

# Package ‘omXplore’

January 30, 2026

**Type** Package

**Title** Vizualization tools for 'omics' datasets with R

**Version** 1.4.2

**Description** This package contains a collection of functions (written as shiny modules) for the visualisation and the statistical analysis of omics data. These plots can be displayed individually or embedded in a global Shiny module.  
Additionally, it is possible to integrate third party modules to the main interface of the package omXplore.

**License** Artistic-2.0

**Depends** R (>= 4.5.0), methods

**Imports** DT, shiny, MSnbase, PSMATCH, SummarizedExperiment, MultiAssayExperiment, shinyBS, shinyjs, shinyjqui, RColorBrewer, gplots, highcharter, visNetwork, tibble, grDevices, stats, utils, htmlwidgets, vioplot, graphics, FactoMineR, dendextend, dplyr, factoextra, tidyr, nipals

**Suggests** knitr, rmarkdown, BiocStyle, testthat, Matrix, graph

**biocViews** Software, ShinyApps, MassSpectrometry, DataRepresentation, GUI, QualityControl

**NeedsCompilation** no

**Collate** 'mod\_explore\_graphs.R' 'Infos\_adjacencyMatrix.R' 'Prostar\_1x.R' 'convert\_to\_mae.R' 'doc-data.R' 'external\_apps\_examples.R' 'get\_pep\_prot\_CC.R' 'mae\_accessors.R' 'metacell\_utils.R' 'mod\_colorLegend.R' 'modules.R' 'omXplore\_cc.R' 'omXplore\_corrmatrix.R' 'omXplore\_density.R' 'omXplore\_format\_DT.R' 'omXplore\_heatmap.R' 'omXplore\_intensity.R' 'omXplore\_PCA\_nipals.R' 'omXplore\_pca.R' 'omXplore\_plots\_tracking.R' 'omXplore\_tabExplorer.R' 'omXplore\_variance.R' 'omXplore\_view\_dataset.R' 'palette.R' 'plot\_boxplot.R' 'plot\_heatmap.R' 'plot\_pca.R' 'plot\_violin.R' 'utils.R' 'zzz.R'

**RoxygenNote** 7.3.3

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<https://edyp-lab.github.io/omXplore/>

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|                  |                            |
|------------------|----------------------------|
| <i>accessors</i> | <i>Accessors functions</i> |
|------------------|----------------------------|

---

## Description

Functions used to access the additional plots in the instances of the class `MultiAssayExperiment`.

## Usage

```
get_adjacencyMatrix(object, ...)

## S4 method for signature 'SummarizedExperiment'
get_adjacencyMatrix(object)

get_design(object, ...)

## S4 method for signature 'MultiAssayExperiment'
get_design(object)

get_group(object, ...)

## S4 method for signature 'MultiAssayExperiment'
get_group(object)

get_metacell(object, ...)

## S4 method for signature 'SummarizedExperiment'
get_metacell(object, slot.name = c("metacell", "qMetacell"))

get_cc(object, ...)

## S4 method for signature 'SummarizedExperiment'
get_cc(object)

get_parentProtId(object, ...)

## S4 method for signature 'SummarizedExperiment'
get_parentProtId(object)

get_colID(object, ...)

## S4 method for signature 'SummarizedExperiment'
get_colID(object)

get_type(object, ...)

## S4 method for signature 'SummarizedExperiment'
```

```

get_type(object)

get_pkg_version(object, ...)

## S4 method for signature 'SummarizedExperiment'
get_pkg_version(object)

```

## Arguments

|                        |                                                                                                          |
|------------------------|----------------------------------------------------------------------------------------------------------|
| <code>object</code>    | An instance of class <code>SummarizedExperiment</code> .                                                 |
| <code>...</code>       | Additional parameters                                                                                    |
| <code>slot.name</code> | The name of the slot dedicated to cell metadata to search. Default values are 'metacell' and 'qMetacell' |

## Value

See individual method description for the return value.

If exists, the slot value requested.

A `DataFrame` containing the adjacency matrix of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

A `data.frame` containing the metadata of the dataset

## Examples

```

## -----
## Accessing slots from a MSnSet dataset
## -----
data(sub_R25)
se1 <- sub_R25[[1]]
parentProtId <- get_parentProtId(se1)
colID <- get_colID(se1)
type <- get_type(se1)
metacell <- get_metacell(se1)
conds <- get_group(sub_R25)

```

---

|                  |                                        |
|------------------|----------------------------------------|
| BuildColorStyles | <i>Build color style for DT tables</i> |
|------------------|----------------------------------------|

---

**Description**

This function builds a list which is used for styling DT tables with the function `DT::styleEqual()`

**Usage**

```
BuildColorStyles(type)
```

**Arguments**

|      |                                                               |
|------|---------------------------------------------------------------|
| type | The type of dataset. Available values are protein and peptide |
|------|---------------------------------------------------------------|

**Value**

A list

**Examples**

```
NULL
```

---

|                      |                                                      |
|----------------------|------------------------------------------------------|
| Build_enriched_qdata | <i>Builds enriched assay with cell metadata info</i> |
|----------------------|------------------------------------------------------|

---

**Description**

If the cell metadata exists in the object of class `SummarizedExperiment`, then these information are added to the quantitative data so as to use styles with the functions of the package DT.

