## Package 'sccore'

February 23, 2024

Title Core Utilities for Single-Cell RNA-Seq

Version 1.0.5

Description Core utilities for single-cell RNA-seq data analysis. Contained within are utility functions for working with differential expression (DE) matrices and count matrices, a collection of functions for manipulating and plotting data via 'ggplot2', and functions to work with cell graphs and cell embeddings. Graph-based methods include embedding kNN cell graphs into a UMAP <doi:10.21105/joss.00861>, collapsing vertices of each cluster in the graph, and propagating graph labels.

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Encoding UTF-8
LazyData true

**Imports** dplyr, ggplot2, ggrepel, graphics, grDevices, igraph, irlba, magrittr, Matrix, methods, parallel, pbmcapply, pROC, Rcpp, rlang, scales, stats, tibble, utils, uwot, withr

**Depends** R (>= 3.5.0)

**Suggests** ggrastr (>= 0.1.7), jsonlite, rmumps, testthat

RoxygenNote 7.2.3

LinkingTo Rcpp, RcppArmadillo, RcppProgress, RcppEigen

NeedsCompilation yes

URL https://github.com/kharchenkolab/sccore

BugReports https://github.com/kharchenkolab/sccore/issues

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Repository CRAN

**Date/Publication** 2024-02-23 00:00:02 UTC

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adjacent Vertices 3

adjacentVertices

List of adjacent vertices from igraph object

## **Description**

List of adjacent vertices from igraph object

## Usage

```
adjacentVertices(edge_verts)
```

## Arguments

```
edge_verts
```

edge vertices of igraph graph object

#### Value

list of adjacent vertices

#### **Examples**

```
## Not run:
edges <- igraph::as_edgelist(conosGraph)
adjacentVertices(edges)
## End(Not run)</pre>
```

```
adjacent_vertex_weights
```

List of adjacent vertex weights from igraph object

## Description

List of adjacent vertex weights from igraph object

## Usage

```
adjacent_vertex_weights(edge_verts, edge_weights)
```

## **Arguments**

```
edge_verts edge vertices of igraph graph object edge_weights edge weights of igraph graph object
```

## Value

list of adjacent vertices

#### **Examples**

```
## Not run:
edges <- igraph::as_edgelist(conosGraph)
edge.weights <- igraph::edge.attributes(conosGraph)$weight
adjacent_vertex_weights(edges, edge.weights)
## End(Not run)</pre>
```

appendSpecificityMetricsToDE

Append specificity metrics to DE

#### **Description**

Append specificity metrics to DE

#### Usage

```
appendSpecificityMetricsToDE(
   de.df,
   clusters,
   cluster.id,
   p2.counts,
   low.expression.threshold = 0,
   append.auc = FALSE
)
```

#### **Arguments**

de.df data.frame of differential expression values

clusters factor of clusters

cluster.id names of 'clusters' factor. If a cluster.id doesn't exist in cluster names, an error

is thrown.

p2. counts counts from Pagoda2, refer to <a href="https://github.com/kharchenkolab/pagoda2">https://github.com/kharchenkolab/pagoda2</a>

low.expression.threshold

numeric Threshold to remove expression values (default=0). Values under this

threshold are discarded.

append.auc boolean If TRUE, append AUC values (default=FALSE)

#### Value

data.frame of differential expression values with metrics attached

as\_factor 5

as\_factor

convert character vector into a factor with names "values" and "levels"

## Description

convert character vector into a factor with names "values" and "levels"

## Usage

```
as_factor(vals)
```

## Arguments

vals

vector of values to evaluate

## Value

factor with names "values" and "levels"

cellAnnotations

Conos cell annotations

## Description

Conos cell annotations

## Usage

cellAnnotations

#### **Format**

An object of class character of length 3000.

checkPackageInstalled Check whether a package is installed and suggest how to install from CRAN, Bioconductor, or other external source

#### Description

Check whether a package is installed and suggest how to install from CRAN, Bioconductor, or other external source

#### Usage

```
checkPackageInstalled(
  pkgs,
  details = "to run this function",
  install.help = NULL,
  bioc = FALSE,
  cran = FALSE
)
```

#### **Arguments**

pkgs character Package name(s)

details character Helper text (default = "to run this function")

install.help character Additional information on how to install package (default = NULL) logical Package(s) is/are available from Bioconductor (default = FALSE) cran logical Package(s) is/are available from CRAN (default = FALSE)

#### **Examples**

```
## Not run:
checkPackageInstalled("sccore", cran = TRUE)
## End(Not run)
```

collapseCellsByType

Collapse count matrices by cell type, given min/max number of cells

#### **Description**

Collapse count matrices by cell type, given min/max number of cells

#### Usage

```
collapseCellsByType(cm, groups, min.cell.count = 10, max.cell.count = Inf)
```

collapseGraphPaga 7

#### **Arguments**

cm count matrix

groups factor specifying cell types

min.cell.count numeric Minimum number of cells to include (default=10)

max.cell.count numeric Maximum number of cells to include (default=Inf). If Inf, there is no

maximum.

#### Value

Subsetted factor of collapsed cells by type, with NA cells omitted

collapseGraphPaga Collapse graph using PAGA 1.2 algorithm,

Wolf et al 2019, Genome Biology (2019) <a href="https://genomebiology.biomedcentral.com/articles/10.1186/s13059-">https://genomebiology.biomedcentral.com/articles/10.1186/s13059-</a>

019-1663-x>

#### **Description**

Collapse graph using PAGA 1.2 algorithm, Wolf et al 2019, Genome Biology (2019) <a href="https://genomebiology.biomedcentral.c019-1663-x">https://genomebiology.biomedcentral.c019-1663-x</a>

#### Usage

```
collapseGraphPaga(graph, groups, linearize = TRUE, winsorize = FALSE)
```

#### **Arguments**

graph igraph graph object Graph to be collapsed

groups factor on vertices describing cluster assignment (can specify integer vertex ids,

or character vertex names which will be matched)

linearize should normally be always TRUE (default=TRUE)

winsorize winsorize final connectivity statistics value (default=FALSE) Note: Original

PAGA has it as always TRUE, but in this case there is no way to distinguish

level of connectivity for highly connected groups.

