

# Package ‘iSEEu’

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**License** MIT + file LICENSE

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AggregatedDotPlot	<i>The AggregatedDotPlot class</i>
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---

## Description

Implements an aggregated dot plot where each feature/group combination is represented by a dot. The color of the dot scales with the mean assay value across all samples for a given group, while the size of the dot scales with the proportion of non-zero values across samples in that group.

## Slot overview

The following slots control the choice of features:

- CustomRows, a logical scalar indicating whether custom rows in CustomRowsText should be used. If TRUE, the feature identities are extracted from the CustomRowsText slot; otherwise they are defined from a transmitted row selection. Defaults to TRUE.

- CustomRowsText, a string containing the names of the features of interest, typically corresponding to the row names of the [SummarizedExperiment](#). Names should be new-line separated within this string. Defaults to the name of the first row in the SummarizedExperiment.

The following slots control the specification of groups:

- ColumnDataLabel, a string specifying the name of the [colData](#) field to use to group cells. The chosen field should correspond to a categorical factor. Defaults to the first categorical field.
- ColumnDataFacet, a string specifying the name of the [colData](#) field to use for faceting. The chosen field should correspond to a categorical factor. Defaults to "---", i.e., no faceting.

The following slots control the choice of assay values:

- Assay, a string specifying the name of the assay containing continuous values, to use for calculating the mean and the proportion of non-zero values. Defaults to the first valid assay name.

The following slots control the visualization parameters:

- VisualBoxOpen, a logical scalar indicating whether the visual parameter box should be open on initialization. Defaults to FALSE.
- VisualChoices, a character vector specifying the visualization options to show. Defaults to "Color" but can also include "Transform" and "Legend".

The following slots control the transformation of the mean values:

- MeanNonZeroes, a logical scalar indicating whether the mean should only be computed over non-zero values. Defaults to FALSE.
- Center, a logical scalar indicating whether the means for each feature should be centered across all groups. Defaults to FALSE.
- Scale, a logical scalar indicating whether the standard deviation for each feature across all groups should be scaled to unity. Defaults to FALSE.

The following slots control the color:

- UseCustomColormap, a logical scalar indicating whether to use a custom color scale. Defaults to FALSE, in which case the application-wide color scale defined by [ExperimentColorMap](#) is used.
- CustomColorLow, a string specifying the low color (i.e., at an average of zero) for a custom scale. Defaults to "grey".
- CustomColorHigh, a string specifying the high color for a custom scale. Defaults to "red".
- CenteredColormap, a string specifying the divergent colormap to use when Center is TRUE. Defaults to "blue < grey < orange"; other choices are "purple < black < yellow", "blue < grey < red" and "green < grey < red".

In addition, this class inherits all slots from its parent [Panel](#) class.

## Constructor

`AggregatedDotPlot(...)` creates an instance of a `AggregatedDotPlot` class, where any slot and its value can be passed to ... as a named argument.

## Supported methods

In the following code snippets, `x` is an instance of an `AggregatedDotPlot` class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x)` adds a "AggregatedDotPlot" entry containing `continuous.assay.names` and `discrete.colData.names`.
- `.refineParameters(x, se)` returns `x` after setting "Assay", "ColumnDataLabel" and "ColumnDataFacet" to valid values. If continuous assays or discrete `colData` variables are not available, NULL is returned instead.

For defining the interface:

- `.defineInterface(x, se, select_info)` creates an interface to modify the various parameters in the slots, mostly by calling the parent method and adding another visualization parameter box.
- `.defineDataInterface(x, se, select_info)` creates an interface to modify the data-related parameters, i.e., those that affect the position of the points.
- `.defineOutput(x)` defines the output HTML element.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.fullName(x)` will return "Aggregated dot plot".
- `.hideInterface(x)` will return TRUE for UI elements related to multiple row selections.

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pobjects, robjects)` will create all relevant observers for the UI elements.

For generating output:

- `.generateOutput(x, se, all_memory, all_contents)` will return the aggregated dot plot as a `ggplot` object, along with the commands used for its creation.
- `.renderOutput(x, se, output, pobjects, robjects)` will render the aggregated dot plot onto the interface.
- `.exportOutput(x, se, all_memory, all_contents)` will save the aggregated dot plot to a PDF file named after `x`, returning the path to the new file.

For providing documentation:

- `.definePanelTour(x)` will return a `data.frame` to be used in **rintrojs** as a panel-specific tour.

## Author(s)

Aaron Lun

## See Also

[Panel](#), for the immediate parent class.

[ComplexHeatmapPlot](#), for another panel with multi-row visualization capability.

## Examples

```
library(scRNAseq)

# Example data ----
sce <- ReprocessedAllenData(assays="tophat_counts")
class(sce)

library(scater)
sce <- logNormCounts(sce, exprs_values="tophat_counts")

# launch the app itself ----
if (interactive()) {
  iSEE(sce, initial=list(
    AggregatedDotPlot(ColumnDataLabel="Primary.Type")
  ))
}
```

---

createGeneSetCommands *Create gene set commands*

---

## Description

Create the commands required to populate [FeatureSetTables](#) with commonly used gene sets.

## Usage

```
createGeneSetCommands(
  collections = c("GO", "KEGG"),
  organism = "org.Hs.eg.db",
  identifier = "ENTREZID"
)
```

## Arguments

<code>collections</code>	Character vectors specifying the gene set collections of interest.
<code>organism</code>	String containing the <b>org.*.eg.db</b> package to use to extract mappings of gene sets to gene IDs.
<code>identifier</code>	String specifying the identifier to use to extract IDs for the organism package.

## Details

GO terms are extracted using the "GOALL" mode, which extracts both direct and indirect children of each term. A description for each GO term is extracted using the **GO.db** package.

Mappings of genes to KEGG pathway are extracted from the organism package using the "PATH" term. Unfortunately, this is not up to date due to the licensing around KEGG terms. Descriptions for each pathway are extracted from <http://rest.kegg.jp/list/pathway>.

The output of this function can be used as the commands argument of [registerFeatureSetCommands](#). It is also used by default in the [FeatureSetTable](#) constructor when no collections are registered.

**Value**

A list of two character vectors describing how to create collections and retrieve gene sets. This follows the expectations for commands in [registerFeatureSetCommands](#).

**Author(s)**

Aaron Lun

**See Also**

[FeatureSetTable](#), where the commands are intended for use.  
[registerFeatureSetCommands](#), to use the commands globally.

**Examples**

```
out <- createGeneSetCommands()
cat(out$collections['GO'], "\n")
cat(out$sets['GO'], "\n")
```

---

defunct

*Defunct functions*


---

**Description**

Pretty much as it says here. These functions are all defunct and should not be used.

