# Package 'openCyto'

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Type Package

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<b>Description</b> This package is designed to facilitate the automated gating methods in sequential way to mimic the manual gating strategy.
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A function to tell wether a gating method is polyFunctions

# Description

A function to tell wether a gating method is polyFunctions

#### Usage

```
.isPolyfunctional(gm)
```

## **Arguments**

gm

an object that extends gtMethod

.prior\_flowClust1d

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

## **Description**

We elicit data-driven prior parameters from a flowSet object for a specified channel. For each sample in the flowSet object, we apply a kernel-density estimator (KDE) and identify its local maxima (peaks). We then aggregate these peaks to elicit a prior parameters for each of K mixture components.

#### Usage

```
.prior_flowClust1d(flow_set, channel, K = NULL, hclust_height = NULL,
 clust_method = c("kmeans", "hclust"), hclust_method = "complete",
 artificial = NULL, nu0 = 4, w0 = 10, adjust = 2, min = -200,
 max = NULL, vague = TRUE)
```

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#### **Arguments**

flow\_set a flowSet object

channel the channel in the flowSet from which we elicit the prior parameters for the

Student's t mixture

K the number of mixture components to identify. By default, this value is NULL

and determined automatically

hclust\_height the height of the hclust tree of peaks, where the should be cut By default, we

use the median of the distances between adjacent peaks. If a value is specified,

we pass it directly to cutree.

clust\_method the method used to cluster peaks together when for prior elicitation. By default,

kmeans is used. However, if K is not specified, hclust will be used instead.

hclust\_method the agglomeration method used in the hierarchical clustering. This value is

passed directly to hclust. Default is complete linkage.

artificial a numeric vector containing prior means for artificial mixture components. The

remaining prior parameters for the artificial components are copied directly from the most informative prior component elicited. If NULL (default), no artificial

prior components are added.

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.

adjust the bandwidth to use in the kernel density estimation. See density for more

information.

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.

vague logical Whether to elicit a vague prior. If TRUE, we first calculate the median of

standard deviations from all flowFrames. Then, we divide the overall standard

deviation by the number of groups to the scale the standard deviation.

#### **Details**

Here, we outline the approach used for prior elicitation. First, we apply a KDE to each sample and extract all of its peaks (local maxima). It is important to note that different samples may have a different number of peaks. Our goal then is to align the peaks before aggregating the information across all samples. To do this, we utilize a technique similar to the peak probability contrasts (PPC) method from Tibshirani et al (2004). Effectively, we apply hierarchical clustering to the peaks from all samples to find clusters of peaks. We compute the sample mean and variance of the peaks within each cluster to elicit the prior means and its hyperprior variance, respectively, for a flowClust mixture component. We elicit the prior variance for each mixture component by first assigning the observations within each sample to the nearest prior mean. Then, we compute the variance of the observations within each cluster. Finally, we average the variances corresponding to each mixture component across all samples in the flowSet object.

Following Tibshirani et al. (2004), we cluster the peaks from each sample using complete-linkage hierarchical clustering. The linkage type can be changed via the hclust\_method argument. This argument is passed directly to hclust.

.prior\_kmeans 5

To cluster the peaks, we must cut the hierarchical tree by selecting either a value for K or by providing a height of the tree to cut. By default, we cut the tree using as the height the median of the distances between adjacent peaks within each sample. This value can be changed via the hclust\_height argument and, if provided, will be passed to cutree. Also, by default, the number of mixture components K is NULL and is ignored. However, if K is provided, then it has priority over hclust\_height and is passed instead directly to cutree.

To ensure that the KDEs are smooth, we recommend that the bandwidth set in the adjust argument be sufficiently large. We have defaulted this value to 2. If the bandwidth is not large enough, the KDE may contain numerous bumps, resulting in erroneous peaks.

#### Value

list of prior parameters

#### References

Tibshirani, R et al. (2004), "Sample classification from protein mass spectrometry, by 'peak probability contrasts'," Bioinformatics, 20, 17, 3034-3044. http://bioinformatics.oxfordjournals.org/content/20/17/3034.

