flowWorkspace: A Package for Importing flowJo Workspaces into R

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1 Purpose

The purpose of this package is to provide functionality to import relatively simple flowJo workspaces into R. By this we mean, accessing the samples, groups, transformations, compensation matrices, gates, and population statistics in the flowJo workspace, and replicating these using (primarily) flowCore functionality.

2 Why Another flowJo Workspace Import Package?

There was a need to import flow Jo workspaces into R for comparative gating. The flow Flow Jo package did not meet our needs. Many groups have legacy data with associated flow Jo XML workspace files in version 2.0 format that they would like to access using Bio Conductor's tools. Hopefully this package will fill that need.

3 Support

This package supports importing of **Version 2.0 XML workspaces only**. We cannot import **.jo** files directly. You will have to save them in XML workspace format, and ensure that that format is *workspace version 2.0*. The package has been tested and works with files generated using flow Jo version 9.1 on Mac OS X. XML generated by older versions of *flowJo* on windows should work as well. We do not yet support *flowJo*'s **Chimera** XML schema, though that support will be provided in the future.

The package supports import of only a subset of the features present in a flow Jo workspace. The package allows importing of sample and group names, gating hierarchy, compensation matrices, data transformation functions, a subset of gates, and population counts.

BooleanGates are now supported by flowWorkspace.

4 Data Structures

The following section walks through opening and importing a flow Jo workspace.

4.1 Loading the library

Simply call:

```
> library(flowWorkspace)
```

The library depends on numerous other packages, including graph, XML, Rgraphviz, flowCore, flowViz, RBGL.

4.2 Opening a Workspace

We represent flowJo workspaces using flowJoWorkspace objects. We only need to know the path to, and filename of the flowJo workspace.

```
> d<-system.file("extdata",package="flowWorkspaceData");
> wsfile<-list.files(d,pattern="A2004Analysis.xml",full=T)

In order to open this workspace we call:
> ws<-openWorkspace(wsfile)
> ws

FlowJo Workspace Version 2.0
File location: /home/biocbuild/bbs-3.1-bioc/R/library/flowWorkspaceData/extdata
File name: A2004Analysis.xml
Workspace is open.

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

We see that this a version 2.0 workspace file. It's location and filename are printed. Additionally, you are notified that the workspace file is open. This refers to the fact that the XML document is internally represented using 'C' data structures from the XML package. After importing the file, the workspace must be explicitly closed using closeWorkspace() in order to free up that memory.

4.3 Parsing the Workspace

With the workspace file open, we have not yet imported the XML document. The next step parses the XML workspace and creates R data structures to represent some of the information therein. Specifically, by calling parse-Workspace() the user will be presented with a list of groups in the workspace file and need to choose one group to import. Why only one? Because of the way flow Jo handles data transformation and compensation. Each group of samples is associated with a compensation matrix and specific data transformation. These are applied to all samples in the group. When a particular group of samples is imported, the package generates a GatingHierarchy for each sample, describing the set of gates applied to the data (note: polygons, rectangles, quadrants, and ovals and boolean gates are supported). The set of GatingHierarchies for the group of samples is stored in a GatingSet object. Calling parseWorkspace() is quite verbose, informing the user as each gate is created. The parsing can also be done non-interactively by specifying which group to import directly in the function call (either an index or a group name). An additional optional argument execute=T/F specifies whether you want to load, compensate, transform the data and compute statistics immediately after parsing the XML tree.

- > G<-parseWorkspace(ws,name=1,path=ws@path,isNcdf=FALSE,cleanup=FALSE,keep.indices=TR
- > #Lots of output here suppressed for the vignette.

When is Ncdf flag is set TRUE, the data is stored in ncdf format on disk.

> G

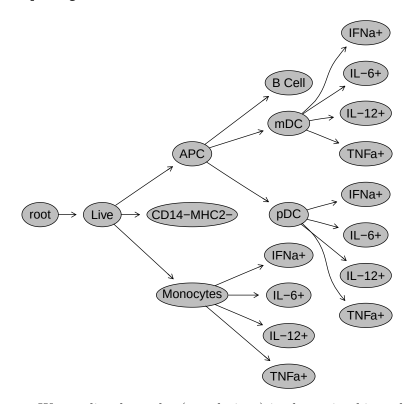
A GatingSet with 2 samples

We have generated a GatingSet with 2 samples, each of which has 19 associated gates. Subsets of gating hierarchies can be accessed using the standard R subset syntax.

At this point we have parsed the workspace file and generate the gating hierarchy associated with each sample imported from the file. The data have been loaded, compensated, and transformed in the workspace, and gating has been executed. The resulting *GatingSet* contains a replicated analysis of the original flowJo workspace.

We can plot the gating hierarchy for a given sample:

