Package 'openCyto'

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Type Package Title Hierarchical Gating Pipeline for flow cytometry data Version 2.2.0 Date 2012-06-11 Author Mike Jiang, John Ramey, Greg Finak, Raphael Gottardo Maintainer Mike Jiang <pre>wji.ang2@fhcrc.org>Jake Wagner <jpwagner@fhcrc.org></jpwagner@fhcrc.org></pre> Description This package is designed to facilitate the automated gating methods in sequential way to mimic the manual gating strategy. License Artistic-2.0 LazyLoad yes Imports methods,Biobase,BiocGenerics,gtools,flowCore(>= 1.99.17),flowViz,ncdfFlow(>= 2.11.34),flowWorkspace(>= 3.99.1),flowStats(>= 3.99.1),flowClust(>= 3.99.1),flowStats(>= 3.99.1),flowClust(>= 3.91.14),MASS,clue.plyr,RBGL.graph,data.table,ks,RColorBrewer,lattice,rrcov,R.utils Suggests flowWorkspaceData, knitr, testthat, utils, tools, parallel, ggcyto, CytoML biocViews ImmunoOncology, FlowCytometry, DataImport, Preprocessing, DataRepresentation Encoding UTF-8 VignetteBuilder knitr LinkingTo Rcpp RoxygenNote 7.1.1 git_url https://git.bioconductor.org/packages/openCyto git_branch RELEASE_3_12 git_last_commit_date 2020-10-27 Date/Publication 2021-03-29 R topics documented: as.data.table	,
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46

Description

It is the inverse function of gatingTemplate constructor.

Usage

```
## S3 method for class 'gatingTemplate'
as.data.table(x, keep.rownames = FALSE)
```

Arguments

```
x gating Template objectkeep.rownames not used
```

Value

a data.table

boolMethod-class

A class to represent a boolean gating method.

Description

It extends refGate class.

4 fcEllipsoidGate

dims, gtMethod-method get gating method dimensions

Description

get gating method dimensions

Usage

```
## S4 method for signature 'gtMethod'
dims(x)
```

Arguments

x gtMethod

dummyMethod-class

A class to represent a dummy gating method that does nothing but serves as reference to be refered by other population

Description

It is generated automatically by the csv template preprocessing to handle the gating function that returns multiple gates.

fcEllipsoidGate

 $constuctor for \; {\tt fcEllipsoidGate}$

Description

```
constuctor\ for\ fc Ellipsoid Gate
```

Usage

```
fcEllipsoidGate(x, priors, posts)
```

Arguments

x a ellipsoidGate object
priors a list storing priors
posts a list storing posteriors

fcEllipsoidGate-class 5

 ${\tt fcEllipsoidGate-class} \ \ \textit{a concrete class that reprents the ellipsoidGate generated by flowClust}$

Description

It stores priors and posteriors as well as the actual ellipsoidGate.

fcFilter-class

a virtual class that represents the gating result generated by flowClust gating function

Description

Bascially it extends flowCore 'filter classes to have extra slot to store priors and posteriors

fcFilterList

 $constuctor\ for\ fcFilterList$

Description

constuctor for fcFilterList

Usage

fcFilterList(x)

Arguments

Χ

list of fcFilter (i.e. fcPolygonGate or fcRectangleGate)

fcFilterList-class

a class that extends filterList class.

Description

Each filter in the filterList must extends the fcFilter class

6 fcRectangleGate

 ${\it fcPolygonGate}$

 $constuctor for \ {\tt fcPolygonGate}$

Description

constuctor for fcPolygonGate

Usage

```
fcPolygonGate(x, priors, posts)
```

Arguments

x a polygonGate objectpriors a list storing priorsposts a list storing posteriors

 ${\it fc} Polygon {\it Gate-class}$

a concrete class that reprents the polygonGate generated by flowClust

Description

It stores priors and posteriors as well as the actual polygonGate.

 ${\tt fcRectangleGate}$

 $constuctor for \; {\tt fcRectangleGate}$

Description

 $constuctor \ for \ fc {\tt RectangleGate}$

Usage

```
fcRectangleGate(x, priors, posts)
```

Arguments

x a rectangleGate object
priors a list storing priors
posts a list storing posteriors

fcRectangleGate-class 7

fcRectangleGate-class a concrete class that reprents the rectangleGate generated by flow-Clust

Description

It stores priors and posteriors as well as the actual rectangleGate.

fcTree

constructor of fcTree

Description

It adds an extra node data slot "fList" (which is a filterList object) to the gatingTemplate

Usage

fcTree(gt)

Arguments

gt

a gatingTemplate object

fcTree-class

A class to represent a flowClust tree.

Description

It is a graphNEL used as a container to store priors and posteriors for each flowClust gate that can be visualized for the purpose of fine-tunning parameters for flowClust algorithm

gate_flowclust_1d

Applies flowClust to 1 feature to determine a cutpoint between the minimum cluster and all other clusters.

Description

We cluster the observations in fr into K clusters.

8 gate_flowclust_1d

Usage

```
gate_flowclust_1d(
  fr,
 params,
 filterId = "",
 K = NULL
  trans = 0,
 min.count = -1,
 max.count = -1,
 nstart = 1,
 prior = NULL,
 criterion = c("BIC", "ICL"),
  cutpoint_method = c("boundary", "min_density", "quantile", "posterior_mean",
    "prior_density"),
  neg_cluster = 1,
  cutpoint_min = NULL,
  cutpoint_max = NULL,
 min = NULL,
 max = NULL,
  quantile = 0.99,
  quantile_interval = c(0, 10),
 plot = FALSE,
 debug = FALSE,
)
```

Arguments

fr a flowFrame object

params character channel to be gated on

filterId A character string that identifies the filter created.

K the number of clusters to find

trans, min.count, max.count, nstart

some flowClust parameters. see flowClust

prior list of prior parameters for the Bayesian flowClust. If NULL, no prior is used.

criterion a character string stating the criterion used to choose the best model. May take

either "BIC" or "ICL". This argument is only relevant when K is NULL or if

length(K) > 1. The value selected is passed to flowClust.

cutpoint_method

How should the cutpoint be chosen from the fitted flowClust model? See De-

tails.

neg_cluster integer. The index of the negative cluster. The cutpoint is computed between

clusters neg_cluster and neg_cluster + 1.

cutpoint_min numeric value that sets a minimum thresold for the cutpoint. If a value is pro-

vided, any cutpoint below this value will be set to the given minimum value. If

NULL (default), there is no minimum cutpoint value.

vided, any cutpoint above this value will be set to the given maximum value. If

NULL (default), there is no maximum cutpoint value.

gate_flowclust_2d 9

min a numeric value that sets the lower bound for data filtering. If NULL (default), no

truncation is applied.

max a numeric value that sets the upper bound for data filtering. If NULL (default), no

truncation is applied.

quantile the quantile for which we will find the cutpoint using the quantile cutpoint_method.

