# Package 'CytoMDS'

# December 5, 2024

**Title** Low Dimensions projection of cytometry samples

Version 1.2.0

Description This package implements a low dimensional visualization of a set of cytometry samples, in order to visually assess the 'distances' between them. This, in turn, can greatly help the user to identify quality issues like batch effects or outlier samples, and/or check the presence of potential sample clusters that might align with the exeprimental design. The CytoMDS algorithm combines, on the one hand, the concept of Earth Mover's Distance (EMD), a.k.a. Wasserstein metric and, on the other hand, the Multi Dimensional Scaling (MDS) algorithm for the low dimensional projection.

Also, the package provides some diagnostic tools for both checking the quality of the MDS projection, as well as tools to help with the interpretation of the axes of the projection.

**License** GPL-3 **Encoding** UTF-8

**Roxygen** list(markdown = TRUE)

RoxygenNote 7.3.2

BugReports https://github.com/UCLouvain-CBIO/CytoMDS/issues

URL https://uclouvain-cbio.github.io/CytoMDS

**biocViews** FlowCytometry, QualityControl, DimensionReduction, MultidimensionalScaling, Software, Visualization

Collate 'CytoMDS-package.R' 'stats.R' 'ggplots.R' 'MDS-class.R'

**Depends** R (>= 4.4)

**Imports** methods, stats, rlang, pracma, withr, flowCore, reshape2, ggplot2, ggrepel, ggforce, patchwork, transport, smacof, BiocParallel, CytoPipeline

**Suggests** testthat (>= 3.0.0), vdiffr, diffviewer, knitr, rmarkdown, BiocStyle, HDCytoData

VignetteBuilder knitr
Config/testthat/edition 3
git\_url https://git.bioconductor.org/packages/CytoMDS
git\_branch RELEASE\_3\_20

git\_last\_commit 66f50ab

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CytoMDS-package

CytoMDS: Low Dimensions projection of cytometry samples

# Description

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This package implements a low dimensional visualization of a set of cytometry samples, in order to visually assess the 'distances' between them. This, in turn, can greatly help the user to identify quality issues like batch effects or outlier samples, and/or check the presence of potential sample clusters that might align with the exeprimental design. The CytoMDS algorithm combines, on the one hand, the concept of Earth Mover's Distance (EMD), a.k.a. Wasserstein metric and, on the other hand, the Multi Dimensional Scaling (MDS) algorithm for the low dimensional projection. Also, the package provides some diagnostic tools for both checking the quality of the MDS projection, as well as tools to help with the interpretation of the axes of the projection.

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#### See Also

Useful links:

- https://uclouvain-cbio.github.io/CytoMDS
- Report bugs at https://github.com/UCLouvain-CBIO/CytoMDS/issues

channelSummaryStats

Summary statistics per channel computation

#### **Description**

Computation of summary statistic for selected channels, for all flowFrames of a flowSet, or for all expression matrices of a list. This method provides three different input modes:

- the user provides directly a flowCore::flowSet loaded in memory (RAM)
- the user provides directly a list of expression matrices of which the column names are the channel/marker names
- the user provides (1.) a number of samples nSamples; (2.) an ad-hoc function that takes as input an index between 1 and nSamples, and codes the method to load the corresponding expression matrix in memory;

# Usage

```
channelSummaryStats(
    X,
    loadExprMatrixFUN = NULL,
    loadExprMatrixFUNArgs = NULL,
    channels = NULL,
    statFUNs = stats::median,
    verbose = FALSE,
    BPPARAM = BiocParallel::SerialParam(),
    BPOPTIONS = BiocParallel::bpoptions(packages = c("flowCore"))
)
```

#### **Arguments**

Х

can be:

- a flowCore::flowSet
- a list of expression matrices (Double matrix with named columns)
- the number of samples (integer >=1)

loadExprMatrixFUN

the function used to translate an integer index into an expression matrix. In other words, the function should code how to load the indexth expression matrix into memory. IMPORTANT: the expression matrix index should be the first function argument and should be named exprMatrixIndex.