```
> gh<-G[[1]]
> plot(gh)
```



We can list the nodes (populations) in the gating hierarchy:

- > nodelist<-getNodes(gh, path = 1)</pre>
- > nodelist

[1]	"root"	"Live"	"APC"	"B Cell"	"mDC"
[6]	"IFNa+"	"IL-6+"	"IL-12+"	"TNFa+"	"pDC"
[11]	"IFNa+"	"IL-6+"	"IL-12+"	"TNFa+"	"CD14-MHC2-"
[16]	"Monocytes"	"IFNa+"	"IL-6+"	"IL-12+"	"TNFa+"

Note that the number preceding the period in the node names is just an identifier to help uniquely label populations in the gating hierarchy. It does

not represent any information about population statistics. When the node name itself is already unique, such as "Live", "B Cell", the prefix is ommitted by default. We can force the prefix to be added to each node by setting prefix to "all".

> getNodes(gh, prefix="all", path = 1)

```
"Live"
                                  "APC"
                                                              "mDC"
 [1] "root"
                                                "B Cell"
 [6] "IFNa+"
                   "IL-6+"
                                                "TNFa+"
                                                              "pDC"
                                  "IL-12+"
[11] "IFNa+"
                   "IL-6+"
                                  "IL-12+"
                                                "TNFa+"
                                                              "CD14-MHC2-"
                                  "IL-6+"
                                                              "TNFa+"
[16] "Monocytes"
                   "IFNa+"
                                                "IL-12+"
```

Alternatively, the the full path of the node can be displayed by setting path to "full".

> getNodes(gh, path = "full")

```
[1] "root"
                                                         "/Live/APC"
                               "/Live"
 [4] "/Live/APC/B Cell"
                               "/Live/APC/mDC"
                                                         "/Live/APC/mDC/IFNa+"
 [7] "/Live/APC/mDC/IL-6+"
                               "/Live/APC/mDC/IL-12+"
                                                         "/Live/APC/mDC/TNFa+"
[10] "/Live/APC/pDC"
                               "/Live/APC/pDC/IFNa+"
                                                         "/Live/APC/pDC/IL-6+"
[13] "/Live/APC/pDC/IL-12+"
                               "/Live/APC/pDC/TNFa+"
                                                         "/Live/CD14-MHC2-"
                               "/Live/Monocytes/IFNa+"
[16] "/Live/Monocytes"
                                                         "/Live/Monocytes/IL-6+"
[19] "/Live/Monocytes/IL-12+" "/Live/Monocytes/TNFa+"
```

We can get a specific gate definition:

```
> node<-nodelist[3]
> g<-getGate(gh,node)
> g
```

Polygonal gate 'APC' with 14 vertices in dimensions <PerCP-CY5-5-A> and <PE-CY7-A>

We can get the population proportion (relative to its parent) for a single population:

```
> getProp(gh, node)
```

[1] 0.08402716

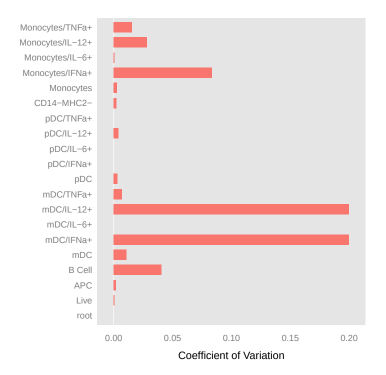
Or we can retrieve the population statistics for all populations in the sample:

> getPopStats(gh)

	<pre>flowCore.freq</pre>	${\tt flowJo.freq}$	<pre>flowJo.count</pre>	${\tt flowCore.count}$	node
1:	1.000000000	1.000000000	61832	61832	root
2:	0.800297581	0.801235606	49542	49484	Live
3:	0.084027160	0.083585645	4141	4158	APC
4:	0.592352092	0.548418256	2271	2463	B Cell
5:	0.123376623	0.121226757	502	513	mDC
6:	0.005847953	0.003984064	2	3	mDC/IFNa+
7:	0.042884990	0.043824701	22	22	mDC/IL-6+
8:	0.005847953	0.003984064	2	3	mDC/IL-12+
9:	0.140350877	0.141434263	71	72	mDC/TNFa+
10:	0.107984608	0.107703453	446	449	pDC
11:	0.002227171	0.002242152	1	1	pDC/IFNa+
12:	0.000000000	0.000000000	0	0	pDC/IL-6+
13:	0.561247216	0.560538117	250	252	pDC/IL-12+
14:	0.000000000	0.000000000	0	0	pDC/TNFa+
15:	0.544074852	0.540854225	26795	26923	CD14-MHC2-
16:	0.058928138	0.059161923	2931	2916	Monocytes
17:	0.003772291	0.004435346	13	11	Monocytes/IFNa+
18:	0.237654321	0.236779256	694	693	Monocytes/IL-6+
19:	0.047325103	0.049812351	146	138	Monocytes/IL-12+
20:	0.250685871	0.257250085	754	731	Monocytes/TNFa+

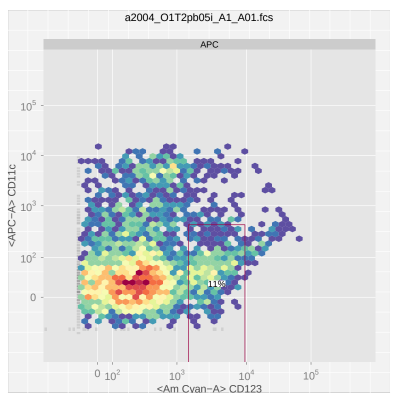
We can plot the coefficients of variation between the counts derived using flowJo and flowCore for each population:

> print(plotPopCV(gh))



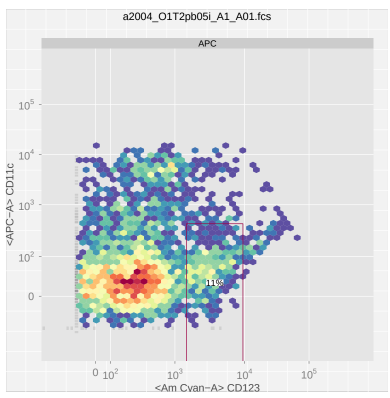
We can plot individual gates: note the scale of the transformed axes. Second argument can be either a numeric index

- > plotGate(gh, 10)
- or unique node name,
- > plotGate(gh, "pDC")
- or a full/parital gating path,
- > plotGate(gh, "APC/pDC")



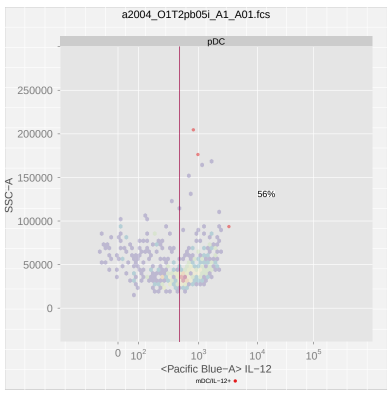
hexbin can be enabled to speed up plotting.

> plotGate(gh, "pDC", xbin =32)



 ${\tt overlay}$ can be used to plot another cell population on top of the existing gates specified by y.

> plotGate(gh,y = "pDC/IL-12+", xbin =32, overlay = "mDC/IL-12+")



overlay can be either a numeric or character scalar indicating the index or gating path of a gate/population within the GatingHierarchy or a logical vector that indicates the cell event indices representing a sub-cell population.

If we have metadata associated with the experiment, it can be attached to the GatingSet.

