

# Package ‘debrowser’

October 16, 2019

**Type** Package

**Title** Interactive Differential Expression Analysis Browser

**Version** 1.12.3

**Date** 2018-11-20

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**Description** Bioinformatics platform containing interactive plots and tables for differential gene and region expression studies. Allows visualizing expression data much more deeply in an interactive and faster way. By changing the parameters, users can easily discover different parts of the data that like never have been done before. Manually creating and looking these plots takes time. With DEBrowser users can prepare plots without writing any code. Differential expression, PCA and clustering analysis are made on site and the results are shown in various plots such as scatter, bar, box, volcano, ma plots and Heatmaps.

**Depends** R (>= 3.5.0),

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**LazyData** true

**Imports** shiny, jsonlite, shinyjs, shinydashboard, shinyBS, gplots, DT, ggplot2, RColorBrewer, annotate, AnnotationDbi, DESeq2, DOSE, igraph, grDevices, graphics, stats, utils, GenomicRanges, IRanges, S4Vectors, SummarizedExperiment, stringi, reshape2, d3heatmap, org.Hs.eg.db, org.Mm.eg.db, limma, edgeR, clusterProfiler, methods, sva, RCurl, enrichplot, colourpicker, plotly, heatmaply, Harman, pathview

**RoxygenNote** 6.1.1

**Encoding** UTF-8

**Suggests** testthat, rmarkdown, knitr, R.rsp

**VignetteBuilder** knitr, R.rsp

**URL** <https://github.com/UMMS-Biocore/debrower>

**BugReports** <https://github.com/UMMS-Biocore/debrower/issues/new>

**biocViews** Sequencing, ChIPSeq, RNASeq, DifferentialExpression, GeneExpression, Clustering, ImmunoOncology

**git\_url** <https://git.bioconductor.org/packages/debrowser>

**git\_branch** RELEASE\_3\_9

**git\_last\_commit** a21fc72

**git\_last\_commit\_date** 2019-08-09

**Date/Publication** 2019-10-15

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`actionButtonDE`*Buttons including Action Buttons and Event Buttons***Description**

Creates an action button whose value is initially zero, and increments by one each time it is pressed.

**Usage**

```
actionButtonDE(inputId, label, styleclass = "", size = "",  
               block = FALSE, icon = NULL, css.class = "", ...)
```

**Arguments**

|                         |   |
|-------------------------|---|
| <code>inputId</code>    | Specifies the input slot that will be used to access the value.   |
| <code>label</code>      | The contents of the button—usually a text label, but you could also use any other HTML, like an image.                |
| <code>styleclass</code> | The Bootstrap styling class of the button—options are primary, info, success, warning, danger, inverse, link or blank |
| <code>size</code>       | The size of the button—options are large, small, mini   |
| <code>block</code>      | Whether the button should fill the block  |
| <code>icon</code>       | Display an icon for the button  |
| <code>css.class</code>  | Any additional CSS class one wishes to add to the action button   |
| <code>...</code>        | Other argument to feed into shiny::actionButton   |

**Examples**

```
actionButtonDE("goDE", "Go to DE Analysis")
```

`addDataCols`*addDataCols***Description**

add additional data columns to de results

**Usage**

```
addDataCols(data = NULL, de_res = NULL, cols = NULL, conds = NULL)
```

**Arguments**

|                      |                |
|----------------------|----------------|
| <code>data,</code>   | loaded dataset |
| <code>de_res,</code> | de results     |
| <code>cols,</code>   | columns        |
| <code>conds,</code>  | inputconds     |

**Value**

data

**Examples**

```
x <- addDataCols()
```

---

addID

*addID*

---

**Description**

Adds an id to the data frame being used.

**Usage**

```
addID(data = NULL)
```

**Arguments**

data, loaded dataset

**Value**

data

**Examples**

```
x <- addID()
```

---

all2all

*all2all*

---

**Description**

Prepares all2all scatter plots for given datasets.

**Usage**

```
all2all(data, cex = 2)
```

**Arguments**

data, data that have the sample names in the header.  
cex text size

**Value**

all2all scatter plots

**Examples**

```
plot<-all2all(mtcars)
```

|                   |                          |
|-------------------|--------------------------|
| all2allControlsUI | <i>all2allControlsUI</i> |
|-------------------|--------------------------|

**Description**

Generates the controls in the left menu for an all2all plot

**Usage**

```
all2allControlsUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

**Value**

returns the controls for left menu

**Note**

```
all2allControlsUI
```

**Examples**

```
x <- all2allControlsUI("bar")
```

|              |                     |
|--------------|---------------------|
| applyFilters | <i>applyFilters</i> |
|--------------|---------------------|

**Description**

Applies filters based on user selected parameters to be displayed within the DEBrowser.

**Usage**

```
applyFilters(filt_data = NULL, cols = NULL, condns = NULL,
            input = NULL)
```

**Arguments**

|            |                     |
|------------|---------------------|
| filt_data, | loaded dataset      |
| cols,      | selected samples    |
| condns,    | selected conditions |
| input,     | input parameters    |

*applyFiltersNew*

9

**Value**

data

**Examples**

```
x <- applyFilters()
```

---

*applyFiltersNew*

*applyFiltersNew*

---

**Description**

Apply filters based on foldChange cutoff and padj value. This function adds a "Legend" column with "Up", "Down" or "NS" values for visualization.

**Usage**

```
applyFiltersNew(data = NULL, input = NULL)
```

**Arguments**

|        |                  |
|--------|------------------|
| data,  | loaded dataset   |
| input, | input parameters |

**Value**

data

**Examples**

```
x <- applyFiltersNew()
```

---

*applyFiltersToMergedComparison*

*applyFiltersToMergedComparison*

---

**Description**

Gathers the merged comparison data to be used within the DEBrowser.

**Usage**

```
applyFiltersToMergedComparison(merged = NULL, nc = NULL,
                               input = NULL)
```

**Arguments**

|         |                           |
|---------|---------------------------|
| merged, | merged data               |
| nc,     | the number of comparisons |
| input,  | input params              |

**Value**

data

**Examples**

```
x <- applyFiltersToMergedComparison()
```

**barMainPlotControlsUI** *barMainPlotControlsUI*

**Description**

Generates the controls in the left menu for a bar main plot

**Usage**

```
barMainPlotControlsUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

**Value**

returns the controls for left menu

**Note**

```
barMainPlotControlsUI
```

**Examples**

```
x <- barMainPlotControlsUI("bar")
```

---

|               |  |
|---------------|--|
| batchEffectUI | <i>batchEffectUI</i> Creates a panel to coorect batch effect |
|---------------|--|

---

**Description**

batchEffectUI Creates a panel to coorect batch effect

**Usage**

```
batchEffectUI(id)
```

**Arguments**

id, namespace id

**Value**

panel

**Examples**

```
x <- batchEffectUI("batcheffect")
```

---

---

|             |                    |
|-------------|--------------------|
| batchMethod | <i>batchMethod</i> |
|-------------|--------------------|

---

**Description**

select batch effect method

**Usage**

```
batchMethod(id)
```

**Arguments**

id, namespace id

**Value**

radio control

**Note**

```
batchMethod
```

**Examples**

```
x <- batchMethod("batch")
```

`BoxMainPlotControlsUI` *BoxMainPlotControlsUI*

### Description

Generates the controls in the left menu for a Box main plot

### Usage

```
BoxMainPlotControlsUI(id)
```

### Arguments

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

### Value

returns the controls for left menu

### Note

```
BoxMainPlotControlsUI
```

### Examples

```
x <- BoxMainPlotControlsUI("box")
```

`changeClusterOrder` *changeClusterOrder*

### Description

change order of K-means clusters

### Usage

```
changeClusterOrder(order = NULL, cld = NULL)
```

### Arguments

|         |       |
|---------|-------|
| order , | order |
| cld ,   | data  |

### Value

heatmap plot area

### Note

```
changeClusterOrder
```

**Examples**

```
x <- changeClusterOrder()
```

---

|                |                       |
|----------------|-----------------------|
| checkCountData | <i>checkCountData</i> |
|----------------|-----------------------|

---

**Description**

Returns if there is a problem in the count data.

**Usage**

```
checkCountData(input = NULL)
```

**Arguments**

|        |        |
|--------|--------|
| input, | inputs |
|--------|--------|

**Value**

error if there is a problem about the loaded data

**Note**

```
checkCountData
```

**Examples**

```
x <- checkCountData()
```

---

|               |                      |
|---------------|----------------------|
| checkMetaData | <i>checkMetaData</i> |
|---------------|----------------------|

---

**Description**

Returns if there is a problem in the count data.

**Usage**

```
checkMetaData(input = NULL, counttable = NULL)
```

**Arguments**

|             |            |
|-------------|------------|
| input,      | input      |
| counttable, | counttable |

**Value**

error if there is a problem about the loaded data

**Note**

```
checkMetaData
```

**Examples**

```
x <- checkMetaData()
```

---

**clusterData**

---

*clusterData*

---

**Description**

Gathers the Cluster analysis data to be used within the GO Term plots.

**Usage**

```
clusterData(dat = NULL)
```

**Arguments**

|      |                     |
|------|---------------------|
| dat, | the data to cluster |
|------|---------------------|

**Value**

clustered data

**Note**

```
clusterData
```

**Examples**

```
mycluster <- clusterData()
```

---

**clustFunParamsUI**

---

*clustFunParamsUI*

---

**Description**

get cluster function parameter control

**Usage**

```
clustFunParamsUI()
```

**Value**

cluster params

**Note**

```
clustFunParamsUI
```

**Examples**

```
x <- clustFunParamsUI()
```

---

```
compareClust
```

```
compareClust
```

---

**Description**

C.compares the clustered data to be displayed within the GO Term plots.