## Value

collapsed graph

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collapseGraphSum

Collapse Graph By Sum

#### **Description**

Collapse Graph By Sum

#### Usage

```
collapseGraphSum(graph, groups, normalize = TRUE)
```

#### **Arguments**

graph igraph object Graph to be collapsed

groups factor on vertices describing cluster assignment (can specify integer vertex ids,

or character vertex names which will be matched)

normalize boolean Whether to recalculate edge weight as observed/expected (default=TRUE)

#### Value

collapsed graph

#### **Examples**

```
collapsed = collapseGraphPaga(conosGraph, igraph::V(conosGraph), linearize=TRUE, winsorize=FALSE)
```

colSumByFactor

Calculates factor-stratified sums for each column

## Description

Calculates factor-stratified sums for each column

#### Usage

```
colSumByFactor(sY, rowSel)
```

#### **Arguments**

sy sparse matrix (dgCmatrix)

rowSel integer factor. Note that the 0-th column will return sums for any NA values; 0

or negative values will be omitted

conos Cluster ListValue Matrix conosClusterList Conos clusters list Description Conos clusters list Usage conosClusterList **Format** An object of class list of length 2.  ${\tt conosGraph}$ Conos graph

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## Description

Conos graph

## Usage

 ${\tt conosGraph}$ 

#### **Format**

An object of class igraph of length 100.

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## Description

Dot plot adapted from Seurat:::DotPlot, see ?Seurat:::DotPlot for details

## Usage

```
dotPlot(
 markers,
  count.matrix,
  cell.groups,
 marker.colour = "black",
  cluster.colour = "black",
  xlab = "Marker",
  ylab = "Cluster",
  n.cores = 1,
  text.angle = 45,
  gene.order = NULL,
  cols = c("blue", "red"),
  col.min = -2.5,
  col.max = 2.5,
  dot.min = 0,
  dot.scale = 6,
  scale.by = "radius",
  scale.center = FALSE,
  scale.min = NA,
  scale.max = NA,
  verbose = FALSE,
)
```

## Arguments

| markers        | Vector of gene markers to plot   |
|----------------|--|
| count.matrix   | Merged count matrix, cells in rows and genes in columns                |
| cell.groups    | Named factor containing cell groups (clusters) and cell names as names |
| marker.colour  | Character or numeric vector (default="black")                          |
| cluster.colour | Character or numeric vector (default="black")                          |
| xlab           | string X-axis title (default="Marker")                                 |
| ylab           | string Y-axis title (default="Cluster")                                |
| n.cores        | integer Number of cores (default=1)                                    |
| text.angle     | numeric Angle of text displayed (default=45)                           |

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| gene.order   | Either factor of genes passed to dplyr::mutate(levels=gene.order), or a boolean. (default=NULL) If TRUE, gene.order is set to the unique markers. If FALSE, gene.order is set to NULL. If NULL, the argument is ignored. |
|--------------|--|
| cols         | Colors to plot (default=c("blue", "red")). The name of a palette from 'RColorBrewer::brewer.pal.info', a pair of colors defining a gradient, or 3+ colors defining multiple gradients (if 'split.by' is set).            |
| col.min      | numeric Minimum scaled average expression threshold (default=-2.5). Everything smaller will be set to this.  |
| col.max      | numeric Maximum scaled average expression threshold (default=2.5). Everything larger will be set to this.  |
| dot.min      | numeric The fraction of cells at which to draw the smallest dot (default=0). All cell groups with less than this expressing the given gene will have no dot drawn.   |
| dot.scale    | numeric Scale the size of the points, similar to cex (default=6)   |
| scale.by     | string Scale the size of the points by 'size' or by 'radius' (default="radius")  |
| scale.center | boolean Center scaling, see 'scale()' argument 'center' (default=FALSE)  |
| scale.min    | numeric Set lower limit for scaling, use NA for default (default=NA)   |
| scale.max    | numeric Set upper limit for scaling, use NA for default (default=NA)   |
| verbose      | boolean Verbose output (default=TRUE)  |
|              | Additional inputs passed to sccore::plapply(), see man for description.  |

#### Value

ggplot2 object

## **Examples**

```
library(dplyr)
## Create merged count matrix
## In this example, cms is a list of count matrices from, e.g., Cellranger count,
## where cells are in columns and genes in rows
## cm <- sccore:::mergeCountMatrices(cms, transposed = FALSE) %>% Matrix::t()
## If coming from Conos, this can be extracted like so
## cm <- conos.obj$getJointCountMatrix(raw = FALSE) # Either normalized or raw values can be used
## Here, we create a random sparse matrix
cm <- Matrix::rsparsematrix(30,3,0.5) %>% abs(.) %>%
            `dimnames<-`(list(1:30,c("gene1","gene2","gene3")))
## Create marker vector
markers <- c("gene1", "gene2", "gene3")</pre>
## Additionally, color vectors can be included.
## These should have the same length as the input (markers, cell groups)
## Otherwise, they are recycled
col.markers <- c("black","black","red") # or c(1,1,2)</pre>
col.clusters <- c("black","red","black") # or c(1,2,1)</pre>
```

embeddingColorsPlot

```
## Create annotation vector
annotation <- c(rep("cluster1",10),rep("cluster2",10),rep("cluster3",10)) %>%
    factor() %>% setNames(1:30)

## Plot. Here, the expression colours range from gray (low expression) to purple (high expression)
sccore:::dotPlot(markers = markers, count.matrix = cm, cell.groups = annotation,
    marker.colour = col.markers, cluster.colour = col.clusters, cols=c("gray", "purple"))
```

embeddingColorsPlot

Set colors for embedding plot. Used primarily in embeddingPlot().

#### **Description**

Set colors for embedding plot. Used primarily in embeddingPlot().

## Usage

```
embeddingColorsPlot(
  plot.df,
  colors,
  groups = NULL,
  geom_point_w = ggplot2::geom_point,
  gradient.range.quantile = 1,
  color.range = "symmetric",
  legend.title = NULL,
  palette = NULL,
  plot.na = TRUE
)
```

## Arguments

plot.df data.frame for plotting. In embeddingPlot(), this is a tibble from tibble::rownames\_to\_column().

colors vector of numbers, which must be shown with point colors, names contain cell

names (default=NULL). This argument is ignored if groups are provided.

groups vector of cluster labels, names contain cell names (default=NULL)

geom\_point\_w function to work with geom\_point layer from ggplot2 (default=ggplot2::geom\_point)

gradient.range.quantile

Winsorization quantile for the numeric colors and gene gradient (default=1)

color.range controls range, in which colors are estimated (default="symmetric"). Pass "all"

to estimate range based on all values of "colors", pass "data" to estimate it only based on colors, presented in the embedding. Alternatively you can pass vector

of length 2 with (min, max) values.

legend.title legend title (default=NULL)

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palette vector or list or function (default=NULL). Accepts number of colors and return

list of colors (i.e. see 'colorRampPalette') (default=NULL)

plot.na boolean/numeric Whether to plot points, for which groups / colors are missed

(default=is.null(subgroups), i.e. FALSE). If plot.na passed a numeric value below 0, the NA symbols are plotted below the cells. Otherwise if values >=0, they're plotted above the cells. Note that this argument is FALSE if 'subgroups'

is NULL

#### Value

ggplot2 object

embeddingGroupPlot

Plotting function for cluster labels, names contain cell names. Used primarily in embeddingPlot().

