

Package ‘flowUtils’

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Type Package

Title Utilities for flow cytometry

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Description Provides utilities for flow cytometry data.

Depends R (>= 2.2.0)

Imports Biobase, graph, methods, stats, utils, corpcor, RUnit, XML,
flowCore (>= 1.32.0)

Suggests gatingMLData

Collate AllClasses.R gatingML.R helperFunctions.R gate-methods.R
transforms.R parameter-methods.R compensation.R
workflow2FlowJo.R writeGatingML.R zzz.R

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biocViews ImmunoOncology, Infrastructure, FlowCytometry,
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URL <https://github.com/jspidlen/flowUtils>

BugReports <https://github.com/jspidlen/flowUtils/issues>

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`flowUtils-package` *Utilities for flow cytometry data*

Description

This package includes functions to import gates, transformations and compensations defined in compliance with Gating-ML specification version 1.5 and 2.0. This package depends on the flowCore package for methods to evaluate the Gating-ML files read into the workspace.

The main features of this package provide compatibility to the data standards defined by the Gating-ML specification version 1.5 and 2.0.

The package also includes a Test Suite, which allows the user to test whether the implementation of gates and transformations are in compliance with the Gating-ML standard.

Author(s)

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References

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Cytometry A. 2008 Dec; 73A(12):1151–7. doi: 10.1002/cyto.a.20637.
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<http://flowcyt.sf.net/gating/Gating-ML.v1.5.081030.Compliance-tests.081030.zip>

See Also

[flowCore](#)

`read.gatingML` *Function to parse a Gating-ML XML file into objects in the R environment*

Description

This function parses a Gating-ML XML file defined in compliance with the Gating-ML recommendation into objects in the R environment, which can then be evaluated using functions provided by the flowCore package.

Usage

```
read.gatingML(file, flowEnv, ...)
```

Arguments

file	Gating-ML XML file describing gates, transformations and/or compensations
flowEnv	environment into which the R objects created from the Gating-ML XML file are to be stored
...	additional arguments that are passed to the methods

Details

The Gating-ML specification has been developed as an interchange format for the description of gates relevant to a flow cytometry experiment. Presently, we can read Gating-ML versions 1.5 and 2.0 of the specification. Version 2.0 is the most recent at the time of this writing.

Author(s)

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References

- Spidlen J, ISAC DSTF, Brinkman RR. 2014.
Gating-ML 2.0. International Society for Advancement of Cytometry (ISAC) standard for representing gating descriptions in flow cytometry.
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<http://flowcyt.sf.net/gating/Gating-ML.v1.5.081030.Compliance-tests.081030.zip>

See Also

[write.gatingML](#)

Examples

```
library("flowCore")  
  
#####  
# Gating-ML 2.0 example #  
#####  
flowEnv <- new.env()  
  
fcsFile <- system.file("extdata/Gm12/FCSFiles",  
  "data1.fcs", package="gatingMLData")  
fcs <- read.FCS(fcsFile,  
  transformation="linearize-with-PnG-scaling")
```

```

gateFile <- system.file("extdata/Gml2/Gating-MLFiles",
  "gates1.xml", package="gatingMLData")
read.gatingML(gateFile, flowEnv)
ls(flowEnv)

result = filter(fcs, flowEnv$Polygon1)
summary(result)

#####
# Gating-ML 1.5 example #
#####
flowEnv <- new.env()

fcsFile <- system.file("extdata/List-modeDataFiles",
  "fcs2_int16_13367ev_8par_GvHD.fcs", package="gatingMLData")
fcs <- read.FCS(fcsFile, transformation=FALSE)

gateFile <- system.file("extdata/Gating-MLFiles",
  "02CtSRectangular.xml", package="gatingMLData")
read.gatingML(gateFile, flowEnv)
ls(flowEnv)

result <- filter(fcs, flowEnv$CtSR_03)
summary(result)

```

testGatingMLCompliance*Function to perform all the Gating-ML compliance tests***Description**

This function performs the Gating-ML compliance tests. Either Gating-ML 1.5 or Gating-ML 2.0 compatibility may be checked. The Gating-ML XML files, FCS data files and the expected results provided by the `gatingMLData` are utilized in performing the compliance tests. The results obtained are compared with the expected results and an HTML summary report is generated.

