nrow, program = character(0),
  event.type = character(0), chromosome = character(0),
  strand = character(0), id = character(0))
```

Arguments

| | |
|------------|---|
| nrow | Integer: Number of rows |
| program | Character: Program used to get the junctions |
| event.type | Character: Event type of the respective events |
| chromosome | Character: Chromosome of the junctions |
| strand | Character: positive ("+") or negative ("-") strand of the event |
| id | Character: events' ID |

Value

A data frame with the junctions coordinate names pre-filled with NAs

Examples

```
psichomics::createJunctionsTemplate(nrow = 8)
```

| | |
|---------------|---|
| createTooltip | <i>Create the interface for the tooltip of a plot</i> |
|---------------|---|

Description

Create the interface for the tooltip of a plot

Usage

```
createTooltip(df, hover, x, y)
```

Arguments

| | |
|-------|--|
| df | Data frame |
| hover | Mouse hover information for a given plot as retrieved from hoverOpts |
| x | Character: name of the variable used for the X axis |
| y | Character: name of the variable used for the Y axis |

Value

HTML elements

| | |
|------------|--|
| dataServer | <i>Server logic of the data module</i> |
|------------|--|

Description

Server logic of the data module

Usage

```
dataServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

Part of the server logic related to this tab

| | |
|--------|--|
| dataUI | <i>User interface of the data module</i> |
|--------|--|

Description

User interface of the data module

Usage

```
dataUI(id, tab)
```

Arguments

| | |
|-----|------------------------|
| id | Character: identifier |
| tab | Function to create tab |

Value

HTML elements

| | |
|--------------|--|
| diffAnalyses | <i>Perform selected statistical analyses on multiple splicing events</i> |
|--------------|--|

Description

Perform selected statistical analyses on multiple splicing events

Usage

```
diffAnalyses(psi, groups = NULL, analyses = c("wilcoxRankSum", "ttest",
      "kruskal", "levene", "fligner"), pvalueAdjust = "BH",
      progress = echoProgress)
```

Arguments

| | |
|--------------|---|
| psi | Data frame or matrix: alternative splicing event quantification |
| groups | Character: group of each sample from the alternative splicing event quantification (if NULL, sample types are used instead, e.g. normal, tumour and metastasis) |
| analyses | Character: analyses to perform (see Details) |
| pvalueAdjust | Character: method used to adjust p-values (see Details) |
| progress | Function to track the progress |

Details

The following statistical analyses may be performed by including the respective string in the `analysis` argument:

- `ttest` - Unpaired t-test (2 groups)
- `wilcoxRankSum` - Wilcoxon Rank Sum test (2 groups)
- `kruskal` - Kruskal test (2 or more groups)
- `levene` - Levene's test (2 or more groups)
- `fligner` - Fligner-Killeen test (2 or more groups)
- `density` - Sample distribution per group (only usable through the visual interface)

The following methods for p-value adjustment are supported by using the respective string in the `pvalueAdjust` argument:

- `none`: do not adjust p-values
- `BH`: Benjamini-Hochberg's method (false discovery rate)
- `BY`: Benjamini-Yekutieli's method (false discovery rate)
- `bonferroni`: Bonferroni correction (family-wise error rate)
- `holm`: Holm's method (family-wise error rate)
- `hochberg`: Hochberg's method (family-wise error rate)
- `hommel`: Hommel's method (family-wise error rate)

Value

Table of statistical analyses

Examples

```
# Calculate PSI for skipped exon (SE) and mutually exclusive (MXE) events
eventType <- c("SE", "MXE")
annot <- readRDS("ex_splicing_annotation.RDS")
junctionQuant <- readRDS("ex_junctionQuant.RDS")

psi <- quantifySplicing(annot, junctionQuant, eventType=c("SE", "MXE"))
group <- c(rep("Normal", 3), rep("Tumour", 3))
diffAnalyses(psi, group)
```

diffSplicingEventServer

Server logic for the analyses of a single alternative splicing event

Description

Server logic for the analyses of a single alternative splicing event

Usage

```
diffSplicingEventServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

diffSplicingEventUI *Interface for the analysis of an alternative splicing event*

Description

Interface for the analysis of an alternative splicing event

Usage

```
diffSplicingEventUI(id)
```

Arguments

| | |
|----|-----------------------|
| id | Character: identifier |
|----|-----------------------|

Value

Character with the HTML interface

diffSplicingServer *Server logic for the differential splicing analyses*

Description

Server logic for the differential splicing analyses

Usage

```
diffSplicingServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

diffSplicingTableServer

Server logic of the exploratory differential analyses

Description

Server logic of the exploratory differential analyses

Usage

```
diffSplicingTableServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

diffSplicingTableUI *Interface for differential analyses on all splicing events*

Description

Interface for differential analyses on all splicing events

Usage

```
diffSplicingTableUI(id)
```

Arguments

| | |
|----|-----------------------|
| id | Character: identifier |
|----|-----------------------|

Value

HTML elements

| | |
|----------------|--|
| diffSplicingUI | <i>User interface for the differential splicing analyses</i> |
|----------------|--|

Description

User interface for the differential splicing analyses

Usage

```
diffSplicingUI(id, tab)
```

Arguments

| | |
|-----|-----------------------------------|
| id | Character: identifier |
| tab | Function to process HTML elements |

Value

HTML element as character

| | |
|------------|--------------------------------------|
| disableTab | <i>Disable a tab from the navbar</i> |
|------------|--------------------------------------|

Description

Disable a tab from the navbar

Usage

```
disableTab(tab)
```

Arguments

| | |
|-----|---------------------------|
| tab | Character: tab to disable |
|-----|---------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|---------------|--|
| downloadFiles | <i>Download files to a given directory</i> |
|---------------|--|

Description

Download files to a given directory

Usage

```
downloadFiles(url, folder, progress = echoProgress,
              download = download.file, ...)
```

Arguments

| | |
|----------|---|
| url | Character: download links |
| folder | Character: directory to store the downloaded archives |
| progress | Function to show the progress (default is to print progress to console) |
| download | Function to use to download files |
| ... | Extra parameters passed to the download function |

Value

Invisible TRUE if every file was successfully downloaded

Examples

```
## Not run:
url <- paste0("https://unsplash.it/400/300/?image=", 570:572)
downloadFiles(url, "~/Pictures")

# Download without printing to console
downloadFiles(url, "~/Pictures", quiet = TRUE)

## End(Not run)
```

| | |
|--------------|---|
| echoProgress | <i>Echo progress to console using cat</i> |
|--------------|---|

Description

Echo progress to console using cat

Usage

```
echoProgress(..., console = TRUE)
```

Arguments

| | |
|---------|--|
| ... | Strings to print to console |
| console | Boolean: print to console? TRUE by default |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-----------|-------------------------------------|
| enableTab | <i>Enable a tab from the navbar</i> |
|-----------|-------------------------------------|

Description

Enable a tab from the navbar

Usage

```
enableTab(tab)
```

Arguments

| | |
|-----|--------------------------|
| tab | Character: tab to enable |
|-----|--------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|--|
| endProcess | <i>Signal the program that a process has ended</i> |
|------------|--|

Description

Style button to show processing is not occurring. Also, close the progress bar (if TRUE) and print the difference between the current time and a given time (if given time is not NULL)

Usage

```
endProcess(id, time = NULL, closeProgressBar = TRUE)
```

Arguments

| | |
|------------------|--|
| id | Character: button identifier |
| time | POSIXct: start time needed to show the interval time (if NULL, the time interval is not displayed) |
| closeProgressBar | Boolean: close progress bar? TRUE by default |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------------|---|
| ensemblToUniprot | <i>Convert a protein's Ensembl identifier to UniProt identifier</i> |
|------------------|---|

Description

Convert a protein's Ensembl identifier to UniProt identifier

Usage

```
ensemblToUniprot(protein)
```

Arguments

| | |
|---------|---------------------------------------|
| protein | Character: Ensembl protein identifier |
|---------|---------------------------------------|

Value

UniProt protein identifier

Examples

```
ensemblToUniprot("ENSP00000445929")
```

| | |
|-------------|---|
| errorDialog | <i>Error alert in the style of a dialog box with a button</i> |
|-------------|---|

Description

Error alert in the style of a dialog box with a button

Usage

```
errorDialog(description, ..., buttonLabel = NULL, buttonIcon = NULL,  
            buttonId = NULL, id = NULL)
```

Arguments

| | |
|-------------|--|
| description | Character: description |
| ... | Extra parameters when creating the alert |
| buttonLabel | Character: button label (NULL by default) |
| buttonIcon | Character: button icon (NULL by default) |
| buttonId | Character: button identifier (NULL by default) |
| id | Character: identifier (NULL by default) |

Value

HTML elements

| | |
|--------|--|
| escape | <i>Escape symbols for use in regular expressions</i> |
|--------|--|

Description

Escape symbols for use in regular expressions

Usage

```
escape(...)
```

Arguments

... Characters to be pasted with no space

Value

Escaped string

| | |
|------------------|-----------------------------------|
| eventPlotOptions | <i>Options for event plotting</i> |
|------------------|-----------------------------------|

Description

Options for event plotting

Usage

```
eventPlotOptions(session, df)
```

Arguments

session Shiny session

df Data frame

Value

HTML elements

`export_highcharts` *Add an exporting feature to a highcharts object*

Description

Add an exporting feature to a highcharts object

Usage

```
export_highcharts(hc, fill = "transparent", text = "Export")
```

Arguments

| | |
|-------------------|------------------------|
| <code>hc</code> | A highcharts object |
| <code>fill</code> | Character: colour fill |
| <code>text</code> | Character: button text |

Value

A highcharts object with an export button

`filterGroups` *Filter groups with less data points than the threshold*

Description

Groups containing a number of non-missing values less than the threshold are discarded.

Usage

```
filterGroups(vector, group, threshold = 1)
```

Arguments

| | |
|------------------------|--|
| <code>vector</code> | Unnamed elements |
| <code>group</code> | Character: group of the elements |
| <code>threshold</code> | Integer: number of valid non-missing values by group |

Value

Named vector with filtered elements from valid groups. The group of the respective element is given in the name.

Examples

```
# Removes groups with less than two elements
filterGroups(1:4, c("A", "B", "B", "D"), threshold=2)
```

| | |
|--------------|---|
| firebrowseUI | <i>User interface of the TCGA/Firebrowse loader</i> |
|--------------|---|

Description

User interface of the TCGA/Firebrowse loader

Usage

```
firebrowseUI(id, panel)
```

Arguments

| | |
|-------|-------------------------------|
| id | Character: identifier |
| panel | Function to enclose interface |

Value

HTML of the interface

| | |
|--------|---|
| fisher | <i>Perform Fisher's exact test and return interface to show the results</i> |
|--------|---|

Description

Perform Fisher's exact test and return interface to show the results

Usage

```
fisher(psi, groups)
```

Arguments

| | |
|--------|---|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |

Value

HTML elements

| | |
|---------|--|
| fligner | <i>Perform Fligner-Killeen test and return interface to show the results</i> |
|---------|--|

Description

Perform Fligner-Killeen test and return interface to show the results

Usage

```
fligner(psi, groups, stat = NULL)
```

Arguments

| | |
|--------|--|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |
| stat | Data frame or matrix: values of the analyses to be performed (if NULL, the analyses will be performed) |

Value

HTML elements

| | |
|------------------|-----------------------------|
| getActiveDataset | <i>Get selected dataset</i> |
|------------------|-----------------------------|

Description

Get selected dataset

Usage

```
getActiveDataset()
```

Value

List of data frames

| | |
|--------------------|--|
| getAssemblyVersion | <i>Get the assembly version of a data category</i> |
|--------------------|--|

Description

Get the assembly version of a data category

Usage

```
getAssemblyVersion(category = getCategory())
```

Arguments

| | |
|----------|--|
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |
|----------|--|

Value

Character value with the assembly version

Note

Needs to be called inside a reactive function

| | |
|-------------------|---|
| getAutoNavigation | <i>Get if history browsing is automatic</i> |
|-------------------|---|

Description

Get if history browsing is automatic

Usage

```
getAutoNavigation()
```

Value

Boolean: is navigation of browser history automatic?

| | |
|---------------|--------------------------------------|
| getCategories | <i>Get available data categories</i> |
|---------------|--------------------------------------|

Description

Get available data categories

Usage

```
getCategories()
```

Value

Name of all data categories

| | |
|-------------|-----------------------------------|
| getCategory | <i>Get selected data category</i> |
|-------------|-----------------------------------|

Description

Get selected data category

Usage

```
getCategory()
```

Value

Name of selected data category

| | |
|-----------------|---|
| getCategoryData | <i>Get data of selected data category</i> |
|-----------------|---|

Description

Get data of selected data category

Usage

```
getCategoryData()
```

Value

If category is selected, returns the respective data as a data frame; otherwise, returns NULL

| | |
|-----------------|---|
| getClinicalData | <i>Get clinical data of the data category</i> |
|-----------------|---|

Description

Get clinical data of the data category

Usage

```
getClinicalData()
```

Value

Data frame with clinical data

| | |
|----------------------|--|
| getClinicalMatchFrom | <i>Get clinical matches from a given data type</i> |
|----------------------|--|

Description

Get clinical matches from a given data type

Usage

```
getClinicalMatchFrom(dataset, category = getCategory())
```

Arguments

| | |
|----------|--|
| dataset | Character: data set (e.g. "Junction quantification") |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

Integer with clinical matches to a given dataset

Note

Needs to be called inside a reactive function

| | |
|----------------|--|
| getColumnsTime | <i>Retrieve the time for given columns in a clinical dataset</i> |
|----------------|--|

Description

Retrieve the time for given columns in a clinical dataset

Usage

```
getColumnsTime(clinical, event, timeStart, timeStop = NULL,
               followup = "days_to_last_followup")
```

Arguments

| | |
|-----------|--|
| clinical | Data frame: clinical data |
| event | Character: name of column containing time of the event of interest |
| timeStart | Character: name of column containing starting time of the interval or follow up time |
| timeStop | Character: name of column containing ending time of the interval |
| followup | Character: name of column containing follow up time |

Value

Data frame containing the time for the given columns

| | |
|----------|-----------------------------------|
| getCores | <i>Get number of cores to use</i> |
|----------|-----------------------------------|

Description

Get number of cores to use

Usage

```
getCores()
```

Value

Numeric value with the number of cores to use

| | |
|---------|------------------------|
| getData | <i>Get global data</i> |
|---------|------------------------|

Description

Get global data

Usage

```
getData()
```

Value

Variable containing all data of interest

| | |
|-------------|---|
| getDataRows | <i>Get rows of a data frame between two row indexes</i> |
|-------------|---|

Description

Get rows of a data frame between two row indexes

Usage

```
getDataRows(i, data, firstRow, lastRow)
```

Arguments

| | |
|-----------------------|---|
| <code>i</code> | Integer: current iteration |
| <code>data</code> | Data.frame: contains the data of interest |
| <code>firstRow</code> | Vector of integers: First row index of interest; value must be less than the respective last row index and less than the number of rows in the data frame |
| <code>lastRow</code> | Vector of integers: Last row index of interest; value must be higher than the respective first row index and less than the number of rows in the data frame |

Details

For a given iteration `i`, returns data from `firstRow[i]` to `lastRow[i]`

Value

Data frame subset from two row indexes (returns NA if the first row index is NA)

`getDifferentialAnalyses`*Get the table of differential analyses of a data category*

Description

Get the table of differential analyses of a data category

Usage

```
getDifferentialAnalyses(category = getCategory())
```

Arguments

`category` Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Data frame of differential analyses

Note

Needs to be called inside a reactive function

`getDifferentialAnalysesFiltered`*Get the filtered table of differential analyses of a data category*

Description

Get the filtered table of differential analyses of a data category

Usage

```
getDifferentialAnalysesFiltered(category = getCategory())
```

Arguments

`category` Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Filtered data frame of differential analyses

Note

Needs to be called inside a reactive function

getDifferentialAnalysesHighlightedEvents

Get highlighted events from differential analyses of a data category

Description

Get highlighted events from differential analyses of a data category

Usage

```
getDifferentialAnalysesHighlightedEvents(category = getCategory())
```

Arguments

category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Integer of indexes relative to a table of differential analyses

Note

Needs to be called inside a reactive function

getDifferentialAnalysesSelected

Get selected points in the differential analysis table of a data category

Description

Get selected points in the differential analysis table of a data category

Usage

```
getDifferentialAnalysesSelected(category = getCategory())
```

Arguments

category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Integer containing index of selected points

Note

Needs to be called inside a reactive function

`getDifferentialAnalysesSurvival`*Get the table of differential analyses' survival data of a data category*

Description

Get the table of differential analyses' survival data of a data category

Usage