**Usage**

```
compareClust(dat = NULL, ont = "CC", org = "org.Hs.eg.db",
             fun = "enrichGO", title = "Ontology Distribution Comparison",
             pvalueCutoff = 0.01)
```

**Arguments**

|               |                          |
|---------------|--------------------------|
| dat,          | data to compare clusters |
| ont,          | the ontology to use      |
| org,          | the organism used        |
| fun,          | fun                      |
| title,        | title of the comparison  |
| pvalueCutoff, | pvalueCutoff             |

**Value**

compared cluster

**Note**

```
compareClust
```

**Examples**

```
x <- compareClust()
```

`condSelectUI`

*condSelectUI Creates a panel to select samples for each condition*

**Description**

`condSelectUI` Creates a panel to select samples for each condition

**Usage**

```
condSelectUI()
```

**Value**

panel

**Examples**

```
x <- condSelectUI()
```

`correctCombat`

*Correct Batch Effect using Combat in sva package*

**Description**

Batch effect correction

**Usage**

```
correctCombat(input = NULL, idata = NULL, metadata = NULL)
```

**Arguments**

|                         |              |
|-------------------------|--------------|
| <code>input</code> ,    | input values |
| <code>idata</code> ,    | data         |
| <code>metadata</code> , | metadata     |

**Value**

data

**Examples**

```
x<-correctCombat ()
```

---

|               |  |
|---------------|--|
| correctHarman | <i>Correct Batch Effect using Harman</i> |
|---------------|--|

---

**Description**

Batch effect correction

**Usage**

```
correctHarman(input = NULL, idata = NULL, metadata = NULL)
```

**Arguments**

|           |              |
|-----------|--------------|
| input,    | input values |
| idata,    | data         |
| metadata, | metadata     |

**Value**

data

**Examples**

```
x<-correctHarman ()
```

---

---

|                |                       |
|----------------|-----------------------|
| customColorsUI | <i>customColorsUI</i> |
|----------------|-----------------------|

---

**Description**

get Custom Color controls

**Usage**

```
customColorsUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace ID |
|-----|--------------|

**Value**

color range

**Note**

```
getColRng
```

**Examples**

```
x <- customColorsUI("heatmap")
```

---

cutOffSelectionUI      *cutOffSelectionUI*

---

**Description**

Gathers the cut off selection for DE analysis

**Usage**

cutOffSelectionUI(id)

**Arguments**

id,                namespace id

**Value**

returns the left menu according to the selected tab;

**Note**

cutOffSelectionUI

**Examples**

x <- cutOffSelectionUI("cutoff")

---

dataLCFUI      *dataLCFUI Creates a panel to filter low count genes and regions*

---

**Description**

dataLCFUI Creates a panel to filter low count genes and regions

**Usage**

dataLCFUI(id)

**Arguments**

id,                namespace id

**Value**

panel

**Examples**

x <- dataLCFUI("lcf")

---

dataLoadUI                  *dataLoadUI*

---

**Description**

Creates a panel to upload the data

**Usage**

```
dataLoadUI(id)
```

**Arguments**

id,                        namespace id

**Value**

panel

**Examples**

```
x <- dataLoadUI("load")
```

---

---

debrowserall2all                  *debrowserall2all*

---

**Description**

Module for a bar plot that can be used in data prep, main plots low count removal modules or any desired module

**Usage**

```
debrowserall2all(input, output, session, data = NULL, cex = 2)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |
| cex,     | the size of the dots                     |

**Value**

all2all plot

**Examples**

```
x <- debrowserall2all()
```

**debrowserbarmainplot** *debrowserbarmainplot*

### Description

Module for a bar plot that can be used in data prep, main plots low count removal modules or any desired module

### Usage

```
debrowserbarmainplot(input, output, session, data = NULL, cols = NULL,
                     conds = NULL, key = NULL)
```

### Arguments

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |
| cols,    | columns                                  |
| conds,   | conditions                               |
| key,     | the gene or region name                  |

### Value

density plot

### Examples

```
x <- debrowserbarmainplot()
```

**debrowserbatcheffect** *debrowserbatcheffect*

### Description

Module to correct batch effect

### Usage

```
debrowserbatcheffect(input, output, session, ldata = NULL)
```

### Arguments

|          |                 |
|----------|-----------------|
| input,   | input variables |
| output,  | output objects  |
| session, | session         |
| ldata,   | loaded data     |

**Value**

main plot  
panel

**Examples**

```
x <- debrowserbatcheffect()
```

---

```
debrowserboxmainplot    debrowserboxmainplot
```

---

**Description**

Module for a box plot that can be used in DEanalysis main part and used heatmaps

**Usage**

```
debrowserboxmainplot(input = NULL, output = NULL, session = NULL,  
                      data = NULL, cols = NULL, cond = NULL, key = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |
| cols,    | columns                                  |
| cond,    | conditions                               |
| key,     | the gene or region name                  |

**Value**

density plot

**Examples**

```
x <- debrowserboxmainplot()
```

**debrowsercondselect**      *debrowsercondselect*

### Description

Condition selection This is not a module. Module construction didn't used here, just use it as functions not in a module.

### Usage

```
debrowsercondselect(input = NULL, output = NULL, session = NULL,
                     data = NULL, metadata = NULL)
```

### Arguments

|           |                 |
|-----------|-----------------|
| input,    | input variables |
| output,   | output objects  |
| session,  | session         |
| data,     | count data      |
| metadata, | metadata        |

### Value

|           |
|-----------|
| main plot |
| panel     |

### Examples

```
x <- debrowsercondselect()
```

**debrowserdataload**      *debrowserdataload*

### Description

Module to load count data and metadata

### Usage

```
debrowserdataload(input = NULL, output = NULL, session = NULL,
                   nextpagebutton = NULL)
```

### Arguments

|                 |   |
|-----------------|---|
| input,          | input variables   |
| output,         | output objects  |
| session,        | session   |
| nextpagebutton, | the name of the next page button after loading the data |

**Value**

```
main plot  
panel
```

**Examples**

```
x <- debrowserdataload()
```

---

```
debrowserdeanalysis      debrowserdeanalysis
```

---

**Description**

Module to perform and visualize DE results.

**Usage**

```
debrowserdeanalysis(input = NULL, output = NULL, session = NULL,  
                     data = NULL, columns = NULL, cond = NULL, params = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |
| columns, | columns                                  |
| cond,    | conditions                               |
| params,  | de parameters                            |

**Value**

DE panel

**Examples**

```
x <- debrowserdeanalysis()
```

**debrowserdensityplot** *debrowserdensityplot*

### Description

Module for a density plot that can be used in data prep and low count removal modules

### Usage

```
debrowserdensityplot(input = NULL, output = NULL, session = NULL,
                      data = NULL)
```

### Arguments

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |

### Value

density plot

### Examples

```
x <- debrowserdensityplot()
```

**debrowserheatmap** *debrowserheatmap*

### Description

Heatmap module to create interactive heatmaps and get selected list from a heatmap

### Usage

```
debrowserheatmap(input, output, session, expdata = NULL)
```

### Arguments

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| expdata, | a matrix that includes expression values |

### Value

heatmapply plot

**Examples**

```
x <- debrowserheatmap()
```

---

debrowserhistogram      *debrowserhistogram*

---

**Description**

Module for a histogram that can be used in data prep and low count removal modules

**Usage**

```
debrowserhistogram(input = NULL, output = NULL, session = NULL,  
                   data = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |

**Value**

histogram

**Examples**

```
x <- debrowserhistogram()
```

---

debrowserIQRplot      *debrowserIQRplot*

---

**Description**

Module for an IQR plot that can be used in data prep and low count removal modules

**Usage**

```
debrowserIQRplot(input = NULL, output = NULL, session = NULL,  
                   data = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |

**Value**

IQR

**Examples**

```
x <- debrowserIQRplot()
```

---

```
debrowserlowcountfilter  
debrowserlowcountfilter
```

---

**Description**

Module to filter low count genes/regions

**Usage**

```
debrowserlowcountfilter(input = NULL, output = NULL, session = NULL,  
ldata = NULL)
```

**Arguments**

|          |                 |
|----------|-----------------|
| input,   | input variables |
| output,  | output objects  |
| session, | session         |
| ldata,   | loaded data     |

**Value**

main plot

panel

**Examples**

```
x <- debrowserlowcountfilter()
```

---

```
debrowsermainplot      debrowsermainplot
```

---

**Description**

Module for a scatter, volcano and ma plots that are going to be used as a mainplot in debrowser

**Usage**

```
debrowsermainplot(input = NULL, output = NULL, session = NULL,  
                  data = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| output,  | output objects                           |
| session, | session                                  |
| data,    | a matrix that includes expression values |

**Value**

|           |
|-----------|
| main plot |
| panel     |

**Examples**

```
x <- debrowsermainplot()
```

---

```
debrowserpcaplot      debrowserpcaplot
```

---

**Description**

Module for a pca plot with its loadings as a mainplot in debrowser

**Usage**

```
debrowserpcaplot(input = NULL, output = NULL, session = NULL,  
                  pcadata = NULL, metadata = NULL)
```

**Arguments**

|           |  |
|-----------|--|
| input,    | input variables                          |
| output,   | output objects                           |
| session,  | session                                  |
| pcadata,  | a matrix that includes expression values |
| metadata, | metadata to color the plots              |

**Value**

main plot  
panel

**Examples**

```
x <- debrowserpcaplot()
```

---

dendControlsUI                  *dendControlsUI*

---

**Description**

get distance metric parameters

**Usage**

```
dendControlsUI(id, dendtype = "Row")
```

**Arguments**

id,                    module ID  
dendtype,            Row or Col

**Value**

controls

**Note**

```
dendControlsUI
```

**Examples**

```
x <- dendControlsUI("heatmap")
```

---

```
densityPlotControlsUI densityPlotControlsUI
```

---

**Description**

Generates the controls in the left menu for a densityPlot

**Usage**

```
densityPlotControlsUI(id)
```

**Arguments**

id, namespace id

**Value**

returns the left menu

**Note**

```
densityPlotControlsUI
```

**Examples**

```
x <- densityPlotControlsUI("density")
```

---

---

```
deServer deServer
```

---

**Description**

Sets up shinyServer to be able to run DEBrowser interactively.

**Usage**

```
deServer(input, output, session)
```

**Arguments**

input, input params from UI  
output, output params to UI  
session, session variable

**Value**

the panel for main plots;

**Note**

```
deServer
```

**Examples**

```
deServer
```

---

deUI

*deUI*

---

**Description**

Creates a shinyUI to be able to run DEBrowser interactively.

**Usage**

```
deUI()
```

**Value**

the panel for main plots;

**Note**

```
deUI
```

**Examples**

```
x<-deUI()
```

---

*distFunParamsUI*

*distFunParamsUI*

---

**Description**

get distance metric parameters

**Usage**

```
distFunParamsUI()
```

**Value**

funParams

**Note**

```
distFunParamsUI
```

**Examples**

```
x <- distFunParamsUI()
```

---

`drawKEGG`*drawKEGG***Description**

draw KEGG patwhay with expression values

**Usage**

```
drawKEGG(input = NULL, dat = NULL, pid = NULL)
```

**Arguments**

|        |                   |
|--------|-------------------|
| input, | input             |
| dat,   | expression matrix |
| pid,   | pathway id        |

**Value**

enriched DO

**Note**

drawKEGG

**Examples**

```
x <- drawKEGG()
```

---

`drawPCAExplained`

*Creates a more detailed plot using the PCA results from the selected dataset.*

---

**Description**

Creates a more detailed plot using the PCA results from the selected dataset.

**Usage**

```
drawPCAExplained(explainedData = NULL)
```

**Arguments**

|                |               |
|----------------|---------------|
| explainedData, |               |
|                | selected data |

**Value**

explained plot

**Examples**

```
x <- drawPCAExplained()
```

**fileTypes**

*fileTypes*

**Description**

Returns fileTypes that are going to be used in creating fileUpload UI

**Usage**

