# Package 'CytoML'

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Type Package

Title A GatingML Interface for Cross Platform Cytometry Data Sharing

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**Description** Uses platform-specific implemenations of the GatingML2.0 standard to exchange gated cytometry data with other software platforms.

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LazyData TRUE

Imports cytolib(>= 2.3.10), flowCore (>= 1.99.10), flowWorkspace (>= 4.1.8), openCyto (>= 1.99.2), XML, data.table, jsonlite, RBGL, Rgraphviz, Biobase, methods, graph, graphics, utils, jsonlite, dplyr, grDevices, methods, ggcyto (>= 1.11.4), yaml, stats, tibble

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Suggests testthat, flowWorkspaceData , knitr, rmarkdown, parallel

VignetteBuilder knitr

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BugReports https://github.com/RGLab/CytoML/issues

URL https://github.com/RGLab/CytoML

Collate 'AllClasses.R' 'GatingSet2cytobank.R' 'GatingSet2flowJo.R' 'gate-methods.R' 'compensation.R' 'cpp11.R' 'cytobank2GatingSet.R' 'cytobankExperiment.R' 'cytolibml\_bin\_path.R' 'flowJoWorkspace\_Methods.R' 'diva2GatingSet.R' 'flowUtils\_functions.R' 'gatingML.R' 'read.gatingML.cytobank.R' 'graphGML\_methods.R' 'helperFunctions.R' 'parameter-methods.R' 'transforms.R' 'utils.R' 'writeGatingML.R'

SystemRequirements xml2, GNU make, C++11

Contents

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addCustomInfo add customInfo nodes to each gate node and add BooleanAndGates

## Description

add customInfo nodes to each gate node and add BooleanAndGates

# Usage

addCustomInfo(root, gs, flowEnv, cytobank.default.scale = TRUE, showHidden)

# Arguments

root	the root node of the XML	
gs	a GatingSet object	
flowEnv	the environment that stores the information parsed by 'read.GatingML'.	
cytobank.default.scale		
	logical flag indicating whether to use the default Cytobank asinhtGml2 settings. Currently it should be set to TRUE in order for gates to be displayed properly in Cytobank because cytobank currently does not parse the global scale settings from GatingML.	
showHidden	whether to include the hidden population nodes in the output	

#### Value

XML root node

ce\_get\_channels Extract channels from cytobank\_experiment

## Description

Extract channels from cytobank\_experiment

## Usage

```
ce_get_channels(x, panel_name = NULL)
```

#### Arguments

х	A cytobank_experiment object
panel_name	select panel to process

ce\_get\_compensations Obtain the spillover matrices for the samples in a Cytobank experiment

# Description

Obtain the spillover matrices for the samples in a Cytobank experiment

# Usage

ce\_get\_compensations(x)

#### Arguments

x A cytobank\_experiment object

#### Value

A named list of spillover matrices

ce\_get\_markers Extract markers from cytobank\_experiment

#### Description

Extract markers from cytobank\_experiment

# Usage

```
ce_get_markers(x, panel_name = NULL)
```

#### Arguments

х	A cytobank_experiment object
panel_name	select panel to process

ce_get_panels	Obtain counts of the number of samples associated with each marker
	panel in a Cytobank experiment

# Description

Obtain counts of the number of samples associated with each marker panel in a Cytobank experiment

# Usage

```
ce_get_panels(x)
```

# Arguments

х

cytobank\_experiment object

#### Value

A tibble of panels with sample counts

<pre>ce_get_samples</pre>	Obtain a mapping between the samples and marker panels in a Cyto-
	bank experiment

# Description

Obtain a mapping between the samples and marker panels in a Cytobank experiment

# Usage

```
ce_get_samples(x)
```

#### Arguments

x A cytobank\_experiment object

# Value

A tibble with rows containing sample names and their associated panel names

#### ce\_get\_transformations

Obtain the transformations associated with each channel in a Cytobank experiment

#### Description

Obtain the transformations associated with each channel in a Cytobank experiment

## Usage

```
ce_get_transformations(x, panel_name = NULL)
```

#### Arguments

x	A cytobank_experiment object
panel_name	select panel to process

#### Value

A transformerList object containing transformation objects for each transformed channel

```
compensate,GatingSet,graphGML-method
~
```

compensate a GatingSet based on the compensation information stored in graphGML object

## Description

compensate a GatingSet based on the compensation information stored in graphGML object

## Usage

```
## S4 method for signature 'GatingSet,graphGML'
compensate(x, spillover, ...)
```

#### Arguments

х	GatingSet
spillover	graphGML
	unused.