If the cutpoint_method is not set to quantile, this argument is ignored.

quantile_interval

a vector of length 2 containing the end-points of the interval of values to find the quantile cutpoint. If the cutpoint_method is not set to quantile, this argument

is ignored.

plot logical value indicating that the fitted flowClust model should be plotted along

with the cutpoint

debug logical indicating whether to carry the prior and posterious with the gate for

debugging purpose. Default is FALSE.

... additional arguments that are passed to flowClust

Details

By default, the cutpoint is chosen to be the boundary of the first two clusters. That is, between the first two cluster centroids, we find the midpoint between the largest observation from the first cluster and the smallest observations from the second cluster. Alternatively, if the cutpoint_method is min_density, then the cutpoint is the point at which the density between the first and second smallest cluster centroids is minimum.

Value

a rectangleGate object consisting of all values beyond the cutpoint calculated

Examples

```
## Not run:
  gate <- gate_flowclust_1d(fr, params = "APC-A", K =2) # fr is a flowFrame
## End(Not run)</pre>
```

gate_flowclust_2d Automatic identification of a population of interest via flowClust based on two markers

Description

We cluster the observations in fr into K clusters. We set the cutpoint to be the point at which the density between the first and second smallest cluster centroids is minimum.

10 gate_flowclust_2d

Usage

```
gate_flowclust_2d(
  fr,
  xChannel,
  yChannel,
  filterId = "",
  K = 2,
  usePrior = "no",
  prior = list(NA),
  trans = 0,
  min.count = -1,
  max.count = -1,
  nstart = 1,
  plot = FALSE,
  target = NULL,
  transitional = FALSE,
  quantile = 0.9,
  translation = 0.25,
  transitional_angle = NULL,
  min = NULL,
  max = NULL,
)
```

Arguments

fr

a flowFrame object xChannel, yChannel character specifying channels to be gated on filterId A character string that identifies the filter created. the number of clusters to find usePrior Should we use the Bayesian version of flowClust? Answers are "yes", "no", or "vague". The answer is passed along to flowClust. prior list of prior parameters for the Bayesian version of flowClust. If usePrior is set to no, then the list is unused. trans, min.count, max.count, nstart some flowClust parameters. see flowClust plot a logical value indicating if the fitted mixture model should be plotted. By default, no. a numeric vector of length 2 (number of dimensions) containing the location of target the cluster of interest. See details. logical value indicating if a transitional gate should be constructed from the transitional target flowClust cluster. By default, no. quantile the contour level of the target cluster from the flowClust fit to construct the gate a numeric value between 0 and 1 used to position a transitional gate if transitional translation

= TRUE. This argument is ignored if transitional = FALSE. See details

gate_mindensity 11

the angle (in radians) of the transitional gate. It is also used to determine which quadrant the final gate resides in. See details. Ignored if transitional = FALSE.

min A vector of length 2. Truncate observations less than this minimum value. The first value truncates the xChannel, and the second value truncates the yChannel. By default, this vector is NULL and is ignored.

max A vector of length 2. Truncate observations greater than this maximum value. The first value truncates the xChannel, and the second value truncates the yChannel. By default, this vector is NULL and is ignored.

... additional arguments that are passed to flowClust

Details

The cluster for the population of interest is selected as the one with cluster centroid nearest the target in Euclidean distance. By default, the largest cluster (i.e., the cluster with the largest proportion of observations) is selected as the population of interest.

We also provide the option of constructing a transitional gate from the selected population of interest. The location of the gate can be controlled with the translation argument, which translates the gate along the major axis of the targest cluster as a function of the appropriate chi-squared coefficient. The larger translation is, the more gate is shifted in a positive direction. Furthermore, the width of the transitional gate can be controlled with the quantile argument.

The direction of the transitional gate can be controlled with the transitional_angle argument. By default, it is NULL, and we use the eigenvector of the target cluster that points towards the first quadrant (has positive slope). If transitional_angle is specified, we rotate the eigenvectors so that the angle between the x-axis (with the cluster centroid as the origin) and the major eigenvector (i.e., the eigenvector with the larger eigenvalue) is transitional_angle. So based on range that the angle falls in, the final rectangleGate will be constructed at the corresponding quadrant. i.e. Clockwise, [0,pi/2] UR, (pi/2, pi] LR, (pi, 3/2 * pi] LL, (3/2 * pi, 2 * pi] UL

Value

a polygonGate object containing the contour (ellipse) for 2D gating.

Examples

Description

We fit a kernel density estimator to the cells in the flowFrame and identify the two largest peaks. We then select as the cutpoint the value at which the minimum density is attained between the two peaks of interest.

12 gate_mindensity

Usage

```
gate_mindensity(
   fr,
   channel,
   filterId = "",
   positive = TRUE,
   pivot = FALSE,
   gate_range = NULL,
   min = NULL,
   max = NULL,
   peaks = NULL,
   ...
)
```

Arguments

fr a flowFrame object

channel TODO filterId TODO

positive If TRUE, then the gate consists of the entire real line to the right of the cutpoint.

Otherwise, the gate is the entire real line to the left of the cutpoint. (Default:

TRUE)

pivot logical value. If TRUE, we choose as the two peaks the largest peak and its

neighboring peak. See details.

gate_range numeric vector of length 2. If given, this sets the bounds on the gate applied. If

no gate is found within this range, we set the gate to the minimum value within this range if positive is TRUE and the maximum value of the range otherwise.

a numeric value that sets the lower boundary for data filtering

max a numeric value that sets the upper boundary for data filtering

peaks numeric vector. If not given , then perform peak detection first by .find_peaks

... Additional arguments for peak detection.

Details

min

In the default case, the two peaks of interest are the two largest peaks obtained from the link{density} function. However, if pivot is TRUE, we choose the largest peak and its neighboring peak as the two peaks of interest. In this case, the neighboring peak is the peak immediately to the left of the largest peak if positive is TRUE. Otherwise, the neighboring peak is selected as the peak to the right.

In the special case that there is only one peak, we are conservative and set the cutpoint as the min(x) if positive is TRUE, and the max(x) otherwise.

