

# Package: stops (via r-universe)

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**Title** Structure Optimized Proximity Scaling

**Version** 1.8-2

**Maintainer** Thomas Rusch <thomas.rusch@wu.ac.at>

**Description** Methods that use flexible variants of multidimensional scaling (MDS) which incorporate parametric nonlinear distance transformations and trade-off the goodness-of-fit fit with structure considerations to find optimal hyperparameters, also known as structure optimized proximity scaling (STOPS) (Rusch, Mair & Hornik, 2023, <doi:10.1007/s11222-022-10197-w>). The package contains various functions, wrappers, methods and classes for fitting, plotting and displaying different 1-way MDS models with ratio, interval, ordinal optimal scaling in a STOPS framework. These cover essentially the functionality of the package smacofx, including Torgerson (classical) scaling with power transformations of dissimilarities, SMACOF MDS with powers of dissimilarities, Sammon mapping with powers of dissimilarities, elastic scaling with powers of dissimilarities, spherical SMACOF with powers of dissimilarities, (ALSCAL) s-stress MDS with powers of dissimilarities, r-stress MDS, MDS with powers of dissimilarities and configuration distances, elastic scaling powers of dissimilarities and configuration distances, Sammon mapping powers of dissimilarities and configuration distances, power stress MDS (POST-MDS), approximate power stress, Box-Cox MDS, local MDS, Isomap, curvilinear component analysis (CLCA), curvilinear distance analysis (CLDA) and sparsified (power) multidimensional scaling and (power) multidimensional distance analysis (experimental models from smacofx influenced by CLCA). All of these models can also be fit by optimizing over hyperparameters based on goodness-of-fit fit only (i.e., no structure considerations). The package further contains functions for optimization, specifically the adaptive Luus-Jaakola algorithm and a wrapper for Bayesian optimization with treed Gaussian process with jumps to linear models, and functions for various c-structuredness indices.

**Depends** R (>= 3.5.0), smacofx

**Imports** acepack, clue, cmaes, cordillera, dfoptim, DiceOptim,  
DiceKriging, energy, minerva, nloptr, pomp, pso, registry,  
scagnostics, smacof, tgp, vegan

**Enhances** stats

**Suggests** R.rsp

**License** GPL-2 | GPL-3

**LazyData** true

**URL** <https://r-forge.r-project.org/projects/stops/>

**VignetteBuilder** R.rsp

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**NeedsCompilation** no

**Author** Thomas Rusch [aut, cre]  
(<<https://orcid.org/0000-0002-7773-2096>>), Patrick Mair [aut]  
(<<https://orcid.org/0000-0003-0100-6511>>), Kurt Hornik [ctb]  
(<<https://orcid.org/0000-0003-4198-9911>>)

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BankingCrisesDistances

*Banking Crises Distances*


---

### Description

Matrix of Jaccard distances between 70 countries (Hungary and Greece were combined to be the same observation) based on their binary time series of having had a banking crises in a year from 1800 to 2010 or not. See `data(bankingCrises)` in package `Ecdat` for more info. The last column is Reinhart & Rogoffs classification as a low (3), middle- (2) or high-income country (1).

### Format

A 69 x 70 matrix.

### Source

`data(bankingCrises)` in library(`Ecdat`)

---

biplotmds.stops

*S3 method for stops objects*


---

### Description

S3 method for stops objects

### Usage

```
## S3 method for class 'stops'
biplotmds(object, extvar, scale = TRUE)
```

### Arguments

<code>object</code>	An object of class <code>stops</code>
<code>extvar</code>	Data frame with external variables.
<code>scale</code>	if <code>'TRUE'</code> external variables are standardized internally.

### Details

If a model for individual differences is provided, the external variables are regressed on the group stimulus space configurations. For objects returned from `'biplotmds'` we use the `plot` method in [biplotmds](#). In the biplot called with `plot()` only the relative length of the vectors and their direction matters. Using the `vecscale` argument in `plot()` the user can control for the relative length of the vectors. If `'vecscale = NULL'`, the `'vecscale()'` function from the `'candisc'` package is used which tries to automatically calculate the scale factor so that the vectors approximately fill the same space as the configuration. In this method `vecscale` should usually be smaller than the one used in `smacof` by a factor of 0.1.

