

Package ‘ggcyto’

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Type Package

Title Visualize Cytometry data with ggplot

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Description With the dedicated `fortify` method implemented for `flowSet`, `ncdfFlowSet` and `GatingSet` classes, both raw and gated flow cytometry data can be plotted directly with `ggplot`. `ggcyto` wrapper and some customized layers also make it easy to add gates and population statistics to the plot.

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'compute_stats.R' 'fortify.R' 'fortify_fs.R' 'geom_gate.R'
'geom_hvline.R' 'geom_overlay.R' 'geom_stats.R'
'getFlowFrame.R' 'ggcyto.R' 'ggcyto_GatingLayout.R'
'ggcyto_GatingSet.R' 'ggcyto_flowSet.R' 'labs.R' 'ggcyto_par.R'
'ggplot_data_frame.R' 'merge.quad.gates.R' 'replace_data.R'
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as.ggplot	<i>It fortifies the data, fills some default settings and returns a regular ggplot object.</i>
-----------	--

Description

The orginal data format is preserved during the ggcyo constructor because they still need to be used during the plot building process. This function is usually called automatically in the print/plot method of ggcyto. Sometime it is useful to coerce it to ggplot explicitly by user so that it can be used as a regular ggplot object.

Usage

```
as.ggplot(x, pre_binning = FALSE)
```

Arguments

<code>x</code>	ggcyto object with the data that has not yet been fortified to data.frame.
<code>pre_binning</code>	whether to pass the binned data to ggplot to avoid the overhead to scaling the original raw data for geom_hex layer

Value

ggplot object

Examples

```
data(GvHD)
fs <- GvHD[1:3]
#construct the `ggcyto` object (inherits from `ggplot` class)
p <- ggcyo(fs, aes(x = `FSC-H`)) + geom_histogram()
class(p) # a ggcyo object
p$data # data has not been fortified
p1 <- as.ggplot(p) # convert it to a ggplot object explicitly
class(p1)
p1$data # data is fortified
```

autoplot.flowSet	<i>Plot cytometry data in one or two dimension with the ggcyo package.</i>
------------------	--

Description

Overloaded autoplot methods for the cytometry data structure: flowFrame or flowSet, Gatinghierarchy, GatingSet. It plots the cytometry data with geom_histogram, geom_density or geom_hex. When autoplot is called on a GatingSet/Gatinghierarchy, the second argument should be a gate or population node. And the dimensions(channels/markers) are deduced from the gate dimensions.

Usage

```

## S3 method for class 'flowSet'
autoplots(object, x, y = NULL, bins = 30, ...)

## S3 method for class 'ncdfFlowList'
autoplots(object, ...)

## S3 method for class 'flowFrame'
autoplots(object, x, ...)

## S3 method for class 'GatingSetList'
autoplots(object, ...)

## S3 method for class 'GatingSet'
autoplots(
  object,
  gate,
  x = NULL,
  y = "SSC-A",
  bins = 30,
  axis_inverse_trans = TRUE,
  ...
)

## S3 method for class 'GatingHierarchy'
autoplots(
  object,
  gate,
  y = "SSC-A",
  bool = FALSE,
  arrange.main = sampleNames(object),
  arrange = TRUE,
  merge = TRUE,
  projections = list(),
  strip.text = c("parent", "gate"),
  path = "auto",
  ...
)

```

Arguments

object	The data source. A core cytometry data structure. A flowFrame, flowSet, GatingSet or GatingHierarchy object
x	define the x dimension of the plot (not used when object is a GatingSet). When object is a flowFrame, it can be missing, which plots 1d density plot on all the channels.
y	define the y dimension of the plot. Default is NULL, which means 1d density-plot.
bins	passed to geom_hex
...	other arguments passed to ggplot
gate	the gate to be plotted

axis_x_inverse_trans	logical flag indicating whether to add <code>axis_x_inverse_trans</code> and <code>axis_x_inverse_trans</code> layers.
bool	whether to plot boolean gates
arrange.main	the main title of the arranged plots
arrange	whether to use arrangeGrob to put multiple plots in the same page
merge	whether to merge multiple gates into the same panel when they share the same parent and projections
projections	a list of customized projections
strip.text	either "parent" (the parent population name) or "gate" (the gate name). The latter usually is used when merge is FALSE
path	the gating path format (passed to <code>gs_get_pop_paths</code>)

Value

a ggcyto object

Examples

```
library(flowCore)
data(GvHD)
fs <- GvHD[subset(pData(GvHD), Patient %in% 5:7 & Visit %in% c(5:6))[, "name"]]

#1d- density plot
autoplot(fs, x = "SSC-H")

#1d- density plot on all channels
autoplot(fs[[1]])

#2d plot: default geom_hex plot
autoplot(fs, x = 'FSC-H', y ='SSC-H')

#autplot for GatingSet
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
autoplot(gs, "CD3+")
#display axis values in transformed scale
autoplot(gs, "CD3+", axis_inverse_trans = FALSE)

#autplot for GatingHierarchy
gh <- gs[[1]]
autoplot(gh) # by default the strip.text shows the parent population

#To display the gate name
#autoplot(gh , strip.text = "gate")
```

`axis_x_inverse_trans` *Display ggcyto axis labels using their raw values (as stored in the data structure)*

Description

It is essentially a dummy continuous scale and will be instantiated by '+.ggcyto_GatingSet' with 'breaks' and 'labels' customized.