 ${\tt loadExprMatrixFUNArgs}$ 

(optional) a named list containing additional input parameters of loadExprMatrixFUN()

channels

which channels needs to be included:

- if it is a character vector, it can refer to either the channel names, or the marker names
- if it is a numeric vector, it refers to the indices of channels in fs
- if NULL, all scatter and fluorescent channels of fs #' will be selected.

statFUNs a list (possibly of length one) of functions to call to calculate the statistics, or

a simple function. This list can be named, in that case, these names will be

transfered to the returned list.

verbose if TRUE, output a message after each single statistics calculation

BPPARAM sets the BPPARAM back-end to be used for the computation. If not provided, will

use BiocParallel::SerialParam() (no task parallelization)

BPOPTIONS sets the BPOPTIONS to be passed to bplapply() function. Note that if you

use a SnowParams back-end, you need to specify all the packages that need to be loaded for the different CytoProcessingStep to work properly (visibility of functions). As a minimum, the flowCore package needs to be loaded. (hence

the default BPOPTIONS = bpoptions(packages = c("flowCore")))

#### Value

a list of named statistic matrices. In each stat matrix, the columns are the channel statistics for all flowFrames of the flowSet. Exception: if only one stat function (and not a list) is passed in statFUNs, the return value is simplified to the stat matrix itself.

```
library(CytoPipeline)
data(OMIP021Samples)
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
channelsOrMarkers <- c("FSC-A", "SSC-A", "BV785 - CD3")
# calculate mean for each 4 selected channels, for each 2 samples
channelMeans <- channelSummaryStats(</pre>
    OMIP021Trans,
    channels = channelsOrMarkers,
    statFUNs = mean)
# calculate median AND std deviation
# for each 4 selected channels, for each 2 samples
channelMedians <- channelSummaryStats(</pre>
```

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computeMetricMDS

metric MDS projection of sample

# **Description**

Multi-dimensional scaling projection of samples, using a distance matrix as an input. The MDS algorithm is not the classical MDS (cmdscale alike, aka Torgerson's algorithm), but is the SMA-COF algorithm for metric distances that are not necessarily euclidean. After having obtained the projections on the nDim dimensions, we always apply svd decomposition to visualize as first axes the ones that contain the most variance of the projected dataset in nDim dimensions. Instead of being provided directly by the user, the nDim parameter can otherwise be found iteratively by finding the minimum nDim parameter that allows the projection to reach a target pseudo RSquare. If this is the case, the maxDim parameter is used to avoid looking for too big projection spaces.

# Usage

```
computeMetricMDS(
  pwDist,
  nDim = NULL,
  seed = NULL,
  targetPseudoRSq = 0.95,
  maxDim = 128,
  ...
)
```

#### **Arguments**

. . .

	pwDist	(nSamples rows, nSamples columns), previously calculated pairwise distances between samples, must be provided as a full symmetric square matrix, with 0. diagonal
	nDim	number of dimensions of projection, as input to SMACOF algorithm if not provided, will be found iteratively using $targetPseudoRSq$
	seed	seed to be set when launching SMACOF algorithm (e.g. when init is set to "random" but not only) $ \\$
targetPseudoRSq		
		target pseudo RSquare to be reached (only used when nDim is set to NULL)
	maxDim	in case $\ensuremath{nDim}$ is found iteratively, maximum number of dimensions the search procedure is allowed to explore

additional parameters passed to SMACOF algorithm

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#### Value

a list with six elements:

- \$pwDist the initial pair-wise distance (same as input)
- \$proj the final configuration, i.e. the projected data matrix (nSamples rows, nDim columns) in nDim dimensions
- \$projDist the distance matrix of projected data
- · stress the global stress loss function final value obtained from the SMACOF algorithm
- spp the stress per point obtained from the SMACOF algorithm, i.e. the contribution of each point to the stress loss function
- \$RSq R squares, for each d, from 1 to nDim: the (pseudo) R square when taking all dims from 1 to d.
- \$GoF Goodness of fit, for each d, from 1 to nDim: the goodness of fit indicator (b/w 0 and 1) when taking all dims from 1 to d. Note pseudo R square and goodness of fit indicators are essentially the same indicator, only the definition of total sum of squares differ:
- for pseudo RSq: TSS is calculated using the mean pairwise distance as minimum
- for goodness of fit: TSS is calculated using 0 as minimum

```
library(CytoPipeline)
data(OMIP021Samples)
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
# As there are only 2 samples in OMIP021Samples dataset,
# we create artificial samples that are random combinations of both samples
ffList <- c(
    flowCore::flowSet_to_list(OMIP021Trans),
    lapply(3:5,
           FUN = function(i) {
               aggregateAndSample(
                    OMIP021Trans,
                    seed = 10*i,
                    nTotalEvents = 5000)[,1:22]
           }))
fsNames <- c("Donor1", "Donor2", paste0("Agg",1:3))</pre>
names(ffList) <- fsNames</pre>
fsAll <- as(ffList,"flowSet")</pre>
```