**Value**

Returns an object belonging to classes 'mlm' and 'mdsbi'. See 'lm' for details. R2vec: Vector containing the R2 values. See also [biplotmds](#) for the plot method.

---

bootmds.stops	<i>MDS Bootstrap for stops objects</i>
---------------	--

---

**Description**

Performs a bootstrap on an MDS solution. It works for derived dissimilarities only, i.e. generated by the call `dist(data)`. The original data matrix needs to be provided, as well as the type of dissimilarity measure used to compute the input dissimilarities (note we cannot as of yet have any dissimilarity matrix).

**Usage**

```
## S3 method for class 'stops'
bootmds(
  object,
  data,
  method.dat = "pearson",
  nrep = 100,
  alpha = 0.05,
  verbose = FALSE,
  ...
)
```

**Arguments**

<code>object</code>	Object of class stops or pcops.
<code>data</code>	Initial data (before dissimilarity computation).
<code>method.dat</code>	Dissimilarity computation used as MDS input. This must be one of "pearson", "spearman", "kendall", "euclidean", "maximum", "manhattan", "canberra", "binary".
<code>nrep</code>	Number of bootstrap replications.
<code>alpha</code>	Alpha level for confidence ellipsoids.
<code>verbose</code>	If 'TRUE', bootstrap index is printed out.
<code>...</code>	Additional arguments needed for dissimilarity computation as specified in <a href="#">sim2diss</a> .

**Details**

In order to examine the stability solution of an MDS, a bootstrap on the raw data can be performed. This results in confidence ellipses in the configuration plot. The ellipses are returned as list which allows users to produce (and further customize) the plot by hand. See [bootmds](#) for more.

**Value**

An object of class 'smacofboot', see [bootmds](#). With values

- cov: Covariances for ellipse computation
- bootconf: Configurations bootstrap samples
- stressvec: Bootstrap stress values
- bootci: Stress bootstrap percentile confidence interval
- spp: Stress per point (based on stress.en)
- stab: Stability coefficient

**Examples**

```

dats <- na.omit(PVQ40[,1:5])
diss <- dist(t(dats)) ## Euclidean distances
fit <- stops(diss,loss="rstress",itmax=5,lower=0.2,upper=3)
set.seed(123)
resboot <- bootmds(fit, dats, method.dat = "euclidean", nrep = 2)
resboot

```

---

coef.stops

*S3 coef method for stops objects*


---

**Description**

S3 coef method for stops objects

**Usage**

```

## S3 method for class 'stops'
coef(object, ...)

```

**Arguments**

object	object of class stops
...	additional arguments

**Value**

a vector of hyperparameters theta

---

c_association	<i>c-association calculates the c-association based on the maximal information coefficient We define c-association as the aggregated association between any two columns in confs</i>
---------------	---

---

## Description

c-association calculates the c-association based on the maximal information coefficient We define c-association as the aggregated association between any two columns in confs

## Usage

```
c_association(
  confs,
  aggr = NULL,
  alpha = 0.6,
  C = 15,
  var.thr = 1e-05,
  zeta = NULL
)
```

## Arguments

confs	a numeric matrix or data frame
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.
alpha	an optional number of cells allowed in the X-by-Y search-grid. Default value is 0.6
C	an optional number determining the starting point of the X-by-Y search-grid. When trying to partition the x-axis into X columns, the algorithm will start with at most C X clumps. Default value is 15.
var.thr	minimum value allowed for the variance of the input variables, since mine can not be computed in case of variance close to 0. Default value is 1e-5.
zeta	integer in [0,1] (?). If NULL (default) it is set to 1-MIC. It can be set to zero for noiseless functions, but the default choice is the most appropriate parametrization for general cases (as stated in Reshef et al). It provides robustness.