Usage

```
axis_x_inverse_trans(...)

axis_y_inverse_trans(...)
```

Arguments

...	common continuous scale parameters passed to 'continuous_scale' (not used currently)
-----	--

Value

a raw_scale object that inherits scale class.

Examples

```
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
p <- ggcryo(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)
p <- p + geom_gate("CD4") + geom_stats() #plot CD4 gate and it is stats
p
p + axis_x_inverse_trans() #inverse transform the x axis into raw scale
```

compute_stats

compute the statistics of the cell population defined by gates

Description

It calls the underlining stats routine and merge it with the label position calculated by stat_position as well as the pData of flowSet.

Usage

```
compute_stats(fs = NULL, gates, type = "percent", value = NULL, ...)
```

Arguments

fs	flowSet. can be NULL when precalculated 'value' is provided
gates	a list of filters
type	a vector of strings to specify the stats types. can be any or multiple values of "percent", "count", "gate_name", or "MFI" (MFI is currently not supported yet).
value	the pre-calculated stats value. when supplied, the stats computing is skipped.
...	other arguments passed to stat_position function

Details

This function is usually not called directly by user but used by ggcryo when geom_stat layer is added.

Value

a data.table that contains percent and centroid locations as well as pData that used as data for geom_btext layer.

Examples

```
data(GvHD)
fs <- GvHD[1:4]
rect.g <- rectangleGate(list("FSC-H" = c(300,500), "SSC-H" = c(50,200)), filterId = "P1")
rect.gates <- sapply(sampleNames(fs), function(sn)rect.g)
compute_stats(fs, rect.gates)
compute_stats(fs, rect.gates, type = c("gate_name", "percent"))
```

`flowCore_asinht_trans` *Inverse hyperbolic sine transformation(flowCore version).*

Description

Used to construct inverse hyperbolic sine transform object.

Usage

```
flowCore_asinht_trans(..., n = 6, equal.space = FALSE)
```

Arguments

...	parameters passed to arcsinhTransform
n	desired number of breaks (the actual number will be different depending on the data range)
equal.space	whether breaks at equal-spaced intervals

Value

asinht transformation object

Examples

```
trans.obj <- flowCore_asinht_trans(equal.space = TRUE)
data <- 1:1e3
brks.func <- trans.obj[["breaks"]]
brks <- brks.func(data)
brks # fasinh space displayed at raw data scale

#transform it to verify it is equal-spaced at transformed scale
trans.func <- trans.obj[["transform"]]
brks.trans <- trans.func(brks)
brks.trans
```

fortify.ellipsoidGate *Convert a ellipsoidGate to a data.table useful for ggplot*

Description

It interpolates the ellipsoidGate to polygonate before fortifying it.

Usage

```
## S3 method for class 'ellipsoidGate'
fortify(model, data = NULL, ...)
```

Arguments

model	ellipsoidGate
data	data range used for polygon interpolation.
...	not used.

Value

`data.table`

Examples

```
## Defining the gate
cov <- matrix(c(6879, 3612, 3612, 5215), ncol=2,
               dimnames=list(c("FSC-H", "SSC-H"), c("FSC-H", "SSC-H")))
mean <- c("FSC-H"=430, "SSC-H"=175)
eg <- ellipsoidGate(filterId= "myEllipsoidGate", .gate=cov, mean=mean)
fortify(eg)
```

fortify.filterList *Convert a filterList to a data.table useful for ggplot*

Description

It tries to merge with pData that is associated with filterList as attribute 'pd'

Usage

```
## S3 method for class 'filterList'
fortify(model, data = NULL, nPoints = NULL, ...)
```

Arguments

model	filterList
data	not used
nPoints	not used
...	not used.

Value

```
data.table
```

Examples

```
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
gates <- gs_pop_get_gate(gs, "CD4")
gates <- as(gates, "filterList") #must convert list to filterList in order for the method to dispatch properly
fortify(gates)
```

fortify.flowFrame

Convert a flowFrame/flowSet/GatingSet to a ggplot-compatible data.table

Description

It extracts events matrices and appends the pData to it so that ggplot can use the pData for facetting.

Usage

```
## S3 method for class 'flowFrame'
fortify(model, data, ...)

## S3 method for class 'flowSet'
fortify(model, data, ...)

## S3 method for class 'ncdfFlowList'
fortify(model, ...)

## S3 method for class 'GatingSetList'
fortify(model, ...)

## S3 method for class 'GatingSet'
fortify(model, ...)
```

Arguments

model	flowFrame, flowSet or GatingSet
data	not used.
...	not used.