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**EMDDist** 

Calculate Earth Mover's distance between two samples

#### **Description**

Calculate Earth Mover's distance between two samples

# Usage

```
EMDDist(
  x1,
  x2,
  channels = NULL,
  binSize = 0.05,
  minRange = -10,
  maxRange = 10,
  returnAll = FALSE
)
```

#### **Arguments**

can be either a flowCore::flowFrame, or an expression matrix
can be either a flowCore::flowFrame, or an expression matrix
which channels (integer index(ices) or character(s)):

- if it is a character vector, it can refer to either the channel names, or the marker names if x1 and x2 have been provided as flowCore::flowFrame
- if it is a numeric vector, it refers to the indexes of channels in x1
- if NULL: if x1 and x2 are provided as flowCore::flowFrames, all scatter and fluorescent channels of x1 will be selected; if x1 and x2 are provided as expression matrices, all colnames of x1 will be selected.

binSize size of equal bins to approximate the marginal distributions.

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minRange minimum value taken when approximating the marginal distributions

maxRange maximum value taken when approximating the marginal distributions

returnAll If TRUE, distributions and marginal distribution distances are returned as well.

Default = FALSE.

#### Value

the Earth Mover's distance between x1 and x2, which is calculated by summing up all EMD approximates for the marginal distributions of each channel

```
library(CytoPipeline)
data(OMIP021Samples)
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
# distance with itself (all channels at once)
# => should return 0
dist0 <- EMDDist(</pre>
    x1 = OMIP021Trans[[1]],
    x2 = OMIP021Trans[[1]])
# returning only distance, 2 channels
dist1 <- EMDDist(</pre>
    x1 = OMIP021Trans[[1]],
    x2 = OMIP021Trans[[2]],
    channels = c("FSC-A", "SSC-A"))
# using only one channel, passed by marker name
dist2 <- EMDDist(x1 = OMIP021Trans[[1]],</pre>
                 x2 = OMIP021Trans[[2]],
                  channels = c("BV785 - CD3"))
# using only one channel, passed by index
dist3 <- EMDDist(x1 = OMIP021Trans[[1]],</pre>
                  x2 = OMIP021Trans[[2]],
                  channels = 10)
dist2 == dist3
```

```
ggplotMarginalDensities
```

Plot of channel intensity marginal densities

# **Description**

ggplotMarginalDensities uses ggplot2 to draw plots of marginal densities of selected channels of a flowSet. If the flowSet contains several flowFrames, all events are concatenated together. By default, a pseudo Rsquare projection quality indicator, and the number of dimensions of the MDS projection are provided in sub-title

### Usage

```
ggplotMarginalDensities(
    x,
    sampleSubset,
    channels,
    pDataForColour,
    pDataForGroup,
    nEventInSubsample = Inf,
    seed = NULL,
    transList
)
```

# **Arguments**

x a flowCore::flowSet (or a single flowCore::flowFrame)

sampleSubset (optional) a logical vector, of size nrow(pData), which is by construction the nb

of samples, indicating which samples to keep in the plot. Typically it is obtained

through the evaluation of a logical condition on pData rows.

channels (optional)

pDataForColour (optional) which phenoData(fs) variable will be used as colour aesthetic. Should

be a character.

pDataForGroup (optional) which phenoData(fs) variable will be used as group aesthetic. Should

be a character.

nEventInSubsample

how many event to take (per flowFrame of the flowSet).

seed if not null, used in subsampling.

transList a flowCore::transformList that will be applied before plotting.

