# ncdfFlow: Provides netCDF storage based methods and functions for manipulation of flow cytometry data

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#### Abstract

Background The Bioconductor package flowCore is the object model and a collection of standard tools designed for flow cytometry data analysis. The related R packages including data analysis (flowClust, flowMerge, flowMeans,flowTrans,flowStats), visualization (flowViz) and quality control (flowQ) use the flowCore infrastructure to deal with flow cytometry data. However the flowFrame or flowSet which represent a single or a set of FCS files are memory-resident data structures and require the entire data elements to remain in memory in order to perform all kinds of the data manipulations. Hundreds or thousands of datasets generated by high throughput instruments can easily hit the memory limit if they are imported as the flowSet or flowFrames in R. It presents a challenge to scientists and bioinformaticians who use the R tools described above to perform statistical data analysis on a regular computer. We propose a new R object model and related functions to address this problem. The new model ncdfFlowSet inherit most of data structures from flowSet. It stores the large volume of event-level data on disk and only keeps the file handler and meta data in memory. Thus the memory consumption is significantly reduced. NetCDF is used as the data formats because it is self-describing, machine-independent and specifically optimized for storing and accessing array-oriented scientific data. With the compression and chunking features introduced by the new release of netCDF4, the new model is able to maintain high performance of data processing.

Most of the functions and methods including transformation, compensation, gating and subsetting methods for flowSet are extended to ncdfFlowSet (spillover,normalize and workflow methods of flowCore are currently not supported yet.). Thus the change of data structure is almost transparent to the users of flowCore-based R packages. keywords Flow cytometry, high throughput, netCDF, flowSet, ncdfFlowSet

## 1 Representing Flow Cytometry Data with ncdfFlowSet

ncdfFlow reppresents a flow cytometry data model that is very similar to the flowSet structure. The only difference is that the event-based 2-D data matrices from multiple samples of the same experiment are stored as one single 3D data matrix on disk in ncdf format. Each sample can be accessed efficiently from the 3-D matrix as a data chunk or slice and further manipulated in memory.

The basic unit of manipulation in ncdfFlow is the ncdfFlowSet. It inherites all the slots from flowSet. However, the flowFrame objects stored in the "frames" slot of a ncdfFlowSet object do not host the data matrix.Instead, their the "exprs" lots are kepted empty and the

actual data are stored in the 3-D data matrix on disk and only the file name is stored in "file" slot of ncdfFlowSet. Thus ncdfFlowSet reduces the memory requirements and meanwhile ensures the consistent data structure with flowSet.

### 2 Creating a ncdfFlowSet

We provide a function to read FCS files into a ncdfFlowSet object:

```
> path<-system.file("extdata","compdata","data",package="flowCore")
> files<-list.files(path,full.names=TRUE)[1:3]
> nc1 <- read.ncdfFlowSet(files=files,ncdfFile="ncfsTest.nc")
> nc1

An ncdfFlowSet with 3 samples.
flowSetId :
NCDF file : ncfsTest.nc
An object of class "AnnotatedDataFrame"
  rowNames: 060909.001 060909.002 060909.003
  varLabels: name
  varMetadata: labelDescription

column names:
  FSC-H, SSC-H, FL1-H, FL2-H, FL3-H, FL1-A, FL4-H
```

As we see, the contructor function is very similar to the *flowSet* execpt that it requires a filename for the ncdf file.

```
> fs1 <- read.flowSet(files=files)</pre>
```

Note that an ncdf file that stores the actual data is generated and saved on disk once a ncdfFlowSet is created. Users need to explicitely call the ncfsUnlink method to remove the file before delete the object from memory by rm.

```
> ncfsUnlink(nc1)
> rm(nc1)
```

Sometime it is necessary to save both data matrix and meta data on disk and reload the ncdfFlowSet object later. In order to do this, isSaveMeta argument has to be set to TRUE when the ncdfFlowSet is created by read.ncdfFlowSet.ncdfFlowSet\_sync can be also called to save the meta data explicitly in cdf file.

```
> nc1 <- read.ncdfFlowSet(files=files,ncdfFile="ncfsTest.nc",isSaveMeta=TRUE)
> rm(nc1)
> nc1<-ncdfFlowSet_open("ncfsTest.nc")
> nc1[[1]]
> ncfsUnlink(nc1)
> rm(nc1)
```

Users can also create an empty ncdfFlowSet first and add data slices later by assigning argument isWriteSlice as FALSE.

