# QUALIFIER: Quality Assessment of Gated Flow Cytometry Data

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#### Abstract

Background The current flowQ package does the quality assessment on the ungated FCM data. However, there is a need to identify deviant samples by monitoring the consistencies of the underlying statistical properties of different gated cell populations (such as white blood cells, lymphacytes, monocytes etc). The current package was also not designed for dealing with large datasets. To meet these needs, We developed QUALIFIER package using the gating template created in flowJo and performing QA checks on different gated populations. It divides the data preprocessing from the actual outlier detection process so that the statistics are calcuated all at once and the outlier detection and visualization can be done more efficiently and interactively. ncdfFlow is used to solve the memory limit issue for large datasets.

**keywords** Flow cytometry, Quality Assessment, high throughput, svg, flowWorkspace, ncdfFlowSet

# 1 Parsing the QA gating template

parseWorkspace function from flowWorkspace package is used to parse the flowJo workspace. Since we are only interested in the gating strategy (gates defined in the template), execute can be set as FALSE to skip the actual gating process. useInternal is set as TRUE to use internal structure (c++) for faster parsing.

- > ws<-openWorkspace("./data/QA\_MFI\_RBC\_bounary\_eventsV3.xml")##should replace with your own x
- > GT<-parseWorkspace(ws,execute=FALSE,useInternal=TRUE)

# 2 Apply the gating template to the data

Then gating hierarchy gh\_template containing the actual template gates is extracted from the result GatingSet "GT".

> gh\_template<-GT[[1]]

The GatingSet method here is the contructor that takes a gatingHierarchy object as the template and a list of FCS file names (filenames) that need the quality assurance. The result "G" is the GatingSet that contains the gated data and some of the cell population statistics that can be viewed by getPopStats method.

```
> ##datapath is the path where FCS files stores
> G<-GatingSet(gh_template,filenames,path="./data")
> getPopStats(G[[1]])
```

Optionally, isNcdf can be set to TRUE to support netCDF storage of large flow datasets when memory resource is limited.

## 3 Calculating the statistics

After flow data is gated, the statistics of the gated data need to be extracted and saved before the QA checks. db is an environment that serves as a container storing all the QA data and results. Firsly, initDB function initializes and creates the data structures for QA. Then qaPreprocess is a convenient wrapper that calls underlining routines (getQAStats,saveToDB) to calculates/extracts statistics of gated cell populations and save them along with the gating set and the FCS meta data. metaFile is a csv file that contains the extra annoations about each FCS file.It should at least have one column speficifying FCS file names.

fcs.colname and date.colname are the arguments to identify the columns in the meta data (the flat table stored as a csv file) that specifies the FCS filenames and the dates (to be formatted as to "%m/%d/%y"). Once the preprocessing steps are finished, the data are ready for quality assessments.

# 4 Defining qaTasks

read.qaTask reads external csv spreadsheet that contains the descriptions of each QA task and creates a list of qaTask objects.

```
> checkListFile<-file.path(system.file("data",package="QUALIFIER"),"qaCheckList.csv.gz")
> qaTask.list<-read.qaTask(db,checkListFile=checkListFile)

[1] "7 qaTask created and saved in db!"
> qaTask.list[1:2]

$MFIOverTime
qaTask: MFIOverTime
Level : Assay
Description : Fluorescence stability over time
```

population: MFI

Default formula :MFI ~ RecdDt | channel \* stain

<environment: 0x7f2e744cb198>

Plot type: xyplot

\$horiz
[1] FALSE

\$scales

\$scales\$format
[1] "%m/%d/%y"

\$NumberOfEvents

qaTask: NumberOfEvents

Level : Tube

Description: Number of Events Collected

population: root

Default formula :count ~ RecdDt | Tube

<environment: 0x7f2e7446c6c0>

Plot type: xyplot

\$horiz
[1] FALSE

\$scales

\$scales\$format
[1] "%m/%d/%y"

The qaTask can also be created individually by the contructor makeQaTask.

