Package ‘stream’

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animation ............................ Animates the plotting of a DSD and the clustering process

Description

Generates an animation of a data stream or a data stream clustering. **Note:** You need to install package animation, and, if necessary, the libraries required for package magick.

Usage

animate_data(dsd, horizon = 100, n = 1000, wait = .1, plot.args = NULL, ...)
animate_cluster(dsc, dsd, measure = NULL, horizon = 100, n = 1000,
  type=c("auto", "micro", "macro"), assign="micro",
  assignmentMethod=c("auto","model", "nn"),
  noise = c("class", "exclude"),
  wait=.1, plot.args = NULL, ...)

Arguments

dsd ............... a DSD object
dsc .......... a DSC object
horizon ............... the number of points displayed at once/used for evaluation.
n ............... the number of points to be plotted
measure ............... the evaluation measure that should be graphed below the animation
type ............... evaluate "micro" or "macro"-clusters? "auto" chooses micro if dsc is of class DSC_micro and no macro is given. Otherwise macro is used.
assign ............... assign new points to the closest "micro" or "macro"-cluster to calculate the evaluation measure.
assignmentMethod ............... how to assign data points to micro-clusters. Options are "model" and "nn" (nearest neighbor). "auto" uses model if available and nn otherwise.
noise ............... how to handle noise for calculating the evaluation measure (as a separate class or excluded).
wait ............... the time interval between each frame
plot.args ............... a list with plotting parameters for the clusters.
... ............... extra arguments are added to plot.args.
Details

Animations are recorded using the library animation and can be replayed (which gives a smoother experience since there is no more computation done) and saved in various formats (see Examples section below).

Author(s)

Michael Hahsler

See Also

evaluate_cluster for stream evaluation without animation. See ani.replay for replaying and saving animations.

Examples

```r
## Not run:
stream <- DSD_Benchmark(1)
animate_data(stream, horizon=100, n=5000, xlim=c(0,1), ylim=c(0,1))

### animations can be replayed with the animation package
library(animation)
animation::ani.options(interval=.1) ## change speed
ani.replay()

### animations can also be saved as HTML, animated gifs, etc.
saveHTML(ani.replay())

### animate the clustering process with evaluation
### Note: we choose to exclude noise points from the evaluation
### measure calculation, even if the algorithm would assign
### them to a cluster.
reset_stream(stream)
dbstream <- DSC_DBSTREAM(r=.04, lambda=.1, gaptime=100, Cm=3,
shared_density=TRUE, alpha=.2)
animate_cluster(dbstream, stream, horizon=100, n=5000,
measure="crand", type="macro", assign="micro", noise = "exclude",
plot.args = list(xlim=c(0,1), ylim=c(0,1), shared = TRUE))

## End(Not run)
```

DSC  
---

Data Stream Clusterer Base Classes

Description

Abstract base classes for all DSC (Data Stream Clusterer) and DSC_R classes.
The DSC and DSC_R classes cannot be instantiated (calling DSC() or DSC_R() produces errors), but they serve as a base class from which other DSC classes inherit.

Class DSC provides several generic functions that can operate on all DSC subclasses: print(), plot(), nclusters() to get the current number of clusters, get_centers() to get the cluster centers, and get_weights() to get the cluster weights (if implemented). get_centers() and get_weights() are typically overwritten by subclasses of DSC. DSC_R provides these functions for R-based DSC implementations.

Since DSC objects often contain external pointers, regular saving and reading operations will fail. Use saveDSC() and readDSC() which will serialize the objects first appropriately.

Author(s)

Michael Hahsler

See Also

animate_cluster, cluster, evaluate, get_assignment, get_centers, get_weights, get_copy, microToMacro, nclusters, plot, prune_clusters, recluster, readDSC, saveDSC
Description

BICO maintains a tree which is inspired by the clustering tree of BIRCH, a SIGMOD Test of Time award-winning clustering algorithm. Each node in the tree represents a subset of these points. Instead of storing all points as individual objects, only the number of points, the sum and the squared sum of the subset’s points are stored as key features of each subset. Points are inserted into exactly one node.

Usage

```
DSC_BICO(k = 5, space = 10, p = 10, iterations = 10)
```

Arguments

- `k`: number of centres
- `space`: coreset size
- `p`: number of random projections used for nearest neighbour search in first level
- `iterations`: number of repetitions for the kmeans++ procedure in the offline component

Details

In this implementation, the nearest neighbour search on the first level of the tree is sped up by projecting all points to random 1-d subspaces. The first estimation of the optimal clustering cost is computed in a buffer phase at the beginning of the algorithm. This implementation interfaces the original C++ implementation available here: [http://ls2-www.cs.tu-dortmund.de/grav/de/bico](http://ls2-www.cs.tu-dortmund.de/grav/de/bico). For micro-clustering, the algorithm computes the coreset of the stream. Recustering is performed by using the kmeans++ algorithm on the coreset.

Author(s)

R-Interface: Matthias Carnein (<Matthias.Carnein@uni-muenster.de>), Dennis Assenmacher.

C-Implementation: Hendrik Fichtenberger, Marc Gille, Melanie Schmidt, Chris Schwiegelshohn, Christian Sohler.

References


Examples

```
stream <- DSD_Gaussians(k = 3, d = 2)
BICO <- DSC_BICO(k = 3, p = 10, space = 100, iterations = 10)
update(BICO, stream, n = 500)
plot(BICO, stream, type = "both")
```
**Description**

BIRCH builds a balanced-tree of Clustering Features (CFs) to summarize the stream. A CF is a tuple \((n, LS, SS)\) which represents a cluster by storing the number of elements \(n\), their linear sum \(LS\) and their squared sum \(SS\). Each new observation descends the tree by following its closest CF until a leaf node is reached. It is either merged into its closest leaf-CF or inserted as a new one. All leaf-CFs form the micro-clusters. Rebuilding the tree is realized by inserting all leaf-CF nodes into a new tree structure with an increased treshold.

**Usage**

\[
\text{DSC\_BIRCH}(\text{treshold, branching, maxLeaf, maxMem = 0, outlierThreshold = 0.25})
\]

**Arguments**

- `treshold` : treshold used to check whether a new datapoint can be absorbed or not
- `branching` : branching factor (maximum amount of child nodes for a nonleaf node) of the CF-Tree.
- `maxLeaf` : maximum number of entries within a leaf node
- `outlierThreshold` : threshold for identifying outliers when rebuilding the CF-Tree
- `maxMem` : memory limitation for the whole CFTree in bytes. Default is 0, indicating no memory restriction.

**Author(s)**

Dennis Assenmacher (<Dennis.Assenmacher@uni-muenster.de>), Matthias Carnein (<Matthias.Carnein@uni-muenster.de>)

**References**


**Examples**

```r
stream <- DSD_Gaussians(k = 3, d = 2)

BIRCH <- DSC\_BIRCH(treshold = .1, branching = 8, maxLeaf = 20)
update(BIRCH, stream, n = 500)

plot(BIRCH, stream)
```
**DSC_DBSCAN**

DBSCAN Macro-clusterer

**Description**

Implements the DBSCAN algorithm for reclustering micro-clusterings.

**Usage**

```
DSC_DBSCAN(eps, MinPts = 5, weighted = TRUE, description=NULL)
```

**Arguments**

- **eps**
  - radius of the eps-neighborhood.
- **MinPts**
  - minimum number of points required in the eps-neighborhood.
- **weighted**
  - logical indicating if a weighted version of DBSCAN should be used.
- **description**
  - optional character string to describe the clustering method.

**Details**

DBSCAN is a weighted extended version of the implementation in `fpc` where each micro-cluster center considered a pseudo point. For weighting we use in the MinPts comparison the sum of weights of the micro-cluster instead of the number.

DBSCAN first finds core points based on the number of other points in its eps-neighborhood. Then core points are joined into clusters using reachability (overlapping eps-neighborhoods).

Note that this clustering cannot be updated iteratively and every time it is used for (re)clustering, the old clustering is deleted.

**Value**

An object of class `DSC_DBSCAN` (a subclass of DSC, DSC_R, DSC_MACRO).

**Author(s)**

Michael Hahsler

**References**


**See Also**

DSC, DSC_MACRO
Examples

# 3 clusters with 5% noise
stream <- DSD_Gaussians(k=3, d=2, noise=0.05)

# Use DBSCAN to recluster micro clusters (a sample)
sample <- DSC_Sample(k=100)
update(sample, stream, 500)

dbscan <- DSC_DBSCAN(eps = .05)
recluster(dbscan, sample)
plot(dbscan, stream, type="both")

# For comparison we can cluster some data with DBSCAN directly
# Note: DBSCAN is not suitable for data streams since it passes over the data
# several times.
dbscan <- DSC_DBSCAN(eps = .05)
update(dbscan, stream, 500)
plot(dbscan, stream)

---

DSC_DBSTREAM  DBSTREAM clustering algorithm

Description

Implements a simple density-based stream clustering algorithm that assigns data points to micro-clusters with a given radius and implements shared-density-based reclustering.

Usage

DSC_DBSTREAM(r, lambda = 0.001, gaptime = 1000L,
       Cm = 3, metric = "Euclidean", shared_density = FALSE,
       alpha=0.1, k=0, minweight = 0)
get_shared_density(x, use_alpha = TRUE)
change_alpha(x, alpha)
get_cluster_assignments(x)

Arguments

- **r**  
The radius of micro-clusters.
- **lambda**  
The lambda used in the fading function.
- **gaptime**  
weak micro-clusters (and weak shared density entries) are removed every gaptime points.
- **Cm**  
minimum weight for a micro-cluster.
- **metric**  
metric used to calculate distances.
- **shared_density**  
Record shared density information. If set to TRUE then shared density is used for reclustering, otherwise reachability is used (overlapping clusters with less than \(r \times (1 - \alpha)\) distance are clustered together).
The number of macro clusters to be returned if macro is true.

k

alpha

For shared density: The minimum proportion of shared points between to clusters to warrant combining them (a suitable value for 2D data is .3). For reachability clustering it is a distance factor.

minweight

The proportion of the total weight a macro-cluster needs to have not to be noise (between 0 and 1).

x

A DSC_DBSTREAM object to get the shared density information from.

use_alpha

only return shared density if it exceeds alpha.

Details

The DBSTREAM algorithm checks for each new data point in the incoming stream, if it is below the threshold value of dissimilarity value of any existing micro-clusters, and if so, merges the point with the micro-cluster. Otherwise, a new micro-cluster is created to accommodate the new data point.