Value

```
data.table
data.table
data.table
```

Examples

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

attr(gs, "subset") <- "CD4" #must attach subset information to GatingSet object before fortifying it
fortify(gs)

fs <- gs_pop_get_data(gs, "CD8")
fortify(fs)#fs is a flowSet/ncdfFlowSet

fr <- fs[[1]]
fortify(fr)#fr is a flowFrame
```

fortify.polygonGate *Convert a polygonGate to a data.table useful for ggplot*

Description

It converts the boundaries slot into a data.table

Usage

```
## S3 method for class 'polygonGate'
fortify(model, data = NULL, nPoints = NULL, ...)
```

Arguments

model	polygonGate
data	data range used to reset off-bound gate coordinates to prevent interpolating on the extremely large space unnecessarily.
nPoints	not used
...	not used.

Value

data.table

Examples

```
srcut <- matrix(c(300,300,600,600,50,300,300,50),ncol=2,nrow=4)
colnames(srcut) <- c("FSC-H","SSC-H")
pg <- polygonGate(filterId="nonDebris", .gate= srcut)
fortify(pg)
```

`fortify.rectangleGate` *Convert a rectangleGate to a data.table useful for ggplot*

Description

For 2d rectangelGate, it is converted to a polygonGate first and then dispatch to the fortify method for polygonGate. for 1d, uses geom_vline/hline format.

Usage

```
## S3 method for class 'rectangleGate'
fortify(model, data = NULL, ...)
```

Arguments

model	rectangleGate
data	data range used for polygon interpolation.
...	not used.

Value

`data.table`

Examples

```
#2d rectangleGate
rect.g <- rectangleGate(list("FSC-H" = c(300,500), "SSC-H" = c(50,200)))
fortify(rect.g)
#1d gate
rg <- rectangleGate(list("FSC-H" = c(300,500)))
fortify(rg)
```

`fortify_fs` *Fortify a model into flowSet object*

Description

The method provides a universe interface to convert a generic R object into a flowSet useful for ggcryo

Usage

```
fortify_fs(model, data, ...)
## S3 method for class 'flowSet'
fortify_fs(model, data, ...)
## Default S3 method:
fortify_fs(model, data, ...)
```

```
## S3 method for class 'flowFrame'
fortify_fs(model, data, ...)

## S3 method for class 'GatingSetList'
fortify_fs(model, data, ...)

## S3 method for class 'GatingSet'
fortify_fs(model, data, ...)
```

Arguments

model	flow object(flowFrame or GatingSet) to be converted to flowSet. when it is a GatingSet, it must contain the subset information stored as 'subset' attribute.
data	original dataset, if needed
...	other arguments passed to methods

Value

a flowSet/ncdfFlowSet object

Examples

```
data(GvHD)
fr <- GvHD[[1]]
fortify_fs(fr)

dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
attr(gs, "subset") <- "CD4"
fortify_fs(gs)
```

geom_gate

Add a gate layer to a ggcyto plot.

Description

When 'data' is a gate (or flowCore filter) or a list of gates or a filterList object. When it is used directly with 'ggplot', pdata of the flow data must be supplied through 'pd' argument explicitly in order for the gates to be dispatched to each panel. However It is not necessary when used with 'ggcyto' wrapper since the latter will attach pData automatically.

Usage

```
geom_gate(data, ...)

## S3 method for class 'filterList'
geom_gate(data, pd, nPoints = 100, ...)

## S3 method for class 'filter'
geom_gate(data, mapping = NULL, fill = NA, colour = "red", nPoints = 100, ...)
```

Arguments

data	a filter (Currently only rectangleGate (1d or 2d), polygonGate, ellipsoidGate are supported.) or a list of these gates or filterList or character specifying a gated cell population in the GatingSet
...	other arguments
pd	pData (data.frame) that has rownames represents the sample names used as key to be merged with filterList
nPoints	used for interpolating polygonGates to prevent them from losing shape when truncated by axis limits
mapping	The aesthetic mapping
fill	fill color for the gate. Not filled by default.
colour	default is red

Details

When 'data' is a character, it construct an abstract geom layer for a character that represents nodes in a Gating tree and will be instantiated later as a specific geom_gate layer or layers based on the gates extracted from the given GatingSet object.

Value

a geom_gate layer

Examples

```
data(GvHD)
fs <- GvHD[subset(pData(GvHD), Patient %in% 5:7 & Visit %in% c(5:6))[, "name"]]
p <- ggcryo(fs, aes(x = `FSC-H`, y = `SSC-H`))
p <- p + geom_hex(bins = 128)
rect.g <- rectangleGate(list("FSC-H" = c(300, 500), "SSC-H" = c(50, 200)))
#constructor for a list of filters
rect.gates <- sapply(sampleNames(fs), function(sn)rect.g)
p + geom_gate(rect.gates)

dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
p <- ggcryo(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)
# add gate layer by gate name
p + geom_gate("CD4")
```

geom_hvline

Vertical or horizontal line.

Description

This geom is based on the source code of '[geom_hline](#)' and '[geom_vline](#)'.

Usage

