Package 'WeightedCluster'

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Title Clustering of Weighted Data

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Suggests RUnit, knitr, isotone, vegan, lattice
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VignetteBuilder knitr
Description Clusters state sequences and weighted data. It provides an optimized weighted PAM algorithm as well as functions for aggregating replicated cases, computing cluster quality measures for a range of clustering solutions and plotting clusters of state sequences.
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as.clustrange	Build a clustrange object to compare different clustering solutions.
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Description

Build a clustrange object to compare different clustering solutions.

Usage

Arguments

object	The object to convert such as a data.frame.
diss	A dissimilarity matrix or a dist object (see dist).
weights	Optional numerical vector containing weights.
R	Optional number of bootstrap that can be used to build confidence intervals.
samplesize	Size of bootstrap sample. Default to sum of weights.
ncluster	Integer. Maximum number of cluster. The range will include all clustering solution starting from two to ncluster.
x	A clustrange object to be plotted.
stat	Character. The list of statistics to plot or "noCH" to plot all statistics except "CH" and "CHsq" or "all" for all statistics. See wcClusterQuality for a list of possible values. It is also possible to use "RHC" to plot the quality measure 1-HC. Unlike HC, RHC should be maximized as all other quality measures.
legendpos	Character. legend position, see legend.
norm	Character. Normalization method of the statistics can be one of "none" (no normalization), "range" (given as (value -min)/(max-min), "zscore" (adjusted by mean and standard deviation) or "zscoremed" (adjusted by median and median of the difference to the median).
withlegend	Logical. If FALSE, the legend is not plotted.
lwd	Numeric. Line width, see par.
col	A vector of line colors, see par. If NULL, a default set of color is used.
xlab	x axis label.
ylab	y axis label.

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conf.int	Confidence to build the confidence interval (default: 0.9).
ci.method	Method used to build the confidence interval (only if bootstrap has been used, see R above). One of "none" (do not plot confidence interval), "norm" (based on normal approximation), "perc" (based on percentile).)
ci.alpha	alpha color value used to plot the interval.
line	Which value should be plotted by the line? One of "t0" (value for actual sample), "mean" (average over all bootstraps), "median" (median over all bootstraps).
	Additionnal parameters passed to/from methods.

Details

as.clustrange convert objects to clustrange objects. clustrange objects contains a list of clustering solution with associated statistics and can be used to find the optimal clustering solution.

If object is a data. frame or a matrix, each column should be a clustering solution to be evaluated.

If object is an helust or twins objects (i.e. hierarchical clustering output, see helust, diana or agnes), the function compute all clustering solution ranging from two to neluster and compute the associated statistics.

Value

An object of class clustrange with the following elements:

```
clustering: A data.frame of all clustering solutions.
stats: A matrix containing the clustering statistics of each cluster solution.
```

See Also

See also wcKMedRange, wcClusterQuality.

Examples

```
data(mvad)
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)

## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)

## COmpute distance using Hamming distance
diss <- seqdist(mvad.seq, method="HAM")

## Ward clustering
wardCluster <- hclust(as.dist(diss), method="ward", members=aggMvad$aggWeights)

## Computing clustrange from Ward clustering
wardRange <- as.clustrange(wardCluster, diss=diss,
weights=aggMvad$aggWeights, ncluster=15)

## Plot all statistics (standardized)</pre>
```

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```
plot(wardRange, stat="all", norm="zscoremed", lwd=3)
## Plot HC, RHC and ASW
plot(wardRange, stat=c("HC", "RHC", "ASWw"), norm="zscore", lwd=3)
```

as.seqtree

Convert a hierarchical clustering object to a seqtree object.

Description

Convert a hierarchical clustering object to a seqtree object which can then be displayed using seqtreedisplay.

Usage

```
as.seqtree(object, seqdata, diss, weighted=TRUE, ...)
## S3 method for class 'twins'
as.seqtree(object, seqdata, diss, weighted=TRUE, ncluster, ...)
## S3 method for class 'hclust'
as.seqtree(object, seqdata, diss, weighted=TRUE, ncluster, ...)
```

Arguments

object An object to be converted to a seqtree.

segdata State sequence object.

diss A dissimilarity matrix or a dist object (see dist)

weighted Logical. If TRUE, weights of the seqdata object are taken to build the tree.

ncluster Maximum number of cluster. The tree will be builded until this number of clus-

ter.

... Additionnal parameters passed to/from methods.

Details

By default as. seqtree try to convert the object to a data. frame assuming that it contains a list of nested clustering solutions. Be aware that seqtree and as. seqtree only support binary splits.

If object is an helust or twins objects (i.e. hierarchical clustering output, see helust, diana or agnes), the function returns a seqtree object reproducing the agglomerative schedulde.

Value

A seqtree object.

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Examples