# Value

```
a ggplot object
```

```
library(CytoPipeline)
data(OMIP021Samples)
```

```
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
# As there are only 2 samples in OMIP021Samples dataset,
# we create artificial samples that are random combinations of both samples
ffList <- c(
    flowCore::flowSet_to_list(OMIP021Trans),
    lapply(3:5,
           FUN = function(i) {
                aggregateAndSample(
                    OMIP021Trans,
                    seed = 10*i,
                    nTotalEvents = 5000)[,1:22]
           }))
fsNames <- c("Donor1", "Donor2", paste0("Agg",1:3))</pre>
names(ffList) <- fsNames</pre>
fsAll <- as(ffList, "flowSet")</pre>
flowCore::pData(fsAll)$grpId <- factor(c("D1", "D2", rep("Agg", 3)))</pre>
flowCore::pData(fsAll)$lbl <- paste0("S", 1:5)</pre>
# plot densities, all samples together
p <- ggplotMarginalDensities(fsAll)</pre>
# plot densities, per sample
p <- ggplotMarginalDensities(fsAll, pDataForGroup = "lbl")</pre>
# plot densities, per sample and coloured by group
p <- ggplotMarginalDensities(</pre>
    fsAll,
    pDataForGroup = "lb1",
    pDataForColour = "grpId")
```

 ${\tt ggplotSampleMDS}$ 

Plot of Metric MDS object

# **Description**

ggplotSampleMDS uses ggplot2 to provide plots of Metric MDS results. By default, a pseudo Rsquare projection quality indicator, and the number of dimensions of the MDS projection are provided in sub-title

#### Usage

```
ggplotSampleMDS(
 mdsObj,
 pData,
  sampleSubset,
  projectionAxes = c(1, 2),
  biplot = FALSE,
 biplotType = c("correlation", "regression"),
  extVariables,
  pDataForColour,
 pDataForShape,
 pDataForLabel,
 pDataForAdditionalLabelling,
  sizeReflectingStress = FALSE,
  title = "Multi Dimensional Scaling",
  displayPointLabels = TRUE,
 pointLabelSize = 3.88,
  repelPointLabels = TRUE,
  displayArrowLabels = TRUE,
  arrowLabelSize = 3.88,
  repelArrowLabels = FALSE,
  arrowThreshold = 0.8,
  flipXAxis = FALSE,
  flipYAxis = FALSE,
  displayPseudoRSq = TRUE,
)
```

# **Arguments**

mdsObj a MDS object, output of the computeMetricMDS() method.

pData (optional) a data.frame providing user input sample data. These can be design of

experiment variables, phenotype data per sample,... and will be used to highlight

sample categories in the plot and/or for subsetting.

sampleSubset (optional) a logical vector, of size nrow(pData), which is by construction the nb

of samples, indicating which samples to keep in the plot. Typically it is obtained

through the evaluation of a logical condition on pData rows.

projectionAxes which two axes should be plotted (should be a numeric vector of length 2)

biplot if TRUE, adds projection of external variables

biplotType type of biplot used:

• if "correlation", projection of external variables will be according to Pearson correlations w.r.t. projection axes (arrow x & y coordinates)

• if "regression", a linear regression of external variables using the 2 projection axes as explanatory variables is performed, and the projection of external variables will be according to regression coefficients (arrow direction) and R square of regression (arrow size)

extVariables

are used to generate a biplot these are the external variables that will be used in the biplot. They should be provided as a matrix with named columns corresponding to the variables. The number of rows should be the same as the number of samples. The matrix might contain some NA's, in that case only complete rows will be used to calculate biplot arrows.

pDataForColour (optional) which pData variable will be used as colour aesthetic. Should be a character.

pDataForShape (optional) which pData variable will be used as shape aesthetic. Should be a character.

pDataForLabel (optional) which pData variable will be used as point labels in the plot. Should be a character. If missing, point labels will be set equal to point names defined

in MDS object (if not NULL, otherwise no labels will be set).

pDataForAdditionalLabelling

(optional) which pData variable(s) will be add to the ggplot mapping, as to make them available for *plotly* tooltipping. Should be an array of character of maximum length 3. Note this works only if biplot=FALSE, as biplots contain circle and arrows that are currently not supported under ggplotly.

