

# Package ‘flowQ’

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**Type** Package

**Title** Quality control for flow cytometry

**Version** 1.18.0

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**Description** Provides quality control and quality assessment tools for flow cytometry data.

**Depends** R (>= 2.10.0), methods, BiocGenerics (>= 0.1.3), outliers,lattice, flowViz, mvoutlier, bioDist, parody, RColorBrewer,latticeExtra

**Imports** methods, BiocGenerics, geneplotter, flowCore, flowViz, IRanges

**Suggests** flowStats

**SystemRequirements** ImageMagick

**License** Artistic-2.0

**biocViews** Infrastructure, FlowCytometry, CellBasedAssays

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flowQ-package	<i>Quality control for flow cytometry</i>
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## Description

Functions and methods for quality control and QA of flow cytometry data. This package heavily depends on the flowCore package.

## Details

Package:	flowQ
Type:	Package
Version:	0.2.1
Date:	2006-11-16
License:	Artistic

Quality control is an important aspect when dealing with large amounts of complex high-throughput data. This package comprises functionality for efficient QA of flow cytometry data.

## Author(s)

Maintainer: Florian Hahne <f.hahne@dkfz.de> Authors: R. Gentleman, F. Hahne, J. Kettman, N. Le Meur, M. Tang

## References

references go here

**See Also**

[flowCore](#)

**Examples**

```
## examples go here
```

---

aggregatorList-class      *Class "aggregatorList"*

---

**Description**

A list of qaAggregators

**Details**

This class directly extends class "list" and is intended to exclusively hold objects of class qaAggregator, where each list item represents the outcome of a QA subprocess for a single [flowFrame](#). It mainly exists to allow for method dispatch and should never be populated manually; instead, use the constructor qaAggregatorList which checks for valid objects.

**Objects from the Class**

Objects should be created using the constructor:

aggregatorList(...), where ... are objects inheriting from class [qaAggregator](#) or a list of such objects.

**Slots**

.Data: Object of class "list", the list data

**Extends**

Class "[list](#)", from data part. Class "[vector](#)", by class "list", distance 2.

**Methods**

**initialize** signature(.Object = "aggregatorList"): constructor

**show** signature(object = "aggregatorList"): print object details

**Author(s)**

Florian Hahne

**See Also**

[qaGraph](#), [writeQAReport](#), [qaProcess](#), [qaAggregator](#)

**Examples**

```
showClass("aggregatorList")
```

---

binaryAggregator-class    *Class "binaryAggregator"*

---

### Description

Abstraction of a binary type of aggregator with possible states "passed" and "not passed"

### Objects from the Class

Objects can be created by calls of the form `new("binaryAggregator", ...)`, or using the constructor `binaryAggregator(passed)`, where `passed` is a logical scalar.

### Slots

`passed`: Object of class "logical" indicating whether the process has passed the QA requirements

### Extends

Class [qaAggregator](#), directly.

### Methods

**show** signature(object = "binaryAggregator"): print object details

**writeLines** signature(text = "binaryAggregator", con = "file", sep = "missing", use-Bytes="missing"): write to HTML file connection

### Author(s)

Florian Hahne

### See Also

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [discreteAggregator](#), [factorAggregator](#), [numericAggregator](#), [stringAggregator](#), [rangeAggregator](#)

### Examples

```
showClass("binaryAggregator")
```

---

discreteAggregator-class *Class "discreteAggregator"*

---

### Description

Abstraction of a discrete type of aggregator with possible states "passed", "not passed" and "warning"

### Objects from the Class

Objects can be created by calls of the form `new("discreteAggregator", ...)` or using the constructor `discreteAggregator(x)`, where `x` is an integer value in `[0,1,2]` or a factor with levels 0, 1 and 2.

### Slots

`x`: Object of class "factor" One in 0 (not passes), 1 (passed) or 2 (warning)

`passed`: Object of class "logical" indicating whether the process has passed the QA requirements (not evaluated)

### Extends

Class [qaAggregator](#), directly.

### Methods

**show** signature(object = "discreteAggregator"): print object details

**writeLines** signature(text = "discreteAggregator", con = "file", sep = "missing", useBytes="missing"): write to HTML file connection

### Author(s)

Florian Hahne

### See Also

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [binaryAggregator](#), [factorAggregator](#), [numericAggregator](#), [stringAggregator](#), [rangeAggregator](#)

### Examples

```
showClass("discreteAggregator")
```

---

`evaluateProcess`*Evaluate QA processes*

---

**Description**

Re-evaluate an object of class `qaProcess`, e.g. for the case that a threshold value has changed.

**Usage**

```
evaluateProcess(process, thresh, ...)
```

**Arguments**

<code>process</code>	An object of class <code>qaProcess</code> .
<code>thresh</code>	The new threshold on which the process is to be evaluated.
<code>...</code>	Further arguments that are passed on to the individual functions for each QA process type.

**Details**

It is sometimes useful to update the state of aggregators in a `qaProcess`, for instances after changing the threshold value, without having to recompute all images, which can be very time consuming.

**Value**

An updated object of class `qaProcess`

**Note**

This function needs to be extended for new types of `qaProcess`.

**Author(s)**

Florian Hahne

**See Also**

`qaProcess`, `writeQAReport`

**Examples**

```
## Not run:  
data(GvHD)  
dest <- tempdir()  
qp1 <- qaProcess.timeline(GvHD[1:3], channel="FL1-H", outdir=dest,  
  cutoff=1)  
evaluateProcess(qp1, thresh=4)  
  
## End(Not run)
```

---

factorAggregator-class    *Class "factorAggregator"*

---

### Description

Abstraction of a factor type of aggregator with possible states coded by the factor levels

### Objects from the Class

Objects can be created by calls of the form `new("factorAggregator", ...)` or using the constructor `factorAggregator(x, passed)`, where `x` is a factor, or an object which can be coerced to a factor, and `passed` is a logical scalar.