## Value

a numeric value; association (aggregated maximal information coefficient MIC, see [mine](#))

## Examples

```
x<-seq(-3,3,length.out=200)
y<-sqrt(3^2-x^2)
z<- sin(y-x)
```

```
confs<-cbind(x,y,z)
c_association(confs)
```

---

c_clumpiness	<i>c-clumpiness</i>
--------------	---------------------

---

**Description**

Measures the c-clumpiness structure

**Usage**

```
c_clumpiness(conf, aggr = NULL)
```

**Arguments**

- conf            A numeric matrix.
- aggr           the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; clumpiness (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacof::smacofSym(delts)$conf
plot(conf,pch=19,asp=1)
c_clumpiness(conf)
```

---

c_clusteredness	<i>c-clusteredness calculates c-clusteredness as the OPTICS cordillera. The higher the more clustered.</i>
-----------------	--

---

**Description**

c-clusteredness calculates c-clusteredness as the OPTICS cordillera. The higher the more clustered.



**Usage**

```

c_clusteredness(
  confs,
  voidarg = NULL,
  minpts = 2,
  q = 2,
  epsilon = 2 * max(dist(confs)),
  distmeth = "euclidean",
  dmax = NULL,
  digits = 10,
  scale = 0,
  ...
)

```

**Arguments**

confs	a numeric matrix or a dist object
voidarg	a placeholder to allow to pass NULL as strucpar and not interfere with the other arguments
minpts	The minimum number of points that must make up a cluster in OPTICS (corresponds to k in the paper). It is passed to <a href="#">optics</a> where it is called minPts. Defaults to 2.
q	The norm used for the Cordillera. Defaults to 2.
epsilon	The epsilon parameter for OPTICS (called epsilon_max in the paper). Defaults to 2 times the maximum distance between any two points.
distmeth	The distance to be computed if X is not a symmetric matrix or a dist object (otherwise ignored). Defaults to Euclidean distance.
dmax	The winsorization value for the highest allowed reachability. If used for comparisons between different configurations this should be supplied. If no value is supplied, it is NULL (default); then dmax is taken from the data as the either epsilon or the largest reachability, whatever is smaller.
digits	The precision to round the raw Cordillera and the norm factor. Defaults to 10.
scale	Should X be scaled if it is an asymmetric matrix or data frame? Can take values TRUE or FALSE or a numeric value. If TRUE or 1, standardisation is to mean=0 and sd=1. If 2, no centering is applied and scaling of each column is done with the root mean square of each column. If 3, no centering is applied and scaling of all columns is done as X/max(standard deviation(allcolumns)). If 4, no centering is applied and scaling of all columns is done as X/max(rmsq(allcolumns)). If FALSE, 0 or any other numeric value, no standardisation is applied. Defaults to 0.
...	Additional arguments to be passed to <code>cordillera::cordillera</code>

**Value**

a numeric value; clusteredness (see [cordillera](#))

## Examples

```
delts<-smacof::kinshipdelta
dis<-smacofSym(delts)$confdist
c_clusteredness(dis,minpts=3)
```

---

c_complexity	<i>c-complexity</i> Calculates the c-complexity based on the minimum cell number We define c-complexity as the aggregated minimum cell number between any two columns in confs This is one of few c-structuredness indices not between 0 and 1, but can be between 0 and (theoretically) infinity
--------------	---

---

## Description

c-complexity Calculates the c-complexity based on the minimum cell number We define c-complexity as the aggregated minimum cell number between any two columns in confs This is one of few c-structuredness indices not between 0 and 1, but can be between 0 and (theoretically) infinity

## Usage

```
c_complexity(
  confs,
  aggr = NULL,
  alpha = 1,
  C = 15,
  var.thr = 1e-05,
  zeta = NULL
)
```

## Arguments

confs	a numeric matrix or data frame
aggr	the aggregation function for configurations of more than two dimensions. Defaults to min.
alpha	an optional number of cells allowed in the X-by-Y search-grid. Default value is 1
C	an optional number determining the starting point of the X-by-Y search-grid. When trying to partition the x-axis into X columns, the algorithm will start with at most C X clumps. Default value is 15.
var.thr	minimum value allowed for the variance of the input variables, since mine can not be computed in case of variance close to 0. Default value is 1e-5.
zeta	integer in [0,1] (?). If NULL (default) it is set to 1-MIC. It can be set to zero for noiseless functions, but the default choice is the most appropriate parametrization for general cases (as stated in Reshef et al.). It provides robustness.

**Value**

a numeric value; complexity (aggregated minimum cell number MCN, see [mine](#))

**Examples**