# 5 Quality assessment and visualization

qaCheck and plot are the two main methods to perform the quality assessment and visualize the QA results. They both use the information stored in qaTask object and the formula, which is given either explicitly by the argument or implicitly by the qaTask object. It is generally of the form  $y \sim x \mid g1 * g2 * \dots$ , where y is the statistics to be checked in this QA and must be one of the four types:

"MFI": Median Fluorescence Intensity of the cell population specified by qaTask,

"proportion": the percentage of the cell population specified by qaTask in the parent population,

"count": the number of events of the cell population specified by qaTask,

"spike": the variance of intensity over time of each channel, which indicates the stability of the fluorescence intensity.

x specifies the variable plotted on x-axis (such as date) in the plot method.

g1, g2, ... are the conditioning variables, which divide the data into subgroups and apply the outlier detection whitin each individual group or plot the subgroups in different panels. They may also be omitted, in which case the outlier detection is performed in the entire dataset.

For example, RBC Lysis efficiency (percentage of WBC population) check is defined by qaTask.

#### > qaTask.list[["RBCLysis"]]

qaTask: RBCLysis
Level : Tube

Description : Sufficient RBC lysis

population: WBC\_perct

Default formula :proportion ~ RecdDt | Tube

<environment: 0x7f2e74308900>

Plot type: xyplot

\$horiz
[1] FALSE

\$scales
\$scales\$format
[1] "%m/%d/%y"

According to the formula stored in qaTask, it uses the statistical property "proportion" and groups the data by "Tube" (or staining panel). "RecdDt" is reserved for plotting purpose. Cell population is defined as "WBC\_perct"

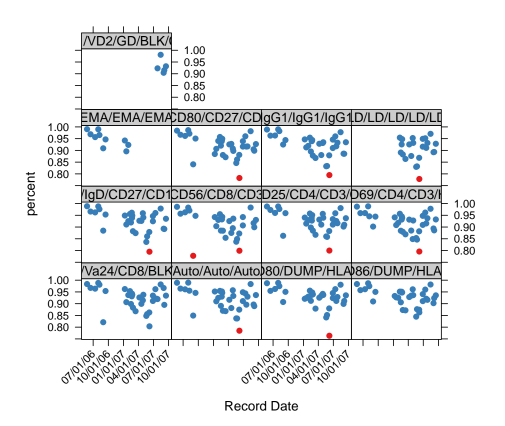
```
> qaCheck(qaTask.list[["RBCLysis"]],outlierfunc=outlier.cutoff,1Bound=0.8)
```

qaCheck reads all the necessary information about the gated data from qaTask object. Users only needs to specificy how the outliers are called. This is done by providing an outlier detection function outlierfunc that takes a numeric vector as input and returns a logical vector as the output. Here "outlier.cutoff" is used and threshold "lBound" is specified ("less than", use uBound for "larger than").

After the outliers are called, the results can be plotted by plot method.

> plot(qaTask.list[["RBCLysis"]],xlab="Record Date",ylab="percent")

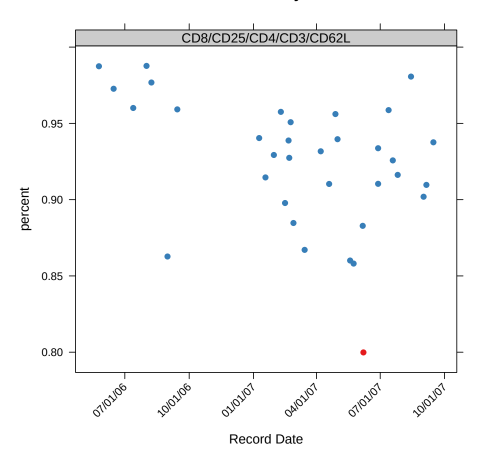
## **Sufficient RBC lysis**



By default all the data are plotted, argument "subset" can be used to visualize a small subset.

> plot(qaTask.list[["RBCLysis"]],subset=Tube=='CD8/CD25/CD4/CD3/CD62L',xlab="Record Date",yla

## **Sufficient RBC lysis**

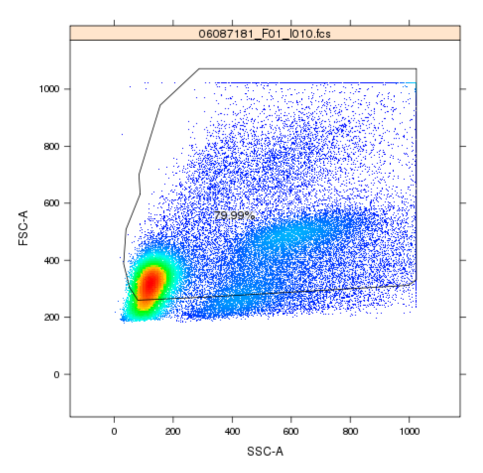


 ${\tt clearCheck}$  is the method to removes the outlier results detected by the previous  ${\tt qaCheck}$  call for the specific  ${\tt qaTask}$ .