```
geom_hvline(
  mapping = NULL,
  data = NULL,
  position = "identity",
  show.legend = FALSE,
  ...
)
```

Arguments

mapping	The aesthetic mapping, usually constructed with <code>aes</code> or <code>aes_string</code> . Only needs to be set at the layer level if you are overriding the plot defaults.
data	A layer specific dataset - only needed if you want to override the plot defaults.
position	The position adjustment to use for overlapping points on this layer
show.legend	should a legend be drawn? (defaults to FALSE)
...	other arguments passed on to <code>layer</code> . This can include aesthetics whose values you want to set, not map. See <code>layer</code> for more details.

Details

The goal is to determine the line to be either vertical or horizontal based on the 1-d data provided in this layer.

Value

a `geom_hvline` layer

Aesthetics

@section Aesthetics: `geom_vline()` understands the following aesthetics (required aesthetics are in bold):

- `xintercept`
- `alpha`
- `colour`
- `group`
- `linetype`
- `size`

Learn more about setting these aesthetics in `vignette("ggplot2-specs")`.

Examples

```
p <- ggplot(mtcars, aes(x = wt, y = mpg)) + geom_point()
# vline
p + geom_hvline(data = data.frame(wt= 3))
# hline
p + geom_hvline(data = data.frame(mpg= 20))
```

geom_overlay	<i>Overlay a population on an existing ggcyto plot analogous to backgating.</i>
--------------	---

Description

It is useful for "backgating" plots.

Usage

```
geom_overlay(data, ...)
```

Arguments

- data** a filter (Currently only rectangleGate (1d or 2d), polygonGate, ellipsoidGate are supported.) or a list of these gates or filterList or character specifying a gated cell population in the GatingSet
- ...** other arguments mapping, The mapping aesthetic mapping data a polygonGate fill polygonGate is not filled by default colour default is red pd pData (data.frame) that has rownames represents the sample names used as key to be merged with filterList

Value

a geom_overlay layer

Examples

```
library(ggcyto)
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
p <- autoplot(gs, "CD3+")

# add a flowSet as the overlay
fs <- gs_pop_get_data(gs, "DPT")
p + geom_overlay(data = fs, size = 0.3, alpha = 0.7)

# add overlay layer by gate name
p + geom_overlay(data = "DNT", size = 0.3, alpha = 0.7)

#add overlay for 1d densityplot
p <- ggcyto(gs, aes(x = CD4), subset = "CD3+") + geom_density(aes(y = ..count..))
p + geom_overlay("DNT", aes(y = ..count..), fill = "red")
```

geom_stats*Add a population statistics layer to a ggcyto plot.*

Description

This is a virtual layer and will be instantiated as geom_label layer within ggcyto.+ operator.

Usage

```
geom_stats(
  gate = NULL,
  ...,
  value = NULL,
  type = "percent",
  negated = FALSE,
  adjust = 0.5,
  label.padding = unit(0.05, "lines"),
  label.size = 0,
  digits = 3
)
```

Arguments

gate	a 'filterList' or character (represent as a population node in GatingSet) if not supplied, ggcyto then tries to parse the gate from the first geom_gate layer.
...	other arguments passed to geom_label layer
value	the pre-calculated stats value. when supplied, the stats computing is skipped.
type	a vector of strings to specify the stats types. can be any or multiple values of "percent", "count", "gate_name", or "MFI" (MFI is currently not supported yet).
negated	whether the gate needs to be negated
adjust	adjust the position of the centroid. from 0 to 1.
label.padding, label.size	arguments passed to geom_label layer
digits	control the stats format

Details

So it is dedicated for ggcyto context and thus cannot be added to ggplot object directly.

Value

a geom_popStats layer

Examples

```
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
p <- ggcyto(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)
p
# add gate and stats layer
```

```

p + geom_gate("CD4") + geom_stats()

# display gate name
p + geom_gate(c("CD4", "CD8")) + geom_stats(type = "gate_name")
# display gate name and percent
p + geom_gate(c("CD4", "CD8")) + geom_stats(type = c("gate_name", "percent"))

```

getFlowFrame*extract flowFrame data structure from the given R object***Description**

Mainly to get the channel and marker information.

Usage

```
getFlowFrame(x)
```

Arguments

x	flowSet, ncdfFlowList, GatingSet, GatingHierarchy, or GatingSetList
---	---

Value

a flowFrame. When x is a ncdfFlowSet or GatingSet that is associated with ncdfFlowSet, the raw event data is not read and an empty flowFrame is returned.

Examples

```

data(GvHD)
fs <- GvHD[1:2]
getFlowFrame(fs)# fs is a flowSet

dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
getFlowFrame(gs)# gs is a GatingSet

```

gcyto-class*Plot cytometry data using the gcyto API***Description**

`gcyto()` initializes a gcyto object that inherits ggplot class. Similarly the `+` operator can be used to add layers to the existing gcyto object.

Usage

```
ggcyto(data = NULL, ...)