```
fileTypes()
```

**Value**

file types

**Note**

```
fileTypes
```

**Examples**

```
x <- fileTypes()
```

**fileUploadBox**

*fileUploadBox*

**Description**

File upload module

**Usage**

```
fileUploadBox(id = NULL, inputId = NULL, label = NULL)
```

**Arguments**

|                       |               |
|-----------------------|---------------|
| <code>id,</code>      | namespace id  |
| <code>inputId,</code> | input file ID |
| <code>label,</code>   | label         |

**Value**

radio control

**Note**

```
fileUploadBox
```

**Examples**

```
x <- fileUploadBox("meta", "metadata", "Metadata")
```

---

```
generateTestData      generateTestData
```

---

**Description**

This generates a test data that is suitable to main plots in debrowser

**Usage**

```
generateTestData(dat = NULL)
```

**Arguments**

dat, DESeq results will be generated for loaded data

**Value**

testData

**Examples**

```
x <- generateTestData()
```

---

```
getAfterLoadMsg      getAfterLoadMsg
```

---

**Description**

Generates and displays the message to be shown after loading data within the DEBrowser.

**Usage**

```
getAfterLoadMsg()
```

**Value**

return After Load Msg

**Note**

```
getAfterLoadMsg
```

**Examples**

```
x <- getAfterLoadMsg()
```

|                               |                         |
|-------------------------------|-------------------------|
| <code>getAll2AllPlotUI</code> | <i>getAll2AllPlotUI</i> |
|-------------------------------|-------------------------|

**Description**

all2all plots UI.

**Usage**

```
getAll2AllPlotUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

**Value**

the panel for all2all plots;

**Note**

```
getAll2AllPlotUI
```

**Examples**

```
x <- getAll2AllPlotUI("bar")
```

|                             |                       |
|-----------------------------|-----------------------|
| <code>getBarMainPlot</code> | <i>getBarMainPlot</i> |
|-----------------------------|-----------------------|

**Description**

Makes Density plots

**Usage**

```
getBarMainPlot(data = NULL, cols = NULL, conds = NULL, key = NULL,
               title = "", input = NULL)
```

**Arguments**

|        |                          |
|--------|--------------------------|
| data,  | count or normalized data |
| cols,  | cols                     |
| conds, | conds                    |
| key,   | key                      |
| title, | title                    |
| input, | input                    |

**Examples**

```
getBarMainPlot()
```

---

```
getBarMainPlotUI      getBarMainPlotUI
```

---

**Description**

main bar plots UI.

**Usage**

```
getBarMainPlotUI(id)
```

**Arguments**

`id,`                  namespace id

**Value**

the panel for Density plots;

**Note**

```
getBarMainPlotUI
```

**Examples**

```
x <- getBarMainPlotUI("bar")
```

---

```
getBoxMainPlot      getBoxMainPlot
```

---

**Description**

Makes Density plots

**Usage**

```
getBoxMainPlot(data = NULL, cols = NULL, cond = NULL, key = NULL,  
              title = "", input = NULL)
```

**Arguments**

|                     |                          |
|---------------------|--------------------------|
| <code>data,</code>  | count or normalized data |
| <code>cols,</code>  | cols                     |
| <code>cond,</code>  | conds                    |
| <code>key,</code>   | key                      |
| <code>title,</code> | title                    |
| <code>input,</code> | input                    |

**Examples**


---

```
getBoxMainPlot()
```

---

|                  |                         |
|------------------|-------------------------|
| getBoxMainPlotUI | <i>getBoxMainPlotUI</i> |
|------------------|-------------------------|

---

**Description**

main Box plots UI.

**Usage**

```
getBoxMainPlotUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

**Value**

the panel for Density plots;

**Note**

```
getBoxMainPlotUI
```

**Examples**

```
x <- getBoxMainPlotUI("box")
```

---

|             |  |
|-------------|--|
| getBTableUI | <i>getBTableUI prepares a Modal to put a table</i> |
|-------------|--|

---

**Description**

getBTableUI prepares a Modal to put a table

**Usage**

```
getBTableUI(name = NULL, label = NULL, trigger = NULL,
           size = "large", modal = NULL)
```

**Arguments**

|          |                              |
|----------|------------------------------|
| name,    | name                         |
| label,   | label                        |
| trigger, | trigger button for the modal |
| size,    | size of the modal            |
| modal,   | modal yes/no                 |

**Value**

the modal

**Examples**

```
x<- getBTableUI()
```

---

getColors

*getColors*

---

**Description**

get colors for the domains

**Usage**

```
getColors(domains = NULL)
```

**Arguments**

domains,            domains to be colored

**Value**

colors

**Examples**

```
x<-getColors()
```

---

getColorShapeSelection

*getColorShapeSelection*

---

**Description**

Generates the fill and shape selection boxes for PCA plots. metadata file has to be loaded in this case

**Usage**

```
getColorShapeSelection(metadata = NULL, input = NULL, session = NULL)
```

**Arguments**

metadata,        metadata table  
input,            input  
session,         session

**Value**

Color and shape selection boxes

**Examples**

```
x <- getColorShapeSelection()
```

|                               |                         |
|-------------------------------|-------------------------|
| <code>getCompSelection</code> | <i>getCompSelection</i> |
|-------------------------------|-------------------------|

**Description**

Gathers the user selected comparison set to be used within the DEBrowser.

**Usage**

```
getCompSelection(name = NULL, count = NULL)
```

**Arguments**

|        |                             |
|--------|-----------------------------|
| name,  | the name of the selectInput |
| count, | comparison count            |

**Note**

```
getCompSelection
```

**Examples**

```
x <- getCompSelection(name="comp", count = 2)
```

|                                   |                             |
|-----------------------------------|-----------------------------|
| <code>getConditionSelector</code> | <i>getConditionSelector</i> |
|-----------------------------------|-----------------------------|

**Description**

Selects user input conditions to run in DESeq.

**Usage**

```
getConditionSelector(num = 0, choices = NULL, selected = NULL)
```

**Arguments**

|           |                                 |
|-----------|---------------------------------|
| num,      | panel that is going to be shown |
| choices,  | sample list                     |
| selected, | selected smaple list            |

**Examples**

```
x <- getConditionSelector()
```

---

```
getConditionSelectorFromMeta  
    getConditionSelectorFromMeta
```

---

## Description

Selects user input conditions to run in DESeq from metadata

## Usage

```
getConditionSelectorFromMeta(metadata = NULL, input = NULL,  
    index = 1, num = 0, choices = NULL, selected = NULL)
```

## Arguments

|           |                 |
|-----------|-----------------|
| metadata, | meta data table |
| input,    | input           |
| index,    | index           |
| num,      | num             |
| choices,  | choices         |
| selected, | selected        |

## Examples

```
x <- getConditionSelectorFromMeta()
```

---

```
getCondMsg          getCondMsg
```

---

## Description

Generates and displays the current conditions and their samples within the DEBrowser.

## Usage

```
getCondMsg(dc = NULL, input = NULL, cols = NULL, cond = NULL)
```

## Arguments

|        |                     |
|--------|---------------------|
| dc,    | columns             |
| input, | selected comparison |
| cols,  | columns             |
| cond,  | selected conditions |

## Value

return conditions

**Note**

```
getCondMsg
```

**Examples**

```
x <- getCondMsg()
```

|                           |                           |
|---------------------------|---------------------------|
| <i>getCutOffSelection</i> | <i>getCutOffSelection</i> |
|---------------------------|---------------------------|

**Description**

Gathers the cut off selection for DE analysis

**Usage**

```
getCutOffSelection(nc = 1)
```

**Arguments**

|     |                             |
|-----|-----------------------------|
| nc, | total number of comparisons |
|-----|-----------------------------|

**Value**

returns the left menu according to the selected tab;

**Note**

```
getCutOffSelection
```

**Examples**

```
x <- getCutOffSelection()
```

|                             |  |
|-----------------------------|--|
| <i>getDataAssesmentText</i> | <i>getDataAssesmentText</i> DataAssesment text |
|-----------------------------|--|

**Description**

*getDataAssesmentText* DataAssesment text

**Usage**

```
getDataAssesmentText()
```

**Value**

help text for data assesment

**Examples**

```
x<- getDataAssesmentText()
```

---

```
getDataForTables      getDataForTables get data to fill up tables tab
```

---

**Description**

getDataForTables get data to fill up tables tab

**Usage**

```
getDataForTables(input = NULL, init_data = NULL, filt_data = NULL,  
                 selected = NULL, getMostVaried = NULL, mergedComp = NULL,  
                 explainedData = NULL)
```

**Arguments**

|                |                       |
|----------------|-----------------------|
| input,         | input parameters      |
| init_data,     | initial dataset       |
| filt_data,     | filt_data             |
| selected,      | selected genes        |
| getMostVaried, | most varied genes     |
| mergedComp,    | merged comparison set |
| explainedData, | pca gene set          |

**Value**

data

**Examples**

```
x <- getDataForTables()
```

---

```
getDataPreparationText  
      getDataPreparationText DataPreparation text
```

---

**Description**

getDataPreparationText DataPreparation text

**Usage**

```
getDataPreparationText()
```

**Value**

help text for data preparation

**Examples**

```
x<- getDataPreparationText()
```

|                                |  |
|--------------------------------|--|
| <code>getDEAnalysisText</code> | <i>getDEAnalysisText DEAnalysis text</i> |
|--------------------------------|--|

**Description**

`getDEAnalysisText DEAnalysis text`

**Usage**

```
getDEAnalysisText()
```

**Value**

help text for DE Analysis

**Examples**

```
x<- getDEAnalysisText()
```

|                             |                       |
|-----------------------------|-----------------------|
| <code>getDensityPlot</code> | <i>getDensityPlot</i> |
|-----------------------------|-----------------------|

**Description**

Makes Density plots

**Usage**

```
getDensityPlot(data = NULL, input = NULL, title = "")
```

**Arguments**

|                     |                          |
|---------------------|--------------------------|
| <code>data,</code>  | count or normalized data |
| <code>input,</code> | input                    |
| <code>title,</code> | title                    |

**Examples**

```
getDensityPlot()
```

---

getDensityPlotUI      *getDensityPlotUI*

---

**Description**

Density plot UI.

**Usage**

```
getDensityPlotUI(id)
```

**Arguments**

id,                namespace id

**Value**

the panel for Density plots;

**Note**

```
getDensityPlotUI
```

**Examples**

```
x <- getDensityPlotUI("density")
```

---

getDEResultsUI      *getDEResultsUI* Creates a panel to visualize DE results

---

**Description**

getDEResultsUI Creates a panel to visualize DE results

**Usage**

```
getDEResultsUI(id)
```

**Arguments**

id,                namespace id

**Value**

panel

**Examples**

```
x <- getDEResultsUI("batcheffect")
```

---

|            |                   |
|------------|-------------------|
| getDomains | <i>getDomains</i> |
|------------|-------------------|

---

**Description**

Get domains for the main plots.

**Usage**