Value

a rectangleGate object based on the minimum density cutpoint

Examples

```
## Not run:
  gate <- gate_mindensity(fr, channel = "APC-A") # fr is a flowFrame
## End(Not run)</pre>
```

gate_mindensity2

gate_mindensity2	An improved version of mindensity used to determines a cutpoint as
	the minimum point of a kernel density estimate between two peaks.

Description

Analogous to the original openCyto::mindensity(), mindensity2 operates on a standard flowFrame. Its behavior is closely modeled on the original mindensity() whenever possible. However, the underlying peak-finding algorithm (improvedMindensity) behaves significantly differently.

Usage

```
gate_mindensity2(
    fr,
    channel,
    filterId = "",
    pivot = FALSE,
    gate_range = NULL,
    min = NULL,
    max = NULL,
    peaks = NULL,
    ...
)
```

Arguments

fr	a flowFrame object
channel	the channel to operate on
filterId	a name to refer to this filter
pivot	logical value. If TRUE, we choose as the two peaks the largest peak and its neighboring peak. See details.
gate_range	numeric vector of length 2. If given, this sets the bounds on the gate applied.
min	a numeric value that sets the lower boundary for data filtering
max	a numeric value that sets the upper boundary for data filtering
peaks	numeric vector. If not given , then perform peak detection first by .find_peaks
	Additional arguments for peak detection.

Value

a rectangleGate object based on the minimum density cutpoint

Author(s)

```
Greg Finak, Phu T. Van
```

Examples

```
## Not run:
gate <- gate_mindensity2(fr, channel = "APC-A") # fr is a flowFrame
## End(Not run)</pre>
```

14 gate_quad_tmix

```
gate_quad_sequential sequential quadrant gating function
```

Description

The order of 1d-gating is determined so that the gates better capture the distributions of flow data.

Usage

```
gate_quad_sequential(fr, channels, gFunc, min = NULL, max = NULL, ...)
```

Arguments

```
fr flowFrame
channels character two channels used for gating
gFunc the name of the 1d-gating function to be used for either dimension
min a numeric vector that sets the lower bounds for data filtering
max a numeric vector that sets the upper bounds for data filtering
other arguments passed to .find_peak (e.g. 'num_peaks' and 'adjust'). see
tailgate
```

Value

a filters that contains four rectangleGates

```
gate_quad_tmix quadGate based on flowClust::tmixFiler
```

Description

This gating method identifies two quadrants (first, and third quadrants) by fitting the data with tmixture model. It is particually useful when the two markers are not well resolved thus the regular quadGate method based on 1d gating will not find the perfect cut points on both dimensions.

Usage

```
gate_quad_tmix(
   fr,
   channels,
   K,
   usePrior = "no",
   prior = list(NA),
   quantile1 = 0.8,
   quantile3 = 0.8,
   trans = 0,
   plot = FALSE,
   ...
)
```

gate_quantile 15

Arguments

fr	flowFrame
channels	character vector specifies two channels
K	see gate_flowclust_2d
usePrior	see gate_flowclust_2d
prior	see gate_flowclust_2d
quantile1	numeric specifies the quantile level (see 'level' in $flowClust$) for the first quadrant (x-y+)
quantile3	numeric specifies the quantile level see 'level' in flowClust for third quadrant $(x+y-)$
trans	see gate_flowclust_2d
plot	logical whether to plot flowClust clustering results
	other arguments passed to flowClust

Value

a filters object that contains four polygonGates following the order of (-+,++,+-,-)

gate_quantile

Determine the cutpoint by the events quantile.

Description

It is possible that the cutpoint calculated by quantile function may not produce the exact the probability set by 'probs' argument if there are not enough cell events to reach that precision. Sometime the difference could be significant.

Usage

```
gate_quantile(
   fr,
   channel,
   probs = 0.999,
   plot = FALSE,
   filterId = "",
   min = NULL,
   max = NULL,
   ...
)
```

Arguments

```
fr
                   a flowFrame object
                   the channel from which the cytokine gate is constructed
channel
                   probabilities passed to 'stats::quantile' function.
probs
plot
                   whether to plot the gate result
filterId
                   the name of the filter
                   a numeric value that sets the lower boundary for data filtering
min
                   a numeric value that sets the upper boundary for data filtering
max
                   additional arguments passed to 'stats::quantile' function.
. . .
```

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Value

```
a rectangleGate
```

Examples

```
## Not run:
  gate <- gate_quantile(fr, Channel = "APC-A", probs = 0.995) # fr is a flowFrame
## End(Not run)</pre>
```

gate_tail

Gates the tail of a density using the derivative of a kernel density estimate

Description

These methods aim to set a one-dimensional gate (cutpoint) near the edge of a peak in the density specified by a channel of a flowFrame to isolate the tail population. They allow two approaches to do this, both beginning by obtaining a smoothened kernel density estimate (KDE) of the original density and then utilizing either its first or second derivative.

We determine a gating cutpoint using either the first or second derivative of the kernel density estimate (KDE) of the x.

Usage

```
gate_tail(
  fr,
  channel,
  filterId = "",
  num_peaks = 1,
  ref_peak = 1,
  strict = TRUE,
  tol = 0.01,
  side = "right",
  min = NULL,
  max = NULL,
  bias = 0,
  positive = TRUE,
.cytokine_cutpoint(
  х,
  num_peaks = 1,
  ref_peak = 1,
  method = c("first_deriv", "second_deriv"),
  tol = 0.01,
  adjust = 1,
  side = "right",
  strict = TRUE,
  plot = FALSE,
```