```
getDifferentialAnalysesSurvival(category = getCategory())
```

Arguments

`category` Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Data frame of differential analyses' survival data

Note

Needs to be called inside a reactive function

`getDifferentialAnalysesZoom`*Get plot coordinates for zooming from differential analyses of a data category*

Description

Get plot coordinates for zooming from differential analyses of a data category

Usage

```
getDifferentialAnalysesZoom(category = getCategory())
```

Arguments

`category` Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Integer of X and Y axes coordinates

Note

Needs to be called inside a reactive function

`getDownloadsFolder` *Get the Downloads folder of the user*

Description

Get the Downloads folder of the user

Usage

`getDownloadsFolder()`

Value

Path to Downloads folder

Examples

`getDownloadsFolder()`

`getEvent` *Get selected alternative splicing event's identifier*

Description

Get selected alternative splicing event's identifier

Usage

`getEvent()`

Value

Alternative splicing event's identifier as a string

`getFirebrowseCohorts` *Query the Firebrowse web API for the cohorts available*

Description

Query the Firebrowse web API for the cohorts available

Usage

`getFirebrowseCohorts(cohort = NULL)`

`getFirehoseCohorts(cohort = NULL)`

Arguments

cohort Character: filter by given cohorts (optional)

Value

Character with cohort abbreviations (as values) and description (as names)

Examples

```
if (isFirebrowseUp()) getFirebrowseCohorts()
```

```
getFirebrowseDataTypes
```

Get data types available from Firebrowse

Description

Get data types available from Firebrowse

Usage

```
getFirebrowseDataTypes()
```

```
getFirehoseDataTypes()
```

Value

Named character vector

Examples

```
getFirebrowseDataTypes()
```

```
getFirebrowseDateFormat
```

Returns the date format used by the Firebrowse web API

Description

Returns the date format used by the Firebrowse web API

Usage

```
getFirebrowseDateFormat()
```

Value

Named list with Fireweb web API's date formats

Examples

```
format <- psychomics::getFirebrowseDateFormat()

# date format to use in a query to Firebrowse web API
format$query

# date format to parse a date in a response from Firebrowse web API
format$response
```

| | |
|--------------------|---|
| getFirebrowseDates | <i>Query the Firebrowse web API for the timestamps of the data available and parse the response</i> |
|--------------------|---|

Description

Query the Firebrowse web API for the timestamps of the data available and parse the response

Usage

```
getFirebrowseDates()

getFirehoseDates()
```

Value

Date with timestamps of the data available

Examples

```
if (isFirebrowseUp()) getFirebrowseDates()
```

| | |
|-----------|----------------------------------|
| getGlobal | <i>Get data from global data</i> |
|-----------|----------------------------------|

Description

Get data from global data

Usage

```
getGlobal(..., sep = "_")
```

Arguments

| | |
|-----|-----------------------------------|
| ... | Arguments to identify a variable |
| sep | Character to separate identifiers |

Value

Data from global data

`getGroupsFrom` *Get groups from a given data type*

Description

Get groups from a given data type

Usage

```
getGroupsFrom(dataset, category = getCategory(), complete = FALSE,
              samples = FALSE)
```

Arguments

| | |
|-----------------------|---|
| <code>dataset</code> | Character: data set (e.g. "Clinical data") |
| <code>category</code> | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |
| <code>complete</code> | Boolean: return all the information on groups (TRUE) or just the group names and respective indexes (FALSE)? FALSE by default |
| <code>samples</code> | Boolean: show groups by samples (TRUE) or patients (FALSE)? FALSE by default |

Value

Matrix with groups of a given dataset

Note

Needs to be called inside a reactive function

`getInclusionLevels` *Get alternative splicing quantification of the selected data category*

Description

Get alternative splicing quantification of the selected data category

Usage

```
getInclusionLevels()
```

Value

Data frame with the alternative splicing quantification

getInclusionLevelsPCA *Get principal component analysis based on inclusion levels*

Description

Get principal component analysis based on inclusion levels

Usage

```
getInclusionLevelsPCA(category = getCategory())
```

Arguments

category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

prcomp object (PCA) of inclusion levels

Note

Needs to be called inside a reactive function

getJunctionQuantification
Get junction quantification data

Description

Get junction quantification data

Usage

```
getJunctionQuantification(category = getCategory())
```

Arguments

category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

List of data frames of junction quantification

Note

Needs to be called inside a reactive function

| | |
|--------------------|--|
| getMatchingSamples | <i>Search samples in the clinical dataset and return the ones matching the given index</i> |
|--------------------|--|

Description

Search samples in the clinical dataset and return the ones matching the given index

Usage

```
getMatchingSamples(index, samples, clinical, rm.NA = TRUE, match = NULL,
  showMatch = FALSE)
```

Arguments

| | |
|-----------|--|
| index | Numeric or list of numeric: patient row indexes |
| samples | Character: samples |
| clinical | Data frame or matrix: clinical dataset |
| rm.NA | Boolean: remove NAs? TRUE by default |
| match | Integer: vector of patient index with the sample identifiers as name to save time (optional) |
| showMatch | Boolean: show matching patient index? FALSE by default |

Value

Names of the matching samples (if showMatch is TRUE, a integer vector with the patient index and the matching samples as names is shown)

Examples

```
patients <- c("GTEX-ABC", "GTEX-DEF", "GTEX-GHI", "GTEX-JKL", "GTEX-MNO")
samples <- paste0(patients, "-sample")
clinical <- data.frame(samples=samples)
rownames(clinical) <- patients
getMatchingSamples(c(1, 4), samples, clinical)
```

| | |
|-------------|---|
| getNumerics | <i>Convert a column to numeric if possible and ignore given columns composed of lists</i> |
|-------------|---|

Description

Convert a column to numeric if possible and ignore given columns composed of lists

Usage

```
getNumerics(table, by = NULL, toNumeric = FALSE)
```

Arguments

| | |
|-----------|---|
| table | Data matrix: table |
| by | Character: column names of interest |
| toNumeric | Boolean: which columns to convert to numeric (FALSE by default) |

Value

Processed data matrix

Examples

```
event <- read.table(text = "ABC123 + 250 300 350
                          DEF456 - 900 800 700")
names(event) <- c("Event ID", "Strand", "C1.end", "A1.end", "A1.start")

# Let's change one column to character
event[ , "C1.end"] <- as.character(event[ , "C1.end"])
is.character(event[ , "C1.end"])

event <- psychomics::getNumerics(event, by = c("Strand", "C1.end", "A1.end",
                                              "A1.start"),
                                toNumeric = c(FALSE, TRUE, TRUE, TRUE))

# Let's check if the same column is now integer
is.numeric(event[ , "C1.end"])
```

getPatientFromSample *Match given sample identifiers and return the respective row in clinical data*

Description

Match given sample identifiers and return the respective row in clinical data

Usage

```
getPatientFromSample(sampleId, patientId)
```

Arguments

| | |
|-----------|--|
| sampleId | Character: sample identifiers |
| patientId | Character: clinical patient identifiers (if a matrix or data frame is given, its row-names will be retrieved as patient identifiers) |

Value

Integer vector of the row number in clinical data corresponding to the given IDs (named with the ID)

Examples

```
patients <- c("GTEX-ABC", "GTEX-DEF", "GTEX-GHI", "GTEX-JKL", "GTEX-MNO")
samples <- paste0(patients, "-sample")
getPatientFromSample(samples, patients)
```

getPatientId *Get the identifier of patients for a given category*

Description

Get the identifier of patients for a given category

Usage

```
getPatientId(category = getCategory())
```

Arguments

category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Character vector with identifier of patients

Note

Needs to be called inside a reactive function

getPrecision *Get number of decimal places*

Description

Get number of decimal places

Usage

```
getPrecision()
```

Value

Numeric value regarding the number of decimal places

| | |
|------------------|--|
| getPSIperPatient | <i>Assign alternative splicing quantification to patients based on their samples</i> |
|------------------|--|

Description

Match filtered samples with clinical patients to retrieve alternative splicing quantification per clinical patient. Only one sample can be matched with one patient. Normal and control samples are filtered out by default.

Usage

```
getPSIperPatient(psi, match, clinical, pattern = c("Normal", "Control"),
  filterOut = TRUE)
```

Arguments

| | |
|-----------|--|
| psi | Data frame or matrix: alternative splicing quantification per samples |
| match | Matrix: match between samples and clinical patients |
| clinical | Data frame or matrix: clinical dataset |
| pattern | Character: pattern to use when filtering sample types (normal and control samples are filtered by default) |
| filterOut | Boolean: filter out (TRUE) or filter in (FALSE) samples with the given pattern; by default, filter out |

Value

Alternative splicing quantification per clinical patients

| | |
|-------------|---|
| getSampleId | <i>Get the identifier of samples for a given category</i> |
|-------------|---|

Description

Get the identifier of samples for a given category

Usage

```
getSampleId(category = getCategory())
```

Arguments

| | |
|----------|--|
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |
|----------|--|

Value

Character vector with identifier of samples

Note

Needs to be called inside a reactive function

| | |
|---------------|---|
| getSampleInfo | <i>Get sample information of the selected data category</i> |
|---------------|---|

Description

Get sample information of the selected data category

Usage

```
getSampleInfo()
```

Value

Data frame with sample information

| | |
|-------------------|--|
| getSelectedGroups | <i>Get selected groups for a given group selection element</i> |
|-------------------|--|

Description

Get selected groups for a given group selection element

Usage

```
getSelectedGroups(input, id, samples = FALSE, dataset = "Clinical data",
  filter = NULL)
```

Arguments

| | |
|---------|--|
| input | Shiny input |
| id | Character: identifier of the group selection element |
| samples | Boolean: show groups by samples (TRUE) or patients (FALSE)? FALSE by default |
| dataset | Character: data set (e.g. "Clinical data") |
| filter | Character: only get groups passed |

Value

List with selected groups (or NULL if no groups were selected)

| | |
|--------------------|---|
| getServerFunctions | <i>Matches server functions from a given loader</i> |
|--------------------|---|

Description

Matches server functions from a given loader

Usage

```
getServerFunctions(loader, ..., priority = NULL)
```

Arguments

| | |
|----------|--|
| loader | Character: loader to run the functions |
| ... | Extra arguments to pass to server functions |
| priority | Character: name of functions to prioritise by the given order; for instance, c("data", "analyses") would load "data", then "analyses" then remaining functions |

Value

Invisible TRUE

| | |
|----------------|---|
| getSignificant | <i>Get number of significant digits</i> |
|----------------|---|

Description

Get number of significant digits

Usage

```
getSignificant()
```

Value

Numeric value regarding the number of significant digits

`getSpecies`*Get the species of a data category*

Description

Get the species of a data category

Usage

```
getSpecies(category = getCategory())
```

Arguments

`category` Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

Character value with the species

Note

Needs to be called inside a reactive function

`getSplicingEventCoordinates`*Returns the coordinates of interest for a given event type*

Description

Returns the coordinates of interest for a given event type

Usage

```
getSplicingEventCoordinates(type, sorting = FALSE)
```

Arguments

`type` Character: alternative splicing event type
`sorting` Boolean: get coordinates used for sorting and comparison between different programs? FALSE by default

Value

Coordinates of interest according to the alternative splicing event type

getSplicingEventTypes *Splicing event types available*

Description

Splicing event types available

Usage

```
getSplicingEventTypes()
```

Value

Named character vector with splicing event types

Examples

```
getSplicingEventTypes()
```

getUiFunctions *Matches user interface (UI) functions from a given loader*

Description

Matches user interface (UI) functions from a given loader

Usage

```
getUiFunctions(ns, loader, ..., priority = NULL)
```

Arguments

| | |
|----------|--|
| ns | Shiny function to create namespaced IDs |
| loader | Character: loader to run the functions |
| ... | Extra arguments to pass to the user interface (UI) functions |
| priority | Character: name of functions to prioritise by the given order; for instance, c("data", "analyses") would load "data", then "analyses" then remaining functions |

Value

List of functions related to the given loader

| | |
|------------------|--------------------------------------|
| getUrltoDownload | <i>Get the URL links to download</i> |
|------------------|--------------------------------------|

Description

Get the URL links to download

Usage

```
getUrltoDownload()
```

Value

Character vector with URLs to download

Note

Needs to be called inside a reactive function

| | |
|----------------|--|
| getValidEvents | <i>Filters the events with valid elements according to the given validator</i> |
|----------------|--|

Description

Filters the events with valid elements according to the given validator

Usage

```
getValidEvents(event, validator, areMultipleExonsValid = FALSE)
```

Arguments

| | |
|-----------------------|---|
| event | Data.frame containing only one event with at least 7 columns as retrieved from the alternative splicing annotation files from MISO (GFF3 files) |
| validator | Character: valid elements for each event |
| areMultipleExonsValid | Boolean: consider runs of exons as valid when comparing with the validator? Default is FALSE (see details) |

Details

areMultipleExonsValid allows to consider runs of exons (i.e. sequences where "exon" occurs consecutively) as valid when comparing with given validator. For example, if the validator is c("gene", "mRNA", "exon") and areMultipleExonsValid = FALSE, this function will only considerate events as valid if they have the exact same elements. If areMultipleExonsValid = TRUE, a valid events could include the elements c("gene", "mRNA", "exon", "exon", "exon").

Value

Data.frame with valid events

Examples

```

event <- read.table(text = "
chr1 SE gene 17233 18061 . - .
chr1 SE dkfd 00000 30000 . - .
chr1 SE mRNA 17233 18061 . - .
chr1 SE exon 17233 17368 . - .
chr1 SE exon 17526 17742 . - .
chr1 SE exon 17915 18061 . - .
chr1 SE mRNA 17233 18061 . - .
chr1 SE exon 17233 17368 . - .
chr1 SE exon 17915 18061 . - .
chr1 SE gene 17233 18061 . - .
chr1 SE mRNA 17233 18061 . - .
chr1 SE exon 17233 17368 . - .
chr1 SE exon 17606 17742 . - .
chr1 SE exon 17915 18061 . - .
chr1 SE mRNA 17233 18061 . - .
chr1 SE exon 17233 17368 . - .
chr1 SE exon 17915 18061 . - .
")
validator <- c("gene", "mRNA", rep("exon", 3), "mRNA", rep("exon", 2))
psychomics::getValidEvents(event, validator)

```

globalSelectize

Create a selectize input available from any page

Description

Create a selectize input available from any page

Usage

```
globalSelectize(id, placeholder)
```

Arguments

| | |
|-------------|------------------------------|
| id | Character: input identifier |
| placeholder | Character: input placeholder |

Value

HTML element for a global selectize input

groupByAttribute *User interface to group by attribute*

Description

User interface to group by attribute

Usage

```
groupByAttribute(ns, dataset, id, example)
```

Arguments

| | |
|---------|---------------------------------------|
| ns | Namespace function |
| dataset | Data frame: dataset of interest |
| id | Character: identifier |
| example | Character: text to show as an example |

Value

HTML elements

groupByExpression *User interface to group by subset expression*

Description

User interface to group by subset expression

Usage

```
groupByExpression(ns, id)
```

Arguments

| | |
|----|-----------------------|
| ns | Namespace function |
| id | Character: identifier |

Value

HTML elements

| | |
|-------------|---|
| groupByGrep | <i>User interface to group by grep expression</i> |
|-------------|---|

Description

User interface to group by grep expression

Usage

```
groupByGrep(ns, dataset, id)
```

Arguments

| | |
|---------|---------------------------------|
| ns | Namespace function |
| dataset | Data frame: dataset of interest |
| id | Character: identifier |

Value

HTML elements

| | |
|-----------|---------------------------------------|
| groupById | <i>User interface to group by row</i> |
|-----------|---------------------------------------|

Description

User interface to group by row

Usage

```
groupById(ns, id, choices)
```

Arguments

| | |
|---------|-----------------------------------|
| ns | Namespace function |
| id | Character: identifier |
| choices | Character: identifier suggestions |

Value

HTML elements

groupPerPatient *Assign one group to each patient*

Description

Assign one group to each patient

Usage

```
groupPerPatient(groups, patients, includeOuterGroup = FALSE,
  outerGroupName = "(Outer data)")
```

Arguments

groups List of integers: clinical groups
 patients Integer: total number of clinical patients (remaining patients will be filled with missing values)
 includeOuterGroup Boolean: join the patients that have no groups?
 outerGroupName Character: name to give to outer group

Value

Character vector where each element corresponds to the group of a clinical patient

Examples