## S3 method for class 'GatingSet'
ggcyto(data, mapping, subset = "_parent_", ...)

## S3 method for class 'GatingSetList'
ggcyto(data, ...)

## S3 method for class 'GatingHierarchy'
ggcyto(data, ...)

## S3 method for class 'flowSet'
ggcyto(data, mapping, filter = NULL, max_nrow_to_plot = 50000, ...)
```

Arguments

data	The data source. A core cytometry data structure. (flowSet, flowFrame, ncdfFlowSet, GatingSet or GatingHierarchy)
...	other arguments passed to specific methods
mapping	default list of aesthetic mappings (these can be colour, size, shape, line type – see individual geom functions for more details)
subset	character that specifies the node path or node name in the case of GatingSet. Default is "_parent_", which will be substituted with the actual node name based on the geom_gate layer to be added later.
filter	a flowcore gate object or a function that takes a flowSet and channels as input and returns a data-dependent flowcore gate. The gate is used to filter the flow data before it is plotted.
max_nrow_to_plot	the maximum number of cells to be plotted. When the actual data exceeds it, The subsampling process will be triggered to speed up plotting. Default is 5e4. To turn off the subsampling, simply set it to a large enough number or Inf.

Details

To invoke ggcyto:

- `ggcyto(fs, aes(x,y,<other aesthetics>))`

Value

ggcyto object

Examples

```
data(GvHD)
fs <- GvHD[1:3]
#construct the `ggcyto` object (inherits from `ggplot` class)
p <- ggcyto(fs, aes(x = 'FSC-H'))
p + geom_histogram()
```

```

# display density/area
p + geom_density()
p + geom_area(stat = "density")

# 2d scatter plot
p <- ggcyto(fs, aes(x = `FSC-H`, y = `SSC-H`))
p + geom_hex(bins = 128)
# do it programatically through aes_string and variables
col1 <- "`FSC-H`" #note that the dimension names with special characters needs to be quoted by backticks
col2 <- "`SSC-H`"
ggcyto(fs, aes_string(col1,col2)) + geom_hex()

## More flowSet examples
fs <- GvHD[subset(pData(GvHD), Patient %in% 5:7 & Visit %in% c(5:6))[,["name"]]]
# 1d histogram/densityplot
p <- ggcyto(fs, aes(x = `FSC-H`))
#facet_wrap(~name)` is used automatically
p1 <- p + geom_histogram()
p1
#overwriting the default faceeting
p1 + facet_grid(Patient~Visit)

#display density
p + geom_density()

#you can use ggridges package to display stacked density plot
require(ggridges)
#stack by fcs file ('name')
p + geom_density_ridges(aes(y = name)) + facet_null() #facet_null is used to remove the default facet_wrap (by 'name')
#or to stack by Visit and facet by patient
p + geom_density_ridges(aes(y = Visit)) + facet_grid(~Patient)

# 2d scatter/dot plot
p <- ggcyto(fs, aes(x = `FSC-H`, y = `SSC-H`))
p <- p + geom_hex(bins = 128)
p

## GatingSet
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
# 2d plot
ggcyto(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)

# 1d plot
ggcyto(gs, aes(x = CD4), subset = "CD3+") + geom_density()

```

ggcyto_add

overloaded '+' method for ggcyto

Description

It tries to copy pData from ggcyto object to the gate layers so that the gate layer does not need to have 'pd' to be supplied explicitly by users. It also calculates population statistics when geom_stats layer is added. It supports addition ggcyto layers such as 'ggcyto_par' and 'labs_cyto'.

Usage

```
e1 + e2
```

Arguments

- e1 An object of class ggcyto or a class inheriting from ggcyto, such as ggcyto_flowSet, ggcyto_GatingSet, or ggcyto_GatingLayout. In the case of ggcyto_GatingLayout, the component of e2 will be added to each subsidiary plot.
- e2 A component to add to e1

Value

ggcyto object

Examples

```
## flowSet
data(GvHD)
fs <- GvHD[subset(pData(GvHD), Patient %in% 5:7 & Visit %in% c(5:6))[, "name"]]
p <- ggcyto(fs, aes(x = `FSC-H`, y = `SSC-H`)) + geom_hex(bins = 128)
#add rectangleGate layer (2d)
rect.g <- rectangleGate(list("FSC-H" = c(300,500), "SSC-H" = c(50,200)))
rect.gates <- sapply(sampleNames(fs), function(sn)rect.g)
p + geom_gate(rect.gates) + geom_stats()

## GatingSet
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
p <- ggcyto(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)
p <- p + geom_gate("CD4") + geom_stats() #plot CD4 gate and it is stats
p
p + axis_x_inverse_trans() #inverse transform the x axis into raw scale

