

Package ‘clustra’

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Title Clustering Longitudinal Trajectories

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Description Clusters longitudinal trajectories over time (can be unequally spaced, unequal length time series and/or partially overlapping series) on a common time axis. Performs k-means clustering on a single continuous variable measured over time, where each mean is defined by a thin plate spline fit to all points in a cluster. Distance is MSE across trajectory points to cluster spline. Provides graphs of derived cluster splines, silhouette plots, and Adjusted Rand Index evaluations of the number of clusters. Scales well to large data with multicore parallelism available to speed computation.

LazyLoad yes

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clustra-package	<i>clustra-package</i>
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Description

Clusters medical trajectories (unequally spaced and unequal lengths) aligned by an intervention time. Performs k-means clustering, where each mean is a thin plate spline fit to all points in a cluster. Distance is MSE across trajectory points to cluster spline. Provides silhouette plots and Adjusted Rand Index evaluations of the number of clusters. Scales well to large data with multicore parallelism available to speed computation.

Author(s)

George Ostrouchov, David Gagnon, Hanna Gerlovin

allpair_RandIndex	<i>allpair_RandIndex: helper for replicated cluster comparison</i>
-------------------	--

Description

Runs [RandIndex](#) for all pairs of cluster results in its list input and produces a matrix for use by [rand_plot](#). Understands replicates within k values.

Usage

```
allpair_RandIndex(results)
```

Arguments

results	A list with each element packed internally by the clustra_rand function with elements: <ul style="list-style-type: none"> • k - number of clusters • rep - replicate number • deviance - final deviance • group - integer cluster assignments
---------	---

Value

A data frame with [RandIndex](#) for all pairs from trajectories results. The data frame names and its format is intended to be the input for [rand_plot](#). Note that all pairs is the lower triangle plus diagonal of an all-pairs symmetric matrix.

check_df	<i>Checks if non-empty groups have enough data for spline fit degrees of freedom.</i>
----------	---

Description

Checks if non-empty groups have enough data for spline fit degrees of freedom.

Usage

```
check_df(group, loss, data, maxdf)
```

Arguments

group	An integer vector of group membership for each id.
loss	A matrix with rows of computed loss values of each id across all models as columns.
data	A data.table with data. See trajectories .
maxdf	Fitting parameters. See trajectories .

Details

When a group has insufficient data for `maxdf`, its nearest model loss values are set to `Inf`, and new nearest model is assigned. The check repeats until all groups have sufficient data.

Value

Returns the vector of group membership of `id`'s either unchanged or changed to have sufficient data in non-zero groups.

clustra

Cluster longitudinal trajectories over time

Description

The usual top level function for clustering longitudinal trajectories. After initial setup, it calls [trajectories](#) to perform k-means clustering on continuous response measured over time, where each mean is defined by a thin plate spline fit to all points in a cluster. See `clustra_vignette.Rmd` for examples of use.

Usage

```
clustra(
  data,
  k,
  starts = c(1, 0),
  group = NULL,
  maxdf = 30,
  conv = c(10, 0),
  mcores = 1,
  verbose = FALSE
)
```

Arguments

<code>data</code>	Data frame or, preferably, also a <code>data.table</code> with response measurements, one response per observation. Required variables are (<code>id</code> , <code>time</code> , <code>response</code>). Other variables are ignored.
<code>k</code>	Number of clusters
<code>starts</code>	A vector of length two. See start_groups .
<code>group</code>	A vector of initial cluster assignments for unique <code>id</code> 's in <code>data</code> . Normally, this is <code>NULL</code> and good starts are provided by start_groups .
<code>maxdf</code>	Fitting parameters. See trajectories .
<code>conv</code>	Fitting parameters. See trajectories .
<code>mcores</code>	See trajectories .
<code>verbose</code>	Logical to turn on more output during fit iterations.

Value

A list returned by [trajectories](#) plus one more element `ido`, giving the original id numbers.

Examples

```
set.seed(13)
data = gen_traj_data(n_id = c(50, 100), m_obs = 20, s_range = c(-365, -14),
                    e_range = c(0.5*365, 2*365))
cl = clustra(data, k = 2, maxdf = 20, conv = c(5, 0), verbose = TRUE)
tabulate(data$group)
tabulate(data$true_group)
```

clustra_rand

clustra_rand: Rand Index cluster evaluation

Description

Performs [trajectories](#) runs for several k and several random start replicates within k . Returns a data frame with a Rand Index comparison between all pairs of clusterings. This data frame is typically input to [rand_plot](#) to produce a heat map with the Adjusted Rand Index results.