## Value

compensated GatingSet

constructTree

# Description

Reconstruct the population tree from the GateSets

## Usage

```
constructTree(flowEnv, gateInfo)
```

#### Arguments

flowEn∨	the enivornment contains the elements parsed by read.gatingML function
gateInfo	the data.frame contains the gate name, fcs filename parsed by parse.gateInfo
	function

#### Value

a graphNEL represent the population tree. The gate and population name are stored as nodeData in each node.

cytobank\_experiment-methods

Methods for interacting with cytobank\_experiment objects

#### Description

These methods mirror similar accessor methods for the GatingSet class.

## Usage

```
## S4 method for signature 'cytobank_experiment'
markernames(object)
## S4 method for signature 'cytobank_experiment'
colnames(x, do.NULL = "missing", prefix = "missing")
## S4 method for signature 'cytobank_experiment'
sampleNames(object)
## S4 method for signature 'cytobank_experiment'
```

```
pData(object)
```

#### Arguments

object	A cytobank_experiment object
x	cytobank_experiment
do.NULL, prefix	not used

cytobank\_to\_gatingset A wrapper that parses the gatingML and FCS files (or cytobank\_experiment object) into GatingSet

#### Description

A wrapper that parses the gatingML and FCS files (or cytobank\_experiment object) into GatingSet

#### Usage

```
## Default S3 method:
cytobank_to_gatingset(x, FCS, trans = NULL, ...)
## S3 method for class 'cytobank_experiment'
cytobank_to_gatingset(x, panel_id = 1, ...)
```

## Arguments

x	the cytobank_experiment object or the full path of gatingML file
FCS	FCS files to be loaded
trans	a 'transfomerList' object to override the transformations from gatingML files. it is typically used by 'cytobank_experiment' parser(i.e. 'cytobank_to_gatingset.cytobank_experiment' to use the scales info recorded in yaml file.
	other arguments
panel_id	select panel to process

## Value

a GatingSet

#### Examples

```
## Not run:
acsfile <- system.file("extdata/cytobank_experiment.acs", package = "CytoML")
ce <- open_cytobank_experiment(acsfile)
xmlfile <- ce$gatingML
fcsFiles <- list.files(ce$fcsdir, full.names = TRUE)
gs <<- cytobank_to_gatingset(xmlfile, fcsFiles)
library(ggcyto)
autoplot(gs[[1]])
```

## End(Not run)

CytoML-deprecated Deprecated functions in package CytoML.

#### Description

```
GatingSet2cytobank -> gatingset_to_cytobank
GatingSet2flowJo -> gatingset_to_flowjo
cytobankExperiment -> open_cytobank_experiment
cytobank2GatingSet -> cytobank_to_gatingset
parseWorkspace -> flowjo_to_gatingset
getKeywords -> fj_ws_get_keywords
getSamples -> fj_ws_get_samples
getSampleGroups -> fj_ws_get_sample_groups
openDiva -> open_diva_xml
parseWorkspace -> diva_to_gatingset
```

diva\_get\_samples Get a table of samples from a FACSDiva workspace

#### Description

Return a data.frame of sample group information from a FACSDiva workspace

#### Usage

```
diva_get_samples(x)
diva_get_sample_groups(x)
```

#### Arguments

x A diva\_workspace

# Value

A data.frame with columns tub, name, and specimen

diva\_to\_gatingset Parse a FACSDiva Workspace

# Description

Function to parse a FACSDiva Workspace, generate a GatingHierarchy or GatingSet object, and associated flowCore gates.

# Usage

```
diva_to_gatingset(
    obj,
    name = NULL,
    subset = NULL,
    path = obj@path,
    worksheet = c("normal", "global"),
    swap_cols = list(`FSC-H` = "FSC-W", `SSC-H` = "SSC-W"),
    verbose = FALSE,
    ...
)
```

# Arguments

obj	diva_workspace
name	sample group to be parsed, either numeric index or the group name
subset	samples to be imported. either numeric index or the sample name. Default is NULL, which imports all samples.
path	the FCS data path
worksheet	select worksheet to import. either "normal" or "global"
swap_cols	diva seems to swap some data cols during importing fcs to experiments this argument provide a list to tell the parser which cols to be swapped default is list('FSC-H' = 'FSC-W', 'SSC-H' = 'SSC-W')
verbose	whether print more messages during the parsing
	other arguments

diva\_workspace-class An R representation of a BD FACSDiva workspace

# Description

Inherited from flowjo\_workspace-class

#### extend

#### Slots

version: Object of class "character". The version of the XML workspace.

file: Object of class "character". The file name.