## GatingLayout
#autplot for GatingSet
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
gh <- gs[[1]]
p <- autplot(gh)
class(p)
# customize the font size of strip text for each ggcyo plots contained in GatingLayout object
p + theme(strip.text = element_text(size = 14))
```

ggcyto_arrange

Arrange a list of ggplot objects into gtable

Description

It is usually implicitly invoked by print and show method and can be called by user when the further manipulation is needed,

Usage

```
ggcyto_arrange(x, ...)
```

Arguments

- x ggcyto_gate_layout, which is essentially a list of ggplot objects that were previously stored as ggcyto_gate_layout object by autoplot function.
- ... other arguments passed to arrangeGrob

Value

gtable

Examples

```
## Not run:
# get ggcyto_GatingLayout object from first sample
res <- autoplot(gs[[1]], nodes, bins = 64)
class(res)
# arrange it as one-row gtable object
gt <- ggcyto_arrange(res, nrow = 1)
gt
# do the same to the second sample
gt2 <- ggcyto_arrange(autoplot(gs[[2]]), nodes, bins = 64), nrow = 1)
# combine the two and print it on the sampe page
gt3 <- gridExtra::gridExtra_rbind(gt, gt2)
plot(gt3)

## End(Not run)
```

`ggcyto_par_default` *Return The default ggcyto settings*

Description

Return The default ggcyto settings

Usage

```
ggcyto_par_default()
```

Value

a list of default settings for ggcyto

Examples

```
ggcyto_par_default()
```

`ggcyto_par_set` *Set some default parameters for ggcryo*

Description

Use this function to modify ggcryo parameters These are the regular (or to be instantiated as) scales, labs, facet objects. They can be added as a single layer to the plot for the convenience.

Usage

```
ggcyto_par_set(...)
```

Arguments

...	a list of element name, element pairings that modify the existing parameter settings
-----	--

Value

a list of new settings for ggcryo

elements

The individual elements are:

limits	can be "data"(default) or "instrument" or a list of numeric limits for x and y (e.g. <code>list(x = c(0, 4000))</code>)
facet	the regular facet object
hex_fill	default scale_fill_gradientn for geom_hex layer
lab	labs_cryo object

Examples

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

p <- ggcryo(gs, aes(x = CD4, y = CD8), subset = "CD3+")
# 2d plot
p <- p + geom_hex(bins = 64)
p

#use instrument range by overwritting the default limits settings
p + ggcryo_par_set(limits = "instrument")

#manually set limits
myPars <- ggcryo_par_set(limits = list(x = c(0,3.2e3), y = c(-10, 3.5e3)))
p + myPars# or xlim(0,3.2e3) + ylim(-10, 3.5e3)
```

`is.ggcryo` *Reports whether x is a ggcryo object*

Description

Reports whether x is a ggcryo object

Usage

```
is.ggcryo(x)
```

Arguments

x An object to test

Value

TRUE/FALSE

Examples

```
data(GvHD)
fs <- GvHD[1:2]
p <- ggcryo(fs, aes(x = `FSC-H`))
is.ggcryo(p)
```

`is.ggcryo_flowSet` Reports whether x is a ggcryo_flowSet object

Description

Reports whether x is a ggcryo_flowSet object

Usage

```
is.ggcryo_flowSet(x)
```

Arguments

x An object to test

Value

TRUE or FALSE

Examples

```
data(GvHD)
fs <- GvHD[1:2]
p <- ggcryo(fs, aes(x = `FSC-H`))
is.ggcryo_flowSet(p)
```

is.ggcryo_par *Reports whether x is a ggcryo_par object*

Description

Reports whether x is a ggcryo_par object

Usage

```
is.ggcryo_par(x)
```

Arguments

x An object to test

Value

TRUE or FALSE

Examples

```
myPar <- ggcryo_par_set(limits = "instrument")
is.ggcryo_par(myPar)
```

labs_cyto *Change axis labels and legend titles*

Description

The actual labels text will be instantiated when it is added to ggcryo plot.

Usage

```
labs_cyto(labels = "both")
```

Arguments

labels default labels for x, y axis. Can be "channel", "marker", or "both" (default)

Value

a list

Examples

```

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

# default is "both"
p <- ggcryo(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)
p

#use marker name as x,y labs
p + labs_cryo("marker")

#use channel name as x,y labs
p + labs_cryo("channel")

```

marginalFilter *Generate a marginal gate.*

Description

It simply constructs an boundaryFilter that removes the marginal events. It can be passed directly to ggcryo constructor. See the examples for details.