#### Description

Plotting function for cluster labels, names contain cell names. Used primarily in embeddingPlot().

#### Usage

```
embeddingGroupPlot(
  plot.df,
  groups,
  geom_point_w,
  min.cluster.size,
  mark.groups,
  font.size,
  legend.title,
  shuffle.colors,
  palette,
  plot.na,
  ...
)
```

#### **Arguments**

plot.df data.frame for plotting. In embeddingPlot(), this is a tibble from tibble::rownames\_to\_column().

groups vector of cluster labels, names contain cell names (default=NULL)

geom\_point\_w function to work with geom\_point layer from ggplot2 (default=ggplot2::geom\_point)

min.cluster.size

labels for all groups with number of cells fewer than this parameter are considered as missed (default=0). This argument is ignored if groups aren't provided

mark.groups plot cluster labels above points (default=TRUE)

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font size for cluster labels (default=c(3, 7)). It can either be single number for font.size constant font size or pair (min, max) for font size depending on cluster size legend title (default=NULL) legend.title shuffle.colors shuffle colors (default=FALSE) palette vector or list or function (default=NULL). Accepts number of colors and return list of colors (i.e. see 'colorRampPalette') (default=NULL) plot.na boolean/numeric Whether to plot points, for which groups / colors are missed (default=is.null(subgroups), i.e. FALSE). If plot.na passed a numeric value below 0, the NA symbols are plotted below the cells. Otherwise if values >=0, they're plotted above the cells. Note that this argument is FALSE if 'subgroups' is NULL Additional arguments passed to ggplot2::geom\_label\_repel()

#### Value

ggplot2 object

embeddingPlot

Plot embedding with provided labels / colors using ggplot2

## **Description**

Plot embedding with provided labels / colors using ggplot2

#### Usage

```
embeddingPlot(
  embedding,
  groups = NULL,
  colors = NULL,
  subgroups = NULL,
 plot.na = is.null(subgroups),
 min.cluster.size = 0,
 mark.groups = TRUE,
  show.legend = FALSE,
  alpha = 0.4,
  size = 0.8,
  title = NULL,
  plot.theme = NULL,
  palette = NULL,
  color.range = "symmetric",
  font.size = c(3, 7),
  show.ticks = FALSE,
  show.labels = FALSE,
  legend.position = NULL,
  legend.title = NULL,
```

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```
gradient.range.quantile = 1,
raster = FALSE,
raster.dpi = 300,
shuffle.colors = FALSE,
keep.limits = !is.null(subgroups),
...
)
```

#### **Arguments**

embedding two-column matrix with x and y coordinates of the embedding, rownames con-

tain cell names and are used to match coordinates with groups or colors

groups vector of cluster labels, names contain cell names (default=NULL)

colors vector of numbers, which must be shown with point colors, names contain cell

names (default=NULL). This argument is ignored if groups are provided.

subgroups subset of 'groups', selecting the cells for plot (default=NULL). Ignored if 'groups'

is NULL

plot.na boolean/numeric Whether to plot points, for which groups / colors are missed

(default=is.null(subgroups), i.e. FALSE). If plot.na passed a numeric value below 0, the NA symbols are plotted below the cells. Otherwise if values >=0, they're plotted above the cells. Note that this argument is FALSE if 'subgroups'

is NULL

min.cluster.size

labels for all groups with number of cells fewer than this parameter are considered as missed (default=0). This argument is ignored if groups aren't provided

mark.groups plot cluster labels above points (default=TRUE)

show.legend show legend (default=FALSE) alpha opacity level [0, 1] (default=0.4)

size point size (default=0.8) title plot title (default=NULL)

plot. theme for the plot (default=NULL)

palette vector or list or function (default=NULL). Accepts number of colors and return

list of colors (i.e. see 'colorRampPalette') (default=NULL)

color.range controls range, in which colors are estimated (default="symmetric"). Pass "all"

to estimate range based on all values of "colors", pass "data" to estimate it only based on colors, presented in the embedding. Alternatively you can pass vector

of length 2 with (min, max) values.

font.size font size for cluster labels (default=c(3, 7)). It can either be single number for

constant font size or pair (min, max) for font size depending on cluster size

show.ticks show ticks and tick labels (default=FALSE)

show.labels show labels (default=FALSE)

legend.position

vector with (x, y) positions of the legend (default=NULL)

legend.title legend title (default=NULL)

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gradient.range.quantile

Winsorization quantile for the numeric colors and gene gradient (default=1)

raster boolean whether layer with the points be rasterized (default=FALSE). Setting

of this argument to TRUE is useful when you need to export a plot with large

number of points

raster.dpi dpi of the rasterized plot. (default=300). Ignored if raster == FALSE.

shuffle.colors shuffle colors (default=FALSE)

keep.limits Keep axis limits from original plot (default=!is.null(subgroups)). Useful when

plotting subgroups, only meaningful it plot.na=FALSE

Arguments passed on to ggrepel::geom\_label\_repel

mapping Set of aesthetic mappings created by aes or aes\_. If specified and inherit.aes = TRUE (the default), is combined with the default mapping at the top level of the plot. You only need to supply mapping if there isn't a mapping defined for the plot.

data A data frame. If specified, overrides the default data frame defined at the top level of the plot.

stat The statistical transformation to use on the data for this layer, as a string.

position Position adjustment, either as a string, or the result of a call to a position adjustment function.

parse If TRUE, the labels will be parsed into expressions and displayed as described in ?plotmath

box.padding Amount of padding around bounding box, as unit or number. Defaults to 0.25. (Default unit is lines, but other units can be specified by passing unit(x, "units")).

label.padding Amount of padding around label, as unit or number. Defaults to 0.25. (Default unit is lines, but other units can be specified by passing unit(x, "units")).

point.padding Amount of padding around labeled point, as unit or number. Defaults to 0. (Default unit is lines, but other units can be specified by passing unit(x, "units")).

label.r Radius of rounded corners, as unit or number. Defaults to 0.15. (Default unit is lines, but other units can be specified by passing unit(x, "units")).

label.size Size of label border, in mm.

min. segment. length Skip drawing segments shorter than this, as unit or number. Defaults to 0.5. (Default unit is lines, but other units can be specified by passing unit(x, "units")).

arrow specification for arrow heads, as created by arrow

force Force of repulsion between overlapping text labels. Defaults to 1.

force\_pull Force of attraction between a text label and its corresponding data point. Defaults to 1.

max. time Maximum number of seconds to try to resolve overlaps. Defaults to

max.iter Maximum number of iterations to try to resolve overlaps. Defaults to 10000.

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max.overlaps Exclude text labels that overlap too many things. Defaults to 10.

nudge\_x, nudge\_y Horizontal and vertical adjustments to nudge the starting position of each text label. The units for nudge\_x and nudge\_y are the same as for the data units on the x-axis and y-axis.

xlim, ylim Limits for the x and y axes. Text labels will be constrained to these limits. By default, text labels are constrained to the entire plot area.

na.rm If FALSE (the default), removes missing values with a warning. If TRUE silently removes missing values.

direction "both", "x", or "y" — direction in which to adjust position of labels seed Random seed passed to set.seed. Defaults to NA, which means that set.seed will not be called.

verbose If TRUE, some diagnostics of the repel algorithm are printed

inherit.aes If FALSE, overrides the default aesthetics, rather than combining with them. This is most useful for helper functions that define both data and aesthetics and shouldn't inherit behaviour from the default plot specification, e.g. borders.

#### Value

ggplot2 object

#### **Examples**