```
getDomains(filt_data = NULL)
```

**Arguments**

`filt_data`, data to get the domains

**Value**

domains

**Examples**

```
x<-getDomains()
```

---

|         |  |
|---------|--|
| getDown | <i>getDown</i> get down regulated data |
|---------|--|

---

**Description**

getDown get down regulated data

**Usage**

```
getDown(filt_data = NULL)
```

**Arguments**

`filt_data`, `filt_data`

**Value**

data

**Examples**

```
x <- getDown()
```

---

```
getDownloadSection      getDownloadSection
```

---

**Description**

download section button and dataset selection box in the menu for user to download selected data.

**Usage**

```
getDownloadSection(choices = NULL)
```

**Arguments**

choices, main vs. QC section

**Value**

the panel for download section in the menu;

**Note**

```
getDownloadSection
```

**Examples**

```
x<- getDownloadSection()
```

---

```
getEnrichDO      getEnrichDO
```

---

**Description**

Gathers the Enriched DO Term analysis data to be used within the GO Term plots.

**Usage**

```
getEnrichDO(genelist = NULL, pvalueCutoff = 0.01)
```

**Arguments**

genelist, gene list  
pvalueCutoff, the p value cutoff

**Value**

enriched DO

**Note**

```
getEnrichDO
```

**Examples**

```
x <- getEnrichDO()
```

**getEnrichGO**

*getEnrichGO*

**Description**

Gathers the Enriched GO Term analysis data to be used within the GO Term plots.

**Usage**

```
getEnrichGO(genelist = NULL, pvalueCutoff = 0.01,
            org = "org.Hs.eg.db", ont = "CC")
```

**Arguments**

|               |                   |
|---------------|-------------------|
| genelist,     | gene list         |
| pvalueCutoff, | p value cutoff    |
| org,          | the organism used |
| ont,          | the ontology used |

**Value**

Enriched GO

**Note**

`getEnrichGO`

**Examples**

```
x <- getEnrichGO()
```

**getEnrichKEGG**

*getEnrichKEGG*

**Description**

Gathers the Enriched KEGG analysis data to be used within the GO Term plots.

**Usage**

```
getEnrichKEGG(genelist = NULL, pvalueCutoff = 0.01,
              org = "org.Hs.eg.db")
```

**Arguments**

|               |                    |
|---------------|--------------------|
| genelist,     | gene list          |
| pvalueCutoff, | the p value cutoff |
| org,          | the organism used  |

**Value**

Enriched KEGG

**Note**

`getEnrichKEGG`

**Examples**

```
x <- getEnrichKEGG()
```

---

getEntrezIds

*getEntrezIds*

---

**Description**

Gathers the gene list to use for GOTerm analysis.

**Usage**

```
getEntrezIds(genes = NULL, org = "org.Hs.eg.db")
```

**Arguments**

|        |   |
|--------|---|
| genes, | gene list with fold changes                   |
| org,   | organism for gene symbol entrez ID conversion |

**Value**

ENTREZ ID list

**Note**

GOTerm

`getEntrezIds` symbol to ENTREZ ID conversion

**Examples**

```
x <- getEntrezIds()
```

`getEntrezTable`      *getEntrezTable*

### Description

Gathers the entrezIds of the genes in given list and their data

### Usage

```
getEntrezTable(genes = NULL, dat = NULL, org = "org.Hs.eg.db")
```

### Arguments

|        |   |
|--------|---|
| genes, | gene list                                     |
| dat,   | data matrix                                   |
| org,   | organism for gene symbol entrez ID conversion |

### Value

table with the entrez IDs in the rownames

### Note

GOTerm  
getEntrezTable symbol to ENTREZ ID conversion

### Examples

```
x <- getEntrezTable()
```

`getGeneList`      *getGeneList*

### Description

Gathers the gene list to use for GOTerm analysis.

### Usage

```
getGeneList(genes = NULL, org = "org.Hs.eg.db")
```

### Arguments

|        |   |
|--------|---|
| genes, | gene list                                     |
| org,   | organism for gene symbol entrez ID conversion |

### Value

ENTREZ ID list

**Note**

GOTerm  
getGeneList symbol to ENTREZ ID conversion

**Examples**

```
x <- getGeneList(c('OCLN', 'ABCC2'))
```

---

|                |                       |
|----------------|-----------------------|
| getGeneSetData | <i>getGeneSetData</i> |
|----------------|-----------------------|

---

**Description**

Gathers the specified gene set list to be used within the DEBrowser.

**Usage**

```
getGeneSetData(data = NULL, geneset = NULL)
```

**Arguments**

|          |                |
|----------|----------------|
| data,    | loaded dataset |
| geneset, | given gene set |

**Value**

data

**Examples**

```
x <- getGeneSetData()
```

---

|               |                      |
|---------------|----------------------|
| getGOLeftMenu | <i>getGOLeftMenu</i> |
|---------------|----------------------|

---

**Description**

Generates the GO Left menu to be displayed within the DEBrowser.

**Usage**

```
getGOLeftMenu()
```

**Value**

returns the left menu according to the selected tab;

**Note**

```
getGOLeftMenu
```

**Examples**

```
x <- getGOLeftMenu()
```

getGoPanel

*getGoPanel*

**Description**

Creates go term analysis panel within the shiny display.

**Usage**

```
getGoPanel()
```

**Value**

the panel for go term analysis;

**Note**

```
getGoPanel
```

**Examples**

```
x <- getGoPanel()
```

getGOPlots

*getGOPlots*

**Description**

Go term analysis panel. Generates appropriate GO plot based on user selection.

**Usage**

```
getGOPlots(dataset = NULL, input = NULL)
```

**Arguments**

|          |                  |
|----------|------------------|
| dataset, | the dataset used |
| input ,  | input params     |

**Value**

the panel for go plots;

**Note**

```
getGOPlots
```

**Examples**

```
x<- getGOPlots()
```

---

|              |                     |
|--------------|---------------------|
| getHeatmapUI | <i>getHeatmapUI</i> |
|--------------|---------------------|

---

**Description**

Generates the left menu to be used for heatmap plots

**Usage**

```
getHeatmapUI(id)
```

**Arguments**

id,                   module ID

**Value**

heatmap plot area

**Note**

```
getHeatmapUI
```

**Examples**

```
x <- getHeatmapUI("heatmap")
```

---

|               |  |
|---------------|--|
| getHelpButton | <i>getHelpButton prepares a helpbutton for to go to a specific site in the documentation</i> |
|---------------|--|

---

**Description**

getHelpButton prepares a helpbutton for to go to a specific site in the documentation

**Usage**

```
getHelpButton(name = NULL, link = NULL)
```

**Arguments**

name,               name that are going to come after info  
link,               link of the help

**Value**

the info button

**Examples**

```
x<- getHelpButton()
```

|                                 |                           |
|---------------------------------|---------------------------|
| <code>getHideLegendOnOff</code> | <i>getHideLegendOnOff</i> |
|---------------------------------|---------------------------|

**Description**

hide legend

**Usage**

```
getHideLegendOnOff(id = "pca")
```

**Arguments**

`id`, namespace id

**Examples**

```
x <- getHideLegendOnOff("pca")
```

|                             |                       |
|-----------------------------|-----------------------|
| <code>getHistogramUI</code> | <i>getHistogramUI</i> |
|-----------------------------|-----------------------|

**Description**

Histogram plots UI.

**Usage**

```
getHistogramUI(id)
```

**Arguments**

`id`, namespace id

**Value**

the panel for PCA plots;

**Note**

```
getHistogramUI
```

**Examples**

```
x <- getHistogramUI("histogram")
```

---

|              |                                |
|--------------|--------------------------------|
| getIntroText | <i>getIntroText Intro text</i> |
|--------------|--------------------------------|

---

**Description**

getIntroText Intro text

**Usage**

getIntroText()

**Value**

the JS for tab updates

**Examples**

x<- getIntroText()

---

|            |                   |
|------------|-------------------|
| getIQRPlot | <i>getIQRPlot</i> |
|------------|-------------------|

---

**Description**

Makes IQR boxplot plot

**Usage**

getIQRPlot(data = NULL, input = NULL, title = "")

**Arguments**

|        |                          |
|--------|--------------------------|
| data,  | count or normalized data |
| input, | input                    |
| title, | title                    |

**Examples**

getIQRPlot()

---

|              |                     |
|--------------|---------------------|
| getIQRPlotUI | <i>getIQRPlotUI</i> |
|--------------|---------------------|

---

**Description**

IQR plot UI.

**Usage**

```
getIQRPlotUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

**Value**

the panel for IQR plots;

**Note**

```
getIQRPlotUI
```

**Examples**

```
x <- getIQRPlotUI("IQR")
```

---

|           |                  |
|-----------|------------------|
| getJSLine | <i>getJSLine</i> |
|-----------|------------------|

---

**Description**

heatmap JS code for selection functionality

**Usage**

```
getJSLine()
```

**Value**

JS Code

**Examples**

```
x <- getJSLine()
```

---

`getKEGGModal`

*getKEGGModal prepares a modal for KEGG plots*

---

### Description

`getKEGGModal` prepares a modal for KEGG plots

`getKEGGModal` prepares a helpbutton for to go to a specific site in the documentation

### Usage

```
getKEGGModal()
```

```
getKEGGModal()
```

### Value

the info button

the info button

### Examples

```
x<- getKEGGModal()
```

```
x<- getKEGGModal()
```

---

`getLeftMenu`

*getLeftMenu*

---

### Description

Generates the left menu for for plots within the DEBrowser.

### Usage

```
getLeftMenu(input = NULL)
```

### Arguments

`input`,                input values

### Value

returns the left menu according to the selected tab;

### Note

```
getLeftMenu
```

### Examples

```
x <- getLeftMenu()
```

---

getLegendColors      *getLegendColors*

---

**Description**

Generates colors according to the data

**Usage**

```
getLegendColors(Legend = c("up", "down", "NS"))
```

**Arguments**

Legend,      unique Legends

**Value**

mainPlotControls

**Note**

getLegendColors

**Examples**

```
x <- getLegendColors(c("up", "down", "GS", "NS"))
```

---

getLegendRadio      *getLegendRadio*

---

**Description**

Radio buttons for the types in the legend

**Usage**

```
getLegendRadio(id)
```

**Arguments**

id,      namespace id

**Value**

radio control

**Note**

getLegendRadio

**Examples**

```
x <- getLegendRadio("deprog")
```

---

|                 |                        |
|-----------------|------------------------|
| getLegendSelect | <i>getLegendSelect</i> |
|-----------------|------------------------|

---

**Description**

select legend

**Usage**

```
getLegendSelect(id = "pca")
```

**Arguments**

id, namespace id

**Note**

```
getLegendSelect
```

**Examples**

```
x <- getLegendSelect("pca")
```

---

|               |                      |
|---------------|----------------------|
| getLevelOrder | <i>getLevelOrder</i> |
|---------------|----------------------|

---

**Description**

Generates the order of the overlapping points

**Usage**

```
getLevelOrder(Level = c("up", "down", "NS"))
```

**Arguments**

Level, factor levels shown in the legend

**Value**

order

**Note**

```
getLevelOrder
```

**Examples**

```
x <- getLevelOrder(c("up", "down", "GS", "NS"))
```

`getLoadingMsg`

*getLoadingMsg*

**Description**

Creates and displays the loading message/gif to be displayed within the DEBrowser.