**Usage**

```
Build_enriched_qdata(obj)
```

**Arguments**

|     |                                                            |
|-----|------------------------------------------------------------|
| obj | An instance of the class <code>SummarizedExperiment</code> |
|-----|------------------------------------------------------------|

**Value**

A data.frame with new columns corresponding to the cell metadata (if exists)

**Examples**

```
NULL
```

---

`color-legend`*Color legend for DaparToolshed*

---

**Description**

Shows a legend based on the tags in the package 'DaparToolshed'

**Usage**

```
custom_metacell_colors()

colorLegend_ui(id)

colorLegend_server(
  id,
  presentTags = reactive({
    NULL
  }),
  hide.white = TRUE
)

colorLegend(dataIn = SummarizedExperiment::SummarizedExperiment())
```

**Arguments**

|                          |                                                                        |
|--------------------------|------------------------------------------------------------------------|
| <code>id</code>          | A character(1) which is the id of the shiny module.                    |
| <code>presentTags</code> | A vector of character() which correspond to the tags.                  |
| <code>hide.white</code>  | A boolean() to indicate whether the white cells must be hidden or not. |
| <code>dataIn</code>      | An instance of the class SummarizedExperiment.                         |

**Value**

A vector  
NA  
NA  
A shiny app

**Examples**

```
if (interactive()) {
  data(vdata)
  shiny::runApp(colorLegend(vdata[[1]]))
}
```

---

converters

*Convert to enriched MultiAssayExperiment*

---

### Description

The resulting object is an instance of the MultiAssayExperiment class.

### Usage

```
convert_to_mae(obj)

MSnSet_to_mae(obj)

matrix_to_mae(obj)

df_to_mae(obj)

Compute_CC(obj)

QFeatures_to_mae(obj)

SE_to_mae(obj)

MAE_to_mae(obj)

Check_se_Consistency(obj)

list_to_se(ll)

Check_List_consistency(ll)

listOfLists_to_mae(obj, colData = NULL)

listOfSE_to_mae(obj)

Check_MSnSet_Consistency(obj)

matrix_to_se(obj)

df_to_se(obj)

MSnSet_to_se(obj)

Build_X_CC(se)

listOfMSnSet_to_mae(obj)

listOfmatrix_to_mae(obj)

listOfdf_to_mae(obj)
```

**Arguments**

|                      |                                                                        |
|----------------------|------------------------------------------------------------------------|
| <code>obj</code>     | An object compliant with the formats accepted by <code>omXplore</code> |
| <code>ll</code>      | A list                                                                 |
| <code>colData</code> | A <code>data.frame()</code>                                            |
| <code>se</code>      | AN instance of the class <code>SummarizedExperiment</code>             |

**Value**

An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An instance of `SimpleList`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 A `boolean(1)`  
 An enriched instance of the class `SummarizedExperiment`  
 A `boolean(1)`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 A `boolean(1)`  
 An enriched instance of the class `SummarizedExperiment`  
 An enriched instance of the class `SummarizedExperiment`  
 An enriched instance of the class `SummarizedExperiment`  
 An enriched instance of the class `SummarizedExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`  
 An enriched instance of the class `MultiAssayExperiment`

**Examples**

```

#-----
# Conversion of a MultiAssayExperiment instance
#-----
data(miniACC, package = "MultiAssayExperiment")
convert_to_mae(miniACC)

```



---

|            |                                                                                    |
|------------|------------------------------------------------------------------------------------|
| corrmatrix | <i>Displays a correlation matrix of the quantitative data of a numeric matrix.</i> |
|------------|------------------------------------------------------------------------------------|

---

### Description

Displays a correlation matrix of the quantitative data of a numeric matrix.

### Usage

```
omXplore_corrmatrix_ui(id)

omXplore_corrmatrix_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
    1
  })
)

corrMatrix(data, rate = 0.5, showValues = FALSE)

omXplore_corrmatrix(dataIn, i)
```

### Arguments

|            |                                                                              |
|------------|------------------------------------------------------------------------------|
| id         | A character(1) which is the id of the shiny module.                          |
| dataIn     | An instance of the class SummarizedExperiment                                |
| i          | An integer which is the index of the assay in the param dataIn               |
| data       | An object of class 'matrix'                                                  |
| rate       | The rate parameter to control the exponential law for the gradient of colors |
| showValues | A boolean which indicates whether to show values in the correlation plot.    |

### Value

NA  
 NA  
 A plot  
 A shiny app

### Examples

```
if (interactive()) {
  data(vdata)
  omXplore_corrmatrix(vdata, 1)
}
```

---

`customChart`*Customised resetZoom Button of highcharts plots*

---

**Description**

Customised resetZoom Button of highcharts plots

**Usage**

```
customChart(  
  hc,  
  chartType = "scatter",  
  zoomType = "None",  
  width = 0,  
  height = 0  
)
```

**Arguments**

|                        |                                                      |
|------------------------|------------------------------------------------------|
| <code>hc</code>        | A highcharter object                                 |
| <code>chartType</code> | The type of the plot                                 |
| <code>zoomType</code>  | The type of the zoom (one of "x", "y", "xy", "None") |
| <code>width</code>     | The width of the plot                                |
| <code>height</code>    | The height of the plot                               |

**Value**

A highchart plot

**Author(s)**

Samuel Wieczorek

**Examples**

```
if (interactive()) {  
  library(highcharter)  
  hc <- highchart()  
  hc_chart(hc, type = "line")  
  hc_add_series(hc, data = c(29, 71, 40))  
  customChart(hc)  
}  
  
NULL
```

---

|              |                                                                                    |
|--------------|------------------------------------------------------------------------------------|
| density-plot | <i>Displays a correlation matrix of the quantitative data of a numeric matrix.</i> |
|--------------|------------------------------------------------------------------------------------|

---

## Description

Displays a correlation matrix of the quantitative data of a numeric matrix.