Although DSC_DBSTREAM is a micro clustering algorithm, macro clusters and weights are available.

get_cluster_assignments() can be used to extract the MC assignment for each data point clustered during the last update operation (note: update needs to be called with assignments = TRUE and the block size needs to be large enough). The function returns the MC index (in the current set of MCs obtained with, e.g., get_centers()) and as an attribute the permanent MC ids.

plot() for DSC_DBSTREAM has two extra logical parameters called assignment and shared_density which show the assignment area and the shared density graph, respectively.

Value

An object of class DSC_DBSTREAM (subclass of DSC, DSC_R, DSC_Micro).

Author(s)

Michael Hahsler and Matthew Bolanos

References

Michael Hahsler and Matthew Bolanos. Clustering data streams based on shared density between micro-clusters. IEEE Transactions on Knowledge and Data Engineering, 28(6):1449–1461, June 2016

See Also

DSC, DSC_Micro

Examples

```r
set.seed(8)
stream <- DSD_Gaussians(k = 3, noise = 0.05)

# create clusterer with r = 0.05
```
dbstream <- DSC_DBSTREAM(r = .05)
update(dbstream, stream, 1000)
dbstream

# check micro-clusters
nclusters(dbstream)
head(get_centers(dbstream))
plot(dbstream, stream)

# plot macro-clusters
plot(dbstream, stream, type = "both")

# plot micro-clusters with assignment area
plot(dbstream, stream, type = "both", assignment = TRUE)

# DBSTREAM with shared density
dbstream <- DSC_DBSTREAM(r = .05, shared_density = TRUE, Cm=5)
update(dbstream, stream, 1000)
dbstream
plot(dbstream, stream, type = "both")
# plot the shared density graph (several options)
plot(dbstream, stream, type = "both", shared_density = TRUE)
plot(dbstream, stream, type = "micro", shared_density = TRUE)
plot(dbstream, stream, type = "micro", shared_density = TRUE, assignment = TRUE)
plot(dbstream, stream, type = "none", shared_density = TRUE, assignment = TRUE)

# see how micro and macro-clusters relate
# each micro-cluster has an entry with the macro-cluster id
# Note: unassigned micro-clusters (noise) have an NA
microToMacro(dbstream)

# do some evaluation
evaluate(dbstream, stream, measure="purity")
evaluate(dbstream, stream, measure="cRand", type="macro")

# use DBSTREAM for conventional clustering (with assignments = TRUE so we can
# later retrieve the cluster assignments for each point)
data("iris")
dbstream <- DSC_DBSTREAM(r = 1)
update(dbstream, iris[,,-5], assignments = TRUE)
dbstream

cl <- get_cluster_assignments(dbstream)
cl

# micro-clusters
plot(iris[,,-5], col = cl, pch = cl)

# macro-clusters
plot(iris[,,-5], col = microToMacro(dbstream, cl))
DSC_DStream

D-Stream Data Stream Clustering Algorithm

Description

Implements the D-Stream data stream clustering algorithm.

Usage

DSC_DStream(gridsize, lambda = 1e-3, gaptime=1000L,
Cm=3, Cl=.8, attraction=FALSE, epsilon=.3, Cm2=Cm, k=NULL, N = 0)
get_attraction(x, relative=FALSE, grid_type = "dense", dist=FALSE)

Arguments

gridsize     Size of grid cells.
lambda      Fading constant used function to calculate the decay factor $2^{-\lambda}$. (Note: in the paper the authors use lamba to denote the decay factor and not the fading constant!)
gaptime     sporadic grids are removed every gaptime number of points.
Cm          density threshold used to detect dense grids as a proportion of the average expected density ($C_m > 1$). The average density is given by the total weight of the clustering over $N$, the number of grid cells.
Cl          density threshold to detect sporadic grids ($0 > C_l > C_m$). Transitional grids have a density between $C_l$ and $C_m$.
attraction  compute and store information about the attraction between adjacent grids. If TRUE then attraction is used to create macro-clusters, otherwise macro-clusters are created by merging adjacent dense grids.
epsilon     overlap parameter for attraction as a proportion of gridsize.
Cm2         threshold on attraction to join two dense grid cells (as a proportion on the average expected attraction). In the original algorithm $C_{m2}$ is equal to $C_m$.
k           alternative to $C_{m2}$ (not in the original algorithm). Create $k$ clusters based on attraction. In case of more than $k$ unconnected components, closer groups of MCs are joined.
N            Fix the number of grid cells used for the calculation of the density thresholds with $C_l$ and $C_m$. If $N$ is not given (0) then the algorithm tries to determine $N$ from the data. Note that this means that $N$ potentially increases over time and outliers might produce an extremely large value which will lead to a sudden creation of too many dense micro-clusters. The original paper assumed that $N$ is known a priori.
x            DSC_DStream object to get attraction values from.
relative    calculates relative attraction (normalized by the cluster weight).
grid_type   the attraction between what grid types should be returned?
dist        make attraction symmetric and transform into a distance.
Details

D-Stream creates an equally spaced grid and estimates the density in each grid cell using the count of points falling in the cells. Grid cells are classified based on density into dense, transitional and sporadic cells. The density is faded after every new point by a factor of \(2^{-\lambda}\). Every gaptime number of points sporadic grid cells are removed.

For reclustering D-Stream (2007 version) merges adjacent dense grids to form macro-clusters and then assigns adjacent transitional grids to macro-clusters. This behavior is implemented as `attraction=FALSE`.

The 2009 version of the algorithm adds the concept of attraction between grids cells. If `attraction=TRUE` is used then the algorithm produces macro-clusters based on attraction between dense adjacent grids (uses \(Cm2\) which in the original algorithm is equal to \(Cm\)).

For many functions (e.g., `get_centers()`, `plot()`), D-Stream adds a parameter `grid_type` with possible values of "dense", "transitional", "sparse", "all" and "used". This only returns the selected type of grid cells. "used" includes dense and adjacent transitional cells which are used in D-Stream for reclustering.

For plot D-Stream also provides extra parameters "grid" and "grid_type" to show micro-clusters as grid cells (density represented by gray values).

Note that DSC_DStream can at this point not be saved to disk using `save()` or `saveRDS()`. This functionality will be added later!

Value

An object of class `DSC_DStream` (subclass of `DSC`, `DSC_R`, `DSC_Micro`).

Author(s)

Michael Hahsler

References


Li Tu and Yixin Chen. 2009. Stream data clustering based on grid density and attraction. *ACM Transactions on Knowledge Discovery from Data*, 3(3), Article 12 (July 2009), 27 pages.

See Also

`DSC`, `DSC_Micro`

Examples

```r
stream <- DSD_BarsAndGaussians(noise=.05)
plot(stream)

# we set Cm=.8 to pick up the lower density clusters
dstream1 <- DSC_DStream(gridsize=1, Cm=1.5)
update(dstream1, stream, 1000)
dstream1
```
# micro-clusters (these are "used" grid cells)
ncusters(dstream1)
head(get_centers(dstream1))

# plot (DStream provides additional grid visualization)
plot(dstream1, stream)
plot(dstream1, stream, grid=TRUE)

# look only at dense grids
ncusters(dstream1, grid_type="dense")
plot(dstream1, stream, grid=TRUE, grid_type="dense")

# look at transitional and sparse cells
plot(dstream1, stream, grid=TRUE, grid_type="transitional")
plot(dstream1, stream, grid=TRUE, grid_type="sparse")

### Macro-clusters
# standard D-Stream uses reachability
ncusters(dstream1, type="macro")
get_centers(dstream1, type="macro")
plot(dstream1, stream, type="both", grid=TRUE)
evaluate(dstream1, stream, measure="crand", type="macro")

# use attraction for reclustering
dstream2 <- DSC_DStream(gridsize=1, attraction=TRUE, Cm=1.5)
update(dstream2, stream, 1000)
dstream2

plot(dstream2, stream, type="both", grid=TRUE)
evaluate(dstream2, stream, measure="crand", type="macro")

---

**DSC_EA**

**Evolutionary Algorithm**

**Description**

Reclustering using an evolutionary algorithm. This approach was designed for evoStream but can also be used for other micro-clustering algorithms. The evolutionary algorithm uses existing clustering solutions and creates small variations of them by combining and randomly modifying them. The modified solutions can yield better partitions and thus can improve the clustering over time. The evolutionary algorithm is incremental, which allows to improve existing macro-clusters instead of recomputing them every time.

**Usage**

DSC_EA(k, generations = 2000, crossoverRate = 0.8, mutationRate = 0.001, populationSize = 100)
Arguments

- **k**: number of macro-clusters
- **generations**: number of EA generations performed during reclustering
- **crossoverRate**: cross-over rate for the evolutionary algorithm
- **mutationRate**: mutation rate for the evolutionary algorithm
- **populationSize**: number of solutions that the evolutionary algorithm maintains

Author(s)

Matthias Carnein <Matthias.Carnein@uni-muenster.de>

References


Examples

```r
stream <- DSD_Memory(DSD_Gaussians(k = 3, d = 2), 1000)

## online algorithm
dbstream <- DSC_DBSTREAM(r=0.1)

## offline algorithm
EA <- DSC_EA(k=3, generations=1000)

## create pipeline and insert observations
two <- DSC_TwoStage(dbstream, EA)
update(two, stream, n=1000)

## plot resut
reset_stream(stream)
plot(two, stream, type="both")

## if we have time, evaluate additional generations. This can be called at any time, also between observations.
two$macro_dsc$RObj$recluster(2000)

## plot improved result
reset_stream(stream)
plot(two, stream, type="both")

## alternatively: do not create twostage but apply directly
reset_stream(stream)
update(dbstream, stream, n = 1000)
recluster(EA, dbstream)
reset_stream(stream)
plot(EA, stream)
```
Description

Stream clustering algorithm based on evolutionary optimization. The online component uses a simplified version of DBSTREAM to generate micro-clusters. The micro-clusters are then incrementally reclastered using an evolutionary algorithm. Evolutionary algorithms create slight variations by combining and randomly modifying existing solutions. By iteratively selecting better solutions, an evolutionary pressure is created which improves the clustering over time. Since the evolutionary algorithm is incremental, it is possible to apply it between observations, e.g. in the idle time of the stream. Whenever there is idle time, we can call the recluster function of the reference class to improve the macro-clusters (see example). The evolutionary algorithm can also be applied as a traditional reclustering step, or a combination of both. In addition, this implementation also allows to evaluate a fixed number of generations after each observation.

