Automated Gating of Flow Cytometry Data using the Bioconductor openCyto Framework

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What's inside?

Flow cyometry is a method used to gain understanding of cell samples and populations by quantifying scattered and emitted fluorescent light. Signals are captured and analyzed through use of software programs. Flow cytometry analysis consists of gating, a method that dictates which cells will be further analyzed and which will not. Current methods for flow cytometry gating involve manually drawing gates. This process is both time consuming and costly, making automated gating procedures an appealing option. The <code>openCyto</code> package allows users to take manually gated data from flowJo, reproduce those gates in R, and eventually automate the gating process. The goal of this tutorial is to take the user through the process of automated gating analysis.

This tutorial will be useful to anyone who has done manual gating on a sample and wishes to automate the same procedure on additional samples in the future.

The example data used in this tutorial is from Colorado State University's Microbiology, Immunology, and Pathology Department. Alternatively, you can input your own data using the filetypes described in Chapter 3.

Getting Started

Here is an overview of the process to automate flow cytometry data using R's openCyto and what you will need to successfully automate your own flow cytometry analysis. The general steps to accomplish this are as follows:

- 1. Read in a manually gated flow Jo workspace in .wsp file format.
- 2. Parse raw FCS files from the read in workspace.
- 3. Visualize the manual gating template and resulting gates to verify gating scheme.
- 4. Create and read in a .csv gating template.
- 5. Automate gating.
- 6. Visualize automated gating template and gates to verify gating scheme.
- 7. Extract population statistics and relevant information.

This process is completed primarily with the openCyto package but calls upon other packages within the Bioconductor openCyto framework. Packages needed to complete this tutorial are listed at the end of this chapter. Descriptions of each function and R object used for this analysis are below.

2.1 Required Packages and Installation

2.1.1 Package descriptions

Below is a description of each package used in this analysis. Code to install and use these packages will follow. Package descriptions taken from Bioconductor and CRAN.

e	Package Name
This package is designed	openCyto
ce This package allows you to import basic flow Jo workspaces into BioConductor and replicate the gating for	flowWorkspace
Pro	flowCore
Methods and functiona	flowStats
Robu	flowClust
Fast aggregation of large data (e.g. 100GB in RAM), fast ordered joins, fast add/modify/delete	data.table

2.1.2 Installation

Install the following libraries into a new R script. As you will see below, this tutorial uses the development version of openCyto. It is important to use the development version of openCyto to remain up to date on

any changes made by the developers of openCyto. Use the following to ensure the correct packages are installed. Installation will only need to be done once.

To install

RStudio may also prompt you to download XQuartz and XCode based on your computer type, so it may be a good idea to go ahead and also download both.

2.1.3 Load packages

Although installation only needs to be done once, packages will need to be reloaded each time you open an R session. At the beginning of each session, run the following code.

To load

```
library(openCyto)
library(flowWorkspace)
library(data.table)
library(flowCore)
library(flowStats)
library(flowClust)
```

Working with your Manual Gating Scheme

The first step in this process is to bring a pre-existing flow of file into R in order to recreate the gating environment. The remainder of this chapter will detail the following:

- 1. Read in flowJo .wsp file
- 2. Parse FCS files

Workspace is open.

3. Visualize and verify manual gates

3.1 Read in flowJo file

Within flowJo, tranformation, compensation, and gating can be saved as either .xml or .wsp filetypes. This tutorial will only detail steps from a .wsp filetype saved from flowJo. Note that many other tutorials begin from a .xml filetype. Saving analysis within flowJo is detailed here. Your .wsp file will contain samples and groups to be added to the Workspace in R, all gates and analyses, and compensation matrices. Importantly, the .wsp will not save your FCS files. Rather, the path to your files will be saved and can be adjusted later within R.

Before you begin, be sure you have loaded the required packages outlined in the previous chapter.