## Usage

```
omXplore_density_ui(id)

omXplore_density_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
    1
  }),
  pal.name = reactive({
    NULL
  })
)

densityPlot(data, conds = NULL, pal.name = NULL)

omXplore_density(dataIn, i)
```

## Arguments

|                       |                                                                                                                                                                       |
|-----------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>id</code>       | A character(1) which is the id of the shiny module.                                                                                                                   |
| <code>dataIn</code>   | An instance of the class SummarizedExperiment                                                                                                                         |
| <code>i</code>        | An integer which is the index of the assay in the param obj                                                                                                           |
| <code>pal.name</code> | A character(1) which is the name of the palette from the package <a href="#">RColorBrewer::RColorBrewer</a> from which the colors are taken. Default value is 'Set1'. |
| <code>data</code>     | A data.frame() of quantitative data                                                                                                                                   |
| <code>conds</code>    | A vector indicating the name of each sample.                                                                                                                          |

## Value

NA  
 NA  
 A plot  
 A shiny app

**Examples**

```

if (interactive()) {
  data(vdata)
  shiny::runApp(omXplore_density(vdata, 1))
}

if (interactive()) {
  data(vdata)
  qdata <- SummarizedExperiment::assay(vdata[[1]])
  conds <- get_group(vdata)
  densityPlot(qdata, conds)
}

```

ds-cc

*plots\_cc\_ui and plots\_cc\_server***Description**

A shiny Module.

**Usage**

```

omXplore_cc_ui(id)

omXplore_cc_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
    NULL
  })
)

omXplore_cc(dataIn, i)

```

**Arguments**

|                     |                                                             |
|---------------------|-------------------------------------------------------------|
| <code>id</code>     | A character(1) which is the id of the shiny module.         |
| <code>dataIn</code> | An instance of SummarizedExperiment class                   |
| <code>i</code>      | An integer which is the index of the assay in the param obj |

**Value**

A shiny module  
 A shiny plot  
 A shiny module  
 A shiny app  
 A shiny app

## Examples

```
if (interactive()) {
  data(vdata)
  shiny::runApp(omXplore_cc(vdata, 1))
}
```

---

|        |               |
|--------|---------------|
| ds-pca | <i>my_PCA</i> |
|--------|---------------|

---

## Description

Process a PCA, using nipals or FactoMineR, on a quantitative dataset.

This method plots a bar plot which represents the distribution of the number of missing values (NA) per lines (ie proteins).

- `wrapper_pca()`
- `plotPCA_Eigen_hc()`: plots the eigen values of PCA with the highcharts library
- `plotPCA_Eigen()`: plots the eigen values of PCA
- `plotPCA_Var()`
- `plotPCA_Ind()`

## Usage

```
my_PCA(
  X,
  scale.unit = TRUE,
  ncp = min(12, nrow(X) - 1, ncol(X)),
  ind.sup = NULL,
  quanti.sup = NULL,
  quali.sup = NULL,
  row.w = NULL,
  col.w = NULL,
  graph = FALSE,
  axes = c(1, 2),
  approach = "FM",
  gramschmidt = TRUE
)
```

```
omXplore_pca_ui(id)
```

```
omXplore_pca_server(id, dataIn, i)
```

```
omXplore_pca(dataIn, i)
```

```
wrapper_pca(
  qdata,
  group,
  var.scaling = TRUE,
  ncp = NULL,
```

```

    approach = "FM",
    gramschmidt = TRUE
  )

plotPCA_Eigen(res.pca)

plotPCA_Var(res.pca, chosen.axes = c(1, 2))

plotPCA_Ind(res.pca, chosen.axes = c(1, 2))

plotPCA_Eigen_hc(res.pca)

```

### Arguments

|                          |                                                                                                               |
|--------------------------|---------------------------------------------------------------------------------------------------------------|
| <code>X</code>           | a <code>data.frame()</code> of quantitative data                                                              |
| <code>scale.unit</code>  | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>ncp</code>         | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>ind.sup</code>     | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>quanti.sup</code>  | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>quali.sup</code>   | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>row.w</code>       | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>col.w</code>       | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>graph</code>       | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>axes</code>        | See <code>FactoMineR::PCA()</code>                                                                            |
| <code>approach</code>    | a string corresponding to the package to use for PCA (if no NA, default is "FM" for <code>FactoMineR</code> ) |
| <code>gramschmidt</code> | A boolean indicating whether to use Gram-Schmidt orthogonalization or not.                                    |
| <code>id</code>          | A character(1) which is the id of the shiny module.                                                           |
| <code>dataIn</code>      | An instance of the class <code>MultiAssayExperiment</code> .                                                  |
| <code>i</code>           | An integer which is the index of the assay in the param obj                                                   |
| <code>qdata</code>       | A <code>data.frame()</code> of quantitative data                                                              |
| <code>group</code>       | A vector with the name of samples                                                                             |
| <code>var.scaling</code> | A boolean indicating whether to scale the data or not                                                         |
| <code>res.pca</code>     | The result of the function <code>FactoMineR::PCA()</code>                                                     |
| <code>chosen.axes</code> | See the parameter 'axes' of the function <code>factoextra::fviz_pca_var()</code>                              |