**Usage**

```
getLoadingMsg(output = NULL)
```

**Arguments**

|                       |                |
|-----------------------|----------------|
| <code>output</code> , | output message |
|-----------------------|----------------|

**Value**

loading msg

**Note**

```
getLoadingMsg
```

**Examples**

```
x <- getLoadingMsg()
```

`getLogo`

*getLogo*

**Description**

Generates and displays the logo to be shown within DEBrowser.

**Usage**

```
getLogo()
```

**Value**

return logo

**Note**

```
getLogo
```

**Examples**

```
x <- getLogo()
```

---

```
getMainPanel      getMainPanel
```

---

**Description**

main panel for volcano, scatter and maplot. Barplot and box plots are in this page as well.

**Usage**

```
getMainPanel()
```

**Value**

the panel for main plots;

**Note**

```
getMainPanel
```

**Examples**

```
x <- getMainPanel()
```

---

```
getMainPlotsLeftMenu  getMainPlotsLeftMenu
```

---

**Description**

Generates the Main PLots Left menu to be displayed within the DEBrowser.

**Usage**

```
getMainPlotsLeftMenu()
```

**Value**

returns the left menu according to the selected tab;

**Note**

```
getMainPlotsLeftMenu
```

**Examples**

```
x <- getMainPlotsLeftMenu()
```

`getMainPlotUI`      *getMainPlotUI*

### Description

main plot for volcano, scatter and maplot.

### Usage

```
getMainPlotUI(id)
```

### Arguments

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

### Value

the panel for main plots;

### Note

```
getMainPlotUI
```

### Examples

```
x <- getMainPlotUI("main")
```

`getMean`      *getMean*

### Description

Gathers the mean for selected condition.

### Usage

```
getMean(data = NULL, selcols = NULL)
```

### Arguments

|          |            |
|----------|------------|
| data,    | dataset    |
| selcols, | input cols |

### Value

data

### Examples

```
x <- getMean()
```

---

```
getMergedComparison      getMergedComparison
```

---

### Description

Gathers the merged comparison data to be used within the DEBrowser.

### Usage

```
getMergedComparison(dc = NULL, nc = NULL, input = NULL)
```

### Arguments

|        |                           |
|--------|---------------------------|
| dc,    | data container            |
| nc,    | the number of comparisons |
| input, | input params              |

### Value

```
data
```

### Examples

```
x <- getMergedComparison()
```

---

```
getMetaSelector      getMetaSelector
```

---

### Description

Return the sample selection box using meta data table

### Usage

```
getMetaSelector(metadata = NULL, input = NULL, n = 0)
```

### Arguments

|           |                 |
|-----------|-----------------|
| metadata, | meta data table |
| input,    | input params    |
| n,        | the box number  |

### Value

```
meta select box
```

### Examples

```
x<-getMetaSelector()
```

`getMethodDetails`      *get the detail boxes after DE method selected*

### Description

get the detail boxes after DE method selected

### Usage

```
getMethodDetails(num = 0, input = NULL)
```

### Arguments

|        |                                 |
|--------|---------------------------------|
| num,   | panel that is going to be shown |
| input, | user input                      |

### Examples

```
x <- getMethodDetails()
```

`getMostVariedList`      *getMostVariedList*

### Description

Calculates the most varied genes to be used for specific plots within the DEBrowser.

### Usage

```
getMostVariedList(datavar = NULL, cols = NULL, input = NULL)
```

### Arguments

|          |                  |
|----------|------------------|
| datavar, | loaded dataset   |
| cols,    | selected columns |
| input,   | input            |

### Value

data

### Examples

```
x <- getMostVariedList()
```

---

```
getNormalizedMatrix      getNormalizedMatrix
```

---

**Description**

Normalizes the matrix passed to be used within various methods within DEBrowser. Requires edgeR package

**Usage**

```
getNormalizedMatrix(M = NULL, method = "TMM")
```

**Arguments**

|         |  |
|---------|--|
| M,      | numeric matrix                                 |
| method, | normalization method for edgeR. default is TMM |

**Value**

normalized matrix

**Note**

```
getNormalizedMatrix
```

**Examples**

```
x <- getNormalizedMatrix(mtcars)
```

---

```
getOrganism      getOrganism
```

---

**Description**

getOrganism

**Usage**

```
getOrganism(org)
```

**Arguments**

|      |          |
|------|----------|
| org, | organism |
|------|----------|

**Value**

organism name for keg

**Note**

```
getOrganism
```

**Examples**

```
x <- getOrganism()
```

---

|                |                       |
|----------------|-----------------------|
| getOrganismBox | <i>getOrganismBox</i> |
|----------------|-----------------------|

---

**Description**

Get the organism Box.

**Usage**

```
getOrganismBox()
```

**Value**

```
selectInput
```

**Note**

```
getOrganismBox
```

*getOrganismBox* makes the organism box

**Examples**

```
x <- getOrganismBox()
```

---

|                    |                           |
|--------------------|---------------------------|
| getOrganismPathway | <i>getOrganismPathway</i> |
|--------------------|---------------------------|

---

**Description**

```
getOrganismPathway
```

**Usage**

```
getOrganismPathway(org)
```

**Arguments**

|      |          |
|------|----------|
| org, | organism |
|------|----------|

**Value**

organism name for pathway

**Note**

```
getOrganismPathway
```

**Examples**

```
x <- getOrganismPathway()
```

---

getPCAcontrolUpdatesJS *getPCAcontrolUpdatesJS* in the prep menu we have two PCA plots to show how batch effect correction worked. One set of PCA input controls updates two PCA plots with this JS.

---

**Description**

getPCAcontrolUpdatesJS in the prep menu we have two PCA plots to show how batch effect correction worked. One set of PCA input controls updates two PCA plots with this JS.

**Usage**

```
getPCAcontrolUpdatesJS()
```

**Value**

the JS for tab updates

**Examples**

```
x<- getTabUpdateJS()
```

---

getPCAexplained      *getPCAexplained*

---

**Description**

Creates a more detailed plot using the PCA results from the selected dataset.

**Usage**

```
getPCAexplained(datasetInput = NULL, pca_data = NULL, input = NULL)
```

**Arguments**

datasetInput,    selected data  
pca\_data,        from user  
input,            input params

**Value**

explained plot

**Examples**

```
load(system.file("extdata", "demo", "demodata.Rda", package="debrowser"))
input<-c()
input$qcplot<-"pca"
input$col_list<-colnames(demodata[,1:6])
dat <- getNormalizedMatrix(demodata[,1:6])
pca_data <- run_pca(dat)
x <- getPCAExplained(dat, pca_data, input)
```

**getPCAPlotUI**

*getPCAPlotUI*

**Description**

PCA plots UI.

**Usage**

```
getPCAPlotUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

**Value**

the panel for PCA plots;

**Note**

`getPCAPlotUI`

**Examples**

```
x <- getPCAPlotUI("pca")
```

---

|                |                       |
|----------------|-----------------------|
| getPCselection | <i>getPCselection</i> |
|----------------|-----------------------|

---

**Description**

Generates the PC selection number to be used within DEBrowser.

**Usage**

```
getPCselection(id, num = 1, xy = "x")
```

**Arguments**

|      |                     |
|------|---------------------|
| id,  | namespace id        |
| num, | PC selection number |
| xy,  | x or y coordinate   |

**Value**

PC selection for PCA analysis

**Note**

```
getPCselection
```

**Examples**

```
x <- getPCselection("pca")
```

---

|             |                    |
|-------------|--------------------|
| getPlotArea | <i>getPlotArea</i> |
|-------------|--------------------|

---

**Description**

returns plot area either for heatmaply or heatmap.2

**Usage**

```
getPlotArea(input = NULL, session = NULL)
```

**Arguments**

|          |                 |
|----------|-----------------|
| input,   | input variables |
| session, | session         |

**Value**

heatmapply/heatmap.2 plot area

**Examples**

```
x <- getPlotArea()
```

|                 |                        |
|-----------------|------------------------|
| getProgramTitle | <i>getProgramTitle</i> |
|-----------------|------------------------|

**Description**

Generates the title of the program to be displayed within DEBrowser. If it is called in a program, the program title will be hidden

**Usage**

```
getProgramTitle(session = NULL)
```

**Arguments**

session, session var

**Value**

program title

**Note**

```
getProgramTitle
```

**Examples**

```
title<-getProgramTitle()
```

|           |   |
|-----------|---|
| getQAText | <i>getQAText Some questions and answers</i> |
|-----------|---|

**Description**

*getQAText Some questions and answers*

**Usage**

```
getQAText()
```

**Value**

help text for QA

**Examples**

```
x<- getQAText()
```

---

```
getQCLeftMenu
```

*getQCLeftMenu*

---

### Description

Generates the left menu to be used for QC plots within the DEBrowser.

### Usage

```
getQCLeftMenu(input = NULL)
```

### Arguments

input,                input values

### Value

QC left menu

### Note

```
getQCLeftMenu
```

### Examples

```
x <- getQCLeftMenu()
```

---

```
getQCPanel
```

*getQCPanel*

---

### Description

Gathers the conditional panel for QC plots

### Usage

```
getQCPanel(input = NULL)
```

### Arguments

input,                user input

### Value

the panel for QC plots

### Note

```
getQCSection
```

**Examples**

```
x <- getQCPanel()
```

|                               |                         |
|-------------------------------|-------------------------|
| <code>getSampleDetails</code> | <i>getSampleDetails</i> |
|-------------------------------|-------------------------|

**Description**

get sample details

**Usage**

```
getSampleDetails(output = NULL, summary = NULL, details = NULL,
                 data = NULL)
```

**Arguments**

|          |                     |
|----------|---------------------|
| output,  | output              |
| summary, | summary output name |
| details, | details ouput name  |
| data,    | data                |

**Value**

panel

**Examples**

```
x <- getSampleDetails()
```

|                             |                       |
|-----------------------------|-----------------------|
| <code>getSampleNames</code> | <i>getSampleNames</i> |
|-----------------------------|-----------------------|

**Description**

Prepares initial samples to fill condition boxes. it reads the sample names from the data and splits into two.

**Usage**

```
getSampleNames(cnames = NULL, part = 1)
```

**Arguments**

|         |   |
|---------|---|
| cnames, | sample names in the header of a dataset |
| part,   | c(1,2). 1=first half and 2= second half |

**Value**

sample names.

**Examples**

```
x<-getSampleNames()
```

---

*getSamples**getSamples*

---

**Description**

Gathers the sample names to be used within DEBrowser.

**Usage**