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```
auto_tol = FALSE,
...
)
```

Arguments

fr	a flowFrame object
channel	the channel from which the cytokine gate is constructed
filterId	the name of the filter
num_peaks	the number of peaks expected to see. This effectively removes any peaks that are artifacts of smoothing
ref_peak	After num_peaks are found, this argument provides the index of the reference population from which a gate will be obtained. By default, the peak farthest to the left is used.
strict	logical when the actual number of peaks detected is less than ref_peak. an error is reported by default. But if strict is set to FALSE, then the reference peak will be reset to the peak of the far right.
tol	the tolerance value
side	On which side of the density do we want to gate the tail. Valid options are "left" or "right".
min	a numeric value that sets the lower boundary for data filtering
max	a numeric value that sets the upper boundary for data filtering
bias	a numeric value that adds a constant to the calculated cutpoint(threshold). Default is 0.
positive	If FALSE, after finding the cutpoint, the rectangleGate returned will have bounds (-Inf,a] as opposed to [a,Inf). This argument will be ignored if gate_tail is used in a gatingTemplate with gt_gating or via gs_add_gating_method. In those cases, pop = "-" should be used instead.
	additional arguments passed to .deriv_density
Х	a numeric vector used as input data
method	the method used to select the cutpoint. See details.
adjust	the scaling adjustment applied to the bandwidth used in the first derivative of the kernel density estimate
plot	logical specifying whether to plot the peaks found 'right' (default) or 'left'?
auto_tol	when TRUE, it tries to set the tolerance automatically.

Details

The default behavior of the first approach, specified by method = "first_deriv", finds valleys in the first derivative of the KDE and uses the lowest such valley to place the cutpoint on the steep right shoulder of the largest peak in the original density.

The default behavior of the second approach, specified by method = "second_deriv", is to find peaks in the second derivative of the KDE and use the largest such peak to place the cutpoint at the point on the right shoulder of the largest peak in the original density where it is most rapidly flattening (the first derivative is rapidly growing less negative).

Both approaches can be significantly modified from defaults with a number of optional arguments. The num_peaks argument specifes how many peaks should be found in the smoothened KDE and

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ref_peak specifies around which peak the gate's cutpoint should be placed (starting from the leftmost peak). Setting the side argument to "left" modifies the procedure to put the cutpoint on the left side of the reference peak to isolate a left tail. The max and min arguments allow for pre-filtering extreme values in the channel of interest (keeping only observations with channel values less than max and/or more than min). The bandwidth used for kernel density estimation can be proportionally scaled using adjust (e.g. adjust = 0.5 will use a bandwidth that is half of the default). This allows for tuning the level of smoothing applied in the estimation.

Lastly, the tol, auto_tol, and bias arguments allow for adjustments to be made to the cutpoint that would otherwise be returned. tol provides a tolerance value that the absolute value of the KDE derivative at the cutpoint must be under. If the derivative at the original cutpoint is greater than tol in magnitude, the returned cutpoint will be the first point to the right of the original cutpoint (or to the left in the case of side = "left") with corresponding derivative within tol. Thus in practice, a smaller value for tol effectively pushes the cutpoint further down the shoulder of the peak towards the flat tail. tol is set to 0.01 by default but setting auto_tol = TRUE will set the tolerance to a reasonable estimate of 1% of the maximum absolute value of the first derivative of the KDE. tol and auto_tol are only used for method = "first_deriv". Additionally, the bias argument allows for directly shifting the returned cutpoint left or right.

It is also possible to pass additional arguments to control the calculation of the derivative, which will have some effect on the resulting cutpoint determination, but this should usually not be needed. By default the number of grid points for the derivative calculation will be 10,000, but this can be changed with num_points. The default bandwidth can also be directly adjusted with bandwidth, where the final value used will be given by adjust*bandwidth

By default, we compute the first derivative of the kernel density estimate. Next, we determine the lowest valley from the derivative, which corresponds to the density's mode for cytokines. We then contruct a gating cutpoint as the value less than the tolerance value tol in magnitude and is also greater than the lowest valley.

Alternatively, if the method is selected as second_deriv, we select a cutpoint from the second derivative of the KDE. Specifically, we choose the cutpoint as the largest peak of the second derivative of the KDE density which is greater than the reference peak.

Value

a filterList containing the gates (cutpoints) for each sample with the corresponding rectangleGate objects defining the tail as the positive population.

the cutpoint along the x-axis

Examples

```
## Not run:
  gate <- gate_tail(fr, Channel = "APC-A") # fr is a flowFrame
## End(Not run)</pre>
```

gatingTemplate-class a class storing the gating method and population information in a graphNEL object

gatingTemplate-class 19

Description

Each cell population is stored in graph node and is connected with its parent population or its reference node for boolGate or refGate.

It parses the csv file that specifies the gating scheme for a particular staining pannel.

Usage

```
gatingTemplate(x, ...)
## S4 method for signature 'character'
gatingTemplate(
    x,
    name = "default",
    strict = TRUE,
    strip_extra_quotes = FALSE,
    ...
)
```

Arguments

x character csv file name

... other arguments passed to data.table::fread

name character the label of the gating template

strict logical whether to perform validity check(special characters) on the alias col-

umn. By default it is(and should be) turned on for the regular template parsing. But sometime it is useful to turned it off to bypass the check for the dummy nodes(e.g. the csv template generated by 'gh_generate_template' with some

existing boolean gates that has '!' or ':' symbol).

strip_extra_quotes

logical Extra quotes are added to strings by fread. This causes problems with parsing R strings to expressions in some cases. Default FALSE for usual behaviour. TRUE should be passed if parsing gating_args fails.

Details

This csv must have the following columns:

'alias': a name used label the cell population, the path composed by the alias and its precedent nodes (e.g. /root/A/B/alias) has to be uniquely identifiable. So alias can not contain '/' character, which is reserved as path delimiter.

'pop': population patterns of '+/-' or '+/-+/-', which tells the algorithm which side (postive or negative) of 1d gate or which quadrant of 2d gate to be kept.

'parent': the parent population alias, its path has to be uniquely identifiable.

'dims': characters seperated by comma specifying the dimensions(1d or 2d) used for gating. It can be either channel name or stained marker name (or the substrings of channel/marker names as long as they are uniquely identifiable.).

'gating_method': the name of the gating function (e.g. 'flowClust'). It is invoked by a wrapper function that has the identical function name prefixed with a dot.(e.g. '.flowClust')

'gating_args': the named arguments passed to gating function (Note that double quotes are often used as text delimiter by some csv editors. So try to use single quote instead if needed.)

'collapseDataForGating': When TRUE, data is collapsed (within groups if 'groupBy' specified) before gating and the gate is replicated across collapsed samples. When set FALSE (or blank),then 'groupBy' argument is only used by 'preprocessing' and ignored by gating.

'groupBy': If given, samples are split into groups by the unique combinations of study variable (i.e. column names of pData,e.g. "PTID: VISITNO"). when split is numeric, then samples are grouped by every N samples

'preprocessing_method': the name of the preprocessing function(e.g. 'prior_flowclust'). It is invoked by a wrapper function that has the identical function name prefixed with a dot.(e.g. '.prior_flowclust') the preprocessing results are then passed to gating wrapper function through 'pps_res' argument.

'preprocessing_args': the named arguments passed to preprocessing function.

Examples

```
## Not run:
    gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
    plot(gt)
## End(Not run)</pre>
```

Description

get gates saved in fcTree

Usage