Usage

```
testGatingMLCompliance(file = "GatingMLComplianceReport", version = 2.0)
```

Arguments

<code>file</code>	Name of the file in which the generated Gating-ML compliance report is to be saved. The .html extension will be added.
<code>version</code>	The Gating-ML version that is supposed to be checked. Currently, versions 1.5 and 2.0 are supported.

Details

The `testGatingMLCompliance` function depends on the `gatingMLData` data package for performing the compliance tests.

Author(s)

Spidlen J., Gopalakrishnan N.

References

- Spidlen J, ISAC DSTF, Brinkman RR. 2014.
 Gating-ML 2.0. International Society for Advancement of Cytometry (ISAC) standard for representing gating descriptions in flow cytometry.
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Cytometry A. 2008 Dec; 73A(12):1151–7. doi: 10.1002/cyto.a.20637.
- Spidlen J, ISAC DSTF, Brinkman RR. 2008.
 Gating-ML Candidate Recommendation for Gating Description in Flow Cytometry version 1.5.
<http://flowcyt.sf.net/gating/Gating-ML.v1.5.081030.pdf>
<http://flowcyt.sf.net/gating/Gating-ML.v1.5.081030.full.zip>
<http://flowcyt.sf.net/gating/Gating-ML.v1.5.081030.Compliance-tests.081030.zip>

Examples

```
## Not run:
#####
### Performs Gating-ML 1.5 compliance tests and          #
### writes the results to Gating-ML_1.5_Compliance_Report.html #
#####
testGatingMLCompliance("Gating-ML_1.5_Compliance_Report", version = 1.5)

#####
### Performs Gating-ML 2.0 compliance tests and          #
### writes the results to Gating-ML_2.0_Compliance_Report.html #
#####
testGatingMLCompliance("Gating-ML_2.0_Compliance_Report", version = 2.0)

## End(Not run)
```

write.gatingML

Function to write a Gating-ML XML file based on gating and transformation objects stored in an R environment.

Description

This function saves gating and transformation objects stored in an R environment to a Gating-ML 2.0 XML file. The objects expected and supported in the R environment are those that can normally be created by the `read.gatingML` function when a Gating-ML 2.0 XML file is read.

Usage

```
write.gatingML(flowEnv, file = NULL)
```

Arguments

<code>flowEnv</code>	The R environment that is being searched for gating objects and transformations
<code>file</code>	The name of the output Gating-ML XML file. The standard output will be used if <code>file</code> is <code>NULL</code> .

Details

The Gating-ML specification has been developed as an interchange format for the description of gates relevant to a flow cytometry experiment. Presently, `flowUtils` can read Gating-ML versions 1.5 and 2.0 of the specification (see [read.gatingML](#)). Gating-ML version 2.0 only is being used when saving Gating-ML.

Author(s)

Spidlen, J.

References

Spidlen J, ISAC DSTF, Brinkman RR. 2014.
 Gating-ML 2.0. International Society for Advancement of Cytometry (ISAC) standard for representing gating descriptions in flow cytometry.
<http://flowcyt.sf.net/gating/20141009.pdf>
<http://flowcyt.sf.net/gating/20141009.full.zip>

See Also

[read.gatingML](#)