Usage

DSC_evoStream(r, lambda = 0.001, tgap = 100, k = 2,
    crossoverRate = 0.8, mutationRate = 0.001, populationSize = 100,
    initializeAfter = 2 * k, incrementalGenerations = 1,
    reclusterGenerations = 1000)

Arguments

- r: radius threshold for micro-cluster assignment
- lambda: decay rate
- tgap: time-interval between outlier detection and clean-up
- k: number of macro-clusters
- crossoverRate: cross-over rate for the evolutionary algorithm
- mutationRate: mutation rate for the evolutionary algorithm
- populationSize: number of solutions that the evolutionary algorithm maintains
- initializeAfter: number of micro-cluster required for the initialization of the evolutionary algorithm.
- incrementalGenerations: number of EA generations performed after each observation
- reclusterGenerations: number of EA generations performed during reclustering

Author(s)

Matthias Carnein <Matthias.Carnein@uni-muenster.de>
References

Examples

```r
stream <- DSD_Memory(DSD_Gaussians(k = 3, d = 2), 500)

## init evoStream
evoStream <- DSC_evoStream(r=0.05, k=3, incrementalGenerations=1, reclusteringGenerations=500)

## insert observations
update(evoStream, stream, n = 500)

## micro clusters
get_centers(evoStream, type="micro")

## micro weights
get_weights(evoStream, type="micro")

## macro clusters
get_centers(evoStream, type="macro")

## macro weights
get_weights(evoStream, type="macro")

## plot result
reset_stream(stream)
plot(evoStream, stream, type = "both")

## if we have time, evaluate additional generations.
## This can be called at any time, also between observations.
## by default, 1 generation is evaluated after each observation and
## 1000 generations during reclustering but we set it here to 500
evoStream$RObj$recluster(500)

## plot improved result
reset_stream(stream)
plot(evoStream, stream, type="both")

## get assignment of micro to macro clusters
microToMacro(evoStream)
```

---

**DSC_Hierarchical**

**Hierarchical Micro-Cluster Reclusterer**

**Description**

Implementation of hierarchical clustering to recluster a set of micro-clusters.
**Usage**

DSC_Hierarchical(k=NULL, h=NULL, method = "complete", min_weight=NULL, description=NULL)

**Arguments**

- **k**: The number of desired clusters.
- **h**: Height where to cut the dendrogram.
- **method**: the agglomeration method to be used. This should be (an unambiguous abbreviation of) one of "ward", "single", "complete", "average", "mcquitty", "median" or "centroid".
- **min_weight**: micro-clusters with a weight less than this will be ignored for reclustering.
- **description**: optional character string to describe the clustering method.

**Details**

Please refer to `hclust` for more details on the behavior of the algorithm.

Note that this clustering cannot be updated iteratively and every time it is used for (re)clustering, the old clustering is deleted.

**Value**

A list of class `DSC`, `DSC_R`, `DSC_Macro`, and `DSC_Hierarchical`. The list contains the following items:

- **description**: The name of the algorithm in the DSC object.
- **RObj**: The underlying R object.

**Author(s)**

Michael Hahsler

**See Also**

`DSC`, `DSC_Macro`

**Examples**

# Cassini dataset
stream <- DSD_mlbenchGenerator("cassini")

# Use hierarchical clustering to recluster micro-clusters
dbstream <- DSC_DBSTREAM(r=.05)
update(dbstream, stream, 500)

# reclustering using single-link and specifying k
hc <- DSC_Hierarchical(k=3, method="single")
recluster(hc, dbstream)
DSC_Kmeans

hc
plot(hc, stream, type="both")

# re-clustering by specifying height
hc <- DSC_Hierarchical(h=.2, method="single")
recluster(hc, dbstream)
hc
plot(hc, stream, type="both")

# For comparison we use hierarchical clustering directly on the data
# Note: hierarchical clustering is not a data stream clustering algorithm!
hc <- DSC_Hierarchical(k=3, method="single")
update(hc, stream, 500)
plot(hc, stream)

---

DSC_Kmeans  Kmeans Macro-clusterer

Description
Class implements the k-means algorithm for re-clustering a set of micro-clusters.

Usage

DSC_Kmeans(k, weighted = TRUE, iter.max = 10, nstart = 1,
algorithm = c("Hartigan-Wong", "Lloyd", "Forgy", "MacQueen"),
min_weight = NULL, description=NULL)

Arguments

- **k**: either the number of clusters, say k, or a set of initial (distinct) cluster centers. If a number, a random set of (distinct) rows in x is chosen as the initial centers.
- **weighted**: use a weighted k-means (algorithm is ignored).
- **iter.max**: the maximum number of iterations allowed.
- **nstart**: if centers is a number, how many random sets should be chosen?
- **algorithm**: character: may be abbreviated.
- **min_weight**: micro-clusters with a weight less than this will be ignored for re-clustering.
- **description**: optional character string to describe the clustering method.

Details
Please refer to function kmeans in stats for more details on the algorithm.

Note that this clustering cannot be updated iteratively and every time it is used for (re)clustering, the old clustering is deleted.
Value

An object of class DSC_Kmeans (subclass of DSC, DSC_R, DSCMacro)

Author(s)

Michael Hahsler

See Also

DSC, DSCMacro

Examples

```r
stream <- DSD_Gaussians(k=3, noise=0)

# create micro-clusters via sampling
sample <- DSC_Sample(k=20)
update(sample, stream, 500)
sample

# recluster micro-clusters
kmeans <- DSC_Kmeans(k=3)
recluster(kmeans, sample)
plot(kmeans, stream, type="both")

# For comparison we use k-means directly to cluster data
# Note: k-means is not a data stream clustering algorithm
kmeans <- DSC_Kmeans(k=3)
update(kmeans, stream, 500)
plot(kmeans, stream)
```

DSC_Marco

Abstract Class for Macro Clusterers

Description

Abstract class for all DSC Macro Clusterers.

Details

DSC_Macro cannot be instantiated. Calling DSC_Macro() results in an error.

Author(s)

Michael Hahsler

See Also

DSC
Abstract Class for Micro Clusterers

Description
Abstract class for all DSC Micro Clusterers.

Details
DSC_Micro cannot be instantiated. Calling DSC_Micro() results in an error.

Author(s)
Michael Hahsler

See Also
DSC

Reachability Micro-Cluster Reclusterer

Description
Implementation of reachability clustering (based on DBSCAN’s concept of reachability) to recluster a set of micro-clusters. Two micro-clusters are directly reachable if they are within each other’s epsilon-neighborhood (i.e., the distance between the centers is less then epsilon). Two micro-clusters are reachable if they are connected by a chain of pairwise directly reachable micro-clusters. All mutually reachable micro-clusters are put in the same cluster.

Usage
DSC_Reachability(epsilon, min_weight=NULL, description=NULL)

Arguments
- epsilon: radius of the epsilon-neighborhood.
- min_weight: micro-clusters with a weight less than this will be ignored for reclustering.
- description: optional character string to describe the clustering method.

Details
Reachability uses internally DSC_Hierarchical with single link.
Note that this clustering cannot be updated iteratively and every time it is used for (re)clustering, the old clustering is deleted.
Value

An object of class DSC_Reachability. The object contains the following items:

- **description**: The name of the algorithm in the DSC object.
- **Robj**: The underlying R object.

Author(s)

Michael Hahsler

References


See Also

DSC, DSC_Macro

Examples

```r
stream <- DSD_mlbenchGenerator("cassini")

# Recluster micro-clusters from DSC_Sample with reachability
sample <- DSC_Sample(k = 200)
update(sample, stream, 1000)

reach <- DSC_Reachability(epsilon=0.3)
recluster(reach, sample)
plot(reach, stream, type="both")

# For comparison we using reachability clustering directly on data points
# Note: reachability is not a data stream clustering algorithm taking O(n^2)
# time and space.
reach <- DSC_Reachability(epsilon=0.2)
update(reach, stream, 500)
reach
plot(reach, stream)
```
**DSC_Sample**

*Extract a Fixed-size Sample from a Data Stream*

**Description**

Extracts a sample from a data stream using Reservoir Sampling. The sample is stored as a set of micro-clusters to be compatible with other data DSC stream clustering algorithms.

**Usage**

```r
DSC_Sample(k = 100, biased = FALSE)
```

**Arguments**

- `k` the number of points to be sampled from the stream.
- `biased` if `FALSE` then a regular (unbiased) reservoir sampling is used. If `TRUE` then the sample is biased towards keeping more recent data points (see Details section).

**Details**

If `biased=FALSE` then the reservoir sampling algorithm by McLeod and Bellhouse (1983) is used. This sampling makes sure that each data point has the same chance to be sampled. All sampled points will have a weight of 1. Note that this might not be ideal for an evolving stream since very old data points have the same chance to be in the sample as newer points.

If `biased=TRUE` then sampling prefers newer points using the modified reservoir sampling algorithm 2.1 by Aggarwal (2006). New points are always added. They replace a random point in the reservoir with a probability of reservoir size over `k`. This an exponential bias function of $2^{-\lambda}$ with $\lambda = 1/k$.

**Value**

An object of class `DSC_Sample` (subclass of `DSC, DSC_R, DSC_Micro`).

**Author(s)**

Michael Hahsler

**References**


DSC_Static

Create as Static Copy of a Clustering

Description

This representation cannot perform clustering anymore, but it also does not need the supporting data structures. It only stores the cluster centers and weights.

Usage

DSC_Static(x, type=c("auto", "micro", "macro"), k_largest=NULL, min_weight=NULL)
Arguments

- **x**  The clustering (a DSD object) to copy.
- **type**  which clustering to copy.
- **k_largest**  only copy the k largest (highest weight) clusters.
- **min_weight**  only copy clusters with a weight larger or equal to min_weight.

Value

An object of class DSC_Static (sub class of DSC, DSC_R). The list also contains either DSC_Micro or DSC_Macro depending on what type of clustering was copied.

Author(s)

Michael Hahsler

See Also

DSC, DSC_Micro, DSC_Macro

Examples

```r
stream <- DSD_Gaussians(k=3, noise=0.05)
dstream <- DSC_DStream(gridsize=0.05)
update(dstream, stream, 500)
dstream
plot(dstream, stream)

# create a static copy of the clustering
static <- DSC_Static(dstream)
static
plot(static, stream)

# copy only the 5 largest clusters
static2 <- DSC_Static(dstream, k_largest=5)
static2
plot(static2, stream)

# copy all clusters with a weight of at least .3
static3 <- DSC_Static(dstream, min_weight=.3)
static3
plot(static3, stream)
```
DSC_TwoStage  TwoStage Clustering Process

Description
Combines a micro and a macro clustering algorithm into a single process.

Usage
DSC_TwoStage(micro, macro)

Arguments
- micro: Clustering algorithm used in the online stage (DSC_micro)
- macro: Clustering algorithm used for reclustering in the offline stage (DSC_macro)

Details
update() runs the micro-clustering stage and only when macro cluster centers/weights are requested, then the offline stage reclustering is automatically performed.

Value
An object of class DSC_TwoStage (subclass of DSC, DSC_Macro).

