Package ‘slanter’

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meristems

Sample RNA data of similarity between batches of 1000 cells of tomato meristem cells.

Description

This is a simple matrix where each entry is the similarity (correlation) between a pair of batches. Negative correlations were changed to zero to simplify the analysis.

Usage

data(meristems)

Format

A simple square matrix.

Examples

data(meristems)
similarity <- meristems
similarity[similarity < 0] = 0
slanter::sheatmap(meristems, order_data=similarity, show_rownames=FALSE, show_colnames=FALSE)

oclust

Hierarchically cluster ordered data.

Description

Given a distance matrix for sorted objects, compute a hierarchical clustering preserving this order. That is, this is similar to hclust with the constraint that the result’s order is always 1:N.

Usage

oclust(distances, method = "ward.D2", order = NULL, members = NULL)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>distances</td>
<td>A distances object (as created by stats::dist).</td>
</tr>
<tr>
<td>method</td>
<td>The clustering method to use (only ward.D and ward.D2 are supported).</td>
</tr>
<tr>
<td>order</td>
<td>If specified, assume the data will be re-ordered by this order.</td>
</tr>
<tr>
<td>members</td>
<td>Optionally, the number of members for each row/column of the distances (by default, one each).</td>
</tr>
</tbody>
</table>
reorder_frame

Details

If an order is specified, assumes that the data will be re-ordered by this order. That is, the indices in the returned `hclust` object will refer to the post-reorder data locations, **not** to the current data locations.

This can be applied to the results of `slanted_reorder`, to give a "plausible" clustering for the data.

Value

A clustering object (as created by `hclust`).

Examples

```r
clusters <- slanter::oclust(dist(mtcars), order=1:dim(mtcars)[1])
clusters$order
```

reorder_frame

Reorder the rows of a frame.

Description

You’d expect `data[order,]` to "just work". It doesn’t for data frames with a single column, which happens for annotation data, hence the need for this function. Sigh.

Usage

```r
reorder_frame(frame, order)
```

Arguments

- `frame` A data frame to reorder the rows of.
- `order` An array containing indices permutation to apply to the rows.

Value

The data frame with the new row orders.

Examples

```r
df <- data.frame(foo=c(1, 2, 3))
df[c(1,3,2),]
slanter::reorder_frame(df, c(1,3,2))
```
reorder_hclust

Given a clustering of some data, and some ideal order we’d like to use to visualize it, reorder (but do not modify) the clustering to be as consistent as possible with this ideal order.

Description

Given a clustering of some data, and some ideal order we’d like to use to visualize it, reorder (but do not modify) the clustering to be as consistent as possible with this ideal order.

Usage

reorder_hclust(clusters, order)

Arguments

clusters The existing clustering of the data.
order The ideal order we’d like to see the data in.

Value

A reordered clustering which is consistent, wherever possible, the ideal order.

Examples

clusters <- hclust(dist(mtcars))
clusters$order
clusters <- slanter::reorder_hclust(clusters, 1:length(clusters$order))
clusters$order

sheatmap

Plot a heatmap with values as close to the diagonal as possible.

Description

Given a matrix expressing the cross-similarity between two (possibly different) sets of entities, this will reorder it to move the high values close to the diagonal, for a better visualization.
Usage

sheetmap(
  data,
  ...,  
  order_data = NULL,
  annotation_col = NULL,
  annotation_row = NULL,
  order_rows = TRUE,
  order_cols = TRUE,
  squared_order = TRUE,
  same_order = FALSE,
  patch_cols_order = NULL,
  patch_rows_order = NULL,
  discount_outliers = TRUE,
  cluster_rows = TRUE,
  cluster_cols = TRUE,
  oclust_rows = TRUE,
  oclust_cols = TRUE,
  clustering_distance_rows = "euclidian",
  clustering_distance_cols = "euclidian",
  clustering_method = "ward.D2",
  clustering_callback = NA
)

Arguments

data A rectangular matrix to plot, of non-negative values (unless order_data is specified).
...
Additional flags to pass to pheatmap.
order_data An optional matrix of non-negative values of the same size to use for computing the orders.
annotation_col Optional data frame describing each column.
annotation_row Optional data frame describing each row.
order_rows Whether to reorder the rows. Otherwise, use the current order.
order_cols Whether to reorder the columns. Otherwise, use the current order.
squared_order Whether to reorder to minimize the l2 norm (otherwise minimizes the l1 norm).
same_order Whether to apply the same order to both rows and columns (if reordering both). For a square matrix, may also contain 'row' or 'column' to force the order of one axis to apply to both.
patch_cols_order Optional function that may be applied to the columns order, returning a better order.
patch_rows_order Optional function that may be applied to the rows order, returning a better order.
discount_outliers
    Whether to do a final order phase discounting outlier values far from the diagonal.
cluster_rows
    Whether to cluster the rows, or the clustering to use.
cluster_cols
    Whether to cluster the columns, or the clustering to use.
oclust_rows
    Whether to use oclust instead of hclust for the rows (if clustering them).
oclust_cols
    Whether to use oclust instead of hclust for the columns (if clustering them).
clustering_distance_rows
    The default method for computing row distances (by default, euclidian).
clustering_distance_cols
    The default method for computing column distances (by default, euclidian).
clustering_method
    The default method to use for hierarchical clustering (by default, ward.D2 and *not* complete).
clustering_callback
    Is not supported.