.cache: Object of class "environment". An environment for internal use.

path: Object of class "character". The path to the file.

doc: Object of class "XMLInternalDocument". The XML document object.

options: Object of class "integer". The XML parsing options passed to xmlTreeParse.

#### See Also

GatingSet GatingHierarchy

extend

extend the gate to the minimum and maximum limit of both dimensions based on the bounding information.

#### Description

It is equivalent to the behavior of shifting the off-scale boundary events into the gate boundary that is described in bounding transformation section of gatingML standard.

```
extend(
  gate,
  bound,
  data.range = NULL,
  plot = FALSE,
  limits = c("original", "extended")
)
## S3 method for class 'polygonGate'
extend(
  gate,
  bound,
  data.range = NULL,
  plot = FALSE,
  limits = c("original", "extended")
)
## S3 method for class 'rectangleGate'
extend(gate, ...)
## S3 method for class 'ellipsoidGate'
extend(gate, ...)
```

gate	a flowCore filter/gate
bound	numeric matrix representing the bouding information parsed from gatingML. Each row corresponds to a channel. rownames should be the channel names. colnames should be c("min", "max")
data.range	numeric matrix specifying the data limits of each channel. It is used to set the extended value of vertices and must has the same structure as 'bound'. when it is not supplied, c(Machine\$integer.max,Machine\$integer.max) is used.
plot	whether to plot the extended polygon.
limits	character whether to plot in "extended" or "original" gate limits. Default is "orig- inal".
	other arguments

# Details

The advantage of extending gates instead of shifting data are two folds: 1. Avoid the extra computation each time applying or plotting the gates 2. Avoid changing the data distribution caused by adding the gates

Normally this function is not used directly by user but invoked when parsing GatingML file exported from Cytobank.

#### Value

a flowCore filter/gate

#### Examples

```
library(flowCore)
sqrcut <- matrix(c(300,300,600,600,50,300,300,50),ncol=2,nrow=4)
colnames(sqrcut) <- c("FSC-H","SSC-H")
pg <- polygonGate(filterId="nonDebris", sqrcut)
pg
bound <- matrix(c(100,3e3,100,3e3),
    byrow = TRUE, nrow = 2,
    dimnames = list(c("FSC-H", "SSC-H"),
        c("min", "max")))
bound
pg.extened <- extend(pg, bound, plot = TRUE)</pre>
```

fj\_ws\_get\_keywords Get Keywords

## Description

Retrieve keywords associated with a workspace

# Usage

fj\_ws\_get\_keywords(obj, y, ...)

obj	A flowjo_workspace
У	ccharacter or numeric specifying the sample name or sample ID
	other arguments sampNloc a character the location where the sample name is
	specified. See parseWorkspace for more details.

# Details

Retrieve a list of keywords from a flowjo\_workspace

#### Value

A list of keyword - value pairs.

#### Examples

```
## Not run:
    d<-system.file("extdata",package="flowWorkspaceData")
    wsfile<-list.files(d,pattern="manual.xml",full=TRUE)
    ws <- open_flowjo_xml(wsfile)
    fj_ws_get_samples(ws)
    res <- try(fj_ws_get_keywords(ws,"CytoTrol_CytoTrol_1.fcs"), silent = TRUE)
    print(res[[1]])
    fj_ws_get_keywords(ws, 1)
```

```
## End(Not run)
```

fj\_ws\_get\_samples Get a list of samples from a flowJo workspace

## Description

Return a data frame of samples contained in a flowJo workspace

# Usage

```
fj_ws_get_samples(x, group_id = NULL)
```

#### Arguments

х	A flowjo_workspace
group_id	integer specifies the group from which samples are returned

## Details

The samples with 0 populations are excluded. Returns a data.frame of samples in the flowjo\_workspace, including their sampleID, name

# Value

A data.frame with columns sampleID, name

#### Examples

```
## Not run:
    #ws is a flowjo_workspace
    fj_ws_get_samples(ws);
```

```
## End(Not run)
```

fj\_ws\_get\_sample\_groups

Get a table of sample groups from a flowJo workspace

#### Description

Return a data frame of sample group information from a flowJo workspace

## Usage

fj\_ws\_get\_sample\_groups(x)

#### Arguments

х

A flowjo\_workspace object.