.prior_kmeans	Elicits data-driven priors from a flowSet object for specified channels using the K-Means clustering algorithm

#### **Description**

We elicit data-driven prior parameters from a flowSet object for specified channels. For each sample in the flowSet object, we apply kmeans to obtain K clusters. From each cluster, we determine its centroid and the sample covariance matrix. We then aggregate these two sample moments across all samples for each cluster.

#### Usage

```
.prior_kmeans(flow_set, channels, K, nu0 = 4, w0 = 10, nstart = 10,
   pct = 0.1, min = NULL, max = NULL, ...)
```

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 $\frac{1}{2}$
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.
nstart	number of random starts used by kmeans algorithm

6 boolMethod-class

pct	percentage of randomly selected cells in each flowFrame that is used to elicit the prior parameters. The value should must be greater than 0 and less than or equal to 1.
min	a numeric vector that sets the lower bounds for data filtering. If NULL (default), no truncation is applied.
max	a numeric vector that sets the upper bounds for data filtering. If NULL (default), no truncation is applied.
	Additional arguments passed to kmeans

#### **Details**

Because the cluster labels returned from kmeans are arbitrary, we align the clusters based on the centroids that are closest to a randomly selected reference sample. We apply the Hungarian algorithm implemented using the solve\_LSAP function from the clue package to assist with the alignment.

If each frame within flow\_set has a large number of cells, the computational costs of kmeans can be a burden. We provide the option to randomly select pct, a percentage of the cells from each flow frame to which kmeans is applied.

#### Value

list of flowClust prior parameters

```
alias, gtPopulation-method get\ population\ alias
```

# Description

get population alias

#### Usage

```
## S4 method for signature gtPopulation
alias(object)
```

## **Arguments**

object gtPopulation

boolMethod-class A class to represent a boolean gating method.

## **Description**

It extends refGate class.

dims,gtMethod-method

dims, gtMethod-method get gating method dimensions

# Description

get gating method dimensions

# Usage

```
## S4 method for signature gtMethod
dims(object)
```

# Arguments

object gtMethod

dummyMethod-class

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

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## **Description**

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

fcEllipsoidGate

constuctor for fcEllipsoidGate

## **Description**

constuctor for fcEllipsoidGate

#### Usage

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

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

8 fcFilterList-class

 ${\tt fcEllipsoidGate-class}$  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

Х

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

fcPolygonGate 9

fcPolygonGate constuctor for fcPolygonGate

# Description

 $constuctor\ for\ fc Polygon Gate$ 

## Usage

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

## **Arguments**

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

fcPolygonGate-class

a concrete class that reprents the polygonGate generated by flowClust

# Description

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

fcRectangleGate

constuctor for fcRectangleGate

# Description

constuctor for fcRectangleGate

# Usage

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

```
x a rectangleGate objectpriors a list storing priorsposts a list storing posteriors
```

10 fcTree-class

 $\begin{tabular}{ll} fc Rectangle Gate-class & a concrete class that reprents the rectangle Gate generated by flow-clust \\ \hline \\ Clust \\ \hline \end{tabular}$ 

# 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

flowClust.1d

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

# Description

We cluster the observations in fr into K clusters.

## Usage

```
flowClust.1d(fr, params, filterId = "", K = NULL, trans = 0,
  positive = TRUE, 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, ...)
```

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	numeric indicating whether the Box-Cox transformation parameter is estimated from the data. May take 0 (no estimation), 1 (estimation) or 2 (cluster-specific estimation). NOTE: for the Bayesian version of $flowClust$ , this value cannot be 2.
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: $TRUE$ )
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 Details.
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 provided, any cutpoint below this value will be set to the given minimum value. If NULL (default), there is no minimum cutpoint value.
cutpoint_max	numeric value that sets a maximum thresold for the cutpoint. If a value is provided, any cutpoint above this value will be set to the given maximum value. If NULL (default), there is no maximum cutpoint value.