Details

If you have an a-priori order for the rows and/or columns, you can prevent reordering either or both by specifying order_rows=FALSE and/or order_cols=FALSE. Otherwise, slanted_orders is used to compute the "ideal" slanted order for the data.

By default, the rows and columns are ordered independently from each other. If the matrix is asymmetric but square (e.g., a matrix of weights of a directed graph such as a K-nearest-neighbors graph), then you can can specify same_order=TRUE to force both rows and columns to the same order. You can also specify same_order='row' to force the columns to use the same order as the rows, or same_order='column' to force the rows to use the same order as the columns.

You can also specify a patch_cols_order and/or a ‘patch_rows_order‘ function that takes the computed "ideal" order and returns a patched order. For example, this can be used to force special values (such as "outliers") to the side of the heatmap.

There are four options for controlling clustering:

* By default, sheatmap will generate a clustering tree using oclust, to generate the "best" clustering that is also compatible with the slanted order.

* Request that sheatmap will use the same hclust as pheatmap (e.g., oclust_rows=FALSE). In this case, the tree is reordered to be the "most compatible" with the target slanted order. That is, sheatmap will invoke reorder_hclust so that, for each node of the tree, the order of the two sub-trees will be chosen to best match the target slanted order. The end result need not be identical to the slanted order, but is as close as possible given the hclust clustering tree.

* Specify an explicit clustering (e.g., cluster_rows=hclust(...)). In this case, sheatmap will again merely reorder the tree but will not modify it.

In addition, you can give this function any of the pheatmap flags, and it will just pass them on. This allows full control over the diagram's features.

Note that clustering_callback is not supported. In addition, the default clustering_method here is ward.D2 instead of complete, since the only methods supported by oclust are ward.D and ward.D2.
Value

Whatever pheatmap returns.

Examples

slanter::sheatmap(cor(t(mtcars)))
slanter::sheatmap(cor(t(mtcars)), oclust_rows=FALSE, oclust_cols=FALSE)
pheatmap::pheatmap(cor(t(mtcars)))

Description

For a matrix expressing the cross-similarity between two (possibly different) sets of entities, this produces better results than clustering (e.g. as done by pheatmap). This is because clustering does not care about the order of each two sub-partitions. That is, clustering is as happy with ((2,1),(4,3)) as it is with the more sensible ((1,2),(3,4)). As a result, visualizations of similarities using naive clustering can be misleading.

Usage

slanted_orders(
  data,
  order_rows = TRUE,
  order_cols = TRUE,
  squared_order = TRUE,
  same_order = FALSE,
  discount_outliers = TRUE,
  max_spin_count = 10
)

Arguments

data A rectangular matrix containing non-negative values.
order_rows Whether to reorder the rows.
order_cols Whether to reorder the columns.
squared_order Whether to reorder to minimize the l2 norm (otherwise minimizes the l1 norm).
same_order Whether to apply the same order to both rows and columns.
discount_outliers Whether to do a final order phase discounting outlier values far from the diagonal.
max_spin_count How many times to retry improving the solution before giving up.
Value

A list with two keys, rows and cols, which contain the order.

Examples

```r
slanter::slanted_orders(cor(t(mtcars)))
```

---

**slanted_reorder**  
Reorder data rows and columns to move high values close to the diagonal.

---

Description

Given a matrix expressing the cross-similarity between two (possibly different) sets of entities, this uses slanted_orders to compute the "best" order for visualizing the matrix, then returns the reordered data. Commonly used in pheatmap(slanted_reorder(data),...), and of course heatmap does this internally for you.

Usage

```r
slanted_reorder(
  data,
  order_data = NULL,
  order_rows = TRUE,
  order_cols = TRUE,
  squared_order = TRUE,
  same_order = FALSE,
  discount_outliers = TRUE
)
```

Arguments

data  A rectangular matrix to reorder, of non-negative values (unless order_data is specified).

order_data  An optional matrix of non-negative values of the same size to use for computing the orders.

order_rows  Whether to reorder the rows.

order_cols  Whether to reorder the columns.

squared_order  Whether to reorder to minimize the l2 norm (otherwise minimizes the l1 norm).

same_order  Whether to apply the same order to both rows and columns.

discount_outliers  Whether to do a final order phase discounting outlier values far from the diagonal.
**slanted_reorder**

**Value**

A matrix of the same shape whose rows and columns are a permutation of the input.

**Examples**

```r
slanter::slanted_reorder(cor(t(mtcars)))
```
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