**Usage**

```
.getAcceptablePValueFields(...)
.getAcceptableAveAbFields(...)
.getAcceptableLogFCFields(...)
.setAcceptablePValueFields(...)
.setAcceptableAveAbFields(...)
.setAcceptableLogFCFields(...)
```

**Arguments**

... Ignored.

**Value**

All functions error out with a defunct message pointing towards its replacement (if available).

**Author(s)**

Aaron Lun

---

DynamicMarkerTable-class

*Dynamic marker table*


---

## Description

A table that dynamically identifies marker genes for a selected subset of samples. Comparisons are made between the active selection in the transmitting panel and (i) all non-selected points, if no saved selections are available; or (ii) each subset of points in each saved selection.

## Slot overview

The following slots control the test procedure:

- `LogFC`, a numeric scalar indicating the log-fold change threshold to test against. Defaults to zero.
- `TestMethod`, string indicating the test to use (based on the `findMarkers` function from **scran**). This can be "t" (default), "wilcox" or "binom".
- `Assay`, string indicating the assay to use for testing. Defaults to the first named assay in the `SummarizedExperiment`.

The following slots control the rendered table:

- `ExtraFields`, a character vector containing names of `rowData` columns to be included in the table. Set to the output of `getTableExtraFields`. This cannot be changed once the application starts.

In addition, this class inherits all slots from its parent [RowTable](#), [Table](#) and [Panel](#) classes.

## Constructor

`DynamicMarkerTable(...)` creates an instance of a `DynamicMarkerTable` class, where any slot and its value can be passed to ... as a named argument.

## Supported methods

In the following code snippets, `x` is an instance of a [DynamicMarkerTable](#) class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x)` adds a "DynamicMarkerTable" entry containing `valid.assay.names` and `valid.rowdata.names`. This will also call the equivalent [RowTable](#) method.
- `.refineParameters(x, se)` returns `x` after setting "Assay" to the first valid value. This will also call the equivalent [RowTable](#) method for further refinements to `x`. If valid assay names are not available, `NULL` is returned instead. Any "ExtraFields" are intersected with the valid `rowData` names.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.
- `.panelColor(x)` will return the specified default color for this panel class.

- `.fullName(x)` will return "Dynamic marker table".
- `.hideInterface(x)` will return TRUE for UI elements related to multiple row selections, otherwise calling the method for [RowTable](#).

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pObjects, rObjects)` sets up observers for all new slots described above, as well as in the parent classes via the [RowTable](#) method.

For creating the table:

- `.generateTable(x, envir)` will create a data.frame of newly computed statistics in `envir`. The method will return the commands required to do so.

For documentation:

- `.definePanelTour(x)` returns an data.frame containing the steps of a panel-specific tour.

## Examples

```
library(scRNAseq)
library(scater)

sce <- ReprocessedAllenData(assays="tophat_counts")
sce <- logNormCounts(sce, exprs_values="tophat_counts")
sce <- runPCA(sce, ncomponents=4)
sce <- runTSNE(sce)

dst <- DynamicMarkerTable(PanelId=1L, PanelWidth=8L,
  ColumnSelectionSource="ReducedDimensionPlot1")

rdp <- ReducedDimensionPlot(PanelId=1L,
  ColorByFeatureSource="DynamicMarkerTable1")

if (interactive()) {
  iSEE(sce, initial=list(rdp, dst))
}
```

---

DynamicReducedDimensionPlot-class

*Dynamic reduced dimension plot*

---

## Description

A dimensionality reduction plot that dynamically recomputes the coordinates for the samples, based on the selected subset of samples (and possibly features) in transmitting panels. All samples in active and saved multiple selections are used here.



## Slot overview

The following slots control the thresholds used in the visualization:

- `Type`, a string specifying the type of dimensionality reduction method to use. This can be "PCA" (default), "TSNE" or "UMAP", which uses the relevant functions from the **scater** package.
- `NGenes`, an integer scalar specifying the number of highly variable genes to use in the dimensionality reduction. Only used if an explicit selection of features is not made in the app. Defaults to 1000.
- `Assay`, string indicating the assay to use for the calculations. Defaults to the first named assay in the `SummarizedExperiment`.

In addition, this class inherits all slots from its parent [ColumnDotPlot](#), [DotPlot](#) and [Panel](#) classes.

## Constructor

`DynamicReducedDimensionPlot(...)` creates an instance of a `DynamicReducedDimensionPlot` class, where any slot and its value can be passed to ... as a named argument.

## Supported methods

In the following code snippets, `x` is an instance of a [DynamicReducedDimensionPlot](#) class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x)` adds a "DynamicReducedDimensionPlot" entry containing `valid.assay.names`. This will also call the equivalent [ColumnDotPlot](#) method.
- `.refineParameters(x, se)` returns `x` after setting "Assay" to the first valid value. This will also call the equivalent [ColumnDotPlot](#) method for further refinements to `x`. If valid assay names are not available, `NULL` is returned instead.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.fullName(x)` will return "Dynamic reduced dimension plot".

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pobjects, robjects)` sets up observers for all new slots described above, as well as in the parent classes via the [ColumnDotPlot](#) method.

For creating the plot:

- `.generateDotPlotData(x, envir)` will create a `data.frame` of newly computed coordinates in `envir`. The method will return the commands required to do so as well as a list of labels.

For handling multiple selections:

- `.multiSelectionInvalidated(x)` will always return `TRUE`, as any change in the upstream selection of points will alter the coordinates and invalidate any brush/lasso on `x`.

For documentation:

- `.definePanelTour(x)` returns a `data.frame` containing the steps of a panel-specific tour.

**Author(s)**

Aaron Lun

**Examples**

```
library(scRNAseq)
library(scater)

sce <- ReprocessedAllenData(assays="tophat_counts")
sce <- logNormCounts(sce, exprs_values="tophat_counts")
sce <- runPCA(sce, ncomponents=4)
sce <- runTSNE(sce)

drdp <- DynamicReducedDimensionPlot(PanelId=1L, Assay="logcounts",
  ColumnSelectionSource="ReducedDimensionPlot1")

if (interactive()) {
  iSEE(sce, initial=list(ReducedDimensionPlot(PanelId=1L), drdp))
}
```

---

FeatureSetTable-class *Feature set table*

---

**Description**

A table where each row is itself a feature set and can be clicked to transmit a multiple feature selection to another panel. This relies on feature set collections that have been registered in the input [SummarizedExperiment](#), see [registerFeatureSetCollections](#) for more details. If no collections have been registered, we default to the GO and KEGG collections from [createGeneSetCommands](#).

**Slot overview**

The following slots control the feature sets in use:

- Collection, string specifying the type of feature set collection to show. Defaults to the first registered collection in the SummarizedExperiment.