```
library(sccore)
embeddingPlot(umapEmbedding, show.ticks=TRUE, show.labels=TRUE, title="UMAP embedding")
```

 ${\sf embedGraphUmap}$ 

Embed a graph into a UMAP, Uniform Manifold Approximation and Projection for Dimension Reduction, <a href="https://github.com/lmcinnes/umap">https://github.com/lmcinnes/umap</a>, <a href="https://github.com/lmcinnes/umap">doi:10.21105/joss.00861</a>

## **Description**

Embed a graph into a UMAP, Uniform Manifold Approximation and Projection for Dimension Reduction, <a href="https://github.com/lmcinnes/umap">https://github.com/lmcinnes/umap</a>, <a href="https://github.com/lmcinnes/umap">doi:10.21105/joss.00861</a>

#### Usage

```
embedGraphUmap(
  graph,
  min.prob = 0.001,
  min.visited.verts = 1000,
  n.cores = 1,
  max.hitting.nn.num = 0,
```

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```
max.commute.nn.num = 0,
min.prob.lower = 1e-07,
n.neighbors = 40,
n.epochs = 1000,
spread = 15,
min.dist = 0.001,
return.all = FALSE,
n.sgd.cores = n.cores,
verbose = TRUE,
...
)
```

#### **Arguments**

graph input igraph object

min.prob numeric Minimum probability for proximity when calculating hitting time per

neighbors (default=1e-3)

min.visited.verts

numeric Minimum number of vertices visted when calculating hitting time per

neighbors (default=1000)

n.cores numeric Number of cores to use (default=1)

max.hitting.nn.num

numeric Maximum adjacencies for calculating hitting time per neighbor, hit-

ting\_time\_per\_neighbors() (default=0)

max.commute.nn.num

numeric Maximum adjacencies for calculating commute time per neighbor, com-

mute\_time\_per\_node() (default=0)

min.prob.lower numeric Probability threshold to continue iteration in depth first search hitting

time, dfs\_hitting\_time() (default=1e-7)

n.neighbors numeric Number of neighbors (default=40)

n.epochs numeric Number of epochs to use during the optimization of the embedded co-

ordinates (default=1000). See 'n\_epochs' in uwot::umap()

spread numeric The effective scale of embedded points (numeric default=15). See

'spread' in uwot::umap()

min.dist numeric The effective minimum distance between embedded points (default=0.001).

See 'min.dist' in uwot::umap()

return.all boolean If TRUE, return list(adj.info=adj.info, commute.times=commute.times,

umap=umap). Otherwise, just return UMAP(default=FALSE)

n.sgd.cores numeric Number of cores to use during stochastic gradient descent. If set to >

1, then results will not be reproducible, even if 'set.seed' is called with a fixed seed before running (default=n\_threads) See 'n\_sgd\_threads' in uwot::umap()

verbose boolean Verbose output (default=TRUE)

. . . Additional arguments passed to embedKnnGraph()

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#### Value

resulting UMAP embedding

| embedKnnGraph | Embed a k-nearest neighbor (kNN) graph within a UMAP. Used within embedGraphUmap(). Please see McInnes et al <doi:10.21105 joss.00861=""> for the UMAP description and implementation.</doi:10.21105> |
|---------------|---|
|---------------|---|

## Description

Embed a k-nearest neighbor (kNN) graph within a UMAP. Used within embedGraphUmap(). Please see McInnes et al <doi:10.21105/joss.00861> for the UMAP description and implementation.

## Usage

```
embedKnnGraph(
  commute.times,
  n.neighbors,
  names = NULL,
  n.cores = 1,
  n.epochs = 1000,
  spread = 15,
  min.dist = 0.001,
  n.sgd.cores = n.cores,
  target.dims = 2,
  verbose = TRUE,
  ...
)
```

#### **Arguments**

| commute.times | graph commute times from get_nearest_neighbors(). The definition of commute_time(u, v) is the expected time starting at $u = to$ reach $v$ and then return to $v$ . |
|---------------|---|
| n.neighbors   | numeric Number of neighbors   |
| names         | vector of names for UMAP rownames (default=NULL)  |
| n.cores       | numeric Number of cores to use (except during stochastic gradient descent) (default=1). See 'n_threads' in uwot::umap()   |
| n.epochs      | numeric Number of epochs to use during the optimization of the embedded coordinates (default=1000). See 'n_epochs' in uwot::umap()                                  |
| spread        | numeric The effective scale of embedded points (numeric default=15). See 'spread' in uwot::umap()   |
| min.dist      | numeric The effective minimum distance between embedded points (default=0.001). See 'min.dist' in uwot::umap()  |