```
getSamples(cnames = NULL, index = 1)
```

**Arguments**

|         |   |
|---------|---|
| cnames, | names of the samples                    |
| index,  | starting column in a tab separated file |

**Value**

choices

**Examples**

```
x <- getSamples()
```

---

*getSearchData**getSearchData*

---

**Description**

search the geneset in the tables and return it

**Usage**

```
getSearchData(dat = NULL, input = NULL)
```

**Arguments**

|        |              |
|--------|--------------|
| dat,   | table data   |
| input, | input params |

**Value**

```
data
```

**Examples**

```
x <- getSearchData()
```

**getSelectedCols**

*getSelectedCols*

**Description**

gets selected columns

**Usage**

```
getSelectedCols(data = NULL, datasetInput = NULL, input = NULL)
```

**Arguments**

|               |                   |
|---------------|-------------------|
| data,         | all loaded data   |
| datasetInput, | selected dataset  |
| input,        | user input params |

**Examples**

```
getSelectedCols()
```

**getSelectedDatasetInput**

*getSelectedDatasetInput*

**Description**

Gathers the user selected dataset output to be displayed.

**Usage**

```
getSelectedDatasetInput(rdata = NULL, getSelected = NULL,
getMostVaried = NULL, mergedComparison = NULL, input = NULL)
```

*getSelectInputBox*

73

### Arguments

|                   |                        |
|-------------------|------------------------|
| rdata,            | filtered dataset       |
| getSelected,      | selected data          |
| getMostVaried,    | most varied data       |
| mergedComparison, | merged comparison data |
| input,            | input parameters       |

### Value

data

### Examples

```
x <- getSelectedDatasetInput()
```

---

getSelectInputBox      *getSelectInputBox*

---

### Description

Selects user input conditions to run in DESeq.

### Usage

```
getSelectInputBox(id = NULL, name = NULL, num = 0, choices = NULL,  
                  selected = NULL, cw = 2)
```

### Arguments

|           |                                 |
|-----------|---------------------------------|
| id,       | input id                        |
| name,     | label of the box                |
| num,      | panel that is going to be shown |
| choices,  | sample list                     |
| selected, | selected sample list            |
| cw,       | column width                    |

### Examples

```
x <- getSelectInputBox()
```

`getSelHeat`*getSelHeat***Description**

heatmap selection functionality

**Usage**

```
getSelHeat(expdata = NULL, input = NULL)
```

**Arguments**

|                        |                |
|------------------------|----------------|
| <code>expdata</code> , | selected genes |
| <code>input</code> ,   | input params   |

**Value**

plot

**Examples**

```
x <- getSelHeat()
```

`getShapeColor`*getShapeColor***Description**

Generates the fill and shape selection boxes for PCA plots. metadata file has to be loaded in this case

**Usage**

```
getShapeColor(input = NULL)
```

**Arguments**

|                      |              |
|----------------------|--------------|
| <code>input</code> , | input values |
|----------------------|--------------|

**Value**

Color and shape from selection boxes or defaults

**Examples**

```
x <- getShapeColor()
```

---

getStartPlotsMsg      *getStartPlotsMsg*

---

**Description**

Generates and displays the starting message to be shown once the user has first seen the main plots page within DEBrowser.

**Usage**

```
getStartPlotsMsg()
```

**Value**

return start plot msg

**Note**

```
getStartPlotsMsg
```

**Examples**

```
x <- getStartPlotsMsg()
```

---

getStartupMsg      *getStartupMsg*

---

**Description**

Generates and displays the starting message within DEBrowser.

**Usage**

```
getStartupMsg()
```

**Value**

return startup msg

**Note**

```
getStartupMsg
```

**Examples**

```
x <- getStartupMsg()
```

|                              |                        |
|------------------------------|------------------------|
| <code>getTableDetails</code> | <i>getTableDetails</i> |
|------------------------------|------------------------|

### Description

get table details To be able to put a table into two lines are necessary; into the server part; `getTableDetails(output, session, "dataname", data, modal=TRUE)` into the ui part; `uiOutput(ns("dataname"))`

### Usage

```
getTableDetails(output = NULL, session = NULL, tablename = NULL,
               data = NULL, modal = NULL)
```

### Arguments

|            |   |
|------------|---|
| output,    | output  |
| session,   | session   |
| tablename, | table name  |
| data,      | matrix data   |
| modal,     | if it is true, the matrix is going to be in a modal |

### Value

panel

### Examples

```
x <- getTableDetails()
```

|                            |  |
|----------------------------|--|
| <code>getTableModal</code> | <i>getTableModal</i> prepares table modal for KEGG |
|----------------------------|--|

### Description

`getTableModal` prepares table modal for KEGG

### Usage

```
getTableModal()
```

### Value

the info button

### Examples

```
x<- getTableModal()
```

---

|               |                      |
|---------------|----------------------|
| getTableStyle | <i>getTableStyle</i> |
|---------------|----------------------|

---

### Description

User defined selection that selects the style of table to display within the DEBrowser.

### Usage

```
getTableStyle(dat = NULL, input = NULL, padj = c("padj"),
  foldChange = c("foldChange"), DEsection = TRUE)
```

### Arguments

|             |  |
|-------------|--|
| dat,        | dataset  |
| input,      | input params                                     |
| padj,       | the name of the padj value column in the dataset |
| foldChange, | the name of the foldChange column in the dataset |
| DEsection,  | if it is in DESection or not                     |

### Note

```
getTableStyle
```

### Examples

```
x <- getTableStyle()
```

---

|                |   |
|----------------|---|
| getTabUpdateJS | <i>getTabUpdateJS prepmenu tab and discovery menu tab updates</i> |
|----------------|---|

---

### Description

getTabUpdateJS prepmenu tab and discovery menu tab updates

### Usage

```
getTabUpdateJS()
```

### Value

the JS for tab updates

### Examples

```
x<- getTabUpdateJS()
```

|              |                     |
|--------------|---------------------|
| getTextOnOff | <i>getTextOnOff</i> |
|--------------|---------------------|

### Description

text on PCA plot on and off

### Usage

```
getTextOnOff(id = "pca")
```

### Arguments

|     |              |
|-----|--------------|
| id, | namespace id |
|-----|--------------|

### Note

`getTextOnOff`

### Examples

```
x <- getTextOnOff("pca")
```

|       |                                    |
|-------|------------------------------------|
| setUp | <i>setUp</i> get up regulated data |
|-------|------------------------------------|

### Description

`setUp` get up regulated data

### Usage

```
setUp(filt_data = NULL)
```

### Arguments

|            |           |
|------------|-----------|
| filt_data, | filt_data |
|------------|-----------|

### Value

data

### Examples

```
x <- setUp()
```

---

`getUpDown`

*getUpDown get up+down regulated data*

---

**Description**

getUpDown get up+down regulated data

**Usage**

```
getUpDown(filt_data = NULL)
```

**Arguments**

filt\_data,       filt\_data

**Value**

data

**Examples**

```
x <- getUpDown()
```

---

---

`getVariationData`

*getVariationData*

---

**Description**

Adds an id to the data frame being used.

**Usage**

```
getVariationData(inputdata = NULL, cols = NULL, cond = NULL,
key = NULL)
```

**Arguments**

inputdata,       dataset  
cols,           columns  
cond,           conditions  
key,            gene or region name

**Value**

plotdata

**Examples**

```
x <- getVariationData()
```

```
get_conditions_given_selection
    get_conditions_given_selection
```

### Description

Return the two set of conditions given the selection of meta select box

### Usage

```
get_conditions_given_selection(metadata = NULL, selection = NULL)
```

### Arguments

|            |                 |
|------------|-----------------|
| metadata,  | meta data table |
| selection, | selection       |

### Value

meta select box

### Examples

```
x<-get_conditions_given_selection()
```

```
heatmapControlsUI      heatmapControlsUI
```

### Description

Generates the left menu to be used for heatmap plots

### Usage

```
heatmapControlsUI(id)
```

### Arguments

|     |           |
|-----|-----------|
| id, | module ID |
|-----|-----------|

### Value

HeatmapControls

### Note

```
heatmapControlsUI
```

### Examples

```
x <- heatmapControlsUI("heatmap")
```

---

heatmapJScode

---

*heatmapJScode*

---

**Description**

heatmap JS code for selection functionality

**Usage**

```
heatmapJScode()
```

**Value**

JS Code

**Examples**

```
x <- heatmapJScode()
```

---

---

heatmapServer

---

*heatmapServer*

---

**Description**

Sets up shinyServer to be able to run heatmapServer interactively.

**Usage**

```
heatmapServer(input, output, session)
```

**Arguments**

|          |                      |
|----------|----------------------|
| input,   | input params from UI |
| output,  | output params to UI  |
| session, | session variable     |

**Value**

the panel for main plots;

**Note**

```
heatmapServer
```

**Examples**

```
heatmapServer
```

**heatmapUI***heatmapUI***Description**

Creates a shinyUI to be able to run DEBrowser interactively.

**Usage**

```
heatmapUI(input, output, session)
```

**Arguments**

|                        |                 |
|------------------------|-----------------|
| <code>input</code> ,   | input variables |
| <code>output</code> ,  | output objects  |
| <code>session</code> , | session         |

**Value**

the panel for heatmapUI;

**Note**

```
heatmapUI
```

**Examples**

```
x<-heatmapUI()
```

**hideObj***hideObj***Description**

Hides a shiny object.

**Usage**

```
hideObj(btns = NULL)
```

**Arguments**

|                      |                                    |
|----------------------|------------------------------------|
| <code>bttns</code> , | hide group of objects with shinyjs |
|----------------------|------------------------------------|

**Examples**

```
x <- hideObj()
```

---

histogramControlsUI      *histogramControlsUI*

---

**Description**

Generates the controls in the left menu for a histogram

**Usage**

`histogramControlsUI(id)`

**Arguments**

`id`,                namespace id

**Value**

returns the left menu

**Note**

`histogramControlsUI`

**Examples**

`x <- histogramControlsUI("histogram")`

---

---

installpack      *installpack*

---

**Description**

install packages if they don't exist display.

**Usage**

`installpack(package_name = NULL)`

**Arguments**

`package_name`,    package name to be installed

**Note**

`installpack`

**Examples**

`x <- installpack()`

---

IQRPlotControlsUI      *IQRPlotControlsUI*

---

**Description**

Generates the controls in the left menu for an IQR plot#'

**Usage**

```
IQRPlotControlsUI(id)
```

**Arguments**

id,                        namespace id

**Value**

returns the left menu

**Note**

```
IQRPlotControlsUI
```

**Examples**

```
x <- IQRPlotControlsUI("IQR")
```

---

kmeansControlsUI      *kmeansControlsUI*

---

**Description**

get kmeans controls

**Usage**

```
kmeansControlsUI(id)
```

**Arguments**

id,                        module ID

**Value**

controls

**Note**