### Slots

`x`: Object of class "factor" coding the outcome state

`passed`: Object of class "logical" indicating whether the process has passed the QA requirements

### Extends

Class "[qaAggregator](#)", directly.

### Methods

**show** signature(object = "factorAggregator"): print object details

**writeLines** signature(text = "factorAggregator", con = "file", sep = "missing", useBytes = "missing"): write to HTML file connection

### Author(s)

Florian Hahne

### See Also

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [discreteAggregator](#), [binaryAggregator](#), [numericAggregator](#), [stringAggregator](#), [rangeAggregator](#),

### Examples

```
showClass("factorAggregator")
```

locateDuplicatedParameters

*Locate Duplicated Parameters*

---

### Description

Identifies dyes that are repeated across multiple flowSets using information provided in the description field of each flowFrame.

### Usage

```
locateDuplicatedParameters(flowList)
```

### Arguments

flowList            A list of [flowSet](#)

### Details

This function is used internally by the qaProcess functions to identify dyes that are repeated across multiple aliquots. The function takes in a list of flowSet and identifies stains that are repeated across multiple flowSets (aliquots). These parameters are expected to have the same distribution across aliquots and can be used for Quality control. The method looks for the information provided in the description field of each flowFrame.

### Value

An object of class [qaProcess](#).

### Author(s)

Nishant Gopalakrishnan

### See Also

[normalizeSets](#)

---

normailzeSets

*Locate Duplicated Parameters*

---

### Description

Aligns up peaks for parameters across multiple flowSets using the [warpSet](#) method from the flowStats package.

### Usage

```
normalizeSets(flowList,dupes,peaks=NULL)
```



**Arguments**

flowList	A list of flowSet
dupes	Stains that are to be aligned across multiple flowSet
peaks	Number of peaks expected in the distribution. If NULL is passed as input, The number of peaks to align are estimated from the distribution

**Details**

This function can be used to normalize dye information across multiple flowSets so that the peaks in the distributions align. The method makes use of information provided in the description field of each flowFrame to align up floursecense nformation from multiple flowSet.

**Value**

A list of objects of class [flowSet](#).

**Author(s)**

Nishant Gopalakrishnan

**See Also**

[locateDuplicatedParameters warpSet](#)

---

numericAggregator-class *Class "numericAggregator"*

---

**Description**

Abstraction of a numeric type of aggregator for which possible states are coded by a numeric value

**Objects from the Class**

Objects can be created by calls of the form `new("numericAggregator", ...)` or using the constructor `numericAggregator(x, passed)`, where x is a numeric scalar, and passed is a logical scalar.

**Slots**

**x:** Object of class "numeric" coding the outcome state.

**passed:** Object of class "logical" indicating whether the process has passed the QA requirements.

**Extends**

Class [qaAggregator](#), directly.

**Methods**

**show** signature(object = "numericAggregator"): print object details

**writeLines** signature(text = "numericAggregator", con = "file", sep = "missing", useBytes = "missing"): write to HTML file connection

**Author(s)**

Florian Hahne

**See Also**

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [discreteAggregator](#), [factorAggregator](#), [binaryAggregator](#), [stringAggregator](#), [rangeAggregator](#)

**Examples**

```
showClass("numericAggregator")
```

---

outlier-class

*Virtual Class "outlier"*

---

**Description**

A class to represent outlier tests

**Objects from the Class**

A virtual Class: No objects may be created from it.

**Slots**

test: Object of class "character" ~~

parameters: Object of class "ANY" ~~

**Methods**

No methods defined with class "outlier" in the signature.

**References**

~put references to the literature/web site here ~

**Examples**

```
## showClass("outlier")
```

---

outlierResult-class      *Class "outlierResult"*

---

**Description**

A class to hold the results of an outlier test

**Objects from the Class**

Objects can be created by calls of the form `new("outlierResult", ...)`. *~~ describe objects here ~~*

**Slots**

`frameId`: Object of class "character" *~~*

`filterDetails`: Object of class "list" *~~*

`test`: Object of class "character" *~~*

`parameters`: Object of class "ANY" *~~*

**Extends**

Class "[outlier](#)", directly.

**Methods**

No methods defined with class "outlierResult" in the signature.

**References**

*~put references to the literature/web site here ~*

**Examples**

```
#showClass("outlierResult")
```

---

outliers-methods      *~~ Methods for Function outliers in Package 'GlobalEnv' ~~*

---

**Description**

*~~ Methods for function outliers in Package 'GlobalEnv' ~~*

**Methods**

`x = "flowSet"` Describe this method here

---

preProcessFlowList	<i>Preprocess a list of flowSets to add empty flowFrames for patients in which certain aliquots are missing</i>
--------------------	---

---

**Description**

The preProcessFlowList function takes in a list of flowSets as inputs and adds empty flowFrames for patients in which data from certain aliquots are missing. The function is to be used to modify datasets to conform to a format that can be utilized by the multiple sample [qaProcess](#) functions.

**Usage**

```
preProcessFlowList(flowList)
```

**Arguments**

flowList            A list of flowSet in which flowFrames for certain patients are missing

**Details**

This function can be used to modify a list of flowSets to fill in empty flowFrames for patients for which samples are missing.

**Value**

A list of objects of class flowSet.

**Author(s)**

Nishant Gopalakrishnan

**See Also**

[locateDuplicatedParameters](#)

---

qaAggregator-class	<i>Abstraction of the possible outcomes of a QA process</i>
--------------------	---

---

**Description**

Virtual parent class for different types of QA aggregators

**Details**

In the context of this package, qaAggregators are objects that hold the outcome of a QA process. Each subclass implements its own writeLine method, which creates the appropriate HTML code for a graphical representation of the object.