20 extendMatrix

n.sgd.cores

numeric Number of cores to use during stochastic gradient descent. If set to >
1, then results will not be reproducible, even if 'set.seed' is called with a fixed seed before running (default=n.cores) See 'n\_sgd\_threads' in uwot::umap()

target.dims

numeric Dimensions for 'n\_components' in uwot::umap(n\_components=target.dims) (default=2)

verbose

boolean Verbose output (default=TRUE)

arguments passed to uwot::umap()

#### Value

resulting kNN graph embedding within a UMAP

extendMatrix Extend matrix to include new columns in matrix

## **Description**

Extend matrix to include new columns in matrix

#### Usage

```
extendMatrix(mtx, col.names)
```

#### **Arguments**

mtx Matrix

col.names Columns that should be included in matrix

#### Value

Matrix with new columns but rows retained

## Examples

```
library(dplyr)
gene.union <- lapply(conosClusterList, colnames) %>% Reduce(union, .)
extendMatrix(conosClusterList[[1]], col.names=gene.union)
```

fac2col 21

fac2col

Utility function to translate a factor into colors

#### **Description**

Utility function to translate a factor into colors

#### Usage

```
fac2col(
    X,
    s = 1,
    v = 1,
    shuffle = FALSE,
    min.group.size = 1,
    return.details = FALSE,
    unclassified.cell.color = "gray50",
    level.colors = NULL
)
```

#### **Arguments**

```
input factor
Χ
                  numeric The "saturation" to be used to complete the HSV color descriptions
s
                  (default=1) See ?rainbow in Palettes, grDevices
                  numeric The "value" to be used to complete the HSV color descriptions (de-
                  fault=1) See ?rainbow in Palettes, grDevices
shuffle
                  boolean If TRUE, shuffles columns with shuffle(columns) (default=FALSE)
min.group.size integer Exclude groups of size less than the min.group.size (default=1)
return.details boolean If TRUE, returns a list list(colors=y, palette=col). Otherwise, just re-
                  turns the factor (default=FALSE)
unclassified.cell.color
                  Color for unclassified cells (default='gray50')
level.colors
                  (default=NULL)
```

#### Value

vector or list of colors

#### **Examples**

```
genes = factor(c("BRAF", "NPC1", "PAX3", "BRCA2", "FMR1"))
fac2col(genes)
```

22 getClusterGraph

| fac2palette | Encodes logic of how to handle named-vector and functional palettes.  Used primarily within embeddingGroupPlot() |
|-------------|--|
|             |  |

## **Description**

Encodes logic of how to handle named-vector and functional palettes. Used primarily within embeddingGroupPlot()

#### Usage

```
fac2palette(groups, palette, unclassified.cell.color = "gray50")
```

## **Arguments**

```
groups vector of cluster labels, names contain cell names

palette vector or list or function (default=NULL). Accepts number of colors and return list of colors (i.e. see 'colorRampPalette')

unclassified.cell.color

Color for unclassified cells (default='gray50')
```

#### Value

vector or palette

getClusterGraph

Collapse vertices belonging to each cluster in a graph

## Description

Collapse vertices belonging to each cluster in a graph

## Usage

```
getClusterGraph(
  graph,
  groups,
  method = "sum",
  plot = FALSE,
  node.scale = 50,
  edge.scale = 50,
  edge.alpha = 0.3,
  seed = 1,
  ...
)
```

get\_nearest\_neighbors 23

## Arguments

| graph      | igraph graph object Graph to be collapsed  |
|------------|--|
| groups     | factor on vertices describing cluster assignment (can specify integer vertex ids, or character vertex names which will be matched) |
| method     | string Method to be used, either "sum" or "paga" (default="sum")   |
| plot       | boolean Whether to show collapsed graph plot (default=FALSE)   |
| node.scale | numeric Scaling to control value of 'vertex.size' in plot.igraph() (default=50)  |
| edge.scale | numeric Scaling to control value of 'edge.width' in plot.igraph() (default=50)   |
| edge.alpha | numeric Scaling to control value of 'alpha.f' in adjustcolor() within plot.igraph() (default=0.3)                                  |
| seed       | numeric Set seed via set.seed() for plotting (default=1)   |
| • • •      | arguments passed to collapseGraphSum()   |
|            |  |

#### Value

collapsed graph

## **Examples**

```
cluster.graph = getClusterGraph(conosGraph, igraph::V(conosGraph))
```

 ${\tt get\_nearest\_neighbors} \ \ \textit{Get nearest neighbors method on graph}$ 

## Description

Get nearest neighbors method on graph

## Usage

```
get_nearest_neighbors(
  adjacency_list,
  transition_probabilities,
  n_verts = 0L,
  n_cores = 1L,
  min_prob = 0.001,
  min_visited_verts = 1000L,
  min_prob_lower = 1e-05,
  max_hitting_nn_num = 0L,
  max_commute_nn_num = 0L,
  verbose = TRUE
)
```

24 graphToAdjList

#### **Arguments**

adjacency\_list igraph adjacency list

transition\_probabilities

vector of transition probabilites

n\_verts numeric Number of vertices (default=0)

n\_cores numeric Number of cores to use (default=1)

min\_prob numeric Minimum probability for proximity when calculating hitting time per

neighbors (default=1e-3)

min\_visited\_verts

numeric Minimum number of vertices visted when calculating hitting time per

neighbors (default=1000)

min\_prob\_lower numeric Probability threshold to continue iteration in depth first search hitting

time, dfs\_hitting\_time() (default=1e-5)

max\_hitting\_nn\_num

numeric Maximum adjacencies for calculating hitting time per neighbor, hit-

ting\_time\_per\_neighbors() (default=0)

max\_commute\_nn\_num

numeric Maximum adjacencies for calculating commute time per neighbor, com-

mute\_time\_per\_node() (default=0)

verbose boolean Whether to have verbose output (default=TRUE)

#### Value

list of commute times based on adjacencies

graphToAdjList

Convert igraph graph into an adjacency list

#### **Description**

Convert igraph graph into an adjacency list

#### **Usage**

graphToAdjList(graph)