The following slots control the table selections:

- Selected, a string containing the name of the currently selected gene set. Defaults to "", i.e., no selection.
- Search, a string containing the regular expression for the global search. Defaults to "", i.e., no search.
- SearchColumns, a character vector where each entry contains the search string for each column. Defaults to an empty character vector, i.e., no search.

In addition, this class inherits all slots from its parent [Panel](#) class.

**Constructor**

`FeatureSetTable(...)` creates an instance of a `FeatureSetTable` class, where any slot and its value can be passed to ... as a named argument.

## Supported methods

In the following code snippets, `x` is an instance of a [FeatureSetTable](#) class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x)` adds a "FeatureSetTable" entry containing `available.sets`, a named list of DataFrames containing information about the individual gene sets for each collection. This will also call the equivalent [Panel](#) method.
- `.refineParameters(x, se)` replaces NA values in `Collection` with the first valid collection. It also replaces NA values for `Selected` with the first valid set in the chosen collection. This will also call the equivalent [Panel](#) method.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.fullName(x)` will return "Gene set table".
- `.hideInterface(x)` will return TRUE for UI elements related to multiple selections, otherwise calling the method for [Panel](#).
- `.defineOutput(x)` will return a HTML element containing a [datatable](#) widget.

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pObjects, rObjects)` sets up observers for all new slots described above, as well as in the parent classes via the [Panel](#) method.

For creating the table:

- `.generateOutput(x, envir)` will create a data.frame of gene set descriptions in `envir`. It will also return the commands required to do so and the name of the variable corresponding to said data.frame.
- `.renderOutput(x, se, ..., output, pObjects, rObjects)` will add a [datatable](#) widget to the output, which is used to render the aforementioned data.frame.

For controlling the multiple selections:

- `.multiSelectionDimension(x)` returns "row".
- `.multiSelectionCommands(x, index)` returns a string specifying the commands to be used to extract the identities of the genes in the currently selected set. `index` is ignored.
- `.multiSelectionActive(x)` returns the name of the currently selected gene set, unless no selection is made, in which case NULL is returned.
- `.multiSelectionClear(x)` returns `x` but with the `Selected` slot replaced by an empty string.
- `.multiSelectionAvailable(x, contents)` returns `contents$available`, which is set to the number of features in `se`.

For documentation:

- `.definePanelTour(x)` returns an data.frame containing the steps of a panel-specific tour.

## Author(s)

Aaron Lun

## Examples

```
library(scRNAseq)
sce <- LunSpikeInData(location=FALSE)

library(scater)
sce <- logNormCounts(sce)

library(scran)
rowData(sce) <- cbind(rowData(sce), modelGeneVarWithSpikes(sce, "ERCC"))

cmds <- createGeneSetCommands(collections="GO",
                             organism="org.Mm.eg.db", identifier="ENSEMBL")
sce <- registerFeatureSetCommands(sce, cmds)

# Setting up the application.
gst <- FeatureSetTable(PanelId=1L)

rdp <- RowDataPlot(RowSelectionSource="FeatureSetTable1",
                  ColorBy="Row selection",
                  XAxis="Row data", XAxisRowData="mean", YAxis="total")

rdt <- RowDataTable(RowSelectionSource="FeatureSetTable1")

if (interactive()) {
  iSEE(sce, initial=list(gst, rdp, rdt))
}
```

---

GeneSetTable-class	<i>Gene set table</i>
--------------------	-----------------------

---

## Description

A table where each row is a gene set and can be clicked to transmit a multiple feature selection to another panel. This has been deprecated in favor of the simpler [FeatureSetTable](#).

## Slot overview

The following slots control the type of gene sets to show:

- `Type`, string specifying the type of gene set collection to show. Defaults to "GO".

The following slots control the table selections:

- `Selected`, a string containing the name of the currently selected gene set. Defaults to "", i.e., no selection.
- `Search`, a string containing the regular expression for the global search. Defaults to "", i.e., no search.
- `SearchColumns`, a character vector where each entry contains the search string for each column. Defaults to an empty character vector, i.e., no search.

In addition, this class inherits all slots from its parent [Panel](#) class.

## Constructor

`GeneSetTable(...)` creates an instance of a `GeneSetTable` class, where any slot and its value can be passed to ... as a named argument.

## Supported methods

In the following code snippets, `x` is an instance of a `GeneSetTable` class. Refer to the documentation for each method for more details on the remaining arguments.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.fullName(x)` will return "Gene set table".
- `.hideInterface(x)` will return TRUE for UI elements related to multiple selections, otherwise calling the method for `Panel`.
- `.defineOutput(x)` will return a HTML element containing a `datatable` widget.

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pobjects, robjects)` sets up observers for all new slots described above, as well as in the parent classes via the `Panel` method.

For creating the table:

- `.generateOutput(x, envir)` will create a data.frame of gene set descriptions in `envir`, based on the `mode="show"` output of `.getGeneSetCommands`. It will also return the commands required to do so and the name of the variable corresponding to said data.frame.
- `.renderOutput(x, se, ..., output, pobjects, robjects)` will add a `datatable` widget to the output, which is used to render the aforementioned data.frame.

For controlling the multiple selections:

- `.multiSelectionDimension(x)` returns "row".
- `.multiSelectionCommands(x, index)` returns a string specifying the commands to be used to extract the identities of the genes in the currently selected set, based on the `mode="extract"` output of `.getGeneSetCommands`. `index` is ignored.
- `.multiSelectionActive(x)` returns the name of the currently selected gene set, unless no selection is made, in which case NULL is returned.
- `.multiSelectionClear(x)` returns `x` but with the Selected slot replaced by an empty string.
- `.multiSelectionAvailable(x, contents)` returns `contents$available`, which is set to the number of features in `se`.

## Author(s)

Aaron Lun

**Examples**

```

library(scRNAseq)
sce <- LunSpikeInData(location=FALSE)

library(scater)
sce <- logNormCounts(sce)

library(scran)
rowData(sce) <- cbind(rowData(sce), modelGeneVarWithSpikes(sce, "ERCC"))

# This defaults to 'org.Hs.eg.db' with 'ENTREZID'.
.setOrganism("org.Mm.eg.db")
.setIdentifierType("ENSEMBL")
gst <- GeneSetTable(PanelId=1L)

rdp <- RowDataPlot(RowSelectionSource="GeneSetTable1",
  ColorBy="Row selection",
  XAxis="Row data", XAxisRowData="mean", YAxis="total")

rdt <- RowDataTable(RowSelectionSource="GeneSetTable1")

if (interactive()) {
  iSEE(sce, initial=list(gst, rdp, rdt))
}

```

getPValuePattern

*Global DE prefixes***Description**

Get or set patterns for acceptable names of [rowData](#) columns related to a differential expression analysis. These functions are deprecated; use their counterparts in [?"registerPValuePatterns"](#) instead.