> nc1 <- read.ncdfFlowSet(files=files,ncdfFile="ncfsTest.nc",isWriteSlice= FALSE)
> nc1[[1]]

flowFrame object '060909.001' with 0 cells and 7 observables:

	name	desc	range	minRange	maxRange
\$P1	${\tt FSC-H}$	FSC-Height	1024	0	1023
\$P2	SSC-H	${\tt SSC-Height}$	1024	0	1023
\$P3	FL1-H	<na></na>	1024	1	10000
\$P4	FL2-H	<na></na>	1024	1	10000
\$P5	FL3-H	<na></na>	1024	1	10000
\$P6	FL1-A	<na></na>	1024	0	1023
\$P7	FL4-H	<na></na>	1024	1	10000
	_	_			

139 keywords are stored in the 'description' slot

As we see here, before add the actual flowFrame by addFrame, the flowFrame object returned by [[ has 0 events.

- > targetSampleName<-sampleNames(fs1)[1]</pre>
- > addFrame(nc1,fs1[[1]],targetSampleName)
- > nc1[[1]]

flowFrame object '060909.001'

with 10000 cells and 7 observables:

	name	desc	range	${\tt minRange}$	maxRange
\$P1	FSC-H	FSC-Height	1024	0	1023
\$P2	SSC-H	SSC-Height	1024	0	1023
\$P3	FL1-H	<na></na>	1024	1	10000
\$P4	FL2-H	<na></na>	1024	1	10000
\$P5	FL3-H	<na></na>	1024	1	10000
\$P6	FL1-A	<na></na>	1024	0	1023
\$P7	FL4-H	<na></na>	1024	1	10000

141 keywords are stored in the 'description' slot

> nc1[[2]]

flowFrame object '060909.002'

with 0 cells and 7 observables:

	name		desc	range	${\tt minRange}$	maxRange
\$P1	${\tt FSC-H}$	${\tt Forward}$	${\tt Scatter}$	1024	0	1023
\$P2	${\tt SSC-H}$	Side	Scatter	1024	0	1023
\$P3	FL1-H		FITC	1024	1	10000
\$P4	FL2-H		<na></na>	1024	1	10000

\$P5	FL3-H		<na< th=""><th><i>4</i>&gt;</th><th>1024</th><th>Ŀ</th><th>1</th><th>10000</th></na<>	<i>4</i> >	1024	Ŀ	1	10000
\$P6	FL1-A		< N A	<i>\</i> >	1024		0	1023
\$P7	FL4-H		< N A	<i>\</i> >	1024	Ŀ	1	10000
144	keywords	are	stored	in	the	'descrip	tion'	slot

Note that it is important to always use sample name to specify the target position in the data matrix where the actual is added. Because the sample name is the identifier used to index the data matrix.

Sometime it is helpful to copy the structure from an existing ncdfFlow object and then add the data slice to the empty ncdfFlow cloned by clone.ncdfFlowSet.

- > nc2<-clone.ncdfFlowSet(nc1,"clone.nc")
  > nc2[[1]]
- > addFrame(nc2,fs1[[1]],sampleNames(fs1)[1])
- > nc2[[1]]

We also provide the coerce function to convert the flowSet to a ncdfFlowSet.

- > data(GvHD)
- > GvHD <- GvHD[pData(GvHD)\$Patient %in% 6:7][1:4]
- > nc1<-ncdfFlowSet(GvHD)
- [1] "ncdfFlowSet saved in /tmp/RtmpIMzn1g/ncfs4310195c54da.nc"

Or coerce a ncdfFlowSet to flowSet

> fs1<-NcdfFlowSetToFlowSet(nc1,top=2)

Note that ncdfFlowSet is designed to store large datasets and it is not recommened to corece the entire ncdfFlowSet to flowSet. Usually users want to select a subset from ncdfFlowSet by [ and convert the subetted data. Sometimes it is helpful to randomly select a cerntain number of flowFrames from the entire datasets represented by by ncdfFlowSet to have a preview of the data. The arugment "top" can be used here for this purpose.