12 flowClust.2d

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_interv	val
	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
	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**

## 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.

#### Usage

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

flowClust.2d

a flowFrame object

#### **Arguments**

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 numeric indicating whether the Box-Cox transformation parameter is estimated from the data. May take 0 (no estimation), 1 (estimation) or 2 (cluster-specific estimation). NOTE: for the Bayesian version of flowClust, this value cannot plot a logical value indicating if the fitted mixture model should be plotted. By default, no. a numeric vector of length K containing the location of the cluster of interest. target See details. transitional logical value indicating if a transitional gate should be constructed from the target flowClust cluster. By default, no. quantile the contour level of the target cluster from the flowClust fit to construct the translation a numeric value between 0 and 1 used to position a transitional gate if transitional = TRUE. This argument is ignored if transitional = FALSE. See details

transitional\_angle

the angle (in radians) of the transitional gate. 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.

14 gating

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.

#### Value

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

#### **Examples**

```
## Not run:
  gate <- flowClust.2d(fr, xChannel = "FSC-A", xChannel = "SSC-A", K = 3) # fr is a flowFrame
## End(Not run)</pre>
```

gating

Applies gatingTemplate to one GatingSet.

# Description

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

## Usage

```
gating(x, y, ...)
## S4 method for signature gatingTemplate,GatingSet
gating(x, y, env_fct = NULL, ...)
## S4 method for signature gatingTemplate,GatingSetList
gating(x, y, env_fct = NULL, ...)
```

#### **Arguments**

x a gatingTemplate object
y a GatingSet object
env\_fct a environment that contains fcTree object named as 'fct'. If NULL (by default), no fcTree will be constructed. It is currently reserved for the internal debugging.

. . .

• 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.
- 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
    gating(gt, gs)
    gating(gt, gs, stop.at = "v") #proceed the gating until population v
    gating(gt, gs, start = "v") # start from v
    gating(gt, gs, parallel_type = "multicore", mc.cores = 8) #parallel gating using multicore
    #parallel gating by using cluster
    cl1 <- makeCluster (8, type = "MPI")
    gating(gt, gs, parallel_type = "cluster", cl = cl1)
    stopCluster ( cl1 )

## End(Not run)</pre>
```

```
gating, boolMethod, GatingSet-method apply \; a \; {\tt boolMethod} \; to \; the \; {\tt GatingSet}
```

#### **Description**

apply a boolMethod to the GatingSet

## Usage

```
## S4 method for signature boolMethod, GatingSet gating(x, y, \dots)
```

```
x boolMethod
... other arguments
y GatingSet
```

```
{\it gating,gtMethod,GatingSet-method} \\ apply \ a \ {\it gtMethod} \ to \ the \ {\it GatingSet}
```

# Description

```
apply a gtMethod to the GatingSet
```

# Usage

```
## S4 method for signature gtMethod,GatingSet
gating(x, y, ...)
```

## **Arguments**

... other arguments
x gtMethod
y GatingSet

```
{\it gating}, {\it polyFunctions}, {\it GatingSet-method} \\ apply~a~{\it polyFunctions}~{\it gating}~{\it method}~{\it to}~{\it the}~{\it GatingSet}
```

# Description

It generates a batch of boolMethods based on the expression defined in polyFunctions objects. It is a convenience way to generate different boolean combinations of cytokine gates.

## Usage

```
## S4 method for signature polyFunctions, GatingSet gating(x, y, ...)
```

```
x polyFunctions
... other arguments
y GatingSet
```

```
\label{eq:gatingSet-method} apply \, a \; {\tt refGate} \; to \; the \; {\tt GatingSet}
```

## **Description**

```
apply a refGate to the GatingSet
```

## Usage

```
## S4 method for signature refGate,GatingSet
gating(x, y, ...)
```

#### **Arguments**

```
x refGate
... other arguments
y GatingSet
```

 $\begin{array}{ll} {\it gating Template-class} & a \ class \ storing \ the \ gating \ method \ and \ population \ information \ in \ a \\ {\it graph NEL object} \end{array}$ 

# 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", ...)
```

```
x character csv file name
name character the label of the gating template
... other arguments passed to data.table::fread
```

#### **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 'A+/-' or 'A+/-B+/-', which tells the algorithm which side (postive or negative) of 1d gate or which quadrant of 2d gate to be kept. when it is in the form of 'A+/-B+/-', 'A' and 'B' should be the full name (or a substring as long as it is unquiely matched) of either channel or marker of the flow data.