**Usage**

```

getPValuePattern()

getLogFCPattern()

getAveAbPattern()

setPValuePattern(value)

setLogFCPattern(value)

setAveAbPattern(value)

```

**Arguments**

**value** A character vector containing the acceptable prefixes for each statistic.

## Details

These utilities allow users to easily get and set the patterns of acceptable fields in all [VolcanoPlots](#), [MAPlots](#) and [LogFCLogFCPlots](#) at once. Any global settings only take effect (i) during setup of the [iSEE](#) application and (ii) if the first panel of each class does not have existing values in the "PValueFields", "LogFCFields" or "AveAbFields" slots (which take precedence if present).

Each of these global settings are treated as *patterns* for partial matching. For the "PValue" pattern, columns with the names "PValue.X" and "X.PValue" will be considered acceptable matches. All partial matching must be exact - regular expressions are not supported.

## Value

getPValuePattern returns the patterns for acceptable column names for p-values.

getLogFCPattern returns the patterns for acceptable column names for log-fold changes.

getAveAbPattern returns the patterns for acceptable column names for the average abundances.

The corresponding setters set the global parts for each statistic and return NULL invisibly.

## Author(s)

Aaron Lun

## See Also

[VolcanoPlot](#), [MAPlot](#) and [LogFCLogFCPlot](#), which are affected by these globals.

## Examples

```
old <- getPValuePattern()

setPValuePattern(LETTERS)
getPValuePattern()

setPValuePattern(old)
```

---

getTableExtraFields	<i>Global extra table fields</i>
---------------------	----------------------------------

---

## Description

Get or set the names of the extra fields to include in a table.

## Usage

```
getTableExtraFields()

setTableExtraFields(value)
```

## Arguments

value                    A character vector containing the names of extra fields to include.

**Details**

These utilities allow users to easily set the feature set commands for all [DynamicMarkerTables](#) at once. Any global settings only take effect (i) during setup of the [iSEE](#) application and (ii) if the first [DynamicMarkerTable](#) does not have an existing values in the "TableExtraFields" slots.

**Value**

`getTableExtraFields` returns the current global extra table fields.

`setTableExtraFields` will set the current global extra table fields and return NULL invisibly.

**Author(s)**

Aaron Lun

**Examples**

```
old <- getTableExtraFields()

setTableExtraFields(LETTERS)
getTableExtraFields()

setTableExtraFields(old)
```

---

iSEEu-pkg

*iSEEu: iSEE Universe*


---

**Description**

iSEEu is a package that provides modes and panels for iSEE, allowing easy configuration of iSEE applications.

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**See Also**

Useful links:

- <https://github.com/iSEE/iSEEu>
- Report bugs at <https://github.com/iSEE/iSEEu/issues>



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LogFCLogFCPlot-class    *The LogFCLogFCPlot class*


---

## Description

The LogFCLogFCPlot is a [RowDataPlot](#) subclass that is dedicated to creating a scatter plot of two log-fold changes. Each axis contains the log-fold change for a differential expression analysis and each point represents a feature.

## Slot overview

The following slots control the thresholds used in the visualization:

- `XPValueField`, a string specifying the field of `rowData` containing the p-values for the x-axis comparison.
- `YPValueField`, a string specifying the field of `rowData` containing the p-values for the y-axis comparison.
- `PValueThreshold`, a numeric scalar in (0, 1] specifying the threshold to use on the (adjusted) p-value. Defaults to 0.05.
- `LogFCThreshold`, a non-negative numeric scalar specifying the threshold to use on the log-fold change. Defaults to 0.
- `PValueCorrection`, a string specifying the multiple testing correction to apply. Defaults to "BH", but can take any value from `p.adjust.methods`.

In addition, this class inherits all slots from its parent [RowDataPlot](#), [RowDotPlot](#), [DotPlot](#) and [Panel](#) classes.

## Constructor

`LogFCLogFCPlot(...)` creates an instance of a LogFCLogFCPlot class, where any slot and its value can be passed to ... as a named argument.

Users are expected to load relevant statistics into the `rowData` of a [SummarizedExperiment](#). There should be two columns for the p-values from each comparison - and another two for the corresponding log-fold changes - for each gene/row, see Examples. The expected column names (and how to tune them) are listed at `?registerPValueFields`.

## Supported methods

In the following code snippets, `x` is an instance of a [RowDataPlot](#) class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x, se)` returns `se` after being loaded with class-specific constants. This includes `"valid.p.fields"` and `"valid.lfc.fields"`, character vectors containing the names of valid `rowData` columns for the p-values and log-fold changes, respectively.
- `.refineParameters(x, se)` returns `x` after setting `XAxis="Row data"` as well as `"PValuePattern"` and `"LogFCPattern"` to their corresponding cached values. This will also call the equivalent [RowDataPlot](#) method for further refinements to `x`. If valid p-value and log-fold change fields are not available, NULL is returned instead.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.allowableXAxisChoices(x, se)` returns a character vector specifying the acceptable log-fold change-related variables in `rowData(se)` that can be used as choices for the x-axis.
- `.allowableYAxisChoices(x, se)` returns a character vector specifying the acceptable log-fold change-related variables in `rowData(se)` that can be used as choices for the y-axis.
- `.hideInterface(x, field)` will return TRUE for `field="XAxis"`, otherwise it will call the `RowDataPlot` method.
- `.fullName(x)` will return "LogFC-logFC plot".

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pObjects, rObjects)` sets up observers for all new slots described above, as well as in the parent classes via the `RowDataPlot` method.

For creating the plot:

- `.generateDotPlotData(x, envir)` will create a data.frame of row metadata variables in `envir`. This contains the two sets of log-fold changes on both axes, plus an extra field specifying whether or not the feature was considered to be significantly up or down. The method will return the commands required to do so as well as a list of labels.
- `.prioritizeDotPlotData(x, envir)` will create variables in `envir` marking the priority of points. Significant features receive higher priority (i.e., are plotted over their non-significant counterparts) and are less aggressively downsampled when `Downsample=TRUE`. The method will return the commands required to do this as well as a logical scalar indicating that rescaling of downsampling resolution is performed.
- `.colorByNoneDotPlotField(x)` will return a string specifying the field of the data.frame (generated by `.generateDotPlotData`) containing the significance information. This is to be used for coloring when `ColorBy="None"`.
- `.colorByNoneDotPlotScale(x)` will return a string containing a **ggplot2** command to add a default color scale when `ColorBy="None"`.
- `.generateDotPlot(x, labels, envir)` returns a list containing plot and commands, using the initial `ColumnDataPlot ggplot` and adding horizontal lines demarcating the log-fold change threshold.