```
groups <- list(1:3, 4:7, 8:10)
names(groups) <- paste("Stage", 1:3)
groupPerPatient(groups)
```

groupPerSample *Assign one group to each sample*

Description

Assign one group to each sample

Usage

```
groupPerSample(groups, samples, includeOuterGroup = FALSE,
  outerGroupName = "(Outer data)")
```

Arguments

groups List of characters: list of samples
 samples Character: all available samples
 includeOuterGroup Boolean: join the patients that have no groups?
 outerGroupName Character: name to give to outer group

Value

Character vector where each element corresponds to the group of a sample

Examples

```
groups <- list(letters[1:3], letters[10:12], letters[5:8])
names(groups) <- paste("Stage", 1:3)
samples <- letters
groupPerSample(groups, samples)
```

| | |
|--------------|--|
| groupsServer | <i>Server function for data grouping</i> |
|--------------|--|

Description

Server function for data grouping

Usage

```
groupsServer(input, output, session, datasetName)
```

Arguments

| | |
|-------------|----------------------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |
| datasetName | Character: name of dataset |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------------|---|
| groupsServerOnce | <i>Server function for data grouping (one call)</i> |
|------------------|---|

Description

These functions only run once instead of running for every instance of groups

Usage

```
groupsServerOnce(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|----------|---|
| groupsUI | <i>Creates UI elements for the grouping feature</i> |
|----------|---|

Description

Creates UI elements for the grouping feature

Usage

```
groupsUI(id)
```

Arguments

| | |
|----|-----------------------|
| id | Character: identifier |
|----|-----------------------|

Value

HTML elements

| | |
|----------------|--|
| gtexDataServer | <i>Server logic to load GTEEx data</i> |
|----------------|--|

Description

Server logic to load GTEEx data

Usage

```
gtexDataServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|------------------------------------|
| gtexDataUI | <i>Interface to load GTEX data</i> |
|------------|------------------------------------|

Description

Interface to load GTEX data

Usage

```
gtexDataUI(id, panel)
```

Arguments

| | |
|-------|-------------------------------------|
| id | Character: namespace identifier |
| panel | Function to deal with the interface |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|----------------|--|
| hchart.survfit | <i>Plot survival curves using Highcharts</i> |
|----------------|--|

Description

Plot survival curves using Highcharts

Usage

```
## S3 method for class 'survfit'
hchart(object, ..., fun = NULL, markTimes = TRUE,
        symbol = "plus", markerColor = "black", ranges = FALSE,
        rangesOpacity = 0.3)
```

Arguments

| | |
|---------------|--|
| object | A survfit object as returned from the <code>survfit</code> function |
| ... | Extra parameters to pass to <code>hc_add_series</code> function |
| fun | Name of function or function used to transform the survival curve: <code>log</code> will put y axis on log scale, <code>event</code> plots cumulative events ($f(y) = 1-y$), <code>cumhaz</code> plots the cumulative hazard function ($f(y) = -\log(y)$), and <code>cloglog</code> creates a complimentary log-log survival plot ($f(y) = \log(-\log(y))$) along with log scale for the x-axis. |
| markTimes | Label curves marked at each censoring time? TRUE by default |
| symbol | Symbol to use as marker (plus sign by default) |
| markerColor | Color of the marker ("black" by default); use NULL to use the respective color of each series |
| ranges | Plot interval ranges? FALSE by default |
| rangesOpacity | Opacity of the interval ranges (0.3 by default) |

Value

Highcharts object to plot survival curves

Examples

```
# Plot Kaplan-Meier curves
require("survival")
require("highcharter")
leukemia.surv <- survfit(Surv(time, status) ~ x, data = aml)
hchart(leukemia.surv)

# Plot the cumulative hazard function
lsurv2 <- survfit(Surv(time, status) ~ x, aml, type='fleming')
hchart(lsurv2, fun="cumhaz")

# Plot the fit of a Cox proportional hazards regression model
fit <- coxph(Surv(futime, fustat) ~ age, data = ovarian)
ovarian.surv <- survfit(fit, newdata=data.frame(age=60))
hchart(ovarian.surv, ranges = TRUE)
```

hc_scatter

Create scatter plot

Description

Create a scatter plot using highcharter

Usage

```
hc_scatter(hc, x, y, z = NULL, label = NULL, showInLegend = FALSE, ...)
```

Arguments

| | |
|--------------|--|
| hc | Highchart object |
| x | Numeric: X axis |
| y | Numeric: Y axis |
| z | Numeric: Z axis to set the bubble size (optional) |
| label | Character: data label for each point (optional) |
| showInLegend | Boolean: show the data in the legend box? FALSE by default |
| ... | Extra attributes of the data series to plot |

Value

Highchart object containing information for a scatter plot

| | |
|------------|--------------------------------------|
| helpServer | <i>Server logic of the help menu</i> |
|------------|--------------------------------------|

Description

Server logic of the help menu

Usage

```
helpServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|--------|--|
| helpUI | <i>User interface of the help menu</i> |
|--------|--|

Description

User interface of the help menu

Usage

```
helpUI(id, tab)
```

Arguments

| | |
|-----|-------------------------|
| id | Character: identifier |
| tab | Function to create tabs |

Value

HTML elements

inclusionLevelsInterface

Interface to quantify alternative splicing

Description

Interface to quantify alternative splicing

Usage

inclusionLevelsInterface(ns)

Arguments

ns Namespace function

Value

HTML elements

inclusionLevelsServer *Server logic of the alternative splicing event quantification module*

Description

Server logic of the alternative splicing event quantification module

Usage

inclusionLevelsServer(input, output, session)

Arguments

input Shiny input
output Shiny output
session Shiny session

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-------------------|--|
| inclusionLevelsUI | <i>Interface of the alternative splicing event quantification module</i> |
|-------------------|--|

Description

Interface of the alternative splicing event quantification module

Usage

```
inclusionLevelsUI(id, panel)
```

Arguments

| | |
|-------|-----------------------------------|
| id | Character: identifier |
| panel | Function to process HTML elements |

Value

HTML elements

| | |
|------------|---------------------|
| infoServer | <i>Server logic</i> |
|------------|---------------------|

Description

Server logic

Usage

```
infoServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|--------|-------------------------------------|
| infoUI | <i>Information's user interface</i> |
|--------|-------------------------------------|

Description

Information's user interface

Usage

```
infoUI(id)
```

Arguments

| | |
|----|-----------------------|
| id | Character: identifier |
|----|-----------------------|

Value

HTML elements

| | |
|------------|---|
| insideFile | <i>Get psychomics file inside a given directory</i> |
|------------|---|

Description

Get psychomics file inside a given directory

Usage

```
insideFile(...)
```

Arguments

| | |
|-----|--|
| ... | character vectors, specifying subdirectory and file(s) within some package. The default, none, returns the root of the package. Wildcards are not supported. |
|-----|--|

Value

Loaded file

| | |
|----------|-----------------------------------|
| is.whole | <i>Check if a number is whole</i> |
|----------|-----------------------------------|

Description

Check if a number is whole

Usage

```
is.whole(x, tol = .Machine$double.eps^0.5)
```

Arguments

| | |
|-----|--|
| x | Object to be tested |
| tol | Numeric: tolerance used for comparison |

Value

TRUE if number is whole; otherwise, FALSE

| | |
|----------------|---|
| isFirebrowseUp | <i>Check if the Firebrowse web API is running</i> |
|----------------|---|

Description

The Firebrowse web API is running if it returns the status condition 200; if this is not the status code obtained from the API, the function will raise a warning with the status code and a brief explanation.

Usage

```
isFirebrowseUp()
```

```
isFirehoseUp()
```

Value

Invisible TRUE if the Firebrowse web API is working; otherwise, raises a warning

Examples

```
isFirebrowseUp()
```

| | |
|-------------------|---|
| joinEventsPerType | <i>Full outer join all given events based on select columns</i> |
|-------------------|---|

Description

Full outer join all given events based on select columns

Usage

```
joinEventsPerType(events, types)
```

Arguments

| | |
|--------|---|
| events | Data frame or matrix: alternative splicing events |
| types | Character: alternative splicing types |

Value

List of events joined by alternative splicing event type

| | |
|----------------|--|
| junctionString | <i>String used to search for matches in a junction quantification file</i> |
|----------------|--|

Description

String used to search for matches in a junction quantification file

Usage

```
junctionString(chr, strand, junc5, junc3, showStrand)
```

Arguments

| | |
|------------|--------------------------|
| chr | Character: chromosome |
| strand | Character: strand |
| junc5 | Integer: 5' end junction |
| junc3 | Integer: 3' end junction |
| showStrand | Boolean: include strand? |

Value

Formatted character string

| | |
|---------|--|
| kruskal | <i>Perform Kruskal's test and return interface to show the results</i> |
|---------|--|

Description

Perform Kruskal's test and return interface to show the results

Usage

```
kruskal(psi, groups, stat = NULL)
```

Arguments

| | |
|--------|--|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |
| stat | Data frame or matrix: values of the analyses to be performed (if NULL, the analyses will be performed) |

Value

HTML elements

| | |
|--------------------|--|
| labelBasedOnCutoff | <i>Label groups based on a given cut-off</i> |
|--------------------|--|

Description

Label groups based on a given cut-off

Usage

```
labelBasedOnCutoff(data, cutoff, label = NULL, gte = TRUE)
```

Arguments

| | |
|--------|--|
| data | Numeric: test data |
| cutoff | Numeric: test cutoff |
| label | Character: label to prefix group names (NULL by default) |
| gte | Boolean: test with greater than or equal to cutoff (TRUE) or use less than or equal to cutoff (FALSE)? TRUE by default |

Value

Labeled groups

Examples

```
labelBasedOnCutoff(data=c(1, 0, 0, 1, 0, 1), cutoff=0.5)

labelBasedOnCutoff(data=c(1, 0, 0, 1, 0, 1), cutoff=0.5, "Ratio")

# Use "greater than" instead of "greater than or equal to"
labelBasedOnCutoff(data=c(1, 0, 0, 0.5, 0, 1), cutoff=0.5, gte=FALSE)
```

| | |
|--------|---|
| levene | <i>Perform Levene's test and return interface to show the results</i> |
|--------|---|

Description

Perform Levene's test and return interface to show the results

Usage

```
levene(psi, groups, stat = NULL)
```

Arguments

| | |
|--------|--|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |
| stat | Data frame or matrix: values of the analyses to be performed (if NULL, the analyses will be performed) |

Value

HTML elements

| | |
|------------|----------------------|
| leveneTest | <i>Levene's test</i> |
|------------|----------------------|

Description

Performs a Levene's test to assess the equality of variances

Usage

```
leveneTest(x, g, centers = median)
```

Arguments

| | |
|---------|---|
| x | a numeric vector of data values, or a list of numeric data vectors. Non-numeric elements of a list will be coerced, with a warning. |
| g | a vector or factor object giving the group for the corresponding elements of x. Ignored with a warning if x is a list. |
| centers | Function used to calculate how much values spread (median by default; another common function used is mean) |

Value

A list with class "htest" containing the following components:

| | |
|-----------|--|
| statistic | the value of the test statistic with a name describing it. |
| p.value | the p-value for the test. |
| method | the type of test applied. |
| data.name | a character string giving the names of the data. |

Examples

```
vals <- sample(30, replace=TRUE)
group <- lapply(list("A", "B", "C"), rep, 10)
group <- unlist(group)
psychomics:::leveneTest(vals, group)

## Using Levene's test based on the mean
psychomics:::leveneTest(vals, group, mean)
```

| | |
|--------------------|---|
| listAllAnnotations | <i>List alternative splicing annotation files available, as well as custom annotation</i> |
|--------------------|---|

Description

List alternative splicing annotation files available, as well as custom annotation

Usage

```
listAllAnnotations(...)
```

Arguments

| | |
|-----|--------------------------|
| ... | Custom annotation loaded |
|-----|--------------------------|

Value

Named character vector with splicing annotation files available#'

Examples

```
psychomics:::listAllAnnotations()
```

```
listSplicingAnnotations
```

List the alternative splicing annotation files available

Description

List the alternative splicing annotation files available

Usage

```
listSplicingAnnotations()
```

Value

Named character vector with splicing annotation files available

Examples

```
listSplicingAnnotations()
```

```
loadAnnotation
```

Load alternative splicing annotation from AnnotationHub

Description

Load alternative splicing annotation from AnnotationHub

Usage

```
loadAnnotation(annotation)
```

Arguments

annotation Character: annotation to load

Value

List of data frames containing the alternative splicing annotation per event type

Examples

```
human <- listSplicingAnnotations()[[1]]
## Not run:
annot <- loadAnnotation(human)

## End(Not run)
```

| | |
|--------|---|
| loadBy | <i>Check if a given function should be loaded by the calling module</i> |
|--------|---|

Description

Check if a given function should be loaded by the calling module

Usage

```
loadBy(loader, FUN)
```

Arguments

| | |
|--------|---|
| loader | Character: name of the file responsible to load such function |
| FUN | Function |

Value

Boolean vector

| | |
|-----------------|---|
| loadedDataModal | <i>Create a modal warning the user of already loaded data</i> |
|-----------------|---|

Description

Create a modal warning the user of already loaded data

Usage

```
loadedDataModal(session, modalId, replaceButtonId, keepButtonId)
```

Arguments

| | |
|-----------------|---|
| session | Shiny session |
| modalId | Character: identifier of the modal |
| replaceButtonId | Character: identifier of the button to replace data |
| keepButtonId | Character: identifier of the button to append data |

Value

HTML elements for a warning modal reminding data is loaded

| | |
|----------|---|
| loadFile | <i>Loads a file according to its format</i> |
|----------|---|

Description

Loads a file according to its format

Usage

```
loadFile(format, file)
```

Arguments

| | |
|--------|---------------------------------|
| format | Environment: format of the file |
| file | Character: file to load |

Details

The resulting data frame includes the attribute "tablename" with the name of the data frame

Value

Data frame with the loaded file

| | |
|-----------------|---------------------------|
| loadFileFormats | <i>Loads file formats</i> |
|-----------------|---------------------------|

Description

Loads file formats

Usage

```
loadFileFormats()
```

Value

Loaded file formats available

| | |
|--------------------|---|
| loadFirebrowseData | <i>Downloads and processes data from the Firebrowse web API and loads it into R</i> |
|--------------------|---|

Description

Downloads and processes data from the Firebrowse web API and loads it into R

Usage

```
loadFirebrowseData(folder = NULL, data = NULL, exclude = c(".aux.",  
  ".mage-tab.", "MANIFEST.txt"), ..., progress = echoProgress,  
  download = TRUE)
```

```
loadFirehoseData(folder = NULL, data = NULL, exclude = c(".aux.",  
  ".mage-tab.", "MANIFEST.txt"), ..., progress = echoProgress,  
  download = TRUE)
```

Arguments

| | |
|----------|---|
| folder | Character: directory to store the downloaded archives (by default, it saves in the user's "Downloads" folder) |
| data | Character: data to load |
| exclude | Character: files and folders to exclude from downloading and from loading into R (by default, it excludes ".aux.", ".mage-tab." and "MANIFEST.TXT" files) |
| ... | Extra parameters to be passed to queryFirebrowseData |
| progress | Function to show the progress (default is to print progress to console) |
| download | Boolean: download missing files through the function <code>download.file</code> (TRUE by default) |

Value

URL of missing files ("missing" class) if files need to be downloaded and if the argument `download` is FALSE; else, a list with loaded data

Examples

```
## Not run:  
loadFirebrowseData(cohort = "ACC", data_type = "Clinical")  
  
## End(Not run)
```

loadFirebrowseFolders *Load Firebrowse folders*

Description

Loads the files present in each folder as a data.frame.

Usage

```
loadFirebrowseFolders(folder, exclude = "", progress = echoProgress)
```

```
loadFirehoseFolders(folder, exclude = "", progress = echoProgress)
```

Arguments

| | |
|----------|---|
| folder | Character: folder(s) in which to look for Firebrowse files |
| exclude | Character: files to exclude from the loading |
| progress | Function to show the progress (default is to print progress to console) |

Value

List with loaded data.frames

Note

For faster execution, this function uses the readr library. This function ignores subfolders of the given folder (which means that files inside subfolders are NOT loaded).

loadGtexData *Load GTEX data given input*

Description

Load GTEX data given input

Usage

```
loadGtexData(input, replace = TRUE)
```

Arguments

| | |
|---------|---|
| input | Shiny input |
| replace | Boolean: replace loaded data? TRUE by default |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|----------------|-------------------------|
| loadLocalFiles | <i>Load local files</i> |
|----------------|-------------------------|

Description

Load local files

Usage