Examples

```
library("flowCore")

#####
# Read a Gating-ML file and write the objects back in Gating-ML #
#####
flowEnv <- new.env()
gateFile <- system.file("extdata/Gml2/Gating-MLFiles",
  "gates1.xml", package="gatingMLData")
read.gatingML(gateFile, flowEnv)
ls(flowEnv)
write.gatingML(flowEnv)

#####
# Create a quad gate and write it to Gating-ML #
#####
flowEnv=new.env()
myQuad <- quadGate(filterId = "myQuad", "FSC-A" = 15000,
  "SSC-A" = 16000)
flowEnv[['myQuad']] <- myQuad
write.gatingML(flowEnv)
#####
# If we wanted the output to a file instead: #
#####
gatingOutputFile <- tempfile(fileext=".gating-ml2.xml")
write.gatingML(flowEnv, gatingOutputFile)
```

```

#####
# Again a quad gate, but now adding compensation #
#####
flowEnv=new.env()
myCompQuad <- quadGate(filterId = "myCompQuad", "PE-A" = 100,
  "PerCP-Cy5-5-A" = 200)
compPars = list(
  compensatedParameter(parameters="PE-A", spillRefId="SpillFromFCS",
    transformationId=paste("PE-A", "_compensated_according_to_FCS",
    sep=""), searchEnv=flowEnv),
  compensatedParameter(parameters="PerCP-Cy5-5-A", spillRefId="SpillFromFCS",
    transformationId=paste("PerCP-Cy5-5-A", "_compensated_according_to_FCS",
    sep=""), searchEnv=flowEnv)
)
myCompQuad@parameters = new("parameters", compPars)
flowEnv[['myCompQuad']] <- myCompQuad
write.gatingML(flowEnv)

#####
# Again a quad gate, but now adding a scaling transformation #
#####
flowEnv=new.env()
myTrQuad <- quadGate(filterId = "myTrQuad", "APC-A" = 0.5, "APC-Cy7-A" = 0.5)
trArcSinH1 = asinhtGml2(parameters = "APC-A",
  T = 1000, M = 4.5, A = 0, transformationId="trArcSinH1")
trLogicle1 = logicleGml2(parameters = "APC-Cy7-A",
  T = 1000, W = 0.5, M = 4.5, A = 0, transformationId="trLogicle1")
flowEnv[['trArcSinH1']] <- trArcSinH1
flowEnv[['trLogicle1']] <- trLogicle1
trPars = list(
  transformReference("trArcSinH1", flowEnv),
  transformReference("trLogicle1", flowEnv)
)
myTrQuad@parameters = new("parameters", trPars)
flowEnv[['myTrQuad']] <- myTrQuad
write.gatingML(flowEnv)

#####
# Now, we will be adding both scaling transformation and compensation #
# Also demonstrating what happens if 'bad' characters are part of the #
# name                                     #
#####
flowEnv=new.env()
myTrCompQuad <- quadGate(filterId = "myTr!Comp Quad", "APC-A" = 0.5,
  "APC-Cy7-A" = 0.5)
trArcSinH2 = asinhtGml2(parameters = "APC-A",
  T = 1000, M = 4, A = 0, transformationId="trArcSinH2")
trLogicle2 = logicleGml2(parameters = "APC-Cy7-A",
  T = 1000, W = 0.3, M = 4.5, A = 0, transformationId="trLogicle2")
trArcSinH2@parameters = compensatedParameter(parameters="APC-A",
  spillRefId="SpillFromFCS", transformationId=paste("FL3-H",
  "_compensated_according_to_FCS", sep=""), searchEnv=flowEnv)
trLogicle2@parameters = compensatedParameter(parameters="APC-Cy7-A",
  spillRefId="SpillFromFCS", transformationId=paste("FL4-H",
  "_compensated_according_to_FCS", sep=""), searchEnv=flowEnv)

