Package 'flowClean'

September 15, 2024		
Version 1.42.0		
Title flowClean		
Description A quality control tool for flow cytometry data based on compositional data analysis.		
Author Kipper Fletez-Brant		
Maintainer Kipper Fletez-Brant <cafletezbrant@gmail.com></cafletezbrant@gmail.com>		
Depends R ($>= 2.15.0$), flowCore		
Imports bit, changepoint, sfsmisc		
Suggests flowViz, grid, gridExtra		
License Artistic-2.0		
LazyLoad yes		
biocViews FlowCytometry, QualityControl, ImmunoOncology		
NeedsCompilation no		
git_url https://git.bioconductor.org/packages/flowClean		
git_branch RELEASE_3_19		
git_last_commit 12d491a		
git_last_commit_date 2024-04-30		
Repository Bioconductor 3.19		
Date/Publication 2024-09-15		
Contents		
clean		
Index		

2 clean

clean	clean. For cleaning flow cytometry data.	

Description

This function uses compositional data analysis to identify errant collection events.

Usage

```
clean(fF, vectMarkers, filePrefixWithDir, ext, binSize=0.01,
   nCellCutoff=500, announce=TRUE, cutoff="median", diagnostic=FALSE, fcMax=1.3)
```

Arguments

fF flowFrame object containing experimental data to be cleaned.

vectMarkers A vector of indices representing flow parameters to be examined. These are con-

sidered as columns in the data matrix in which cells are rows and parameters are columns. Generally this vector excludes indices for various 'scatter' parameters

(e.g. 'FSC-A')

filePrefixWithDir

A string containing at least the desired name for the output flow file generated.

Can include directory structure and folder ('/' or '\') characters.

ext The file extension for the output flow file.

binSize A number in [0,1]; represents the fraction of duration of collection per bin.

nCellCutoff An integer; represents the minimum number of cells a population must have to

be included in analysis.

cutoff Method for determining threshold for parameter. Can be "median" (default) or

in [0, 1], which is interpreted as a percentile. Integers > 1 will be interpreted as

the fluorescence value to be used for a threshold.

announce Print completion messages.

fcMax Maximum allowable increase relative to presumed 'good' data.

announce If TRUE, will print message to screen if errors detected.

diagnostic If TRUE, will make PNG of populations in time bins, and save with same prefix

as specified in filePrefixWithDir.

returnVector If desired, only return vector indicating if a given cell is 'good' or 'bad'.

nstable The number of stable populations required to be observed during the duration of

an experiment. Default is 5.

Author(s)

Kipper Fletez-Brant

synPerturbed 3

References

Fletez-Brant C, Spidlen J, Brinkman R, Roederer M and Chattopadhyay P. flowClean: Automated identification and removal of fluorescence anomalies in flow cytometry data. Cytometry Part A, 2016.

See Also

The package vignette.

Examples

```
data(synPerturbed)
synPerturbed.c <- clean(synPerturbed, vectMarkers=c(5:17),
  filePrefixWithDir="sampleName", ext="fcs")</pre>
```

synPerturbed

Synthetically Perturbed FCS.

Description

This is a FCS file in which a subset of one parameter was artificially perturbed so as to have a much higher fluorescent intensity than the remainder of the parameter's observations.

Format

A flowFrame with 17 observables and 76466 cells.

Details

Cells during a specific time period had their fluorescent intensities increased on channel < V705-A>.

Examples

data(synPerturbed)

Index