```
loadLocalFiles(folder, ignore = c(".aux.", ".mage-tab."), name = "Data",
  progress = echoProgress)
```

Arguments

| | |
|----------|---|
| folder | Character: path to folder containing files of interest |
| ignore | Character: skip folders and filenames that match the expression |
| name | Character: name of the category containing all loaded datasets |
| progress | Function to keep track of the progress |

Value

List of data frames from valid files

Examples

```
## Not run:
folder <- "~/Downloads/ACC 2016"
data <- loadLocalFiles(folder)

ignore <- c(".aux.", ".mage-tab.", "junction quantification")
loadLocalFiles(folder, ignore)

## End(Not run)
```

| | |
|-----------------|--|
| localDataServer | <i>Server logic to load local data</i> |
|-----------------|--|

Description

Server logic to load local data

Usage

```
localDataServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-------------|-------------------------------------|
| localDataUI | <i>Interface to load local data</i> |
|-------------|-------------------------------------|

Description

Interface to load local data

Usage

```
localDataUI(id, panel)
```

Arguments

| | |
|-------|-------------------------------------|
| id | Character: namespace identifier |
| panel | Function to deal with the interface |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------------|---|
| missingDataModal | <i>Missing information modal template</i> |
|------------------|---|

Description

Missing information modal template

Usage

```
missingDataModal(session, dataType, buttonId)
```

```
loadRequiredData(modal = NULL)
```

```
missingDataGuide(dataType)
```

Arguments

| | |
|----------|---|
| session | Shiny session |
| dataType | Character: type of data missing |
| buttonId | Character: identifier of button to take user to load missing data |
| modal | Character: modal identifier |

Value

NULL (this function is used to modify the Shiny session's state)

Examples

```
## Not run:
  session <- session$ns
  buttonInput <- "takeMeThere"
  buttonId <- ns(buttonInput)
  dataType <- "Inclusion levels"
  missingDataModal(session, buttonId, dataType)
  observeEvent(input[[buttonInput]], missingDataGuide(dataType))

## End(Not run)
```

 modTabPanel

Modified tabPanel function to show icon and title

Description

Modified tabPanel function to show icon and title

Usage

```
modTabPanel(title, ..., icon = NULL, menu = FALSE)
```

Arguments

| | |
|-------|--|
| title | Character: title of the tab |
| ... | HTML elements to pass to tab |
| icon | Character: name of the icon |
| menu | Boolean: create a dropdown menu-like tab? FALSE by default |

Value

HTML interface for a tab panel

Note

Icon is hidden at small viewports

 navSelectize

Create a special selectize input in the navigatin bar

Description

Create a special selectize input in the navigatin bar

Usage

```
navSelectize(id, label, placeholder = label)
```

Arguments

| | |
|-------------|------------------------------|
| id | Character: input identifier |
| label | Character: input label |
| placeholder | Character: input placeholder |

Value

HTML element to be included in a navigation bar

| | |
|--------|---|
| noinfo | <i>Interface when no information could be retrieved</i> |
|--------|---|

Description

Interface when no information could be retrieved

Usage

```
noinfo(output,
  description = paste("No information available for the gene associated with this event."),
  ...)
```

Arguments

| | |
|--------------------|---|
| output | Shiny output |
| description | Character: description of the message to show to the user |
| ... | Arguments passed on to <code>errorDialog</code> |
| id | Character: identifier (NULL by default) |
| buttonId | Character: button identifier (NULL by default) |
| buttonLabel | Character: button label (NULL by default) |
| buttonIcon | Character: button icon (NULL by default) |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-----------------|---------------------------------|
| operateOnGroups | <i>Set operations on groups</i> |
|-----------------|---------------------------------|

Description

This function can be used on groups to merge, intersect, subtract, etc.

Usage

```
operateOnGroups(input, session, FUN, buttonId, symbol = " ", datasetName,
  sharedData = sharedData)
```

Arguments

| | |
|-------------|--|
| input | Shiny input |
| session | Shiny session |
| FUN | Function: operation to set |
| buttonId | Character: ID of the button to trigger operation |
| symbol | Character: operation symbol |
| datasetName | Character: name of dataset |
| sharedData | Shiny app's global variable |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------------|--|
| optimalPSIcutoff | <i>Calculate optimal alternative splicing quantification cut-off to separate survival curves</i> |
|------------------|--|

Description

Calculate optimal alternative splicing quantification cut-off to separate survival curves

Usage

```
optimalPSIcutoff(clinical, psi, censoring, event, timeStart, timeStop = NULL,
  followup = "days_to_last_followup", session = NULL, filter = TRUE,
  survTime = NULL)
```

Arguments

| | |
|-----------|--|
| clinical | Data frame: clinical data |
| psi | Numeric: PSI values to test against the cut-off |
| censoring | Character: censor using "left", "right", "interval" or "interval2" |
| event | Character: name of column containing time of the event of interest |
| timeStart | Character: name of column containing starting time of the interval or follow up time |
| timeStop | Character: name of column containing ending time of the interval |
| followup | Character: name of column containing follow up time |
| session | Shiny session (only used for the visual interface) |
| filter | Boolean or numeric: elements to use (all by default) |
| survTime | survTime object: times to follow up, time start, time stop and event (optional) |

Details

timeStop is only considered if censoring is either interval or interval2

Value

Optimal alternative splicing quantification cut-off

Examples

```
clinical <- read.table(text = "2549  NA ii  female
                             840  NA i   female
                             NA 1204 iv  male
                             NA  383 iv  female
                             1293  NA iii male
                             NA 1355 ii  male")
names(clinical) <- c("patient.days_to_last_followup",
                    "patient.days_to_death",
                    "patient.stage_event.pathologic_stage",
                    "patient.gender")
timeStart <- "days_to_death"
event     <- "days_to_death"

psi <- c(0.1, 0.2, 0.9, 1, 0.2, 0.6)
opt <- optimalPSIcutoff(clinical, psi, "right", event, timeStart)
```

optimSurvDiff

Optimal survival difference given an inclusion level cut-off for a specific alternative splicing event

Description

Optimal survival difference given an inclusion level cut-off for a specific alternative splicing event

Usage

```
optimSurvDiff(session, input, output)
```

Arguments

| | |
|---------|---------------|
| session | Shiny session |
| input | Shiny input |
| output | Shiny output |

Value

NULL (this function is used to modify the Shiny session's state) Calculate optimal survival cut-off for the inclusion levels of a given alternative splicing event

parseDateResponse *Parse the date from a response*

Description

Parse the date from a response

Usage

```
parseDateResponse(string)
```

Arguments

| | |
|--------|------------------|
| string | Character: dates |
|--------|------------------|

Value

Parsed date

parseFirebrowseMetadata *Query the Firebrowse web API for metadata and parse the response*

Description

Query the Firebrowse web API for metadata and parse the response

Usage

```
parseFirebrowseMetadata(type, ...)
```

```
parseFirehoseMetadata(type, ...)
```

Arguments

| | |
|------|---|
| type | Character: metadata to retrieve |
| ... | Character: parameters to pass to query (optional) |

Value

List with parsed JSON response

Examples

```
psychomics:::parseFirebrowseMetadata("Dates")
psychomics:::parseFirebrowseMetadata("Centers")
psychomics:::parseFirebrowseMetadata("HeartBeat")

# Get the abbreviation and description of all cohorts available
psychomics:::parseFirebrowseMetadata("Cohorts")
# Get the abbreviation and description of the selected cohorts
psychomics:::parseFirebrowseMetadata("Cohorts", cohort = c("ACC", "BRCA"))
```

| | |
|----------------|--|
| parseMatsEvent | <i>Parse alternative splicing events from MATS</i> |
|----------------|--|

Description

Parse alternative splicing events from MATS

Usage

```
parseMatsEvent(event, event_type)
```

Arguments

| | |
|------------|---|
| event | Data frame row: MATS splicing event |
| event_type | Character: Type of event to parse (see details) |

Details

The following event types can be parsed:

- **SE**: Skipped exon
- **MXE**: Mutually exclusive exons
- **RI**: Retained intron
- **A3SS**: Alternative 3' splice site
- **A5SS**: Alternative 5' splice site

Value

List containing the event attributes and junctions

Examples

```
# MATS event (alternative 3' splice site)
event <- read.table(text = "
  2 ENSG00000166012 TAF1D chr11 - 93466515 93466671 93466515 93466563 93467790 93467826
  5 ENSG00000166012 TAF1D chr11 - 93466515 93466671 93466515 93466585 93467790 93467826
  6 ENSG00000166012 TAF1D chr11 - 93466515 93466585 93466515 93466563 93467790 93467826
")
psychomics:::parseMatsEvent(event, "A3SS")
```

| | |
|------------------|---|
| parseMatsGeneric | <i>Parse junctions of an alternative splicing event from MATS according to event type</i> |
|------------------|---|

Description

Parse junctions of an alternative splicing event from MATS according to event type

Usage

```
parseMatsGeneric(junctions, strand, coords, plus_pos, minus_pos)
```

```
parseMatsSE(junctions, strand)
```

```
parseMatsMXE(junctions, strand)
```

```
parseMatsRI(junctions, strand)
```

```
parseMatsA3SS(junctions, strand)
```

```
parseMatsA5SS(junctions, strand)
```

```
parseMatsAFE(junctions, strand)
```

```
parseMatsALE(junctions, strand)
```

Arguments

| | |
|-----------|---|
| junctions | Integer: event's junctions |
| strand | Character: strand of the event |
| coords | Character: names of the alternative splicing coordinates |
| plus_pos | Integer: match of each junction in the respective coordinate for the plus strand |
| minus_pos | Integer: match of each junction in the respective coordinate for the minus strand |

Details

The following event types are ready to be parsed:

- **SE** (skipped exon)
- **MXE** (mutually exclusive exon)
- **RI** (intron retention)
- **A5SS** (alternative 5' splice site)
- **A3SS** (alternative 3' splice site)
- **AFE** (alternative first exon)
- **ALE** (alternative last exon)

You can use `parseMatsGeneric` to parse other event types.

Value

Data frame with parsed junctions

See Also

[parseMatsEvent](#)

Examples

```
# Parse generic event (in this case, an exon skipping event)
junctions <- read.table(text=
  "79685787 79685910 79685796 79685910 79679566 79679751")
coords <- c("A1.start", "A1.end",
            "C1.start", "C1.end",
            "C2.start", "C2.end")
plus <- c(1:6)
minus <- c(2:1, 6:3)
psychomics:::parseMatsGeneric(junctions, strand = "+", coords, plus, minus)

# Parse exon skipping event
junctions <- read.table(text=
  "79685787 79685910 79685796 79685910 79679566 79679751")
psychomics:::parseMatsSE(junctions, strand = "+")

# Parse mutually exclusive exon event
junctions <- read.table(text=
  "158282161 158282276 158282689 158282804 158281047 158281295 158283950 158284199")
psychomics:::parseMatsMXE(junctions, strand = "+")

# Parse intron retention event
junctions <- read.table(text=
  "15929853 15932100 15929853 15930016 15930687 15932100")
psychomics:::parseMatsRI(junctions, strand = "+")

# Parse alternative 3' splicing site event
junctions <- read.table(text=
  "79685787 79685910 79685796 79685910 79679566 79679751")
psychomics:::parseMatsA3SS(junctions, strand = "+")

# Parse alternative 5' splicing site event
junctions <- read.table(text=
  "102884421 102884501 102884421 102884489 102884812 102885881")
psychomics:::parseMatsA5SS(junctions, strand = "+")

# Parse alternative first exon event
junctions <- read.table(text=
  "16308723 16308879 16308967 16309119 16314269 16314426")
psychomics:::parseMatsAFE(junctions, strand = "+")

# Parse alternative last exon event
junctions <- read.table(text=
  "111858645 111858828 111851063 111851921 111850441 111850543")
psychomics:::parseMatsAFE(junctions, strand = "+")
```

| | |
|----------------|--|
| parseMisoEvent | <i>Parse an alternative splicing event from MISO</i> |
|----------------|--|

Description

Parse an alternative splicing event from MISO

Usage

```
parseMisoEvent(event)
```

Arguments

| | |
|-------|---|
| event | Data.frame containing only one event with at least 7 columns as retrieved from the alternative splicing annotation files from MISO (GFF3 files) |
|-------|---|

Details

More information about MISO available at <http://miso.readthedocs.org>

Value

List with event attributes and junction positions for the exons (depends on the events)

Examples

```
# example of alternative splicing event: skipped exon (SE)
event <- read.table(text = "
chr1 SE gene 16854 18061 . - .
chr1 SE mRNA 16854 18061 . - .
chr1 SE exon 16854 17055 . - .
chr1 SE exon 17233 17742 . - .
chr1 SE exon 17915 18061 . - .
chr1 SE mRNA 16854 18061 . - .
chr1 SE exon 16854 17955 . - .
chr1 SE exon 17915 18061 . - .")
psychomics:::parseMisoEvent(event)
```

| | |
|------------------|--|
| parseMisoEventID | <i>Match MISO's splicing event IDs with the IDs present in the alternative splicing annotation file and get events in a data frame</i> |
|------------------|--|

Description

Match MISO's splicing event IDs with the IDs present in the alternative splicing annotation file and get events in a data frame

Usage

```
parseMisoEventID(eventID, annotation, IDcolumn)
```

Arguments

| | |
|------------|---|
| eventID | Character: alternative event IDs |
| annotation | Data.frame: alternative event annotation file |
| IDcolumn | Integer: index of the column with the event ID's in the alternative event annotation file |

Details

For faster execution times, provide a vector of event IDs.

For more information about MISO, see <http://miso.readthedocs.org>.

Value

Data frame of the matching events (or NA when nothing is matched)

Note

If possible, it's recommend to use smaller subsets of the alternative events' annotation instead of all data for faster runs. For example, when trying to match only skipped exons event IDs, only use the annotation of skipped exons instead of using a mega annotation with all event types.

Examples

```
eventID <- c("114785@uc001sok.1@uc001soj.1", "114784@uc001bxm.1@uc001bxn.1")
# the annotation is one of the GFF3 files needed to run MISO
gff3 <- system.file("extdata", "miso_AS_annot_example.gff3",
  package="psychomics")
annotation <- read.delim(gff3, header=FALSE, comment.char="#")
IDcolumn <- 9
psychomics:::parseMisoEventID(eventID, annotation, IDcolumn)
```

parseMisoGeneric

Parse junctions of an event from MISO according to event type

Description

Parse junctions of an event from MISO according to event type

Usage

```
parseMisoGeneric(event, validator, eventType, coord, plusIndex, minusIndex)
```

```
parseMisoSE(event)
```

```
parseMisoMXE(event)
```

```
parseMisoRI(event, strand)
```

```
parseMisoA5SS(event)
```

```
parseMisoA3SS(event, plusIndex, minusIndex)
```

```
parseMisoTandemUTR(event, minusIndex)
```

```
parseMisoAFE(event)
```

```
parseMisoALE(event)
```

Arguments

| | |
|------------|---|
| event | Data.frame containing only one event with at least 7 columns as retrieved from the alternative splicing annotation files from MISO (GFF3 files) |
| validator | Character: valid elements for each event |
| eventType | Character: event type (see details for available events) |
| coord | Character: coordinate positions to fill |
| plusIndex | Integer: index of the coordinates for a plus strand event |
| minusIndex | Integer: index of the coordinates for a minus strand event |
| strand | Character: "+" or "-" strand |

Details

The following event types are available to be parsed:

- **SE** (exon skipping)
- **MXE** (mutually exclusive exon)
- **RI** (intron retention)
- **A5SS** (alternative 5' splice site)
- **A3SS** (alternative 3' splice site)
- **AFE** (alternative first exon)
- **ALE** (alternative last exon)
- **Tandem UTR**

Value

List of parsed junctions

See Also

[parseMisoEvent](#)