```
## S4 method for signature 'fcTree,character'
getGate(obj, y, ...)
```

Arguments

```
obj fcTree

y character node name

... other arguments (not used)
```

```
getNodes,fcTree-method
```

get nodes from fcTree

Description

```
get nodes from fcTree
```

Usage

```
## S4 method for signature 'fcTree'
getNodes(x, y)
```

Arguments

```
x fcTree
```

y character node name

 $\begin{array}{ll} {\tt gh_generate_template} & {\tt generate} \ a \ partially \ complete \ csv \ template \ from \ the \ existing \ gating \\ & {\tt hierarchy} \end{array}$

Description

To ease the process of replicating the existing (usually a manual one) gating schemes, this function populate an empty gating template with the 'alias', 'pop', 'parent' and 'dims' columns that exacted from an GatingHierarchy, and leave the other columns (e.g. 'gating_method') blank. So users can make changes to that template instead of writing from scratch.

Usage

```
gh_generate_template(gh)
```

Arguments

gh

a GatingHierarchy likely parsed from a xml workspace

Value

a gating template in data. frame format that requires further edition after output to csv

Examples

```
library(flowWorkspace)
dataDir <- system.file("extdata",package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual",full = TRUE))
gh_generate_template(gs[[1]])</pre>
```

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```
groupBy,gtMethod-method
```

get the grouping variable for the gating method

Description

When specified, the flow data is grouped by the grouping variable (column names in pData). Within each group, when isCollapse is set to TRUE, the gating method is applied to the collapsed data. Otherwise, it is done indepentently for each individual sample(flowFrame). Grouping variable is also used by preprocessing method.

Usage

```
## S4 method for signature 'gtMethod'
groupBy(object)
```

Arguments

object gtMethod

```
gs_add_gating_method apply a gating method to the GatingSet
```

Description

When interacting with the existing gated data, this function provides an alternative way to interact with the GatingSet by supplying the gating description directly through arguments without the need to write the complete csv gating template.

Usage

```
gs_add_gating_method(
    gs,
    alias = "*",
    pop = "+",
    parent,
    dims = NA,
    gating_method,
    gating_args = NA,
    collapseDataForGating = NA,
    groupBy = NA,
    preprocessing_method = NA,
    preprocessing_args = NA,
    strip_extra_quotes = FALSE,
    ...
)
```

Arguments

gs GatingSet or GatingSetList
alias, pop, parent, dims, gating_method, gating_args, collapseDataForGating, groupBy, preprocessing_
see details in gatingTemplate

strip_extra_quotes

logical Extra quotes are added to strings by fread. This causes problems with parsing R strings to expressions in some cases. Default FALSE for usual behaviour. TRUE should be passed if parsing gating_args fails.

.. other arguments

- mc.cores passed to multicore package for parallel computing
- parallel_type character specifying the parallel type. The valid options are "none", "multicore", "cluster".
- cl cluster object passed to parallel package (when parallel_type is "cluster")

Details

Calls to gs_add_gating_method can also be easily reversed with gs_remove_gating_method. Note, however, that it is not possible to differentiate between different GatingSet objects loaded from the same directory with load_gs within a session. Thus, to guarantee a clean history for gs_remove_gating_method, it is necessary to call gs_add_gating_method_init on the loaded GatingSet immediately after re-loading it. See the documentation for gs_add_gating_method_init for more details. This will not be an issue for GatingSet objects created directly using the constructor.

See Also

```
gs_remove_gating_method gs_add_gating_method_init
```

Examples

```
## Not run:
# add quad gates
gs_add_gating_method(gs, gating_method = "mindensity", dims = "CCR7,CD45RA", parent = "cd4-cd8+", pop = "CCR7+
# polyfunctional gates (boolean combinations of exsiting marginal gates)
gs_add_gating_method(gs, gating_method = "polyFunctions", parent = "cd8", gating_args = "cd8/IFNg:cd8/IL2:cd8.
#boolGate method
gs_add_gating_method(gs, alias = "IL2orIFNg", gating_method = "boolGate", parent = "cd4", gating_args = "cd4/Id
## End(Not run)
```

```
gs_add_gating_method_init
```

 ${\it Clear\ history\ of\ gs_add_gating_method\ } calls\ for\ a\ given\ {\it GatingSet}$ $or\ {\it GatingSetList}$

Description

Repeated calls to the <code>load_gs</code> method in the same session will yield indistinguishable objects that can result in overlapping history of <code>gs_add_gating_method</code> calls. This method allows for the history to be cleared if the user would like to reload the <code>GatingSet</code> and start fresh. Calling <code>gs_add_gating_method_init</code> without an argument will clear the entire <code>gs_add_gating_method</code> history.

Usage

```
gs_add_gating_method_init(gs)
```

Arguments

gs

a GatingSet or GatingSetList. Can be omitted to clean entire gs_add_gating_method history.

Examples

```
## Not run:
# load in a GatingSet
gs <- load_gs(path)
# Add some nodes using gs_add_gating_method
gs_add_gating_method(gs, gating_method = "mindensity", dims = "CCR7,CD45RA", parent = "cd4-cd8+", pop = "CCR7+.
gs_add_gating_method(gs, gating_method = "polyFunctions", parent = "cd8", gating_args = "cd8/IFNg:cd8/IL2:cd8,
# Remove the effect of the last gs_add_gating_method call using gs_remove_gating_method (note that the first ca
gs_remove_gating_method(gs)
# Re-load the GatingSet to start over
gs <- load_gs(path)
# At this point, gs will still see the history of the first gs_add_gating_method call above
# which will cause problems for later calls to gs_remove_gating_method.
# To fix that, just call gs_add_gating_method_init() to start a clean history
gs_add_gating_method_init(gs)
{\tt\#}~{\tt Now~you~can~continue~using~gs\_add\_gating\_method~and~gs\_remove\_gating\_method~from~scratch}
gs_add_gating_method(gs, gating_method = "mindensity", dims = "CCR7,CD45RA", parent = "cd4-cd8+", pop = "CCR7+.
## End(Not run)
```

```
gs_remove_gating_method
```

```
Reverse the action of gating methods applied via gs_add_gating_method
```

Description

This function provides an easy way to remove the gates and nodes created by the most recent call to <code>gs_add_gating_method</code> on the specified <code>GatingSet</code> or <code>GatingSetList</code>, with a separate history being maintained for each such object. <code>gs_remove_gating_method</code> allows for repeated use, effectively serving as a multi-level undo function for <code>gs_add_gating_method</code>.

gtMethod-class 25

Usage

```
gs_remove_gating_method(gs)
```

Arguments

gs

The GatingSet or GatingSetList for which the most recent gs_add_gating_method call should be reversed.

