Package `flowBin`

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**Type** Package

**Title** Combining multitube flow cytometry data by binning

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**Description** Software to combine flow cytometry data that has been multiplexed into multiple tubes with common markers between them, by establishing common bins across tubes in terms of the common markers, then determining expression within each tube for each bin in terms of the tube-specific markers.

**License** Artistic-2.0

**LazyLoad** yes

**Imports** class, limma, snow, BiocGenerics

**Depends** methods, flowCore, flowFP, R (>= 2.10)

**Suggests** parallel

**biocViews** ImmunoOncology, CellBasedAssays, FlowCytometry

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  'checkQNorm.R' 'flowBin.R' 'flowFPBin.R'
  'FlowSample-accessors.R' 'getBinExpr.R' 'kMeansBin.R'
  'mapClustersKNN.R' 'quantileNormalise.R' 'removeSparseBins.R'

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**Description**

Multitube AML sample as example data for flowBin

**Format**

A flowSample containing 7 tubes with 3 common parameters and 4 measure parameters per tube.

**Source**

FlowRepository.org accession FR-FCM-ZZYA

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**BinnedFlowSample**

A FlowSample, but with binning information for each tube

**Description**

clust.labels: list of cluster label vectors, one for each tube
Note: all slots can be get and set using accessor methods, for example `bin.pars(myFlowSet) <- c(1,2,5)`
checkQNorm

Function to check the quantile normalisation of a FlowSample using flowFP

Description
Function to check the quantile normalisation of a FlowSample using flowFP

Details
object and normed.object are compared using flowFP binning, to assess the deviation in bin counts between the two.

Value
list containing two matrices of standard deviations across bins (rows) vs tubes (columns) for before (sd.before) and after (sd.after).

Examples
data(aml.sample)
normed.sample <- quantileNormalise(aml.sample)
qnorm.check <- checkQNorm(aml.sample, normed.sample, do.plot=FALSE)
show(qnorm.check)

eventsInBins

Count number of events for each tube in each bin

Description
Useful for QA of bin mapping

flowBin

function to run the entire flowBin pipeline

Description
Takes a list of flowFrames representing tubes from a single flow cytometry sample, and combines them using binning of events in terms of common markers across tubes.
flowBin(tube.list, bin.pars, control.tubes = vector(),
measure.pars = NULL,
sample.name = "Unnamed Flow Expr Set",
bin.method = "kmeans", expr.method = "medianFI",
sparse.bin.thresh = 0.001, dequantize = T,
snow.cluster = NULL, n.bins = 128, scale.expr = F,
do.qnorm = T, return.bins = F)

Arguments

tube.list a list of flowFrames, one for each tube to combine
bin.pars a numerical vector indicating which flow parameters in the each flowFrame to use for combining tubes. These should be the same markers assayed across every tube.
control.tubes a vector indicating which tubes in tube.list to use for negative controls. May be empty.
measure.pars a list of which parameters to measure expression for, with one vector for each tube. If left NULL, this defaults to all parameters other than those specified as bin.pars
sample.name name of this flowSample, for convenience (defaults to 'Unnamed Flow Expr Set')
bin.method The method to use for creating bins. The two options are "kmeans" for k-means clustering and nearest-neighbour mapping of bins. or "flowFP" for flowFP binning and direct mapping of bin boundaries across tubes.
expr.method The method to use to compute bin expression across tubes. This defaults to MFI of the cells belonging to that bin in each tube. Other options are
sparse.bin.thresh
Bins which contain fewer than this proportion of total events in any tube will be excluded as outliers. Defaults to 0.001
dequantize If TRUE, adds a small (region of 1e-8) value to flow data to help break ties when binning.
snow.cluster A cluster created using the snow package, which flowBin will use to speed up computation. If NULL, flowBin will execute in serial mode.
n.bins Number of bins to use. Note that this must be a power of 2 if flowFP is selected as binning method.
scale.expr If TRUE, the resulting expression values will be scaled to (0,1) using the ranges specified in the flowFrames in tube.list.
do.qnorm If TRUE, the binning markers will be quantile normalized prior to binning.
return.bins If TRUE, return a BinnedFlowExprSet containing the bins themselves as well as the expression for each bin.
flowFPBin

Value

A matrix containing expression values for each bin in terms of each marker across all tubes. If return.bins is set TRUE, then a list containing a BinnedFlowExprSet followed by the expression matrix is returned.

Examples

data(aml.sample)
tube.combined <- flowBin(aml.sample@tube.set,
  bin.pars=aml.sample@bin.pars,
  bin.method='flowFP',
  control.tubes=aml.sample@control.tubes,
  expr.method='medianFIDist', scale.expr=TRUE)
heatmap(tube.combined, scale='none')

Description

Bin sample using flowFP binning

Arguments

- object: flowSample to bin
- n.bins=128: number of bins to use. This should be a power of 2, and will be rounded down to the nearest power of 2 if not.
- snow.cluster=NULL: Optional snow cluster to use for parallel execution.
- dequantize=TRUE: If TRUE, adds a small (region of 1e-8) value to flow data to help break ties when binning.

