

Package ‘OperaMate’

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Title An R package of Data Importing, Processing and Analysis for
Opera High Content Screening System

Author Chenglin Liu

Maintainer Chenglin Liu <cliu@sjtu.edu.cn>

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Suggests BiocStyle

Description OperaMate is a flexible R package dealing with the data
generated by PerkinElmer's Opera High Content Screening System.
The functions include the data importing, normalization and
quality control, hit detection and function analysis.

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biocViews Preprocessing, CellBasedAssays, Normalization,
QualityControl

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R topics documented:

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| | |
|----------------|---------------------------|
| cellData-class | <i>The cellData class</i> |
|----------------|---------------------------|

Description

The main class used in OperaMate to hold all levels of experiment data of a specific type.

Usage

```
cellData(name, positive.ctr = character(0), negative.ctr = character(0),
  expwell = character(0),
  norm.method = getOption("opm.normalization.method"),
  QC.threshold = getOption("opm.QC.threshold"))

## S4 method for signature 'cellData,character,ANY'
x[i]

## S4 method for signature 'cellData'
show(object)
```

Arguments

| | |
|--------------|---|
| name | character, the analyzed item |
| positive.ctr | a character vector, the positive control well IDs, e.g. c("E05", "E06") |
| negative.ctr | a character vector, the positive control well IDs, e.g. c("B05", "B06") |
| expwell | include all wells except control and neglect.well if NULL |
| norm.method | character the normalization method. |
| QC.threshold | numeric, the thresholds in the quality control. |
| x | a cellData object |
| i | a requested slot name |
| object | a cellData class |

Slots

| | |
|-------------|---|
| name | character, one parameter in the Columbus system report. |
| posctrwell | a character vector, the positive control well IDs, e.g. B05. |
| negctrwell | a character vector, the negative control well IDs, e.g. B05. |
| expwell | a character vector, the sample well IDs, e.g. C15. |
| cellNum | matrix, cell numbers |
| origin.data | a numeric matrix, the raw data matrix with rows the well IDs and columns the plate IDs. |
| norm.data | a numeric matrix, the normalized data. |

`qc.data` a numeric matrix, the data after quality control, with the rows are "barcode:wellID" and columns are the data of all replicated samples and their means, and if they have passed the quality control.

`norm.method` character the normalization method.

`QC.threshold` numeric, the thresholds in the quality control.

`plate.quality` a logical matrix, the quality data with the rows are the barcode and columns are the replicateIDs.

`plate.quality.data` a list of plate correlations and plate z' factors

`Sig` a list of the following components:

- `SigMat`: a logic matrix marking the high and low expressed hits
- `threshold`: the threshold of the high and low expressed hits
- `stats`: the numbers of the high and low expressed hits
- `pvalue`: the pvalue of each sample by t tests

Methods

Constructor `cellData(name, positive.ctr = character(0), negative.ctr = character(0))`

Show `signature(object="cellLoad")`. Displays object content as text.

Accessor `x[i]`. `x`: a cellData object; `i`: character, a cellData slot name.

Examples

```
oneCell <- cellData(name = "Average Intensity of Nuclei",
  positive.ctr = c("H02", "J02", "L02"),
  negative.ctr = c("C23", "E23", "G23"))
oneCell
oneCell["name"]
```

cellLoad

Data importing

Description

Extracts data of a specific type in a list of `expData` objects to initialize a `cellData` object.

Usage

```
cellLoad(object, lstPlates, ...)

## S4 method for signature 'cellData'
cellLoad(object, lstPlates, positive.ctr = NULL,
  negative.ctr = NULL, neglect.well = NULL, expwell = NULL)
```

Arguments

object a cellData object
 lstPlates a list of expData objects
 ... other parameters
 positive.ctr a character vector, the positive control well IDs, e.g. c("E05", "E06")
 negative.ctr a character vector, the positive control well IDs, e.g. c("B05", "B06")
 neglect.well a character vector, the neglect wells. Accept regular expression, e.g. c("*02", "*23")
 expwell include all wells except control and neglect.well if NULL

Details

negative.ctr accept regular expression

Value

a cellData object, with initialized slot `origin.data`

Examples

```

data(platemap)
platemap$Path <- file.path(
  system.file("Test", package = "OperaMate"), platemap$Path)
data(demoCell)
datapath <- file.path(system.file("Test", package = "OperaMate"), "Matrix")
lstPlates <- loadAll(cellformat = "Matrix", datapath = datapath)
oneCell <- cellLoad(oneCell, lstPlates, neglect.well = c("*02", "*23"))
str(oneCell["origin.data"])

```

cellMean

Mean of two cellData objects

Description

Merges the intensities in nucleus and cytoplasm to their averages for signature detection.