#### Details

Note that the samples with 0 populations are also included (since count populations requires traversing xml for all samples thus can be expensive) Returns a table of samples and groups defined in the flowJo workspace

# Value

A data.frame containing the groupName, groupID, and sampleID for each sample in the workspace. Each sample may be associated with multiple groups.

# See Also

flowjo\_workspace-class flowjo\_to\_gatingset

## Examples

```
## Not run:
    #ws is a flowjo_workspace
    fj_ws_get_sample_groups(ws);
```

## End(Not run)

flowjo\_workspace-class

An R representation of a flowJo workspace.

#### Description

Objects can be created by calls of the form new("flowjo\_workspace.xml", ...).

# Slots

doc: Object of class "externalptr".

## See Also

GatingSet GatingHierarchy

## Examples

```
require(flowWorkspaceData)
d<-system.file("extdata",package="flowWorkspaceData")
wsfile<-list.files(d,pattern="A2004Analysis.xml",full=TRUE)
ws <- open_flowjo_xml(wsfile);
ws
fj_ws_get_samples(ws)</pre>
```

gatingset\_to\_cytobank Convert a GatingSet to a Cytobank-compatible gatingML

# Description

this function retrieves the gates from GatingSet and writes a customed GatingML-2.0 file that can be imported into cytobank.

```
gatingset_to_cytobank(
  gs,
  outFile,
  showHidden = FALSE,
  cytobank.default.scale = TRUE,
  ...
)
```

gs	a GatingSet object
outFile	a file name
showHidden	whether to include the hidden population nodes in the output
cytobank.defau	lt.scale
	logical flag indicating whether to use the default Cytobank asinhtGml2 settings. Currently it should be set to TRUE in order for gates to be displayed properly in Cytobank because cytobank currently does not parse the global scale settings from GatingML.
	rescale.gate default is TRUE. which means the gate is rescaled to the new scale that is understandable by cytobank. It is recommended not to change this behavior unless user wants to export to a gatingML file used for other purpose other than being imported into cytobank.

#### Details

The process can be divided into four steps: 1. Read in gate geometry, compensation and transformation from gatingSet 2. Rescale gate boundaries with flowjo\_biexp() so gates can be displayed properly in Cytobank 3. Save gates and hierarchy structure to R environment 4. Write environment out to gatingML using write.GatingML()

## Value

nothing

## Examples

```
library(flowWorkspace)
```

```
dataDir <- system.file("extdata",package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual",full = TRUE))</pre>
```

```
gs_pop_remove(gs, "CD8")
```

#output to cytobank
outFile <- tempfile(fileext = ".xml")
gatingset\_to\_cytobank(gs, outFile) #type by default is 'cytobank'</pre>

gatingset\_to\_flowjo Convert a GatingSet to flowJo workspace

#### Description

It is a R wrapper for the docker app (https://hub.docker.com/r/rglab/gs-to-flowjo)

```
gatingset_to_flowjo(gs, outFile, showHidden = FALSE, docker_img = NULL, ...)
```

gs	a GatingSet object or a folder contains the GatingSet archive (generated by pre- vious save_gs call)
outFile	the workspace file path to write
showHidden	whether to export hidden gates. Default is FALSE
docker_img	the docker image that does the actual work
	other arguments passed to save_gs

# Details

Docker images for gatingset\_to\_flowjo will be maintained at https://gallery.ecr.aws/x4k5d9i7/cytoverse/gs-to-wsp

docker pull public.ecr.aws/x4k5d9i7/cytoverse/gs-to-wsp:latest

#### Value

nothing

## Examples

```
## Not run:
library(flowWorkspace)
```

```
path <- system.file("extdata",package="flowWorkspaceData")
gs_path <- list.files(path, pattern = "gs_manual",full = TRUE)
gs <- load_gs(gs_path)</pre>
```

```
#output to flowJo
outFile <- tempfile(fileext = ".wsp")
gatingset_to_flowjo(gs, outFile)</pre>
```

```
#or directly use the archive as the input (to avoid the extra copying inside of the wrapper)
gatingset_to_flowjo(gs_path, outFile)
```