Usage

```
clustra_rand(
  data,
  k,
  mcores,
  replicates = 10,
  maxdf = 30,
  conv = c(10, 0),
  save = FALSE,
  verbose = FALSE
)
```

Arguments

<code>data</code>	The data (see clustra description).
<code>k</code>	Vector of k values to try.
<code>mcores</code>	Number of cores for replicate parallelism via <code>mclapply</code> .
<code>replicates</code>	Number of replicates for each k .
<code>maxdf</code>	Fitting parameters. See <code>link{trajectories}</code> .
<code>conv</code>	Fitting parameters. See <code>link{trajectories}</code> .
<code>save</code>	Logical. When TRUE, save all results as file <code>results.Rdata</code> .
<code>verbose</code>	Logical. When TRUE, information about each run of <code>clustra</code> is printed.

Value

See [allpair_RandIndex](#).

clustra_sil	<i>clustra_sil: Prepare silhouette plot data for several k or for a previous clustra run</i>
-------------	--

Description

Performs [clustra](#) runs for several k and prepares silhouette plot data. Computes a proxy silhouette index based on distances to cluster centers rather than trajectory pairs. The cost is essentially that of running [clustra](#) for several k as this information is available directly from [clustra](#). Can also reuse a previous [clustra](#) run and produce data for a single silhouette plot.

Usage

```
clustra_sil(
  data,
  k = NULL,
  mcores = 1,
  maxdf = 30,
  conv = c(10, 0),
  save = FALSE,
  verbose = FALSE
)
```

Arguments

data	Either a data.frame (data parameter of trajectories) or the output from a clustra run. See Details.
k	Vector of k values to try. If output from clustra is the data parameter, k can be left NULL or set to the number of clusters used.
mcores	See trajectories .
maxdf	Fitting parameters. See trajectories .
conv	Fitting parameters. See trajectories .
save	Logical. When TRUE, save all results as file <code>clustra_sil.Rdata</code> .
verbose	Logical. When TRUE, information about each run of clustra is printed.

Details

When given the raw data as the first parameter (input data parameter of [trajectories](#)), k can also specify a vector of cluster numbers to run [clustra](#) and then produce silhouette plots for each of them. Alternatively, the input can be the output from a [clustra](#) run, in which case data for a single silhouette plot will be made without rerunning [clustra](#).

Value

Invisibly returns a list of length `length(k)`, where each element is a matrix with `nrow(data)` rows and three columns `cluster`, `neighbor`, `silhouette`. This list of matrices can be used to draw a silhouette plot.

delttime	<i>Timing function</i>
----------	------------------------

Description

Timing function

Usage

```
delttime(ltime = proc.time()["elapsed"], text = NULL)
```

Arguments

<code>ltime</code>	Result of last call to <code>delttime</code> .
<code>text</code>	Text to display along with elapsed time.

Value

"elapsed" component of current `proc.time`.

gendata	<i>gendata</i>
---------	----------------

Description

Generates data for up to three trajectory clusters

Usage

```
gendata(n_id, m_obs, s_range, e_range, min_obs, reference, noise)
```

Arguments

<code>n_id</code>	See parameters of gen_traj_data .
<code>m_obs</code>	See parameters of gen_traj_data .
<code>s_range</code>	See parameters of gen_traj_data .
<code>e_range</code>	See parameters of gen_traj_data .
<code>min_obs</code>	See parameters of gen_traj_data .
<code>reference</code>	See parameters of gen_traj_data .
<code>noise</code>	See parameters of gen_traj_data .

Details

Time support of each id is at least $s \dots 0 \dots e$, where s is in `s_range` and e is in `e_range`.

Value

A list of length `sum(n_id)`, where each element is a matrix output by `oneid`.

gen_traj_data

Data Generators

Description

Generates a collection of longitudinal responses with possibly varying lengths and varying numbers of observations. Support is `start \dots 0 \dots end`, where `start` \sim `uniform(s_range)` and `end` \sim `uniform(e_range)`, so that all trajectories are aligned at 0 but can start and end at different times. Zero is the intervention time.

Usage

```
gen_traj_data(
  n_id,
  m_obs,
  s_range,
  e_range,
  reference = 100,
  noise = c(0, abs(reference/20)),
  min_obs = 3
)
```

Arguments

<code>n_id</code>	Vector whose length is the number of clusters, giving the number of id's to generate in each cluster.
<code>m_obs</code>	Mean number of observation per id. Provides lambda parameter in <code>rpois</code> .
<code>s_range</code>	A vector of length 2, giving the min and max limits of uniformly generated start observation time.
<code>e_range</code>	A vector of length 2, giving the min and max limits of uniformly generated end observation time.
<code>reference</code>	A nominal response value (for example, blood pressure is near 100, which is the default)
<code>noise</code>	Vector of length 2 giving the <i>mean</i> and <i>sd</i> of added $N(\text{mean}, \text{sd})$ noise.
<code>min_obs</code>	Minimum number of observations in addition to zero time observation.

