# Package 'clustree'

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Type Package

**Title** Visualise Clusterings at Different Resolutions

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Maintainer Luke Zappia < luke.zappia@mcri.edu.au>

**Description** Deciding what resolution to use can be a difficult question when approaching a clustering analysis. One way to approach this problem is to look at how samples move as the number of clusters increases. This package allows you to produce clustering trees, a visualisation for interrogating clusterings as resolution increases.

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Author Luke Zappia [aut, cre] (<a href="https://orcid.org/0000-0001-7744-8565">https://orcid.org/0000-0001-7744-8565</a>),
Alicia Oshlack [aut] (<a href="https://orcid.org/0000-0001-9788-5690">https://orcid.org/0000-0001-9788-5690</a>),
Andrea Rau [ctb],
Paul Hoffman [ctb] (<a href="https://orcid.org/0000-0002-7693-8957">https://orcid.org/0000-0002-7693-8957</a>)

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## **Description**

Deciding what resolution to use can be a difficult question when approaching a clustering analysis. One way to approach this problem is to look at how samples move as the number of clusters increases. This package allows you to produce clustering trees, a visualisation for interrogating clusterings as resolution increases.

add\_node\_labels

Add node labels

## **Description**

Add node labels to a clustering tree plot with the specified aesthetics.

```
add_node_labels(node_label, node_colour, node_label_size,
    node_label_colour, node_label_nudge, allowed)
```

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## **Arguments**

node\_label the name of a metadata column for node labels

node\_colour either a value indicating a colour to use for all nodes or the name of a metadata

column to colour nodes by

node\_label\_size

size of node label text

node\_label\_colour

colour of node\_label text

node\_label\_nudge

numeric value giving nudge in y direction for node labels

allowed vector of allowed node attributes to use as aesthetics

add\_node\_points

Add node points

## **Description**

Add node points to a clustering tree plot with the specified aesthetics.

## Usage

```
add_node_points(node_colour, node_size, node_alpha, allowed)
```

## **Arguments**

node_colour	either a value indicating a colour to use for all nodes or the name of a metadata column to colour nodes by
node_size	either a numeric value giving the size of all nodes or the name of a metadata column to use for node sizes
node_alpha	either a numeric value giving the alpha of all nodes or the name of a metadata column to use for node transparency
allowed	vector of allowed node attributes to use as aesthetics

aggr\_metadata

Aggregate metadata

## **Description**

Aggregate a metadata column to get a summarized value for a cluster node

## Usage

```
aggr_metadata(node_data, col_name, col_aggr, metadata, is_cluster)
```

## **Arguments**

node_data	data.frame containing information about a set of cluster nodes
col_name	the name of the metadata column to aggregate
col_aggr	string naming a function used to aggregate the column
metadata	data.frame providing metadata on samples
is_cluster	logical vector indicating which rows of metadata are in the node to be summa-

## rized

### Value

data.frame with aggregated data

```
assert_colour_node_aes
```

Assert colour node aesthetics

## **Description**

Raise error if an incorrect set of colour node parameters has been supplied.

```
assert_colour_node_aes(node_aes_name, prefix, metadata, node_aes,
  node_aes_aggr, min, max)
```

assert\_node\_aes 5

### **Arguments**

node\_aes\_name name of the node aesthetic to check

prefix string indicating columns containing clustering information

metadata data.frame containing metadata on each sample that can be used as node aes-

thetics

node\_aes value of the node aesthetic to check

node\_aes\_aggr aggregation function associated with the node aesthetic

min minimum numeric value allowed
max maximum numeric value allowed

assert\_node\_aes

Assert node aesthetics

## **Description**

Raise error if an incorrect set of node parameters has been supplied.

## **Usage**

```
assert_node_aes(node_aes_name, prefix, metadata, node_aes, node_aes_aggr)
```

### Arguments

node\_aes\_name name of the node aesthetic to check

prefix string indicating columns containing clustering information

metadata data.frame containing metadata on each sample that can be used as node aes-

thetics

node\_aes value of the node aesthetic to check

node\_aes\_aggr aggregation function associated with the node aesthetic

assert\_numeric\_node\_aes

Assert numeric node aesthetics

## **Description**

Raise error if an incorrect set of numeric node parameters has been supplied.

```
assert_numeric_node_aes(node_aes_name, prefix, metadata, node_aes,
  node_aes_aggr, min, max)
```