Once all packages are loaded, save the .wsp file path as an R object called wsfile. Next, use openWorkspace() with your R object created in the prior step to open the .wsp file in R. Save this as an R object. Here, this was saved as ws and is of flowJoWorkspace class. Here is an example of saving and opening your .wsp filetype in R. Please ensure that ws is saved as a flowWorkspace object containing groups of samples before proceeding.

```
wsfile <- "./tutorial/group1_v_group2.wsp"

ws <- openWorkspace(wsfile)

print(ws)

## FlowJo Workspace Version 20.0

## File location: ./tutorial

## File name: group1_v_group2.wsp</pre>
```

```
##
## Groups in Workspace
## Name Num.Samples
## 1 All Samples 10
## 2 Samples 10
```

3.2 Parse FCS files

The next step is to read in raw FCS files. FCS files contain data from the cytometer. Standards for FCS files are listed here.

Raw FCS files are read using the parseWorkspace function. This function will read the FCS files and transform, compensate, and gate according to parameters defined from the .wsp flowJo workspace, which is now saved as an R object of class flowWorkspace. The parseWorkspace call requires the object that results from running openWorkspace. Here, we named this object ws. The function parseWorkspace() also requires the name of the samples to read in. To list sample names, use the getSampleGroups() function on your flowWorkspace class object. Other options may be customized based on particular needs. A new R object named gating_set is then created and will be a GatingSet object. The isNcdf = TRUE call saves this output to disk rather that into memory because the files are large. Here is an example of parsing FCS files. As this function runs, you will see several messages appear as the FCS files are loaded and the manual gating scheme is replicated. After this, attributes() is used to examine the data.

```
gating_set <- parseWorkspace(ws, name = "Samples", path = "./tutorial/group1_v_group2", isNcdf = TRUE,
## windows version of flowJo workspace recognized.
## version X
attributes(gating_set)</pre>
```

3.3 Visualize and Verify

It is helpful to now visualize both the gating template and gates on a subset of the data in order to verify the gating scheme. This will ensure consistency between the flowJo workspace and the manual gates recreated in R. First, save a subset of the gating_set as follows. The following saves the first FCS file of gating_set as gh. Since each FCS file corresponds to an individual experiment, this saves the first experiment of the group.

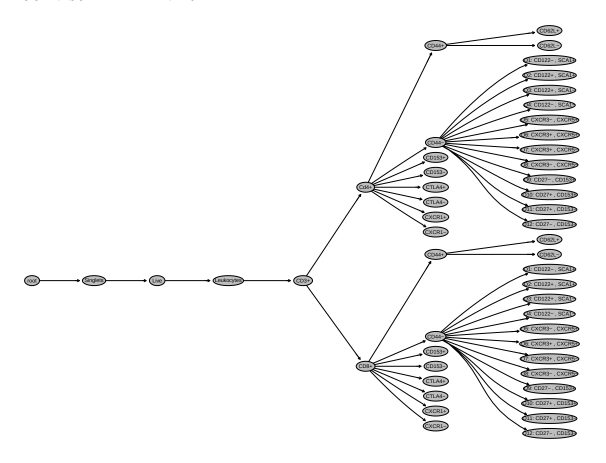
```
gh <- gating_set[[1]]
print(gh)

## Sample: X_group1_1
## GatingHierarchy with 51 gates</pre>
```

3.3.1 plot()

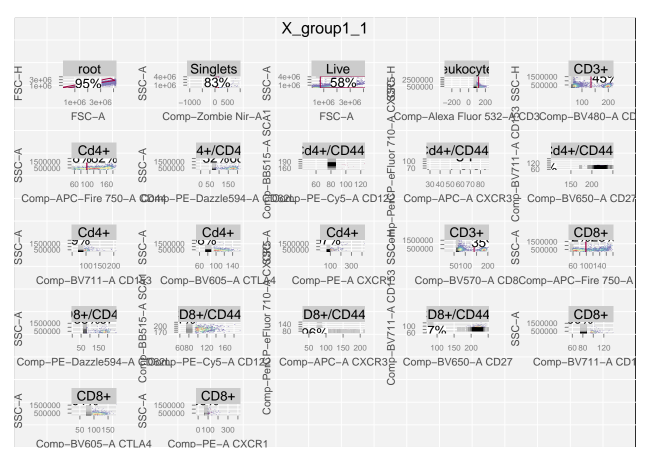
The plot() function will visualize the current gating hierarchy when applied to an object of class GatingHierarchy. This can be done for the entire gating hierarchy or a specific population as seen below.

```
plot(gh)
```