For documentation:

- `.definePanelTour(x)` returns an data.frame containing the steps of a panel-specific tour.
- `.getDotPlotColorHelp(x, color_choices)` returns a function that generates an **rintrojs** tour for the color choice UI.

### Author(s)

Aaron Lun

### See Also

[RowDataPlot](#), for the base class.

## Examples

```
# Making up some results:
se <- SummarizedExperiment(matrix(rnorm(10000), 1000, 10))
rownames(se) <- paste0("GENE_", seq_len(nrow(se)))
rowData(se)$PValue1 <- runif(nrow(se))
rowData(se)$LogFC1 <- rnorm(nrow(se))
rowData(se)$PValue2 <- runif(nrow(se))
rowData(se)$LogFC2 <- rnorm(nrow(se))

if (interactive()) {
  iSEE(se, initial=list(LogFCLogFCPlot(XAxisRowData="LogFC1", YAxis="LogFC2",
    XPValueField="PValue1", YPValueField="PValue2")))
}
```

---

MAPlot-class

*The MAPlot class*


---

## Description

The MAPlot is a [RowDataPlot](#) subclass that is dedicated to creating a MA plot. It retrieves the log-fold change and average abundance and creates a row-based plot where each point represents a feature.

## Slot overview

The following slots control the thresholds used in the visualization:

- `PValueField`, a string specifying the field of [rowData](#) containing the p-values.
- `PValueThreshold`, a numeric scalar in (0, 1] specifying the threshold to use on the (adjusted) p-value. Defaults to 0.05.
- `LogFCThreshold`, a non-negative numeric scalar specifying the threshold to use on the log-fold change. Defaults to 0.
- `PValueCorrection`, a string specifying the multiple testing correction to apply. Defaults to "BH", but can take any value from [p.adjust.methods](#).

In addition, this class inherits all slots from its parent [RowDataPlot](#), [RowDotPlot](#), [DotPlot](#) and [Panel](#) classes.

## Constructor

`MAPlot(...)` creates an instance of a MAPlot class, where any slot and its value can be passed to ... as a named argument.

Users are expected to load relevant statistics into the `rowData` of a [SummarizedExperiment](#). This panel expects one or more columns containing the p-values, log-fold changes and average abundances for each gene/row - see Examples. The expected column names (and how to tune them) are listed at [?"registerPValueFields"](#).

## Supported methods

In the following code snippets, `x` is an instance of a `RowDataPlot` class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x, se)` returns `se` after being loaded with class-specific constants. This includes `"valid.p.fields"`, `"valid.ab.fields"` and `"valid.lfc.fields"`, which are character vectors containing the names of valid `rowData` columns for the p-values, average abundances and log-fold changes, respectively.
- `.refineParameters(x, se)` returns `x` after setting `XAxis="Row data"` and the various `*Pattern` fields to their cached values. This will also call the equivalent `RowDataPlot` method for further refinements to `x`. If valid p-value, abundance and log-fold change fields are not available, `NULL` is returned instead.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.allowableXAxisChoices(x, se)` returns a character vector specifying the acceptable average abundance-related variables in `rowData(se)` that can be used as choices for the x-axis.
- `.allowableYAxisChoices(x, se)` returns a character vector specifying the acceptable log-fold change-related variables in `rowData(se)` that can be used as choices for the y-axis.
- `.hideInterface(x, field)` will return `TRUE` for `field="XAxis"`, otherwise it will call the `RowDataPlot` method.
- `.fullName(x)` will return `"MA plot"`.

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pObjects, rObjects)` sets up observers for all new slots described above, as well as in the parent classes via the `RowDataPlot` method.

For creating the plot:

- `.generateDotPlotData(x, envir)` will create a data.frame of row metadata variables in `envir`. This should contain average abundances on the x-axis and log-fold changes on the y-axis, in addition to an extra field specifying whether or not the feature was considered to be significantly up or down. The method will return the commands required to do so as well as a list of labels.
- `.prioritizeDotPlotData(x, envir)` will create variables in `envir` marking the priority of points. Significant features receive higher priority (i.e., are plotted over their non-significant counterparts) and are less aggressively downsampled when `Downsample=TRUE`. The method will return the commands required to do this as well as a logical scalar indicating that rescaling of downsampling resolution is performed.
- `.colorByNoneDotPlotField(x)` will return a string specifying the field of the data.frame (generated by `.generateDotPlotData`) containing the significance information. This is to be used for coloring when `ColorBy="None"`.
- `.colorByNoneDotPlotScale(x)` will return a string containing a `ggplot2` command to add a default color scale when `ColorBy="None"`.
- `.generateDotPlot(x, labels, envir)` returns a list containing plot and commands, using the initial `ColumnDataPlot ggplot` and adding horizontal lines demarcating the log-fold change threshold.

For documentation:

- [.definePanelTour](#)(x) returns an data.frame containing the steps of a panel-specific tour.
- [.getDotPlotColorHelp](#)(x, color\_choices) returns a function that generates an **rintrojs** tour for the color choice UI.

### Author(s)

Aaron Lun

### See Also

[RowDataPlot](#), for the base class.

### Examples

```
# Making up some results:
se <- SummarizedExperiment(matrix(rnorm(10000), 1000, 10))
rownames(se) <- paste0("GENE_", seq_len(nrow(se)))
rowData(se)$PValue <- runif(nrow(se))
rowData(se)$LogFC <- rnorm(nrow(se))
rowData(se)$AveExpr <- rnorm(nrow(se))

if (interactive()) {
  iSEE(se, initial=list(MAPlot()))
}
```

---

MarkdownBoard-class      *The MarkdownBoard class*

---

### Description

The MarkdownBoard class renders user-supplied Markdown into HTML to display inside the app. This is useful for displaying information alongside other panels, or for users to jot down their own notes.

### Slot overview

The following slots are relevant to the rendered content:

- Content, a string containing Markdown-formatted text. This will be rendered to HTML for display inside the app.

In addition, this class inherits all slots from its parent [Panel](#) class.

### Constructor

`MarkdownBoard(...)` creates an instance of a MarkdownBoard class, where any slot and its value can be passed to ... as a named argument.

## Supported methods

In the following code snippets, `x` is an instance of a [RowDataPlot](#) class. Refer to the documentation for each method for more details on the remaining arguments.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for editing the Content.
- `.panelColor(x)` will return the specified default color for this panel class.
- `.hideInterface(x, field)` will return TRUE for all selection-related parameters.
- `.fullName(x)` will return "Volcano plot".

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pobjects, robjects)` sets up observers for all new slots described above, as well as in the parent classes via the [RowDataPlot](#) method.