Examples

```
# skipped exon event (SE)
event <- read.table(text = "
chr1 SE gene 16854 18061 . - .
chr1 SE mRNA 16854 18061 . - .
chr1 SE exon 16854 17055 . - .
chr1 SE exon 17233 17742 . - .
chr1 SE exon 17915 18061 . - .
chr1 SE mRNA 16854 18061 . - .
chr1 SE exon 16854 17955 . - .
chr1 SE exon 17915 18061 . - .")
```

```

psychomics:::parseMisoSE(event)

# mutually exclusive exon (MXE) event
event <- read.table(text = "
chr1 MXE gene 764383 788090 . + .
chr1 MXE mRNA 764383 788090 . + .
chr1 MXE exon 764383 764484 . + .
chr1 MXE exon 776580 776753 . + .
chr1 MXE exon 787307 788090 . + .
chr1 MXE mRNA 764383 788090 . + .
chr1 MXE exon 764383 764484 . + .
chr1 MXE exon 783034 783186 . + .
chr1 MXE exon 787307 788090 . + .")
psychomics:::parseMisoMXE(event)

# intron retention (RI) event
event <- read.table(text = "
chr1 RI gene 17233 17742 . - .
chr1 RI mRNA 17233 17742 . - .
chr1 RI exon 17233 17742 . - .
chr1 RI mRNA 17233 17742 . - .
chr1 RI exon 17233 17364 . - .
chr1 RI exon 17601 17742 . - .")
psychomics:::parseMisoRI(event)

# alternative 5' splice site (A5SS) event
event <- read.table(text = "
chr1 A5SS gene 17233 17742 . - .
chr1 A5SS mRNA 17233 17742 . - .
chr1 A5SS exon 17233 17368 . - .
chr1 A5SS exon 17526 17742 . - .
chr1 A5SS mRNA 17233 17742 . - .
chr1 A5SS exon 17233 17368 . - .
chr1 A5SS exon 17606 17742 . - .")
psychomics:::parseMisoA5SS(event)

# alternative 3' splice site (A3SS) event
event <- read.table(text = "
chr1 A3SS gene 15796 16765 . - .
chr1 A3SS mRNA 15796 16765 . - .
chr1 A3SS exon 15796 15947 . - .
chr1 A3SS exon 16607 16765 . - .
chr1 A3SS mRNA 15796 16765 . - .
chr1 A3SS exon 15796 15942 . - .
chr1 A3SS exon 16607 16765 . - .")
psychomics:::parseMisoA3SS(event)

# Tandem UTR event
event <- read.table(text = "
chr19 TandemUTR gene 10663759 10664625 . - .
chr19 TandemUTR mRNA 10663759 10664625 . - .
chr19 TandemUTR exon 10663759 10664625 . - .
chr19 TandemUTR mRNA 10664223 10664625 . - .
chr19 TandemUTR exon 10664223 10664625 . - .")
psychomics:::parseMisoTandemUTR(event)

# alternative first exon (AFE) event

```

```

event <- read.table(text = "
chr12 AFE gene 57916659 57920171 . + .
chr12 AFE mRNA 57919131 57920171 . + .
chr12 AFE exon 57919131 57920171 . + .
chr12 AFE mRNA 57916659 57918199 . + .
chr12 AFE exon 57916659 57916794 . + .
chr12 AFE exon 57917812 57917875 . + .
chr12 AFE exon 57918063 57918199 . + .")
psychomics:::parseMisoAFE(event)

# alternative last exon (ALE) event
event <- read.table(text = "
chr6 ALE gene 30620579 30822593 . + .
chr6 ALE mRNA 30822190 30822593 . + .
chr6 ALE exon 30822190 30822593 . + .
chr6 ALE mRNA 30620579 30620982 . + .
chr6 ALE exon 30620579 30620982 . + .")
psychomics:::parseMisoALE(event)

```

parseMisoId

Parse MISO's alternative splicing event identifier

Description

Parse MISO's alternative splicing event identifier

Usage

```
parseMisoId(id)
```

Arguments

id Character: MISO alternative splicing event identifier

Value

Character with the parsed ID

Examples

```

id <- paste0(
  "ID=ENSMUSG00000026150.chr1:82723803:82723911:+@chr1:82724642:82724813:",
  "+@chr1:82725791:82726011:+.B;Parent=ENSMUSG00000026150.chr1:82723803:",
  "82723911:+@chr1:82724642:82724813:+@chr1:82725791:82726011:+")
psychomics:::parseMisoId(id)

```

parseSampleGroups *Return the type of a given sample*

Description

Return the type of a given sample

Usage

```
parseSampleGroups(sample, filename = system.file("extdata",
  "TCGAsampleType.RDS", package = "psychomics"))
```

Arguments

| | |
|----------|---|
| sample | Character: ID of the sample |
| filename | Character: path to RDS file containing corresponding type |

Value

Types of the TCGA samples

Examples

```
parseSampleGroups(c("TCGA-01A-Tumour", "TCGA-10B-Normal"))
```

parseSplicingEvent *Parse an alternative splicing event based on a given identifier*

Description

Parse an alternative splicing event based on a given identifier

Usage

```
parseSplicingEvent(event, char = FALSE)
```

Arguments

| | |
|-------|---|
| event | Character: event identifier |
| char | Boolean: return a single character instead of list with parsed values? FALSE by default |

Value

Parsed event

Examples

```
events <- c("SE_1_-_123_456_789_1024_TST",
  "MX_3+_473_578_686_736_834_937_HEY/YOU")
parseSplicingEvent(events)
```

parseSuppaAnnotation *Get events from alternative splicing annotation*

Description

Get events from alternative splicing annotation

Usage

```
parseSuppaAnnotation(folder, types = c("SE", "AF", "AL", "MX", "A5", "A3",
  "RI"), genome = "hg19")
```

```
parseVastToolsAnnotation(folder, types = c("ALT3", "ALT5", "COMBI", "IR",
  "MERGE3m", "MIC", "EXSK", "MULTI"), genome = "Hsa", complexEvents = FALSE)
```

```
parseMisoAnnotation(folder, types = c("SE", "AFE", "ALE", "MXE", "A5SS",
  "A3SS", "RI", "TandemUTR"), genome = "hg19")
```

```
parseMatsAnnotation(folder, types = c("SE", "AFE", "ALE", "MXE", "A5SS",
  "A3SS", "RI"), genome = "fromGTF", novelEvents = TRUE)
```

Arguments

| | |
|---------------|--|
| folder | Character: path to folder |
| types | Character: type of events to retrieve (depends on the program of origin; see details) |
| genome | Character: genome of interest (for instance, "hg19"; depends on the program of origin) |
| complexEvents | Boolean: should complex events in A3SS and A5SS be parsed? FALSE by default |
| novelEvents | Boolean: parse events dedected due to novel splice sites (TRUE by default) |

Details

Type of parseable events:

- Alternative 3' splice site
- Alternative 5' splice site
- Alternative first exon
- Alternative last exon
- Skipped exon (may include skipped micro-exons)
- Mutually exclusive exon
- Retained intron
- Tandem UTR

Value

Retrieve data frame with events based on a given alternative splicing annotation

Examples

```

# Load sample files
folder <- "extdata/eventsAnnotSample/suppa_output/suppaEvents"
suppaOutput <- system.file(folder, package="psychomics")

suppa <- parseSuppaAnnotation(suppaOutput)
# Load sample files
folder <- "extdata/eventsAnnotSample/VASTDB/Hsa/TEMPLATES"
vastToolsOutput <- system.file(folder, package="psychomics")

vast <- parseVastToolsAnnotation(vastToolsOutput)
# Load sample files
folder <- "extdata/eventsAnnotSample/miso_annotation"
misoOutput <- system.file(folder, package="psychomics")

miso <- parseMisoAnnotation(misoOutput)
# Load sample files
folder <- "extdata/eventsAnnotSample/mats_output/AEvents"
matsOutput <- system.file(folder, package="psychomics")

mats <- parseMatsAnnotation(matsOutput)

# Do not parse novel events
mats <- parseMatsAnnotation(matsOutput, novelEvents=FALSE)

```

parseSuppaEvent

Parses splicing events of a specific event type from SUPPA

Description

Parses splicing events of a specific event type from SUPPA

Usage

```
parseSuppaEvent(event)
```

Arguments

event Character vector: Splicing event attributes and junction positions

Details

More information about SUPPA available at <https://bitbucket.org/regulatorygenomicsupf/suppa>

The following event types are available to be parsed:

- **SE** (skipped exon)
- **RI** (intron retention)
- **MX** (mutually exclusive exons)
- **A5** (alternative 5' splice site)
- **A3** (alternative 3' splice site)
- **AL** (alternative last exon)
- **AF** (alternative first exon)

Value

List with the event attributes (chromosome, strand, event type and the position of the exon boundaries)

Note

It only allows to parse one event type at once.

Examples

```
event <- "ENSG00000000419;A3:20:49557492-49557642:49557470-49557642:-"
psychomics:::parseSuppaEvent(event)
```

parseSuppaGeneric *Parse junctions of an event from SUPPA*

Description

Parse junctions of an event from SUPPA

Usage

```
parseSuppaGeneric(junctions, strand, coords, plus_pos, minus_pos)
parseSuppaSE(junctions, strand)
parseSuppaRI(junctions, strand)
parseSuppaALE(junctions, strand)
parseSuppaAFE(junctions, strand)
parseSuppaMXE(junctions, strand)
parseSuppaA3SS(junctions, strand)
parseSuppaA5SS(junctions, strand)
```

Arguments

| | |
|-----------|--|
| junctions | List of integers: exon-exon junctions of an event |
| strand | Character: positive ("+") or negative ("-") strand |
| coords | Character: coordinate positions to fill |
| plus_pos | Integer: index of the coordinates for a plus strand event |
| minus_pos | Integer: index of the coordinates for a minus strand event |

Details

The following event types are available to be parsed:

- **SE** (exon skipping)
- **RI** (intron retention)
- **MXE** (mutually exclusive exons)
- **A5SS** (alternative 5' splice site)
- **A3SS** (alternative 3' splice site)
- **ALE** (alternative last exon)
- **AFE** (alternative first exon)

Value

Data frame of parsed junctions

See Also

[parseSuppaEvent](#)

Examples

```
# Parse generic event (in this case, an exon skipping event)
junctions <- read.table(text = "169768099 169770024 169770112 169771762")
coords <- c("C1.end", "A1.start", "A1.end", "C2.start")
plus <- 1:4
minus <- 1:4
psychomics:::parseSuppaGeneric(junctions, strand = "+", coords, plus, minus)

junctions <- read.table(text = "169768099 169770024 169770112 169771762")
psychomics:::parseSuppaSE(junctions, "+")

junctions <- read.table(text = "196709749 196709922 196711005 196711181")
psychomics:::parseSuppaRI(junctions, "+")

junctions <- read.table(
  text = "24790610 24792494 24792800 24790610 24795476 24795797")
psychomics:::parseSuppaALE(junctions, "+")

junctions <- read.table(
  text = "169763871 169764046 169767998 169764550 169765124 169767998")
psychomics:::parseSuppaAFE(junctions, "+")

junctions <- read.table(
  text = "202060671 202068453 202068489 202073793 202060671 202072798 202072906 202073793")
psychomics:::parseSuppaMXE(junctions, "+")

junctions <- read.table(text = "169772450 169773216 169772450 169773253")
psychomics:::parseSuppaA3SS(junctions, "+")

junctions <- read.table(text = "50193276 50197008 50192997 50197008")
psychomics:::parseSuppaA5SS(junctions, "+")
```

parseTcgaSampleInfo *Parse and prepare sample information from TCGA samples*

Description

Parse and prepare sample information from TCGA samples

Usage

```
parseTcgaSampleInfo(samples, category = getCategory())
```

Arguments

| | |
|----------|--|
| samples | Character: sample identifiers |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

Data frame containing metadata associated with each TCGA sample

parseUniprotXML *Parse XML from Uniprot's RESTful service*

Description

Parse XML from Uniprot's RESTful service

Usage

```
parseUniprotXML(xml)
```

Arguments

| | |
|-----|-----------------------|
| xml | response from Uniprot |
|-----|-----------------------|

Value

List containing protein length and data frame of protein features

```
parseUrlsFromFirebrowseResponse
```

Retrieve URLs from a response to a Firebrowse data query

Description

Retrieve URLs from a response to a Firebrowse data query

Usage

```
parseUrlsFromFirebrowseResponse(res)
```

```
parseUrlsFromFirehoseResponse(res)
```

Arguments

res Response from http::GET to a Firebrowse data query

Value

Named character with URLs

Examples

```
res <- psychomics:::queryFirebrowseData(cohort = "ACC")
url <- psychomics:::parseUrlsFromFirebrowseResponse(res)
```

```
parseValidFile
```

Parse file given a list of file formats

Description

Tries to recognise the file format and parses the content of the given file accordingly.

Usage

```
parseValidFile(file, formats)
```

Arguments

file Character: file to parse
 formats List of file formats to check

Details

The resulting data frame includes the attribute "tablename" with the name of the data frame

Value

Data frame with the contents of the given file if the file format is recognised; otherwise, returns NULL

parseVastToolsEvent *Parses an alternative splicing event from VAST-TOOLS*

Description

Parses an alternative splicing event from VAST-TOOLS

Usage

```
parseVastToolsEvent(event)
```

Arguments

event Data.frame: VAST-TOOLS event containing gene symbol, event ID, length, junctions coordinates, event type and inclusion levels for both samples

Details

Junctions are parsed from

Value

List with the event attributes (chromosome, strand, event type and the position of the exon boundaries)

Note

Only supports to parse one event at a time.

Examples

```
event <- read.table(text =
"NFYA HsaEX0042823 chr6:41046768-41046903 136 chr6:41040823,41046768-41046903,41051785 C2 0 N 0 N"
)
psychomics:::parseVastToolsEvent(event)
```

parseVastToolsSE *Parse junctions of an event from VAST-TOOLS according to event type*

Description

Parse junctions of an event from VAST-TOOLS according to event type

Usage

```
parseVastToolsSE(junctions)
```

```
parseVastToolsRI(junctions, strand)
```

```
parseVastToolsA3SS(junctions)
```

```
parseVastToolsA5SS(junctions)
```

Arguments

| | |
|-----------|---|
| junctions | Data.frame or matrix: exon-exon junctions of alternative splicing events (it must have 4 columns) |
| strand | Character: positive (+) or negative (-) strand |

Details

The following event types are available to be parsed:

- **SE** (skipped exon)
- **RI** (intron retention)
- **A5SS** (alternative 5' splice site)
- **A3SS** (alternative 3' splice site)

Value

List of parsed junctions

See Also

[parseVastToolsEvent](#)

Examples

```
junctions <- read.table(text = "41040823 41046768 41046903 41051785")
psychomics:::parseVastToolsSE(junctions)

# these functions are vectorised!
junctions <- read.table(text = "41040823 41046768 41046903 41051785
                               58864658 58864693 58864294 58864563")
psychomics:::parseVastToolsSE(junctions)

junctions <- read.table(text = "58864658 58864693 58864294 58864563")
psychomics:::parseVastToolsRI(junctions, strand = "+")

junctions <- rbind(
  c(36276385, list(c(36277798, 36277315)), 36277974),
  c(7133604, 7133377, list(c(7133474, 7133456)))
)
psychomics:::parseVastToolsA3SS(junctions)

junctions <- rbind(
  c(74650610, list(c(74650654, 74650658)), 74650982),
  c(list(c(49557666, 49557642), 49557746, 49557470))
)
psychomics:::parseVastToolsA5SS(junctions)
```

| | |
|-----------|--|
| pcaServer | <i>Server logic for the principal component analysis</i> |
|-----------|--|

Description

Server logic for the principal component analysis

Usage

```
pcaServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-------|---|
| pcaUI | <i>User interface of the principal component analysis</i> |
|-------|---|

Description

User interface of the principal component analysis

Usage

```
pcaUI(id)
```

Arguments

| | |
|----|-----------------------|
| id | Character: identifier |
|----|-----------------------|

Value

HTML element

| | |
|------------|---|
| performPCA | <i>Perform principal component analysis after processing missing values from data frame</i> |
|------------|---|

Description

Perform principal component analysis after processing missing values from data frame

Usage

```
performPCA(data, center = TRUE, scale. = FALSE, naTolerance = 0)
```

Arguments

| | |
|-------------|--|
| data | Data frame: data |
| center | a logical value indicating whether the variables should be shifted to be zero centered. Alternately, a vector of length equal the number of columns of x can be supplied. The value is passed to <code>scale</code> . |
| scale. | a logical value indicating whether the variables should be scaled to have unit variance before the analysis takes place. The default is FALSE for consistency with S, but in general scaling is advisable. Alternatively, a vector of length equal the number of columns of x can be supplied. The value is passed to <code>scale</code> . |
| naTolerance | Integer: percentage of NA tolerance |

Value

PCA result in a `prcomp` object

Examples

```
performPCA(USArrests)
```

| | |
|------------------|---|
| plotDistribution | <i>Plot distribution through a density plot</i> |
|------------------|---|

Description

The tooltip shows the median, variance, max, min and number of non-NA samples of each data series.

Usage