sizeReflectingStress

if TRUE, size of points will appear proportional to stress by point, i.e. the bigger the sample point appears, the less accurate its representation is (in terms of distances w.r.t. other points)

title title to give to the plot

displayPointLabels

if TRUE, displays labels attached to points (see pDataForLabels for the setting of the label values)

pointLabelSize size of point labels (default: 3.88 as in geom\_text())

repelPointLabels

if TRUE, uses ggrepel::geom\_text\_repel() instead of ggplot2::geom\_text() (try to split the labels such that they do not overlap) for the points

displayArrowLabels

if TRUE, displays arrows labels (only with biplot)

arrowLabelSize size of arrow labels (default: 3.88 as in geom\_text())

repelArrowLabels

if TRUE, uses ggrepel::geom\_text\_repel() instead of ggplot2::geom\_text() for the arrows (only with hinlet)

for the arrows (only with biplot)

arrowThreshold (only with biplot), arrows will be made barely visible if their length is (in abso-

lute value) less than this threshold.

flipXAxis if TRUE, take the opposite of x values (provided as it might ease low dimen-

sional projection comparisons)

flipYAxis if TRUE, take the opposite of y values (provided as it might ease low dimen-

sional projection comparisons)

displayPseudoRSq

if TRUE, display pseudo RSquare in subtitle, on top of nb of dimensions

... additional parameters passed to ggrepel::geom\_text\_repel() (if used)

## Value

a ggplot object

# See Also

ggplotSampleMDSWrapBiplots, ggplotSampleMDSShepard, computeMetricMDS

```
library(CytoPipeline)
data(OMIP021Samples)
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
# As there are only 2 samples in OMIP021Samples dataset,
# we create artificial samples that are random combinations of both samples
ffList <- c(
    flowCore::flowSet_to_list(OMIP021Trans),
    lapply(3:5,
           FUN = function(i) {
                aggregateAndSample(
                    OMIP021Trans,
                    seed = 10*i,
                    nTotalEvents = 5000)[,1:22]
           }))
fsNames <- c("Donor1", "Donor2", paste0("Agg",1:3))</pre>
names(ffList) <- fsNames</pre>
fsAll <- as(ffList,"flowSet")</pre>
flowCore::pData(fsAll)$type <- factor(c("real", "real", rep("synthetic", 3)))</pre>
flowCore::pData(fsAll)$grpId <- factor(c("D1", "D2", rep("Agg", 3)))</pre>
# calculate all pairwise distances
pwDist <- pairwiseEMDDist(fsAll,</pre>
                               channels = c("FSC-A", "SSC-A"),
                               verbose = FALSE)
# compute Metric MDS object with explicit number of dimensions
mdsObj <- computeMetricMDS(pwDist, nDim = 4, seed = 0)</pre>
dim \leftarrow nDim(mdsObj) # should be 4
#' # compute Metric MDS object by reaching a target pseudo RSquare
mdsObj2 <- computeMetricMDS(pwDist, seed = 0, targetPseudoRSq = 0.999)</pre>
# plot mds projection on axes 1 and 2,
```

```
# use 'grpId' for colour, 'type' for shape, and no label
p_12 <- ggplotSampleMDS(</pre>
    mdsObj = mdsObj,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(1,2),
    pDataForColour = "grpId",
    pDataForShape = "type")
# plot mds projection on axes 3 and 4,
# use 'grpId' for colour, and 'name' as point label
p_34 <- ggplotSampleMDS(</pre>
    mdsObj = mdsObj,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(3,4),
    pDataForColour = "grpId",
    pDataForLabel = "name")
# plot mds projection on axes 1 and 2,
# use 'group' for colour, 'type' for shape, and 'name' as point label
# have sample point size reflecting 'stress'
# i.e. quality of projection w.r.t. distances to other points
p12_Stress <- ggplotSampleMDS(</pre>
    mdsObj = mdsObj,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(1,2),
    pDataForColour = "grpId",
    pDataForLabel = "name",
    pDataForShape = "type",
    sizeReflectingStress = TRUE)
# try to associate axes with median of each channel
# => use bi-plot
extVars <- channelSummaryStats(</pre>
    fsAll,
    channels = c("FSC-A", "SSC-A"),
    statFUNs = stats::median)
bp_12 <- ggplotSampleMDS(</pre>
    mdsObj = mdsObj,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(1,2),
    biplot = TRUE,
    extVariables = extVars,
    pDataForColour = "grpId",
    pDataForShape = "type",
    seed = 0)
bp_34 <- ggplotSampleMDS(</pre>
    mdsObj = mdsObj,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(3,4),
    biplot = TRUE,
```

```
extVariables = extVars,
pDataForColour = "grpId",
pDataForLabel = "name",
seed = 0)
```

ggplotSampleMDSShepard

Plot of Metric MDS object - Shepard diagram

# **Description**

ggplotSampleMDSShepard uses ggplot2 to provide plot of Metric MDS results. Shepard diagram provides a scatter plot of :