Value

A BinnedFlowSample

Examples

data(aml.sample)
normed.sample <- quantileNormalise(aml.sample)
res <- flowFPBin(normed.sample)
FlowSample

A class similar to flowSet, but with extra information needed by flow-Bin

**Description**

- **name**: character string - name of the object
- **tube.set**: list of flowFrames containing raw flow data.
- **control.tubes**: Integer vector indicating which tubes in the list (if any) to use as negative controls. May be empty.
- **bin.pars**: Integer vector indicating which parameters to use for binning. These must be in the same position in all tubes.
- **measure.pars**: list of integer vectors indicating which parameters to use for measurement. These must be specified per tube. Note: all slots can be get and set using accessor methods, for example bin.pars(myFlowSet) <- c(1,2,5)

**getBinExpr**

Calculate the expression of each bin in a BinnedFlowSample in terms of the measurement markers

**Description**

getBinExpr main function definition

**Arguments**

- **method**: Method to use to compute expression, passed as a string. Defaults to medianFI, which takes the simple median of each bin, and does not require control tubes. Other options available are medianFIDist, which uses medians with the median of the negative control subtracted out, and propPos which sets a threshold at the 98th percentile of the negative control and determines what proportion of cells lie above that.
- **include.bin.medians**: logical, specifies whether to compute the medians of each bin in terms of the binning markers and include them in the result or not. Defaults to T.
- **scale**: logical specifying whether to scale the results to the interval (0,1). If T (default), then all medians will be divided by the range for that marker as specified in the flowFrame.

**Value**

A numeric matrix containing expression values, with bins as rows and markers as columns
**kMeansBin**

**Bin sample using K-means binning**

**Description**

Bin sample using K-means binning

**Arguments**

- `object`  
  flowSample to bin
- `n.bins=128`  
  number of bins to use. This should be a power of 2, and will be rounded down to the nearest power of 2 if not.
- `n.neighbours=1`  
  number of neighbours to use for KNN mapping of bins from clustered tube
- `snow.cluster=NULL`  
  Optional snow cluster to use for parallel execution.
- `random.seed=101`  
  Random seed to set to make K-means clustering deterministic.
- `dequantize=T`  
  If TRUE, adds a small (region of 1e-8) value to flow data to help break ties when binning.

**Details**

Runs K-means clustering on the binning markers in the first tube of the data set. These clusters are then mapped to the other tubes using K-nearest neighbours.

**Value**

A BinnedFlowSample

**Examples**

```r
data(amlSample)
normed.sample <- quantileNormalise(aml.sample)
binned.sample <- flowFPBin(normed.sample)
binned.sample <- removeSparseBins(binned.sample, 0.001)
bin.expr <- getBinExpr(binned.sample)
heatmap(bin.expr, scale='none')
```
**mapBinsKNN**

*Internal function to map bins by KNN*

**Description**

Internal function to map bins by KNN

**Arguments**

- `object`: flowSample to map the bins of
- `tube.1.labels`: integer vector of bin labels for the events in tube 1
- `n.neighbours`=1: number of neighbours to use for KNN mapping of bins from clustered tube
- `snow.cluster=NULL`: Optional snow cluster to use for parallel execution.
- `dequant=T`: If TRUE, adds a small (region of 1e-8) value to flow data to help break ties when binning.

**Details**

Takes a FlowSample and labels for the events in tube 1, and maps these to all other tubes.

**Value**

a BinnedFlowSample

**Examples**

```r
data(aml.sample)
tube1.expr <- exprs(tube.set(aml.sample)[[1]])
kmeans.res <- kmeans(tube1.expr, 100)
kmeans.labels <- kmeans.res$cluster

#Now create a binnedFlowExprSet using the cluster labels for tube 1
clustered.sample <- mapBinsKNN(aml.sample, kmeans.labels)
sort(table(bin.labels(clustered.sample)[[3]]))
```

---

**quantileNormalise**

*quantileNormalise normalise binning paramaters across all tubes of a flowSample*

**Description**

Since the binning parameters are the same across tubes, and samples each tube is an aliquot from the same sample, these should have the same underlying distribution. Hence, quantile normalisation can be used to force this to be so, removing technical variation.
removeSparseBins

Examples

```r
data(amlSample)
normed.sample <- quantileNormalise(aml.sample)
qnorm.check <- checkQNorm(aml.sample, normed.sample, do.plot=FALSE)
show(qnorm.check)
```

removeSparseBins

Remove bins from a BinnedFlowSample with few events in them

Description

Remove bins from a BinnedFlowSample with few events in them

Arguments

- `object`: the BinnedFlowSample to act on
- `cutoff.prop=NULL`: the minimum proportion that a bin must contain to be kept. If NULL, only bins with no events in at least one tube will be removed.

Details

This is important to do prior to calculating bin expression, as bins containing 2 or less events, for example, cannot have their median computed.

Value

a BinnedFlowSample with sparse bins removed

show-methods

Methods to view flowBin objects

Description

Methods for function `show` in Package flowBin

Methods

- `signature(object = "BinnedFlowExprSet")`: Show number of bins and samples for a BinnedFlowExprSet.
- `signature(object = "CVResult")`: Show various statistics stored in a CVResult object.
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