Value

A data table with one response per row and four columns: id, time, response, and true_group.

Details

Generate longitudinal data for a response variable. Trajectories start at time uniformly distributed in s_range and end at time uniformly distributed in e_range. Number of observations in a trajectory is Poisson(m_obs). The result is a number of trajectories, all starting at time 0, with different time spans, and with independently different numbers of observations within the time spans. Each trajectory follows a randomly selected response function with added N(mean, sd) error.

Examples

```
data = gen_traj_data(n_id = c(50, 100), m_obs = 20, s_range = c(-365, -14),
                    e_range = c(0.5*365, 2*365))
head(data)
tail(data)
```

mse_g

Loss functions

Description

mse_g() Computes mean-squared error. mxe_g() Computes maximum absolute error.

Usage

```
mse_g(pred, id, response)
```

```
mxe_g(pred, id, response)
```

Arguments

pred	Vector of predicted values.
id	Integer vector of group assignments.
response	Vector of response values.

Value

A numeric value. For mse_g(), returns the mean-squared error. For mxe_g(), returns the maximum absolute error.

oneid	<i>Generates data for one id</i>
-------	----------------------------------

Description

Generates data for one id

Usage

```
oneid(id, n_obs, type, start, end, smin, emax, reference, noise)
```

Arguments

id	A unique integer.
n_obs	An integer number of observations to produce.
type	Response type, 1 is constant, 2 is a sin curve portion, and 3 is a sigmoid portion.
start	Negative integer giving time of first observation.
end	Positive integer giving time of last observation.
smin	The smallest possible start value among all ids. Currently not used.
emax	The largest possible end value among all ids. Used to scale sin and sigmoid support.
reference	A response value for constant response. Also used to scale sin and sigmoid responses.
noise	Standard deviation of zero mean Gaussian noise added to response functions.

Value

An n_obs by 4 matrix with columns id, time, response, true_group.

pred_g	<i>Function to predict for new data based on fitted gam object.</i>
--------	---

Description

Function to predict for new data based on fitted gam object.

Usage

```
pred_g(tps, newdata)
```

Arguments

tps	Output structure of bam .
newdata	See clustra description of data parameter.

Value

A numeric vector of predicted values corresponding to rows of newdata. If gam object is NULL, NULL is returned instead.

rand_plot	<i>Matrix plot of Rand Index comparison of replicated clusters</i>
-----------	--

Description

Matrix plot of Rand Index comparison of replicated clusters

Usage

```
rand_plot(rand_pairs, name = NULL)
```

Arguments

rand_pairs	A data frame result of allpair_RandIndex
name	Character string file name for pdf plot. If omitted or NULL, plot will render to current graphics device.

Value

Invisible. Full path name of file with plot.

Author(s)

Wei-Chen Chen and George Ostrouchov

References

Wei-chen Chen, George Ostrouchov, David Pugmire, Prabhat, and Michael Wehner. 2013. A Parallel EM Algorithm for Model-Based Clustering Applied to the Exploration of Large Spatio-Temporal Data. *Technometrics*, 55:4, 513-523.

Sorts replicates within cluster K Assumes K starts from 2

start_groups	<i>Function to assign starting groups.</i>
--------------	--

Description

If only one start, a random assignment is done. If more than one start, picks tps fit with smallest deviance after one iteration among random starts. Choosing from samples increases diversity of fits (sum of distances between group fits). Then classifies all ids based on fit from best sample.

Usage

```
start_groups(data, k, starts, maxdf, conv, mcores = 1, verbose = FALSE)
```

Arguments

data	Data.table with response measurements, one per observation. Column names are id, time, response, group. Note that ids are assumed sequential starting from 1. This affects expanding group numbers to ids.
k	Number of clusters (groups).
starts	A vector of length two, giving the number of start values to try and the number of ids per cluster to evaluate the starts (If the number of ids is less than 1, use all data and do not subset data for starts.). The default is <code>c(1, 0)</code> , meaning that one random start is used with all the data. The following are experimental at this time: If more than one start is requested, the most diverse after one trajectories iteration on a sample of data is used. Diversity is measured as sum of pairwise distances between models on a time grid of $2 \times \text{maxdf}$ points.
maxdf	Fitting parameters. See trajectories .
conv	Fitting parameters. See trajectories .
mcores	See trajectories .
verbose	Turn on more output for debugging.