**Objects from the Class**

A virtual Class: No objects may be created from it.

**Slots**

**passed:** Object of class "logical" indicating whether the process has passed the QA requirements

**Methods**

No methods defined with class "qaAggregator" in the signature.

**Author(s)**

Florian Hahne

**See Also**

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [binaryAggregator](#), [discreteAggregator](#), [factorAggregator](#), [numericAggregator](#), [stringAggregator](#), [rangeAggregator](#)

**Examples**

```
showClass("qaAggregator")
```

---

 qaGraph-class

*Class "qaGraph"*


---

**Description**

Abstraction of the graphical output created in the cause of a QA process

**Objects from the Class**

Objects should be created using the constructor:

`qaGraph(fileName, imageDir, width, empty=FALSE, pdf=TRUE)`, where `fileName` is a path to an image file, `imageDir` is the destination path for the images and the optional argument `width` is the final width to which the bitmap images are converted. The `pdf` argument controls whether vectorized versions of the image are to be produced. For the special case of an empty object without any images one can use option `empty=TRUE`, in which case the constructor ignores all other arguments.

During object instantiation the image file will be converted, resized and copied if necessary.

**Slots**

**fileNames:** Object of class "character" The paths to the image files, both the vectorized and unvectorized versions

**dimensions:** Object of class "matrix" The dimensions of the image files, both for the vectorized and unvectorized version

**types:** Object of class "character" The file extensions for both versions

**id:** Object of class "character" A unique identifier for the images

**Methods**

**initialize** signature(.Object = "qaGraph"): constructor

**names** signature(x = "qaGraph"): returns the file name of the bitmap version of the image

**show** signature(object = "qaGraph"): print object details

**Author(s)**

Florian Hahne

**See Also**

[qaGraphList](#), [writeQAReport](#), [qaProcess](#)

**Examples**

```
showClass("qaGraph")
```

---

qaGraphList-class	<i>Class "qaGraphList"</i>
-------------------	----------------------------

---

**Description**

A list of [qaGraph](#) objects

**Details**

This class directly extends class "list" and is intended to exclusively hold objects of class [qaGraph](#), where each list item represents the graphical output of a QA subprocess for a single [flowFrame](#). It mainly exists to allow for method dispatch and should never be populated manually; instead, use the constructor `qaGraphList` which makes sure, that all image files are converted into the appropriate types and sizes and copied to the expected file location.

**Objects from the Class**

Objects should be created using the constructor:

`qaGraphList(imageFiles, imageDir, width, pdf=TRUE)`, where `imageFiles` are paths to image files, `imageDir` is the destination path for the images and `width` is the final width to which the bitmap images are converted. `pdf` can be used to toggle the generation of vectorized versions of the images.

**Slots**

.Data: Object of class "list", the list data

**Extends**

Class "[list](#)", from data part. Class "[vector](#)", by class "list", distance 2.

**Methods**

**initialize** signature(.Object = "qaGraphList"): constructor  
**show** signature(object = "qaGraphList"): print object details

**Author(s)**

Florian Hahne

**See Also**

[qaGraph](#), [writeQAReport](#), [qaProcess](#)

**Examples**

```
showClass("qaGraphList")
```

---

 qaProcess-class

*Abstraction of the results of a QA process*


---

**Description**

QA processes create graphical output which can be bundled in a single HTML document. This class stores all information that is needed by [writeQAReport](#) to produce such HTML reports.

**Objects from the Class**

Objects should be created using the constructor functions. See [qaProcess.timeline](#) and [qaProcess.marginevents](#) for details. When writing new QA process functions, the constructors [qaProcessFrame](#) and `qaProcess` should be used. The latter expects the mandatory arguments `id`, `type` and `frameProcesses` and also accepts the optional arguments `name` and `summaryGraph`. See the vignette of this package for details.

**Slots**

**id:** Object of class "character", the objects unique identifier  
**name:** Object of class "character", the name of the process  
**type:** Object of class "character", the type of process  
**frameIDs:** Object of class "character", the identifiers of the [flowSets](#) to which the subprocesses are linked  
**summaryGraph:** Object of class "qaGraph", a graphical summary of the process's outcome  
**frameProcesses:** Object of class "list", more detailed information for each [flowFrame](#)

**Methods**

**initialize** signature(.Object = "qaProcess"): constructor

**Author(s)**

Florian Hahne

**See Also**

[qaGraphList](#), [writeQAReport](#), [qaProcessFrame](#)

**Examples**

```
showClass("qaProcess")
```

---

`qaProcess.2DStatsPlot`    *Creates a QA process for 2D summary statistic comparisons across multiple samples*

---

**Description**

This function takes a list of [flowSet](#) as input and creates all necessary output for a '2DStatsPlot' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```
qaProcess.2DStatsPlot(flowList,dyes=c("FSC-A","SSC-A"),
  outdir="QAReport",thresh=0.25,func=mean,
  sum.dimensions=NULL,det.dimensions= NULL,pdf=TRUE,
  name="2DStats",...)
```

**Arguments**

<code>flowList</code>	A list of <a href="#">flowSet</a>
<code>dyes</code>	Flow parameters to be compared across multiple samples
<code>outdir</code>	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory every time.
<code>thresh</code>	A numeric value between 0 and 1 which is used as a threshold by the outlier detection algorithm
<code>func</code>	Summary statistic function to be applied to each parameter in a <code>flowFrame</code> (mean,median etc)
<code>sum.dimensions</code>	The dimensions of summary plot generated in inches, default is NULL where the dimensions are automatically selected
<code>det.dimensions</code>	the dimensions of each sub image generated by the QA process in inches, default is NULL where the dimensions are automatically selected.
<code>pdf</code>	boolean value determining if pdf files or jpeg images will be produced by the QA process
<code>name</code>	The name of the process used for the headings in the HTML output.
<code>...</code>	Further arguments.

**Details**

QA processes of type '2DStatsPlot' detects differences in the value of a summary statistic such as the mean, median etc across multiple samples.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).



**Value**

An object of class [qaProcess](#).

**Author(s)**

Nishant Gopalakrishnan

**See Also**

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.marginevents](#), [qaProcess.DensityPlot](#), [qaProcess.ECDFPlot](#)

**Examples**