- on the x axis, the high dimensional pairwise distances between each sample pairs
- on the y axis, the corresponding pairwise distances in the obtained low dimensional projection

# Usage

```
ggplotSampleMDSShepard(
  mdsObj,
  nDim,
  title = "Multi Dimensional Scaling - Shepard's diagram",
  pointSize = 0.5,
  lineWidth = 0.5,
  displayPseudoRSq = TRUE
)
```

# **Arguments**

mdsObj a MDS object, output of the computeMetricMDS() method.

nDim (optional) number of dimensions to use when calculating Shepard's diagram

and pseudoRSquare. If missing, it will be set equal to the number of projection

dimensions as calculated in mdsObj

title title to give to the plot
pointSize point size in plot
lineWidth line width in plot

 ${\tt displayPseudoRSq}$ 

if TRUE, display pseudo RSquare in subtitle, on top of nb of dimensions

# Value

a ggplot object

# See Also

ggplotSampleMDS, computeMetricMDS

```
library(CytoPipeline)
data(OMIP021Samples)
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
ffList <- c(
    flowCore::flowSet_to_list(OMIP021Trans),
    lapply(3:5,
            FUN = function(i) {
                 aggregateAndSample(
                     OMIP021Trans,
                     seed = 10*i,
                     nTotalEvents = 5000)[,1:22]
            }))
fsNames <- c("Donor1", "Donor2", paste0("Agg",1:3))
names(ffList) <- fsNames</pre>
fsAll <- as(ffList, "flowSet")</pre>
flowCore::pData(fsAll)$type <- factor(c("real", "real", rep("synthetic", 3)))</pre>
flowCore::pData(fsAll)$grpId <- factor(c("D1", "D2", rep("Agg", 3)))</pre>
# calculate all pairwise distances
pwDist <- pairwiseEMDDist(fsAll,</pre>
                                channels = c("FSC-A", "SSC-A"),
                                verbose = FALSE)
# compute Metric MDS object with explicit number of dimensions
mdsObj <- computeMetricMDS(pwDist, nDim = 4, seed = 0)</pre>
dim <- nDim(mdsObj) # should be 4
\ensuremath{\mbox{\#'}} \ensuremath{\mbox{\#'}} \ensuremath{\mbox{\#'}} tompute Metric MDS object by reaching a target pseudo RSquare
mdsObj2 <- computeMetricMDS(pwDist, seed = 0, targetPseudoRSq = 0.999)</pre>
# Shepard diagrams
p2D <- ggplotSampleMDSShepard(</pre>
    mdsObj,
    nDim = 2,
```

```
pointSize = 1,
  title = "Shepard with 2 dimensions")

p3D <- ggplotSampleMDSShepard(
  mdsObj,
  nDim = 3,
  title = "Shepard with 3 dimensions")
  #'

pDefD <- ggplotSampleMDSShepard(
  mdsObj,
  title = "Shepard with default nb of dimensions")</pre>
```

ggplotSampleMDSWrapBiplots

SampleMDS biplot wrapping

# Description

ggplotSampleMDSWrapBiplots calls ggplotSampleMDS repeatly to generate biplots with different sets of external variables and align them in a grid using the patchwork package, in a similar fashion as ggplot2::facet\_wrap() does.

### Usage

```
ggplotSampleMDSWrapBiplots(
  mdsObj,
  extVariableList,
  ncol = NULL,
  nrow = NULL,
  byrow = NULL,
  displayLegend = TRUE,
   ...
)
```

# Arguments

```
mdsObj a MDS object, output of the computeMetricMDS() method
extVariableList

should be a named list of external variable matrices Each element of the list
should be a matrix with named columns corresponding to the variables. The
number of rows should be the same as the number of samples.

ncol passed to patchwork::wrap plots()
```

```
ncol passed to patchwork::wrap_plots()
nrow passed to patchwork::wrap_plots()
byrow passed to patchwork::wrap_plots()
displayLegend if FALSE, will de-active the legend display
... additional parameters passed to ggplotSampleMDS() (if used)
```