```
plotDistribution(psi, groups, rug = TRUE, vLine = TRUE, ..., title = NULL)
```

Arguments

| | |
|--------|---|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |
| rug | Boolean: include rug plot to better visualise data distribution (TRUE by default) |
| vLine | Boolean: include vertical plot lines to indicate the mean and median of each group even when those groups are omitted |
| ... | Extra parameters passed to density to create the kernel density estimates |
| title | Character: plot title |

Value

Highcharter object with density plot

Examples

```
data <- sample(20, rep=TRUE)/20
groups <- c(rep("A", 10), rep("B", 10))
plotDistribution(data, groups)
```

plotMiniSurvivalCurves

Perform and plot survival curves

Description

Perform and plot survival curves

Usage

```
plotMiniSurvivalCurves(i, input, index, survParams, clinical, match, psi,
  censoring, event, timeStart, timeStop)
```

Arguments

| | |
|------------|--|
| i | Integer: index of the survival curves plot of interest |
| input | Shiny input |
| index | Integer: index of event to draw |
| survParams | List of parameters to plot survival curves |
| clinical | Data frame: clinical data |
| match | Integer: samples matched with clinical patients |
| psi | Data frame or matrix: alternative splicing quantification |
| censoring | Character: censor using "left", "right", "interval" or "interval2" |
| event | Character: name of column containing time of the event of interest |
| timeStart | Character: name of column containing starting time of the interval or follow up time |
| timeStop | Character: name of column containing ending time of the interval |

Value

A highchart object to plot

plotPCA *Create a scatterplot from a PCA object*

Description

Create a scatterplot from a PCA object

Usage

```
plotPCA(pca, pcX = 1, pcY = 2, groups = NULL, individuals = TRUE,
        loadings = FALSE)
```

Arguments

| | |
|-------------|--|
| pca | prcomp object |
| pcX | Character: name of the xAxis of interest from the PCA |
| pcY | Character: name of the yAxis of interest from the PCA |
| groups | Matrix: groups to plot indicating the index of interest of the samples (use clinical or sample groups) |
| individuals | Boolean: plot PCA individuals (TRUE by default) |
| loadings | Boolean: plot PCA loadings/rotations (FALSE by default) |

Value

Scatterplot as an Highcharter object

Examples

```
pca <- prcomp(USArrests, scale=TRUE)
plotPCA(pca)
plotPCA(pca, pcX=2, pcY=3)

# Plot both individuals and loadings
plotPCA(pca, pcX=2, pcY=3, loadings=TRUE)
```

plotPointsStyle *Interface to modify the style of the plot points*

Description

Interface to modify the style of the plot points

Usage

```
plotPointsStyle(ns, id, description, help = NULL, size = 2,
               colour = "black", alpha = 1)
```

Arguments

| | |
|-------------|--|
| ns | Namespace function |
| id | Character: identifier |
| description | Character: display text for user |
| help | Character: extra text to help the user |
| size | Integer: default size (2 by default) |
| colour | Character: default colour ("black" by default) |
| alpha | Numeric: default transparency value; (opaque by default) |

Value

HTML elements

| | |
|-------------|------------------------------|
| plotProtein | <i>Plot protein features</i> |
|-------------|------------------------------|

Description

Plot protein features

Usage

```
plotProtein(molecule)
```

Arguments

| | |
|----------|---|
| molecule | Character: UniProt protein or Ensembl transcript identifier |
|----------|---|

Value

highchart object

Examples

```
## Not run:  
protein <- "P38398"  
plotProtein(protein)  
  
transcript <- "ENST00000488540"  
plotProtein(transcript)  
  
## End(Not run)
```

plotSurvivalCurves *Plot survival curves*

Description

Plot survival curves

Usage

```
plotSurvivalCurves(surv, mark = TRUE, interval = FALSE, pvalue = NULL,
  title = "Survival analysis", scale = NULL)
```

Arguments

| | |
|----------|---|
| surv | Survival object |
| mark | Boolean: mark times? TRUE by default |
| interval | Boolean: show interval ranges? FALSE by default |
| pvalue | Numeric: p-value of the survival curves |
| title | Character: plot title |
| scale | Character: time scale; default is "days" |

Value

Plot of survival curves

Examples

```
require("survival")
fit <- survfit(Surv(time, status) ~ x, data = aml)
plotSurvivalCurves(fit)
```

plottableXranges *HTML code to plot a X-ranges series*

Description

HTML code to plot a X-ranges series

Usage

```
plottableXranges(hc, shiny = FALSE)
```

Arguments

| | |
|-------|---|
| hc | highcharter object |
| shiny | Boolean: is the function running in a Shiny session? FALSE by default |

Value

HTML elements

| | |
|-----------------|-------------------------|
| plotTranscripts | <i>Plot transcripts</i> |
|-----------------|-------------------------|

Description

Plot transcripts

Usage

```
plotTranscripts(info, eventPosition = NULL, shiny = FALSE)
```

Arguments

| | |
|---------------|---|
| info | Information retrieved from ENSEMBL |
| eventPosition | Numeric: coordinates of the alternative splicing event; NULL by default |
| shiny | Boolean: is the function running in a Shiny session? FALSE by default |

Value

NULL (this function is used to modify the Shiny session's state)

Examples

```
event <- "SE_12_-_7985318_7984360_7984200_7982602_SLC2A14"  
info <- queryEnsemblByEvent(event, species="human", assembly="hg19")  
pos <- parseSplicingEvent(event)$pos[[1]]  
## Not run:  
plotTranscripts(info, pos)  
  
## End(Not run)
```

| | |
|--------------|---|
| plotVariance | <i>Create the explained variance plot</i> |
|--------------|---|

Description

Create the explained variance plot

Usage

```
plotVariance(pca)
```

Arguments

| | |
|-----|------------|
| pca | PCA values |
|-----|------------|

Value

Plot variance as an Highcharter object

Examples

```
pca <- prcomp(USArrests)
plotVariance(pca)
```

```
prepareAnnotationFromEvents
```

Prepare annotation from alternative splicing events

Description

In case more than one data frame with alternative splicing events is given, the events are cross-referenced according to the chromosome, strand and relevant coordinates per event type (see details).

Usage

```
prepareAnnotationFromEvents(...)
```

Arguments

... Data frame(s) of alternative splicing events to include in the annotation

Details

Events from two or more data frames are cross-referenced based on each event's chromosome, strand and specific coordinates relevant for each event type:

- Skipped exon: constitutive exon 1 end, alternative exon (start and end) and constitutive exon 2 start
- Mutually exclusive exon: constitutive exon 1 end, alternative exon 1 and 2 (start and end) and constitutive exon 2 start
- Alternative 5' splice site: constitutive exon 1 end, alternative exon 1 end and constitutive exon 2 start
- Alternative first exon: same as alternative 5' splice site
- Alternative 3' splice site: constitutive exon 1 end, alternative exon 1 start and constitutive exon 2 start
- Alternative last exon: same as alternative 3' splice site

Value

List of data frames with the annotation from different data frames joined by event type

Note

When cross-referencing events, gene information is discarded.

Examples

```
# Load sample files (SUPPA annotation)
folder <- "extdata/eventsAnnotSample/suppa_output/suppaEvents"
suppaOutput <- system.file(folder, package="psychomics")

# Parse and prepare SUPPA annotation
suppa <- parseSuppaAnnotation(suppaOutput)
annot <- prepareAnnotationFromEvents(suppa)

# Load sample files (rMATS annotation)
folder <- "extdata/eventsAnnotSample/mats_output/ASEvents/"
matsOutput <- system.file(folder, package="psychomics")

# Parse rMATS annotation and prepare combined annotation from rMATS and SUPPA
mats <- parseMatsAnnotation(matsOutput)
annot <- prepareAnnotationFromEvents(suppa, mats)
```

```
prepareFirebrowseArchives
```

Prepares Firebrowse archives in a given directory

Description

Checks Firebrowse archives' integrity using the MD5 files, extracts the content of the archives, moves the content to newly-created folders and removes the original downloaded archives.

Usage

```
prepareFirebrowseArchives(archive, md5, folder, outdir)
```

```
prepareFirehoseArchives(archive, md5, folder, outdir)
```

Arguments

| | |
|---------|---|
| archive | Character: path to downloaded archives |
| md5 | Character: path to MD5 files of each archive |
| folder | Character: master directory where every archive will be extracted |
| outdir | Character: subdirectories where to move the extracted content |

Value

Invisible TRUE if successful

Examples

```
file <- paste0(
  "~/Downloads",
  "ACC/20151101/gdac.broadinstitute.org_ACC.",
  "Merge_Clinical.Level_1.2015110100.0.0.tar.gz")
md5 <- paste0(file, ".md5")
## Not run:
prepareFirebrowseArchives(archive = file, md5 = paste0(file, ".md5"))
```

```
## End(Not run)
```

```
processButton      Style button used to initiate a process
```

Description

Style button used to initiate a process

Usage

```
processButton(id, label, ..., class = "btn-primary")
```

Arguments

| | |
|-------|---|
| id | Character: button identifier |
| label | Character: label |
| ... | Extra parameters to pass to <code>actionButton</code> |
| class | Character: class |

Value

HTML for a button

```
processDatasetNames Process dataset names
```

Description

Process dataset names

Usage

```
processDatasetNames(data)
```

Arguments

| | |
|------|------------------------------|
| data | List of lists of data frames |
|------|------------------------------|

Details

Avoid duplicated names and append the technology used for junction quantification

Value

Processed list of lists of data frames

| | |
|-----------------|---|
| processSurvData | <i>Process survival data to calculate survival curves</i> |
|-----------------|---|

Description

Process survival data to calculate survival curves

Usage

```
processSurvData(event, timeStart, timeStop, followup, group, clinical,
  survTime = NULL)
```

Arguments

| | |
|-----------|--|
| event | Character: name of column containing time of the event of interest |
| timeStart | Character: name of column containing starting time of the interval or follow up time |
| timeStop | Character: name of column containing ending time of the interval |
| followup | Character: name of column containing follow up time |
| group | Character: group of each individual |
| clinical | Data frame: clinical data |
| survTime | survTime object: Times to follow up, time start, time stop and event (optional) |

Details

The event time will only be used to determine whether the event has occurred (1) or not (0) in case of missing values.

If survTime is NULL, the survival times will be fetch from the clinical dataset according to the names given in timeStart, timeStop, event and followup. This can became quite slow when using the function in a for loop. If these variables are constant, consider running the function [getColumnnsTime](#) to retrieve the time of such columns once and hand the result to the survTime argument of this function.

Value

Data frame with terms needed to calculate survival curves

| | |
|-----------------|---|
| processSurvival | <i>Check if survival analyses successfully completed or returned errors</i> |
|-----------------|---|

Description

Check if survival analyses successfully completed or returned errors

Usage

```
processSurvival(session, ...)
```

Arguments

session Shiny session
 ... Arguments to pass to function processSurvTerms

Value

List with survival analysis results

processSurvTerms *Process survival curves terms to calculate survival curves*

Description

Process survival curves terms to calculate survival curves

Usage

```
processSurvTerms(clinical, censoring, event, timeStart, timeStop = NULL,
  group = NULL, formulaStr = NULL, coxph = FALSE, scale = "days",
  followup = "days_to_last_followup", survTime = NULL)
```

Arguments

clinical Data frame: clinical data
 censoring Character: censor using "left", "right", "interval" or "interval2"
 event Character: name of column containing time of the event of interest
 timeStart Character: name of column containing starting time of the interval or follow up time
 timeStop Character: name of column containing ending time of the interval
 group Character: group of each individual
 formulaStr Character: formula to use
 coxph Boolean: fit a Cox proportional hazards regression model? FALSE by default
 scale Character: rescale the survival time to "days", "weeks", "months" or "years"
 followup Character: name of column containing follow up time
 survTime survTime object: times to follow up, time start, time stop and event (optional)

Details

timeStop is only considered if censoring is either interval or interval2

If survTime is NULL, the survival times will be fetch from the clinical dataset according to the names given in timeStart, timeStop, event and followup. This can became quite slow when using the function in a for loop. If these variables are constant, consider running the function [getColumnTime](#) to retrieve the time of such columns once and hand the result to the survTime argument of this function.

Value

A list with a formula object and a data frame with terms needed to calculate survival curves

Examples

```
clinical <- read.table(text = "2549  NA ii  female
                             840  NA i   female
                             NA 1204 iv  male
                             NA  383 iv  female
                             1293  NA iii male
                             NA 1355 ii  male")
names(clinical) <- c("patient.days_to_last_followup",
                    "patient.days_to_death",
                    "patient.stage_event.pathologic_stage",
                    "patient.gender")
timeStart <- "days_to_death"
event <- "days_to_death"
formulaStr <- "patient.stage_event.pathologic_stage + patient.gender"
survTerms <- processSurvTerms(clinical, censoring="right", event, timeStart,
                              formulaStr=formulaStr)
```

psychomics

Start graphical interface of PSICHOMICS

Description

Start graphical interface of PSICHOMICS

Usage

```
psychomics(..., reset = FALSE)
```

Arguments

| | |
|-------|---|
| ... | Parameters to pass to the function runApp |
| reset | Boolean: reset Shiny session? FALSE by default; requires the package devtools to reset data |

Value

NULL (this function is used to modify the Shiny session's state)

Examples

```
## Not run:
psychomics()

## End(Not run)
```

| | |
|----------|--|
| pubmedUI | <i>Return the interface of relevant PubMed articles for a given gene</i> |
|----------|--|

Description

Return the interface of relevant PubMed articles for a given gene

Usage

```
pubmedUI(gene, ...)
```

Arguments

| | |
|------|---|
| gene | Character: gene |
| ... | Arguments to pass to queryPubMed function |

Value

HTML interface of relevant PubMed articles

| | |
|------------------|---|
| quantifySplicing | <i>Quantify alternative splicing events</i> |
|------------------|---|

Description

Quantify alternative splicing events

Usage

```
quantifySplicing(annotation, junctionQuant, eventType = c("SE", "MXE", "ALE",
  "AFE", "A3SS", "A5SS"), minReads = 10, progress = echoProgress)
```

Arguments

| | |
|---------------|---|
| annotation | List of data frames: annotation for each alternative splicing event type |
| junctionQuant | Data frame: junction quantification |
| eventType | Character: splicing event types to quantify |
| minReads | Integer: minimum of read counts to consider a junction read in calculations |
| progress | Function to track the progress |

Value

Data frame with the quantification of the alternative splicing events

Examples

```
# Calculate PSI for skipped exon (SE) and mutually exclusive (MXE) events
annot <- readRDS("ex_splicing_annotation.RDS")
junctionQuant <- readRDS("ex_junctionQuant.RDS")

psi <- quantifySplicing(annot, junctionQuant, eventType=c("SE", "MXE"))
```

queryEnsembl *Query the Ensembl REST API*

Description

Query the Ensembl REST API

Usage

```
queryEnsembl(path, query, grch37 = TRUE)
```

Arguments

| | |
|--------|--|
| path | Character: API path |
| query | Character: API query |
| grch37 | Boolean: query the Ensembl GRCh37 API? TRUE by default; otherwise, query the most recent API |

Value

Parsed response or NULL if there's no response

Examples

```
path <- "overlap/region/human/7:140424943-140624564"
query <- list(feature = "gene")
psychomics:::queryEnsembl(path, query, grch37 = TRUE)

path <- "lookup/symbol/human/BRCA2"
query <- list(expand=1)
psychomics:::queryEnsembl(path, query, grch37 = TRUE)
```

queryEnsemblByEvent *Query information from Ensembl by a given alternative splicing event*

Description

Query information from Ensembl by a given alternative splicing event

Usage

```
queryEnsemblByEvent(event, ...)
```

Arguments

| | |
|-------|--|
| event | Character: alternative splicing event identifier |
| ... | Arguments to pass to queryEnsemblByGene |

Value

Information from Ensembl

Examples

```
event <- c("SE_17_-_41251792_41249306_41249261_41246877_BRCA1")
queryEnsemblByEvent(event, species="human", assembly="hg19")
```

queryEnsemblByGene *Query information from Ensembl by a given gene*

Description

Query information from Ensembl by a given gene

Usage

```
queryEnsemblByGene(gene, species = NULL, assembly = NULL)
```

Arguments

| | |
|----------|---|
| gene | Character: gene identifier |
| species | Character: species (can be NULL when handling an ENSEMBL identifier) |
| assembly | Character: assembly version (can be NULL when handling an ENSEMBL identifier) |

Value

Information from Ensembl

Examples

```
queryEnsemblByGene("BRCA1", "human", "hg19")
queryEnsemblByGene("ENSG00000139618")
```

queryFirebrowseData *Query the Firebrowse web API for TCGA data*

Description

Query the Firebrowse web API for TCGA data

Usage