Usage

```

cellMean(cell1, cell2, name)

## S4 method for signature 'cellData,cellData,character'
cellMean(cell1, cell2, name)

```

Arguments

cell1 one cellData object
 cell2 another celldata object
 name the name of mean cellData object

Value

the mean cellData object

Examples

```
data(demoCell)
meanCell <- cellMean(oneCell, oneCell, "meanCell")
meanCell
```

cellNorm

Data normalization

Description

Normalizes raw data based on different normalization methods.

Usage

```
cellNorm(object, norm.method)

## S4 method for signature 'cellData'
cellNorm(object,
  norm.method = getOption("opm.normalization.method"))
```

Arguments

`object` a cellData object
`norm.method` `getOption("opm.normalization.method")`

Details

Method description: "MP" employs the median polish algorithm which divides data by the median of their plates and wells recursively, while "PMed" only divides data by the median of their plates; "Z" subtracts data by their plate medians, and then divides by the median absolute deviations; "Ctr" divides data by the mean of their plate negative controls; "None" avoids the data normalization in this step. The first three methods are based on the assumption that most samples display no biological effects in the assay be analyzed. They are often more effective than "Ctr" method as to the high throughput screening.

Value

a celldata object with initialized slot `norm.data`

Examples

```
data(demoCell)
oneCell <- cellNorm(oneCell, norm.method = "MP")
str(oneCell["norm.data"])
```

| | |
|-------------|-------------------------|
| cellNumLoad | <i>Load cell number</i> |
|-------------|-------------------------|

Description

Load cell number

Usage

```
cellNumLoad(object, object.cellnum)

## S4 method for signature 'cellData,cellData'
cellNumLoad(object, object.cellnum)
```

Arguments

| | |
|----------------|------------------------------------|
| object | a cellData object |
| object.cellnum | a cellData object for cell numbers |

Value

a cellData object, with initialized slot cellNum

Examples

```
data(demoCell)
data(demoCellNum)
oneCell <- cellNumLoad(oneCell, oneCellNum)
str(oneCell["cellNum"])
```

| | |
|--------|------------------------|
| cellQC | <i>Quality control</i> |
|--------|------------------------|

Description

Checks quality of all plates and then wells.

Usage

```
cellQC(object, qcType = NULL, qc.threshold = NULL,
  replace.badPlateData = TRUE, plot = TRUE,
  outpath = getOption("opm.outpath"), ...)

## S4 method for signature 'cellData'
cellQC(object, qcType = getOption("opm.QC.type"),
  qc.threshold = getOption("opm.QC.threshold"),
  replace.badPlateData = getOption("opm.replace.badPlateData"), plot = TRUE,
  outpath = getOption("opm.outpath"), ...)
```

Arguments

| | |
|----------------------|---|
| object | a cellData object |
| qcType | the type of quality control |
| qc.threshold | quality control thresholds |
| replace.badPlateData | if TRUE, replace the values of bad plate by their replicates |
| plot | if TRUE, plot figures |
| outpath | directory of output figures, default: <code>getOption("opm.outpath")</code> |
| ... | arguments for the graphic device |

Details

Requires three or more replicated samples.

qcType include `c("plateCorrelation", "wellSd", "zFactor", "cellNumber")`, An example of qc.threshold is `c(correlation = 0.8, zfactor = 0.5, cellnumber = 50)`.

Value

a cellData object with intialized slot `qc.data`, `plate.quality` and `plate.quality.data`.

Examples

```
data(demoCell)
op <- options("device")
options("device" = "png")
oneCell <- cellQC(oneCell, qcType = c("plateCorrelation", "wellSd", "cellNumber"),
qc.threshold = c(correlation = 0.7), outpath = tempdir())
options(op)
str(oneCell["qc.data"])
str(oneCell["plate.quality"])
```

cellSig

Hit identification

Description

Detects samples those are most different from the negative controls.

Usage

```
cellSig(object, method = c("stable", "ksd", "kmsd"), th = NULL,
thPval = 0.05, digits = 3, adjust.method = p.adjust.methods,
plot = TRUE, outpath = getOption("opm.outpath"), ...)
```

```
## S4 method for signature 'cellData'
cellSig(object, method = c("stable", "ksd", "kmsd"),
th = NULL, thPval = 0.05, digits = 3,
adjust.method = p.adjust.methods, plot = TRUE,
outpath = getOption("opm.outpath"), ...)
```

Arguments

| | |
|---------------|---|
| object | a cellData object |
| method | method = c("stable","ksd","kmsd"). Details are referred in the vignette. |
| th | numeric, the thresholds. It can be one threshold for both high and low expressed hit or two thresholds for each respectively. |
| thPval | numeric, threshold of pvalues in the t-test between the sample and control replicates |
| digits | integer, the number of digits used to show the thresholds |
| adjust.method | pvalue correction method |
| plot | plot QQ-plot when method is "stable" if TRUE. |
| outpath | directory of output figures, default: getOption("opm.outpath") |
| ... | arguments of the graphic device |

Value

a cellData object with initialized slot Sig.