For rendering the display:

- `.defineOutput(x)` will return a UI element to display the HTML.
- `.renderOutput(x, se, ..., output, pobjects, robjects)` will add reactive expressions to render the HTML.
- `.generateOutput(x, se, all_memory, all_contents)` will render the Markdown to HTML via the `rmarkdown` package, returning a string containing the rendered content in the text element of the output list. The Markdown-formatted content is converted into an R comment for code tracking purposes.
- `.exportOutput(x, se, all_memory, all_contents)` will create a HTML containing the rendered Markdown, and return a string containing the path to that HTML.

For documentation:

- `.definePanelTour(x)` returns an data.frame containing the steps of a panel-specific tour. Not that there's a great deal to say here.

## Author(s)

Aaron Lun

## See Also

[Panel](#), for the base class.

## Examples

```
if (interactive()) {
  iSEE(SummarizedExperiment(), initial=list(MarkdownBoard()))
}
```

---

modeEmpty

*App pre-configured to launch with no visible panel*


---

**Description**

This mode launches an app that does not display any panel.

**Usage**

```
modeEmpty(...)
```

**Arguments**

... Arguments passed to [iSEE\(\)](#).

**Details**

This mode presents the advantage to launch an interface in a minimal amount of time, as it does not need to render any panel when the interface is launched. Users can then use the "Organize panels" widget to select panels to display in the interface.

**Value**

A Shiny app object is returned.

**Examples**

```
example("SingleCellExperiment")
rownames(sce) <- paste0("G", 1:200)
colnames(sce) <- paste0("C", 1:100)

app <- modeEmpty(sce)
if (interactive()) {
  shiny::runApp(app, port=1234)
}
```

---

modeGating

*App pre-configured to link multiple feature assay plots*


---

**Description**

This mode launches a Shiny App preconfigured with multiple chain-linked feature expression plots for interactive data exploration of the [SingleCellExperiment](#) or [SummarizedExperiment](#) object.

**Usage**

```
modeGating(se, features, plotAssay = NA_character_, ..., plotWidth = 4)
```

**Arguments**

se	An object that coercible to <a href="#">SingleCellExperiment-class</a>
features	data.frame with columns named x and y that define the features on the axes of the linked plots. Plots are serially linked from the first row to the last.
plotAssay	The assay (one of assayNames(se)) to use for the plots (character vector of length either 1 or equal to nrow(features)).
...	Additional arguments passed to <a href="#">iSEE()</a> .
plotWidth	The grid width of linked plots (numeric vector of length either 1 or equal to nrow(features))

**Value**

A Shiny app object is returned.

**Examples**

```
library(scRNAseq)

# Example data ----
sce <- ReprocessedAllenData(assays="tophat_counts")
class(sce)

library(scater)
sce <- logNormCounts(sce, exprs_values="tophat_counts")

# Select top variable genes ----

plot_count <- 6
rv <- rowVars(assay(sce, "tophat_counts"))
top_var <- head(order(rv, decreasing=TRUE), plot_count*2)
top_var_genes <- rownames(sce)[top_var]

plot_features <- data.frame(
  x=head(top_var_genes, plot_count),
  y=tail(top_var_genes, plot_count),
  stringsAsFactors=FALSE
)

# launch the app itself ----

app <- modeGating(sce, features = plot_features)
if (interactive()) {
  shiny::runApp(app, port=1234)
}
```

---

modeReducedDim

*App pre-configured to compare multiple reduced dimension plots*


---

**Description**

This mode launches a Shiny App preconfigured with multiple linked reduced dimension plots for interactive data exploration of the [SingleCellExperiment](#) object.



**Usage**

```
modeReducedDim(
  se,
  includeNames = reducedDimNames(se),
  colorBy = NULL,
  ...,
  plotWidth = NULL
)
```

**Arguments**

<code>se</code>	An object that coercible to <a href="#">SingleCellExperiment</a>
<code>includeNames</code>	Character vector with the names of reduced dimensions to display as individual panels. The default uses all available in <code>reducedDimNames(se)</code> .
<code>colorBy</code>	Character scalar controlling coloring of cells. Must match either to one of <code>colnames(colData(se))</code> or <code>rownames(se)</code> . If coloring by a <code>colData</code> column, a column data plot is opened in addition to the reduced dimension panels. If coloring by a feature, a row statistics table is opened in addition to the reduced dimension panels, from which the latter are receiving the color.
<code>...</code>	Additional arguments passed to <a href="#">iSEE</a> .
<code>plotWidth</code>	The grid width of linked plots (numeric vector of length either 1 or equal to <code>length(includeNames)</code> ). The total width of the window is 12, so <code>plotWidth = 4</code> for example will show three panels per row. If <code>plotWidth = NULL</code> (the default), a value will be estimated depending on the number of reduced dimension panels.

**Value**

A Shiny app object is returned.

**Examples**

```
library(scRNAseq)

# Example data ----
sce <- ReprocessedAllenData(assays="tophat_counts")
class(sce)

library(scater)
sce <- logNormCounts(sce, exprs_values="tophat_counts")
sce <- runPCA(sce, ncomponents = 30)
sce <- runTSNE(sce)
sce <- runUMAP(sce)
reducedDimNames(sce)

# launch the app ----
# ... coloring by a column data variable
app <- modeReducedDim(sce, colorBy = "Primary.Type")
if (interactive()) {
  shiny::runApp(app, port=1234)
}
# ... coloring by a feature
app <- modeReducedDim(sce, colorBy = "Scnn1a")
if (interactive()) {
```

```

    shiny::runApp(app, port=1234)
  }

```

---

registerDEFields	<i>Register DE-related fields</i>
------------------	-----------------------------------

---

## Description

Register the names of fields containing various DE statistics, to populate the user interface of DE-related [Panels](#).

## Usage

```

registerPValueFields(se, fields)

registerAveAbFields(se, fields)

registerLogFCFields(se, fields)

registerPValuePatterns(se, patterns)

registerAveAbPatterns(se, patterns)

registerLogFCPatterns(se, patterns)

getPValueFields(se)

getAveAbFields(se)

getLogFCFields(se)

getPValuePatterns(se, defaults = c("PValue", "p.value", "pval"))

getAveAbPatterns(se, defaults = c("AveExpr", "logCPM"))

getLogFCPatterns(se, defaults = c("logFC", "LogFC"))

```

## Arguments

se	A <a href="#">SummarizedExperiment</a> to be visualized with various DE-related Panels. This is expected to have a number of DE-related fields in its <a href="#">rowData</a> .
fields	A character vector containing the names of the relevant fields containing the DE statistics. Alternatively NULL to remove any existing setting.
patterns	A character vector containing partial names, to match against the colnames of the <a href="#">rowData</a> to identify relevant fields containing DE statistics. Alternatively NULL to remove any existing setting.
defaults	Character vector specifying the default patterns to provide when no patterns were registered in se.