```
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)</pre>
## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)</pre>
## COmpute distance using Hamming distance
diss <- seqdist(mvad.seq, method="HAM")</pre>
## Ward clustering
wardCluster <- hclust(as.dist(diss), method="ward", members=aggMvad$weight)</pre>
st <- as.seqtree(wardCluster, seqdata=mvad.seq, diss=diss, weighted=TRUE, ncluster=10)
print(st)
## You typically want to run (You need to install GraphViz before)
## seqtreedisplay(st, type="d", border=NA)
```

seqclustname

Automatic labeling of cluster using sequence medoids

Description

This function automatically name the cluster using the sequence medoid of each cluster.

Usage

```
seqclustname(seqdata, group, diss, weighted = TRUE, perc = FALSE)
```

Arguments

seqdata State sequence object (see seqdef). A vector of clustering membership. group diss a dissimilarity matrix or a dist object.

Logical. If TRUE, weights of the sequata object are taken to find the medoids. weighted perc

Logical. If TRUE, the percentage of sequences in each cluster is added to the

label of each group.

Value

A factor of clustering membership. The labels are defined using sequences medoids and optionnaly percentage of case in each cluster.

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Examples

```
data(mvad)
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)

## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)

## Computing Hamming distance between sequence
diss <- seqdist(mvad.seq, method="HAM")

## KMedoids using PAMonce method (clustering only)
clust5 <- wcKMedoids(diss, k=5, weights=aggMvad$aggWeights)

clust5.labels <- seqclustname(mvad.seq, clust5$clustering, diss=diss, perc=TRUE)
seqdplot(mvad.seq, group=clust5.labels)</pre>
```

wcAggregateCases

Aggregate identical cases.

Description

Function to aggregate identical cases.

Usage

```
wcAggregateCases(x, weights = NULL, ...)
## S3 method for class 'data.frame'
wcAggregateCases(x, weights=NULL, ...)
## S3 method for class 'matrix'
wcAggregateCases(x, weights=NULL, ...)
## S3 method for class 'wcAggregateCases'
print(x, ...)
```

Arguments

x The object to aggregate.

weights Numeric. An optional case weights vector.

. . . Optional additionnal arguments.

Value

A wcAggregateCases object with the following components:

aggIndex Index of the unique cases in the original object data.

aggWeights Aggregated case weights

disaggIndex Index of the original object data in the unique cases.

disaggWeights Original weights used.

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Examples

```
data(mvad)
## Taking only the father unemployment and
## success at the end of compulsory schooling.
myData <- mvad[ , c("funemp", "gcse5eq")]
## Computing aggregated cases informations
ac <- wcAggregateCases(myData, weights=mvad$weight)
print(ac)
## Retrieving unique cases in the original data set
uniqueData <- myData[ac$aggIndex, ]
## Table from original data
table.orig <- xtabs(mvad$weight~funemp+gcse5eq, data=myData)
## Table from aggregated data
table.agg <- xtabs(ac$aggWeights~funemp+gcse5eq, data=uniqueData)
## Both table are equal, no information is lost
## (only the call command is different)
all(table.orig == table.agg)</pre>
```

wcClusterQuality

Cluster quality statistics

Description

Compute several quality statistics of a given clustering solution.

Usage

```
wcClusterQuality(diss, clustering, weights = NULL)
```

Arguments

diss A dissimilarity matrix or a dist object (see dist)

clustering Factor. A vector of clustering membership.
weights optional numerical vector containing weights.

Details

Compute several quality statistics of a given clustering solution. See value for details.

Value

```
A list with two elements stats and ASW:
```

stats with the following statistics:

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PBC Point Biserial Correlation. Correlation between the given distance matrice and a distance which equal to zero for individuals in the same cluster and one otherwise.

HG Hubert's Gamma. Same as previous but using Kendall's Gamma coefficient.

HGSD Hubert's Gamma (Somers'D). Same as previous but using Somers' D coefficient.

ASW Average Silhouette width (observation).

ASWw Average Silhouette width (weighted).

CH Calinski-Harabasz index (Pseudo F statistics computed from distances).

R2 Share of the discrepancy explained by the clustering solution.

CHsq Calinski-Harabasz index (Pseudo F statistics computed from *squared* distances).

R2sq Share of the discrepancy explained by the clustering solution (computed using *squared* distances).

HC Hubert's C coefficient.

ASW: The Average Silhouette Width of each cluster, one column for each ASW measure.

Examples

```
data(mvad)
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)

## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)
## Computing Hamming distance between sequence
diss <- seqdist(mvad.seq, method="HAM")