> clearCheck(qaTask.list[["RBCLysis"]])

With scatterPlot flag set as true and subset properly specified plot method can generate scatter plots for the selected FCS files,

> plot(qaTask.list[["RBCLysis"]],subset=name=='06087181\_F01\_I010.fcs',scatterPlot=TRUE)



x term in the formula is normally ignored in qaCheck. However, when "plotType" of the qaTask is "bwplot", it is used as the conditioning variable that divides the data into subgroups, within which the outlierfunc is applied.

#### > qaTask.list[["MNC"]]

qaTask: MNC
Level : Assay

Description : Consistency of Lymphocyte/MNC Gate

population: MNC

Default formula :proportion ~ coresampleid

<environment: 0x7f2e74409e38>

Plot type: bwplot

\$horiz
[1] FALSE

\$scales

\$scales\$format

[1] "%m/%d/%y"

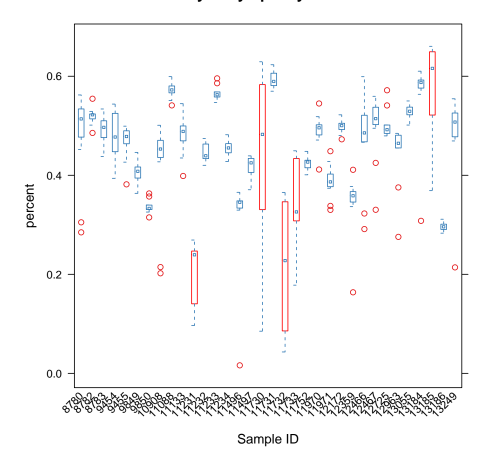
This qaTask detects the significant variance of MNC cell population percentage among aliquots, which have the same "coresampleid". Plot type of this object tells the method to group data by "coresampleid".

> qaCheck(qaTask.list[["MNC"]],z.cutoff=1.5)

Interquartile Range based outlier detection function is used to detect outliers

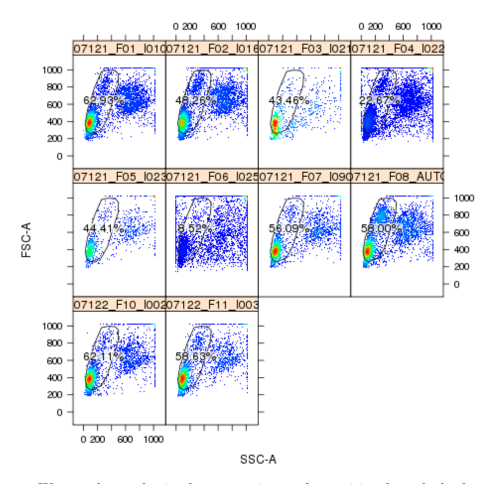
> plot(qaTask.list[["MNC"]],proportion~factor(coresampleid),xlab="Sample ID",ylab="percent")

#### Consistency of Lymphocyte/MNC Gate



The red circles in the boxplot indicate the possible outlier samples and the box of red color indicates the entire sample group has significant variance and is marked as the group outlier. The formula supplied here in the plot method overwrites the one stored in the qaTask object, thus change the way of viewing the data.

Again, with scatterPlot and subset arguments, scatter plots can be generated for the selected FCS files or sample groups,



We can also apply simple aggregation to the statistics through the formula.

#### > qaTask.list[["BoundaryEvents"]]

qaTask: BoundaryEvents

Level : Channel

Description : Off-scale Boundary Events

population: margin

Default formula :proportion ~ RecdDt | channel

<environment: 0x7f2e74206318>

Plot type: xyplot

\$horiz
[1] FALSE

\$scales

\$scales\$format

[1] "%m/%d/%y"

Here the default formula only extracts the "proportion" from each individual channel. In order to check the total percentage of boundary events of all channels for each fcs file, we can