3.3.2 plotGate()

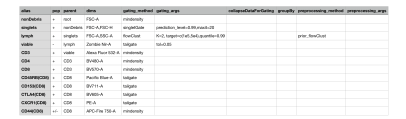
The plotGate() function will gate the designated subset of your data according to parameters replicated from flowJo. This must also be called on an object of class GatingHierarchy.



^{**}Note the use of flowWorkspace.par.set() here. Chapter 5 of this tutorial will discuss customizations such as this one.

Create .csv

The creation of a .csv gating template is arguably the most important step to automating flow cytometry analysis. The .csv template that you create will tell <code>openCyto</code> how to gate your data. Included with this tutorial is a partial .csv template that can be used to gate the sample data. Generally, .csv templates will look like this:



4.1 .csv Gating Template Structure

In the gating template, each row corresponds to a single cell population and the method used to gate that population. When read into R, the .csv will direct gating based on parameters listed in each row and column. The .csv must contain 10 predefined columns as seen here:



4.1.1 alias

The first column must be titled alias. This is where you will put your cell population names. Remember, each row corresponds to a single cell population. Population names in the alias column must be unique.

alias
nonDebris
singlets
lymph
viable
CD3
CD4
CD8
CD45RB(CD8)
CD153(CD8)
CTLA4(CD8)
CXCR1(CD8)
CD44(CD8)

4.1.2 pop

The second column must be titled pop. This column will contain a + or - to designate which subset or quadrant will be gated. A + will gate the positive subset while a - will gate the negative. This column can only contain strings of + and -, so do not use any characters as separators for quadrant gates.

	_
pop	
+	
+	
+	
-	
+	
+	
+	
+	
+	
+	
+	

4.1.3 parent

The third column must be titled parent. This column refers to the parent cell population, or where the current cell population originates from. Similar to the alias column, parent names must be unique. This column cannot contain any commas, otherwise openCyto will assume the population has multiple parents and you will get an error message.

parent
root
nonDebris
singlets
lymph
viable
CD3
CD3
CD8

4.1.4 Remaining template columns

preprocessing_args- arguments for preprocessing function

dims- channel or marker names for gating gating_method- gating function (supported options listed above) quadrantGate ${\bf range Gate}$ quantileGate mindensity tailgate cytokine ${\rm flowClust}$ boundary singletGate transitionalplolyfunctionalityGate flowDensity gating_args- arguments to be passed to gating function collapseDataforGating- data is collapsed and replicated across all samples groupBy- used to group samples into unique combinations preprocessing_method- preprocessing function