## KMedoids using PAMonce method (clustering only)
clust5 <- wcKMedoids(diss, k=5, weights=aggMvad$aggWeights, cluster.only=TRUE)

## Compute the silhouette of each observation
qual <- wcClusterQuality(diss, clust5, weights=aggMvad$aggWeights)

print(qual)</pre>
```

wcCmpCluster

Automatic comparison of clustering methods.

Description

Automatically compute different clustering solutions and associated quality measures to help identifying the best one.

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Usage

Arguments

diss A dissimilarity matrix or a dist object (see dist).

weights Optional numerical vector containing weights.

maxcluster Integer. Maximum number of cluster. The range will include all clustering

solution starting from two to ncluster.

method A vector of hierarchical clustering methods to compute or "all" for all meth-

ods. Possible values include "ward", "single", "complete", "average", "mcquitty", "median", "centroid" (using hclust), "pam" (using wcKMedRange), "diana" (only for unweighted datasets using diana), "beta.flexible" (only for un-

weighted datasets using agnes)

pam.combine Logical. Should we try all combinations of hierarchical and PAM clustering?

x A clustrangefamily object to plot or print object A clustrangefamily object to summarize

max.rank Integer. The different number of solution to print/summarize

group One of "stat" or "method". If "stat", plots are grouped by statistics, other-

wise by clustering methods.

stat Character. The list of statistics to plot or "noCH" to plot all statistics except

"CH" and "CHsq" or "all" for all statistics. See wcClusterQuality for a list of possible values. It is also possible to use "RHC" to plot the quality measure 1-HC. Unlike HC, RHC should be maximized as all other quality measures.

norm Character. Normalization method of the statistics can be one of "none" (no

normalization), "range" (given as (value -min)/(max-min), "zscore" (adjusted by mean and standard deviation) or "zscoremed" (adjusted by median and median

of the difference to the median).

withlegend Logical. If FALSE, the legend is not plotted.

lwd Numeric. Line width, see par.

col A vector of line colors, see par. If NULL, a default set of color is used.

legend.prop When withlegend=TRUE, sets the proportion of the graphic area used for plot-

ting the legend. Default value is set according to the place (bottom or right of

the graphic area) where the legend is plotted. Values from 0 to 1.

rows, cols optional arguments to arrange plots.

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```
x lab
y axis label.
main
main title of the plot.
...
Additionnal parameters passed to lines.
```

Value

An object of class clustrangefamily with the following elements:

Method name: the results of as.clustrange objects under each method name (see argument method for a list of possible values)

allstats: A matrix containing the clustering statistics for each cluster solution and method.

param: The parameters set when the function was called.

See Also

```
See Also as.clustrange
```

Examples

wcKMedoids

K-Medoids or PAM clustering of weighted data.

Description

K-Medoids or PAM clustering of weighted data.

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Usage

```
wcKMedoids(diss, k, weights=NULL, npass = 1, initialclust=NULL,
method="PAMonce", cluster.only = FALSE, debuglevel=0)
```

Arguments

diss A dissimilarity matrix or a dist object (see dist).

k Integer. The number of cluster.

weights Numeric. Optional numerical vector containing case weights.

npass Integer. Number of random start solution to test.

initialclust An integer vector, a factor, an "helust" or a "twins" object. Can be either the

index of the initial medoids (length should equal to k) or a vector specifying an initial clustering solution (length should then be equal to the number of observation.). If initialclust is an "hclust" or a "twins" object, then the initial

clustering solution is taken from the hierarchical clustering in k groups.

method Character. One of "KMedoids", "PAM" or "PAMonce" (default). See details.

cluster.only Logical. If FALSE, the quality of the retained solution is computed.

debuglevel Integer. If greater than zero, print some debugging messages.

Details

K-Medoids algorithms aim at finding the best partition of the data in a k predefined number of groups. Based on a dissimilarity matrix, those algorithms seeks to minimize the (weighted) sum of distance to the medoid of each group. The medoid is defined as the observation that minimize the sum of distance to the other observations of this group. The function wcKMedoids support three differents algorithms specified using the method argument:

- "KMedoids" Start with a random solution and then iteratively adapt the medoids using an algorithm similar to kmeans. Part of the code is inspired (but completely rewritten) by the C clustering library (see de Hoon et al. 2010). If you use this solution, you should set npass>1 to try several solution.
- "PAM" See pam in the cluster library. This code is based on the one available in the cluster library (Maechler et al. 2011). The advantage over the previous method is that it try to minimize a global criteria instead of a local one.
- "PAMonce" Same as previous but with two optimizations. First, the optimization presented by Reynolds et al. 2006. Second, only evaluate possible swap if the dissimilarity is greater than zero. This algorithm is used by default.

wcKMedoids works differently according to the diss argument. It may be faster using a matrix but require more memory (since all distances are stored twice). All combination between method and diss argument are possible, except for the "PAM" algorithm were only distance matrix may be used (use the "PAMonce" algorithm instead).