### Value

```

res.pca a "PCA"  "list" object
NA
NA
A shiny app
The result of the function FactoMineR::PCA()
A plot
A plot
A plot
A plot

```

**Author(s)**

Samuel Wieczorek, Enora Fremy

**Examples**

```
if (interactive()) {
  data(vdata)
  obj <- vdata[[1]]
  res.pca <- my_PCA(SummarizedExperiment::assay(obj), approach = "FM")
  plotPCA_Eigen(res.pca)
  plotPCA_Var(res.pca)
  plotPCA_Eigen_hc(res.pca)
  plotPCA_Ind(res.pca)
}

if (interactive()) {
  data(vdata)
  library(shiny)
  library(dplyr)
  library(highcharter)
  # Replace missing values for the example
  sel <- is.na(SummarizedExperiment::assay(vdata, 1))
  SummarizedExperiment::assay(vdata[[1]])[sel] <- 0
  SummarizedExperiment::assay(vdata[[1]])[1, 1] <- NA
  omXplore_pca(vdata, 1)
}

if (interactive()) {
  data(vdata)
  obj <- vdata[[1]]
  res.pca <- wrapper_pca(qdata = SummarizedExperiment::assay(obj), group = get_group(obj))
  plotPCA_Eigen(res.pca)
  plotPCA_Var(res.pca)
  plotPCA_Eigen_hc(res.pca)
  plotPCA_Ind(res.pca)
}
```

---

ds-view

*Bar plot of missing values per lines using highcharter.*


---

**Description**

This method plots a bar plot which represents the distribution of the number of missing values (NA) per lines (i.e. proteins).

**Usage**

```
view_dataset_ui(id)

view_dataset_server(
  id,
  dataIn = reactive({
```

```

    NULL
  }},
  addons = list(),
  verbose = FALSE
)

view_dataset(dataIn = NULL, addons = NULL)

```

### Arguments

|         |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 |
|---------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| id      | A character(1) for the 'id' of the shiny module. It must be the same as for the '*_ui' function.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| dataIn  | An instance of the class MultiAssayExperiment.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| addons  | A list to configure the other shiny apps to integrate. Each item correspond to one package: <ul style="list-style-type: none"> <li>• the name of the slot is the name of the package</li> <li>• the content of the slot is a vector composed of the generic name of the shiny app. Each of the apps listed here must be an exported app of the package. For example, given the value <code>addons = list(testPkg = c('foo', 'foo2'))</code>. That means that the package called "testPkg" must provide the four functions: <code>foo1_ui()</code>, <code>foo1_server()</code> and <code>foo2_ui()</code>, <code>foo2_server()</code></li> </ul> |
| verbose | A boolean for verbose mode. Default is FALSE.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   |

### Details

- distribution of the missing values per line,
- a bar plot which represents the distribution of the number of missing values (NA) per lines (i.e. proteins) and per conditions,
- Histogram of missing values.
- Variance : Builds a densityplot of the CV of entities in numeric matrix. The CV is calculated for each condition present in the dataset (see the slot 'Condition' in the `colData()` DataFrame)
- Heatmap:

The function [heatmapD\(\)](#)

The function `[]` is inspired from the function 'heatmap.2' that displays a numeric matrix. For more information, please refer to the help of the heatmap.2 function.

### Value

NA

NA

NA

A shiny application which wraps the functions `view_dataset_ui()` and the `view_dataset_server()`

### Missing values

#' - distribution of the missing values per line,

- a bar plot which represents the distribution of the number of missing values (NA) per lines (ie proteins) and per conditions,
- Histogram of missing values.



**Author(s)**

Samuel Wieczorek, Enora Fremy

**Examples**

```
if (interactive()) {
  library(shiny)
  library(omXplore)
  data(vdata)
  addons <- list(omXplore = c("extFoo1", "extFoo2"))
  runApp(omXplore::view_dataset(vdata, addons))

  omXplore::view_dataset(vdata)
}

if (interactive()) {
  data(vdata)
  view_dataset(vdata)
}
```

---

external\_app

*External module example*


---

**Description**

Example for an external shiny module, well structured to be run within a workflow for MagellanNTK

**Usage**

```
extFoo1_ui(id)

extFoo1_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
    NULL
  })
)

extFoo1(dataIn, i)

extFoo2_ui(id)

extFoo2_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
```

```

      NULL
    })
  )

  extFoo2(dataIn, i)

```

### Arguments

|                     |                                                             |
|---------------------|-------------------------------------------------------------|
| <code>id</code>     | A character(1) which is the id of the shiny module.         |
| <code>dataIn</code> | An object of instance MultiAssayExperiment                  |
| <code>i</code>      | An integer which is the index of the assay in the param obj |

### Value

```

NA
NA
NA
A shiny app
NA
NA
A shiny app

```

### Examples

```

if (interactive()) {
  data(vdata)
  app1 <- extFoo1(vdata, 1)
  app2 <- extFoo2(vdata, 1)
  shiny::runApp(app1)
  shiny::runApp(app2)
}

```

---

FormatDataForDT

Constructs a dataset suitable to use with the module format\_DT.

---

### Description

This function builds the skeleton of a dataset which can be used by the module formatDT. It creates additional columns to be used to style the table. to colors cells.