```
> d<-data.frame(sample=factor(c("sample 1", "sample 2")),treatment=factor(c("sample",
> pd<-pData(G)
> pd<-cbind(pd,d)
> pData(G)<-pd
> pData(G);
name sample treatment
```

sample

control

We can retrieve the subset of data associated with a node:

a2004_01T2pb05i_A1_A01.fcs a2004_01T2pb05i_A1_A01.fcs sample 1

a2004_01T2pb05i_A2_A02.fcs a2004_01T2pb05i_A2_A02.fcs sample 2

> getData(gh,node)

flowFrame object 'a2004_01T2pb05i_A1_A01.fcs'
with 4158 cells and 23 observables:

name	desc	range	minRange	maxRange
\$P1 FSC-A	<na></na>	262254.000	-111.00000	262143.000
\$P2 FSC-H	<na></na>	262143.000	0.00000	262143.000
\$P3 FSC-W	<na></na>	262143.000	0.00000	262143.000
\$P4 SSC-A	<na></na>	262254.000	-111.00000	262143.000
\$P5 SSC-H	<na></na>	262143.000	0.00000	262143.000
\$P6 SSC-W	<na></na>	262143.000	0.00000	262143.000
\$P7 <am cyan-a=""></am>	CD123	3661.959	435.34379	4097.303
\$P8 Am Cyan-H	CD123	3641.837	455.00000	4096.837
<pre>\$P9 <pacific blue-a=""></pacific></pre>	IL-12	3927.974	169.60860	4097.582
\$P10 Pacific Blue-H	IL-12	3641.837	455.00000	4096.837
\$P11 <apc-a></apc-a>	CD11c	4405.818	-308.01302	4097.805
\$P12 APC-H	CD11c	3641.837	455.00000	4096.837
\$P13 <apc-cy7-a></apc-cy7-a>	IL-6	3714.446	382.93207	4097.378
\$P14 APC-CY7-H	IL-6	3641.837	455.00000	4096.837
\$P15 <alexa 700-a=""></alexa>	TNFa	3712.753	384.62271	4097.376
\$P16 Alexa 700-H	TNFa	3641.837	455.00000	4096.837
\$P17 <fitc-a></fitc-a>	IFNa	4180.519	-82.81306	4097.706
\$P18 FITC-H	IFNa	3641.837	455.00000	4096.837
<pre>\$P19 <percp-cy5-5-a></percp-cy5-5-a></pre>	MHCII	4942.398	-844.59317	4097.805
\$P20 PerCP-CY5-5-H	MHCII	3641.837	455.00000	4096.837
\$P21 <pe-cy7-a></pe-cy7-a>	CD14	4942.398	-844.59317	4097.805
\$P22 PE-CY7-H	CD14	3641.837	455.00000	4096.837
\$P23 Time	<na></na>	9918.400	89.00000	10007.400
323 keywords are store				

When applied to the GatingSet, a flowSet or ncdfFlowSet is returned.

> getData(G, node);

A flowSet with 2 experiments.

An object of class 'AnnotatedDataFrame'

 $\verb"rowNames: a2004_O1T2pb05i_A1_A01.fcs a2004_O1T2pb05i_A2_A02.fcs"$

varLabels: name sample treatment
varMetadata: labelDescription

column names:

FSC-A FSC-H FSC-W SSC-A SSC-H SSC-W <Am Cyan-A> Am Cyan-H <Pacific Blue-A> Pacific

Or we can retrieve the indices specifying if an event is included inside or outside a gate using:

> table(getIndices(gh,node))

FALSE TRUE 57674 4158

The indices returned are relative to the parent population (member of parent AND member of current gate), so they reflect the true hierarchical gating structure.

If we wish to do compensation or transformation manually, we can retrieve all the compensation matrices from the workspace:

> C<-getCompensationMatrices(gh);</pre>

> C

Compensation object 'defaultCompensation':

	Am Cyan-A	Pacific Blue-A	APC-A	APC-CY7-A	Alexa 700-A
Am Cyan-A	1.00000	0.04800	0.000000	0.0000	0.00000
Pacific Blue-A	0.38600	1.00000	0.000529	0.0000	0.00000
APC-A	0.00642	0.00235	1.000000	0.0611	0.19800
APC-CY7-A	0.03270	0.02460	0.084000	1.0000	0.02870
Alexa 700-A	0.07030	0.05800	0.016200	0.3990	1.00000
FITC-A	0.74500	0.02090	0.001870	0.0000	0.00000
PerCP-CY5-5-A	0.00368	0.00178	0.015300	0.0269	0.07690
PE-CY7-A	0.01330	0.00948	0.000951	0.1380	0.00182
	FITC-A	PerCP-CY5-5-A F	PE-CY7-A		
Am Cyan-A	0.028500	0.00104	0.00000		
Pacific Blue-A	0.000546	0.00000	0.00000		
APC-A	-0.000611	0.00776	0.00076		
APC-CY7-A	0.002690	0.00304	0.01010		
Alexa 700-A	0.001530	0.10800	0.00679		
FITC-A	1.000000	0.04180	0.00281		
PerCP-CY5-5-A	0.000000	1.00000	0.07030		
PE-CY7-A	0.002340	0.03360	1.00000		

Or we can retrieve transformations:

> names(T)

> T<-getTransformations(gh)