```
data(qData)
dest <- tempdir()
par<-c("FSC-A","SSC-A","CD4","CD8")
resMean <- qaProcess.2DStatsPlot(qData,dyes=par,
                               outdir=dest, func=mean, thresh=0.28,pdf=TRUE)
resMean
```

---

`qaProcess.BoundaryPlot` *Creates a QA process for comparison of the percentage of boundary events for the specified parameters across multiple samples*

---

**Description**

This function takes a list of [flowSet](#) as input and creates all necessary output for a 'BoundaryPlot' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```
qaProcess.BoundaryPlot(flowList,dyes=NULL,
                       outdir="QAReport",cutoff=3,sum.dimensions=NULL,
                       det.dimensions=NULL,pdf=TRUE,name="Boundary",
                       side="both",...)
```

**Arguments**

<code>flowList</code>	A list of <a href="#">flowSet</a>
<code>dyes</code>	Flow parameters to be compared across multiple samples. If set to NULL, all parameters that are duplicated across the list of <a href="#">flowSet</a> are identified and compared.
<code>outdir</code>	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory every time.
<code>cutoff</code>	A numeric value for the percentage of boundary events that is used by the process to identify samples that failed the QA process

sum.dimensions	The dimensions of summary plot generated in inches, default is NULL where the dimensions are automatically selected
det.dimensions	the dimensions of each sub image generated by the QA process in inches, default is NULL where the dimensions are automatically selected.
pdf	boolean value determining if pdf files or jpeg images will be produced by the QA process
name	The name of the process used for the headings in the HTML output.
side	Object of class "character". The margin on which to evaluate the filter. Either upper for the upper margin or lower for the lower margin or both for both margins.
...	Further arguments.

### Details

QA processes of type 'BoundaryPlot' helps identify samples that have a large number of boundary events that must be removed by further gating before detailed analysis of the data.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

### Value

An object of class [qaProcess](#).

### Author(s)

Nishant Gopalakrishnan

### See Also

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.DensityPlot](#), [qaProcess.ECDFPlot](#)

### Examples

```
data(qData)
dest <- tempdir()
resBoundary <- qaProcess.BoundaryPlot(qData, dyes=c("FSC-A", "CD3"),
                                     outdir=dest, cutoff=3, pdf=TRUE)
resBoundary
```

---

qaProcess.cellnumber      *Create QA process of type 'cellnumber'*

---

### Description

This function takes a [flowSet](#) as input and creates all necessary output for a 'cellnumber' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```
qaProcess.cellnumber(set, grouping=NULL, outdir, cFactor=2,
                    absolute.value=NULL, two.sided = FALSE,
                    name="cell number", sum.dimensions=NULL,
                    pdf=TRUE, ...)
```

**Arguments**

set	A <a href="#">flowSet</a> .
grouping	A character vector defining one of the variables in the phenoData of set used as a grouping variable. If this argument is used and if absolute.value is NULL, outlier detection will be performed within groups rather than across all samples.
outdir	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory for all of them.
cFactor	The outlier threshold at which the QA criterion is considered to have failed. This is essentially the factor of standard deviations away from the average number of cells per sample, either in both directions if two.sided=TRUE or only towards smaller event numbers if two.sided=FALSE.
absolute.value	An absolute event count below which the QA criterion is considered to be failed. If this argument is not NULL, cFactor and two.sided are ignored.
two.sided	Perform a two-sided outlier detection, i.e., report both unproportionally high and low event numbers.
name	The name of the process used for the headings in the HTML output.
sum.dimensions	The dimensions of the pdf deviced in inches used for the summary plot.
pdf	Logical indicating whether to create vectorized versions of images for this quality process. This should be set to FALSE if disk space is critical, since the pdf versions of the image consume much more space on the hard drive compared to the bitmap version.
...	Further arguments.

**Details**

QA processes of type 'cellnumber' detect aberations in the number of events per sample. These are either determined dynamically as outliers from the typical distribution of event counts for the whole set, or, if absolute.value is not NULL, by an absolute cutoff value. If there is a natural grouping among the samples, this can be specified using the grouping argument. In this case, the outlier detection will be performed within its respective group for a particular sample.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

**Value**

An object of class [qaProcess](#).

**Author(s)**

Florian Hahne

**See Also**

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.marginevents](#), [qaProcess.timeflow](#), [qaProcess.timeline](#)

**Examples**

```
## Not run:
data(GvHD)
dest <- file.path(tempdir(), "flowQ")
qp <- qaProcess.cellnumber(GvHD, outdir=dest, cFactor=2)
qp

## End(Not run)
```

---

qaProcess.DensityPlot	<i>Creates a QA process for comparisons of density plots across multiple samples</i>
-----------------------	--

---

**Description**

This function takes a list of [flowSet](#) as input and creates all necessary outputs for a 'DensityPlot' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```
qaProcess.DensityPlot(flowList,dyes=NULL,outdir="QAReport",
  alpha=0.05,absolute.value=NULL,sum.dimensions=NULL,
  det.dimensions=NULL, pdf=TRUE,name="Density",...)
```

**Arguments**

flowList	A list of <a href="#">flowSet</a>
dyes	Flow parameters to be compared across multiple samples. If set to NULL, all parameters that are duplicated across the list of <a href="#">flowSet</a> are identified and compared.
outdir	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory every time.
alpha	Outlier mislabeling rate
absolute.value	If provided, the absolute.value is used for the outlier detection and the alpha argument is ignored. default is NULL
sum.dimensions	The dimensions of summary plot generated in inches, default is NULL where the dimensions are automatically selected
det.dimensions	the dimensions of each sub image generated by the QA process in inches, default is NULL where the dimensions are automatically selected.
pdf	boolean value determining if pdf files or jpeg images will be produced by the QA process
name	The name of the process used for the headings in the HTML output.
...	Further arguments.