4.2 Creating the Template

The gating template can be created manually or assisted by the use of the templateGen() function. The function templateGen() will input the alias, pop, parent, and dims columns and the rest must be completed manually. To use templateGen(), you must input a GatingHierarchy object. In this example, that is gh, the subset created from gating_set.

```
gt <- templateGen(gh)
head(gt)</pre>
```

```
##
           alias
                         pop
                                                             parent
## 1
       Singlets
                   Singlets
                                                               root
## 2
           Live
                       Live
                                                          /Singlets
## 3 Leukocytes Leukocytes
                                                    /Singlets/Live
           CD3+
                        CD3+
                                        /Singlets/Live/Leukocytes
## 4
## 5
            CD8+
                        CD8+
                                   /Singlets/Live/Leukocytes/CD3+
## 6
         CXCR1-
                     CXCR1- /Singlets/Live/Leukocytes/CD3+/CD8+
##
                                dims gating method gating args
## 1
                        FSC-A, FSC-H
                                               <NA>
                                                            <NA>
## 2
                 Comp-Zombie Nir-A
                                               <NA>
                                                            <NA>
## 3
                        FSC-A, SSC-A
                                               <NA>
                                                            <NA>
## 4 Comp-Alexa Fluor 532-A,SSC-H
                                               <NA>
                                                            <NA>
                Comp-BV570-A, SSC-H
## 5
                                               <NA>
                                                            <NA>
##
  6
                          Comp-PE-A
                                               <NA>
                                                            <NA>
##
     collapseDataForGating groupBy preprocessing_method preprocessing_args
## 1
                                <NA>
                        <NA>
                                                        <NA>
## 2
                        <NA>
                                 <NA>
                                                        <NA>
                                                                             <NA>
## 3
                        <NA>
                                <NA>
                                                        <NA>
                                                                             <NA>
## 4
                        <NA>
                                <NA>
                                                        <NA>
                                                                             <NA>
## 5
                                <NA>
                                                                             <NA>
                        <NA>
                                                        <NA>
## 6
                        <NA>
                                 <NA>
                                                        <NA>
                                                                             <NA>
```

The auto-filled template will generate within the R Console and can then be saved locally with the following code. You will see that all columns besides the first four will contain NA values. These are the values that must be manually input to complete the .csv.

```
write.csv(gt, "gt.csv")
```

If you choose to create the gating template manually, the same conventions must be followed. Start with a blank spreadsheet. Next, fill in the 10 required column names. From there, use the manual gating hierarchy to fill in each cell population alias. Fill in the remainder accordingly.

There will likely be troubleshooting involved in this process. This is a great place to start if you're seeking more information on the gating template. The openCyto GitHub page is also very responsive to issues posted.

Automate Gating

5.1 Load .csv into R

As noted in the previous chapter, there is a sample gating template titled *partial.csv* with the sample data. This may serve as a guide to creating your own. When the .csv gating template is complete, it is then read into R and saved as **gt**, a gating Template object.

```
gt <- gatingTemplate("./tutorial/partial_gt.csv")</pre>
```

The flow cytometry equipment at CSU will compensate and transform the data automatically. Other tutorials may highlight the steps to compensate and transform data, but these are not relevant to CSU at this moment. In the event that equipment changes, it may be necessary to complete compensation and transformation steps to prepare data. More on the current equipment used as CSU here.

5.2 Read in raw FCS files

Now that the GatingTemplate object has been loaded into R, you will need to load in raw FCS files to perform the automated gating on. For gating, these files must be in a GatingSet object type, which requires the following steps. When the path is saved using list.files, a character matrix of file names will be saved. Next, read.ncdfFlowSet will save FCS files as a ncdfFlowSet object. The GatingSet function will then save the FCS files as a GatingSet object. In this form, the FCS files can be input and gated.

```
fcs_files <- list.files(path = "./tutorial/group1_v_group2", full.names = TRUE)
ncfs <- read.ncdfFlowSet(files = fcs_files)
gs_auto <- GatingSet(ncfs)</pre>
```

5.3 Apply Gating

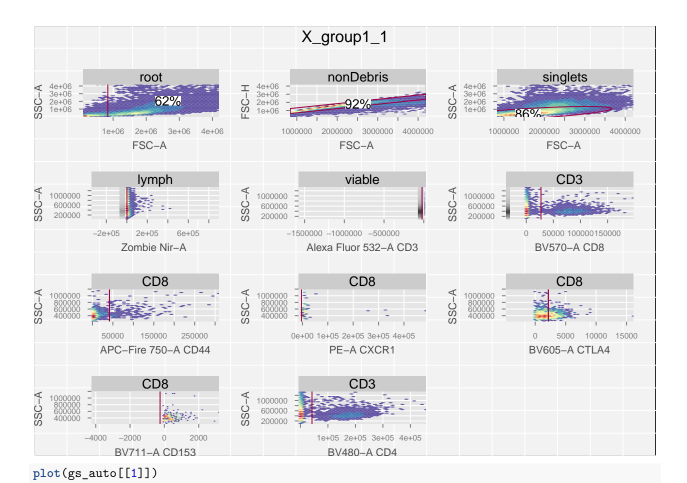
At this point, you now have GatingTemplate and GatingSet object to be used for gating. Apply your GatingTemplate object to the GatingSet object, where x = GatingTemplate object and y = GatingSet object.