## End(Not run)

# Description

get children nodes

```
## S4 method for signature 'graphGML,character'
getChildren(obj, y)
```

obj	graphGML
У	character parent node path

## Value

a graphNEL node

### Examples

```
## Not run:
g <- read.gatingML.cytobank(xmlfile)
getChildren(g, "GateSet_722326")
getParent(g, "GateSet_722326")
```

## End(Not run)

getCompensationMatrices.graphGML

Extract compensation from graphGML object.

#### Description

Extract compensation from graphGML object.

#### Usage

```
## S3 method for class 'graphGML'
getCompensationMatrices(x)
```

#### Arguments

x graphGML

#### Value

compensation object or "FCS" when compensation comes from FCS keywords

## Description

get gate from the node

## Usage

```
## S4 method for signature 'graphGML,character'
getGate(obj, y)
```

obj	graphGML
У	character node path

# Value

the gate information associated with the node

getNodes,graphGML-method

get nodes from graphGML object

# Description

get nodes from graphGML object

# Usage

```
## S4 method for signature 'graphGML'
getNodes(x, y, order = c("default", "bfs", "dfs", "tsort"), only.names = TRUE)
```

### Arguments

х	graphGML
У	character node index. When missing, return all the nodes
order	character specifying the order of nodes. options are "default", "bfs", "dfs", "tsort"
only.names	logical specifying whether user wants to get the entire nodeData or just the name of the population node

## Value

It returns the node names and population names by default. Or return the entire nodeData associated with each node.

## Examples

```
## Not run:
g <- read.gatingML.cytobank(xmlfile)
getNodes(g)
getNodes(g, only.names = FALSE)
## End(Not run)
```

getParent,graphGML,character-method get parent nodes

# Description

get parent nodes

## Usage

## S4 method for signature 'graphGML,character'
getParent(obj, y)

## Arguments

obj	graphGML
У	character child node path

# Value

a graphNEL node

```
getTransformations.graphGML
```

Extract transformations from graphGML object.

#### Description

Extract transformations from graphGML object.

# Usage

```
## S3 method for class 'graphGML'
getTransformations(x, ...)
```

### Arguments

х	graphGML
	not used

## Value

transformerList object

graphGML-class

# Description

Each node corresponds to a population(or GateSet) defined in gatingML file. The actual gate object (both global and tailored gates) is associated with each node as nodeData. Compensation and transformations are stored in graphData slot.

#### Details

The class simply extends the graphNEL class and exists for the purpose of method dispatching.

gs\_compare\_cytobank\_counts

compare the counts to cytobank's exported csv so that the parsing result can be verified.

## Description

compare the counts to cytobank's exported csv so that the parsing result can be verified.

#### Usage

```
gs_compare_cytobank_counts(
  gs,
  file,
  id.vars = c("FCS Filename", "population"),
   ...
)
```

#### Arguments

gs	parsed GatingSet
file	the stats file (contains the populatio counts) exported from cytobank.
id.vars	either "population" or "FCS filename" that tells whether the stats file format is one population per row or FCS file per row.
	arguments passed to data.table::fread function

# Value

a data.table (in long format) that contains the counts from openCyto and Cytobank side by side.

## Examples

```
acsfile <- system.file("extdata/cytobank_experiment.acs", package = "CytoML")
ce <- open_cytobank_experiment(acsfile)
gs <- cytobank_to_gatingset(ce)
## verify the stats are correct
statsfile <- ce$attachments[1]
dt_merged <- gs_compare_cytobank_counts(gs, statsfile, id.vars = "population", skip = "FCS Filename")
all.equal(dt_merged[, count.x], dt_merged[, count.y], tol = 5e-4)</pre>
```

matchPath	Given the leaf node, try to find out if a collection of nodes can be
	matched to a path in a graph(tree) by the bottom-up searching

# Description

Given the leaf node, try to find out if a collection of nodes can be matched to a path in a graph(tree) by the bottom-up searching

## Usage

```
matchPath(g, leaf, nodeSet)
```

#### Arguments

g	graphNEL
leaf	the name of leaf(terminal) node
nodeSet	a set of node names

## Value

TRUE if path is found, FALSE if not path is matched.

open\_cytobank\_experiment

Construct a cytobank\_experiment object from ACS file

#### Description

Construct a cytobank\_experiment object from ACS file

## Usage

open\_cytobank\_experiment(acs, exdir = tempfile())

#### Arguments

acs	ACS file exported from Cytobank
exdir	he directory to extract files to

open\_diva\_xml

## Value

cytobank\_experiment object

open\_diva\_xml open Diva xml workspace

#### Description

open Diva xml workspace

# Usage

open\_diva\_xml(file, options = 0, ...)