Author(s)
Michael Hahsler

See Also
DSC, DSC_Macro

Examples
```
stream <- DSD_Gaussians(k=3)

# Create a clustering process that uses a window for the online stage and
# k-means for the offline stage (reclustering)
win_km <- DSC_TwoStage(
  micro=DSC_Window(horizon=100),
  macro=DSC_Kmeans(k=3)
)
win_km

update(win_km, stream, 200)
win_km
plot(win_km, stream, type="both")
evaluate(win_km, stream, assign="macro")
```
A sliding window from a Data Stream

Description

Implements a sliding window which keeps a fixed amount (window length) of the most recent data points of the stream.

Usage

DSC_Window(horizon = 100, lambda=0)

Arguments

<table>
<thead>
<tr>
<th>Argument</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>horizon</td>
<td>the window length.</td>
</tr>
<tr>
<td>lambda</td>
<td>decay factor damped window model. lambda=0 means no dampening.</td>
</tr>
</tbody>
</table>

Details

If lambda is greater than 0 then the weight uses a damped window model (Zhu and Shasha, 2002). The weight for points in the window follows $2^{-\lambda t}$ where $t$ is the age of the point.

Value

An object of class DSC_Window (subclass of DSC, DSC_R, DSC_Micro).

Author(s)

Michael Hahsler

References


See Also

DSC, DSC_Micro

Examples

```r
stream <- DSD_Gaussians(k=3, d=2, noise=0.05)

window <- DSC_Window(horizon=100)

window

update(window, stream, 200)

window
```
# plot micro-clusters
plot(window, stream)

# animation for a window using a damped window model. The weight decays
# with a half-life of 25
## Not run:
window <- DSC_Window(horizon=25, lambda=1/25)
animate_cluster(window, stream, horizon=1, n=100, xlim=c(0,1), ylim=c(0,1))

## End(Not run)

---

## Description

Abstract base classes for DSD (Data Stream Data Generator).

## Details

The DSD class cannot be instantiated, but it serves as a abstract base class from which all DSD objects inherit.

DSD_R inherits form DSD and is the abstract parent class for DSD implemented in R. To create a new R-based implementation there are only two function that needs to be implemented for a new DSD subclass: A creator function (the name should start with DSD_) and a method get_points() for that class.

DSD provides common functionality like print(), plot(), etc.

Note that calling DSD() or DSD_R() results in an error since both are abstract classes.

## Author(s)

Michael Hahsler

## See Also

animate_data, animate_cluster, cluster, evaluate, get_points, plot, recluster, write_stream

## Examples

# create data stream with three clusters in 3-dimensional space
stream <- DSD_Gaussians(k=3, d=3)

# get points from stream
get_points(stream, n=5)

# get points with true cluster assignment
p <- get_points(stream, n=5, cluster=TRUE)
attr(p, "cluster")
# Data Stream Generator for Bars and Gaussians

### Description

A data stream generator which creates the shape of two bars and two Gaussians clusters with different density.

### Usage

```r
DSD_BarsAndGaussians(angle = NULL, noise = 0)
```

### Arguments

- `angle`: rotation in degrees. `NULL` will produce a random rotation.
- `noise`: The amount of noise that should be added to the output.

### Value

Returns a `DSD_BarsAndGaussians` object.

### Author(s)

Michael Hahsler

### See Also

- `DSD`

### Examples

```r
# create data stream with three clusters in 2D
stream <- DSD_BarsAndGaussians(noise=0.1)

# plotting the data
plot(stream)
```
**Description**

A data stream generator that generates several dynamic streams intended to be benchmarks to compare data stream clustering algorithms.

**Usage**

`DSD_Benchmark(i=1)`

**Arguments**

- `i` number of the benchmark.

**Details**

Currently available benchmarks are 1 and 2.

**Value**

Returns a DSD object.

**Author(s)**

Michael Hahsler

**See Also**

[DSD](#)

**Examples**

```r
stream <- DSD_Benchmark(i=1)
## Not run:
animate_data(stream, n=10000, horizon=100, xlim=c(0,1), ylim=c(0,1))
## End(Not run)
```
DSD_Cubes

**Description**

A data stream generator that produces a data stream with static (hyper) cubes filled uniformly with data points.

**Usage**

DSD_Cubes(k=2, d=2, center, size, p, noise = 0, noise_range)

**Arguments**

- **k**: Determines the number of clusters.
- **d**: Determines the number of dimensions.
- **center**: A matrix of means for each dimension of each cluster.
- **size**: A k times d matrix with the cube dimensions.
- **p**: A vector of probabilities that determines the likelihood of generated a data point from a particular cluster.
- **noise**: Noise probability between 0 and 1. Noise is uniformly distributed within noise range (see below).
- **noise_range**: A matrix with d rows and 2 columns. The first column contains the minimum values and the second column contains the maximum values for noise.

**Value**

Returns a DSD_Cubes object (subclass of DSD_R, DSD).

**Author(s)**

Michael Hahsler

**See Also**

- DSD

**Examples**

```r
# create data stream with three clusters in 3D
stream <- DSD_Cubes(k=3, d=3)

# plotting the data
plot(stream)
```
DSD_Gaussians  

*Mixture of Gaussians Data Stream Generator*

**Description**

A data stream generator that produces a data stream with a mixture of static Gaussians.

**Usage**

DSD_Gaussians(k=2, d=2, mu, sigma, p, separation=0.2, noise=0, noise_range)

**Arguments**

- **k**  
  Determines the number of clusters.
- **d**  
  Determines the number of dimensions.
- **mu**  
  A matrix of means for each dimension of each cluster.
- **sigma**  
  A list of length \( k \) of covariance matrices.
- **p**  
  A vector of probabilities that determines the likelihood of generating a data point from a particular cluster.
- **separation**  
  Minimum distance between cluster centers to reduce overlap between clusters (0–8).
- **noise**  
  Noise probability between 0 and 1. Noise is uniformly distributed within noise range (see below).
- **noise_range**  
  A matrix with \( d \) rows and 2 columns. The first column contains the minimum values and the second column contains the maximum values for noise.

**Details**

DSD_Gaussians creates a mixture of \( k \) \( d \)-dimensional static Gaussians in approximately unit space. The centers \( \mu \) and the covariance matrices \( \sigma \) can be supplied or will be randomly generated. The probability vector \( p \) defines for each cluster the probability that the next data point will be chosen from it (defaults to equal probability).

The generation method is similar to the one suggested by Jain and Dubes (1988).

**Value**

Returns a DSD_Gaussians object (subclass of DSD_R, DSD) which is a list of the defined params. The params are either passed in from the function or created internally. They include:

- **description**  
  A brief description of the DSD object.
- **k**  
  The number of clusters.
- **d**  
  The number of dimensions.
- **mu**  
  The matrix of means of the dimensions in each cluster.
- **sigma**  
  The covariance matrix.
- **p**  
  The probability vector for the clusters.
- **noise**  
  A flag that determines if or if not noise is generated.
**Description**

This class provides a data stream interface for data stored in memory as matrix-like objects (including data frames). All or a portion of the stored data can be replayed several times.

**Usage**

```
DSD_Memory(x, n, k=NA, loop=FALSE,
            class = NULL, description=NULL)
```

**Arguments**

- `x`: A matrix-like object containing the data. If `x` is a DSD object then a data frame for `n` data points from this DSD is created.
- `n`: Number of points used if `x` is a DSD object. If `x` is a matrix-like object then `n` is ignored.

**Examples**

```r
# create data stream with three clusters in 3-dimensional data space
stream1 <- DSD_Gaussians(k=3, d=3)
plot(stream1)

# create data stream with specified clusterpositions, # 20% noise in a given bounding box and # with different densities (1 to 9 between the two clusters)
stream2 <- DSD_Gaussians(k=2, d=2,
                         mu=rbind(c(-.5,-.5), c(.5,.5)),
                         noise=0.2, noise_range=rbind(c(-1,1),c(-1,1)),
                         p=c(.1,.9))
plot(stream2)
```
Optional: The known number of clusters in the data loop
Should the stream start over when it reaches the end? class
Vector with the class/cluster label (only used if x is not a DSD object).
description
character string with a description.

Details
In addition to regular data.frames other matrix-like objects that provide subsetting with the bracket operator can be used. This includes ffdf (large data.frames stored on disk) from package ff and big.matrix from bigmemory.

Value
Returns a DSD_Memory object (subclass of DSD_R, DSD).

Author(s)
Michael Hahsler

See Also
DSD, reset_stream

Examples

# store 1000 points from a stream
stream <- DSD_Gaussians(k=3, d=2)
replayer <- DSD_Memory(stream, k=3, n=1000)
replayer
plot(replayer)

# creating 2 clusterers of different algorithms
dsc1 <- DSC_DBSTREAM(r=0.1)
dsc2 <- DSC_DStream(gridsize=0.1, Cm=1.5)

# clustering the same data in 2 DSC objects
reset_stream(replayer) # resetting the replayer to the first position
update(dsc1, replayer, 500)
reset_stream(replayer)
update(dsc2, replayer, 500)

# plot the resulting clusterings
reset_stream(replayer)
plot(dsc1, replayer, main="DBSTREAM")
reset_stream(replayer)
plot(dsc2, replayer, main="D-Stream")

### use a data.frame to create a stream (3rd col. contains the assignment)
df <- data.frame(x=runif(100), y=runif(100),
    class=sample(1:3, 100, replace=TRUE))
head(df)
stream <- DSD_Memory(df[, c("x", "y")], class=df[,"class"])
stream

---

### Description

Creates an evolving DSD that consists of several MGCs.

### Usage

```r
DSD_MG(dimension = 2, ..., labels=NULL, description=NULL)
add_cluster(x, c, label=NULL)
get_clusters(x)
remove_cluster(x, i)
```

### Arguments

- `dimension` the dimension of the DSD object
- `...` initial set of MGCs
- `x` A `DSD_MG` object.
- `c` The cluster that should be added to the `DSD_MG` object.
- `i` The index of the cluster that should be removed from the `DSD_MG` object.
- `label, labels` integer representing the cluster label. NA represents noise. If labels are not specified, then each new cluster gets a new label.
- `description` An optional string used by `print` to describe the data generator.

### Details

This DSD is able to generate complex datasets that are able to evolve over a period of time. Its behavior is determined by the MGCs it is composed of.