See Also

```
gs_add_gating_method gs_add_gating_method_init
```

Examples

```
## Not run:
# add quad gates
gs_add_gating_method(gs, gating_method = "mindensity", dims = "CCR7,CD45RA", parent = "cd4-cd8+", pop = "CCR7+
# Remove the gates and nodes resulting from that gs_add_gating_method call
gs_remove_gating_method(gs)
## End(Not run)
```

gtMethod-class

A class to represent a gating method.

Description

A gating method object contains the specifics for generating the gates.

Slots

```
name a character specifying the name of the gating method
```

dims a character vector specifying the dimensions (channels or markers) of the gate

args a list specifying the arguments passed to gating function

groupBy a character or integer specifying how to group the data. If character, group the data by the study variables (columns in pData). If integer, group the data by every N samples.

collapse a logical specifying wether to collapse the data within group before gating. it is only valid when groupBy is specified

Examples

```
## Not run:
    gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
    gh_pop_get_gate(gt, '2', '3')
## End(Not run)</pre>
```

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gtPopulation-class	A class to represent a cell population that will be generated by a gating method.
--------------------	---

Description

A class to represent a cell population that will be generated by a gating method.

Slots

```
id numeric unique ID that is consistent with node label of graphNEL in gating templatename character the name of populationalias character the more user friendly name of population
```

Examples

```
## Not run:
    gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
    gt_get_nodes(gt, '2')
## End(Not run)</pre>
```

gtSubsets-class

A class representing a group of cell populations.

Description

It extends gtPopulation class.

gt_gating

Applies a gatingTemplate to a GatingSet.

Description

It loads the gating methods by topological order and applies them to GatingSet.

Usage

```
gt_gating(x, y, ...)
```

gt_get_children 27

Arguments

x a gatingTemplate objecty a GatingSet object

. . .

- start a character that specifies the population (correspoding to 'alias' column in csv template) where the gating process will start from. It is useful to quickly skip some gates and go directly to the target population in the testing run. Default is "root".
- stop.at a character that specifies the population (correspoding to 'alias' column in csv template) where the gating process will stop at. Default is NULL, indicating the end of gating tree.
- keep.helperGatesa logical flag indicating whether to keep the intermediate helper gates that are automatically generated by openCyto. Default is TRUE.
- mc.cores passed to multicore package for parallel computing
- parallel_type character specifying the parallel type. The valid options are "none", "multicore", "cluster".
- cl cluster object passed to parallel package (when parallel_type is "cluster")

Value

Nothing. As the side effect, gates generated by gating methods are saved in GatingSet.

Examples

```
## Not run:
gt <- gatingTemplate(file.path(path, "data/ICStemplate.csv"), "ICS")
gs <- GatingSet(fs) #fs is a flowSet/ncdfFlowSet
gt_gating(gt, gs)
gt_gating(gt, gs, stop.at = "v") #proceed the gating until population 'v'
gt_gating(gt, gs, start = "v") # start from 'v'
gt_gating(gt, gs, parallel_type = "multicore", mc.cores = 8) #parallel gating using multicore
#parallel gating by using cluster
cl1 <- makeCluster (8, type = "MPI")
gt_gating(gt, gs, parallel_type = "cluster", cl = cl1)
stopCluster ( cl1 )
## End(Not run)</pre>
```

gt_get_children

get children nodes

Description

get children nodes

Usage

```
gt_get_children(obj, y)
```

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Arguments

```
obj gatingTemplate
y character parent node path
```

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
gt_get_nodes(gt, "/nonDebris")
gt_get_children(gt, "/nonDebris")
## End(Not run)</pre>
```

gt_get_gate

get gating method from the node

Description

get gating method from the node

Usage

```
gt_get_gate(obj, y, z)
```

Arguments

obj gatingTemplate
y character parent node path
z character child node path

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
gt_get_nodes(gt, only.names = TRUE)
gt_get_nodes(gt, "/nonDebris")
gt_get_children(gt, "/nonDebris")
gt_get_gate(gt, "/nonDebris", "/nonDebris/singlets")
## End(Not run)</pre>
```

gt_get_nodes 29

gt_get_nodes

get nodes from gatingTemplate object

Description

get nodes from gatingTemplate object

Usage

```
gt_get_nodes(
    x,
    y,
    order = c("default", "bfs", "dfs", "tsort"),
    only.names = FALSE
)
```

Arguments

x gatingTemplate

y character node index. When missing, return all the nodes

order character specifying the order of nodes. options are "default", "bfs", "dfs",

"tsort"

only.names logical specifiying whether user wants to get the entire gtPopulation object

or just the name of the population node

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
gt_get_nodes(gt)[1:2]
gt_get_nodes(gt, only.names = TRUE)
gt_get_nodes(gt, "/nonDebris")
## End(Not run)</pre>
```

gt_get_parent

get parent nodes

Description

get parent nodes

Usage

```
gt_get_parent(obj, y, isRef = FALSE)
```

gt_toggle_helpergates

Arguments

obj	gatingTemplate
у	character child node path
isRef	logical whether show the reference node besides the parent node

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
gt_get_nodes(gt, "/nonDebris")
gt_get_parent(gt, "/nonDebris/singlets")
## End(Not run)</pre>
```

 $gt_list_methods$

Print a list of the registered gating methods

Description

Print a list of the registered gating methods

Usage

```
gt_list_methods()
```

Value

Does not return anything. Prints a list of the available gating methods.

```
gt_toggle_helpergates toggle/delete the hidden flag of the helper gates
```

Description

The helper gates are defined as the referred gates in csv template. And all the children of referred gates are also referred gates thus they are considered the helper gates and can usually be hidden to simply the final gating tree.

Usage

```
gt_toggle_helpergates(gt, gs)
gt_get_helpergates(gt, gs)
gt_delete_helpergates(gt, gs)
```

Arguments

gt gatingTemplate object gs GatingSet

Details

Note that delete action is NOT reversible.

Examples

```
## Not run:
gt <- gatingTemplate(gtFile)
#run the gating
gt_gating(gt, gs)
#hide the gates that are not of interest
gt_toggle_helpergates(gt, gs)
#or simply remove them if you are sure they will not be useful in future
gt_delete_helpergates(gt, gs)
## End(Not run)</pre>
```

isCollapse,gtMethod-method

get the flag that determines whether gating method is applied on collapsed data

Description

When TRUE, the flow data(multiple flowFrames) is collapsed into one and the gating method is applied on the collapsed data. Once the gate is generated, it is then replicated and applied to the each single flowFrame.