#### Value

```
a ggplot object
```

#### See Also

ggplotSampleMDS, ggplotSampleMDSShepard, computeMetricMDS

```
library(CytoPipeline)
data(OMIP021Samples)
# estimate scale transformations
# and transform the whole OMIP021Samples
transList <- estimateScaleTransforms(</pre>
    ff = OMIP021Samples[[1]],
    fluoMethod = "estimateLogicle",
    scatterMethod = "linearQuantile",
    scatterRefMarker = "BV785 - CD3")
OMIP021Trans <- CytoPipeline::applyScaleTransforms(</pre>
    OMIP021Samples,
    transList)
# As there are only 2 samples in OMIP021Samples dataset,
# we create artificial samples that are random combinations of both samples
ffList <- c(
    flowCore::flowSet_to_list(OMIP021Trans),
    lapply(3:5,
          FUN = function(i) {
               aggregateAndSample(
                   OMIP021Trans,
                   seed = 10*i,
                   nTotalEvents = 5000)[,1:22]
           }))
fsNames <- c("Donor1", "Donor2", paste0("Agg",1:3))</pre>
names(ffList) <- fsNames</pre>
fsAll <- as(ffList, "flowSet")</pre>
flowCore::pData(fsAll)$grpId <- factor(c("D1", "D2", rep("Agg", 3)))</pre>
# calculate all pairwise distances
pwDist <- pairwiseEMDDist(fsAll,</pre>
                             channels = c("FSC-A", "SSC-A"),
                             verbose = FALSE)
# compute Metric MDS object with explicit number of dimensions
mdsObj <- computeMetricMDS(pwDist, nDim = 4, seed = 0)</pre>
dim \leftarrow nDim(mdsObj) # should be 4
#' # compute Metric MDS object by reaching a target pseudo RSquare
```

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```
mdsObj2 <- computeMetricMDS(pwDist, seed = 0, targetPseudoRSq = 0.999)</pre>
# plot mds projection on axes 1 and 2,
# use 'group' for colour, 'type' for shape, and no label
p_12 <- ggplotSampleMDS(</pre>
    mdsObj = mdsObj,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(1,2),
    pDataForColour = "grpId",
    pDataForShape = "type")
# try to associate axes with median or std deviation of each channel
# => use bi-plots
extVarList <- channelSummaryStats(</pre>
    fsAll,
    channels = c("FSC-A", "SSC-A"),
    statFUNs = c("median" = stats::median,
                  "std.dev" = stats::sd))
bpFull <- ggplotSampleMDSWrapBiplots(</pre>
    mdsObj = mdsObj,
    extVariableList = extVarList,
    pData = flowCore::pData(fsAll),
    projectionAxes = c(1,2),
    pDataForColour = "group",
    pDataForShape = "type",
    seed = 0)
```

MDS-class

MDS class

#### **Description**

Class representing Multi Dimensional Scaling (MDS) projection.

returns the value of the stress criterion, minimized by the SMACOF algorithm.

returns a vector of nPoints dimension, containing the stress indicator per point. The stress minimization criterion can indeed be allocated per represented point. The more the stress of a particular point, the less accurate its distances w.r.t. the other points.

# Usage

```
## $4 method for signature 'MDS'
show(object)

nDim(x)

nPoints(x)

pwDist(x)
```

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```
projections(x)
projDist(x)
stress(x)
spp(x)
eigenVals(x)
pctvar(x)
RSq(x)
RSqVec(x)
GoF(x)
smacofRes(x)
```

# **Arguments**

a MDS object object a MDS object

#### Value

nothing

#### **Slots**

nDim numeric, nb of dimensions of the projection

pwDist An object of class dist storing the triangular relevant part of the symmetric, zero diagonal pairwise distance matrix (nPoints \* nPoints), BEFORE projection.

proj The projection matrix, resulting from MDS

projDist An object of class dist storing the triangular relevant part of the symmetric, zero diagonal pairwise distance matrix (nPoints \* nPoints), AFTER projection.

eigen numeric, vector of nDim length, containing the eigen values of the PCA that is applied after the Smacof algorithm.

pctvar numeric, vector of nDim length, containing the percentage of explained variance per axis.

RSq\_numeric, vector of pseudo R square indicators, as a function of number of dimensions. RSq[nDim] is the global pseudo R square, as displayed on plots.