**Details**

QA processes of type 'DensityPlot' detects differences in the density plots across multiple samples. A summary measure based on the sum of the pairwise KL Distances of density plots is used as a parameter to flag patient panels that are potential outliers.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

**Value**

An object of class [qaProcess](#).

**Author(s)**

Nishant Gopalakrishnan

**See Also**

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.marginevents](#), [qaProcess.ECDFPlot](#), [qaProcess.2DStatsPlot](#)

**Examples**

```
data(qData)
dest <- tempdir()
resDensityFSC <- qaProcess.DensityPlot(qData, dyes=c("FSC-A", "SSC-A"),
  outdir=dest, alpha=0.2, pdf=TRUE)
```

---

qaProcess.ECDFPlot	<i>Creates a QA process for comparisons of ECDF's across multiple samples</i>
--------------------	---

---

**Description**

This function takes a list of [flowSet](#) as input and creates all necessary outputs for an 'ecdfOutliers' type of QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```
qaProcess.ECDFPlot(flowList, dyes=NULL, outdir="QAReport",
  alpha=0.05, absolute.value=NULL, sum.dimensions=NULL,
  det.dimensions=NULL, pdf=TRUE,
  name="ECDF",...)
```



---

qaProcess.KLDistPlot     *Creates a QA process for comparisons of KL Distances across multiple samples*

---

### Description

This function takes a list of [flowSet](#) as input and creates all necessary outputs for an 'KLDistPlot' type of QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

### Usage

```
qaProcess.KLDistPlot(flowList,dyes=NULL,outdir="QAReport",
alpha=0.05, absolute.value=NULL,sum.dimensions=NULL,
det.dimensions=NULL, pdf=TRUE,
name="KLDist",...)
```

### Arguments

flowList	A list of <a href="#">flowSet</a>
dyes	Flow parameters to be compared across multiple samples. If set to NULL, all parameters that are duplicated across the list of <a href="#">flowSet</a> are identified and compared.
outdir	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory every time.
alpha	Outlier mislabeling rate
absolute.value	If provided, the absolute.value is used for the outlier detection and the alpha argument is ignored. default is NULL
sum.dimensions	The dimensions of summary plot generated in inches, default is NULL where the dimensions are automatically selected
det.dimensions	the dimensions of each sub image generated by the QA process in inches, default is NULL where the dimensions are automatically selected.
pdf	boolean value determining if pdf files or jpeg images will be produced by the QA process
name	The name of the process used for the headings in the HTML output.
...	Further arguments.

### Details

QA processes of type 'KLDistPlot' detects differences in the pairwise KL Distances of the flow parameters across multiple samples.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

### Value

An object of class [qaProcess](#).

**Author(s)**

Nishant Gopalakrishnan

**See Also**[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.DensityPlot](#) [qaProcess.2DStatsPlot](#) [qaProcess.BoundaryPlot](#)**Examples**

```

data(qData)
dest <- tempdir()
resKLDist <- qaProcess.KLDistPlot(qData, dyes=c("SSC-A", "FSC-A"),
                               outdir=dest, alpha=0.05, pdf=TRUE)
resKLDist

```

---

qaProcess.marginevents *Create QA process of type 'marginevents'*

---

**Description**

This function takes a [flowSet](#) as input and creates all necessary output for a 'marginevents' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```

qaProcess.marginevents(set, channels=NULL, side="both", grouping=NULL,
                      outdir, cFactor=2, absolute.value=NULL, name="margin events",
                      sum.dimensions=NULL, det.dimensions=c(3,1), pdf=TRUE, ...)

```

**Arguments**

set	A <a href="#">flowSet</a> .
channels	A character vector of channel names for which margin events are to be recorded. Will use all available channels except for the time channel if NULL.
side	Object of class "character". The margin on which to evaluate the filter. Either upper for the upper margin or lower for the lower margin or both for both margins.
grouping	A character vector defining one of the variables in the phenoData of set used as a grouping variable. If this argument is used and if absolute.value is NULL, outlier detection will be performed within groups rather than across all samples.
outdir	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory for all of them.
cFactor	The outlier threshold at which the QA criterion is considered to have failed. This is essentially the factor of standard deviations away from the average number of cells per sample for the respective channel.



absolute.value	An absolute percentage of margin events below which the QA criterion is considered to be failed. If this argument is not NULL, cFactor is ignored.
name	The name of the process used for the headings in the HTML output.
sum.dimensions	The dimensions of summary plot generated in inches, default is NULL where the dimensions are automatically selected
det.dimensions	the dimensions of each sub image generated by the QA process in inches.
pdf	Logical indicating whether to create vectorized versions of images for this quality process. This should be set to FALSE if disk space is critical, since the pdf versions of the image consume much more space on the hard drive compared to the bitmap version. Setting the argument to FALSE will also speed up the process.
...	Further arguments.

### Details

QA processes of type 'marginevents' record the number of events that fall on the margins of the measurement range for each channel. Unproportionally high numbers of such events can indicate problems with the instrument settings or with compensation. These are either determined dynamically as outliers from the typical distribution of margin event percentages for the whole set, or, if absolute.value is not NULL, by an absolute cutoff value. If there is a natural grouping among the samples, this can be specified using the grouping argument. In this case, the outlier detection will be performed within its respective group for a particular sample.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

### Value

An object of class [qaProcess](#).