Usage

```
marginalFilter(fs, dims, ...)
```

Arguments

<code>fs</code>	flowSet (not used.)
<code>dims</code>	the channels involved
<code>...</code>	arguments passed to boundaryFilter

Value

an boundaryFilter

Examples

```

data(GvHD)
fs <- GvHD[1]
chnls <- c("FSC-H", "SSC-H")
#before removign marginal events
summary(fs[, chnls])

# create merginal filter
g <- marginalFilter(fs, chnls)
g

#after remove marginal events
fs.clean <- Subset(fs, g)
summary(fs.clean[, chnls])

```

```
#pass the function directly to ggcryo
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_manual", full = TRUE))
# with marginal events
ggcyto(gs, aes(x = CD4, y = CD8), subset = "CD3+") + geom_hex(bins = 64)

# using marginalFilter to remove these events
ggcyto(gs, aes(x = CD4, y = CD8), subset = "CD3+", filter = marginalFilter) + geom_hex(bins = 64)
```

merge.quad.gates

extend the original flowWorkspace:::mergeGates function to restore quadGate when applicable

Description

For internal usage.

Usage

```
## S3 method for class 'quad.gates'
merge(gh, pops, bool = TRUE)
```

Arguments

gh	a GatingHierarchy
pops	a vector of population names
bool	whether to deal with boolean gate

Value

a nested list of data structure that captures the information of parent, grouped populations (with the same projections) and the reconstructed quadGate object and the respective quadrant pattern

Examples

```
library(flowWorkspace)
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(file.path(dataDir, "gs_manual"))
#get the GatingHierarchy object
gh <- gs[[1]]
pops <- gs_pop_get_children(gh, "CD4")
grps <- ggcryo:::merge.quad.gates(gh, pops)
length(grps) # pops are grouped into two
grps[[1]] # each group is annotated with quadGate information

ggcyto:::merge.quad.gates(gh, gs_pop_get_children(gh, "CD3+")) # cd3 subsets are not coercible to quadgate thus
```

print.ggcryo	<i>Draw ggcryo on current graphics device.</i>
--------------	--

Description

A wrapper for print.ggplot. It converts the ggcryo to conventional ggplot object before printing it. This is usually invoked automatically when a ggcryo object is returned to R console.

Usage

```
## S3 method for class 'ggcyto'
print(x, ...)

## S3 method for class 'ggcyto'
plot(x, ...)

## S3 method for class 'ggcyto'
show(object)
```

Arguments

x	ggcyto object to display
...	other arguments not used by this method
object	ggcyto object

Value

nothing

print.ggcryo_GatingLayout	<i>print method for ggcryo_gate_layout class</i>
---------------------------	--

Description

print method for ggcryo_gate_layout class

Usage

```
## S3 method for class 'ggcyto_GatingLayout'
print(x, ...)

## S3 method for class 'ggcyto_GatingLayout'
show(object)
```

Arguments

- x ggcryo_gate_layout, which is essentially a list of ggplot objects that were previously stored as ggcryo_gate_layout object by autoplot function.
- ... other arguments passed to arrangeGrob
- object ggcryo_GatingLayout

Value

nothing

<i>replace_data</i>	<i>replace current cytometry data</i>
---------------------	---------------------------------------

Description

It essentially reconstructs the entire ggcryo plot object based on the new data and the original mapping and layers recorded in the plot object.

Usage

```
e1 %+% e2
```

Arguments

- e1 the ggcryo object
- e2 the new cytometry data . It can be 'GatingSet' or 'flowSet'.

Value

the new ggcryo object

Examples

```
dataDir <- system.file("extdata", package="flowWorkspaceData")
gs <- load_gs(list.files(dataDir, pattern = "gs_bcell_auto", full = TRUE))
gs1 <- gs[1]
gs2 <- gs[2]

#construct the ggcryo object for gs1
p <- ggcryo(gs1, aes(cd24, cd38)) + geom_hex(bins = 128)
p <- p + geom_gate("Transitional") #add gate
#customize the stats layer
p <- p + geom_stats(type = "count", size = 6, color = "white", fill = "black", adjust = 0.3)
#customize the layer
p <- p + labs_cryo("channel")
#customize the axis limits
p <- p + ggcryo_par_set(limits = "instrument")
#add another population as the overlay dots
p <- p + geom_overlay("IgD-CD27-", col = "black", size = 1.2, alpha = 0.4)
#hide the legend
p <- p + guides(fill=FALSE)
p
```

```
#replace the data with gs2 and see the same visual effect
p %+% gs2
```

scales_flowjo.biexp *Add a flowJo biexponential scale to the x or y axes of a ggcryo plot.*

Description

Add a flowJo biexponential scale to the x or y axes of a ggcryo plot.