'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.

'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**

#### **Description**

get children nodes

## Usage

```
## S4 method for signature gatingTemplate,character
getChildren(obj, y)
```

## **Arguments**

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

# **Examples**

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

# Description

```
get gates saved in fcTree
```

## Usage

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

```
obj fcTree
y character node name
... other arguments (not used)
```

## **Description**

get gating method from the node

## Usage

```
## S4 method for signature gatingTemplate,character
getGate(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"))
getNodes(gt, only.names = TRUE)
getNodes(gt, "/nonDebris")
getChildren(gt, "/nonDebris")
getGate(gt, "/nonDebris", "/nonDebris/singlets")
## End(Not run)</pre>
```

 ${\tt getNodes,fcTree-method}$ 

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

#### **Description**

```
get nodes from gatingTemplate object
```

#### Usage

```
## S4 method for signature gatingTemplate
getNodes(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 specifying 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"))
getNodes(gt)[1:2]
getNodes(gt, only.names = TRUE)
getNodes(gt, "/nonDebris")
## End(Not run)</pre>
```

## **Description**

get parent nodes

## Usage

```
## S4 method for signature gatingTemplate,character
getParent(obj, y, isRef = FALSE)
```

22 gtMethod-class

#### **Arguments**

obj gatingTemplate

y 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"))
getNodes(gt, "/nonDebris")
getParent(gt, "/nonDebris/singlets")
## End(Not run)</pre>
```

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

gtMethod-class

A class to represent a gating method.

## **Description**

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

gtPopulation-class 23

# 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**

#### 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**

#### **Description**

It extends gtPopulation class.

24 listgtMethods

```
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

listgtMethods

Print a list of the registered gating methods

## **Description**

Print a list of the registered gating methods

## Usage

listgtMethods()

## Value

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

mindensity 25

mindensity	Determines a cutpoint as the minimum point of a kernel density esti- mate between two peaks

#### **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.

#### Usage

```
mindensity(flow_frame, channel, filter_id = "", positive = TRUE,
    pivot = FALSE, gate_range = NULL, min = NULL, max = NULL,
    peaks = NULL, ...)
```

#### **Arguments**

flow_frame	a flowFrame object
channel	TODO
filter_id	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.
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.

#### **Details**

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

names, gtMethod-method get gating method name

# Description

get gating method name

## Usage

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

# Arguments

x gtMethod

## **Examples**

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

openCyto 27

```
names,gtPopulation-method
```

get population name

## **Description**

get population name

## Usage

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

## **Arguments**

,

gtPopulation object

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)

Mike Jiang <wjiang2@fhcrc.org>, John Ramey <jramey@fhcrc.org>, Greg Finak <gfinak@fhcrc.org> Maintainer: Mike Jiang <wjiang2@fhcrc.org>

#### See Also

See gating, flowClust.1d, for an overview of gating functions.