### Author(s)

Florian Hahne

### See Also

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [qaProcess.timeflow](#), [qaProcess.cellnumber](#)

### Examples

```
## Not run:
data(GvHD)
dest <- tempdir()
qp <- qaProcess.marginevents(GvHD, channels=c("FSC-H", "SSC-H"),
  outdir=dest)
qp

## End(Not run)
```

---

qaProcess.timeflow      *Create QA process of type 'timeflow'*

---

### Description

This function takes a [flowSet](#) as input and creates all necessary output for a 'timeflow' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

### Usage

```
qaProcess.timeflow(set, outdir, cutoff=2, name="time flow",
sum.dimensions=NULL, det.dimensions=c(3,2), pdf=TRUE, ...)
```

### Arguments

set	A <a href="#">flowSet</a> .
outdir	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory for all of them.
cutoff	The threshold at which the QA criterion is considered to have failed. An absolute value in the timeline deviation score as computed by the <a href="#">timeLinePlot</a> function.
name	The name of the process used for the headings in the HTML output.
sum.dimensions, det.dimensions	The dimensions of the pdf device in inches used for the summary and the detailed plots.
pdf	Logical indicating whether to create vectorized versions of images for this quality process. This should be set to FALSE if disk space is critical, since the pdf versions of the image consume much more space on the hard drive compared to the bitmap version.
...	Further arguments.

### Details

QA processes of type 'timeflow' detect disturbances in the flow of cells over time. These indicate problems with the cell suspension, clogging of the needle or similar issues. If the flow rate is too high, the frequency of measuring cell doublets increases.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

### Value

An object of class [qaProcess](#).

### Author(s)

Florian Hahne

### See Also

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.marginevents](#), [qaProcess.timeline](#), [qaProcess.cellnumber](#)

**Examples**

```
## Not run:
data(GvHD)
dest <- tempdir()
qp <- qaProcess.timeflow(GvHD, outdir=dest, cutoff=1)
qp

## End(Not run)
```

---

qaProcess.timeline      *Create QA process of type 'timeline'*

---

**Description**

This function takes a [flowSet](#) as input and creates all necessary output for a 'timeline' type QA process. Objects created by this function can be laid out as HTML using [writeQAReport](#).

**Usage**

```
qaProcess.timeline(set, channels=NULL, outdir, cutoff=1,
  name="time line", sum.dimensions=NULL, det.dimensions=c(3,2),
  pdf=TRUE, ...)
```

**Arguments**

set	A <a href="#">flowSet</a>
channels	A character vector of channel names for which the qaReport is to be produced. Will use all available channels except for the time channel if NULL.
outdir	The directory to which the graphical output is to be saved. If multiple QA processes are to be combined, make sure to use the same directory for all of them.
cutoff	The threshold at which the QA criterion is considered to be failed. An absolute value in the timeline deviation score as computed by the <a href="#">timeLinePlot</a> function.
name	The name of the process used for the headings in the HTML output.
sum.dimensions, det.dimensions	The dimensions of the pdf device in inches used for the summary and the detailed plots.
pdf	Logical indicating whether to create vectorized versions of images for this quality process. This should be set to FALSE if disk space is critical, since the pdf versions of the image consume much more space on the hard drive compared to the bitmap version.
...	Further arguments.

**Details**

QA processes of type 'timeline' detect unusual patterns in the acquisition of fluorescence and light scatter measurements over time. These are detected dynamically by identifying trends in the signal intensity over time or local changes in the measurement intensities. The underlying hypothesis is that measurement values are acquired randomly, hence there shouldn't be any correlation to time.

For more details on how to layout [qaProcess](#) objects to HTML, see [writeQAReport](#) and [qaReport](#).

**Value**

An object of class [qaProcess](#).

**Author(s)**

Florian Hahne

**See Also**

[writeQAReport](#), [qaReport](#), [qaProcess](#), [qaProcess.marginevents](#) [qaProcess.timeflow](#), [qaProcess.cellnumber](#)

**Examples**

```
## Not run:
data(GvHD)
GvHD <- transform(GvHD, "FL1-H"=asinh('FL1-H'), "FL2-H"=asinh('FL2-H'))
dest <- tempdir()
qp <- qaProcess.timeline(GvHD, channel="FL1-H", outdir=dest, cutoff=1)
qp

## End(Not run)
```

---

qaProcessFrame-class    *Class "qaProcessFrame"*

---

**Description**

Abstraction of subitems within a qQA process

**Details**

This class bundles graphs and aggregators for a single [flowFrame](#). This allows to create processes with subcomponents, where each item in the `frameAggregators` and `frameGraphs` lists corresponds to one subprocess, which can be used, for instance, to create individual plots for each flow channel. For QA processes without subcomponents, these slots would simply not be populated.

## Objects from the Class

Objects should be created using the constructor:

`qaProcessFrame(frameID, summaryAggregator, summaryGraph, frameAggregators, frameGraphs, details)`  
where `frameID` is the ID of the flowFrame the process is linked to, `summaryAggregator` is an object inheriting from class `qaAggregator` which summarizes the outcome, `summaryGraph` is an object of class `qaGraph` which is the overview graph of the process for the whole frame, `details` is a list containing any additional information regarding the QA process, `frameAggregators` is an object of class `aggregatorList` and `frameGraphs` is an object of class `qaGraphList`. The latter two are the collections of aggregators and graphs for each subprocess. Only `frameID` and `summaryAggregator` are mandatory arguments.