Usage

```
scale_x_flowjo_biexp(
  ...,
  maxValue = 262144,
  widthBasis = -10,
  pos = 4.5,
  neg = 0,
  equal.space = FALSE
)

scale_y_flowjo_biexp(
  ...,
  maxValue = 262144,
  widthBasis = -10,
  pos = 4.5,
  neg = 0,
  equal.space = FALSE
)
```

Arguments

...	common continuous scale parameters passed to 'continuous_scale' (not used currently)
maxValue, widthBasis, pos, neg	see 'help(flowjo.biexp')
equal.space	whether to display the breaks in equal.space format

Value

ScaleContinuous object

Examples

```
data(GvHD)
fr <- GvHD[[1]]
p <- ggcryo(fr, aes(x = `FL1-H`)) + geom_density()
#display at raw scale
p
#display at transformed scale
p + scale_x_flowjo_biexp(maxValue = 1e4, widthBasis = 0)
```

`scales_flowjo_fasinh` *Add a flowJo inverse hyperbolic sine scale to the x or y axes of a ggcryo plot.*

Description

Add a flowJo inverse hyperbolic sine scale to the x or y axes of a ggcryo plot.

Usage

```
scale_x_flowjo_fasinh(..., m = 4, t = 1200)
scale_y_flowjo_fasinh(..., m = 4, t = 1200)
```

Arguments

...	common continuous scale parameters passed to 'continuous_scale' (not used currently)
m, t	see 'help(flowjo_fasinh)'

Value

ScaleContinuous object

Examples

```
data(GvHD)
fr <- GvHD[[1]]
p <- ggcryo(fr, aes(x = 'FL1-H')) + geom_density()
#display at raw scale
p
#display at transformed scale
p + scale_x_flowjo_fasinh(t = 1e4)
```

`scale_x_flowCore_fasinh`

Add a flowCore inverse hyperbolic sine scale to the x or y axes of a ggcryo plot.

Description

Add a flowCore inverse hyperbolic sine scale to the x or y axes of a ggcryo plot.

Usage

```
scale_x_flowCore_fasinh(..., a = 1, b = 1, c = 0)
scale_y_flowCore_fasinh(..., a = 1, b = 1, c = 0)
```

Arguments

- ... common continuous scale parameters passed to 'continuous_scale' (not used currently)
- a, b, c see 'help(arcsinhTransform)'

Value

ScaleContinuous object

Examples

```
data(GvHD)
fr <- GvHD[[1]]
p <- ggcyto(fr, aes(x = `FL1-H`)) + geom_density()
#display at raw scale
p
#display at transformed scale
p + scale_x_flowCore_fasinh(a = 2)
```

scale_x_logicle *Add a logicle scale to the x or y axes of a ggcyto plot.*

Description

Add a logicle scale to the x or y axes of a ggcyto plot.

Usage

```
scale_x_logicle(..., w = 0.5, t = 262144, m = 4.5, a = 0)
scale_y_logicle(..., w = 0.5, t = 262144, m = 4.5, a = 0)
```

Arguments

- ... common continuous scale parameters passed to 'continuous_scale' (not used currently)
- w, t, m, a see 'help(logicleTransform)'

Value

ScaleContinuous object

Examples

```
data(GvHD)
fr <- GvHD[[1]]
p <- ggcyto(fr, aes(x = `FL1-H`)) + geom_density()
#display at raw scale
p
#display at transformed scale
p + scale_x_logicle(t = 1e4)
```

stat_position	<i>Compute the positions of the population statistics based on the geometric gate centroid for a ggcyto plot.</i>
----------------------	---

Description

It is usually not called directly by user but mainly used by `compute_stats` function (which is called by `ggcyto` add method when `geom_states` layer is added).

Usage

```
stat_position(gate, ...)

## S3 method for class 'filter'
stat_position(
  gate,
  negated = FALSE,
  adjust = 0.5,
  abs = FALSE,
  data_range = NULL,
  limits = NULL,
  ...
)
```

Arguments

<code>gate</code>	a flowCore filter
<code>...</code>	other arguments
<code>negated</code>	logical indicating whether position needs to be moved to negative side of gate
<code>adjust</code>	adjust the position of the centroid. This can be a length-2 vector with an adjustment in each dimension.
<code>abs</code>	logical
<code>data_range</code>	a two-row data.frame representing the actual data range. Each column is a range for a specific channel. First row is min, Second row is max.
<code>limits</code>	used to fix the gate range

Value

a data.table of gate centroid coordinates

Examples

```
data(GvHD)
fs <- GvHD[1:4]
rect.g <- rectangleGate(list("FSC-H" = c(300,500), "SSC-H" = c(50,200)))
rect.gates <- sapply(sampleNames(fs), function(sn)rect.g)
stat_position(rect.gates)
```

transform-gate	<i>rescale methods for gates</i>
----------------	----------------------------------

Description

rescale the gate coordinates with the transformation provided

Usage

```
transform(`_data`, ...)  
rescale_gate(gate, trans, param)
```

Arguments

_data	the filter or filterList object. Currently support polygonGate, ellipsoidGate, rectangleGate and quadGate.
...	trans the transformation function or transformList object param the parameter/dimension to be transformed. When trans is transformList object, param is not needed since it is derived from transformList.
gate	gate object
trans	the transformation function
param	the parameter/dimension to be transformed.

Value

the transformed filter/filterList object

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