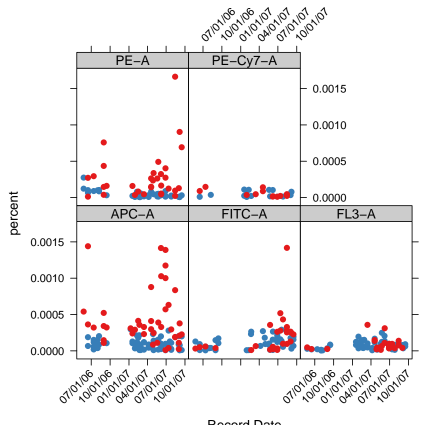
write a new formula by applying aggregation function "sum" to "proportion" and group the data by fcs file ("name" in this case).

```
> qaCheck(qaTask.list[["BoundaryEvents"]]
                  ,sum(proportion) ~ RecdDt | name
                  ,outlierfunc=outlier.cutoff
                  ,uBound=0.0003
```

And we still can visualize the results chanel by chanel.

> plot(qaTask.list[["BoundaryEvents"]],proportion ~ RecdDt | channel,xlab="Record Date",ylab=

## **Off-scale Boundary Events**

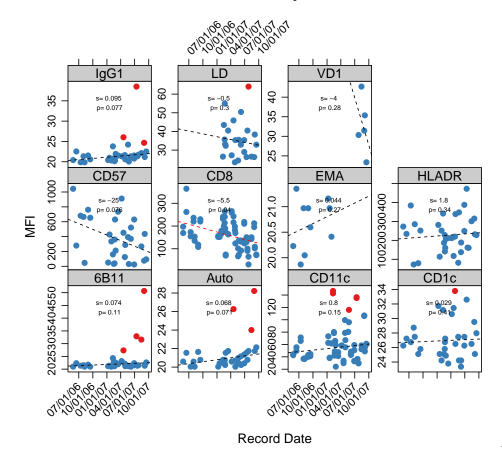


Record Date

Another three examples: QA check of Fluorescence stability overtime using t-distribution based outlier detection function.

```
> qaCheck(qaTask.list[["MFIOverTime"]]
                   ,rFunc=rlm
                   ,z.cutoff=3
                   )
```

#### Fluorescence stability over time

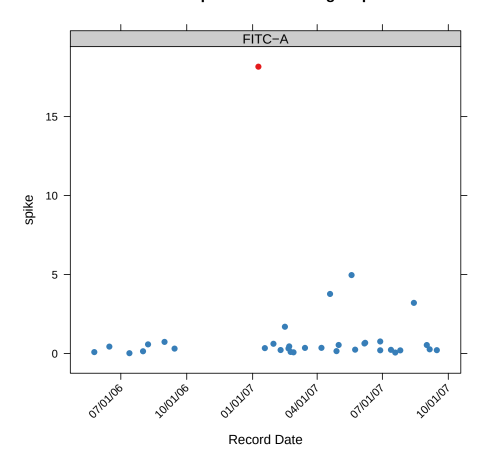


Note that the ro-

bust linear regression is applied in each group in order to capture the significant MFI change over time. The individual outliers within each group are detected based on the residue. All the lattice options (like scales here in this example) can be passed to control the appearance of trellis plot, see the documentation of lattice for more details.

+ )

## Measurement spikes/drifts during acquisition



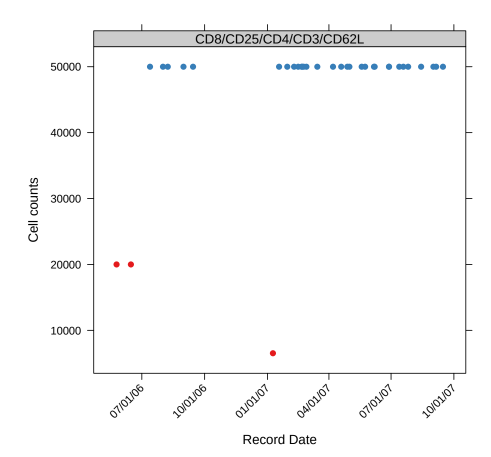
When monitoring the total number of events for each tube, a pre-determined events number can be provided as the threshold to the qaCheck method. tubesEvents pass to the argument of qaCheck could be a one-column data frame or a named list/vector. Threshold values are stored in the column or list/vector and conditioning values stored in rownames or names of the list/vector.