```
" <Alexa 700-H>"
 [1] "A2004-A2005_06i <Alexa 700-A>"
 [3] "A2004-A2005_06i <Am Cyan-A>"
                                         " <Am Cyan-H>"
 [5] "A2004-A2005_06i <APC-A>"
                                         "A2004-A2005_06i <APC-CY7-A>"
 [7] " <APC-CY7-H>"
                                         " <APC-H>"
 [9] "A2004-A2005_06i <FITC-A>"
                                         " <FITC-H>"
[11] "A2004-A2005_06i <Pacific Blue-A>" " <Pacific Blue-H>"
                                    " <PE-CY7-H>"
[13] "A2004-A2005_06i <PE-CY7-A>"
[15] "A2004-A2005_06i <PerCP-CY5-5-A>" " <PerCP-CY5-5-H>"
> T[[1]]
function (x, deriv = 0)
    deriv <- as.integer(deriv)</pre>
    if (deriv < 0 || deriv > 3)
        stop("'deriv' must be between 0 and 3")
    if (deriv > 0) {
        z0 <- double(z$n)</pre>
        z[c("y", "b", "c")] \leftarrow switch(deriv, list(y = z$b, b = 2 *
            z$c, c = 3 * z$d), list(y = 2 * z$c, b = 6 * z$d,
            c = z0), list(y = 6 * z$d, b = z0, c = z0))
        z[["d"]] <- z0
    }
    res <- stats:::.splinefun(x, z)
    if (deriv > 0 \&\& z\$method == 2 \&\& any(ind <- x <= z\$x[1L]))
        res[ind] <- ifelse(deriv == 1, z$y[1L], 0)
    res
}
<environment: 0x86b98f0>
attr(,"type")
[1] "caltbl"
```

getTransformations returns a list of functions to be applied to different dimensions of the data. Above, the transformation is applied to this sample, the appropriate dimension is transformed using a channel–specific function from the list.

The list of samples in a workspace can be accessed by:

```
> getSamples(ws);
```

And the groups can be accessed by:

> getSampleGroups(ws)

```
groupName groupID sampleID
1 All Samples 0 1
2 All Samples 0 2
```

The compID column tells you which compensation matrix to apply to a group of files, and similarly, based on the name of the compensation matrix, which transformations to apply.

4.4 add/remove gates

> data(GvHD)

> getNodes(gh1)

[1] "root"

GatingSet is equivalent to the workFlow in flowCore package ,which provides methods to build a gating tree from raw FCS files and add or remove flowCore gates(or populations) to or from it. Firstly,we start from a flowSet that contains three ungated flow samples:

```
> #select raw flow data
> fs<-GvHD[1:3]
Then we can transform the raw data:
> tf <- transformList(colnames(fs[[1]])[3:6], asinh, transformationId="asinh")
> fs_trans<-transform(fs,tf)
and create a gatingset from the transformed flowSet:
> gs <- GatingSet(fs_trans)
> gs
A GatingSet with 3 samples
> gh1<-gs[[1]]</pre>
```

It now only contains the root node. We can add one rectangleGate:

Note that the gate is added to root node by default if parent is not spec-

ified. Then we add a quadGate to the new population generated by the rectangeGate which is named after filterId of the gate because the name is not specified when add method is called.

[5] "/rectangle/CD15 FITC+CD45 PE-" "/rectangle/CD15 FITC-CD45 PE-"

[3] "/rectangle/CD15 FITC-CD45 PE+" "/rectangle/CD15 FITC+CD45 PE+"

Here quadGate produces four population nodes/populations whose names are named after dimensions of gate if not specified.

Boolean Gate can also be defined and added to GatingSet:

```
> bg<-booleanFilter(`CD15 FITC-CD45 PE+|CD15 FITC+CD45 PE-`)
> bg
```

booleanFilter filter 'CD15 FITC-CD45 PE+|CD15 FITC+CD45 PE-' evaluating the expressio CD15 FITC-CD45 PE+|CD15 FITC+CD45 PE-