### Author(s)

Matthew Bolanos

### See Also

`MGC_Function, MGC_Linear, MGC_Noise, MGC_Random` for details on the different MGC objects.
Examples

```R
### create an empty DSD_MG
stream <- DSD_MG(dim = 2)

### add two clusters
cl <- MGC_Random(density=50, center=c(50,50), parameter=1, randomness = )
add_cluster(stream, cl)

cl2 <- MGC_Noise(density=1, range=rbind(c(-20,120), c(-20,120)))
add_cluster(stream, cl2)

get_clusters(stream)
plot(stream, xlim=c(-20,120), ylim=c(-20,120))

### Not run:
animate_data(stream, n=5000, xlim=c(-20,120), ylim=c(-20,120))

### End(Not run)

### remove cluster 1
remove_cluster(stream, 1)

get_clusters(stream)
plot(stream, xlim=c(-20,120), ylim=c(-20,120))

### create a more complicated cluster structure (using 2 clusters with the same label to form an L shape)
stream <- DSD_MG(dim=2,
  MGC_Static(density=10, center=c(5,.2), par=c(.4,.2), shape=MGC_Shape_Block),
  MGC_Static(density=10, center=c(6,.5), par=c(.2,.4), shape=MGC_Shape_Block),
  MGC_Static(density=5, center=c(.39,.53), par=c(.16,.35), shape=MGC_Shape_Block),
  MGC_Noise(density=1, range=rbind(c(0,1), c(0,1))),
  labels= c(1, 1, 2, NA)
)
plot(stream, xlim=c(0,1), ylim=c(0,1))

### simulate the clustering of a splitting cluster
c1 <- MGC_Linear(dim = 2, keyframelist = list(
    keyframe(time = 1, dens = 20, center = c(0,0), param = 10),
    keyframe(time = 50, dens = 10, center = c(50,50), param = 10),
    keyframe(time = 100, dens = 10, center = c(50,100),param = 10)
))

### Note: Second cluster appear at time=50
c2 <- MGC_Linear(dim = 2, keyframelist = list(
    keyframe(time = 50, dens = 10, center = c(50,50), param = 10),
    keyframe(time = 100, dens = 10, center = c(100,50),param = 10)
)
```
DSD_mlbenchData

Description
Provides a convenient stream interface for data sets from the mlbench package.

Usage
DSD_mlbenchData(data=NULL, loop = FALSE, random = FALSE, scale = FALSE)

Arguments
- data: The name of the dataset from mlbench. If missing then a list of all available data sets is shown and returned.
- loop: A flag that tells the stream to loop or not to loop over the data frame.
- random: A flag that determines if the data should be in a random order.
- scale: A flag that determines if the data should be scaled.

Details
The DSD_mlbenchData class is designed to be a wrapper class for data that is held in memory in either a data frame or matrix form. It is a subclass of DSD_Memory.

Call DSD_mlbenchData with a missing value for data to get a list of all available data sets.

Value
Returns a DSD_mlbenchData object which is also of class DSD_Memory.

Author(s)
Michael Hahsler and Matthew Bolanos
DSD_mlbenchGenerator

See Also
DSD, DSD_Memory, reset_stream

Examples
stream <- DSD_mlbenchData("Shuttle")
stream
plot(stream, n=100)

DSD_mlbenchGenerator  mlbench Data Stream Generator

Description
A data stream generator class that interfaces data generators found in mlbench.

Usage
DSD_mlbenchGenerator(method, ...)

Arguments
method  The name of the mlbench data generator.
...  Parameters for the mlbench data generator.

Details
The DSD_mlbenchGenerator class is designed to be a wrapper class for data created by data generators in the mlbench library.
Call DSD_mlbenchGenerator with missing method to get a list of available methods.

Value
Returns a DSD_mlbenchGenerator object (subclass of DSD_R, DSD) which is a list of the defined parameters. The parameters are either passed in from the function or created internally. They include:
description  The name of the class of the DSD object.
method  The name of the mlbench data generator.
variables  The variables for the mlbench data generator.

Author(s)
John Forrest
See Also

DSD

Examples

stream <- DSD_mlbenchGenerator(method="cassini")

plot(stream, n=500)

DSD_ReadCSV Read a Data Stream from File

Description

A DSD class that reads a data stream from a file or any R connection.

Usage

DSD_ReadCSV(file, k=NA, take=NULL, class=NULL, loop=FALSE,
sep="", header=FALSE, skip=0, colClasses = NA, ...)
close_stream(dsd)

Arguments

file A file/URL or an open connection.
k Number of true clusters, if known.
take indices of columns to extract from the file.
class column index for the class attribute/cluster label. If take is specified then it needs to also include the class/label column.
loop If enabled, the object will loop through the stream when the end has been reached. If disabled, the object will warn the user upon reaching the end.
sep The character string that separates dimensions in data points in the stream.
header Does the first line contain variable names?
skip the number of lines of the data file to skip before beginning to read data.
colClasses A vector of classes to be assumed for the columns passed on to read.table.
... Further arguments are passed on to read.table. This can for example be used for encoding, quotes, etc.
dsd A object of class DSD_ReadCSV.
Details

DSD_ReadCSV uses `read.table()` to read in data from an R connection. The connection is responsible for maintaining where the stream is currently being read from. In general, the connections will consist of files stored on disk but have many other possibilities (see `connection`).

The implementation tries to gracefully deal with slightly corrupted data by dropping points with inconsistent reading and producing a warning. However, this might not always be possible resulting in an error instead.

The position in the file can be reset to the beginning using `reset_stream()`. The connection can be closed using `close_stream()`.

Value

An object of class `DSD_ReadCSV` (subclass of `DSD_R`, `DSD`).

Author(s)

Michael Hahsler

See Also

`DSD_read_csv`, `reset_stream`, `read.table`.

Examples

```r
# creating data and writing it to disk
stream <- DSD_Gaussians(k=3, d=5)
write_stream(stream, "data.txt", n=10, header = TRUE, sep="",")

# reading the same data back (as a loop)
stream2 <- DSD_ReadCSV("data.txt", sep="","", header = TRUE, loop=TRUE)
stream2

# get points (first a single point and then 20 using loop)
get_points(stream2)
get_points(stream2, n=20)

# clean up
close_stream(stream2)
file.remove("data.txt")

# example with a part of the kddcup1999 data (take only cont. variables)
file <- system.file("examples", "kddcup10000.data.gz", package="stream")
stream <- DSD_ReadCSV(gzfile(file),
  take=c(1, 5, 6, 8:11, 13:20, 23:42), class=42, k=7)
stream
get_points(stream, 5, class = TRUE)

# plot 100 points (projected on the first two principal components)
```
DSD_ReadDB

Read a Data Stream from an open DB Query

Description
A DSD class that reads a data stream from an open DB result set from a relational database with using R’s data base interface (DBI).

Usage
DSD_ReadDB(result, k=NA, class=NULL, description=NULL)

Arguments
- result: An open DBI result set.
- k: Number of true clusters, if known.
- class: column index for the class/cluster assignment.
- description: a character string describing the data.

Details
This class provides a streaming interface for result sets from a database with via DBI. You need to connect to the database and submit a SQL query using dbGetQuery() to obtain a result set. Make sure that your query only includes the columns that should be included in the stream. Do not forget to close the result set and the database connection.

Value
An object of class DSD_ReadDB (subclass of DSD_R, DSD).

Author(s)
Michael Hahsler

See Also
DSD, dbGetQuery
Examples

```r
### create a data base with a table with 3 Gaussians
library("RSQlite")
con <- dbConnect(RSQlite::SQLite(), ":memory:"
points <- get_points(DSD_Gaussians(k=3, d=2), 600, class = TRUE)
head(points)

dbWriteTable(con, "gaussians", points)

### prepare a query result set
res <- dbSendQuery(con, "SELECT X1, X2, class FROM gaussians")
res

### create a stream interface to the result set
stream <- DSD_ReadDB(res, k=3, class = 3)

### get points
get_points(stream, 5, class = TRUE)
plot(stream)

### clean up
dbClearResult(res)
dbDisconnect(con)
```

---

**DSD_ScaleStream**  
*Scale a Stream from a DSD*

### Description

Make an unscaled data stream into a scaled data stream.

### Usage

```r
DSD_ScaleStream(dsd, center=TRUE, scale=TRUE, n=1000, reset=FALSE)
```

### Arguments

- **dsd**: A object of class DSD that will be scaled.
- **center**, **scale**: logical or a numeric vector of length equal to the number of columns used for centering/scaling (see function `scale`).
- **n**: The number of points used to creating the centering/scaling
- **reset**: Try to reset the stream to its beginning after taking n points for scaling.

### Details

`scale_stream()` estimates the values for centering and scaling (see `scale` in `base`) using n points from the stream.
**Value**

An object of class DSD.ScaleStream (subclass of DSD_R, DSD).

**Author(s)**

Michael Hahsler

**See Also**

DSD, reset_stream, scale in base.

**Examples**

```r
stream <- DSD_Gaussians(k=3, d=3)
plot(stream)

# scale stream using 100 points
stream_scaled <- DSD_ScaleStream(stream, n=100)
plot(stream_scaled)
```

---

**Description**

A data stream generator that generates a data stream in the shape of a target. It has a single Gaussian cluster in the center and a ring that surrounds it.

**Usage**

```r
DSD_Target(center_sd = 0.05, center_weight = 0.5, ring_r = 0.2,
            ring_sd = 0.02, noise = 0)
```

**Arguments**

- `center_sd`: standard deviation of center
- `center_weight`: proportion of points in center
- `ring_r`: average ring radius
- `ring_sd`: standard deviation of ring radius
- `noise`: proportion of noise

**Details**

DSD_Target is a DSD generator for stream data. It has been implemented entirely in R, so there is no computational overhead with communicating to the Java Runtime Interface (JRI) or native C code. This DSD will produce a singular Gaussian cluster in the center with a ring around it.
Value

Returns a DSD_Target object which is a list of the defined params. The params are either passed in from the function or created internally. They include:

- **description**: A brief description of the DSD object.
- **k**: The number of clusters.
- **d**: The number of dimensions.

Author(s)

Michael Hahsler

See Also

DSD

Examples

```r
# create data stream with three clusters in 2D
stream <- DSD_Target()
# plotting the data
plot(stream)
```

---

**DSDUniformNoise**

*Uniform Noise Data Stream Generator*

Description

This generator produces uniform noise in a d-dimensional unit (hyper) cube.

Usage

```r
DSDUniformNoise(d=2, range=NULL)
```

Arguments

- **d**: Determines the number of dimensions.
- **range**: A matrix with two columns and d rows giving the minimum and maximum for each dimension. Defaults to the range of [0, 1].

Value

Returns a DSDUniformNoise object (subclass of DSD_R, DSD).

Author(s)

Michael Hahsler
See Also

DSD

Examples

# create data stream with three clusters in 2D
stream <- DSD_UniformNoise(d=2)
plot(stream, n=100)

# specify a different range for each dimension
stream <- DSD_UniformNoise(d=3, range=rbind(c(0,1), c(0,10), c(0,5)))
plot(stream, n=100)

Description

Abstract class for frequent pattern mining algorithms for data streams. Currently, stream does not implement frequent pattern mining algorithms.