### 3 Working with metadata

Like flowSet,ncdfFlowSet has an associated AnnotatedDataFrame that provides metadata of experiments. This data frame is accessed and modified via the same methods of flowCore. :

- > phenoData(nc1)
- > pData(nc1)
- > varLabels(nc1)
- > varMetadata(nc1)
- > sampleNames(nc1)
- > keyword(nc1,"FILENAME")
- > identifier(nc1)
- > colnames(nc1)
- > colnames(nc1,prefix="s6a01")
- > length(nc1)
- > getIndices(nc1, "s6a01")

### Manipulating a ncdfFlowSet

You can extract a flowFrame from a ncdfFlowSet object in the same way as flowSet by using the [[ or \$ extraction operators. Note that using the [ extraction operator returns a new ncdfFlowSet that points to the same ncdf file. SO the original ncdf file serves as a data repository and the ncdfFlowSet works as view of the data in this sense.

1023

1023

10000

10000

10000

1023

10000

1023

0

```
> nm<-sampleNames(nc1)[1]
> expr1<-paste("nc1$'",nm,"'",sep="")</pre>
> eval(parse(text=expr1))
flowFrame object 's6a01'
with 2205 cells and 8 observables:
     name
                        desc range minRange maxRange
$P1 FSC-H
                  FSC-Height 1024
                                           0
$P2 SSC-H
                  SSC-Height
                              1024
                                           0
$P3 FL1-H
                   CD15 FITC 1024
                                           1
$P4 FL2-H
                     CD45 PE 1024
                                           1
$P5 FL3-H
                  CD14 PerCP 1024
                                           1
$P6 FL2-A
                         <NA>
                              1024
                                           0
$P7 FL4-H
                    CD33 APC
                              1024
                                           1
```

> nc1[[nm]]

flowFrame object 's6a01'

\$P8 Time Time (102.40 sec.)

with 2205 cells and 8 observables:

	name			desc	range	${\tt minRange}$	${\tt maxRange}$
\$P1	${\tt FSC-H}$		FSC-I	Height	1024	0	1023
\$P2	SSC-H		SSC-I	Height	1024	0	1023
\$P3	FL1-H		CD15	5 FITC	1024	1	10000
\$P4	FL2-H		CI	045 PE	1024	1	10000
\$P5	FL3-H		CD14	${\tt PerCP}$	1024	1	10000
\$P6	FL2-A			<na></na>	1024	0	1023
\$P7	FL4-H		CD3	33 APC	1024	1	10000
\$P8	Time	Time	(102.40	sec.)	1024	0	1023
150	keywoi	rds aı	e store	d in th	ne 'des	scription	' slot

150 keywords are stored in the 'description' slot

1024

- > nm < -sampleNames(nc1)[c(1,3)]
- > nc2<-nc1[nm]
- > summary(nc2)

#### \$s6a01

FSC-H SSC-H FL1-H FL2-H FL3-H FL2-A FL4-H Time 60.0 1.000 1.00 1.000 0.0 1.000 11.0 Min. 0.0

```
1st Qu.
         159.0
                  48.0
                           1.046
                                   35.35
                                            1.000
                                                    6.0
                                                           1.000 40.0
Median
         196.0
                  65.0
                          2.644
                                  160.40
                                            1.383
                                                   36.0
                                                           5.289 57.0
Mean
         220.8
                 108.9
                          57.540
                                  210.10
                                            7.367
                                                   48.7
                                                          16.240 51.9
         264.0
                  97.0
                          7.055
                                  320.90
                                            2.460
                                                   75.0
                                                          20.780 66.0
3rd Qu.
Max.
        1023.0 1023.0 3782.000 1637.00 326.700 516.0 503.300 80.0
$s6a03
         FSC-H SSC-H
                          FL1-H
                                  FL2-H
                                            FL3-H
                                                  FL2-A
                                                            FL4-H
                                                                   Time
          59.0
                           1.000
                                            1.000
                                                            1.000
Min.
                    0
                                    1.0
                                                     0.0
                                                                     0.0
         147.0
1st Qu.
                   49
                           1.000
                                  341.8
                                            1.000
                                                    79.0
                                                            1.165 105.0
Median
         192.0
                   71
                           1.145
                                  526.5
                                            1.070
                                                   124.0
                                                            2.228 215.5
         188.5
                                  543.7
                                            5.400
                                                   127.9
                                                            8.352 233.9
Mean
                  116
                          62.170
3rd Qu.
         226.0
                  119
                          10.200
                                  702.3
                                            2.208
                                                   164.0
                                                            4.834 353.0
        1023.0
                 1023 10000.000 8504.0 7565.000 1023.0 665.400 567.0
Max.
```

flowSet-specific iterator fsApply can also be applied to RobjectncdfFlowSet:

> fsApply(nc1,range)

> rm(nc2)

> fsApply(nc1, each\_col, median)

However, we recmmend to use another iterator ncfsApply designed for the function that returns a flowFrame (such as compensate,transform...). ncfsApply works the same as fsApply execpt that it returns a ncdfFlowSet object with the acutal data stored in cdf to avoid the huge memory consumption. Note that the return value of the function applied in ncfsApply must be a flowFrame object and it should have the same dimensions(channles and events) as the original data.

## 5 Compensation, Transformation and Gating

transform amd compensate for ncdfFlowSet work the same as flowSet.

```
> cfile <- system.file("extdata","compdata","compmatrix", package="flowCore")
> comp.mat <- read.table(cfile, header=TRUE, skip=2, check.names = FALSE)
> comp <- compensation(comp.mat)
> #compensation
> summary(nc1)[[1]]
> nc2<-compensate(nc1, comp)
> summary(nc2)[[1]]
> ncfsUnlink(nc2)
> rm(nc2)
> #transformation
> asinhTrans <- arcsinhTransform(transformationId="ln-transformation", a=1, b=1, c=1)
> nc2 <- transform(nc1, `FSC-H`=asinhTrans(`FSC-H`))
> summary(nc1)[[1]]
> summary(nc2)[[1]]
> ncfsUnlink(nc2)
```

Note that compensation/transformation return the ncdfFlowSet objects that point to the new cdf file containing the compensated/transformed data.

filter for flowSet also works for ncdfFlowSet:

```
> rectGate <- rectangleGate(filterId="nonDebris", "FSC-H"=c(200,Inf))
> fr <- filter(nc1,rectGate)
> summary(fr)
> rg2 <- rectangleGate(filterId="nonDebris", "FSC-H"=c(300,Inf))
> rg3 <- rectangleGate(filterId="nonDebris", "FSC-H"=c(400,Inf))
> flist <- list(rectGate, rg2, rg3)
> names(flist) <- sampleNames(nc1[1:3])
> fr3 <- filter(nc1[1:3], flist)
> summary(fr3[[3]])
```

### 6 Subsetting

> summary(nc2[[1]])

The Subset and split methods for *ncdfFlowSet*:

```
> nc2 <- Subset(nc1,rectGate)</pre>
> summary(nc2[[1]])
        FSC-H SSC-H
                        FL1-H
                                 FL2-H
                                         FL3-H FL2-A
                                                          FL4-H Time
Min.
          200
                   0
                        1.000
                                  1.00
                                         1.000
                                                  0.00
                                                          1.000 11.00
1st Qu.
          230
                  69
                        1.334
                                 22.33
                                         1.000
                                                  3.00
                                                          2.692 40.00
Median
          266
                  87
                        3.372
                                 77.37
                                         1.500
                                                 16.00 12.900 57.00
Mean
          297
                       91.900
                               165.70
                                        10.830
                                                 38.24
                                                        22.020 51.82
                 141
3rd Qu.
          316
                 122
                       18.990
                               223.80
                                         2.551
                                                 53.00
                                                        29.790 66.00
Max.
         1023
                1023 1943.000 1637.00 326.700 516.00 464.200 80.00
> nc2 <- Subset(nc1, fr)</pre>
```

```
FSC-H SSC-H
                                         FL3-H FL2-A
                        FL1-H
                                FL2-H
                                                         FL4-H Time
Min.
          200
                   0
                        1.000
                                 1.00
                                         1.000
                                                 0.00
                                                         1.000 11.00
1st Qu.
          230
                  69
                        1.334
                                22.33
                                         1.000
                                                 3.00
                                                         2.692 40.00
Median
          266
                 87
                        3.372
                                77.37
                                         1.500
                                                16.00 12.900 57.00
          297
                               165.70
                                                38.24
Mean
                 141
                       91.900
                                        10.830
                                                        22.020 51.82
3rd Qu.
                 122
                       18.990
                                         2.551
                                                53.00 29.790 66.00
          316
                               223.80
Max.
         1023
               1023 1943.000 1637.00 326.700 516.00 464.200 80.00
```

```
> rm(nc2)
```