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### Arguments

node\_aes\_name name of the node aesthetic to check

prefix string indicating columns containing clustering information

metadata data.frame containing metadata on each sample that can be used as node aes-

thetics

node\_aes value of the node aesthetic to check

node\_aes\_aggr aggregation function associated with the node aesthetic

min minimum numeric value allowed
max maximum numeric value allowed

build\_tree\_graph

Build tree graph

### **Description**

Build a tree graph from a set of clusterings, metadata and associated aesthetics

### Usage

```
build_tree_graph(clusterings, prefix, count_filter, prop_filter, metadata,
  node_aes_list)
```

### **Arguments**

prop\_filter

clusterings numeric matrix containing clustering information, each column contains clus-

tering at a separate resolution

prefix string indicating columns containing clustering information

count\_filter count threshold for filtering edges in the clustering graph

metadata data.frame containing metadata on each sample that can be used as node aes-

in proportion threshold for filtering edges in the clustering graph

thetics

node\_aes\_list nested list containing node aesthetics

#### Value

tidygraph::tbl\_graph object containing the tree graph

calc\_sc3\_stability 7

calc\_sc3\_stability

Calculate SC3 stability

### **Description**

Calculate the SC3 stability index for every cluster at every resolution in a set of clusterings. The index varies from 0 to 1, where 1 suggests that a cluster is more stable across resolutions. See calc\_sc3\_stability\_cluster() for more details.

## Usage

```
calc_sc3_stability(clusterings)
```

### Arguments

clusterings

numeric matrix containing clustering information, each column contains clus-

tering at a separate resolution

### Value

matrix with stability score for each cluster

```
calc_sc3_stability_cluster
```

Calculate single SC3 stability

### **Description**

Calculate the SC3 stability index for a single cluster in a set of clusterings. The index varies from 0 to 1, where 1 suggests that a cluster is more stable across resolutions.

## Usage

```
calc_sc3_stability_cluster(clusterings, res, cluster)
```

### **Arguments**

clusterings numeric matrix containing clustering information, each column contains clus-

tering at a separate resolution

res resolution of the cluster to calculate stability for

cluster index of the cluster to calculate stability for

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#### **Details**

This index was originally introduced in the SC3 package for clustering single-cell RNA-seq data. Clusters are awarded increased stability if they share the same samples as a cluster at another resolution and penalised at higher resolutions. We use a slightly different notation to describe the score but the results are the same:

$$s(c_{k,i}) = \frac{1}{size(L) + 1} \sum_{l \in L} \sum_{j \in N_l} \frac{size(c_{k,i} \cap c_{l,j})}{size(c_{l,j}) * size(N_l)^2}$$

#### Where:

- c\_{x, y} is cluster y at resolution x
- k is the resolution of the cluster we want to score
- i is the index of the cluster we want to score
- L is the set of all resolutions except k
- 1 is a resolution in L
- $N_1$  is the set of clusters at resolution 1 that share samples with  $c_{k, i}$
- j is a cluster in N\_1

#### Value

SC3 stability index

#### See Also

The documentation for the calculate\_stability function in the SC3 package

### **Description**

Warn if node aesthetic names are incorrect

#### Usage

```
check_node_aes_list(node_aes_list)
```

### **Arguments**

node\_aes\_list List of node aesthetics

#### Value

Corrected node aesthetics list

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clustree

Plot a clustering tree

### **Description**

Creates a plot of a clustering tree showing the relationship between clusterings at different resolutions.

### Usage

```
clustree(x, ...)
## S3 method for class 'matrix'
clustree(x, prefix, suffix = NULL, metadata = NULL,
  count_filter = 0, prop_filter = 0.1, layout = c("tree",
  "sugiyama"), use_core_edges = TRUE, highlight_core = FALSE,
 node_colour = prefix, node_colour_aggr = NULL, node_size = "size",
 node_size_aggr = NULL, node_size_range = c(4, 15), node_alpha = 1,
  node_alpha_aggr = NULL, node_text_size = 3,
  scale_node_text = FALSE, node_text_colour = "black",
 node_label = NULL, node_label_aggr = NULL, node_label_size = 3,
  node_label_nudge = -0.2, edge_width = 1.5, edge_arrow = TRUE,
  edge_arrow_ends = c("last", "first", "both"), show_axis = FALSE,
 return = c("plot", "graph", "layout"), ...)
## S3 method for class 'data.frame'
clustree(x, prefix, ...)
## S3 method for class 'SingleCellExperiment'
clustree(x, prefix, exprs = "counts", ...)
## S3 method for class 'seurat'
clustree(x, prefix = "res.", exprs = c("data",
  "raw.data", "scale.data"), ...)
## S3 method for class 'Seurat'
clustree(x, prefix = paste0(assay, "_snn_res."),
  exprs = c("data", "counts", "scale.data"), assay = NULL, ...)
```