## **Arguments**

graph input igraph object

#### Value

adjacency list, defined by list(idx=adj.list, probabilities=probs, names=edge.list.fact\$levels

heatFilter 25

## **Examples**

```
library(dplyr)
edge.list.fact <- igraph::as_edgelist(conosGraph) %>% as_factor()
edge.list <- matrix(edge.list.fact$values, ncol=2)
n.nodes <- length(igraph::V(conosGraph))
splitVectorByNodes(edge.list[,1], edge.list[,2], n.nodes)</pre>
```

heatFilter

*Graph filter with the heat kernel:*  $f(x) = exp(-\beta|x/\lambda_m - a|^b)$ 

## Description

```
Graph filter with the heat kernel: f(x) = exp(-\beta|x/\lambda_m - a|^b)
```

#### Usage

```
heatFilter(x, 1.max, order = 1, offset = 0, beta = 30)
```

## Arguments

| X      | numeric Values to be filtered. Normally, these are graph laplacian engenvalues.  |
|--------|--|
| 1.max  | numeric Maximum eigenvalue on the graph ( $\lambda_m$ in the equation)   |
| order  | numeric Parameter $b$ in the equation. Larger values correspond to the sharper kernel form (default=1). The values should be positive.   |
| offset | numeric Mean kernel value ( $a$ in the equation), must be in [0:1] (default=0)   |
| beta   | numeric Parameter $\beta$ in the equation. Larger values provide stronger smoothing. $\beta=0$ corresponds to no smoothing (default=30). |

#### Value

```
smoothed values for 'x'
```

## See Also

Other graph smoothing: computeChebyshevCoeffs(), smoothChebyshev(), smoothSignalOnGraph()

26 mergeCountMatrices

| jsDist | Jensen–Shannon distance metric (i.e. the square root of the Jensen–Shannon divergence) between the columns of a dense matrix |
|--------|--|
|        | m  |

#### **Description**

Jensen-Shannon distance metric (i.e. the square root of the Jensen-Shannon divergence) between the columns of a dense matrix m

#### Usage

```
jsDist(m)
```

#### Arguments

m

Input matrix

#### Value

Vectorized version of the lower triangle as an R distance object, stats::dist()

## **Examples**

```
ex = matrix(1:9, nrow = 3, ncol = 3)
jsDist(ex)
```

 ${\tt mergeCountMatrices}$ 

Merge list of count matrices into a common matrix, entering 0s for the missing entries

## **Description**

Merge list of count matrices into a common matrix, entering 0s for the missing entries

#### Usage

```
mergeCountMatrices(cms, transposed = FALSE, ...)
```

#### **Arguments**

cms List of count matrices

transposed boolean Indicate whether 'cms' is transposed, e.g. cells in rows and genes in

columns (default=FALSE)

.. Parameters for 'plapply' function

multi2dend 27

## Value

A merged extended matrix, with 0s for missing entries

## **Examples**

```
mergeCountMatrices(conosClusterList, n.cores=1)
## 12 x 67388 sparse Matrix of class "dgCMatrix"
```

| multi2dend | Translate multilevel segmentation into a dendrogram, with the lowest |
|------------|--|
|            | level of the dendrogram listing the cells                            |

## Description

Translate multilevel segmentation into a dendrogram, with the lowest level of the dendrogram listing the cells

## Usage

```
multi2dend(cl, counts, deep = FALSE, dist = "cor")
```

## Arguments

| cl     | igraph communities object, returned from igraph community detection functions   |
|--------|---|
| counts | dgCmatrix of counts   |
| deep   | $boolean\ If\ TRUE, take\ (cl\ memberships[1,]).\ Otherwise,\ uses\ as.integer (membership(cl))\ (default=FALSE)$   |
| dist   | Distance metric used (default='cor'). Eiether 'cor' for the correlation distance in log10 space, or 'JS' for the Jensen–Shannon distance metric (i.e. the square root of the Jensen–Shannon divergence) |

#### Value

resulting dendrogram

28 plapply

| plapply Parallel, optionally verbose lapply. See ?parallel::mclapply for more info. | plapply |  |
|---|---------|--|
|---|---------|--|

## Description

Parallel, optionally verbose lapply. See ?parallel::mclapply for more info.

## Usage

```
plapply(
  . . . ,
 progress = FALSE,
 n.cores = parallel::detectCores(),
 mc.preschedule = FALSE,
 mc.allow.recursive = TRUE,
  fail.on.error = FALSE
)
```

#### Arguments

Additional arguments passed to mclapply(), lapply(), or pbmcapply::pbmclapply() Show progress bar via pbmcapply::pbmclapply() (default=FALSE). progress Number of cores to use (default=parallel::detectCores()). When n.cores=1, regn.cores ular lapply() is used. Note: doesn't work when progress=TRUE mc.preschedule if set to TRUE then the computation is first divided to (at most) as many jobs are there are cores and then the jobs are started, each job possibly covering more than one value. If set to FALSE then one job is forked for each value of X. The former is better for short computations or large number of values in X, the latter is better for jobs that have high variance of completion time and not too many values of X compared to mc.cores. mc.allow.recursive boolean Unless true, calling mclapply in a child process will use the child and

not fork again (default=TRUE)

fail.on.error

boolean Whether to fail and report and error (using stop()) as long as any of the individual tasks has failed (default =FALSE)

#### Value

list, as returned by lapply

## **Examples**

```
square = function(x){ x**2 }
plapply(1:10, square, n.cores=1, progress=TRUE)
```

propagateLabels 29

| propagateLabels Estimate labeling distribution for each vertex, based on provide bels. | 'ed la- |
|--|---------|
|--|---------|

#### **Description**

Estimate labeling distribution for each vertex, based on provided labels.

## Usage

```
propagateLabels(graph, labels, method = "diffusion", ...)
```

#### **Arguments**

| graph  | igraph graph object  |
|--------|--|
| labels | vector of factor or character labels, named by cell names, used in propagateLabelsSolver() and propagateLabelsDiffusion()  |
| method | string Type of propagation. Either 'diffusion' or 'solver'. (default='diffusion') 'solver' gives better result but has bad asymptotics, so it is inappropriate for datasets > 20k cells. |
|        | additional arguments passed to either propagateLabelsSolver() or propagateLabelsDiffusion()  |

#### Value

matrix with distribution of label probabilities for each vertex by rows.