## Details

DE-related Panels need to find relevant `rowData` fields containing p-values, log-fold changes, etc. to set appropriate defaults in the user interface. These functions allow a user to tune the definition of what those Panels consider to be relevant, which is occasionally necessary if the DE statistics are stored in a `rowData` field with an unusual column name. The idea is to register the relevant fields in `se`, which can then be supplied to `iSEE` with the affected Panels - see Examples.

The registered fields should be the names of appropriate columns in `rowData` containing continuous variables. Columns containing categorical or non-atomic variables will generally be ignored. For each DE statistic, if any fields are registered in `se`, they will be used directly and patterns will be ignored.

The registered patterns are used for partial name matching to the names of appropriate columns of `rowData`. All partial matching must be exact - regular expressions are not supported. Matches can occur anywhere in the name. For example, with "PValue", columns with the names "PValue.X" and "X.PValue" will be considered acceptable matches. If no patterns are supplied, the Panels will use the values in defaults.

## Value

All register functions will return `se`, modified to contain the supplied patterns or fields. These will be used as suggestions by DE-related Panels to identify the relevant fields.

All get functions will return a character vector containing the value set by the corresponding register function; or NULL, if nothing was set.

## Author(s)

Aaron Lun

## Examples

```
# Making up some results with unusual names.
se <- SummarizedExperiment(matrix(rnorm(10000), 1000, 10))
rownames(se) <- paste0("GENE_", seq_len(nrow(se)))
rowData(se)$pvalue <- runif(nrow(se))
rowData(se)$lfc <- rnorm(nrow(se))
rowData(se)$average <- rnorm(nrow(se))

se <- registerPValueFields(se, "pvalue")
getPValueFields(se)
se <- registerAveAbFields(se, "average")
getAveAbFields(se)
se <- registerLogFCFields(se, "lfc")
getLogFCFields(se)

if (interactive()) {
  iSEE(se, initial=list(MAPlot()))
}
```

---

```
registerFeatureSetCollections
  Register feature set collections
```

---

## Description

Register feature set collations and their annotations for display in [FeatureSetTables](#).

## Usage

```
registerFeatureSetCollections(se, collections)

registerFeatureSetCommands(se, commands)

getFeatureSetCollections(se)

getFeatureSetCommands(se)
```

## Arguments

se	The <a href="#">SummarizedExperiment</a> object to be used in <a href="#">iSEE</a> .
collections	A named list containing one or more <a href="#">CharacterList</a> objects. Each entry represents a collection of feature sets (see Details) and should be named.
commands	A named list containing two character vectors of commands to use to generate collections and sets.

## Details

Arbitrary feature sets are challenging as there is no obvious place to store them. `registerFeatureSetCollections` and friends will insert these sets into the [metadata](#) of the `SummarizedExperiment` object, allowing the corresponding getter functions to quickly extract them later within the `iSEE` app.

`collections` should be a named list containing [CharacterList](#) objects. Each [CharacterList](#) represents a collection where each entry is a feature set, i.e., a character vector corresponding to some of the row names of `se`. The [mcols](#) can contain additional per-set fields (e.g., descriptions, enrichment statistics) that will be shown in the [FeatureSetTable](#).

`commands` should be a list containing:

- `collections`, a named character vector where each entry is named after a feature set collection. Each entry should be a string containing R commands to define a `data.frame` named `tab`, where each row is a feature set and the row names are the names of those sets.
- `sets`, a character vector where each entry is named after a feature set collection in the same order as `commands$collections`. Each entry should be a string containing R commands to define a character vector named `selected` containing the identity of all rows of the `SummarizedExperiment` in the set of interest. (These commands can assume that a `.set_id` variable is present containing the name of the chosen feature set, as well as the `se` variable containing the input `SummarizedExperiment` object.)

If neither `collections` nor `commands` are provided, any previously registered content in `se` is removed.

**Value**

For registerFeatureSetCollections and registerFeatureSetCommands, a modified se is returned that contains the feature set collections or commands, respectively. This can be used with [FeatureSetTables](#) in [iSEE](#) calls.

For getFeatureSetCollections, the list of CharacterLists is returned. Alternatively NULL, if no such list was stored by registerFeatureSetCollections.

For getFeatureSetCommands, the list of of commands is returned containing collections and sets. Alternatively NULL, if no such list was stored by registerFeatureSetCommands.

**Author(s)**

Aaron Lun

**Examples**

```
library(scRNAseq)
sce <- LunSpikeInData(location=FALSE)

# Make up some random collections.
random <- CharacterList(
  Aaron = sample(rownames(sce), 10),
  Kevin = sample(rownames(sce), 20),
  Charlotte = sample(rownames(sce), 30),
  Fed = sample(rownames(sce), 40)
)
mcols(random)$p.value <- runif(4)

# Storing the collections inside our SummarizedExperiment.
sce <- registerFeatureSetCollections(sce, list(random=random))
getFeatureSetCollections(sce)

if (interactive()) {
  iSEE(sce, initial=list(FeatureSetTable()))
}
```

---

setFeatureSetCommands *Global feature set commands*

---

**Description**

Set the commands to define the global collection of feature sets. This is deprecated in favor of [registerFeatureSetCommands](#).

**Usage**

```
setFeatureSetCommands(value)
```

**Arguments**

value	A list of two character vectors named "collections" and "sets". Both vectors should be of the same length and have the same names. Vectors should contain R commands to create collections and retrieve sets; see <a href="#">?FeatureSetTable</a> and the output of <a href="#">createGeneSetCommands</a> for details.
-------	---

**Value**

setFeatureSetCommands will set the current global feature set commands and return NULL invisibly.

**Author(s)**

Aaron Lun

**See Also**

[createGeneSetCommands](#), for one method of generating value.

**Examples**

```
old <- getFeatureSetCommands()

new.cmds <- createGeneSetCommands(organism="org.Mm.eg.db",
  identifier="SYMBOL")
setFeatureSetCommands(new.cmds)

getFeatureSetCommands()

setFeatureSetCommands(old)
```

---

utils-geneset

*Gene set utilities*


---

**Description**

Utility functions to control the behavior of the [GeneSetTable](#).

**Usage**

```
.getIdentifierType()

.setIdentifierType(value)

.getOrganism()

.setOrganism(value)

.getGeneSetCommands(collection, mode)

.setGeneSetCommands(value)
```

**Arguments**

value	For .setIdentifierType and .setOrganism, a string containing the type of identifier or organism package to use. For .setGeneSetCommands, a named list containing two character vectors, see Details.
collection	String specifying the gene set collection.
mode	String specifying the mode of operation for the returned commands.