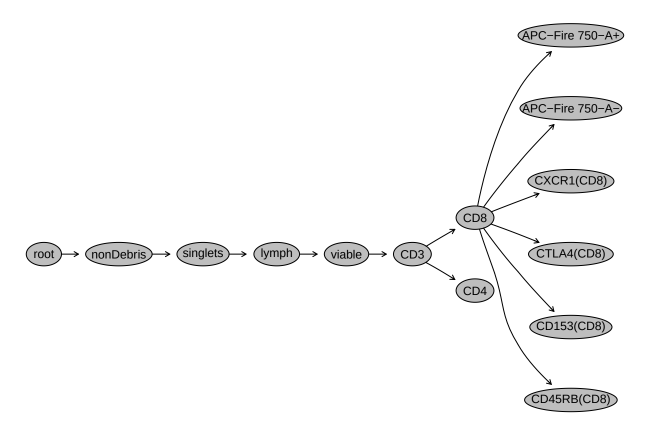
```
gating(x = gt, y = gs_auto)
```

5.4 Plot Automated Gating

Just as before, plot both the gating hierarchy and the automated gates. You may notice extra nodes have been added to the hierarchy. Chapter 5 will highlight additional cusomization to remove unwanted nodes and improve upon visualization.

plotGate(gs_auto[[1]])





5.5 Population Statistics

Both counts and frequencies can be generated for analysis. This can be generate based on the analysis completed in R, or pulled directly from flowJo. To pull from flowJo, simply at flowJo=TRUE to either code chunk below.

Counts

```
head(getPopStats(gs_auto,statistic="count"))
```

```
##
                              Parent Count ParentCount
           name Population
## 1: X_group1_1 nonDebris
                                root 87715
                                                 142158
## 2: X_group1_1
                 singlets nonDebris 80775
                                                  87715
## 3: X_group1_1
                                                  80775
                     lymph singlets 69710
## 4: X_group1_1
                               lymph 40982
                                                  69710
                    viable
## 5: X_group1_1
                        CD3
                               viable 40937
                                                  40982
## 6: X_group1_1
                        CD8
                                  CD3 1888
                                                  40937
```

Frequencies

head(getPopStats(gs_auto,statistic="freq"))

##		name	Population	Parent	Count	${\tt ParentCount}$
##	1:	X_group1_1	nonDebris	root	87715	142158
##	2:	X_group1_1	singlets	${\tt nonDebris}$	80775	87715
##	3:	<pre>X_group1_1</pre>	lymph	singlets	69710	80775
##	4:	X_group1_1	viable	lymph	40982	69710
##	5:	X_group1_1	CD3	viable	40937	40982
##	6:	X group1 1	CD8	CD3	1888	40937

Customization

It is possible that additional customization may be necessary when working with the openCyto framework. Below are three common customizations that will be outlined in this chapter.

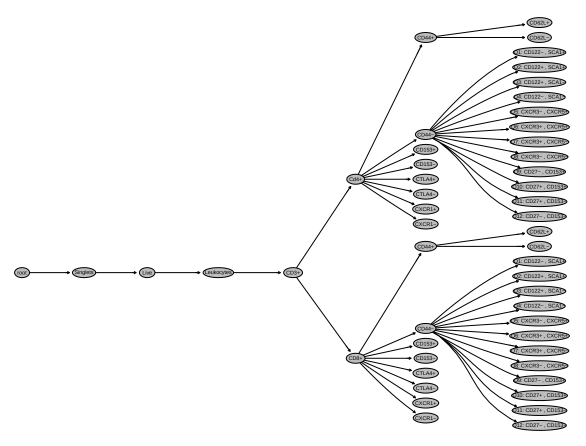
- 1. Hiding unwanted nodes
- 2. Renaming nodes
- 3. Adjusting plots

6.1 Hiding unwanted nodes

When automating analysis, there may be nodes that were not predefined in the .csv gating template or nodes that may not be of interest in your particular analysis. Plotting the gating hierarchy using the plot() function will display this and then nodes can be hidden based on need with the following code. Below is an example of a "full" gating hierarchy and then the same hierarchy with the CD3+ node removed.