### Usage

```
FormatDataForDT(se, digits = 2)
```

### Arguments

|                     |                                                                                                              |
|---------------------|--------------------------------------------------------------------------------------------------------------|
| <code>se</code>     | An instance of the class SummarizedExperiment                                                                |
| <code>digits</code> | An 'integer(1)' to specify the number of digits to display in the tables for numerical values. Default is 2. |

**Value**

A data.frame

---

format\_DT

*formatDT\_ui and formatDT\_server*


---

**Description**

A shiny Module.

See DT package homepage for more details about styling tables. If no style is precised, this module show the raw data. If any style is given, then the dataset must be well configured (I.e. it must contain the correct columns )

**Usage**

```
formatDT_ui(id)

formatDT_server(
  id,
  data = reactive({
    NULL
  }),
  data_nostyle = reactive({
    NULL
  }),
  withDLBtns = FALSE,
  showRownames = FALSE,
  dt_style = reactive({
    NULL
  }),
  filename = "Prostar_export",
  selection = "single"
)

formatDT(data)
```

**Arguments**

|              |                                                                                                                                |
|--------------|--------------------------------------------------------------------------------------------------------------------------------|
| id           | shiny id                                                                                                                       |
| data         | A data.frame                                                                                                                   |
| data_nostyle | A data.frame() to be bind to the main data with no custom style                                                                |
| withDLBtns   | A boolean to indicate whether to display download buttons or not.                                                              |
| showRownames | A boolean to indicate whether to show rownames.                                                                                |
| dt_style     | A list composed of: <ul style="list-style-type: none"> <li>• data : a data.frame</li> <li>• colors : a named vector</li> </ul> |
| filename     | A character(1) which is the default filename for download.                                                                     |
| selection    | A character(1) which indicates the type of selection. Default is 'single'.                                                     |

**Value**

NA

NA

NA

NA

**Examples**

```
if (interactive()) {  
  data(vdata)  
  formatDT(vdata)  
}
```

---

|               |                        |
|---------------|------------------------|
| GetPkgVersion | <i>Package version</i> |
|---------------|------------------------|

---

**Description**

Gets the version number of a package

**Usage**

```
GetPkgVersion(pkg)
```

**Arguments**

|     |                         |
|-----|-------------------------|
| pkg | The name of the package |
|-----|-------------------------|

**Value**

A character(1) with the name of the package and its version number.

**Examples**

```
GetPkgVersion("omXplore")
```

---

`globals`*Global variables*

---

**Description**

Defines the global variables for the package omXplore

**Usage**

```
globals()
```

**Value**

A list

**Examples**

```
globals()
```

---

`intensity-plots`*Displays different intensity plots.*

---

**Description**

Displays different intensity plots.

**Usage**

```
omXplore_intensity_ui(id)

omXplore_intensity_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
    1
  }),
  track.indices = reactive({
    NULL
  }),
  remoteReset = reactive({
    NULL
  }),
  is.enabled = reactive({
    TRUE
  })
)
```

```
omXplore_intensity(dataIn, i = NULL, withTracking = FALSE)

boxPlot(dataIn, conds, legend = NULL, pal = NULL, subset = NULL)

violinPlot(data, conds, subset = NULL, pal.name = "Set1")
```

### Arguments

|                            |                                                                                                                                                                       |
|----------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>id</code>            | A character(1) which is the id of the shiny module.                                                                                                                   |
| <code>dataIn</code>        | xxxx                                                                                                                                                                  |
| <code>i</code>             | An integer which is the index of the assay in the param obj                                                                                                           |
| <code>track.indices</code> | A vector of integers which are the indices of lines to track.                                                                                                         |
| <code>remoteReset</code>   | An integer to activate the reset functions                                                                                                                            |
| <code>is.enabled</code>    | A boolean that indicates whether the widgets should be enabled or disabled. Default is TRUE                                                                           |
| <code>withTracking</code>  | A boolean(1) indicating whether the tracking option is activated or not.                                                                                              |
| <code>conds</code>         | A vector indicating the name of each sample.                                                                                                                          |
| <code>legend</code>        | A vector of the conditions (one condition per sample).                                                                                                                |
| <code>pal</code>           | A basis palette for the boxes which length must be equal to the number of unique conditions in the dataset.                                                           |
| <code>subset</code>        | A integer() vector of index indicating the indices of rows in the dataset to highlight                                                                                |
| <code>data</code>          | xxxx                                                                                                                                                                  |
| <code>pal.name</code>      | A character(1) which is the name of the palette from the package <a href="#">RColorBrewer::RColorBrewer</a> from which the colors are taken. Default value is 'Set1'. |

### Value

NA  
 NA  
 A shiny app  
 A boxplot  
 A violin plot

### Author(s)

Samuel Wieczorek, Anais Courtier, Enora Fremy

### Examples

```
if (interactive()) {
  data(vdata)
  shiny::runApp(omXplore_intensity(vdata, 1))

  data(sub_R25)
  conds <- legend <- SummarizedExperiment::colData(sub_R25)$group
  pal <- ExtendPalette(length(unique(conds)))
  boxPlot(sub_R25[[1]], conds, legend, pal, seq_len(10))
}
```

```

    shiny::runApp(omXplore_intensity(sub_R25, 1, withTracking = TRUE))
  }

```

is.listOf

*Checks the class of a list's slots***Description**

Checks if all slots of the given list are of the same class.