```
queryFirebrowseData(format = "json", date = NULL, cohort = NULL,
  data_type = NULL, tool = NULL, platform = NULL, center = NULL,
  level = NULL, protocol = NULL, page = NULL, page_size = NULL,
  sort_by = NULL)
```

```
queryFirehoseData(format = "json", date = NULL, cohort = NULL,
  data_type = NULL, tool = NULL, platform = NULL, center = NULL,
  level = NULL, protocol = NULL, page = NULL, page_size = NULL,
  sort_by = NULL)
```

Arguments

| | |
|-----------|---|
| format | Character: response format as JSON (default), CSV or TSV |
| date | Character: dates of the data retrieval by Firebrowse (by default, it uses the most recent data available) |
| cohort | Character: abbreviation of the cohorts (by default, returns data for all cohorts) |
| data_type | Character: data types (optional) |
| tool | Character: data produced by the selected Firebrowse tools (optional) |
| platform | Character: data generation platforms (optional) |
| center | Character: data generation centers (optional) |
| level | Integer: data levels (optional) |
| protocol | Character: sample characterization protocols (optional) |
| page | Integer: page of the results to return (optional) |
| page_size | Integer: number of records per page of results; max is 2000 (optional) |
| sort_by | String: column used to sort the data (by default, it sorts by cohort) |

Value

Response from the Firebrowse web API (it needs to be parsed)

Examples

```
cohort <- psychomics::getFirebrowseCohorts()[1]
psychomics::queryFirebrowseData(cohort = cohort, data_type = "mRNASeq")

# Querying for data from a specific date
dates <- psychomics::getFirebrowseDates()
dates <- format(dates, psychomics::getFirebrowseDateFormat()$query)

psychomics::queryFirebrowseData(date = dates[2], cohort = cohort)
```

queryPubMed

Query the PubMed REST API

Description

Query the PubMed REST API

Usage

```
queryPubMed(primary, ..., top = 3, field = "abstract", sort = "relevance")
```

Arguments

| | |
|---------|--|
| primary | Character: primary search term |
| ... | Character: other relevant search terms |
| top | Numeric: number of articles to retrieve (3 by default) |
| field | Character: field of interest where to look for terms ("abstract" by default) |
| sort | Character: sort by a given parameter ("relevance" by default) |

Value

Parsed response

Examples

```
psychomics:::queryPubMed("BRCA1", "cancer", "adrenocortical carcinoma")
```

| | |
|--------------|-----------------------------------|
| queryUniprot | <i>Query the Uniprot REST API</i> |
|--------------|-----------------------------------|

Description

Query the Uniprot REST API

Usage

```
queryUniprot(molecule, format = "xml")
```

Arguments

| | |
|----------|---|
| molecule | Character: protein or transcript to query |
| format | Character: format of the response |

Value

Parsed response

Examples

```
protein <- "P51587"
format <- "xml"
psychomics:::queryUniprot(protein, format)

transcript <- "ENST00000488540"
format <- "xml"
psychomics:::queryUniprot(transcript, format)
```

| | |
|----------|------------------------|
| readFile | <i>Load local file</i> |
|----------|------------------------|

Description

Load local file

Usage

```
readFile(file)
```

Arguments

| | |
|------|-----------------------------|
| file | Character: path to the file |
|------|-----------------------------|

Value

Loaded file

Examples

```
junctionQuant <- readfile("ex_junctionQuant.RDS")
```

| | |
|------------------|---|
| renameDuplicated | <i>Rename vector to avoid duplicated values with another vector</i> |
|------------------|---|

Description

Renames values by adding an index to the end of duplicates. This allows to prepare unique values in two vectors before a merge, for instance.

Usage

```
renameDuplicated(check, comp)
```

Arguments

| | |
|-------|---|
| check | Character: values to rename if duplicated |
| comp | Character: values to compare with |

Value

Character vector with renamed values if duplicated; else, it returns the usual values. It doesn't return the comparator values.

Examples

```
psychomics:::renameDuplicated(check = c("blue", "red"), comp = c("green",  
"blue"))
```

| | |
|--------------|---|
| renameGroups | <i>Rename duplicated names from a new group</i> |
|--------------|---|

Description

Rename duplicated names from a new group

Usage

```
renameGroups(new, old)
```

Arguments

| | |
|-----|-----------------------------|
| new | Matrix: new groups |
| old | Matrix: pre-existing groups |

Value

Character with no duplicated group names

Note

The names of pre-existing groups are not modified.

renderDataTableSparklines

Render a data table with Sparkline HTML elements

Description

Render a data table with Sparkline HTML elements

Usage

```
renderDataTableSparklines(..., options = NULL)
```

Arguments

... Arguments to pass to [renderDataTable](#)
options List of options to pass to [renderDataTable](#)

Details

This slightly modified version of [renderDataTable](#) calls a JavaScript function to convert the Sparkline HTML elements to interactive Highcharts

Value

NULL (this function is used to modify the Shiny session's state)

renderGeneticInfo *Render genetic information*

Description

Render genetic information

Usage

```
renderGeneticInfo(ns, info, species = NULL, assembly = NULL,  
grch37 = FALSE)
```

Arguments

| | |
|----------|---|
| ns | Namespace function |
| info | Information as retrieved from ENSEMBL |
| species | Character: species name (NULL by default) |
| assembly | Character: assembly version (NULL by default) |
| grch37 | Boolean: use version GRCh37 of the genome? FALSE by default |

Value

HTML elements to render gene, protein and transcript annotation

| | |
|-------------------|-----------------------------------|
| renderProteinInfo | <i>Render protein information</i> |
|-------------------|-----------------------------------|

Description

Render protein information

Usage

```
renderProteinInfo(protein, transcript, species, assembly)
```

Arguments

| | |
|------------|--|
| protein | Character: protein identifier |
| transcript | Character: Ensembl identifier of the protein's respective transcript |
| species | Character: species |
| assembly | Character: assembly |

Value

HTML elements

| | |
|---------|---|
| rm.null | <i>Filter NULL elements from vector or list</i> |
|---------|---|

Description

Filter NULL elements from vector or list

Usage

```
rm.null(v)
```

Arguments

| | |
|---|----------------|
| v | Vector or list |
|---|----------------|

Value

Filtered vector or list with no NULL elements; if the input is a vector composed of only NULL elements, it returns a NULL (note that it will returns an empty list if the input is a list with only NULL elements)

| | |
|-------------|--|
| roundDigits | <i>Round by the given number of digits</i> |
|-------------|--|

Description

Round by the given number of digits

Usage

```
roundDigits(n)
```

Arguments

| | |
|---|----------------------------|
| n | Numeric: number to roundhf |
|---|----------------------------|

Value

Formatted number with a given numeric precision

| | |
|--------|-------------------------------|
| rowVar | <i>Sample variance by row</i> |
|--------|-------------------------------|

Description

Calculate the sample variance of each row in the given matrix

Usage

```
rowVar(x, na.rm = FALSE)
```

Arguments

| | |
|-------|--|
| x | Matrix |
| na.rm | Boolean: should the NAs be ignored? FALSE by default |

Value

Variance for each row

selectGroupsServer *Group selection logic*

Description

Group selection logic

Usage

```
selectGroupsServer(session, id)
```

Arguments

| | |
|---------|--|
| session | Shiny session |
| id | Character: identifier of the group selection |

Value

Server logic for group selection

selectGroupsUI *Group selection interface*

Description

Group selection interface

Usage

```
selectGroupsUI(id, label,  
  placeholder = "Click on 'Groups' to create or edit groups",  
  noGroupsLabel = NULL, groupsLabel = NULL)
```

Arguments

| | |
|---------------|---|
| id | Character: identifier of the group selection |
| label | Character: selectize label |
| placeholder | Character: selectize placeholder |
| noGroupsLabel | Character: label to show when no groups may be selected (if NULL, the option to show no groups will not be shown) |
| groupsLabel | Character: label to show to the option of using groups when no groups may be selected |

Value

Interface for group selection

Note

To allow the user to (explicitly) select no groups, pass the noGroupsLabel and groupsLabel arguments.

See Also

selectGroupsServer getSelectedGroups

setActiveDataset *Set active dataset*

Description

Set active dataset

Usage

```
setActiveDataset(dataset)
```

Arguments

dataset Character: dataset

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setAssemblyVersion *Set the assembly version of a data category*

Description

Set the assembly version of a data category

Usage

```
setAssemblyVersion(value, category = getCategory())
```

Arguments

value Character: assembly version
category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|-------------------|---|
| setAutoNavigation | <i>Set if history browsing is automatic</i> |
|-------------------|---|

Description

Set if history browsing is automatic

Usage

```
setAutoNavigation(param)
```

Arguments

| | |
|-------|--|
| param | Boolean: is navigation of browser history automatic? |
|-------|--|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|-------------|--------------------------|
| setCategory | <i>Set data category</i> |
|-------------|--------------------------|

Description

Set data category

Usage

```
setCategory(category)
```

Arguments

| | |
|----------|--------------------------|
| category | Character: data category |
|----------|--------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setClinicalMatchFrom *Set clinical matches from a given data type*

Description

Set clinical matches from a given data type

Usage

```
setClinicalMatchFrom(dataset, matches, category = getCategory())
```

Arguments

| | |
|----------|--|
| dataset | Character: data set (e.g. "Clinical data") |
| matches | Vector of integers: clinical matches of dataset |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setCores *Set number of cores*

Description

Set number of cores

Usage

```
setCores(cores)
```

Arguments

| | |
|-------|----------------------------|
| cores | Character: number of cores |
|-------|----------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|---------|------------------------------------|
| setData | <i>Set data of the global data</i> |
|---------|------------------------------------|

Description

Set data of the global data

Usage

```
setData(data)
```

Arguments

| | |
|------|-------------------------------------|
| data | Data frame or matrix to set as data |
|------|-------------------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|-------------------------|--|
| setDifferentialAnalyses | <i>Set the table of differential analyses of a data category</i> |
|-------------------------|--|

Description

Set the table of differential analyses of a data category

Usage

```
setDifferentialAnalyses(table, category = getCategory())
```

Arguments

| | |
|----------|--|
| table | Character: differential analyses table |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

`setDifferentialAnalysesFiltered`*Set the filtered table of differential analyses of a data category*

Description

Set the filtered table of differential analyses of a data category

Usage

```
setDifferentialAnalysesFiltered(table, category = getCategory())
```

Arguments

| | |
|-----------------------|--|
| <code>table</code> | Character: filtered differential analyses table |
| <code>category</code> | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

`setDifferentialAnalysesHighlightedEvents`*Set highlighted events from differential analyses of a data category*

Description

Set highlighted events from differential analyses of a data category

Usage

```
setDifferentialAnalysesHighlightedEvents(events, category = getCategory())
```

Arguments

| | |
|-----------------------|--|
| <code>events</code> | Integer: indexes relative to a table of differential analyses |
| <code>category</code> | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

`setDifferentialAnalysesSelected`*Set selected points in the differential analysis table of a data category*

Description

Set selected points in the differential analysis table of a data category

Usage

```
setDifferentialAnalysesSelected(points, category = getCategory())
```

Arguments

| | |
|-----------------------|--|
| <code>points</code> | Integer: index of selected points |
| <code>category</code> | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

`setDifferentialAnalysesSurvival`*Set the table of differential analyses' survival data of a data category*

Description

Set the table of differential analyses' survival data of a data category

Usage

```
setDifferentialAnalysesSurvival(table, category = getCategory())
```

Arguments

| | |
|-----------------------|--|
| <code>table</code> | Character: differential analyses' survival data |
| <code>category</code> | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

`setDifferentialAnalysesZoom`*Set plot coordinates for zooming from differential analyses of a data category*

Description

Set plot coordinates for zooming from differential analyses of a data category

Usage

```
setDifferentialAnalysesZoom(zoom, category = getCategory())
```

Arguments

| | |
|-----------------------|--|
| <code>zoom</code> | Integer: X and Y coordinates |
| <code>category</code> | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

`setEvent`*Set event*

Description

Set event

Usage

```
setEvent(event)
```

Arguments

| | |
|--------------------|------------------|
| <code>event</code> | Character: event |
|--------------------|------------------|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|-------------------|---------------------------------|
| setFirebrowseData | <i>Set data from Firebrowse</i> |
|-------------------|---------------------------------|

Description

Set data from Firebrowse

Usage

```
setFirebrowseData(input, output, session, replace = TRUE)
```

Arguments

| | |
|---------|---|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |
| replace | Boolean: replace loaded data? TRUE by default |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|-----------|---|
| setGlobal | <i>Set element as globally accessible</i> |
|-----------|---|

Description

Set element as globally accessible

Usage

```
setGlobal(..., value, sep = "_")
```

Arguments

| | |
|-------|--------------------------------------|
| ... | Arguments to identify a variable |
| value | Any value to attribute to an element |
| sep | Character to separate identifier |

Details

Set element inside the global variable

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setGroupsFrom *Set groups from a given data type*

Description

Set groups from a given data type

Usage

```
setGroupsFrom(dataset, groups, category = getCategory())
```

Arguments

| | |
|----------|--|
| dataset | Character: data set (e.g. "Clinical data") |
| groups | Matrix: groups of dataset |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setInclusionLevels *Set inclusion levels for a given data category*

Description

Set inclusion levels for a given data category

Usage

```
setInclusionLevels(value, category = getCategory())
```

Arguments

| | |
|----------|--|
| value | Data frame or matrix: inclusion levels |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setInclusionLevelsPCA *Get principal component analysis based on inclusion levels*

Description

Get principal component analysis based on inclusion levels

Usage

```
setInclusionLevelsPCA(pca, category = getCategory())
```

Arguments

| | |
|----------|--|
| pca | prcomp object (PCA) of inclusion levels |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

setLocalData *Load local files*

Description

Load local files

Usage

```
setLocalData(input, output, session, replace = TRUE)
```

Arguments

| | |
|---------|---|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |
| replace | Boolean: replace loaded data? TRUE by default |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|--------------|---|
| setPatientId | <i>Set the identifier of patients for a data category</i> |
|--------------|---|

Description

Set the identifier of patients for a data category

Usage

```
setPatientId(value, category = getCategory())
```

Arguments

| | |
|----------|--|
| value | Character: identifier of patients |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|--------------|-------------------------------------|
| setPrecision | <i>Set number of decimal places</i> |
|--------------|-------------------------------------|

Description

Set number of decimal places

Usage

```
setPrecision(precision)
```

Arguments

| | |
|-----------|-----------------------------------|
| precision | Numeric: number of decimal places |
|-----------|-----------------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|-------------|--|
| setSampleId | <i>Set the identifier of samples for a data category</i> |
|-------------|--|

Description

Set the identifier of samples for a data category

Usage

```
setSampleId(value, category = getCategory())
```

Arguments

| | |
|----------|--|
| value | Character: identifier of samples |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|---------------|---|
| setSampleInfo | <i>Set sample information for a given data category</i> |
|---------------|---|

Description

Set sample information for a given data category

Usage

```
setSampleInfo(value, category = getCategory())
```

Arguments

| | |
|----------|--|
| value | Data frame or matrix: sample information |
| category | Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category |

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|----------------|---|
| setSignificant | <i>Set number of significant digits</i> |
|----------------|---|

Description

Set number of significant digits

Usage

```
setSignificant(significant)
```

Arguments

significant Character: number of significant digits

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|------------|---|
| setSpecies | <i>Set the species of a data category</i> |
|------------|---|

Description

Set the species of a data category

Usage

```
setSpecies(value, category = getCategory())
```

Arguments

value Character: species
category Character: data category (e.g. "Carcinoma 2016"); by default, it uses the selected data category

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|------------------|----------------------------------|
| setURLtoDownload | <i>Set URL links to download</i> |
|------------------|----------------------------------|

Description

Set URL links to download

Usage

```
setURLtoDownload(url)
```

Arguments

| | |
|-----|----------------------------------|
| url | Character: URL links to download |
|-----|----------------------------------|

Value

NULL (this function is used to modify the Shiny session's state)

Note

Needs to be called inside a reactive function

| | |
|-----------|--------------------------------|
| showAlert | <i>Show or remove an alert</i> |
|-----------|--------------------------------|

Description

You can also use `errorAlert` and `warningAlert` to use template alerts already stylised to show errors and warnings respectively.

Usage

```
showAlert(session, ..., title = NULL, style = NULL, dismissable = TRUE,  
  alertId = "alert")
```

```
errorAlert(session, ..., title = NULL, dismissable = TRUE,  
  alertId = "alert")
```

```
warningAlert(session, ..., title = NULL, dismissable = TRUE,  
  alertId = "alert")
```