Author(s)

Michael Hahsler

See Also

DST

Data Stream Operator Base Classes

Description

Abstract base classes for all DSO (Data Stream Operator) classes.

Details

The DSO class cannot be instantiated (calling DSO() produces errors), but it serve as a base class from which other DSO classes inherit.

Data stream operators use update() to process new data from the stream. The result of the operator can be obtained via get_points() and get_weights().
Author(s)
Michael Hahsler

See Also
update, get_points, get_weights, DSO_Window, DSO_Sample

---

**DSO_Sample**

*Sampling from a Data Stream (Data Stream Operator)*

**Description**

Extracts a sample form a data stream using Reservoir Sampling.

**Usage**

```r
DSO_Sample(k = 100, biased = FALSE)
```

**Arguments**

- `k`: the number of points to be sampled from the stream.
- `biased`: if FALSE then a regular (unbiased) reservoir sampling is used. If true then the sample is biased towards keeping more recent data points (see Details section).

**Details**

If `biased=FALSE` then the reservoir sampling algorithm by McLeod and Bellhouse (1983) is used. This sampling makes sure that each data point has the same chance to be sampled. All sampled points will have a weight of 1. Note that this might not be ideal for an evolving stream since very old data points have the same chance to be in the sample as newer points.

If `bias=TRUE` then sampling prefers newer points using the modified reservoir sampling algorithm 2.1 by Aggarwal (2006). New points are always added. They replace a random point in the reservoir with a probability of `reservoir size over k`. This an exponential bias function of $2^{-\lambda}$ with $\lambda = 1/k$.

**Value**

An object of class DSO_Sample (subclass of DSO).

**Author(s)**

Michael Hahsler
DSO_Window

References


See Also

DSO

Examples

stream <- DSD_Gaussians(k=3, noise=0.05)
sample <- DSO_Sample(k=20)
update(sample, stream, 500)
sample

# plot points in sample
plot(get_points(sample))

---

DSO_Window  Sliding Window (Data Stream Operator)

Description

 Implements a sliding window data stream operator which keeps a fixed amount (window length) of the most recent data points of the stream.

Usage

DSO_Window(horizon = 100, lambda=0)

Arguments

horizon  the window length.
lambda  decay factor damped window model. lambda=0 means no dampening.

Details

 If lambda is greater than 0 then the weight uses a damped window model (Zhu and Shasha, 2002). The weight for points in the window follows $2^{-\lambda t}$ where $t$ is the age of the point.
Value

An object of class DS0_Window (subclass of DS0).

Author(s)

Michael Hahsler

References


See Also

DSO

Examples

stream <- DSD_Gaussians(k=3, d=2, noise=0.05)

window <- DS0_Window(horizon=100)
window

update(window, stream, 200)
window

# plot points in window
plot(get_points(window))

Abstract Base Class for All Data Stream Mining Tasks

Description

Abstract base class for all data stream mining tasks. Current tasks are data stream clustering DSC, classification on data streams DSClassify and frequent pattern mining on data streams DSFP.

Author(s)

Michael Hahsler

See Also

DSC, DSClassify, DSFP


**Description**

Gets evaluation measures for micro or macro-clusters from a DSC object given the original DSD object.

**Usage**

```r
evaluate(dsc, dsd, measure, n = 100, type=c("auto", "micro", "macro"),
assign="micro", assignmentMethod=c("auto", "model", "nn"),
noise = c("class", "exclude"), ...)
evaluate_cluster(dsc, dsd, measure, n = 1000,
type=c("auto", "micro", "macro"),
assign="micro", assignmentMethod=c("auto", "model", "nn"),
horizon=100, verbose=FALSE, noise = c("class", "exclude"), ...)
```

**Arguments**

- **dsc** The DSC object that the evaluation measure is being requested from.
- **dsd** The DSD object that holds the initial training data for the DSC.
- **measure** Evaluation measure(s) to use. If missing then all available measures are returned.
- **n** The number of data points being requested.
- **type** Use micro- or macro-clusters for evaluation. Auto used the class of dsc to decide.
- **assign** Assign points to micro or macro-clusters?
- **assignmentMethod** How are points assigned to clusters for evaluation (see get_assignment)?
- **horizon** Evaluation is done using horizon many previous points (see detail section).
- **verbose** report progress?
- **noise** how to handle noise points in the data. Options are to treat as a separate class (default) or to exclude them from evaluation.
- **...** Unused arguments are ignored.

**Details**

For evaluation each data points are assigned to its nearest cluster using Euclidean distance to the cluster centers. Then for each cluster the majority class is determined. Based on the majority class several evaluation measures can be computed.

For evaluate_cluster the most commonly used method of prequential error estimation (see Gama, Sebastiao and Rodrigues; 2013). The data points in the horizon are first used to calculate the evaluation measure and then they are used for updating the cluster model. Many evaluation
measures are calculated with code from the packages **cluster**, **clue** and **fpc**. Detailed documentation can be found in these packages (see Section See Also.)

The following information items are available:

- "numMicroClusters" number of micro-clusters
- "numMacroClusters" number of macro-clusters
- "numClasses" number of classes

The following noise-related items are available:

- "noisePredicted" Number data points predicted as noise
- "noiseActual" Number of data points which are actually noise
- "noisePrecision" Precision of the predicting noise (i.e., number of correctly predicted noise points over the total number of points predicted as noise)

The following internal evaluation measures are available:

- "SSQ" within cluster sum of squares. Assigns each non-noise point to its nearest center from the clustering and calculates the sum of squares
- "silhouette" average silhouette width (actual noise points which stay unassigned by the clustering algorithm are removed; regular points that are unassigned by the clustering algorithm will form their own noise cluster) (**cluster**)
- "average.between" average distance between clusters (**fpc**)
- "average.within" average distance within clusters (**fpc**)
- "max.diameter" maximum cluster diameter (**fpc**)
- "min.separation" minimum cluster separation (**fpc**)
- "ave.within.cluster.ss" a generalization of the within clusters sum of squares (half the sum of the within cluster squared dissimilarities divided by the cluster size) (**fpc**)
- "g2" Goodman and Kruskal’s Gamma coefficient (**fpc**)
- "pearsongamma" correlation between distances and a 0-1-vector where 0 means same cluster, 1 means different clusters (**fpc**)
- "dunn" Dunn index (minimum separation / maximum diameter) (**fpc**)
- "dunn2" minimum average dissimilarity between two cluster / maximum average within cluster dissimilarity (**fpc**)
- "entropy" entropy of the distribution of cluster memberships (**fpc**)
- "wb.ratio" average.within/average.between (**fpc**)

The following external evaluation measures are available:

- "precision", "recall", "F1" F1. A true positive (TP) decision assigns two points in the same true cluster also to the same cluster, a true negative (TN) decision assigns two points from two different true clusters to two different clusters. A false positive (FP) decision assigns two points from the same true cluster to two different clusters. A false negative (FN) decision assigns two points from the same true cluster to different clusters.

\[
\text{precision} = \frac{TP}{TP+FP} \\
\text{recall} = \frac{TP}{TP+FN}
\]

The F1 measure is the harmonic mean of precision and recall.
- "purity" Average purity of clusters. The purity of each cluster is the proportion of the points of the majority true group assigned to it (see Cao et al. (2006))
- "Euclidean" Euclidean dissimilarity of the memberships (see Dimitriadou, Weingessel and Hornik (2002)) (clue)
- "Manhattan" Manhattan dissimilarity of the memberships (clue)
- "Rand" Rand index (see Rand (1971)) (clue)
- "cRand" Adjusted Rand index (see Hubert and Arabie (1985)) (clue)
- "NMI" Normalized Mutual Information (see Strehl and Ghosh (2002)) (clue)
- "KP" Katz-Powell index (see Katz and Powell (1953)) (clue)
- "angle" maximal cosine of the angle between the agreements (clue)
- "diag" maximal co-classification rate (clue)
- "FM" Fowlkes and Mallows’s index (see Fowlkes and Mallows (1983)) (clue)
- "Jaccard" Jaccard index (clue)
- "PS" Prediction Strength (see Tibshirani and Walter (2005)) (clue)
- "vi" variation of information (VI) index (fpc)

Many measures are the average over all clusters. For example, purity is the average purity over all clusters.

For DSC_Micro objects, data points are assigned to micro-clusters and then each micro-cluster is evaluated. For DSC_Macro objects, data points by default (assign="micro") also assigned to micro-clusters, but these assignments are translated to macro-clusters. The evaluation is here done for macro-clusters. This is important when macro-clustering is done with algorithms which do not create spherical clusters (e.g., hierarchical clustering with single-linkage or DBSCAN) and this assignment to the macro-clusters directly (i.e., their center) does not make sense.

Using type and assign, the user can select how to assign data points and ad what level (micro or macro) to evaluate.

Many of the above measures are implemented package clue in function cl_agreement().

evaluate_cluster() is used to evaluate an evolving data stream using the method described by Wan et al. (2009). Of the n data points horizon many points are clustered and then the evaluation measure is calculated on the same data points. The idea is to find out if the clustering algorithm was able to adapt to the changing stream.

Value

evaluate returns an object of class stream_eval which is a numeric vector of the values of the requested measures and two attributes, "type" and "assign", to see at what level the evaluation was done.

Author(s)

Michael Hahsler, Matthew Bolanos and John Forrest
References


L. Wan, W.K. Ng, X.H. Dang, P.S. Yu and K. Zhang (2009). Density-Based Clustering of Data Streams at Multiple Resolutions, *ACM Transactions on Knowledge Discovery from Data*, 3(3).

See Also

*animate_cluster, cl_agreement* in *clue*, *cluster.stats* in *fpc*, *silhouette* in *cluster*.

Examples

```r
stream <- DSD_Gaussians(k=3, d=2)

dstream <- DSC_DStream(gridsize=0.05, Cm=1.5)
update(dstream, stream, 500)
plot(dstream, stream)
# Evaluate micro-clusters
# Note: we use here only n=500 points for evaluation to speed up execution
evaluate(dstream, stream, measure=c("numMicro","numMacro","purity","crand", "SSQ"),
        n=100)

# DStream also provides macro clusters. Evaluate macro clusters with type="macro"
plot(dstream, stream, type="macro")
evaluate(dstream, stream, type="macro",
         measure=c("numMicro","numMacro","purity","crand", "SSQ"), n=100)

# Points are by default assigned to the closest micro clusters for evaluation.
# However, points can also be assigned to the closest macro-cluster using
# assign="macro".
evaluate(dstream, stream, type="macro", assign="macro",
         measure=c("numMicro","numMacro","purity","crand", "SSQ"), n=100)
```
# Evaluate an evolving data stream
stream <- DSD_Benchmark(1)
dstream <- DSD_DStream(gridsize=0.05, lambda=0.1)
evaluate_cluster(dstream, stream, type="macro", assign="micro",
               measure=c("numMicro","numMacro","purity","crand"),
               n=600, horizon=100)

## Not run:
# animate the clustering process
reset_stream(stream)
dstream <- DSD_DStream(gridsize=0.05, lambda=0.1)
animate_cluster(dstream, stream, horizon=100, n=5000,
               measure=c("crand"), type="macro", assign="micro",
               plot.args = list(type="both", xlim=c(0,1), ylim=c(0,1)))

## End(Not run)

desc = Assignment Data Points to Clusters

Description
Get the assignment of data points to clusters in a DSC (nearest-neighbor).