**Usage**

```
is.listOf(object, obj.class = NULL)
```

**Arguments**

|           |                                                       |
|-----------|-------------------------------------------------------|
| object    | A list                                                |
| obj.class | The name of the class to search in items of the list. |

**Value**

A character(1) with the name of the package or NULL

**Examples**

```

ll <- as.list(LETTERS[1:3])
is.listOf(ll, "data.frame")
is.listOf(ll, "character")

```

omXplore-modules

*Shiny modules used by omXplore***Description**

These functions are relative to external modules that can be added into omXplore UI:

- `listShinyApps()`: Show the shiny modules recognized by omXplore and ready to be integrated in the UI of the function `view_dataset()`
- `listPlotModules()`: Show the shiny modules function names (only prefixes) recognized by omXplore and ready to use in the UI.
- `addModules()`: Add external shiny module(s) to the R global environment in such a way (specific prefix renaming of the functions) that it can be discovered by the function `view_dataset()` of the package omXplore during its launch.

**Usage**

```
addModules(addons = list())

listShinyApps(location = "both")

listPlotModules(location = "both")
```

**Arguments**

|          |                                                                                                                                                                                                                                                                                 |
|----------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| addons   | A list in which each item: <ul style="list-style-type: none"> <li>• is named by the name of a package containing the modules to add,</li> <li>• contains the name of the shiny modules to integrate (without <code>'_ui'</code> nor <code>'_server'</code> suffixes)</li> </ul> |
| location | A character(0) to indicate which modules to list. Available values are: <code>'builtin'</code> , <code>'external'</code> and <code>'both'</code> (default).                                                                                                                     |

**Value**

NA  
A vector  
A vector

**Examples**

```
listShinyApps()
listPlotModules()

#####
# Integration of a module in the package 'mypackage'
#####
addons <- list(omXplore = c("extFoo1", "extFoo2"))
addModules(addons)
```

---

|                  |                                                                                    |
|------------------|------------------------------------------------------------------------------------|
| omXplore_heatmap | <i>Displays a correlation matrix of the quantitative data of a numeric matrix.</i> |
|------------------|------------------------------------------------------------------------------------|

---

**Description**

This function is a wrapper to `heatmap.2()` that displays assay data in an instance of `SummarizedExperiment`. For more details, see `heatmap.2()`.

**Usage**

```
omXplore_heatmap_ui(id)

omXplore_heatmap_server(
  id,
  dataIn = reactive({
```



```

        NULL
    }},
    i = reactive({
        NULL
    })
)

omXplore_heatmap(dataIn, i)

heatmapD(
  qdata,
  conds,
  distance = "euclidean",
  cluster = "complete",
  dendro = FALSE
)

mv.heatmap(
  x,
  col = grDevices::heat.colors(100),
  srtCol = NULL,
  labCol = NULL,
  labRow = NULL,
  key = TRUE,
  key.title = NULL,
  main = NULL,
  ylab = NULL
)

heatmapForMissingValues(
  x,
  col = NULL,
  srtCol = NULL,
  labCol = NULL,
  labRow = NULL,
  key = TRUE,
  key.title = NULL,
  main = NULL,
  ylab = NULL
)

```

### Arguments

|          |                                                                          |
|----------|--------------------------------------------------------------------------|
| id       | A character(1) which is the id of the shiny module.                      |
| dataIn   | An instance of a class MultiAssayExperiment.                             |
| i        | An integer which is the index of the assay in the param obj              |
| qdata    | A data.frame() of quantitative data.                                     |
| conds    | A vector indicating the name of each sample.                             |
| distance | The distance used by the clustering algorithm to compute the dendrogram. |
| cluster  | the clustering algorithm used to build the dendrogram.                   |
| dendro   | A boolean to indicate fi the dendrogram has to be displayed              |

|           |                                                                     |
|-----------|---------------------------------------------------------------------|
| x         | A matrix or array containing the quantitative data.                 |
| col       | Colors used for the image. Defaults to heat colors (heat.colors).   |
| srtCol    | Angle of column conds, in degrees from horizontal                   |
| labCol    | Character vectors with column conds to use.                         |
| labRow    | Character vectors with row conds to use.                            |
| key       | Logical indicating whether a color-key should be shown.             |
| key.title | Main title of the color key. If set to NA no title will be plotted. |
| main      | Main title; default to none.                                        |
| ylab      | y-axis title; default to none.                                      |

**Value**

NA  
 NA  
 A shiny app  
 A heatmap  
 A heatmap  
 A heatmap

**Author(s)**

Florence Combes, Samuel Wiczorek, Enora Fremy

**Examples**

```
if (interactive()) {
  data(vdata)
  omXplore_heatmap(vdata, 1)
}
```

---

omXplore\_tabExplorer    *Explore MultiAssayExperiment objects.*

---

**Description**

Explore MultiAssayExperiment objects.