```

```

trPars = list(trArcSinH2,trLogic2)
myTrCompQuad@parameters = new("parameters", trPars)
flowEnv[['myTr!Comp Quad']] <- myTrCompQuad
write.gatingML(flowEnv)

#####
# Creating a rectangle gate on a ratio of two parameters and #
# saving the result to a Gating-ML file. #
#####
flowEnv=new.env()
rat1 <- ratio("FSC-A", "SSC-A", transformationId = "rat1")
gate1 <- rectangleGate(filterId="gate1", "rat1"=c(0.8, 1.4))
gate1@parameters = new("parameters", list(rat1))
flowEnv[['gate1']] <- gate1
trArcSinH = asinhtGml2(parameters = "rat2",
    T = 1000, M = 4.5, A = 0, transformationId="trArcSinH")
rat2 <- ratio("FSC-A", "APC-A", transformationId = "rat2")
trArcSinH@parameters = rat2
gate2 <- rectangleGate(filterId="gate2", "rat2"=c(0.6, 1.3))
gate2@parameters = new("parameters", list(trArcSinH))
flowEnv[['gate2']] <- gate2
write.gatingML(flowEnv)

#####
# Example with an ellipse gate on compensated parameters #
#####
flowEnv <- new.env()
covM <- matrix(c(62.5, 37.5, 37.5, 62.5), nrow = 2, byrow=TRUE)
colnames(covM) <- c("FL1-H", "FL2-H")
compPars <- list(
  compensatedParameter(parameters="FL1-H", spillRefId="SpillFromFCS",
    transformationId=paste("FL1-H", "_compensated_according_to_FCS", sep=""),
    searchEnv=flowEnv),
  compensatedParameter(parameters="FL2-H", spillRefId="SpillFromFCS",
    transformationId=paste("FL2-H", "_compensated_according_to_FCS", sep=""),
    searchEnv=flowEnv)
)
myEl <- ellipsoidGate(mean=c(12, 16), distance=1, .gate=covM, filterId="myEl")
myEl@parameters <- new("parameters", compPars)
flowEnv[['myEl']] <- myEl
write.gatingML(flowEnv)

#####
# Creating some Boolean gates and saving the result to a Gating-ML file. #
#####
flowEnv=new.env()
rg1 <- rectangleGate(filterId="rg1", list("FSC-A"=c(0200, 16000),
  "SSC-A"=c(0, 34000)))
rg2 <- rectangleGate(filterId="rg2", list("PE-A"=c(100, 8000),
  "APC-Cy7-A"=c(0, 59000)))
orGate <- new("unionFilter", filterId="orGate",
  filters=list(rg1, rg2))
flowEnv[['orGate']] <- orGate
andGate <- new("intersectFilter", filterId="andGate",
  filters=list(rg1, rg2))
flowEnv[['andGate']] <- andGate
notGate <- new("complementFilter", filterId="notGate",

```

```
filters=list(rg1))
flowEnv[['notGate']] <- notGate
parentGate <- new("subsetFilter", filterId="parentGate",
  filters=list(rg1, rg2))
flowEnv[['parentGate']] <- parentGate
write.gatingML(flowEnv)
#####
# Or if we wanted to write to a file instead... #
#####
gatingOutputFile <- tempfile(fileext=".gating-ml2.xml")
write.gatingML(flowEnv, gatingOutputFile)

#####
# A few of the Gating-ML 1.5 transforms can be converted to Gating-ML 2.0 #
# and therefore be used with the write.gatingML function.                 #
#####
flowEnv=new.env()
trArcSinhGml1.5 = asinht(parameters = "APC-A", a = 1, b = 1,
  transformationId="trArcSinHGml1.5")
gateAsinhGml1.5 <- rectangleGate(filterId="gateAsinhGml1.5",
  "trArcSinHGml1.5"=c(0.3, 4.7))
gateAsinhGml1.5@parameters = new("parameters", list(trArcSinHGml1.5))
flowEnv[['gateAsinhGml1.5']] <- gateAsinhGml1.5
gatingOutputFile <- tempfile(fileext=".gating-ml2.xml")
write.gatingML(flowEnv, gatingOutputFile)
```

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