### Arguments

```
    x object containing clustering data
    ... extra parameters passed to other methods
    prefix string indicating columns containing clustering information
    suffix string at the end of column names containing clustering information
```

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metadata data.frame containing metadata on each sample that can be used as node aesthetics count threshold for filtering edges in the clustering graph count\_filter in proportion threshold for filtering edges in the clustering graph prop\_filter layout string specifying the "tree" or "sugiyama" layout, see igraph::layout\_as\_tree() and igraph::layout\_with\_sugiyama() for details use\_core\_edges logical, whether to only use core tree (edges with maximum in proportion for a node) when creating the graph layout, all (unfiltered) edges will still be displayed highlight\_core logical, whether to increase the edge width of the core network to make it easier to see node\_colour either a value indicating a colour to use for all nodes or the name of a metadata column to colour nodes by node\_colour\_aggr if node\_colour is a column name than a string giving the name of a function to aggregate that column for samples in each cluster node\_size either a numeric value giving the size of all nodes or the name of a metadata column to use for node sizes node\_size\_aggr if node\_size is a column name than a string giving the name of a function to aggregate that column for samples in each cluster node\_size\_range numeric vector of length two giving the maximum and minimum point size for plotting nodes either a numeric value giving the alpha of all nodes or the name of a metadata node\_alpha column to use for node transparency node\_alpha\_aggr if node\_aggr is a column name than a string giving the name of a function to aggregate that column for samples in each cluster node\_text\_size numeric value giving the size of node text if scale\_node\_text is FALSE scale\_node\_text logical indicating whether to scale node text along with the node size node\_text\_colour colour value for node text (and label) node\_label additional label to add to nodes node\_label\_aggr if node\_label is a column name than a string giving the name of a function to aggregate that column for samples in each cluster node\_label\_size numeric value giving the size of node label text node\_label\_nudge numeric value giving nudge in y direction for node labels

> numeric value giving the width of plotted edges logical indicating whether to add an arrow to edges

edge\_width

edge\_arrow

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edge\_arrow\_ends

string indicating which ends of the line to draw arrow heads if edge\_arrow is

TRUE, one of "last", "first", or "both"

show\_axis whether to show resolution axis

return string specifying what to return, either "plot" (a ggplot object), "graph" (a

tbl\_graph object) or "layout" (a ggraph layout object)

exprs source of gene expression information to use as node aesthetics, for SingleCellExperime

objects it must be a name in assayNames(x), for a seurat object it must be one of data, raw.data or scale.data and for a Seurat object it must be one of

data, counts or scale.data

assay name of assay to pull expression and clustering data from for Seurat objects

### **Details**

#### **Data sources**

Plotting a clustering tree requires information about which cluster each sample has been assigned to at different resolutions. This information can be supplied in various forms, as a matrix, data.frame or more specialised object. In all cases the object provided must contain numeric columns with the naming structure PXS where P is a prefix indicating that the column contains clustering information, X is a numeric value indicating the clustering resolution and S is any additional suffix to be removed. For SingleCellExperiment objects this information must be in the colData slot and for Seurat objects it must be in the meta.data slot. For all objects except matrices any additional columns can be used as aesthetics, for matrices an additional metadata data.frame can be supplied if required.

### **Filtering**

Edges in the graph can be filtered by adjusting the count\_filter and prop\_filter parameters. The count\_filter removes any edges that represent less than that number of samples, while the prop\_filter removes edges that represent less than that proportion of cells in the node it points towards.

#### Node aesthetics

The aesthetics of the plotted nodes can be controlled in various ways. By default the colour indicates the clustering resolution, the size indicates the number of samples in that cluster and the transparency is set to 100 Each of these can be set to a specific value or linked to a supplied metadata column. For a SingleCellExperiment or Seurat object the names of genes can also be used. If a metadata column is used than an aggregation function must also be supplied to combine the samples in each cluster. This function must take a vector of values and return a single value.