Usage

```
## S4 method for signature 'gtMethod'
isCollapse(object)
```

Arguments

object gtMethod

Value

logical

```
names, gtMethod-method get gating method name
```

Description

```
get gating method name
```

Usage

```
## S4 method for signature 'gtMethod'
names(x)
```

Arguments

Χ

 ${\tt gtMethod}$

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
gtMthd <- gt_get_gate(gt, "/nonDebris/singlets", "/nonDebris/singlets/lymph")
names(gtMthd)
dims(gtMthd)
parameters(gtMthd)
isCollapse(gtMthd)
groupBy(gtMthd)
gtPop <- gt_get_nodes(gt, "/nonDebris/singlets/lymph/cd3/cd4+cd8-/CD38+")
names(gtPop)
alias(gtPop)
## End(Not run)</pre>
```

```
names,gtPopulation-method
```

get population name

Description

```
get population name
```

Usage

```
## S4 method for signature 'gtPopulation'
names(x)
```

Arguments

Χ

gtPopulation object

openCyto 33

ocRectangleGate-class the class that carries event indices as well

Description

the class that carries event indices as well

ocRectRefGate

 $constructor\ for\ ocRectRefGate$

Description

constructor for ocRectRefGate

Usage

ocRectRefGate(rectGate, boolExprs)

Arguments

 ${\tt rectGate}$

rectangleGate

boolExprs

character boolean expression of reference nodes

 ${\tt ocRectRefGate-class}$

special gate type that mix the rectangleGate with boolean gate

Description

special gate type that mix the rectangleGate with boolean gate

openCyto

Hierarchical Gating Pipeline for flow cytometry data

Description

Hierarchical Gating Pipeline for flow cytometry data.

Details

openCyto is a package designed to facilitate the automated gating methods in sequential way to mimic the manual gating strategy.

Package: openCyto
Type: Package
Version: 1.2.8
Date: 2014-04-10
License: GPL (>= 2)

LazyLoad: yes

Author(s)

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See Also

See gt_gating, gate_flowclust_1d, for an overview of gating functions.

Examples

```
## Not run: gatingTemplate('test.csv')
```

openCyto-deprecated

Deprecated functions in package openCyto.

Description

```
add_pop -> gs_add_gating_method
add_pop_init -> gs_add_gating_method_init
prior_flowClust -> prior_flowclust
templateGen -> gh_generate_template
gate_flowClust_1d -> gate_flowclust_1d
gate_flowClust_2d -> gate_flowclust_2d
quantileGate -> gate_quantile
cytokine -> gate_tail
quadGate.seq -> gate_quad_sequential
quadGate.tmix -> gate_quad_tmix
gating -> gt_gating
getNodes -> gt_get_nodes
getChildren -> gt_get_children
getParent -> gt_get_parent
getGate -> gt_get_gate
listgtMethods -> gt_list_methods
registerPlugins -> register_plugins
remove_pop -> gs_remove_gating_method
tautString -> tautstring
tautStringGate -> gate_tautstring
toggle.helperGates -> gt_toggle_helpergates
get.helperGates -> gt_get_helpergates
delete.helperGates -> gt_delete_helpergates
```

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openCyto.options	Some global options for openCyto See examples for the meaning of	
	these options and how to get/set them.	

Description

Get/set some global options for openCyto

Examples

```
opt <- getOption("openCyto")
#the threshold of minimum cell events required for the gating algorithm to proceed
opt[["gating"]][["minEvents"]]
#to change the threshold
opt[["gating"]][["minEvents"]] <- 100
options(openCyto = opt)

#switch off the validity check flags(Not recommended)
opt[["check.pop"]] <- FALSE
options(openCyto = opt)</pre>
```

```
parameters, gtMethod-method
```

get parameters of the gating method/function

Description

get parameters of the gating method/function

Usage

```
## S4 method for signature 'gtMethod'
parameters(object)
```

Arguments

```
object gtMethod
```

```
plot,fcFilterList,ANY-method plot\;a\;\mathrm{fcFilterList}
```

Description

It is usually called by plot method for fcTree instead of directly by users.

Usage

```
## S4 method for signature 'fcFilterList,ANY'
plot(
    X,
    y,
    samples = NULL,
    posteriors = FALSE,
    xlim = NULL,
    ylim = NULL,
    node = NULL,
    data = NULL,
    breaks = 20,
    lwd = 1,
    ...
)
```

Arguments

```
fcFilterList
Χ
                  character channel name
У
samples
                  character a vector of sample names to be plotted
                  logical indicating whether posteriors should be plotted
posteriors
                  scale settings for x,y axises
xlim, ylim
node
                  character population name associated with the fcFilterList
                  GatingSet object
data
breaks
                  passed to hist
lwd
                  line width
                  other arguments passed to base plot
. . .
```

Examples

```
## Not run:
    env1<-new.env(parent=emptyenv())
    #gt is a gatingTemplate, gs is a GatingSet
    gt_gating(gt,gs,env1) #the flowClust gating results are stored in env1
    plot(env1$fct,"nonDebris",post=T) #plot the priors as well as posteriors for the "nonDebris" gate
## End(Not run)</pre>
```

Description

This provides the priors and posteriors as well as the gates for the purpose of debugging flowClust gating algorithm

Usage

```
## S4 method for signature 'fcTree, character'
plot(x, y, channel = NULL, data = NULL, ...)
```

Arguments

x fcTree
 y character node name in the fcTree
 channel channel.
 data GatingSet that the fcTree is associated with

... other arguments

Description

plot the gating scheme using Rgraphviz

Usage

```
## S4 method for signature 'gatingTemplate,missing' plot(x, y, ...)
```

Arguments

x gatingTemplate object

y either character specifying the root node which can be used to visualize only

the subgraph or missing which display the entire gating scheme

.. other arguments

graphAttr, nodeAttr: graph rendering attributes passed to renderGraph showRef logical: whether to display the reference gates. Sometime it maybe helpful to hide all those reference gates which are not the cell population of interest and used primarily for generating other population nodes.

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
plot(gt) #plot entire tree
plot(gt, "lymph") #only plot the subtree rooted from "lymph"
## End(Not run)</pre>
```

polyFunctions-class

A class to represent a polyFunctions gating method.