## Slots

`id`: Object of class "character", a unique ID of the object

`frameID`: Object of class "character", ID of the flowFrame the process is linked to

`summaryAggregator`: Object of class "qaAggregator", an aggregator summarizing the output of the process

`summaryGraph`: Object of class "qaGraph", a graphical summary of the process

`frameAggregators`: Object of class "aggregatorList" a list of aggregators for the subprocesses

`frameGraphs`: Object of class "qaGraphList" a list of graphical summaries for the subprocesses

`details`: A list for any additional information

## Methods

**initialize** `signature(.Object = "qaProcessFrame")`: constructor

## Author(s)

Florian Hahne

## See Also

[qaGraphList](#), [writeQAReport](#), [qaProcess](#)

## Examples

```
showClass("qaProcessFrame")
```

---

qaProcessSummary	<i>Class "qaProcessSummary"</i>
------------------	---------------------------------

---

## Description

An internal class to represent QA summaries

## Objects from the Class

The class is internal and not ment for interactive use.

**Slots**

panels: Object of class "list"  
 ranges: Object of class "matrix"  
 summary: Object of class "list"  
 mapping: Object of class "list"  
 pnames: Object of class "character"  
 overallSum: Object of class "matrix"

**Methods**

**writeLines** signature(text = "qaGraphList", con = "file"): HTML output  
**show** signature(object = "qaGraphList"): print object details

**Author(s)**

Florian Hahne

---

 qaReport

---

*Create HTML report using one or several QA process function(s)*


---

**Description**

This function combines all graphical output of multiple QA process functions for one [flowSet](#) in a single hyperlinked HTML document.

**Usage**

```
qaReport(set, qaFunctions, outdir = "./qaReport",
          argLists, grouping = NULL, ...)
```

**Arguments**

set	A <a href="#">flowSet</a>
qaFunctions	A character vector of the names of QA process functions to be used
outdir	The directory to which the HTML report is to be saved.
argLists	lists of argument lists for each of the QA process functions specified via qaFunctions
grouping	A character scalar indicating a variable in the <a href="#">flowSet</a> 's phenoData that is used as a grouping factor in the output.
...	Further arguments that are passed on to <a href="#">writeQAReport</a> .

**Details**

This is a simple convenience function to produce HTML QA reports for a single [flowSet](#) given a list of QA process functions. For more fine-grained control use function [writeQAReport](#) directly.

An entry point to the output of this function can be found at outdir/index.html.

**Value**

The function is called for its side effects

**Author(s)**

Florian Hahne

**See Also**

[qaProcess.marginevents](#), [writeQAReport](#), [qaProcess](#), [qaProcess.timeline](#)

**Examples**

```
## Not run:
data(GvHD)
GvHD <- transform(GvHD, "FL1-H"=asinh('FL1-H'), "FL2-H"=asinh('FL2-H'))
dest <- tempdir()
qaReport(GvHD, c("qaProcess.timeline", "qaProcess.marginevents"), dest,
  list(list(channel="FL1-H", cutoff=1), list(channels=c("FL1-H",
    "FL2-H"), cFactor=4)))
browseURL(file.path(dest, "index.html"))

## End(Not run)
```

---

qData

*Data for demonstrating QA processes that compares dye parameters from each patient across multiple aliquots*

---

**Description**

A list of 8 flowSet. Each flowSet has data from four patients and each element in the list corresponds to an aliquot.

**Usage**

```
data(qData)
```

**Format**

The object contains a list of 8 objects of class flowSet, each composed of 4 flowFrames. Each flowFrame corresponds to a sample from an aliquot.

**Details**

This qData dataset contains fluorescence information for samples from four patients. The sample from each patient was separated into eight aliquots and each aliquot was stained using a different combination of dyes. For each patient, some dyes appear more than once in multiple aliquots. These along with the Forward and Side scatter information can be used for Quality control.

---

rangeAggregator-class    *Class "rangeAggregator"*

---

### Description

Abstraction of a range type of aggregator where possible states are within certain ranges (e.g. percentages)

### Objects from the Class

Objects can be created by calls of the form `new("rangeAggregator", ...)` or using the constructor `rangeAggregator(x, min, max, passed)`, where `x`, `min` and `max` are numeric scalars, with `x` in the range of `[min, max]`, and `passed` is a logical scalar.

### Slots

`min`: Object of class "numeric", the range minimum

`max`: Object of class "numeric", the range maximum

`x`: Object of class "numeric", the value within the range

`passed`: Object of class "logical" indicating whether the process has passed the QA requirements

### Extends

Class `numericAggregator`, directly. Class `qaAggregator`, by class "numericAggregator", distance 2.

### Methods

**show** signature(object = "rangeAggregator"): print object details

**writeLines** signature(text = "rangeAggregator", con = "file", sep = "missing", useBytes = "missing"): write to HTML file connection

### Author(s)

Florian Hahne

### See Also

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [discreteAggregator](#), [factorAggregator](#), [numericAggregator](#), [stringAggregator](#), [binaryAggregator](#)

### Examples

```
showClass("rangeAggregator")
```



---

stringAggregator-class    *Class "stringAggregator"*

---

### Description

Abstraction of a string type of aggregator for which possible states are indicated through a textual description

### Objects from the Class

Objects can be created by calls of the form `new("stringAggregator", ...)` or using the constructor `stringAggregator(x, passed)`, where `x` is a character scalar, and `passed` is a logical scalar.

### Slots

`x`: Object of class "character" which is a textual description of the outcome

`passed`: Object of class "logical" indicating whether the process has passed the QA requirements

### Extends

Class "[qaAggregator](#)", directly.

### Methods

**show** signature(object = "stringAggregator"): print object details

**writeLines** signature(text = "stringAggregator", con = "file", sep = "missing", useBytes = "missing"): write to HTML file connection

### Author(s)

Florian Hahne

### See Also

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [discreteAggregator](#), [factorAggregator](#), [numericAggregator](#), [binaryAggregator](#), [rangeAggregator](#)

### Examples

```
showClass("stringAggregator")
```

---

txtFormatQAObject	<i>Formats results contained in QA process objects into a data frame with appropriate row and column names</i>
-------------------	--

---

## Description

Formats information contained in QA process objects into a data frame which can then be easily written to a tab delimited file. Information contained includes the values used in the outlier detection, information whether the particular parameter passed the outlier test and the summary aggregator information.