**Usage**

```
omXplore_tabExplorer_ui(id)

omXplore_tabExplorer_server(
  id,
  dataIn = reactive({
    NULL
  }),
  i = reactive({
```

```

      NULL
    }},
    digits = reactive({
      3
    })
  )

  omXplore_tabExplorer(dataIn, i)

```

### Arguments

|                     |                                                             |
|---------------------|-------------------------------------------------------------|
| <code>id</code>     | A character(1) which is the id of the shiny module.         |
| <code>dataIn</code> | An instance of the class <code>MultiAssayExperiment</code>  |
| <code>i</code>      | An integer which is the index of the assay in the param obj |
| <code>digits</code> | An integer for the number of digits shown in the table      |

### Value

```

NA
NA
NA
A shiny app

```

### Examples

```

if (interactive()) {
  data(vdata)
  shiny::runApp(omXplore_tabExplorer(vdata, 1))
}

NULL

```

---

|         |                             |
|---------|-----------------------------|
| palette | <i>Palette for samples.</i> |
|---------|-----------------------------|

---

### Description

Builds a vector of `#conditions` colors for a set of samples. One color is given for a given condition. This function extends a base palette from the package [RColorBrewer::RColorBrewer](#) to 'n' colors. The colors in the returned palette are always in the same order

### Usage

```

SampleColors(conds, pal.name = "Set1")

ExtendPalette(n, pal.name = "Set1")

GetColorsForConditions(conds, pal = NULL)

```

**Arguments**

|          |                                                                                                                                                                       |
|----------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| conds    | A character() of conditions. The length of the vector is the number of samples in the dataset.                                                                        |
| pal.name | A character(1) which is the name of the palette from the package <a href="#">RColorBrewer::RColorBrewer</a> from which the colors are taken. Default value is 'Set1'. |
| n        | The number of desired colors in the palette                                                                                                                           |
| pal      | A vector of HEX color code that form the basis palette from which to build the complete color vector for the conditions.                                              |

**Value**

A vector  
A vector  
A vector

**Examples**

```
if (interactive()) {
  #-----
  # Builds a palette for a dataset with 3 conditions
  # of 3 samples each.
  #-----

  conds <- c(rep("A", 3), rep("B", 3), rep("C", 3))
  SampleColors(conds)
  SampleColors(conds, "Dark2")

  #-----
  # Extend the default palette to 12 colors
  #-----

  ExtendPalette(12)

  data(vdata)
  conds <- get_group(vdata)
  GetColorsForConditions(conds, ExtendPalette(2))
}
```

---

pep\_prot\_CC

---

Display a CC

---

**Description**

Display a CC  
Connected Components infos

## Usage

```
buildGraph(cc = NULL, meta = NULL)

display.CC.visNet(g = NULL, layout = "layout_with_fr")

plotCCJitter(df, clickFunction = NULL)

GetCCInfos(cc)
```

## Arguments

|               |                                                                                          |
|---------------|------------------------------------------------------------------------------------------|
| cc            | A list of connected component                                                            |
| meta          | A data.frame()                                                                           |
| g             | An instance of a graph                                                                   |
| layout        | A character(1) which is the layout used in visNetwork. Default value is 'layout_with_fr' |
| df            | A data.frame()                                                                           |
| clickFunction | A JS function to determine the behaviour of a click                                      |

## Value

A list

A plot

A plot

A list of three items:

- One\_One: the number of cc composed of one protein and one peptide
- One\_Multi: the number of cc composed of one protein and several peptides
- Multi\_Multi: the number of cc composed of several proteins and several (shared) peptides.

## Author(s)

Thomas Burger, Samuel Wiczorek

## Examples

```
if (interactive()) {
  data(sub_R25)
  se <- sub_R25[[1]]
  g <- buildGraph(get_cc(se)[[1]])
  display.CC.visNet(g)
}

if (interactive()) {
  data(sub_R25)
  GetCCInfos(get_cc(sub_R25[[1]]))
}
```

---

|               |                       |
|---------------|-----------------------|
| pkgs.require2 | <i>Loads packages</i> |
|---------------|-----------------------|

---

**Description**

Checks if a package is available to load it

**Usage**

```
pkgs.require2(ll.deps)
```

**Arguments**

|         |                                                    |
|---------|----------------------------------------------------|
| ll.deps | A character() vector which contains packages names |
|---------|----------------------------------------------------|

**Value**

NA

**Author(s)**

Samuel Wieczorek

**Examples**

```
pkgs.require2("omXplore")
```

---

|               |                      |
|---------------|----------------------|
| plot-variance | <i>Variance plot</i> |
|---------------|----------------------|

---

**Description**

A shiny module which plots the variance of samples

**Usage**

```
omXplore_variance_ui(id)

omXplore_variance_server(id, dataIn, i, pal.name = NULL)

CVDist(dataIn, conds, pal.name = NULL)

omXplore_variance(dataIn, i)
```

**Arguments**

|          |                                                                                                                                                                       |
|----------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| id       | A character(1) which is the id of the shiny module.                                                                                                                   |
| dataIn   | An matrix                                                                                                                                                             |
| i        | An integer which is the index of the assay in the param obj                                                                                                           |
| pal.name | A character(1) which is the name of the palette from the package <a href="#">RColorBrewer::RColorBrewer</a> from which the colors are taken. Default value is 'Set1'. |
| conds    | A vector indicating the name of each sample.                                                                                                                          |

**Value**

NA  
 NA  
 A plot  
 A shiny app

**Examples**

```
if (interactive()) {
  data(vdata)
  shiny::runApp(omXplore_variance(vdata, 1))
}
```

---

|                |                                                    |
|----------------|----------------------------------------------------|
| plots_tracking | <i>plots_tracking_ui and plots_tracking_server</i> |
|----------------|----------------------------------------------------|