#### Arguments

file	xml file
options	argument passed to xmlTreeParse
	arguments passed to xmlTreeParse

#### Value

a diva\_workspace object

## Examples

```
## Not run:
library(flowWorkspace)
library(CytoML)
ws <- open_diva_xml(system.file('extdata/diva/PE_2.xml', package = "flowWorkspaceData"))
ws
diva_get_sample_groups(ws)
gs <- diva_to_gatingset(ws, name = 2, subset = 1)
sampleNames(gs)
gs_get_pop_paths(gs)
plotGate(gs[[1]])
## End(Not run)
```

open\_flowjo\_xml Open/Close a flowJo workspace

#### Description

Open a flowJo workspace and return a flowjo\_workspace object. Close a flowjo\_workspace, destroying the internal representation of the XML document, and freeing the associated memory.

```
open_flowjo_xml(file, options = 0, sample_names_from = "keyword", ...)
```

file	Full path to the XML flowJo workspace file.
options	xml parsing options passed to xmlTreeParse. See http://xmlsoft.org/html/libxml-parser.html#xmlParserOption for details.
sample_names_from	
	character specifying where in the XML workspace file to obtain the sample names, either "keyword" for the included \$FIL keyword for each sample, or "sampleNode" for the name of the sample node
	not used

#### Details

Open an XML flowJo workspace file and return a flowjo\_workspace object. The workspace is represented using a XMLInternalDocument object. Close a flowJoWorkpsace after finishing with it. This is necessary to explicitly clean up the C-based representation of the XML tree. (See the XML package).

## Value

a flowjo\_workspace object.

#### Examples

```
## Not run:
file<-"myworkspace.xml"
ws<-open_flowjo_xml(file);
ws
```

## End(Not run)

parse.gateInfo Parse the cytobank custom\_info for each gate

#### Description

Fcs filename and gate name stored in 'custom\_info' element are beyong the scope of the gatingML standard and thus not covered by the default 'read.gatingML'.

## Usage

```
parse.gateInfo(file, ...)
```

#### Arguments

file	xml file path
	additional arguments passed to the handlers of 'xmlTreeParse'

# Value

a data.frame that contains three columns: id (gateId), name (gate name), fcs (fcs\_file\_filename).

parseWorkspace

## Description

Function to parse a flowJo Workspace, generate a GatingHierarchy or GatingSet object, and associated flowCore gates. The data are not loaded or acted upon until an explicit call to recompute() is made on the GatingHierarchy objects in the GatingSet.

## Usage

```
parseWorkspace(obj, ...)
## S4 method for signature 'flowjo_workspace'
parseWorkspace(obj, ...)
flowjo_to_gatingset(
  ws,
  name = NULL,
  subset = list(),
  execute = TRUE,
  path = "",
  cytoset = NULL,
  backend_dir = tempdir(),
  backend = get_default_backend(),
  includeGates = TRUE,
  additional.keys = "$TOT",
  additional.sampleID = FALSE,
  keywords = character(),
  keywords.source = "XML",
  keyword.ignore.case = FALSE,
  extend_val = 0,
  extend_to = -4000,
  channel.ignore.case = FALSE,
  leaf.bool = TRUE,
  include_empty_tree = FALSE,
  skip_faulty_gate = FALSE,
  compensation = NULL,
  transform = TRUE,
  fcs_file_extension = ".fcs",
  greedy_match = FALSE,
  mc.cores = 1,
)
```

#### Arguments

obj	flowjo_workspace
	Additional arguments to be passed to FCS parser
WS	A flowjo_workspace to be parsed.