Usage
get_assignment(dsc, points, type=c("auto", "micro", "macro"),
               method="auto", ...)

Arguments
dsc The DSC object with the clusters for assignment.
points The points to be assigned as a data.frame.
type Use micro- or macro-clusters in DSC for assignment. Auto used the class of dsc
to decide.
method assignment method "model" uses the assignment model of the underlying algo-
               rithm (unassigned points return NA). "nn" performs nearest neighbor assignment
               using Euclidean distance. "auto" uses model if it is available and defaults to nn
               otherwise.
... Additional arguments are passed on.

Details
Each data point is assigned either using the original model’s assignment rule or Euclidean nearest
neighbor assignment.
get_centers

Value

A vector containing the assignment of each point. NA means that a data point was not assigned to a cluster.

Author(s)

Michael Hahsler

See Also

DSC

Examples

stream <- DSD_Gaussians(k=3, d=2, noise=.05)
dbstream <- DSC_DBSTREAM(r=.1)
update(dbstream, stream, n=100)

# find the assignment for the next 100 points to
# micro-clusters in dsc. This uses the model's assignemnt function
points <- get_points(stream, n=100)
a <- get_assignment(dbstream, points)
a

# show the MC assignment areas. Assigned points as blue circles and
# the unassigned points as red dots
plot(dbstream, stream, assignment= TRUE, type="none")
points(points[!is.na(a),], col="blue")
points(points[is.na(a),], col="red", pch=20)

# use nearest neighbor assignment instead
get_assignment(dbstream, points, method="nn")

get_centers

Get Cluster Centers from a DSC

Description

Gets the cluster centers (micro- or macro-clusters) from a DSC object.

Usage

get_centers(x, type=c("auto", "micro", "macro"), ...)

Arguments

x The DSC object the centers are being requested from.

type get centers for micro- or macro-clusters. Auto used the class of dsc to decide.

... Additional parameters.
get_copy

Value
A data.frame with the micro- or macro-cluster centers as rows.

Author(s)
Michael Hahsler

See Also
DSC

Examples
stream <- DSD_Gaussians(k=3, d=2)
dstream <- DSC_DStream(gridsize=.1)
update(dstream, stream, 500)
dstream

# getting the micro-cluster centers
get_centers(dstream)

# D-Stream also has macro-clusters
get_centers(dstream, type="macro")

---

get_copy  Create a Deep Copy of a DSC Object

Description
DSC objects contain reference classes or Java data structures (for MOA). Therefore, we provide a mechanism to create deep copies.

Usage
get_copy(x)

Arguments
x  The DSC object being copied.

Value
A deep copy of the original DSC.

Author(s)
Michael Hahsler
get_points

Description

Gets points from a DSD object.

Usage

get_points(x, n=1, outofpoints=c("stop", "warn", "ignore"), ...)

Arguments

x
The DSD object.

n
Request up to n points from the stream.

outofpoints
Action taken if less than n data points are available. The default is to stop with an error. For warn and ignore all available (possibly zero) points are returned.

... Additional parameters to pass to get_points() implementations.

Details

Each DSD object has a unique way for returning data points, but they all are called through the generic function, get_points(). This is done by using the S3 class system. See the man page for the specific DSD class on the semantics for each implementation of get_points().

Value

Returns a matrix of x$d columns and n rows.

Author(s)

Michael Hahsler

See Also

DSD

Examples

stream <- DSD_Gaussians()
get_points(stream, 100)
**get_weights**  
*Get Cluster Weights*

**Description**
Get the weights of the clusters in the DSC

**Usage**

```r
get_weights(x, type=c("auto", "micro", "macro"), scale=NULL, ...)
```

**Arguments**
- `x`: The DSC object the weights are being requested from.
- `type`: Return weights of micro- or macro-clusters in `x`. Auto uses the class of `x` to decide.
- `scale`: a range (from, to) to scale the weights. Returns by default the raw weights.
- `...`: Additional arguments are passed on.

**Details**
The cluster weights are typically a function of how many points were assigned to each cluster.

**Value**
A vector containing the weight of each micro-cluster or macro-cluster. Internally the call is delegated to the appropriate `get_microweights()` or `get_macroweights()` method.

**Author(s)**
Michael Hahsler

**See Also**
- `dsc`

**Examples**
```
stream <- DSD_Gaussians(k=3, d=2)
dstream <- DSC_DStream(gridsize=.1)
update(dstream, stream, 500)
dstream

# getting the micro-cluster weights
get_weights(dstream)

# D-Stream also has macro-clusters
get_weights(dstream, type="macro")
```
MGC

Moving Generator Cluster

Description

Creates an evolving cluster for a DSD_MG.

Usage

MGC_Static(density, center, parameter, shape = NULL)
MGC_Function(density, center, parameter, shape = NULL)
MGC_Random(density, center, parameter, randomness = 1, shape = NULL)
MGC_Noise(density, range)
MGC_Linear(dimension = 2, keyframelist = NULL, shape = NULL)
keyframe(time, density, center, parameter, reset = FALSE)
add_keyframe(x, time, density, center, parameter, reset = FALSE)
get_keyframes(x)
remove_keyframe(x, time)

Arguments

center A list that defines the center of the cluster. The list should have a length equal to the dimensionality. For MGC_Function, this list consists of functions that define the movement of the cluster. For MGC_Random, this attribute defines the beginning location for the MGC before it begins moving.
density The density of the cluster. For MGC_Function, this attribute is a function and defines the density of a cluster at a given timestamp.
dimension Dimensionality of the data stream.
keyframelist A list of keyframes to initialize the MGC_Linear object with.
parameter Parameters for the shape. For the default shape MGC_Shape_Gaussian the parameter is the standard deviation, one per dimension. If a single value is specified then it is recycled for all dimensions.
randomness The maximum amount the cluster will move during one time step.
range The area in which the noise should appear.
reset Should the cluster reset to the first keyframe (time 0) after this keyframe is finished?
shape A function creating the shape of the cluster. It gets passed on the parameters argument from above. Available functions are MGC_Shape_Gaussian (the parameters are a vector containing standard deviations) and MGC_Shape_Block (parameters are the dimensions of the uniform block).
time The time stamp the keyframe should be located or which keyframe should be removed.
x An object of class MGC_Linear.
Details

An MGC describes a single cluster for use within an DSD_MG. There are currently four different MGCs that allow a user to express many different behaviors within a single data stream.

An MGC_Linear creates an evolving Gaussian cluster for a DSD_MG who’s behavior is determined by several keyframes. Keyframes can be added and removed.

An MGC_Function allows for a creation of a DSD_MG that is defined by functions of time.

An MGC_Random allows for a creation of a DSD_MG that moves randomly.

An MGC_Noise allows for a creation of noise within a DSD_MG.

Author(s)

Matthew Bolanos

See Also

DSD_MG for details on how to use an MGC within a DSD

Examples

```r
### Two static clusters
stream <- DSD_MG(dim=2,
    MGC_Static(den = 1, center=c(1, 0), par=.1),
    MGC_Static(den = 1, center=c(2, 0), par=.4, shape=MGC_Shape_Block)
)

plot(stream)

### Example of several MGC_Randoms
stream <- DSD_MG(dimension=2,
    MGC_Random(den = 100, center=c(1, 0), par=.1, rand=.1),
    MGC_Random(den = 100, center=c(2, 0), par=.4, shape=MGC_Shape_Block, rand=.1)
)

### Not run:
    animate_data(stream, 2500, xlim=c(0,3), ylim=c(-2,2), horizon=100)

### End(Not run)

### Example of several MGC_Functions
stream <- DSD_MG(dim = 2)

### block-shaped cluster moving from bottom-left to top-right increasing size
c1 <- MGC_Function(    density = function(t){100},
                       parameter = function(t){1*t},
                       center = function(t) c(t,t),
                       shape = MGC_Shape_Block )
add_cluster(stream,c1)
```
### cluster moving in a circle (default shape is Gaussian)
c2 <- MGC_Function(
    density = function(t)(25),
    parameter = function(t)(5),
    center= function(t) c(sin(t/10)*50+50, cos(t/10)*50+50)
)
add_cluster(stream,c2)

# Not run:
animate_data(stream,10000,xlim=c(-20,120),ylim=c(-20,120), horizon=100)

# End(Not run)

### Example of several MGC_Linears: A single cluster splits at time 50 into two.
### Note that c2 starts at time=50!
stream <- DSD_MG(dim = 2)
c1 <- MGC_Linear(dim = 2)
add_keyframe(c1, time=1, dens=50, par=5, center=c(0,0))
add_keyframe(c1, time=50, dens=50, par=5, center=c(50,50))
add_keyframe(c1, time=100, dens=50, par=5, center=c(50,100))
add_cluster(stream,c1)

c2 <- MGC_Linear(dim = 2, shape=MGC_Shape_Block)
add_keyframe(c2, time=50, dens=25, par=c(10,10), center=c(50,50))
add_keyframe(c2, time=100, dens=25, par=c(30,30), center=c(100,50))
add_cluster(stream,c2)

# Not run:
animate_data(stream,5000,xlim=c(0,100),ylim=c(0,100), horiz=100)

# End(Not run)

### two fixed and a moving cluster
stream <- DSD_MG(dim = 2,
    MGC_Static(dens=1, par=.1, center=c(0,0)),
    MGC_Static(dens=1, par=.1, center=c(1,1)),
    MGC_Linear(dim=2,list(
        keyframe(time = 0, dens=1, par=.1, center=c(0,0)),
        keyframe(time = 1000, dens=1, par=.1, center=c(1,1)),
        keyframe(time = 2000, dens=1, par=.1, center=c(0,0), reset=TRUE)
    )))
noise <- MGC_Noise(dens=.1, range=rbind(c(-.2,1.2),c(-.2,1.2)))
add_cluster(stream,noise)

# Not run:
animate_data(stream, n=2000*3.1, xlim=c(-2,1.2), ylim=c(-2,1.2), horiz=200)

# End(Not run)
**microToMacro**

*Translate Micro-cluster IDs to Macro-cluster IDs*

**Description**

Translates micro-cluster ids into macro-cluster ids for a DSC_Macro object.