---

**Description**

This shiny module provides a tool to select

**Usage**

```
plots_tracking_ui(id)

plots_tracking_server(
  id,
  dataIn = reactive({
    NULL
  }),
  remoteReset = reactive({
    NULL
  }),
  is.enabled = reactive({
    TRUE
  })
)

plots_tracking(obj)
```

**Arguments**

|                          |                                                                                      |
|--------------------------|--------------------------------------------------------------------------------------|
| <code>id</code>          | shiny id                                                                             |
| <code>remoteReset</code> | A <code>boolean(1)</code> which indicates whether to show the 'Reset' button or not. |
| <code>is.enabled</code>  | xxx                                                                                  |
| <code>obj</code>         | An instance of the class <code>MultiAssayExperiment</code>                           |

**Value**

NA  
 A list (same structure as the parameter `params`)  
 A shiny app

**Examples**

```
if (interactive()) {
  data(vdata)
  shiny::runApp(plots_tracking(vdata[[1]]))
}

NULL
```

---

Prostar-1x-compatible *Convert datasets exported by the package Prostar*

---

**Description**

Convert datasets exported by the package Prostar

**Usage**

```
SE_Compatibility_with_Prostar_1x(obj, se)

MAE_Compatibility_with_Prostar_1x(obj, mae)
```

**Arguments**

|                  |                                                            |
|------------------|------------------------------------------------------------|
| <code>obj</code> | An instance of the class <code>MSnSet</code>               |
| <code>se</code>  | An instance of the class <code>SummarizedExperiment</code> |
| <code>mae</code> | An instance of the class <code>MultiAssayExperiment</code> |

**Value**

An enriched instance of the class `SummarizedExperiment`  
 An enriched instance of the class `MultiAssayExperiment`

**Examples**

```
data(sub_R25)
```



q\_metadata

*Quantitative metadata vocabulary for entities***Description**

This function gives the vocabulary used for the quantitative metadata of each entity in each condition.

**Usage**

```
metacell.def(level)
```

```
Parent(level, node = NULL)
```

```
Children(level, parent = NULL)
```

```
GetMetacellTags(metacells = NULL, level = NULL, onlyPresent = FALSE)
```

**Arguments**

|             |                                                     |
|-------------|-----------------------------------------------------|
| level       | A string corresponding to the type of object        |
| node        | The name of the node for which one wants its parent |
| parent      | The name of the parent node                         |
| metacells   | A data.frame() representing the cell metadata       |
| onlyPresent | A boolean(1)                                        |

**Value**

A data.frame containing the different tags and corresponding colors for the level given in parameter

A list

A vector

A vector

A vector

**Glossary**

Peptide-level vocabulary

└ 'Any' ||||└ 1.0 'Quantified' ||||└ 1.1 "Quant. by direct id" (color 4, white) ||||└ 1.2 "Quant. by recovery" (color 3, lightgrey) ||||└ 2.0 "Missing" (no color) ||||└ 2.1 "Missing POV" (color 1) ||||└ 2.2 'Missing MEC' (color 2) ||||└ 3.0 'Imputed' ||||└ 3.1 'Imputed POV' (color 1) ||||└ 3.2 'Imputed MEC' (color 2)

Protein-level vocabulary: └ 'Any' ||||└ 1.0 'Quantified' ||||└ 1.1 "Quant. by direct id" (color 4, white) ||||└ 1.2 "Quant. by recovery" (color 3, lightgrey) ||||└ 2.0 "Missing" ||||└ 2.1 "Missing POV" (color 1) ||||└ 2.2 'Missing MEC' (color 2) ||||└ 3.0 'Imputed' ||||└ 3.1 'Imputed POV' (color 1) ||||└ 3.2 'Imputed MEC' (color 2) ||||└ 4.0 'Combined tags' (color 3bis, lightgrey)

**Conversion to the glossary**

A generic conversion

Conversion for Proline datasets

Conversion from Maxquant datasets

**Author(s)**

Thomas Burger, Samuel Wieczorek

Samuel Wieczorek

**Examples**

```
if (interactive()) {
  metacell.def("protein")
  metacell.def("peptide")
}

Parent("protein", "Missing")
Parent("protein", "Missing POV")
Parent("protein", c("Missing POV", "Missing MEC"))
Parent("protein", c("Missing", "Missing POV", "Missing MEC"))

Children("protein", "Missing")
Children("protein", "Missing POV")
Children("protein", c("Missing POV", "Missing MEC"))
Children("protein", c("Missing", "Missing POV", "Missing MEC"))
data(vdata)
metacells <- get_metacell(vdata[[1]])
level <- get_type(vdata[[1]])
GetMetacellTags(metacells, level)
```

---

sub\_R25

---

*Feature example data*


---

**Description**

sub\_R25 is a protein subset of the dataset 'Exp1\_R25\_pept' in the package DAPARdata.

**Format**

An instance of the class MultiAssayExperiment

**Value**

An enriched instance of the class MultiAssayExperiment

**Source**

sub\_R25 was built from the source code available in [inst/scripts/build\\_datasets.R](#)

The DAPARdata package: <https://github.com/edyp-lab/DAPARdata>

---

|       |                             |
|-------|-----------------------------|
| vdata | <i>Feature example data</i> |
|-------|-----------------------------|

---

**Description**

vdata is a small object for testing and demonstration.

**Format**

An instance of the class `MultiAssayExperiment`

**Value**

An enriched instance of the class `MultiAssayExperiment`

**Source**

vdata was built from the source code available in `inst/scripts/build_datasets.R`

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