name	numeric or character. The name or index of the group of samples to be imported. If NULL, the groups are printed to the screen and one can be selected interactively. Usually, multiple groups are defined in the flowJo workspace file.
subset	numeric vector specifying the subset of samples in a group to import. Or a character specifying the FCS filenames to be imported. Or an expression to be passed to 'subset' function to filter samples by 'pData' (Note that the columns referred by the expression must also be explicitly specified in 'keywords' argument)
execute	TRUE FALSE a logical specifying if the gates, transformations, and compensa- tion should be immediately calculated after the flowJo workspace have been imported. TRUE by default.
path	either a character scalar . it is a path to the fcs files that are to be imported. The code will search recursively, so you can point it to a location above the files.
cytoset	a cytoset object that provides the alternative data source other than FCS files. It is useful sometime to preprocess the raw fcs files (e.g. standardize channels using cytoqc package) and then directly use them for flowJo parsing. when cytoset is provided, path argument is ignored.
includeGates	logical Should gates be imported, or just the data with compensation and trans- formation?
additional.keys	
	character vector: The keywords (parsed from FCS header) to be combined(concatenated with "_") with FCS filename to uniquely identify samples. Default is '\$TOT' (total number of cells) and more keywords can be added to make this GUID.
additional.samp	leID
	boolean: A boolean specifying whether to include the flowJo sample ID in a GUID to uniquely identify samples. This can be helpful when the filename or other keywords are not enough to differentiate between samples. Default is FALSE.
keywords	character vector specifying the keywords to be extracted as pData of GatingSet
keywords.source	
	character the place where the keywords are extracted from, can be either "XML" or "FCS"
keyword.ignore.	
	a logical flag indicates whether the keywords matching needs to be case sen- sitive.
extend_val	numeric the threshold that determine wether the gates need to be extended. default is 0. It is triggered when gate coordinates are below this value.
extend_to	numeric the value that gate coordinates are extended to. Default is -4000. Usu- ally this value will be automatically detected according to the real data range. But when the gates needs to be extended without loading the raw data (i.e. execute is set to FALSE), then this hard-coded value is used.
channel.ignore.	case
	a logical flag indicates whether the colnames(channel names) matching needs to be case sensitive (e.g. compensation, gating)
leaf.bool	a logical whether to compute the leaf boolean gates. Default is TRUE. It helps to speed up parsing by turning it off when the statistics of these leaf boolean gates are not important for analysis. (e.g. COMPASS package will calculate them by itself.) If needed, they can be calculated by calling recompute method at later stage.

include_empty_tree	
	a logical whether to include samples that don't have gates.
skip_faulty_ga	ote
	a logical whether to skip the faulty gates so that the parser can still process the rest of gating tree.
compensation	a compensation object, matrix or data.frame or a list of these objects that al- low the customized compensation () to be used instead of the one specified in flowJo workspace or FCS file. When it is a list, its names is supposed to be matched to sample guids (Default is the fcs filename suffixed by \$TOT. See "ad- ditional.keys" arguments for details of guids) When some of the samples don't have the external compensations matched, it will fall back to the flowJo xml or FCS looking for the compensation matrix.
transform	logical to enable/disable transformation of gates and data. Default is TRUE. It is mainly for debug purpose (when the raw gates need to be parsed.), and only valid when execute is FALSE.
<pre>fcs_file_extension</pre>	
	default is ".fcs"
greedy_match	logical: By default, if flowjo_to_gatingset finds multiple FCS files matching a sample by total event count as well as sampleID and/or keywords specified by additional.keys and additional.sampleID, it will return an error listing the duplicate files. If greedy_match is TRUE, the method will simply take the first file with either filename or \$FIL keyword matching the sample name and having the correct number of events.
mc.cores	numeric the number of threads to pass to the C++ parser to run in parallel
h5_dir	the path to write h5 data

#### Details

A flowjo\_workspace is generated with a call to open\_flowjo\_xml(), passing the name of the xml workspace file. This returns a flowjo\_workspace, which can be parsed using the flowjo\_to\_gatingset() method. The function can be called non-interactively by passing the index or name of the group of samples to be imported via flowjo\_to\_gatingset(obj,name=x), where x is either the numeric index, or the name. The subset argument allows one to select a set of files from the chosen sample group. The routine will take the intersection of the files in the sample group, the files specified in subset and the files available on disk, and import them.