**Usage**

```r
microToMacro(x, micro=NULL)
```

**Arguments**

- `x`: a DSC_Macro object that also contains information about micro-clusters.
- `micro`: A vector with micro-cluster ids. If `NULL` then the assignments for all micro-clusters in `x` are returned.

**Value**

A vector of the same length as `micro` with the macro-cluster ids.

**Author(s)**

Michael Hahsler

**See Also**

DSC_Macro

**Examples**

```r
stream <- DSD_Gaussians(k=3, d=2, noise=0.05, p=c(.2,.4,.6))

# recluster a micro-clusters
micro <- DSC_DStream(gridsize=0.05)
update(micro, stream, 500)

macro <- DSC_Kmeans(k=3)
recluster(macro, micro)

# translate all micro-cluster ids
microToMacro(macro)

# plot some data points in gray
plot(stream, col="gray", cex=.5, xlim=c(0,1), ylim=c(0,1))
# add micro-clusters and use the macro-cluster ids as color and weights as size
points(get_centers(macro, type="micro"),
       col=microToMacro(macro),
       cex=get_weights(macro, type="micro", scale=c(.5,3)))
```
# add macro-cluster centers (size is weight)
points(get_centers(macro, type="macro"),
    cex = get_weights(macro, type="macro", scale=c(2,5)),
    pch=3,lwd=3, col=1:3)

## nclusters

### Description

Returns the number of micro-clusters from the DSC object.

### Usage

```r
nclusters(x, type=c("auto", "micro", "macro"), ...)
```

### Arguments

- `x` A DSC object.
- `type` Return the number of micro- or macro-clusters in DSC. Auto used the class of `dsc` to decide.
- `...` Additional arguments are passed on.

### Value

An integer; the number of micro- or macro-clusters in the clustering.

### Author(s)

Michael Hahsler

### See Also

[DSC](#)

### Examples

# setting up the objects
stream <- DSD_Gaussians(k=3)
dstream <- DSC_DStream(gridsize=0.1)
update(dstream, stream, 500)

# retrieving the results
get_centers(dstream)
nclusters(dstream)
Description

Methods to plot data stream data and clusterings.

Usage

```r
## S3 method for class 'DSD'
plot(x, n = 500, col = NULL,
     pch = NULL, ..., method = "pairs", dim = NULL, alpha = 0.6)
## S3 method for class 'DSC'
plot(x, dsd = NULL, n = 500,
     col_points = NULL, col_clusters = c("red", "blue"),
     weights = TRUE, scale = c(1,5), cex = 1, pch = NULL, method = "pairs",
     dim = NULL, type = c("auto", "micro", "macro", "both"),
     assignment = FALSE, ...)
```

Arguments

- `x`: the DSD or DSC object to be plotted.
- `dsd`: a DSD object to plot the data in the background.
- `n`: number of plots taken from the dsd to plot.
- `col, col_points, col_clusters`: colors used for plotting.
- `weights`: the size of the symbols for micro- and macro-clusters represents its weight.
- `scale`: range for the symbol sizes used.
- `cex`: size factor for symbols.
- `pch`: symbol type.
- `method`: method used for plotting: "pairs" (pairs plot), "scatter" (scatter plot) or "pc" (plot first 2 principal components).
- `dim`: an integer vector with the dimensions to plot. If NULL then for methods "pairs" and "pc" all dimensions are used and for "scatter" the first two dimensions are plotted.
- `alpha`: alpha shading used to plot the points.
- `type`: Plot micro- or macro-clusters. Auto used the class of dsc to decide.
- `assignment`: logical; show assignment area of micro-clusters.
- `...`: further arguments are passed on to plot or pairs in `graphics`.

Author(s)

Michael Hahsler
See Also

DSC, DSD

Examples

stream <- DSD_Gaussians(k=3, d=3)

## plot data
plot(stream, n=500)
plot(stream, method="pc", n=500)
plot(stream, method="scatter", dim=c(1,3), n=500)

## create and plot micro-clusters
dstream <- DSC_DStream(gridsize=0.1)
update(dstream, stream, 500)
plot(dstream)

## plot with data, projected on the first two principal components
## and dimensions 2 and 3
plot(dstream, stream)
plot(dstream, stream, method="pc")
plot(dstream, stream, dim=c(2,3))

## plot micro and macro-clusters
plot(dstream, stream, type="both")

---

**prune_clusters**

**Prune Clusters from a Clustering**

**Description**

Creates a (static) copy of a clustering where a fraction of the weight or the number of clusters with the lowest weights were pruned.

**Usage**

prune_clusters(dsc, threshold= 0.05, weight = TRUE)

**Arguments**

- `dsc`: The DSC object to be pruned.
- `threshold`: The numeric vector of probabilities for the quantile.
- `weight`: should a fraction of the total weight in the clustering be pruned? Otherwise a fraction of clusters is pruned.

**Value**

Returns an object of class DSC_Static.
Description

Use a macro clustering algorithm to recluster micro-clusters into a final clustering.

Usage

recluster(macro, micro, type="auto", ...)

Arguments

- **macro**: a macro clustering algorithm (class "DSC_Macro")
- **micro**: a DSC object containing micro-clusters.
- **type**: controls which clustering is used from dsc (typically micro-clusters).
- **...**: additional arguments passed on.

Details

Takes centers and weights of the micro-clusters and applies the macro clustering algorithm.

Value

The object macro is altered and contains the clustering.
reset_stream

**Author(s)**

Michael Hahsler

**Examples**

```r
set.seed(0)
### create a data stream and a micro-clustering
stream <- DSD_Gaussians(k=3, d=3)

sample <- DSC_Sample(k=50)
update(sample, stream, 500)
sample

### recluster using k-means
kmeans <- DSC_Kmeans(k=3)
recluster(kmeans, sample)

### plot clustering
plot(kmeans, stream, main="Macro-clusters (Sampling + k-means")
```

---

**reset_stream**  
*Reset a Data Stream to its Beginning*

**Description**

Resets the counter in a DSD object to the beginning or any other position in the stream.

**Usage**

```r
reset_stream(dsd, pos = 1)
```

**Arguments**

- `dsd`: An object of class a subclass of DSD which implements a reset function.
- `pos`: Position in the stream (the beginning of the stream is position 1).

**Details**

Resets the counter of the stream object. For example, forDSD_Memory, the counter stored in the environment variable is moved back to 1. For DSD_ReadCSV objects, this is done by calling `seek()` on the underlying connection.

**Author(s)**

Michael Hahsler

**See Also**

DSD_ReadCSV, DSD_MG, DSD_ScaleStream, DSD_Memory
Examples

# initializing the objects
stream <- DSD_Gaussians(k=3, d=2)
replayer <- DSD_Memory(stream, 100)
replayer

p <- get_points(replayer, 50)
replayer

# reset replayer to the begining of the stream
reset_stream(replayer)
replayer

# set replayer to position 21
reset_stream(replayer, pos=21)
replayer

save

---

Save and Read DSC Objects

Description

Save and Read DSC objects savely (serializes the underlying data structure). This also works for streamMOA DSC objects.

Usage

```r
saveDSC(object, file, ...)
readDSC(file)
```

Arguments

- `object` a DSC object.
- `file` filename.
- `...` further arguments.

Author(s)

Michael Hahsler

See Also

`saveRDS` and `readRDS`. 
Examples

stream <- DSD_Gaussians(k = 3, noise = 0.05)

# create clusterer with r = 0.05
dbstream1 <- DSC_DBSTREAM(r = .05)
update(dbstream1, stream, 1000)
dbstream1

saveDSC(dbstream1, file="dbstream.Rds")

dbstream2 <- readDSC("dbstream.Rds")
dbstream2

## cleanup
unlink("dbstream.Rds")

update

Update a Data Stream Clustering Model

Description

Update a clustering model by clustering a number of input points from a data stream into a clustering object.

Usage

## S3 method for class 'DSC_R'
update(object, dsd, n = 1, verbose = FALSE, block=10000L, ...)

## S3 method for class 'DSC_TwoStage'
update(object, dsd, n = 1, verbose = FALSE,
       block=10000L, ...)

## S3 method for class 'DSC_Sample'
update(object, dsd, n = 1, verbose = FALSE, ...)

## S3 method for class 'DSC_Window'
update(object, dsd, n = 1, verbose = FALSE, ...)

Arguments

object an object of a subclass of DST (data stream mining task).
dsd     a DSD object (data stream).
n       number of points to cluster.
verbose report progress.
block  maximal number of data points passed on at once to the algorithm. This only is used since R loops are very slow.
...   extra arguments for clusterer.
**write_stream**

**Details**

update takes n times a single data points out of the DSD updates the model in object. Note that update directly modifies the object (which is a reference class) and thus the result does not need to be reassigned to the object name.

**Value**

The updated model is returned invisibly for reassignment (however, this is not necessary).

To obtain the updated model for a DSC (data stream clustering model), call `get_centers()` on the DSC object.

**Author(s)**

Michael Hahsler

**See Also**

DSC, DSD, and animation for producing an animation of the clustering process.

**Examples**

```r
stream <- DSD_Gaussians(k=3)
dstream <- DSC_DStream(gridsize=.05)

update(dstream, stream, 500)
plot(dstream, stream)
```

---

**write_stream**

Write a Data Stream to a File

**Description**

Writes points from a data stream DSD object to a file or a connection.

**Usage**

```r
write_stream(dsd, file, n=100, block=100000L, class=FALSE, append = FALSE, sep=",", header=FALSE, row.names=FALSE, ...)
```

**Arguments**

- **dsd**: The DSD object that will generate the data points for output.
- **file**: A file name or a R connection to be written to.
- **n**: The number of data points to be written.
- **block**: Write stream in blocks to improve file I/O speed.
- **class**: Save the class/cluster labels of the points as the last column.
write_stream

- **sep**: The character that will separate attributes in a data point.
- **append**: Append the data to an existing file.
- **header**: A flag that determines if column names will be output (equivalent to `col_names` in `write.table()`).
- **row.names**: A flag that determines if row names will be output.
- **...**: Additional parameters that are passed to `write.table()`.

**Value**

There is no value returned from this operation.

**Author(s)**

Michael Hahsler

**See Also**

`write.table`, `DSD`

**Examples**

```r
# creating data and writing it to disk
stream <- DSD_Gaussians(k=3, d=5)
write_stream(stream, file="data.txt", n=10, class=TRUE)

# file.show("data.txt")

# clean up
file.remove("data.txt")
```
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