```
x<-seq(-3,3,length.out=200)
y<-sqrt(3^2-x^2)
z<- sin(y-x)
confs<-cbind(x,y,z)
c_complexity(confs)
```

---

c_convexity	<i>c-convexity</i>
-------------	--------------------

---

**Description**

Measures the c-convexity structure

**Usage**

```
c_convexity(conf, aggr = NULL)
```

**Arguments**

conf	A numeric matrix.
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; convexity (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacof::smacofSym(delts)$conf
plot(conf,pch=19,asp=1)
c_convexity(conf)
```

---

c_dependence	<i>c-dependence calculates c-dependence as the aggregated distance correlation of each pair if nonidentical columns</i>
--------------	---

---

### Description

c-dependence calculates c-dependence as the aggregated distance correlation of each pair if non-identical columns

### Usage

```
c_dependence(confs, aggr = NULL, index = 1)
```

### Arguments

confs	a numeric matrix or data frame
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.
index	exponent on Euclidean distance, in (0,2]

### Value

a numeric value; dependence (aggregated distance correlation)

### Examples

```
x<-1:10
y<-2+3*x+rnorm(10)
confs<-cbind(x,y)
c_dependence(confs,1.5)
```

---

c_faithfulness	<i>c-faithfulness calculates the c-faithfulness based on the index by Chen and Buja 2013 (M_adj) with equal input neighbourhoods</i>
----------------	--

---

### Description

c-faithfulness calculates the c-faithfulness based on the index by Chen and Buja 2013 (M\_adj) with equal input neighbourhoods

### Usage

```
c_faithfulness(confs, obsdiss, k = 3, ...)
```

**Arguments**

confs            a numeric matrix or a dist object  
 obsdiss        a symmetric numeric matrix or a dist object. Must be supplied.  
 k               the number of nearest neighbours to be looked at  
 ...             additional arguments passed to dist()

**Value**

a numeric value; faithfulness

**Examples**

```
delts<-smacof::kinshipdelta
dis<-smacofSym(delts)$confdist
c_faithfulness(dis,obsdiss=delts,k=3)
```

---

c_functionality	<i>c-functionality calculates the c-functionality based on the maximum edge value We define c-functionality as the aggregated functionality between any two columns of confs</i>
-----------------	--

---

**Description**

c-functionality calculates the c-functionality based on the maximum edge value We define c-functionality as the aggregated functionality between any two columns of confs

**Usage**

```
c_functionality(
  confs,
  aggr = NULL,
  alpha = 1,
  C = 15,
  var.thr = 1e-05,
  zeta = NULL
)
```

**Arguments**

confs            a numeric matrix or data frame  
 aggr            the aggregation function for configurations of more than two dimensions. Defaults to mean  
 alpha           an optional number of cells allowed in the X-by-Y search-grid. Default value is 1

C	an optional number determining the starting point of the X-by-Y search-grid. When trying to partition the x-axis into X columns, the algorithm will start with at most C X clumps. Default value is 15.
var.thr	minimum value allowed for the variance of the input variables, since mine can not be computed in case of variance close to 0. Default value is 1e-5.
zeta	integer in [0,1] (?). If NULL (default) it is set to 1-MIC. It can be set to zero for noiseless functions, but the default choice is the most appropriate parametrization for general cases (as stated in Reshef et al.). It provides robustness.

### Value

a numeric value; functionality (aggregated maximum edge value MEV, see [mine](#))

### Examples

```
x<-seq(-3,3,length.out=200)
y<-sqrt(3^2-x^2)
z<- sin(y-x)
confs<-cbind(x,y,z)
c_functionality(confs)
```

---

c_hierarchy	<i>c-hierarchy captures how well a partition/ultrametric (obtained by hclust) explains the configuration distances. Uses variance explained for euclidean distances and deviance explained for everything else.</i>
-------------	---

---

### Description

c-hierarchy captures how well a partition/ultrametric (obtained by hclust) explains the configuration distances. Uses variance explained for euclidean distances and deviance explained for everything else.

### Usage

```
c_hierarchy(confs, voidarg = NULL, p = 2, agglmethod = "complete")
```

### Arguments

confs	a numeric matrix
voidarg	a placeholder to allow to pass NULL as strucpar and not interfere with the other arguments
p	the parameter of the Minokwski distances (p=2 euclidean and p=1 is manhattan)
agglmethod	the method used for creating the clustering, see <a href="#">hclust</a> .

