Package 'flowClean'

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Version 1.47.0 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> 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 devel git_last_commit 28921a8 git_last_commit_date 2025-04-15 **Repository** Bioconductor 3.22 Date/Publication 2025-07-18

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Description

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

Usage

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

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.

clean

synPerturbed

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)

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