Value

An integer vector with the index of the medoids associated with each observation.

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References

Maechler, M., P. Rousseeuw, A. Struyf, M. Hubert and K. Hornik (2011). cluster: Cluster Analysis Basics and Extensions. R package version 1.14.1 — For new features, see the 'Changelog' file (in the package source).

Hoon, M. d.; Imoto, S. & Miyano, S. (2010). The C Clustering Library. Manual

See Also

pam in the cluster library, wcClusterQuality, wcKMedRange.

Examples

```
data(mvad)
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)</pre>
## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)</pre>
## Computing Hamming distance between sequence
diss <- segdist(mvad.seg, method="HAM")</pre>
## K-Medoids
clust5 <- wcKMedoids(diss, k=5, weights=aggMvad$aggWeights)</pre>
## clust5$clustering contains index number of each medoids
## Those medoids are
unique(clust5$clustering)
## Print the medoids sequences
print(mvad.seq[unique(clust5$clustering), ], informat="SPS")
## Some info about the clustering
print(clust5)
## Plot sequences according to clustering solution.
segdplot(mvad.seq, group=clust5$clustering)
```

wcKMedRange

Compute wcKMedoids clustering for different number of clusters.

Description

Compute wcKMedoids clustering for different number of clusters.

Usage

```
wcKMedRange(diss, kvals, weights=NULL, R=1, samplesize=NULL, ...)
```

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Arguments

diss	A dissimilarity matrix or a dist object (see dist).
kvals	A numeric vector containing the number of cluster to compute.
weights	Numeric. Optional numerical vector containing case weights.
R	Optional number of bootstrap that can be used to build confidence intervals.
samplesize	Size of bootstrap sample. Default to sum of weights.
	Additionnal parameters passed to wcKMedoids.

Details

Compute a clustrange object using the wcKMedoids method. clustrange objects contains a list of clustering solution with associated statistics and can be used to find the optimal clustering solution.

See as.clustrange for more details.

See Also

```
See as.clustrange.
```

Examples

```
data(mvad)
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)

## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)

## Compute distance using Hamming distance
diss <- seqdist(mvad.seq, method="HAM")

## Pam clustering
pamRange <- wcKMedRange(diss, 2:15)

## Plot all statistics (standardized)
plot(pamRange, stat="all", norm="zscoremed", lwd=3)

## Plotting sequences in 3 groups
seqdplot(mvad.seq, group=pamRange$clustering$cluster3)</pre>
```

wcSilhouetteObs

Compute the silhouette of each object using weighted data.

Description

Compute the silhouette of each object using weighted data.

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Usage

```
wcSilhouetteObs(diss, clustering, weights = NULL, measure="ASW")
```

Arguments

diss A dissimilarity matrix or a dist object (see dist)

clustering Factor. A vector of clustering membership.
weights optional numerical vector containing weights.

measure "ASW" or "ASWw", the measure of the silhouette. See the WeigthedCluster

vignettes.

Details

See the silhouette function in the cluster package for a detailed explanation of the silhouette.

Value

A numeric vector containing the silhouette of each observation.

References

Maechler, M., P. Rousseeuw, A. Struyf, M. Hubert and K. Hornik (2011). cluster: Cluster Analysis Basics and Extensions. R package version 1.14.1 — For new features, see the 'Changelog' file (in the package source).

See Also

See also silhouette.

Examples

```
## Aggregating state sequence
aggMvad <- wcAggregateCases(mvad[, 17:86], weights=mvad$weight)

## Creating state sequence object
mvad.seq <- seqdef(mvad[aggMvad$aggIndex, 17:86], weights=aggMvad$aggWeights)

## Computing Hamming distance between sequence
diss <- seqdist(mvad.seq, method="HAM")

## KMedoids using PAMonce method (clustering only)
clust5 <- wcKMedoids(diss, k=5, weights=aggMvad$aggWeights, cluster.only=TRUE)

## Compute the silhouette of each observation
sil <- wcSilhouetteObs(diss, clust5, weights=aggMvad$aggWeights, measure="ASWw")

## If you want to compute the average silhouette width,
## you should take weights into account
weighted.mean(sil, w=aggMvad$aggWeights)</pre>
```

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```
## Plotting sequences ordred by silhouette width,
## best classified are draw on the top.
seqIplot(mvad.seq, group=clust5, sortv=sil)
```

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