Examples

```
data(demoCell)
op <- options("device")
options("device" = "png")
oneCell <- cellSig(oneCell, method = "stable", th = c(0.05, 0.05),
  outpath = tempdir())
options(op)
names(oneCell["Sig"])
```

cellSigAnalysis *Hits function analysis*

Description

Performs function analysis using gProfileR

Usage

```
cellSigAnalysis(object, genemap, organism, type = c("High", "Low"),
  file = NULL, ...)
```

Arguments

| | |
|----------|--|
| object | a cellData object |
| genemap | a data frame, the well-gene specification table |
| organism | organism name. |
| type | include both high and low expressed hits or one of them. |
| file | the filename of the enrichment table (default: disabled) |
| ... | the arguments of gprofiler. |

Details

genemap must include colnames "Barcode","Well","GeneSymbol". organism name can be referred to g:Profiler tool. For example, human: hsapiens, mouse: mmusculus.

Value

a data frame of the functional report from gProfiler

Examples

```
data(demoCell)
genemap <- read.csv(file.path(system.file("demoData", package = "OperaMate"),
"genemap.csv"), stringsAsFactors = FALSE)
chart <- cellSigAnalysis(oneCell, genemap, organism = "mmusculus")
head(chart)
```

cellSigAnalysisPlot

The barplot of enrichment functions

Description

The barplot of enrichment functions

Usage

```
cellSigAnalysisPlot(chart, prefix = "", type = NULL, fill = "steelblue",
  outpath = getOption("opm.outpath"), ...)
```

Arguments

| | |
|---------|--|
| chart | data frame, the functional annotation chart |
| prefix | character, the prefix of figure name |
| type | selected domains from chart, e.g. BP. |
| fill | color of the bars |
| outpath | directory of output figures, default: getOption("opm.outpath") |
| ... | other arguments for graphical devices |

Value

Invisibly the ggplot2 function for barplot

Examples

```
data(demoCell)
genemap <- read.csv(file.path(system.file("demoData", package = "OperaMate"),
"genemap.csv"), stringsAsFactors = FALSE)
chart <- cellSigAnalysis(oneCell, genemap, organism = "mmusculus")
op <- options("device")
options("device" = "png")
cellSigAnalysisPlot(chart, type = "BP", outpath = tempdir())
options(op)
```

`cellSigPlot`*Hits volcano plot*

Description

Visualizes hits by volcano plot.

Usage

```
cellSigPlot(object, outpath = getOption("opm.outpath"),
  color.highlight = getOption("opm.sig.color.highlight"),
  color.background = getOption("opm.sig.color.background"),
  highlight.label = NULL,
  highlight.label.color = getOption("opm.sig.label.color"), ...)
```

Arguments

| | |
|------------------------------------|--|
| <code>object</code> | a <code>cellData</code> object |
| <code>outpath</code> | directory of the output figures |
| <code>color.highlight</code> | a character specifying the color of the hits |
| <code>color.background</code> | a character specifying the color of the other samples |
| <code>highlight.label</code> | a vector of characters specifying the names of the samples to be highlighted, with the names are the "barcode:wellID". |
| <code>highlight.label.color</code> | a character specifying the color of the labels |
| <code>...</code> | arguments of the graphic device and <code>ggplot2</code> |

Details

Users can highlight a certain samples during plotting.

Value

Invisibly an object of `ggplot`

Examples

```
data(demoCell)
op <- options("device")
options("device" = "png")
labels <- c("Axin1")
names(labels) <- c("DSIMGA04:C07")
cellSigPlot(oneCell, highlight.label = labels, outpath = tempdir())
options(op)
```

`cellViz`*Data visualization*

Description

Visualize data by heatmap or boxplot.