Description

It extends boolMethod class and will be expanded to multiple boolMethod object.

```
pop_add.ocRectangleGate
```

bypass the default flowWorkspace:::.addGate

Description

to support adding gate along with indices without loading flow data and computing to support adding rectangleGate yet gating through boolean operations without loading flow data

Usage

```
## S3 method for class 'ocRectangleGate'
pop_add(gate, gh, recompute, ...)
## S3 method for class 'ocRectRefGate'
pop_add(gate, gh, recompute, ...)
```

Arguments

gate ocRectangleGate or logicalFilterResult

gh GatingHierarchy see add in flowWorkspace package

recompute logical see add in flowWorkspace package

... see add in flowWorkspace package

Details

however it is proven that logical indices are too big to be efficiently passed around

```
posteriors, fcFilter, ANY-method get\ posteriors\ from\ a\ fcFilter\ object
```

Description

```
get posteriors from a fcFilter object
```

Usage

```
## S4 method for signature 'fcFilter,ANY'
posteriors(x, y = "missing")
```

Arguments

```
x fcFilter
```

y character or missing that specifiy which channel to look for

Description

get preprocessing method from the node

Usage

```
## S4 method for signature 'gatingTemplate,character'
ppMethod(obj, y, z)
```

Arguments

```
obj gatingTemplate
y character parent node path
z character child node path
```

Examples

```
## Not run:
gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
ppMethod(gt, "/nonDebris/singlets", "/nonDebris/singlets/lymph")
## End(Not run)</pre>
```

ppMethod-class

A class to represent a preprocessing method.

Description

It extends gtMethod class.

Examples

Description

```
apply a ppMethod to the GatingSet
```

Usage

```
## S4 method for signature 'ppMethod, GatingSet' preprocessing(x, y, ...)
```

Arguments

```
x ppMethod
y GatingSet or GatingSetList
... other arguments
```

```
\label{eq:priors} \textit{priors}, \textit{fcFilter}, \textit{ANY-method} \\ \textit{get priors from } a \; \textit{fcFilter object} \\
```

Description

```
get priors from a fcFilter object
```

Usage

```
## S4 method for signature 'fcFilter,ANY'
priors(x, y = "missing")
```

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Arguments

X	fcFilter object
У	character specifying channel name. if missing then extract priors for all the channels

prior_flowclust

Elicits data-driven priors from a flowSet object for specified channels

Description

We elicit data-driven prior parameters from a flowSet object for specified channels. For each sample in the flowSet object, we apply the given prior_method to elicit the priors parameters.

Usage

```
prior_flowclust(
   flow_set,
   channels,
   prior_method = c("kmeans"),
   K = 2,
   nu0 = 4,
   w0 = c(10, 10),
   shrink = 0.000001,
   ...
)
```

Arguments

flow_set a flowSet object

channels a character vector containing the channels in the flowSet from which we elicit

the prior parameters for the Student's t mixture

prior_method the method to elicit the prior parameters

K the number of mixture components to identify

nu0 prior degrees of freedom of the Student's t mixture components.

w0 the number of prior pseudocounts of the Student's t mixture components. (only

the first element is used and the rest is ignored at the moment)

shrink the amount of eigenvalue shrinkage to add in the case the prior covariance ma-

trices are singular. See details.

... Additional arguments passed to the prior elicitation method selected

Details

Currently, we have implemented only two methods. In the case that one channel is given, we use the kernel-density estimator (KDE) approach for each sample to obtain K peaks from which we elicit prior parameters. Otherwise, if more than one channel is specified, we apply K-Means to each of the samples in the flowSet and aggregate the clusters to elicit the prior parameters.

In the rare case that a prior covariance matrix is singular, we shrink the eigenvalues of the matrix slightly to ensure that it is positive definite. For instance, if the flow_set has two samples, this case can occur. The amount of shrinkage is controlled in shrink.

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Value

list of the necessary prior parameters

refGate-class

A class to represent a reference gating method.

Description

It extends gtMethod class.

Slots

refNodes character specifying the reference nodes

register_plugins

Register a gating or preprocessing function with OpenCyto

Description

Function registers a new gating or preprocessing method with openCyto so that it may be used in the csv template.

Usage

```
register_plugins(fun = NA, methodName, dep = NA, ...)
```

Arguments

fun function to be registered

methodName character name of the gating or preprocessing method

dep character name of the library dependency required for the plugin method to

work.

... other arguments type character specifying the type of registering method.

Should be either "gating" or "preprocessing".

Details

The fun argument should be a wrapper function definition for the gating or preprocessing method. Gating method must have formal arguments:

fraflowFrame

pp_res a pre-processing result

xChannel character (optional)

yChannel character (required)

filterId character

... ellipses for the additional parameters.

Preprocessing method must have formal arguments:

fs a flowSet that stores the flow data (could be subgrouped data if groupBy column is defined in the csv template

```
gs a GatingSet
```

gm a gtMethod object that stores the information from gating method

xChannel character (required)

yChannel character (required)

... ellipses for the additional parameters.

The gating function must return a filter (i.e. polygonGate or other instance) from flowCore. The preprocessing can return anything and it will be passed on to the gating function. So it is up to gating function to use and interpret the results of preprocessing. Not all formal parameters need to be used. Additional arguments are passed via the ... and can be processed in the wrapper

Value

logical TRUE if successful and prints a message. FALSE otherwise.

show, boolMethod-method

show method for boolMethod

Description

show method for boolMethod

Usage

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

Arguments

object

boolMethod

```
show,fcFilter-method show method for fcFilter
```

Description

show method for fcFilter

Usage

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

Arguments

object

fcFilter show method for fcFilter

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Description

show method for gatingTemplate

Usage

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

Arguments

object gatingTemplate

show, gtMethod-method show method for gtMethod

Description

show method for gtMethod

Usage

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

Arguments

object gtMethod show method for gtMethod

tautstring

Taut String Density Estimator Gating

Description

The taut string density estimator gating returns 0, 1, or more gates, depending on how many modes it identifies in the data.

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Usage

```
tautstring(sorted_vector, modeprior = 0)

gate_tautstring(
   fr,
   channel,
   gate_range = NULL,
   min = NULL,
   max = NULL,
   filterId = ""
)
```

Arguments

sorted_vector numeric vector of single cell expresion from a single cytometric channel.

modeprior numeric scalar specifying the expected number of modes. Default 0 (autode-

tect). Rarely should this be set by the user.

fr a flowFrame object channel The channel to gate.

gate_range The range to look for a gate, no truncation occurs.

min The min range of the data to truncate the flowFrame

max The max range of the data to truncate the flowFrame

filterId The id / name of the gate.

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