#### Layout

The clustering tree can be displayed using either the Reingold-Tilford tree layout algorithm or the Sugiyama layout algorithm for layered directed acyclic graphs. These layouts were selected as the are the algorithms available in the igraph package designed for trees. The Reingold-Tilford algorithm places children below their parents while the Sugiyama places nodes in layers while trying to minimise the number of crossing edges. See igraph::layout\_as\_tree() and igraph::layout\_with\_sugiyama() for more details. When use\_core\_edges is TRUE (default) only the core tree of the maximum in proportion edges for each node are used for constructing the layout. This can often lead to more attractive layouts where the core tree is more visible.

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#### Value

a ggplot object (default), a tbl\_graph object or a ggraph layout object depending on the value of return

### **Examples**

```
data(iris_clusts)
clustree(iris_clusts, prefix = "K")
```

clustree\_overlay

Overlay a clustering tree

## **Description**

Creates a plot of a clustering tree overlaid on a scatter plot of individual samples.

```
clustree_overlay(x, ...)
## S3 method for class 'matrix'
clustree_overlay(x, prefix, metadata, x_value, y_value,
  suffix = NULL, count_filter = 0, prop_filter = 0.1,
 node_colour = prefix, node_colour_aggr = NULL, node_size = "size",
 node_size_aggr = NULL, node_size_range = c(4, 15), node_alpha = 1,
 node_alpha_aggr = NULL, edge_width = 1, use_colour = c("edges",
  "points"), alt_colour = "black", point_size = 3, point_alpha = 0.2,
 point_shape = 18, label_nodes = FALSE, label_size = 3,
 plot_sides = FALSE, side_point_jitter = 0.45,
  side_point_offset = 1, ...)
## S3 method for class 'data.frame'
clustree_overlay(x, prefix, ...)
## S3 method for class 'SingleCellExperiment'
clustree_overlay(x, prefix, x_value,
 y_value, exprs = "counts", red_dim = NULL, ...)
## S3 method for class 'seurat'
clustree_overlay(x, x_value, y_value, prefix = "res.",
  exprs = c("data", "raw.data", "scale.data"), red_dim = NULL, ...)
## S3 method for class 'Seurat'
clustree_overlay(x, x_value, y_value,
 prefix = paste0(assay, "_snn_res."), exprs = c("data", "counts",
  "scale.data"), red_dim = NULL, assay = NULL, ...)
```

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### **Arguments**

object containing clustering data Х extra parameters passed to other methods prefix string indicating columns containing clustering information metadata data.frame containing metadata on each sample that can be used as node aesthetics x\_value numeric metadata column to use as the x axis y\_value numeric metadata column to use as the y axis suffix string at the end of column names containing clustering information count filter count threshold for filtering edges in the clustering graph prop\_filter in proportion threshold for filtering edges in the clustering graph node\_colour either a value indicating a colour to use for all nodes or the name of a metadata column to colour nodes by node\_colour\_aggr if node\_colour is a column name than a string giving the name of a function to aggregate that column for samples in each cluster node size either a numeric value giving the size of all nodes or the name of a metadata column to use for node sizes node\_size\_aggr if node\_size is a column name than a string giving the name of a function to aggregate that column for samples in each cluster node\_size\_range numeric vector of length two giving the maximum and minimum point size for plotting nodes node\_alpha either a numeric value giving the alpha of all nodes or the name of a metadata column to use for node transparency node\_alpha\_aggr if node\_aggr is a column name than a string giving the name of a function to aggregate that column for samples in each cluster edge\_width numeric value giving the width of plotted edges use colour one of "edges" or "points" specifying which element to apply the colour aesthetic to alt\_colour colour value to be used for edges or points (whichever is NOT given by use\_colour) point\_size numeric value giving the size of sample points point\_alpha numeric value giving the alpha of sample points numeric value giving the shape of sample points point\_shape label\_nodes logical value indicating whether to add labels to clustering graph nodes label\_size numeric value giving the size of node labels is label\_nodes is TRUE plot\_sides logical value indicating whether to produce side on plots side\_point\_jitter

numeric value giving the y-direction spread of points in side plots

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side\_point\_offset

numeric value giving the y-direction offset for points in side plots

exprs source of gene expression information to use as node aesthetics, for SingleCellExperime

objects it must be a name in assayNames(x), for a seurat object it must be one of data, raw.data or scale.data and for a Seurat object it must be one of

data, counts or scale.data

red\_dim dimensionality reduction to use as a source for x\_value and y\_value

assay name of assay to pull expression and clustering data from for Seurat objects

#### **Details**

#### **Data sources**

Plotting a clustering tree requires information about which cluster each sample has been assigned to at different resolutions. This information can be supplied in various forms, as a matrix, data.frame or more specialised object. In all cases the object provided must contain numeric columns with the naming structure PXS where P is a prefix indicating that the column contains clustering information, X is a numeric value indicating the clustering resolution and S is any additional suffix to be removed. For SingleCellExperiment objects this information must be in the colData slot and for Seurat objects it must be in the meta.data slot. For all objects except matrices any additional columns can be used as aesthetics.