## Usage

```
txtFormatQAObject(qp)
```

## Arguments

qp                    A qaProcess object

## Value

A data.frame containing the values used in outlier detection, thresholds used, information regarding whether the parameter passed/failed the test as well as the summary aggregator information

## Author(s)

Nishant Gopalakrishnan

## See Also

[writeQATextReport](#)

## Examples

```
data(qData)
p1 <- qaProcess.marginevents(qData[[1]], channels=c("FSC-A", "SSC-A"),
  outdir=tempdir(), name=paste("Panel", 1, sep="_"))
res <- txtFormatQAObject(p1)
res
```

---

validProcess	<i>Validate a QAProcess object</i>
--------------	------------------------------------

---

**Description**

Check for integrity and existence of files specified in a [qaProcess](#) object.

**Usage**

```
validProcess(object)
```

**Arguments**

object            A [qaProcess](#) object

**Value**

The function is called for its side effects.

**Author(s)**

Florian Hahne

**See Also**

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#), [writeQAReport](#)

---

writeQAReport	<i>Create HTML report for (lists of) qaProcess objects</i>
---------------	--

---

**Description**

This function combines all graphical output of multiple QA processes for one or several [flowSets](#) in a single hyperlinked HTML document.

**Usage**

```
writeQAReport(set, processes=NULL, globalProcess=NULL,  
              outdir = "./qaReport", grouping = NULL, pagebreaks=TRUE, pdf=TRUE)
```

**Arguments**

set	A <a href="#">flowSet</a> or a list of several <a href="#">flowSets</a> .
processes	A list of <a href="#">qaProcess</a> objects or, in the case of multiple <a href="#">flowSets</a> , a list of lists of <a href="#">qaProcess</a> objects. See below for further details.
globalProcess	A list of <a href="#">qaProcess</a> objects generated by multipanel comparison <a href="#">qaProcess</a> functions, when set is a list of <a href="#">flowSets</a> .
outdir	The directory to which the HTML report is to be saved. Each <a href="#">qaProcess</a> object should have been created in the same directory.
grouping	A character scalar indicating a variable in the <a href="#">flowSet</a> 's <code>phenoData</code> that is used as a grouping factor in the output.
pagebreaks	A logical indicating whether the output should be on one long page, or split over several pages.
pdf	A logical indicating whether vectorized versions of the images should be included. Setting this to TRUE implies that the individual QA processes have also been created with setting <code>pdf=TRUE</code> .

**Details**

Both the information about graphical output generated as part of a QA process as well as the quantitative or qualitative results are stored in objects of class [qaProcess](#). The creation of such objects is abstracted in dedicated constructor function and the user should call these functions directly rather than creating [qaProcess](#) manually. [writeQAReport](#) takes lists of such objects and combines their information in a unified HTML document. A grouping factor can be specified to indicate subgroups of the data. In the case of multiple panels, a list of [flowSets](#) can be given to [writeQAReport](#), and the function expects a list of lists of processes, where each process list is specific to one panel.

An entry point to the output of this function can be found at `outdir/index.html`.

**Value**

The function is mostly called for its side effects, that is, the generation of a HTML quality report in `outdir`. A URL to an entry point for browsing of the report is returned.

**Author(s)**

Florian Hahne

**See Also**

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#)

**Examples**

```
## Not run:
data(GvHD)
GvHD <- transform(GvHD, "FL1-H"=asinh('FL1-H'), "FL2-H"=asinh('FL2-H'))
dest <- tempdir()
qp1 <- qaProcess.timeline(GvHD, channel="FL1-H", outdir=dest, cutoff=1)
qp2 <- qaProcess.marginevents(GvHD, channels=c("FL1-H", "FL2-H"),
  outdir=dest, cFactor=4)
url <- writeQAReport(GvHD, processes=list(qp1, qp2), outdir=dest,
  grouping="Patient")
browseURL(url)
```

```
### End(Not run)
```

---

writeQATextReport      *Create tab delimited text files for (lists of) qaProcess objects*

---

## Description

This function combines all the results of multiple QA processes for one or several [flowSets](#) in a single tab delimited text document.

## Usage

```
writeQATextReport(set, processes=NULL, globalProcess=NULL,  
                  fileName="textReport.txt")
```

## Arguments

set	A <a href="#">flowSet</a> or a list of several <a href="#">flowSets</a> .
processes	A list of <a href="#">qaProcess</a> objects or, in the case of multiple <a href="#">flowSets</a> , a list of lists of <a href="#">qaProcess</a> objects. See below for further details.
globalProcess	A list of <a href="#">qaProcess</a> objects generated by multipanel comparison <a href="#">qaProcess</a> functions, when set is a list of <a href="#">flowSets</a> .
fileName	A character vector for the name of the file.

## Details

Quantitative or qualitative results are stored in objects of class [qaProcess](#). [writeQATextReport](#) takes lists of such objects and combines their information in a tab delimited text document. In the case of multiple panels, a list of [flowSets](#) can be given to [writeQATextReport](#), and the function expects a list of lists of processes, where each process list is specific to one panel.

## Value

The function is mostly called for its side effects, that is, the generation of a tab delimited text file. .

## Author(s)

Nishant Gopalakrishnan

## See Also

[qaProcess.marginevents](#), [qaReport](#), [qaProcess](#), [qaProcess.timeline](#)

**Examples**

```
## Not run:  
data(qData)  
p1 <- qaProcess.marginevents(qData[[1]],  
  channels=c("FSC-A", "SSC-A"), outdir=tempdir(),  
  name=paste("Panel",1,sep="_"))  
writeQATextReport(qData[[1]], processes=list(p1), fileName="test.txt")  
  
## End(Not run)
```

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