Value

An integer vector corresponding to unique ids, giving group number assignments.

tps_g	<i>Fits a thin plate spline to a single group with bam.</i>
-------	---

Description

Fits a thin plate spline to a single group (one list element) in data with [bam](#). Uses data from only on group rather than a zero weights approach. Zero weights would result in incorrect crossvalidation sampling.

Usage

```
tps_g(g, data, maxdf, nthreads)
```

Arguments

<code>g</code>	Integer group number.
<code>data</code>	A list of group-separated data using <code>lapply</code> with <code>data.table::copy(data[group == g])</code> from original data in clustra description.
<code>maxdf</code>	See trajectories description.
<code>nthreads</code>	Controls bam threads.

Value

Returns an object of class "gam". See [bam](#) value. If group data has zero rows, NULL is returned instead.

<code>trajectories</code>	<i>Cluster longitudinal trajectories over time.</i>
---------------------------	---

Description

Performs k-means clustering on continuous response measured over time, where each mean is defined by a thin plate spline fit to all points in a cluster. Typically, this function is called by [clustra](#).

Usage

```
trajectories(
  data,
  k,
  group,
  maxdf,
  conv = c(10, 0),
  mcores = 1,
  verbose = FALSE
)
```

Arguments

<code>data</code>	Data table or data frame with response measurements, one per observation. Column names are <code>id</code> , <code>time</code> , <code>response</code> , <code>group</code> . Note that <code>ids</code> must be sequential starting from 1. This affects expanding group numbers to <code>ids</code> .
<code>k</code>	Number of clusters (groups)
<code>group</code>	Vector of initial group numbers corresponding to <code>ids</code> .

maxdf	Integer. Basis dimension of smooth term. See <code>s</code> function parameter <code>k</code> , in package <code>mgcv</code> .
conv	A vector of length two, <code>c(iter, minchange)</code> , where <code>iter</code> is the maximum number of EM iterations and <code>minchange</code> is the minimum percentage of subjects changing group to continue iterations. Setting <code>minchange</code> to zero continues iterations until no more changes occur or <code>maxiter</code> is reached.
mccores	Integer number of cores to use by <code>mclapply</code> sections. Parallelization is over <code>k</code> , the number of clusters.
verbose	Logical, whether to produce debug output.

Value

A list with components

- `deviance` - The final deviance in each cluster added across clusters.
- `group` - Integer vector of group assignments corresponding to unique `ids`.
- `loss` - Numeric matrix with rows corresponding to unique `ids` and one column for each cluster. Each entry is the mean squared loss for the data in the `id` relative to the cluster model.
- `k` - An integer giving the requested number of clusters.
- `k_cl` - An integer giving the converged number of clusters. Can be smaller than `k` when some clusters become too small for degrees of freedom during convergence.
- `data_group` - An integer vector, giving group assignment as expanded into all `id` time points.
- `tps` - A list with `k_cl` elements, each an object returned by the `mgcv::bam` fit of a cluster thin plate spline model.
- `iterations` - An integer giving the number of iterations taken.
- `counts` - An integer vector giving the number of `ids` in each cluster.
- `counts_df` - An integer vector giving the total number of observations in each cluster (sum of the number of observations for `ids` belonging to the cluster).
- `changes` - An integer, giving the number of `ids` that changed clusters in the last iteration. This is zero if converged.

Author(s)

George Ostrouchov and David Gagnon

traj_rep

Function to run trajectories inside mclapply with one core.

Description

Function to run trajectories inside `mclapply` with one core.

Usage

```
traj_rep(group, data, k, maxdf, conv)
```

Arguments

group	Vector of starting group values for unique id's.
data	The data (see clustra description).
k	Integer number of clusters.
maxdf	Fitting parameters. See trajectories .
conv	Fitting parameters. See trajectories .

Value

See return of [trajectories](#).

xit_report	<i>xit_report</i>
------------	-------------------

Description

Examines trajectories output to name what was concluded, such as convergence, maximum iterations reached, a zero cluster, etc. Multiple conclusions are possible as not all are mutually exclusive.

Usage

```
xit_report(cl, maxdf, conv)
```

Arguments

cl	Output structure from trajectories function
maxdf	Fitting parameters. See trajectories .
conv	Fitting parameters. See trajectories .

Value

NULL or a character vector of exit criteria satisfied.

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