## **Filtering**

Edges in the graph can be filtered by adjusting the count\_filter and prop\_filter parameters. The count\_filter removes any edges that represent less than that number of samples, while the prop\_filter removes edges that represent less than that proportion of cells in the node it points towards.

#### Node aesthetics

The aesthetics of the plotted nodes can be controlled in various ways. By default the colour indicates the clustering resolution, the size indicates the number of samples in that cluster and the transparency is set to 100 Each of these can be set to a specific value or linked to a supplied metadata column. For a SingleCellExperiment or Seurat object the names of genes can also be used. If a metadata column is used than an aggregation function must also be supplied to combine the samples in each cluster. This function must take a vector of values and return a single value.

#### Colour aesthetic

The colour aesthetic can be applied to either edges or sample points by setting use\_colour. If "edges" is selected edges will be coloured according to the clustering resolution they originate at. If "points" is selected they will be coloured according to the cluster they are assigned to at the highest resolution.

## **Dimensionality reductions**

For SingleCellExperiment and Seurat objects precomputed dimensionality reductions can be used for x or y aesthetics. To do so red\_dim must be set to the name of a dimensionality reduction in reducedDimNames(x) (for a SingleCellExperiment) or x@dr (for a Seurat object). x\_value and y\_value can then be set to red\_dimX when red\_dim matches the red\_dim argument and X is the column of the dimensionality reduction to use.

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#### Value

a ggplot object if plot\_sides is FALSE or a list of ggplot objects if plot\_sides is TRUE

## **Examples**

```
data(iris_clusts)
clustree_overlay(iris_clusts, prefix = "K", x_value = "PC1", y_value = "PC2")
```

get\_tree\_edges

Get tree edges

## **Description**

Extract the edges from a set of clusterings

## Usage

```
get_tree_edges(clusterings, prefix)
```

## **Arguments**

clusterings numeric matrix containing clustering information, each column contains clus-

tering at a separate resolution

prefix string indicating columns containing clustering information

### Value

data.frame containing edge information

get\_tree\_nodes

Get tree nodes

## **Description**

Extract the nodes from a set of clusterings and add relevant attributes

```
get_tree_nodes(clusterings, prefix, metadata, node_aes_list)
```

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### **Arguments**

clusterings numeric matrix containing clustering information, each column contains clus-

tering at a separate resolution

prefix string indicating columns containing clustering information

metadata data.frame containing metadata on each sample that can be used as node aes-

thetics

node\_aes\_list nested list containing node aesthetics

#### Value

data.frame containing node information

iris\_clusts

Clustered Iris dataset

## Description

Iris dataset clustered using k-means with a range of values of k

## **Usage**

```
iris_clusts
```

## **Format**

iris\_clusts is a data.frame containing the normal iris dataset with additional columns holding k-means clusterings at different values of k and the first two principal components

### Source

```
set.seed(1)
iris_mat <- as.matrix(iris[1:4])
iris_km <- sapply(1:5, function(x) {
    km <- kmeans(iris_mat, centers = x, iter.max = 100, nstart = 10)
    km$cluster
})
colnames(iris_km) <- paste0("K", 1:5)
iris_clusts <- cbind(iris, iris_km)
iris_pca <- prcomp(iris_clusts[1:4])
iris_clusts$PC1 <- iris_pca$x[, 1]
iris_clusts$PC2 <- iris_pca$x[, 2]</pre>
```

overlay\_node\_points 17

overlay\_node\_points
Overlay node points

## **Description**

Overlay clustering tree nodes on a scatter plot with the specified aesthetics.