## Details

By default, `.getGeneSetCommands` will extract GO and KEGG terms. The organism and identifier type relates to the manner in which this default extraction is performed.

Users can add their own gene set collections by supplying a named list to `.setGeneSetCommands`. Each element of the list should be a named character vector of length two, with names "show" and "extract" - see the return value for what these are. The names of the list should be unique and will be used in the [GeneSetTable](#) interface.

Alternatively, any element of the list may be NULL, in which case it is excluded from the interface. This is useful for setting, e.g., `GO=NULL` to ignore the in-built GO terms.

## Value

`.getIdentifierType` will return the identifier type to use, defaulting to "ENTREZID".

`.getOrganism` will return the organism package to use, defaulting "org.Hs.eg.db".

`.getGeneSetCommands` will return:

- If `mode="show"`, a string containing R commands that create `tab`, a data.frame of all gene sets for a given collection.
- If `mode="extract"`, a format string containing R commands that (after formatting) create `selected`, a character vector of gene identities for the selected gene set. This format string should accept one string argument corresponding to the deparsed name of the gene set.

Each of the setter functions will set the corresponding option and return NULL, invisibly.

## Author(s)

Aaron Lun

## See Also

[GeneSetTable](#), where these functions have their effect.

## Examples

```
.setIdentifierType("ENSEMBLID")
.getIdentifierType()

.setOrganism("org.Mm.eg.db")
.getOrganism()

.getGeneSetCommands("GO", "show")
.getGeneSetCommands("GO", "extract")

.setGeneSetCommands(
  list(AaronRandomCollection=
    c(
      show='tab <- some_function_to_list_my_gene_sets()',
      extract='selected <- some_function_to_get_one_gene_set(%s)'
    )
  )
)

.getGeneSetCommands("AaronRandomCollection", "show")
.getGeneSetCommands("AaronRandomCollection", "extract")
```

---

VolcanoPlot-class      *The VolcanoPlot class*


---

### Description

The VolcanoPlot is a [RowDataPlot](#) subclass that is dedicated to creating a volcano plot. It retrieves the log-fold change and p-value from and creates a row-based plot where each point represents a feature.

### Slot overview

The following slots control the thresholds used in the visualization:

- `PValueThreshold`, a numeric scalar in (0, 1] specifying the threshold to use on the (adjusted) p-value. Defaults to 0.05.
- `LogFCThreshold`, a non-negative numeric scalar specifying the threshold to use on the log-fold change. Defaults to 0.
- `PValueCorrection`, a string specifying the multiple testing correction to apply. Defaults to "BH", but can take any value from [p.adjust.methods](#).

In addition, this class inherits all slots from its parent [RowDataPlot](#), [RowDotPlot](#), [DotPlot](#) and [Panel](#) classes.

### Constructor

`VolcanoPlot(...)` creates an instance of a VolcanoPlot class, where any slot and its value can be passed to ... as a named argument.

Users are expected to load relevant statistics into the `rowData` of a [SummarizedExperiment](#). This panel expects one or more columns containing the p-values and log-fold changes for each gene/row - see Examples. The expected column names (and how to tune them) are listed at [?"registerPValueFields"](#).

### Supported methods

In the following code snippets, `x` is an instance of a [RowDataPlot](#) class. Refer to the documentation for each method for more details on the remaining arguments.

For setting up data values:

- `.cacheCommonInfo(x, se)` returns `se` after being loaded with class-specific constants. This includes `"valid.p.fields"` and `"valid.lfc.fields"`, character vectors containing the names of valid `rowData` columns for the p-values and log-fold changes, respectively.
- `.refineParameters(x, se)` returns `x` after setting `XAxis="Row data"` and the various `*Pattern` fields to their cached values. This will also call the equivalent [RowDataPlot](#) method for further refinements to `x`. If valid p-value and log-fold change fields are not available, NULL is returned instead.

For defining the interface:

- `.defineDataInterface(x, se, select_info)` returns a list of interface elements for manipulating all slots described above.



- `.panelColor(x)` will return the specified default color for this panel class.
- `.allowableXAxisChoices(x, se)` returns a character vector specifying the acceptable log-fold change-related variables in `rowData(se)` that can be used as choices for the x-axis.
- `.allowableYAxisChoices(x, se)` returns a character vector specifying the acceptable p-value-related variables in `rowData(se)` that can be used as choices for the y-axis.
- `.hideInterface(x, field)` will return TRUE for `field="XAxis"`, otherwise it will call the `RowDataPlot` method.
- `.fullName(x)` will return "Volcano plot".

For monitoring reactive expressions:

- `.createObservers(x, se, input, session, pObjects, rObjects)` sets up observers for all new slots described above, as well as in the parent classes via the `RowDataPlot` method.

For creating the plot:

- `.generateDotPlotData(x, envir)` will create a data.frame of row metadata variables in `envir`. This should contain negative log-transformed p-values on the y-axis and log-fold changes on the x-axis, in addition to an extra field specifying whether or not the feature was considered to be significantly up or down. The method will return the commands required to do so as well as a list of labels.
- `.prioritizeDotPlotData(x, envir)` will create variables in `envir` marking the priority of points. Significant features receive higher priority (i.e., are plotted over their non-significant counterparts) and are less aggressively downsampled when `Downsample=TRUE`. The method will return the commands required to do this as well as a logical scalar indicating that rescaling of downsampling resolution is performed.
- `.colorByNoneDotPlotField(x)` will return a string specifying the field of the data.frame (generated by `.generateDotPlotData`) containing the significance information. This is to be used for coloring when `ColorBy="None"`.
- `.colorByNoneDotPlotScale(x)` will return a string containing a **ggplot2** command to add a default color scale when `ColorBy="None"`.
- `.generateDotPlot(x, labels, envir)` returns a list containing plot and commands, using the initial `ColumnDataPlot` **ggplot** and adding vertical lines demarcating the log-fold change threshold.

For documentation:

- `.definePanelTour(x)` returns an data.frame containing the steps of a panel-specific tour.
- `.getDotPlotColorHelp(x, color_choices)` returns a function that generates an **rintrojs** tour for the color choice UI.

### Author(s)

Aaron Lun

### See Also

[RowDataPlot](#), for the base class.

**Examples**

```
# Making up some results:
se <- SummarizedExperiment(matrix(rnorm(10000), 1000, 10))
rownames(se) <- paste0("GENE_", seq_len(nrow(se)))
rowData(se)$PValue <- runif(nrow(se))
rowData(se)$LogFC <- rnorm(nrow(se))
rowData(se)$AveExpr <- rnorm(nrow(se))

if (interactive()) {
  iSEE(se, initial=list(VolcanoPlot()))
}
```

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