#### Value

a GatingSet, which is a wrapper around a list of GatingHierarchy objects, each representing a single sample in the workspace. The GatingHierarchy objects contain graphNEL trees that represent the gating hierarchy of each sample. Each node in the GatingHierarchy has associated data, including the population counts from flowJo, the parent population counts, the flowCore gates generated from the flowJo workspace gate definitions. Data are not yet loaded or acted upon at this stage. To execute the gating of each data file, a call to execute() must be made on each GatingHierarchy object in the GatingSet. This is done automatically by default, and there is no more reason to set this argument to FALSE.

#### See Also

fj\_ws\_get\_sample\_groups,GatingSet

#### Examples

```
## Not run:
 #f is a xml file name of a flowJo workspace
 ws <- open_flowjo_xml(f)</pre>
 #parse the second group
gs <- flowjo_to_gatingset(ws, name = 2); #assume that the fcs files are under the same folder as workspace
 gs <- flowjo_to_gatingset(ws, name = 4</pre>
                        , path = dataDir
                                               #specify the FCS path
                 , subset = "CytoTrol_CytoTrol_1.fcs") #subset the parsing by FCS filename
 gs <- flowjo_to_gatingset(ws, path = dataDir, name = 4</pre>
                 , keywords = c("PATIENT ID", "SAMPLE ID", "$TOT", "EXPERIMENT NAME") #tell the parser to extra-
                 , keywords.source = "XML" # keywords are extracted from xml workspace (alternatively can be see
                 , additional.keys = c("PATIENT ID") #use additional keywords together with FCS filename to uni
                 , execute = F) # parse workspace without the actual gating (can save time if just want to get th
#subset by pData (extracted from keywords)
gs <- flowjo_to_gatingset(ws, path = dataDir, name = 4</pre>
                          , subset = `TUBE NAME` %in% c("CytoTrol_1", "CytoTrol_2")
                          , keywords = "TUBE NAME")
#overide the default compensation defined in xml with the customized compenstations
gs <- flowjo_to_gatingset(ws, name = 2, compensation = comps); #comp is either a compensation object or a list or
```

## End(Not run)

plot,graphGML,missing-method

plot the population tree stored in graphGML.

#### Description

The node with dotted order represents the population that has tailored gates (sample-specific gates) defined.

#### Usage

```
## S4 method for signature 'graphGML,missing'
plot(x, y = "missing", label = c("popName", "gateName"))
```

## Arguments

Х	a graphNEL generated by constructTree function
У	not used
label	specifies what to be dispaled as node label. Can be either 'popName' (population name parsed from GateSets) or 'gateName'(the name of the actual gate associated with each node)

#### range.GatingHierarchy

#### Value

nothing

#### Examples

```
## Not run:
g <- read.gatingML.cytobank(xmlfile)
plot(g)
```

## End(Not run)

range.GatingHierarchy the parameter range from the flow data associated with GatingHierarchy

## Description

the parameter range from the flow data associated with GatingHierarchy

#### Usage

```
## S3 method for class 'GatingHierarchy'
range(..., na.rm = FALSE, type = c("instrument", "data"), raw.scale = FALSE)
```

## Arguments

	GatingHierarchy object
na.rm	not used
type	character of "instrument" or "data" indicating whether to retrieve the instrument or the actual data range
raw.scale	logical whether convert the range from transformed scale to raw scale

#### Value

matrix

#### Examples

```
## Not run:
    range(gh, type = "data")#return data range
    range(gh) #return instrument range
    range(gh, raw.scale = TRUE) #inverse transform the range to the raw scale
## End(Not run)
```

read.gatingML.cytobank

#### Description

The Default parser (read.gatingML) does not parse the population tree as well as the custom information from cytobank. (e.g. gate name, fcs filename).

#### Usage

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

#### Arguments

file	Gating-ML XML file
	additional arguments passed to the handlers of 'xmlTreeParse'

# Value

a graphGML that represents the population tree. The gate and population name are stored in node-Data of each node. Compensation and transformations are stored in graphData.

#### Examples

```
## Not run:
g <- read.gatingML.cytobank(xml) #parse the population tree
#plot(g) #visualize it
```

## End(Not run)

show,graphGML-method show method for graphGML

#### Description

show method for graphGML

#### Usage

```
## S4 method for signature 'graphGML'
show(object)
```

#### Arguments

object graphGML

#### Value

nothing

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