## Usage

```
overlay_node_points(nodes, x_value, y_value, node_colour, node_size,
    node_alpha)
```

## Arguments

nodes	data.frame describing nodes
x_value	column of nodes to use for the x position
y_value	column of nodes to use for the y position
node_colour	either a value indicating a colour to use for all nodes or the name of a metadata column to colour nodes by
node_size	either a numeric value giving the size of all nodes or the name of a metadata column to use for node sizes
node_alpha	either a numeric value giving the alpha of all nodes or the name of a metadata column to use for node transparency

## **Description**

Plot the side view of a clustree overlay plot. If the ordinary plot shows the tree from above this plot shows it from the side, highlighting either the x or y dimension and the clustering resolution.

```
plot_overlay_side(nodes, edges, points, prefix, side_value, graph_attr,
  node_size_range, edge_width, use_colour, alt_colour, point_size,
  point_alpha, point_shape, label_nodes, label_size, y_jitter, y_offset)
```

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## **Arguments**

nodes	data.frame describing nodes
edges	data.frame describing edges
points	data.frame describing points
prefix	string indicating columns containing clustering information
side_value	string giving the metadata column to use for the x axis
graph_attr	list describing graph attributes
node_size_range	
	numeric vector of length two giving the maximum and minimum point size for plotting nodes
edge_width	numeric value giving the width of plotted edges
use_colour	one of "edges" or "points" specifying which element to apply the colour aesthetic to
alt_colour	colour value to be used for edges or points (whichever is NOT given by use_colour)
point_size	numeric value giving the size of sample points
point_alpha	numeric value giving the alpha of sample points
point_shape	numeric value giving the shape of sample points
label_nodes	logical value indicating whether to add labels to clustering graph nodes
label_size	numeric value giving the size of node labels is label_nodes is TRUE
$y_{-}$ jitter	numeric value giving the y-direction spread of points in side plots
y_offset	numeric value giving the y-direction offset for points in side plots

## Value

## RETURN\_DESCRIPTION

sc_example	Simulated scRNA-seq dataset	
------------	-----------------------------	--

## Description

A simulated scRNA-seq dataset generated using the splatter package and clustered using the SC3 and Seurat packages.

## Usage

sc\_example

### **Format**

 $sc\_example$  is a list holding a simulated scRNA-seq dataset. Items in the list included the simulated counts, normalised log counts, tSNE dimensionality reduction and cell assignments from SC3 and Seurat clustering.

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#### Source

```
# Simulation
library("splatter") # Version 1.2.1
sim <- splatSimulate(batchCells = 200, nGenes = 10000,</pre>
                      group.prob = c(0.4, 0.2, 0.2, 0.15, 0.05),
                      de.prob = c(0.1, 0.2, 0.05, 0.1, 0.05),
                      method = "groups", seed = 1)
sim_counts <- counts(sim)[1:1000, ]</pre>
# SC3 Clustering
library("SC3") # Version 1.7.6
library("scater") # Version 1.6.2
sim_sc3 <- SingleCellExperiment(assays = list(counts = sim_counts))</pre>
rowData(sim_sc3)$feature_symbol <- rownames(sim_counts)</pre>
sim_sc3 <- normalise(sim_sc3)</pre>
sim_sc3 \leftarrow sc3(sim_sc3, ks = 1:8, biology = FALSE, n_cores = 1)
sim_sc3 <- runTSNE(sim_sc3)</pre>
# Seurat Clustering
library("Seurat") # Version 2.2.0
sim_seurat <- CreateSeuratObject(sim_counts)</pre>
sim_seurat <- NormalizeData(sim_seurat, display.progress = FALSE)</pre>
sim_seurat <- FindVariableGenes(sim_seurat, do.plot = FALSE,</pre>
                                  display.progress = FALSE)
sim_seurat <- ScaleData(sim_seurat, display.progress = FALSE)</pre>
sim_seurat <- RunPCA(sim_seurat, do.print = FALSE)</pre>
sim_seurat <- FindClusters(sim_seurat, dims.use = 1:6,</pre>
                             resolution = seq(0, 1, 0.1),
                             print.output = FALSE)
sc_example <- list(counts = counts(sim_sc3),</pre>
                    tsne = reducedDim(sim_sc3),
                    sc3_clusters = colData(sim_sc3),
                    seurat_clusters = sim_seurat@meta.data)
```

store\_node\_aes

Store node aesthetics

### **Description**

Store the names of node attributes to use as aesthetics as graph attributes

20 store\_node\_aes

## Usage

```
store_node_aes(graph, node_aes_list, metadata)
```

## Arguments

graph graph to store attributes in

node\_aes\_list nested list containing node aesthetics

metadata data.frame containing metadata that can be used as aesthetics

## Value

graph with additional attributes

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