## **Examples**

```
propagateLabels(conosGraph, labels=cellAnnotations)
```

```
propagateLabelsDiffusion
```

Estimate labeling distribution for each vertex, based on provided labels using a Random Walk on graph

## Description

Estimate labeling distribution for each vertex, based on provided labels using a Random Walk on graph

#### Usage

```
propagateLabelsDiffusion(
  graph,
  labels,
  max.iters = 100,
  diffusion.fading = 10,
  diffusion.fading.const = 0.1,
  tol = 0.025,
  fixed.initial.labels = TRUE,
  verbose = TRUE
)
```

#### **Arguments**

graph igraph graph object Graph input labels vector of factor or character labels, named by cell names integer Maximal number of iterations (default=100) max.iters diffusion.fading numeric Constant used for diffusion on the graph, exp(-diffusion.fading \* (edge\_length + diffusion.fading.const)) (default=10.0) diffusion.fading.const numeric Another constant used for diffusion on the graph, exp(-diffusion.fading \* (edge\_length + diffusion.fading.const)) (default=0.1) numeric Absolute tolerance as a stopping criteria (default=0.025) tol fixed.initial.labels boolean Prohibit changes of initial labels during diffusion (default=TRUE) boolean Verbose mode (default=TRUE) verbose

#### Value

matrix from input graph, with labels propagated

#### **Examples**

```
propagateLabelsDiffusion(conosGraph, labels=cellAnnotations)
```

```
propagateLabelsSolver Propagate labels using Zhu, Ghahramani, Lafferty (2003) algorithm, "Semi-Supervised Learning Using Gaussian Fields and Harmonic Functions" <a href="http://mlg.eng.cam.ac.uk/zoubin/papers/zgl.pdf">http://mlg.eng.cam.ac.uk/zoubin/papers/zgl.pdf</a>
```

#### **Description**

Propagate labels using Zhu, Ghahramani, Lafferty (2003) algorithm, "Semi-Supervised Learning Using Gaussian Fields and Harmonic Functions" <a href="http://mlg.eng.cam.ac.uk/zoubin/papers/zgl.pdf">http://mlg.eng.cam.ac.uk/zoubin/papers/zgl.pdf</a>

propagate\_labels 31

#### Usage

```
propagateLabelsSolver(graph, labels, solver = "mumps")
```

## Arguments

graph igraph object Graph input

labels vector of factor or character labels, named by cell names

solver Method of solver to use (default="mumps"), either "Matrix" or "mumps" (i.e.

"rmumps::Rmumps")

#### Value

result from Matrix::solve() or rmumps::Rmumps

#### **Examples**

```
propagateLabelsSolver(conosGraph, labels=cellAnnotations)
```

propagate\_labels

Label propagation

## Description

Label propagation

## Usage

```
propagate_labels(
  edge_verts,
  edge_weights,
  vert_labels,
  max_n_iters = 10L,
  verbose = TRUE,
  diffusion_fading = 10,
  diffusion_fading_const = 0.5,
  tol = 0.005,
  fixed_initial_labels = FALSE
)
```

32 saveDeAsJson

## **Arguments**

```
edge_verts
                  edge vertices of igraph graph object
                  edge weights of igraph graph object
edge_weights
vert_labels
                  vector of factor or character labels, named by cell names
max_n_iters
                  integer Maximal number of iterations (default=10)
verbose
                  boolean Verbose mode (default=TRUE)
diffusion fading
                  numeric Constant used for diffusion on the graph, exp(-diffusion.fading * (edge_length
                  + diffusion.fading.const)) (default=10.0)
diffusion_fading_const
                  numeric Another constant used for diffusion on the graph, exp(-diffusion.fading
                  * (edge_length + diffusion.fading.const)) (default=0.5)
tol
                  numeric Absolute tolerance as a stopping criteria (default=5e-3)
fixed_initial_labels
                  boolean Prohibit changes of initial labels during diffusion (default=FALSE)
```

#### Value

matrix from input graph, with labels propagated

saveDeAsJson Save DE results as JSON tables for viewing in browser

Description

Save DE results as JSON tables for viewing in browser

#### Usage

```
saveDeAsJson(
  de.raw,
  sample.groups = NULL,
  saveprefix = NULL,
  dir.name = "JSON",
  gene.metadata = NULL,
  verbose = TRUE
)
```

#### **Arguments**

de.raw List of DE results from e.g. cacoa, conos

sample groups Sample groups as named list, each element containing a vector of samples. Can

be retrieved from e.g. package cacoa (default=NULL)

saveprefix Prefix for created files (default=NULL)

setMinMax 33

dir.name Name for directory with results. If it doesn't exist, it will be created. To disable,

set as NULL (default="JSON")

gene.metadata (default=NULL) # Needs explanation

verbose Show progress (default=TRUE)

#### Value

JSON files, table of content, and viewer files for viewing DE results in browser

#### **Examples**

```
## Not run:
saveDeAsJson(de.raw, sample.groups)
## End(Not run)
## The results can be viewed in a webbrowser by opening toc.html
```

setMinMax Set range for values in object. Changes values outside of range to min

or max. Adapted from Seurat::MinMax

#### **Description**

Set range for values in object. Changes values outside of range to min or max. Adapted from Seurat::MinMax

#### Usage

```
setMinMax(obj, min, max)
```

## **Arguments**

obj Object to manipulate
min Minimum value
max Maximum value

#### Value

An object with the same dimensions as input but with altered range in values

#### **Examples**

```
example_matrix = matrix(rep(c(1:5), 3), 5)
setMinMax(example_matrix, 2, 4)
```

smoothSignalOnGraph

Smooth Signal on Graph

#### Description

Smooth Signal on Graph

#### Usage

```
smoothSignalOnGraph(
  signal,
  filter,
  graph = NULL,
  lap = NULL,
  l.max = NULL,
  m = 50,
  ...
)
```

#### **Arguments**

```
signal
                  signal to be smoothed
filter
                  function that accepts signal 'x' and the maximal Laplacian eigenvalue 'l.max'.
                  See heatFilter as an example.
                  igraph object with the graph (default=NULL)
graph
lap
                  graph laplacian (default=NULL). If NULL, 'lap' estimated from graph.
1.max
                  maximal eigenvalue of 'lap' (default=NULL). If NULL, estimated from 'lap'.
                  numeric Maximum order of Chebyshev coeff to compute (default=50)
                  Arguments passed on to smoothChebyshev
                  n.cores numeric Number of cores for parallel run (default=1)
                  progress. chunks numeric Number of chunks per core for estimating progress
                      (default=5). Large values are not suggested, as it may bring overhead.
                 progress boolean Flag on whether progress must be shown (default=TRUE,
                      i.e. 'progress.chunks > 1')
```

## See Also

Other graph smoothing: computeChebyshevCoeffs(), heatFilter(), smoothChebyshev()

smooth\_count\_matrix 35

smooth\_count\_matrix

Smooth gene expression, used primarily within conos::correctGenes. Used to smooth gene expression values in order to better represent the graph structure. Use diffusion of expression on graph with the equation dv = exp(-a \* (v + b))