Usage

```
cellViz(object, data.type = c("raw", "norm"), plot = c("heatmap",  
  "boxplot"), outpath = getOption("opm.outpath"), multiplot = FALSE,  
  plateID = NULL, tag = NULL, ctr.excluded = TRUE, ...)
```

Arguments

| | |
|---------------------------|---|
| <code>object</code> | a <code>cellData</code> object |
| <code>data.type</code> | <code>c("raw", "norm)</code> , visualizing both types by default |
| <code>plot</code> | <code>c("heatmap", "boxplot")</code> |
| <code>outpath</code> | directory of output figures, default: <code>getOption("opm.outpath")</code> |
| <code>multiplot</code> | logical, the output images are placed in one figure or not |
| <code>plateID</code> | numeric or character |
| <code>tag</code> | character, unique tag for one figure |
| <code>ctr.excluded</code> | logical, if controls are included in the visualization |
| <code>...</code> | other arguments for graphical devices and <code>pheatmap</code> |

Details

By visualizing the raw data, users can observe the batch effects as a large region of distinguishing color in heatmap or biased distribution by boxplots. Users can also visualize the normalized data for comparison.

Value

Invisibly a list of the values returned by `pheatmap` and `ggplot2` function for boxplot

Examples

```
data(demoCell)  
op <- options("device")  
options("device" = "png")  
cellViz(oneCell, data.type = c("raw", "norm"), plateID = 1:6, outpath = tempdir())  
cellViz(oneCell, data.type = c("raw", "norm"), plateID = 1, outpath = tempdir())  
options(op)
```

demoData

Examples of tables and cellData objects

Description

oneCellNum

Value

platemap: a data frame

oneCell: a cellData object

oneCellNum: a cellData object

platemap

Description The experiment information of each Columbus analysis report. This table is required only if the report formats are not standardized. See [loadAll](#) for more information.

Format data.frame with the following required column names:

FileName: character, the name of the report.

Format: character, only "Tab" and "Matrix" are supported in the current version.

Barcode: character, the barcode of the plates.

RepID: character, the ID to distinguish the replicated plates.

Path: character, the full path of the report.

oneCell

Description oneCell is a cellData object used in the examples of the package.

oneCellNum

Description oneCellNum is a cellData object storing the cell numbers.

Examples

```
data(platemap)
str(platemap)
data(demoCell)
oneCell
data(demoCellNum)
```

 expData-class *The expData class*

Description

The expData class is a container to store data imported from one Columbus system report

Constructor method of expData class.

Show method

Usage

```
expData(name, path, rep.id, exp.id, format)

## S4 method for signature 'expData'
show(object)

## S4 method for signature 'expData,ANY,ANY'
x[i]

dataLoad(object, data, wellID)

## S4 method for signature 'expData'
dataLoad(object, data, wellID)
```

Arguments

| | |
|--------|--|
| name | character, the plate ID (barcode-replicateID), e.g. DSIMGA03-s1. |
| path | character, the path of the Columbus system report. |
| rep.id | character, replicateID, e.g. s1. |
| exp.id | character, barcode, e.g. DSIMGA03. |
| format | character, format of the Columbus system report. |
| object | a expData class |
| x | a expData object |
| i | a requested slot name |
| data | the vectorized raw data matrix of one plate of each type. |
| wellID | a character vector, the well IDs. |

Slots

| | |
|--------|--|
| name | character, the plate ID (barcode-replicateID), e.g. DSIMGA03-s1. |
| path | character, the path of the Columbus system report. |
| rep.id | character, replicateID, e.g. s1. |
| exp.id | character, barcode, e.g. DSIMGA03. |
| data | a list of vectors, the vectorized raw data matrix of one plate of each type. |
| format | character, format of the Columbus system report. |
| wellID | a character vector, the well IDs. |

Methods

Constructor `expData(name, path, rep.id, exp.id, format)`.

Show `signature(object = "expData")`. Displays object content as text.

Accessor `x[i]`. `x`: an `expData` object; `i`: character, an `expData` slot name.

dataLoad `dataLoad(object, data, wellID)`

Examples

```
onePlate <- expData(name = "130504-s1-02.txt",
  path = file.path(system.file("Test", package = "OperaMate"),
    "Matrix", "130504-s1-02.txt"),
  rep.id = "s1",
  exp.id = "DSIMGA02",
  format = "Matrix")
onePlate
onePlate["name"]
```

generateReport

Report generation

Description

Summarizes all results in the list of `cellData` objects, and writes out a report to file.

Usage

```
generateReport(lstCells, genemap = NULL, verbose = FALSE, file = NULL,
  outpath = getOption("opm.outpath"), plot = TRUE, ...)
```

Arguments

| | |
|-----------------------|---|
| <code>lstCells</code> | a list of <code>cellData</code> objects |
| <code>genemap</code> | a data frame, the well-gene specification table |
| <code>verbose</code> | logical, detailed data will be provided if TRUE |
| <code>file</code> | the path of the file to generate to |
| <code>outpath</code> | a character string naming the location the figures to generate to |
| <code>plot</code> | if TRUE, plot barplot |
| <code>...</code> | arguments of the graphic device |

Details

This function summarizes the information from all `cellData` objects, and visualizes the number of the hists if required.