```
removeAlert(output, alertId = "alert")
```

Arguments

| | |
|-------------|---|
| session | Shiny session |
| ... | Arguments to render as elements of alert |
| title | Character: title of the alert (optional) |
| style | Character: style of the alert ("alert-danger", "alert-warning" or NULL) |
| dismissable | Boolean: is the alert dismissable? TRUE by default |
| alertId | Character: alert identifier |
| output | Shiny output |

Value

NULL (this function is used to modify the Shiny session's state)

See Also

showModal

| | |
|-----------------|-----------------------------|
| showGroupsTable | <i>Present groups table</i> |
|-----------------|-----------------------------|

Description

Present groups table

Usage

```
showGroupsTable(datasetName)
```

Arguments

| | |
|-------------|----------------------------|
| datasetName | Character: name of dataset |
|-------------|----------------------------|

Value

Matrix with groups ordered (or NULL if no groups exist)

| | |
|--------------|---|
| signifDigits | <i>Get number of significant digits</i> |
|--------------|---|

Description

Get number of significant digits

Usage

```
signifDigits(n)
```

Arguments

| | |
|---|--------------------------|
| n | Numeric: number to round |
|---|--------------------------|

Value

Formatted number with a given number of significant digits

| | |
|--------------------|---|
| singleDiffAnalyses | <i>Perform statistical analysis on a given splicing event</i> |
|--------------------|---|

Description

Perform statistical analyses on a given vector containing elements from different groups

Usage

```
singleDiffAnalyses(vector, group, threshold = 1, step = 100,
  analyses = c("wilcoxRankSum", "ttest", "kruskal", "levene", "fligner"))
```

Arguments

| | |
|-----------|--|
| vector | Numeric |
| group | Character: group of each element in the vector |
| threshold | Integer: minimum number of data points to perform analysis in a group (default is 1) |
| step | Numeric: number of events before the progress bar is updated (a bigger number allows for a faster execution) |
| analyses | Character: analyses to perform (see "Details") |

Details

The following statistical analyses may be performed by including the respective string in the analysis argument:

- ttest - Unpaired t-test (2 groups)
- wilcoxRankSum - Wilcoxon Rank Sum test (2 groups)
- kruskal - Kruskal test (2 or more groups)
- levene - Levene's test (2 or more groups)
- fligner - Fligner-Killeen test (2 or more groups)

Value

A row from a data frame with the results

| | |
|-----------------|--|
| sortCoordinates | <i>Sort coordinates for some event types</i> |
|-----------------|--|

Description

Some programs sort the coordinates of specific event types differently. To make them all comparable across programs, the coordinates are ordered by increasing (plus strand) or decreasing order (minus strand)

Usage

```
sortCoordinates(events)
```

Arguments

| | |
|--------|--|
| events | List of data frames with alternative splicing events for a given program |
|--------|--|

Value

List of data frames with alternative splicing events for a given program

| | |
|----------|---|
| spearman | <i>Perform Spearman's test and return interface to show the results</i> |
|----------|---|

Description

Perform Spearman's test and return interface to show the results

Usage

```
spearman(psi, groups)
```

Arguments

| | |
|--------|---|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |

Value

HTML elements

| | |
|--------------|--|
| startProcess | <i>Signal the program that a process is starting</i> |
|--------------|--|

Description

Style button to show processing is in progress

Usage

```
startProcess(id)
```

Arguments

| | |
|----|------------------------------|
| id | Character: button identifier |
|----|------------------------------|

Value

Start time of the process

| | |
|---------------|---------------------------------|
| startProgress | <i>Create a progress object</i> |
|---------------|---------------------------------|

Description

Create a progress object

Usage

```
startProgress(message, divisions, global = sharedData)
```

Arguments

| | |
|-----------|--|
| message | Character: progress message |
| divisions | Integer: number of divisions in the progress bar |
| global | Shiny's global variable |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|-------------------------------|
| styleModal | <i>Style and show a modal</i> |
|------------|-------------------------------|

Description

You can also use `errorModal` and `warningModal` to use template modals already stylised to show errors and warnings, respectively.

Usage

```
styleModal(session, title, ..., style = NULL,  
           iconName = "exclamation-circle", footer = NULL, echo = FALSE,  
           size = "medium", dismissButton = TRUE)
```

```
errorModal(session, title, ..., size = "small", footer = NULL)
```

```
warningModal(session, title, ..., size = "small", footer = NULL)
```

```
infoModal(session, title, ..., size = "small", footer = NULL)
```

Arguments

| | |
|----------------------------|---|
| <code>session</code> | Current Shiny session |
| <code>title</code> | Character: modal title |
| <code>...</code> | Extra arguments to pass to <code>shiny::modalDialog</code> |
| <code>style</code> | Character: style of the modal (NULL, "warning", "error" or "info"; NULL by default) |
| <code>iconName</code> | Character: FontAwesome icon name to appear with the title |
| <code>footer</code> | HTML elements to use in footer |
| <code>echo</code> | Boolean: print to console? FALSE by default |
| <code>size</code> | Character: size of the modal - "medium" (default), "small" or "large" |
| <code>dismissButton</code> | Boolean: show dismiss button in footer? TRUE by default |

Value

NULL (this function is used to modify the Shiny session's state)

See Also

`showAlert`

| | |
|--------------------|---|
| survdiff.survTerms | <i>Test difference between two or more survival curves using processed survival terms</i> |
|--------------------|---|

Description

Test difference between two or more survival curves using processed survival terms

Usage

```
survdiff.survTerms(survTerms, ...)
```

Arguments

| | |
|-----------|--|
| survTerms | survTerms object: processed survival terms |
| ... | Extra arguments passed to survdiff |

Value

an object of class "survfit". See survfit.object for details. Methods defined for survfit objects are print, plot, lines, and points.

Examples

```
clinical <- read.table(text = "2549  NA ii  female
                             840  NA i   female
                             NA 1204 iv  male
                             NA  383 iv  female
                             1293  NA iii male
                             NA 1355 ii  male")
names(clinical) <- c("patient.days_to_last_followup",
                    "patient.days_to_death",
                    "patient.stage_event.pathologic_stage",
                    "patient.gender")
timeStart <- "days_to_death"
event <- "days_to_death"
formulaStr <- "patient.stage_event.pathologic_stage + patient.gender"
survTerms <- processSurvTerms(clinical, censoring="right", event, timeStart,
                              formulaStr=formulaStr)
survdiff.survTerms(survTerms)
```

| | |
|-------------------|--|
| survfit.survTerms | <i>Compute estimate of a survival curve using processed survival terms</i> |
|-------------------|--|

Description

Compute estimate of a survival curve using processed survival terms

Usage

```
## S3 method for class 'survTerms'
survfit(survTerms, ...)
```

Arguments

survTerms survTerms object: processed survival terms
 ... Extra arguments passed to survfit

Value

an object of class "survfit". See survfit.object for details. Methods defined for survfit objects are print, plot, lines, and points.

Examples

```
clinical <- read.table(text = "2549  NA ii  female
                             840  NA i   female
                             NA 1204 iv  male
                             NA  383 iv  female
                             1293  NA iii male
                             NA 1355 ii  male")
names(clinical) <- c("patient.days_to_last_followup",
                    "patient.days_to_death",
                    "patient.stage_event.pathologic_stage",
                    "patient.gender")
timeStart <- "days_to_death"
event     <- "days_to_death"
formulaStr <- "patient.stage_event.pathologic_stage + patient.gender"
survTerms <- processSurvTerms(clinical, censoring="right", event, timeStart,
                              formulaStr=formulaStr)

require("survival")
survfit(survTerms)
```

 survivalServer

Server logic of survival analysis

Description

Server logic of survival analysis

Usage

```
survivalServer(input, output, session)
```

Arguments

input Shiny input
 output Shiny output
 session Shiny session

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|--|
| survivalUI | <i>User interface of survival analysis</i> |
|------------|--|

Description

User interface of survival analysis

Usage

```
survivalUI(id)
```

Arguments

| | |
|----|---------------------------------|
| id | Character: namespace identifier |
|----|---------------------------------|

Value

Character with HTML

| | |
|------------|---|
| tabDataset | <i>Creates a tabPanel template for a datatable with a title and description</i> |
|------------|---|

Description

Creates a tabPanel template for a datatable with a title and description

Usage

```
tabDataset(ns, title, tableId, columns, visCols, data, description = NULL)
```

Arguments

| | |
|-------------|--|
| ns | Namespace function |
| title | Character: tab title |
| tableId | Character: id of the datatable |
| columns | Character: column names of the datatable |
| visCols | Boolean: visible columns |
| data | Data frame: dataset of interest |
| description | Character: description of the table (optional) |

Value

The HTML code for a tabPanel template

| | |
|----------------|---------------------------------|
| templateServer | <i>Server logic of template</i> |
|----------------|---------------------------------|

Description

Server logic of template

Usage

```
templateServer(input, output, session)
```

Arguments

| | |
|---------|---------------|
| input | Shiny input |
| output | Shiny output |
| session | Shiny session |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|-----------------------------------|
| templateUI | <i>User interface of template</i> |
|------------|-----------------------------------|

Description

User interface of template

Usage

```
templateUI(id)
```

Arguments

| | |
|----|---------------------------------|
| id | Character: namespace identifier |
|----|---------------------------------|

Value

HTML elements for the interface of the template

| | |
|--------------|---|
| testSurvival | <i>Test the survival difference between survival groups</i> |
|--------------|---|

Description

Test the survival difference between survival groups

Usage

```
testSurvival(survTerms, ...)
```

Arguments

| | |
|-----------|--|
| survTerms | survTerms object: processed survival terms |
| ... | Extra arguments passed to survdiff |

Value

p-value of the survival difference or NA

Note

Instead of raising errors, an NA is returned

Examples

```
require("survival")
data <- aml
timeStart <- "event"
event <- "event"
followup <- "time"
data$event <- NA
data$event[aml$status == 1] <- aml$time[aml$status == 1]
censoring <- "right"
formulaStr <- "x"
survTerms <- processSurvTerms(data, censoring=censoring, event=event,
                              timeStart=timeStart, followup=followup,
                              formulaStr=formulaStr)
testSurvival(survTerms)
```

| | |
|--------------------|--|
| testSurvivalCutoff | <i>Test the survival difference between two survival groups given a cutoff</i> |
|--------------------|--|

Description

Test the survival difference between two survival groups given a cutoff

Usage

```
testSurvivalCutoff(cutoff, data, filter = TRUE, clinical, ...,
                  session = NULL)
```

Arguments

| | |
|----------|---|
| cutoff | Numeric: Cut-off of interest |
| data | Numeric: elements of interest to test against the cut-off |
| filter | Boolean or numeric: elements to use (all by default) |
| clinical | Data frame: clinical data |
| ... | Arguments to pass to processSurvTerms |
| session | Shiny session |

Value

p-value of the survival difference

| | |
|-----------------|---|
| textSuggestions | <i>Create script for autocompletion of text input</i> |
|-----------------|---|

Description

Uses the JavaScript library jquery.textcomplete

Usage

```
textSuggestions(id, words, novalue = "No matching value", char = " ")
```

Arguments

| | |
|---------|---|
| id | Character: input ID |
| words | Character: words to suggest |
| novalue | Character: string when there's no matching values |
| char | Character to succeed accepted word |

Value

HTML string with the JavaScript script prepared to run

Examples

```
words <- c("tumor_stage", "age", "gender")
psychomics:::textSuggestions("textareaid", words)
```

| | |
|----------------|---|
| timePerPatient | <i>Get all columns matching a given string and return a single vector with the max time for each patient if available</i> |
|----------------|---|

Description

Get all columns matching a given string and return a single vector with the max time for each patient if available

Usage

```
timePerPatient(col, clinical)
```

Arguments

| | |
|----------|-------------------------------|
| col | Character: column of interest |
| clinical | Data.frame: clinical data |

Value

Numeric vector with days recorded for columns of interest

| | |
|---------------|-------------------------------------|
| transformData | <i>Transform data in data frame</i> |
|---------------|-------------------------------------|

Description

Transform data in data frame

Usage

```
transformData(input, df, x, y)
```

Arguments

| | |
|-------|------------------------|
| input | Shiny input |
| df | Data frame |
| x | Character: column name |
| y | Character: column name |

Value

Data frame with transformed data in new columns and respective name of created columns

| | |
|------------------|--|
| transformOptions | <i>Show variable transformation(s)</i> |
|------------------|--|

Description

Show variable transformation(s)

Usage

```
transformOptions(label, type = NULL)
```

Arguments

| | |
|-------|---|
| label | Character: label to display |
| type | Character: show the variable transformation for the chosen type; NULL (by default) to show all variable transformations |

Value

Character labelling variable transformation(s)

| | |
|-----------------|---|
| transformValues | <i>Transform values as per a given type of transformation</i> |
|-----------------|---|

Description

Transform values as per a given type of transformation

Usage

```
transformValues(val, type, avoidZero = TRUE)
```

Arguments

| | |
|-----------|--|
| val | Integer: values to transform |
| type | Character: type of transformation |
| avoidZero | Boolean: add the smallest non-zero number available to zero values; avoids returning infinity values during Log transformation (which are not plotted); useful for preserving p-values of 0, for instance; TRUE by default |

Value

Integer containing transformed values

| | |
|----------------|-------------------------------------|
| trimWhitespace | <i>Trims whitespace from a word</i> |
|----------------|-------------------------------------|

Description

Trims whitespace from a word

Usage

```
trimWhitespace(word)
```

Arguments

| | |
|------|-------------------|
| word | Character to trim |
|------|-------------------|

Value

Character without whitespace

Examples

```
psychomics::trimWhitespace("  hey  there  ")
psychomics::trimWhitespace(c("pineapple  ", "one two three",
                             " sunken  ship  "))
```

| | |
|-------|--|
| ttest | <i>Perform unpaired t-test analysis and return interface to show the results</i> |
|-------|--|

Description

Perform unpaired t-test analysis and return interface to show the results

Usage

```
ttest(psi, groups, stat = NULL)
```

Arguments

| | |
|--------|--|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |
| stat | Data frame or matrix: values of the analyses to be performed (if NULL, the analyses will be performed) |

Value

HTML elements

| | |
|----------|--|
| uniqueBy | <i>Check unique rows of a data frame based on a set of its columns</i> |
|----------|--|

Description

Check unique rows of a data frame based on a set of its columns

Usage

```
uniqueBy(data, ...)
```

Arguments

| | |
|------|----------------------|
| data | Data frame or matrix |
| ... | Name of columns |

Value

Data frame with unique values based on set of columns

| | |
|----------------------|--|
| updateClinicalParams | <i>Update available clinical attributes when the clinical data changes</i> |
|----------------------|--|

Description

Update available clinical attributes when the clinical data changes

Usage

```
updateClinicalParams(session, clinical)
```

Arguments

| | |
|----------|---------------------------|
| session | Shiny session |
| clinical | Data frame: clinical data |

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|----------------|---------------------------------|
| updateProgress | <i>Update a progress object</i> |
|----------------|---------------------------------|

Description

Update a progress object

Usage

```
updateProgress(message = "Hang in there", value = NULL, max = NULL,
  detail = NULL, divisions = NULL, global = sharedData, console = TRUE)
```

Arguments

| | |
|-----------|--|
| message | Character: progress message |
| value | Integer: current progress value |
| max | Integer: maximum progress value |
| detail | Character: detailed message |
| divisions | Integer: number of divisions in the progress bar |
| global | Shiny's global variable |
| console | Boolean: print message to console? (TRUE by default) |

Details

If divisions isn't NULL, a progress bar is started with the given divisions. If value is NULL, the progress bar will be incremented by one; otherwise, the progress bar will be incremented by the integer given in value.

Value

NULL (this function is used to modify the Shiny session's state)

| | |
|------------|---|
| vennEvents | <i>Compare the number of events from the different programs in a Venn diagram</i> |
|------------|---|

Description

Compare the number of events from the different programs in a Venn diagram

Usage

```
vennEvents(join, eventType)
```

Arguments

| | |
|-----------|-----------------------------|
| join | List of lists of data frame |
| eventType | Character: type of event |

Value

Venn diagrams for a given event type

| | |
|--------|---|
| wilcox | <i>Perform Wilcoxon analysis and return interface to show the results</i> |
|--------|---|

Description

Perform Wilcoxon analysis and return interface to show the results

Usage

```
wilcox(psi, groups, stat = NULL)
```

Arguments

| | |
|--------|--|
| psi | Numeric: quantification of one alternative splicing event |
| groups | Character: group of each PSI index |
| stat | Data frame or matrix: values of the analyses to be performed (if NULL, the analyses will be performed) |

Value

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