#### > tubesEvents

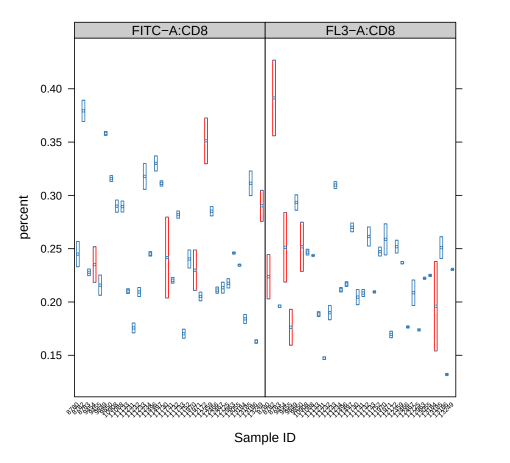
	events
CD8/CD25/CD4/CD3/CD62L	50000
CD1c/IgD/CD27/CD19/IgM	150000
HLADR/CD80/CD27/CD19/CD86	150000
CD57/CD56/CD8/CD3/CD14	20000
6B11/Va24/CD8/BLK/CD4	50000
CD8/CD69/CD4/CD3/HLADR	20000
IgG1/IgG1/IgG1/IgG1	20000
Auto/Auto/Auto/Auto	20000

```
CD11c/CD80/DUMP/HLADr/CD123 400000
CD11c/CD86/DUMP/HLADr/CD123 400000
LD/LD/LD/LD/LD
                              20000
VD1/VD2/GD/BLK/CD3
                             250000
> qaCheck(qaTask.list[["NumberOfEvents"]]
                  ,formula=count ~ RecdDt | Tube
                  ,outlierfunc=outlier.cutoff
                   ,1Bound=0.8*tubesEvents
[1] "done!"
> plot(qaTask.list[["NumberOfEvents"]]
                   ,subset=Tube=='CD8/CD25/CD4/CD3/CD62L'
                   ,xlab="Record Date"
                   ,ylab="Cell counts"
          )
```

#### **Number of Events Collected**



## **Consistency of redundant Staining Across sample aliquots**



# 6 Creating quality assessment report

Besides the interactive visualization provided by plot method,we also provide one routine to generate all plots in one report. This function reads the QA results calculated by qaCheck and the meta information of each QA task provided in spreadsheet qaCheckList and generate the summary tables and svg plots. Svg plots provide tooltips containing the detail information about each sample as and hyperlinks of densityplot for each individual FCS file.

plotAll is the argument to control the plotting of the individual scatter plot for each FCS file. If TRUE, scatter plots for all the FCS files are generated. If FALSE, only the FCS files marked as outliers will be plotted. When it is set to "none", no scatter plot will be generated, which is helpful to provide a quick preview of the html report. subset is the filter to only generate the report on a subset of flow data.

Note that if there is need to adjust the QA plot for each individual qaTask in the report, the arguments must be stored in qaTask before qaReport method is called. If there is no outlier is detected, the qaTask is not plotted by default. In order to change this setting, htmlReport can be set as TRUE to plot the qaTask regardlessly. This is particually useful for the task that tracks the longitudinal trend instead of the individual outliers. Besides the lattice arguments that can be configured by qpar (for the summary xyplot and bwplot) and scatterPar (for the individual FCS xyplot). xlog,ylog flag can be set for scatter plot to transform the flow data to log scale for the proper display of the stained channels.

```
> htmlReport(qaTask.list[["MFIOverTime"]])<-TRUE
> rFunc(qaTask.list[["MFIOverTime"]])<-rlm
> scatterPar(qaTask.list[["MFIOverTime"]])<-list(xlog=TRUE)
> scatterPar(qaTask.list[["BoundaryEvents"]])<-list(xlog=TRUE)
> scatterPar(qaTask.list[["RedundantStain"]])<-list(xlog=TRUE)
> qpar(qaTask.list[["RedundantStain"]])<-list(scales=list(x=list(relation="free")))</pre>
```

#### 7 Conclusion

Here is the example of the QA report.

By the formula-based qaCheck and plot methods, different QA tasks can be defined and performed in a generic way. And plot only reads the outlier detection results pre-calculated by qaCheck, which reduces the cost of interactive visualization.

Two kinds of lattice plots are currently supported:xyplot and bwplot(boxplot), depending on the plotType in qaTask object. When the output path is provided by dest, the svg plot is generated. In svg plot, each dot or box (or only the one marked as outliers) is annotated by the tooltip or hyperlink, which further points to the individual density plot of the gated population.