Value

a data frame with annotated information of each well

Examples

```

data(demoCell)
genemap <- read.csv(file.path(system.file("demoData", package = "OperaMate"),
"genemap.csv"), stringsAsFactors = FALSE)
report <- generateReport(list(oneCell), genemap, verbose = FALSE,
plot = FALSE)
str(report)

```

loadAll

*Data importing***Description**

Initializes a list of `expData` objects from the Columbus system reports.

Usage

```

loadAll(cellformat = NULL, datapath = "./",
        egFilename = getOption("opm.filename.example"), well.digits = 2,
        platemap = NULL)

```

Arguments

| | |
|--------------------------|--|
| <code>cellformat</code> | character specifying the format of the reports. Enable when <code>platemap</code> is <code>NULL</code> . |
| <code>datapath</code> | character specifying the location of the reports. Enable when <code>platemap</code> is <code>NULL</code> . |
| <code>egFilename</code> | a file name example |
| <code>well.digits</code> | the digits of the well column in the well-gene |
| <code>platemap</code> | data frame. See an example as platemap . |

Details

To facility the automatic file name parsing, the reports obtained from Columbus system should be of the same format, and located under the same directory. Users can obtain this plate specification table for further modification. An example of the table can be referred by [platemap](#). After modification, users can submit a plate specification data frame to parameter `platemap`. The data format supported for the reports are "Tab" and "Matrix". If the reports are of other cellformats, you can specify its `cellformat` and rewrite the function `parseTemplate` to import the data separately.

An example of `egFilename = list(eg.filename = "0205-s2-01.txt", rep.id = "s2", exp.id = "01", sep = "-", barcode = "DSIMGA01")`. `well.digits`: In the well-gene specification file, if the well ID is B1, B2, ..., B11, the `well.digit = 1`; while B01, B02, ..., B11, the `well.digit = 2`; and B001, B002, ..., B011, the `well.digit = 3`.

Value

a list of `expData` objects

Examples

```

# Data frame \code{platemap} provided
data(platemap)
platemap$Path <- file.path(
  system.file("Test", package = "OperaMate"), platemap$Path)
lstPlates <- loadAll(platemap = platemap)
#
# Consistent file name format
datapath <- file.path(system.file("Test", package = "OperaMate"), "Tab")
egFilename <- list(eg.filename = "Tab.130504-s1-01.txt",
  rep.id = "s1", exp.id = "01", sep = "-",
  barcode = "DSIMGA01")
lstPlates <- loadAll(cellformat = "Tab", datapath = datapath,
  egFilename = egFilename, well.digits = 2)
#
lstPlates[[1]]

```

nameParser

Plate information extraction

Description

Extract plate information from file names.

Usage

```
nameParser(vec.files, egFilename)
```

Arguments

`vec.files` a vector of file names
`egFilename` a file name example

Details

An example of `egFilename = list(eg.filename = "0205-s2-01.txt", rep.id = "s2", exp.id = "01", sep = "-", barcode = "DSIMGA01")`.

Value

a data frame of PlateID, RepID, and Barcode

operaMate *Data process and analysis pipeline*

Description

A systematical pipeline for opera data importing, normalization, quality control, hit detection, analysis, and visualization.

Usage

```
operaMate(configFile, gDevice = "png", ...)
```

Arguments

| | |
|------------|--|
| configFile | the location of the file specifying all parameters |
| gDevice | the graphics device |
| ... | addition arguments for graphics devices |

Value

a list of three components: a list of cellData objects, the annotated table of each well, and the enrichment analysis table

Examples

```
configFile <- file.path(system.file("demoData", package = "OperaMate"), "demoParam.txt")
operaReport <- operaMate(configFile, gDevice = "png")
head(operaReport$report)
```

parseTemplete *Data extraction from one report*

Description

Extracts data in the report to the slot data in the expData object. An inner function of [loadAll](#).

Usage

```
parseTemplete(onePlate, well.digits = 2)
```

Arguments

| | |
|-------------|---|
| onePlate | an expData object |
| well.digits | the digits of the well column in the well-gene specification file |

Value

an expData object with initialized slot data.

Examples

```
datapath <- file.path(system.file("Test", package = "OperaMate"), "Tab")
1stPlates <- loadAll(cellformat = "Tab", datapath = datapath )
onePlate <- parseTemplate(1stPlates[[1]])
```

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