Full Hierarchy

plot(gh)



CD3+ Removed Hierarchy

To remove nodes, first save the unwanted nodes as an R object named **nodesToHide**. Next, use the code following the lapply() function, only replacing **gs** with your GatingSet object name.

```
nodesToHide <- "CD3+"
lapply(nodesToHide, function(thisNode)setNode(gh, thisNode, FALSE))
## [[1]]
## NULL</pre>
```

6.2 Renaming nodes

Rename nodes based on your preferences with the following code. Within the setNode function, the first input is the current cell population name and the second is the desired change.

```
setNode("Live", "Viable")
plot(gh)
```

6.3 Adjusting plots

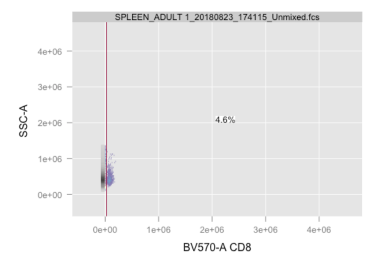
6.3.1 Adjust plot axes

As seen in chapter 2, it may be necessary to adjust the plot axes in order to best view the gates. This is done using the code below. Setting xlim and ylim to "data" adjusts plot based on the actual data range,

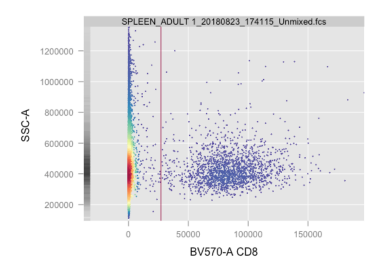
rather than instrument specifications. Custom ranges can also be input numerically.

Here is a comparison of xlim and ylim set as "instrument" and then "data".

Instrument



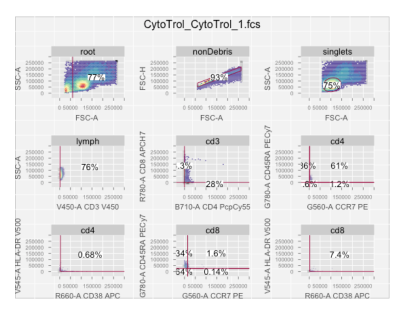
Data



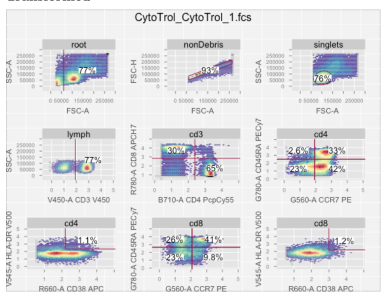
6.3.2 Transform data for better visualization

Although data will not be altered in any way, transformation may allow for better visualization. The most common form of transformation for flow cytometry analysis is biexponential. Below is a comparison of gates without transformation and gates that have been transformed.

Without Transformation



Transformed



Appendix

7.1 Function and R Object Definitions

Function /Object Name	Definition
Function/Object Name	
wsfile	flowJo .wsp file location
openWorkspace()	function used to read in **wsfile**
ws	read in data from flowJo
parseWorkspace()	function to extract FCS files from **ws**
gating_set	parsed FCS files to be gated
clone()	function used to create a clone of **gating_set**
gh	subset of gating_set
gt	.csv gating template
templateGen()	function used to generate a .csv template from existing manual gates
gatingTemplate()	function used to read in .csv template
gating()	function used to apply gates to a gating set
plot()	function to visualize gating tree
plotGate()	function to visualize gates