<sup>&</sup>gt; morphGate <- norm2Filter("FSC-H", "SSC-H", filterId = "MorphologyGate",scale = 2)

<sup>&</sup>gt; smaller <- Subset(nc1[c(1,3)], morphGate,c("FSC-H", "SSC-H"))</pre>

<sup>&</sup>gt; smaller[[1]]

#### > nc1[[1]]

flowFrame object 's6a01' with 2205 cells and 8 observables:

	name		desc	range	minRange	${\tt maxRange}$	
\$P1	FSC-H		FSC-Height	1024	0	1023	
\$P2	SSC-H		SSC-Height	1024	0	1023	
\$P3	FL1-H		CD15 FITC	1024	1	10000	
\$P4	FL2-H		CD45 PE	1024	1	10000	
\$P5	FL3-H		CD14 PerCP	1024	1	10000	
\$P6	FL2-A		<na></na>	1024	0	1023	
\$P7	FL4-H		CD33 APC	1024	1	10000	
\$P8	Time	Time	(102.40 sec.)	1024	0	1023	
150 keywords are stored in the !description! slot							

150 keywords are stored in the 'description' slot

#### > rm(smaller)

Note that Subset does not create the new ncdf file (the same as extraction operator [). So we need to be careful about using ncfsUnlink to delete the subsetted data since it points to the same ncdf file that is also used by the original ncdfFlowSet object.

split returns a ncdfFlowList object which is a container of multiple ncdfFlowSet objects.

- > ##splitting by a gate
- > qGate <- quadGate(filterId="qg", "FSC-H"=200, "SSC-H"=400)</pre>
- > fr<-filter(nc1,qGate)
- > ncList<-split(nc1,fr)
- > ncList

An ncdfFlowList with 4 ncdfFlowSets containing 4 unique samples.

#### > nc1[[1]]

flowFrame object 's6a01'

with 2205 cells and 8 observables:

	name	desc	range	minRange	maxRange
\$P1	FSC-H	FSC-Height	1024	0	1023
\$P2	SSC-H	SSC-Height	1024	0	1023
\$P3	FL1-H	CD15 FITC	1024	1	10000
\$P4	FL2-H	CD45 PE	1024	1	10000

```
$P5 FL3-H
                CD14 PerCP 1024
                                        1
                                             10000
                                              1023
$P6 FL2-A
                       <NA> 1024
                                        0
$P7 FL4-H
                   CD33 APC 1024
                                        1
                                             10000
$P8 Time Time (102.40 sec.) 1024
                                              1023
150 keywords are stored in the 'description' slot
```

### > ncList[[2]][[1]]

flowFrame object 's6a01'

with 36 cells and 8 observables:

	name			desc	range	${\tt minRange}$	${\tt maxRange}$
\$P1	FSC-H		FSC-H	Height	1024	0	1023
\$P2	SSC-H		SSC-I	Height	1024	0	1023
\$P3	FL1-H		CD15	5 FITC	1024	1	10000
\$P4	FL2-H		CI	)45 PE	1024	1	10000
\$P5	FL3-H		CD14	${\tt PerCP}$	1024	1	10000
\$P6	FL2-A			<na></na>	1024	0	1023
\$P7	FL4-H		CD3	33 APC	1024	1	10000
\$P8	Time	Time	(102.40	sec.)	1024	0	1023
150 keywords are stored in the 'description' slot							

## > ncList[[1]][[1]]

flowFrame object 's6a01'

with 74 cells and 8 observables:

	name		desc	range	minRange	maxRange
\$P1	${\tt FSC-H}$		FSC-Height	1024	0	1023
\$P2	SSC-H		SSC-Height	1024	0	1023
\$P3	FL1-H		CD15 FITC	1024	1	10000
\$P4	FL2-H		CD45 PE	1024	1	10000
\$P5	FL3-H		CD14 PerCP	1024	1	10000
\$P6	FL2-A		<na></na>	1024	0	1023
\$P7	FL4-H		CD33 APC	1024	1	10000
\$P8	Time	Time	(102.40 sec.)	1024	0	1023

150 keywords are stored in the 'description' slot

Keep in mind that the ncdfFlowSet objects contained in ncdfFlowList share the same ncdf file as the original ncdfFlowSet. So deleting ncdfFlowList by ncfsUnlink is also going to make the data not accessible to the original ncdfFlowSet.