#### **Description**

Smooth gene expression, used primarily within conos::correctGenes. Used to smooth gene expression values in order to better represent the graph structure. Use diffusion of expression on graph with the equation  $dv = \exp(-a * (v + b))$ 

#### Usage

```
smooth_count_matrix(
  edge_verts,
  edge_weights,
  count_matrix,
  is_label_fixed,
  max_n_iters = 10L,
  diffusion_fading = 1,
  diffusion_fading_const = 0.1,
  tol = 0.001,
  verbose = TRUE,
  normalize = FALSE
)
```

## **Arguments**

```
edge_verts
                  edge vertices of igraph graph object
edge_weights
                  edge weights of igraph graph object
count_matrix
                  gene count matrix
is_label_fixed boolean Whether label is fixed
max_n_iters
                  integer Maximal number of iterations (default=10)
diffusion_fading
                  numeric Constant used for diffusion on the graph, exp(-diffusion.fading * (edge_length
                  + diffusion.fading.const)) (default=1.0)
diffusion_fading_const
                  numeric Another constant used for diffusion on the graph, exp(-diffusion.fading
                  * (edge_length + diffusion.fading.const)) (default=0.1)
tol
                  numeric Absolute tolerance as a stopping criteria (default=1e-3)
                  boolean Verbose mode (default=TRUE)
verbose
normalize
                  boolean Whether to normalize values (default=FALSE)
```

36 splitVectorByNodes

#### Value

matrix from input graph, with labels propagated

sn

Set names equal to values, a stats::setNames wrapper function

## Description

Set names equal to values, a stats::setNames wrapper function

#### Usage

sn(x)

## **Arguments**

Х

an object for which names attribute will be meaningful

#### Value

An object with names assigned equal to values

## **Examples**

```
vec = c(1, 2, 3, 4)

sn(vec)
```

 ${\tt splitVectorByNodes}$ 

splitVectorByNodes

## Description

```
splitVectorByNodes
```

#### Usage

```
splitVectorByNodes(vec, nodes, n.nodes)
```

#### **Arguments**

vec input vector to be divided

nodes nodes used to divide the vector 'vec' via split()
n.nodes numeric The number of nodes for splitting

styleEmbeddingPlot 37

#### Value

list from vec with names given by the nodes

#### **Examples**

```
adjList = graphToAdjList(conosGraph)
print(names(adjList))
## [1] "idx" "probabilities" "names"
length(adjList$names)
## [1] 12000
```

styleEmbeddingPlot

Set plot.theme, legend, ticks for embedding plot. Used primarily in embedding Plot().

## Description

Set plot.theme, legend, ticks for embedding plot. Used primarily in embeddingPlot().

#### Usage

```
styleEmbeddingPlot(
   gg,
   plot.theme = NULL,
   title = NULL,
   legend.position = NULL,
   show.legend = TRUE,
   show.ticks = TRUE,
   show.labels = TRUE,
   relabel.axis = TRUE
)
```

## Arguments

```
ggplot2 object to plot
plot.theme
                 theme for the plot (default=NULL)
                 plot title (default=NULL)
title
legend.position
                  vector with (x, y) positions of the legend (default=NULL)
                 show legend (default=TRUE)
show.legend
show.ticks
                  show ticks and tick labels (default=TRUE)
show.labels
                 show labels (default=TRUE)
relabel.axis
                 boolean If TRUE, relabel axes with ggplot2::labs(x='Component 1', y='Component
                 2') (default=TRUE)
```

38 val2col

#### Value

ggplot2 object

umapEmbedding

UMAP embedding

#### **Description**

**UMAP** embedding

#### Usage

umapEmbedding

#### **Format**

An object of class matrix (inherits from array) with 12000 rows and 2 columns.

val2col

*Utility function to translate values into colors.* 

## **Description**

Utility function to translate values into colors.

#### **Usage**

```
val2col(x, gradientPalette = NULL, zlim = NULL, gradient.range.quantile = 0.95)
```

#### **Arguments**

```
x input values
```

gradientPalette

gradient palette (default=NULL). If NULL, use colorRampPalette(c('gray90','red'), space = "Lab")(1024) if the values are non-negative; otherwise colorRamp-

Palette(c("blue", "grey90", "red"), space = "Lab")(1024) is used

zlim

a two-value vector specifying limits of the values that should correspond to the

extremes of the color gradient

gradient.range.quantile

extreme quantiles of values that should be trimmed prior to color mapping (default=0.95)

#### **Examples**

```
colors <- val2col( rnorm(10) )</pre>
```

val2ggcol 39

| val2ggcol | Helper function to return a ggplot color gradient for a numeric vector $ggplot(aes(color=x,),) + val2ggcol(x)$ |
|-----------|--|
|           |  |

## Description

Helper function to return a ggplot color gradient for a numeric vector ggplot(aes(color=x, ...), ...) + val2ggcol(x)

## Usage

```
val2ggcol(
  values,
  gradient.range.quantile = 1,
  color.range = "symmetric",
  palette = NULL,
  midpoint = NULL,
  oob = scales::squish,
  return.fill = FALSE,
  ...
)
```

## Arguments

| values<br>gradient.range | values by which the color gradient is determined quantile   |
|--------------------------|---|
|                          | numeric Trimming quantile (default=1). Either a single number or two numbers - for lower and upper quantile.  |
| color.range              | either a vector of two values explicitly specifying the values corresponding to the start/end of the gradient, or string "symmetric" or "all" (default="symmetric"). "symmetric": range will fit data, but will be symmetrized around zeros, "all": gradient will match the span of the range of the data (after gradient.range.quantile) |
| palette                  | an optional palette (default=NULL). The default becomes blue-gray90-red; if the values do not straddle 0, then truncated gradients (blue-gray90 or gray90-red) will be used   |
| midpoint                 | optional midpoint (default=NULL). Set for the center of the resulting range by default  |
| oob                      | function to determine what to do with the values outside of the range (default =scales::squish). Refer to 'oob' parameter in ggplot   |
| return.fill              | boolean Whether to return fill gradients instead of color (default=FALSE)   |
| •••                      | additional arguments are passed to ggplot2::scale_color_gradient* functions, i.e. scale_color_gradient(), scale_color_gradient(), scale_color_gradientn()   |

## Value

```
ggplot2::scale_colour_gradient object
```

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