```
kmeansControlsUI
```

**Examples**

```
x <- kmeansControlsUI("heatmap")
```

---

lcfMetRadio

*lcfMetRadio*

---

**Description**

Radio buttons for low count removal methods

**Usage**

`lcfMetRadio(id)`

**Arguments**

`id`, namespace id

**Value**

radio control

**Note**

`lcfMetRadio`

**Examples**

`x <- lcfMetRadio("lcf")`

---

---

loadpack

*loadpack*

---

**Description**

load packages

**Usage**

`loadpack(package_name = NULL)`

**Arguments**

`package_name`, package name to be loaded

**Note**

`loadpack`

**Examples**

`x <- loadpack()`

`mainPlotControlsUI`      *mainPlotControlsUI*

### Description

Generates the left menu to be used for main plots

### Usage

```
mainPlotControlsUI(id)
```

### Arguments

`id`,                    module ID

### Value

`mainPlotControls`

### Note

```
mainPlotControlsUI
```

### Examples

```
x <- mainPlotControlsUI("main")
```

`mainScatterNew`      *mainScatterNew*

### Description

Creates the main scatter, volcano or MA plot to be displayed within the main panel.

### Usage

```
mainScatterNew(input = NULL, data = NULL, source = NULL)
```

### Arguments

|                       |  |
|-----------------------|--|
| <code>input</code> ,  | input params   |
| <code>data</code> ,   | dataframe that has log2FoldChange and log10padj values |
| <code>source</code> , | for event triggering to select genes                   |

### Value

scatter, volcano or MA plot

### Examples

```
x <- mainScatterNew()
```

---

`niceKmeans`*niceKmeans*

---

**Description**

Generates hierarchically clustered K-means clusters

**Usage**

```
niceKmeans(df = NULL, input = NULL, iter.max = 1000, nstart = 100)
```

**Arguments**

|                         |                                     |
|-------------------------|-------------------------------------|
| <code>df</code> ,       | data                                |
| <code>input</code> ,    | user inputs                         |
| <code>iter.max</code> , | max iteration for kmeans clustering |
| <code>nstart</code> ,   | n for kmeans clustering             |

**Value**

heatmap plot area

**Note**

`niceKmeans`

**Examples**

```
x <- niceKmeans()
```

---

`normalizationMethods`    *normalizationMethods*

---

**Description**

Select box to select normalization method prior to batch effect correction

**Usage**

```
normalizationMethods(id)
```

**Arguments**

|                   |              |
|-------------------|--------------|
| <code>id</code> , | namespace id |
|-------------------|--------------|

**Value**

radio control

**Note**

```
normalizationMethods
```

**Examples**

```
x <- normalizationMethods("batch")
```

**palUI**

*palUI*

**Description**

get pallete

**Usage**

```
palUI(id)
```

**Arguments**

|     |              |
|-----|--------------|
| id, | namespace ID |
|-----|--------------|

**Value**

pals

**Note**

```
palUI
```

**Examples**

```
x <- palUI("heatmap")
```

**panel.cor**

*panel.cor*

**Description**

Prepares the correlations for the all2all plot.

**Usage**

```
panel.cor(x, y, prefix = "rho=", cex.cor = 2, ...)
```

**Arguments**

|          |                       |
|----------|-----------------------|
| x,       | numeric vector x      |
| y,       | numeric vector y      |
| prefix,  | prefix for the text   |
| cex.cor, | correlation font size |
| ... ,    | additional parameters |

**Value**

all2all correlation plots

**Examples**

```
panel.cor(c(1,2,3), c(4,5,6))
```

---

panel.hist

*panel.hist*

---

**Description**

Prepares the histogram for the all2all plot.

**Usage**

```
panel.hist(x, ...)
```

**Arguments**

|       |   |
|-------|---|
| x,    | a vector of values for which the histogram is desired |
| ... , | any additional params                                 |

**Value**

all2all histogram plots

**Examples**

```
panel.hist(1)
```

pcaPlotControlsUI      *pcaPlotControlsUI*

### Description

Generates the PCA PLots Left menu to be displayed within the DEBrowser.

### Usage

```
pcaPlotControlsUI(id = "pca")
```

### Arguments

**id**,                namespace id

### Value

returns the left menu according to the selected tab;

### Note

```
pcaPlotControlsUI
```

### Examples

```
x <- pcaPlotControlsUI("pca")
```

plotData      *plotData*

### Description

prepare plot data for mainplots

### Usage

```
plotData(pdata = NULL, input = NULL)
```

### Arguments

**pdata**,                data  
**input**,                input

### Value

prepdata

### Note

```
plotData
```

**Examples**

```
x <- plotData()
```

---

plotMarginsUI

*plotMarginsUI*

---

**Description**

Margins module for plotly plots

**Usage**

```
plotMarginsUI(id, t = 20, b = 100, l = 100, r = 20)
```

**Arguments**

|     |               |
|-----|---------------|
| id, | id            |
| t,  | top margin    |
| b,  | bottom margin |
| l,  | left margin   |
| r,  | right margin  |

**Value**

size and margins controls

**Note**

```
plotMarginsUI
```

**Examples**

```
x <- plotMarginsUI("heatmap")
```

---

plotSizeMarginsUI

*plotSizeMarginsUI*

---

**Description**

Size and margins module for plotly plots

**Usage**

```
plotSizeMarginsUI(id, w = 800, h = 640, t = 20, b = 100, l = 100,  
r = 20)
```

**Arguments**

|                   |                            |
|-------------------|----------------------------|
| <code>id</code> , | <code>id</code>            |
| <code>w</code> ,  | <code>width</code>         |
| <code>h</code> ,  | <code>height</code>        |
| <code>t</code> ,  | <code>top margin</code>    |
| <code>b</code> ,  | <code>bottom margin</code> |
| <code>l</code> ,  | <code>left margin</code>   |
| <code>r</code> ,  | <code>right margin</code>  |

**Value**

size and margins controls

**Note**

`plotSizeMarginsUI`

**Examples**

```
x <- plotSizeMarginsUI("heatmap")
```

`plotSizeUI`

*plotSizeUI*

**Description**

Size module for plotly plots

**Usage**

```
plotSizeUI(id, w = 800, h = 600)
```

**Arguments**

|                   |                     |
|-------------------|---------------------|
| <code>id</code> , | <code>id</code>     |
| <code>w</code> ,  | <code>width</code>  |
| <code>h</code> ,  | <code>height</code> |

**Value**

size and margins controls

**Note**

`plotSizeUI`

**Examples**

```
x <- plotSizeUI("heatmap")
```

---

*plotTypeUI**plotTypeUI*

---

**Description**

Plot download type

**Usage**

```
plotTypeUI(id)
```

**Arguments**

|     |    |
|-----|----|
| id, | id |
|-----|----|

**Value**

size and margins controls

**Note**

```
plotTypeUI
```

**Examples**

```
x <- plotTypeUI("heatmap")
```

---

*plot\_pca**plot\_pca*

---

**Description**

Plots the PCA results for the selected dataset.

**Usage**

```
plot_pca(dat = NULL, pcx = 1, pcy = 2, metadata = NULL,
          color = NULL, shape = NULL, size = NULL, textonoff = "Off",
          legendSelect = "samples", input = NULL)
```

**Arguments**

|           |                 |
|-----------|-----------------|
| dat,      | data            |
| pcx,      | x axis label    |
| pcy,      | y axis label    |
| metadata, | additional data |
| color,    | color for plot  |
| shape,    | shape for plot  |

```

size,           size of the plot
textonoff,      text on off
legendSelect,   select legend
input,          input param

```

**Value**

pca list

**Examples**

```

load(system.file("extdata", "demo", "demodata.Rda",
                package="debrowser"))
metadata<-cbind(colnames(demodata[,1:6]),
                 colnames(demodata[,1:6]),
                 c(rep("Cond1",3), rep("Cond2",3)))
colnames(metadata)<-c("samples", "color", "shape")

a <- plot_pca(getNormalizedMatrix(
  demodata[rowSums(demodata[,1:6])>10,1:6]),
  metadata = metadata, color = "samples",
  size = 5, shape = "shape")

```

**prepDataContainer**      *prepDataContainer*

**Description**

Prepares the data container that stores values used within DESeq.

**Usage**

```
prepDataContainer(data = NULL, counter = NULL, input = NULL)
```

**Arguments**

```

data,           loaded dataset
counter,       the number of comparisons
input,          input parameters

```

**Value**

data

**Examples**

```
x <- prepDataContainer()
```

---

```
prepDEOutput
```

*prepDEOutput*

---

**Description**

Prepares the output data from DE analysis to be used within DEBrowser

**Usage**

```
prepDEOutput(data = NULL, cols = NULL, cond = NULL,  
            inputconds = NULL, i = NULL, input = NULL)
```

**Arguments**

|             |                            |
|-------------|----------------------------|
| data,       | loaded dataset             |
| cols,       | columns                    |
| cond,       | conds                      |
| inputconds, | inputconds                 |
| i,          | selected comparison number |
| input,      | input                      |

**Value**

```
data
```

**Examples**

```
x <- prepDEOutput()
```

---

```
prepGroup
```

*prepGroup*

---

**Description**

prepare group table

**Usage**

```
prepGroup(conds = NULL, cols = NULL)
```

**Arguments**

|        |            |
|--------|------------|
| conds, | inputconds |
| cols,  | columns    |

**Value**

```
data
```

**Examples**

```
x <- prepGroup()
```

---

prepHeatData

*prepHeatData*

---

**Description**

scales the data

**Usage**

```
prepHeatData(expdata = NULL, input = NULL)
```

**Arguments**

|          |  |
|----------|--|
| expdata, | a matrix that includes expression values |
| input,   | input variables                          |

**Value**

heatdata

**Examples**

```
x <- prepHeatData()
```

---

prepPCADat

*prepPCADat*

---

**Description**

prepares pca data with metadata. If metadata doesn't exists it puts all the samples into a single group; "Conds".

**Usage**

```
prepPCADat(pca_data = NULL, metadata = NULL, input = NULL, pcx = 1,
pcy = 2)
```

**Arguments**

|           |                      |
|-----------|----------------------|
| pca_data, | pca run results      |
| metadata, | additional meta data |
| input,    | input                |
| pcx,      | x axis label         |
| pcy,      | y axis label         |

**Value**

Color and shape from selection boxes or defaults

**Examples**

```
x <- prepPCADat()
```

---

push

*push*

---

**Description**

Push an object to the list.

**Usage**

```
push(1, ...)
```

**Arguments**

|      |                                    |
|------|------------------------------------|
| 1,   | that are going to push to the list |
| ..., | list object                        |

**Value**

combined list

**Examples**

```
mylist <- list()
newlist <- push ( 1, mylist )
```

---

removeCols

*removeCols*

---

**Description**

remove unnecessary columns

**Usage**