#### **Examples**

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

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

```
{\it plot}, {\it fcFilterList}, {\it ANY-method} \\ {\it plot}~a~{\it 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**

x fcFilterList

y character channel name

samples character a vector of sample names to be plotted

posteriors logical indicating whether posteriors should be plotted

xlim, ylim scale settings for x,y axises

lwd line width

```
breaks passed to hist

data GatingSet object

node character population name associated with the fcFilterList

... other arguments passed to base plot
```

# **Examples**

## **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, ...)
```

X	fcTree
У	character node name in the fcTree
channel	character specifying the channel.
data	${\tt GatingSet}\ that\ the\ {\tt fcTree}\ is\ associated\ with$
	other arguments

30 polyFunctions-class

```
{\it plot}, {\it gating Template}, {\it missing-method} \\ {\it plot the gating scheme}
```

## **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.

```
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 specify which channel to look for

```
\label{eq:ppMethod} ppMethod, gating Template, character-method\\ \textit{get preprocessing method from the node}
```

#### **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**

```
## Not run:
    gt <- gatingTemplate(system.file("extdata/gating_template/tcell.csv",package = "openCyto"))
    ppMethod(gt, 3, 4)
## End(Not run)</pre>
```

```
preprocessing, pp {\tt Method}, {\tt GatingSet-method} \\ apply \ a \ pp {\tt Method} \ to \ the \ {\tt GatingSet}
```

# Description

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

# Usage

```
## S4 method for signature ppMethod, GatingSet preprocessing(x, y, \ldots)
```

# Arguments

```
x ppMethod
```

y GatingSet or GatingSetList

... other arguments

#### **Description**

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

#### Usage

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

# Arguments

fcFilter	object
	fcFilter

y 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 = 1e-06, ...)
```

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 matrices are singular. See details.
	Additional arguments passed to the prior elicitation method selected

34 quadGate.seq

#### **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.

#### Value

list of the necessary prior parameters

quadGate.seq	sequential quadrant gating function

## Description

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

#### Usage

```
quadGate.seq(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

quadGate.tmix 35

	o .	
duad	(aate	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

```
quadGate.tmix(fr, channels, K, usePrior = "yes", prior = list(NA),
  quantile1 = 0.8, quantile3 = 0.8, trans = 0, B = 10, ...)
```

#### **Arguments**

fr	flowFrame
channels	character vector specifies two channels
usePrior	see flowClust.2d
K	see flowClust.2d
prior	see flowClust.2d
trans	see flowClust.2d
В	see flowClust.2d
quantile1	numeric specifies the quantile level (see 'level' in $\mbox{flowClust})$ for the first quadrant $(x\mbox{-}y\mbox{+})$
quantile3	numeric specifies the quantile level see 'level' in flowClust for third quadrant $(x+y-)$
	other arguments passed to flowClust

#### Value

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

refGate-class

A class to represent a reference gating method.

## **Description**

It extends gtMethod class.

# **Slots**

refNodes character specifying the reference nodes

36 registerPlugins

registerPlugins

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

```
registerPlugins(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:

fr a flowFrame

pp\_res a pre-processing result

xChannel character (optional)

yChannel character (required)

filter\_id 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

show,boolMethod-method 37

# 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

Any R object

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

# Description

show method for fcFilter

# Usage

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

# Arguments

object

Any R object

```
show, {\tt gatingTemplate-method} \\ show\ method\ for\ gatingTemplate
```

# Description

show method for gatingTemplate

# Usage

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

# Arguments

object

Any R object

```
{\tt show,gtMethod-method} \quad \textit{show method for gtMethod}
```

# Description

show method for gtMethod

# Usage

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

# Arguments

object

Any R object

tailgate 39

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

# Description

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

# Usage

```
tailgate(fr, channel, filter_id = "", num_peaks = 1, ref_peak = 1,
    strict = TRUE, tol = 0.01, positive = TRUE, side = "right", ...)

cytokine(fr, channel, filter_id = "", num_peaks = 1, ref_peak = 1,
    tol = 0.01, positive = TRUE, side = "right", ...)
```

# Arguments

fr	a flowFrame object
channel	the channel from which the cytokine gate is constructed
filter_id	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.
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 used to construct the cytokine gate from the derivative of the kernel density estimate
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)
side	On which side of the density do we want to gate the tail, the right (default) or left?
	additional arguments.

# Value

a filterList containing the gates (cutpoints) for each sample

# Examples

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

40 templateGen

templateGen	generate a partially complete csv template from the existing gating hierarchy

# 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

```
templateGen(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

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