```
> nodeID2<-add(gs,bg,parent="rectangle")
> nodeID2
```

[1] 7

> getNodes(gh1)

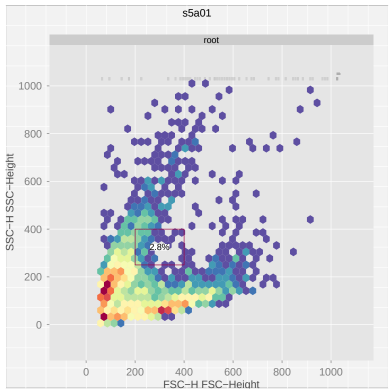
- [1] "root"
- [2] "/rectangle"
- [3] "/rectangle/CD15 FITC-CD45 PE+"
- [4] "/rectangle/CD15 FITC+CD45 PE+"
- [5] "/rectangle/CD15 FITC+CD45 PE-"
- [6] "/rectangle/CD15 FITC-CD45 PE-"
- [7] "/rectangle/CD15 FITC-CD45 PE+|CD15 FITC+CD45 PE-"

Now the gating tree is finished but the actual gating is done by **recompute** method:

> recompute(gs)

After gating is finished, gating results can be visualized by plotGate method:

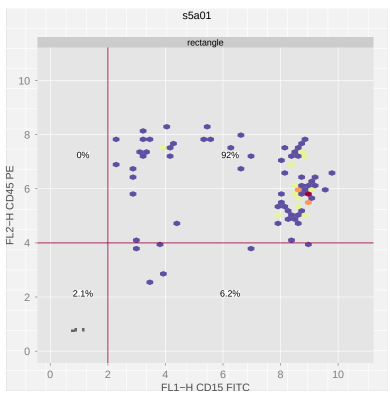
> plotGate(gh1, "rectangle") #plot one Gate



The second ar-

gument of plotGate is used to specify node index/name. Multiple gates can be plotted on the same pannel:

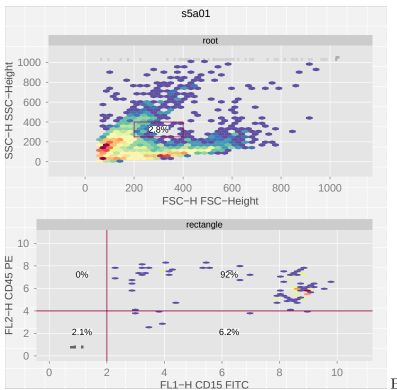
> plotGate(gh1,nodeIDs)



We may also want

to plot all the gates without specifying the gate index:

> plotGate(gh1)

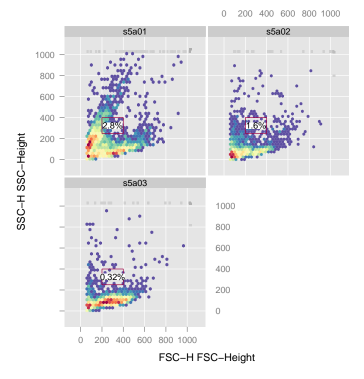


FL1-H CD15 FITC Boolean gate is skipped by default and can be enabled by:

> plotGate(gh1,bool=TRUE)

Note that smoothing may be applied automatically if there are not enough events after gating Sometime it is more useful to compare gates across samples using lattice plot by applying plotGate to a GatingSet:

> plotGate(gs,getNodes(gs)[nodeID])



The gating hierarchy is plotted by:

> plot(gh1,bool=TRUE)

If we want to remove one node, simply:

- > Rm('rectangle', gs)
 > getNodes(gh1)
- [1] "root"

As we see,removing one node causes all its descendants to be removed as well.

4.5 archive and clone

Oftentime, we need to save a GatingSet including the gated flow data, gates and populations to disk and reload it later on. It can be achieved by:

save_gs(gs,file = "~/gs")
G1 <- load_gs(file = "~/gs")</pre>

We also provide the clone method to make a full copy of an existing GatingSet:

Note that the GatingSet contains environment slots and external pointer that point to the internal C data structure. So make sure to use these methods in order to save or make a copy of existing object. The regular R assignment (<-) or save routine doesn't work as expected for the GatingSet object.

4.6 Exporting to FlowJo OSX 9.2

The exportAsFlowJoXML function can be used to export a flowCore::workFlow as an XML workspace for FlowJo 9.2 OSX. If flowWorkspace has been used to import an existing FlowJo workspace, flowWorkspace2flowCore can be used to obtain a workFlow for exporting. Currently this function can export one workFlow at a time.

4.7 Deprecated Functionality

The following behaviour is no longer supported and has been replace by more extensive netCDF support via the ncdfFlow package. If you have particularly large data files (millions of events), then you won't want to keep the data around, nor the indices for gate membership. Instead, pass the options cleanup=TRUE, keep.indices=FALSE to the execute() function, and the data will be scrubbed after computing population statistics. With future improvements making use of the netCDF framework, and bitvector representations of population memberships; this will improve memory usage in high-throughput unsupervised analysis settings.

5 Troubleshooting

If this package is throwing errors when parsing your workspace, and you are certain your workspace is version 2.0, contact the package author. If you can send your workspace by email, we can test, debug, and fix the package so that it works for you. Our goal is to provide a tool that works, and that people find useful.

6 Future Improvements

We are working on support for flowJo XML workspaces exported from the Windows version of flowJo. Efforts are underway to integrate GatingSet and GatingHierarchy objects more closely with the rest of the flow infrastructure.