```
removeCols(cols = NULL, dat = NULL)
```

**Arguments**

|       |  |
|-------|--|
| cols, | columns that are going to be removed from data frame |
| dat,  | data   |

**Value**

data

**Examples**

```
x <- removeCols()
```

**removeExtraCols**

*removeExtraCols*

**Description**

remove extra columns for QC plots

**Usage**

```
removeExtraCols(dat = NULL)
```

**Arguments**

dat, selected data

**Examples**

```
removeExtraCols()
```

**round\_vals**

*round\_vals*

**Description**

Plot PCA results.

**Usage**

```
round_vals(1)
```

**Arguments**

1, the value

**Value**

round value

**Examples**

```
x<-round_vals(5.1323223)
```

---

`runDE`*runDE*

---

## Description

Run DE algorithms on the selected parameters. Output is to be used for the interactive display.

## Usage

```
runDE(data = NULL, columns = NULL, cond = NULL, params = NULL)
```

## Arguments

|          |  |
|----------|--|
| data,    | A matrix that includes all the expression raw counts, rownames has to be the gene, isoform or region names/IDs       |
| columns, | is a vector that includes the columns that are going to be analyzed. These columns has to match with the given data. |
| cond,    | experimental conditions. The order has to match with the column order  |
| params,  | all params for the DE methods  |

## Value

de results

## Examples

```
x <- runDE()
```

---

`runDESeq2`*runDESeq2*

---

## Description

Run DESeq2 algorithm on the selected conditions. Output is to be used for the interactive display.

## Usage

```
runDESeq2(data = NULL, columns = NULL, cond = NULL, params = NULL)
```

## Arguments

|          |  |
|----------|--|
| data,    | A matrix that includes all the expression raw counts, rownames has to be the gene, isoform or region names/IDs       |
| columns, | is a vector that includes the columns that are going to be analyzed. These columns has to match with the given data. |
| cond,    | experimental conditions. The order has to match with the column order  |

params,  
 fitType: either "parametric", "local", or "mean" for the type of fitting of dispersions to the mean intensity. See estimateDispersions for description. betaPrior: whether or not to put a zero-mean normal prior on the non-intercept coefficients. See nbinomWaldTest for description of the calculation of the beta prior. By default, the beta prior is used only for the Wald test, but can also be specified for the likelihood ratio test. testType: either "Wald" or "LRT", which will then use either Wald significance tests (defined by nbinomWaldTest), or the likelihood ratio test on the difference in deviance between a full and reduced model formula (defined by nbinomLRT) rowsum.filter: regions/genes/isoforms with total count (across all samples) below this value will be filtered out

### Value

deseq2 results

### Examples

```
x <- runDESeq2()
```

**runEdgeR**

*runEdgeR*

### Description

Run EdgeR algorithm on the selected conditions. Output is to be used for the interactive display.

### Usage

```
runEdgeR(data = NULL, columns = NULL, condns = NULL, params = NULL)
```

### Arguments

data, A matrix that includes all the expression raw counts, rownames has to be the gene, isoform or region names/IDs  
 columns, is a vector that includes the columns that are going to be analyzed. These columns has to match with the given data.  
 condns, experimental conditions. The order has to match with the column order  
 params, normfact: Calculate normalization factors to scale the raw library sizes. Values can be "TMM", "RLE", "upperquartile", "none". dispersion: either a numeric vector of dispersions or a character string indicating that dispersions should be taken from the data object. If a numeric vector, then can be either of length one or of length equal to the number of genes. Allowable character values are "common", "trended", "tagwise" or "auto". Default behavior ("auto" is to use most complex dispersions found in data object. testType: exactTest or glmLRT. exactTest: Computes p-values for differential abundance for each gene between two digital libraries, conditioning on the total count for each gene. The counts in each group as a proportion of the whole are assumed to follow a binomial distribution. glmLRT: Fit a negative binomial generalized log-linear model to the read counts for each gene. Conduct genewise statistical tests for a given coefficient or coefficient contrast. rowsum.filter: regions/genes/isoforms with total count (across all samples) below this value will be filtered out

**Value**

edgeR results

**Examples**

```
x <- runEdgeR()
```

---

*runHeatmap*

*runHeatmap*

---

**Description**

Creates a heatmap based on the user selected parameters within shiny

**Usage**

```
runHeatmap(input = NULL, session = NULL, expdata = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| session, | session                                  |
| expdata, | a matrix that includes expression values |

**Value**

heatmapply plot

**Examples**

```
x <- runHeatmap()
```

---

*runHeatmap2*

*runHeatmap2*

---

**Description**

Creates a heatmap based on the user selected parameters within shiny

**Usage**

```
runHeatmap2(input = NULL, session = NULL, expdata = NULL)
```

**Arguments**

|          |  |
|----------|--|
| input,   | input variables                          |
| session, | session                                  |
| expdata, | a matrix that includes expression values |

**Value**

```
heatmap.2
```

**Examples**

```
x <- runHeatmap2()
```

**runLimma**

*runLimma*

**Description**

Run Limma algorithm on the selected conditions. Output is to be used for the interactive display.

**Usage**

```
runLimma(data = NULL, columns = NULL, cond = NULL, params = NULL)
```

**Arguments**

- |          |   |
|----------|---|
| data,    | A matrix that includes all the expression raw counts, rownames has to be the gene, isoform or region names/IDs  |
| columns, | is a vector that includes the columns that are going to be analyzed. These columns has to match with the given data.  |
| cond,    | experimental conditions. The order has to match with the column order   |
| params,  | normfact: Calculate normalization factors to scale the raw library sizes. Values can be "TMM","RLE","upperquartile","none". fitType, fitting method; "ls" for least squares or "robust" for robust regression normBet: Normalizes expression intensities so that the intensities or log-ratios have similar distributions across a set of arrays. rowsum.filter: regions/genes/isoforms with total count (across all samples) below this value will be filtered out |

**Value**

Limma results

**Examples**

```
x <- runLimma()
```

---

`run_pca``run_pca`

---

### Description

Runs PCA on the selected dataset.

### Usage

```
run_pca(x = NULL, retx = TRUE, center = TRUE, scale = TRUE)
```

### Arguments

|         |  |
|---------|--|
| x,      | dataframe with experiment data           |
| retx,   | specifies if the data should be returned |
| center, | center the PCA (Boolean)                 |
| scale,  | scale the PCA (Boolean)                  |

### Value

pca list

### Examples

```
load(system.file("extdata", "demo", "demodata.Rda",
  package="debrowser"))
pca_data<-run_pca(getNormalizedMatrix(
  demodata[rowSums(demodata[,1:6])>10,1:6]))
```

---

`selectConditions``selectConditions`

---

### Description

Selects user input conditions, multiple if present, to be used in DESeq.

### Usage

```
selectConditions(Dataset = NULL, metadata = NULL,
  choicecounter = NULL, session = NULL, input = NULL)
```

### Arguments

|                |   |
|----------------|---|
| Dataset,       | used dataset                              |
| metadata,      | metadatatable to select from metadata     |
| choicecounter, | choicecounter to add multiple comparisons |
| session,       | session                                   |
| input,         | input params                              |

**Value**

the panel for go plots;

**Note**

`selectConditions`

**Examples**

```
x<- selectConditions()
```

`selectedInput`

*selectedInput*

**Description**

Selects user input conditions to run in DESeq.

**Usage**

```
selectedInput(id = NULL, num = 0, default = NULL, input = NULL)
```

**Arguments**

|                        |                                 |
|------------------------|---------------------------------|
| <code>id</code> ,      | input id                        |
| <code>num</code> ,     | panel that is going to be shown |
| <code>default</code> , | default text                    |
| <code>input</code> ,   | input params                    |

**Examples**

```
x <- selectedInput()
```

`selectGroupInfo`

*selectGroupInfo*

**Description**

Group info column selection. This can be used in batch effect or coloring the groups in the plots.

**Usage**

```
selectGroupInfo(metadata = NULL, input = NULL,
                selectname = "groupselect", label = "Group info")
```

**Arguments**

|             |                         |
|-------------|-------------------------|
| metadata,   | metadata                |
| input,      | input values            |
| selectname, | name of the select box  |
| label,      | label of the select box |

**Note**

`selectGroupInfo`

**Examples**

```
x <- selectGroupInfo()
```

---

sepRadio

*sepRadio*

---

**Description**

Radio button for separators

**Usage**

`sepRadio(id, name)`

**Arguments**

|       |           |
|-------|-----------|
| id,   | module id |
| name, | name      |

**Value**

radio control

**Note**

`sepRadio`

**Examples**

```
x <- sepRadio("meta", "metadata")
```

---

|                       |  |
|-----------------------|--|
| <code>setBatch</code> | <i>setBatch to skip batch effect correction batch variable set with the filter results</i> |
|-----------------------|--|

---

**Description**

`setBatch` to skip batch effect correction batch variable set with the filter results

**Usage**

```
setBatch(fd = NULL)
```

**Arguments**

`fd`,                filtered data

**Value**

`fd` data

**Examples**

```
x <- setBatch()
```

---

|                      |                |
|----------------------|----------------|
| <code>showObj</code> | <i>showObj</i> |
|----------------------|----------------|

---

**Description**

Displays a shiny object.

**Usage**

```
showObj(btns = NULL)
```

**Arguments**

`btns`,                show group of objects with shinyjs

**Examples**

```
x <- showObj()
```

---

`startDEBrowser`*startDEBrowser*

---

**Description**

Starts the DEBrowser to be able to run interactively.

**Usage**`startDEBrowser()`**Value**

the app

**Note**`startDEBrowser`**Examples**`startDEBrowser()`

---

`startHeatmap`*startHeatmap*

---

**Description**

Starts the DEBrowser heatmap

**Usage**`startHeatmap()`**Value**

the app

**Note**`startHeatmap`**Examples**`startHeatmap()`

|               |                      |
|---------------|----------------------|
| textareaInput | <i>textareaInput</i> |
|---------------|----------------------|

### Description

Generates a text area input to be used for gene selection within the DEBrowser.

### Usage

```
textareaInput(id, label, value, rows = 20, cols = 35,
             class = "form-control")
```

### Arguments

|        |                      |
|--------|----------------------|
| id,    | id of the control    |
| label, | label of the control |
| value, | initial value        |
| rows,  | the # of rows        |
| cols,  | the # of cols        |
| class, | css class            |

### Examples

```
x <- textareaInput("genesetarea", "Gene Set",
                    "Fgf21", rows = 5, cols = 35)
```

|              |                     |
|--------------|---------------------|
| togglePanels | <i>togglePanels</i> |
|--------------|---------------------|

### Description

User defined toggle to display which panels are to be shown within DEBrowser.

### Usage

```
togglePanels(num = NULL, nums = NULL, session = NULL)
```

### Arguments

|          |                |
|----------|----------------|
| num,     | selected panel |
| nums,    | all panels     |
| session, | session info   |

### Note

```
togglePanels
```

### Examples

```
x <- togglePanels()
```

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