Package ‘ksharp’

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Title Cluster Sharpening
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Description Clustering typically assigns data points into discrete groups, but the clusters can sometimes be indistinct. Cluster sharpening adjusts an existing clustering to create contrast between groups. This package provides a general interface for cluster sharpening along with several implementations based on different excision criteria.
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**kdata.1**

*Toy dataset with two convex groups with partial overlap*

**Description**

Toy dataset with two convex groups with partial overlap

**Usage**

`data(kdata.1)`

**Format**

Matrix with two columns: D1, D2

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**kdata.2**

*Toy dataset with two non-overlapping and non-spherical groups*

**Description**

Toy dataset with two non-overlapping and non-spherical groups

**Usage**

`data(kdata.2)`

**Format**

Matrix with two columns: D1, D2
**kdata.3**  
*Toy dataset with three groups*

**Description**
Toy dataset with three groups

**Usage**
data(kdata.3)

**Format**
matrix with two columns: D1, D2

---

**kdata.4**  
*Toy dataset with four groups atop a wide area of noise points*

**Description**
Toy dataset with four groups atop a wide area of noise points

**Usage**
data(kdata.4)

**Format**
matrix with two columns: D1, D2

---

**ksharp**  
*Sharpen a clustering*

**Description**
Each data point in a clustering is assigned to a cluster, but some data points may lie in ambiguous zones between two or more clusters, or far from other points. Cluster sharpening assigns these border points into a separate noise group, thereby creating more stark distinctions between groups.
Usage

ksharp(
  x,
  threshold = 0.1,
  data = NULL,
  method = c("silhouette", "neighbor", "medoid"),
  threshold.abs = NULL
)

Arguments

x           clustering object; several types of inputs are acceptable, including objects of
            class kmeans, pam, and self-made lists with a component "cluster".
threshold    numeric; the fraction of points to place in noise group
data         matrix, raw data corresponding to clustering x; must be present when sharpening
             for the first time or if data is not present within x.
method       character, determines method used for sharpening
threshold.abs numeric; absolute-value of threshold for sharpening. When non-NULL, this
             value overrides value in argument 'threshold'

Details

Noise points are assigned to a group with cluster index 0. This is analogous behavior to output
produced by dbscan.

Value

clustering object based on input x, with adjusted cluster assignments and additional list components
with sharpness measures. Cluster assignments are placed in $cluster and excised data points are
given a cluster index of 0. Original cluster assignments are saved in $cluster.original. Sharpness
measures are stored in components $silinfo, $medinfo, and $neiinfo, although these details may
change in future versions of the package.

Examples

# prepare iris dataset for analysis
iris.data = iris[, 1:4]
rownames(iris.data) = paste0("iris_", seq_len(nrow(iris.data)))

# cluster the dataset into three groups
iris.clustered = kmeans(iris.data, centers=3)
table(iris.clustered$cluster)

# sharpen the clustering by excluding 10% of the data points
iris.sharp = ksharp(iris.clustered, threshold=0.1, data=iris.data)
table(iris.sharp$cluster)

# visualize cluster assignments
medinfo

iris.pca = prcomp(iris.data)$x[,1:2]
plot(iris.pca, col=iris$Species, pch=ifelse(iris.sharp$cluster==0, 1, 19))

Description

Analogous in structure to silinfo and neiinfo, it computes a "widths" matrix assessing how well each data point belongs to its cluster. Here, this measure is the ratio of two distances: in the numerator, the distance from the point to the nearest cluster center, and in the denominator, from the point to its own cluster center.

Usage

medinfo(cluster, data, silwidths)

Arguments

cluster named vector
data matrix with raw data
silwidths matrix with silhouette widths

Value

list with component widths. The widths object is a matrix with one row per data item, with column med_ratio holding the sharpness measure.

Examples

# construct a manual clustering of the iris dataset
iris.data = iris[, 1:4]
rownames(iris.data) = paste0("iris_", seq_len(nrow(iris.data)))
iris.dist = dist(iris.data)
iris.clusters = setNames(as.integer(iris$Species), rownames(iris.data))

# compute sharpness values based on medoids
iris.silinfo = silinfo(iris.clusters, iris.dist)
medinfo(iris.clusters, iris.data, iris.silinfo$widths)
neiinfo  
Compute info on ‘neighbor widths’

Description
This function provides information on how well each data point belongs to its cluster. For each query point, the function considers n of its nearest neighbors. The neighbor widths are defined as the fraction of those neighbors that belong to the same cluster as the query point. These values are termed ‘widths’ in analogy to silhouette widths, another measure of cluster membership.

Usage
neiinfo(cluster, dist)

Arguments
- cluster: vector with assignments of data elements to clusters
- dist: distance object or matrix

Details
The function follows a similar signature as silinfo from this package.

Value
list with component widths. The widths object is a matrix with one row per data item, with column neighborhood holding the sharpness value.

Examples

```r
# construct a manual clustering of the iris dataset
iris.data = iris[, 1:4]
rownames(iris.data) = paste0("iris_", seq_len(nrow(iris)))
iris.dist = dist(iris.data)
iris.clusters = setNames(as.integer(iris$Species), rownames(iris.data))

# compute neighbor-based sharpness widths
neiinfo(iris.clusters, iris.dist)
```
silinfo

Compute info on silhouette widths

Description

This function provides information on how well each data point belongs to its cluster. For each
query point, the function considers the average distance to other members of the same cluster and
the average distance to members of another, nearest, cluster. The widths are defined as the

Usage

silinfo(cluster, dist)

Arguments

cluster vector with assignments of data elements to clusters
dist distance object or matrix

Details

The function signature is very similar to cluster::silhouette but the implementation has important
differences. This implementation requires both the dist object and and cluster vector must have
names. This prevents accidental assignment of silhouette widths to the wrong elements.

Value

list, analogous to object within output from cluster::pam. In particular, the list has a component
widths. The widths object is matrix with one row per data item, with column sil_width holding the
silhouette width.

Examples

# construct a manual clustering of the iris dataset
iris.data = iris[, 1:4]
rownames(iris.data) = paste0("iris_", seq_len(nrow(iris.data)))
iris.dist = dist(iris.data)
iris.clusters = setNames(as.integer(iris$Species), rownames(iris.data))

# compute sharpness values based on silhouette widths
silinfo(iris.clusters, iris.dist)
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