GOF numeric, vector of goodness of fit indicators, as a function of number of dimensions. GOF[nDim] is the global goodness of fit.

smacofRes an object of class 'smacofB' containing the algorithmic optimization results, for example stress and stress per point, as returned by smacof::smacofSym() method.

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#### **Examples**

```
nHD <- 10
nLD <- 2
nPoints <- 20

# generate uniformly distributed points in 10 dimensions
points <- matrix(
    data = runif(n = nPoints * nHD),
    nrow = nPoints)

# calculate euclidian distances
pwDist <- dist(points)

# compute Metric MDS object by reaching a target pseudo RSquare
mdsObj <- computeMetricMDS(pwDist, targetPseudoRSq = 0.95)
show(mdsObj)</pre>
```

pairwiseEMDDist

Pairwise Earth Mover's Distance calculation

#### **Description**

Computation of all EMD between pairs of flowFrames belonging to a flowSet. This method provides three different input modes:

- the user provides directly a flowCore::flowSet loaded in memory (RAM).
- the user provides directly a list of expression matrices loaded in RAM, of which the column names are the channel/marker names
- the user provides (1.) a number of samples nSamples; (2.) an ad-hoc function that takes as input an index between 1 and nSamples, and codes the method to load the corresponding expression matrix in memory; Optional row and column ranges can be provided to limit the calculation to a specific rectangle of the matrix. These i.e. can be specified as a way to split heavy calculations of large distance matrices on several computation nodes.

#### Usage

```
pairwiseEMDDist(
    x,
    rowRange = c(1, nSamples),
    colRange = c(min(rowRange), nSamples),
    loadExprMatrixFUN = NULL,
    loadExprMatrixFUNArgs = NULL,
    channels = NULL,
    verbose = FALSE,
    BPPARAM = BiocParallel::SerialParam(),
    BPOPTIONS = BiocParallel::bpoptions(packages = c("flowCore")),
    binSize = 0.05,
    minRange = -10,
```

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```
maxRange = 10
)
```

#### **Arguments**

x can be:

• a flowCore::flowSet

• a list of expression matrices (Double matrix with named columns)

• the number of samples (integer >=1)

rowRange the range of rows of the distance matrix to be calculated

colRange the range of columns of the distance matrix to be calculated

loadExprMatrixFUN

the function used to translate an integer index into an expression matrix. In other words, the function should code how to load the indexth expression matrix into memory. IMPORTANT: the expression matrix index should be the first function argument and should be named exprMatrixIndex.

loadExprMatrixFUNArgs

(optional) a named list containing additional input parameters of loadExprMatrixFUN()

channels which channels (integer index(ices) or character(s)):

 if it is a character vector, it can refer to either the channel names, or the marker names

• if it is a numeric vector, it refers to the indexes of channels in fs

• if NULL all scatter and fluorescent channels of fs #' will be selected

verbose if TRUE, output a message after each single distance calculation

BPPARAM sets the BPPARAM back-end to be used for the computation. If not provided, will

use BiocParallel::SerialParam() (no task parallelization)

BPOPTIONS sets the BPOPTIONS to be passed to bplapply() function. Note that if you

use a SnowParams back-end, you need to specify all the packages that need to be loaded for the different CytoProcessingStep to work properly (visibility of functions). As a minimum, the flowCore package needs to be loaded. (hence

the default BPOPTIONS = bpoptions(packages = c("flowCore")))

binSize size of equal bins to approximate the marginal distributions.

minRange minimum value taken when approximating the marginal distributions maxRange maximum value taken when approximating the marginal distributions

#### Value

a distance matrix of pairwise distances (full symmetric with 0. diagonal)

```
library(CytoPipeline)

data(OMIP021Samples)

# estimate scale transformations
# and transform the whole OMIP021Samples

transList <- estimateScaleTransforms(</pre>
```

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```
ff = OMIP021Samples[[1]],
  fluoMethod = "estimateLogicle",
  scatterMethod = "linearQuantile",
  scatterRefMarker = "BV785 - CD3")

OMIP021Trans <- CytoPipeline::applyScaleTransforms(
    OMIP021Samples,
    transList)

# calculate pairwise distances using only FSC-A & SSC-A channels
pwDist <- pairwiseEMDDist(
    x = OMIP021Trans,
    channels = c("FSC-A", "SSC-A"))</pre>
```

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