### Value

a numeric value; hierarchy (see [cl\\_validity](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacofSym(delts)$conf
c_hierarchy(conf,p=2,agglmethod="single")
```

---

c_inequality	<i>c-inequality Calculates c-inequality (as in an economic measure of inequality) as Pearsons coefficient of variation of the fitted distance matrix. This can help with avoiding degenerate solutions. This is one of few c-structuredness indices not between 0 and 1, but 0 and infinity.</i>
--------------	--

---

**Description**

c-inequality Calculates c-inequality (as in an economic measure of inequality) as Pearsons coefficient of variation of the fitted distance matrix. This can help with avoiding degenerate solutions. This is one of few c-structuredness indices not between 0 and 1, but 0 and infinity.

**Usage**

```
c_inequality(confs, ...)
```

**Arguments**

confs	a numeric matrix or data frame
...	additional arguments (don't do anything)

**Value**

a numeric value; inequality (Pearsons coefficient of variation of the fitted distance matrix)

**Examples**

```
x<-1:10
y<-2+3*x+rnorm(10)
z<- sin(y-x)
confs<-cbind(z,y,x)
c_inequality(confs)
```

---

c_linearity	<i>c-linearity calculates c-linearity as the aggregated multiple correlation of all columns of the configuration.</i>
-------------	---

---

### Description

c-linearity calculates c-linearity as the aggregated multiple correlation of all columns of the configuration.

### Usage

```
c_linearity(confs, aggr = NULL)
```

### Arguments

confs	a numeric matrix or data frame
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

### Value

a numeric value; linearity (aggregated multiple correlation of all columns of the configuration)

### Examples

```
x<-1:10
y<-2+3*x+rnorm(10)
z<- sin(y-x)
confs<-cbind(z,y,x)
c_linearity(confs)
```

---

c_manifoldness	<i>c-manifoldness calculates c-manifoldness as the aggregated maximal correlation coefficient (i.e., Pearson correlation of the ACE transformed variables) of all pairwise combinations of two different columns in confs. If there is an NA (happens usually when the optimal transformation of any variable is a constant and therefore the covariance is 0 but also one of the sds in the denominator), it gets skipped.</i>
----------------	---

---

### Description

c-manifoldness calculates c-manifoldness as the aggregated maximal correlation coefficient (i.e., Pearson correlation of the ACE transformed variables) of all pairwise combinations of two different columns in confs. If there is an NA (happens usually when the optimal transformation of any variable is a constant and therefore the covariance is 0 but also one of the sds in the denominator), it gets skipped.



**Usage**

```
c_manifoldness(confs, aggr = NULL)
```

**Arguments**

confs	a numeric matrix or data frame
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; manifoldness (aggregated maximal correlation, correlation of ACE tranformed x and y, see [ace](#))

**Examples**

```
x<--100:100
y<-sqrt(100^2-x^2)
confs<-cbind(x,y)
c_manifoldness(confs)
```

---

c_mine	<i>wrapper for getting the mine coefficients</i>
--------	--

---

**Description**

wrapper for getting the mine coefficients

**Usage**

```
c_mine(confs, master = NULL, alpha = 0.6, C = 15, var.thr = 1e-05, zeta = NULL)
```

**Arguments**

confs	a numeric matrix or data frame with two columns
master	the master column
alpha	an optional number of cells allowed in the X-by-Y search-grid. Default value is 0.6
C	an optional number determining the starting point of the X-by-Y search-grid. When trying to partition the x-axis into X columns, the algorithm will start with at most C X clumps. Default value is 15.
var.thr	minimum value allowed for the variance of the input variables, since mine can not be computed in case of variance close to 0. Default value is 1e-5.
zeta	integer in [0,1] (?). If NULL (default) it is set to 1-MIC. It can be set to zero for noiseless functions, but the default choice is the most appropriate parametrization for general cases (as stated in Reshef et al. SOM; they call it epsilon in the paper). It provides robustness.

---

c_nonmonotonicity	<i>c-nonmonotonicity calculates the c-nonmonotonicity based on the maximum asymmetric score We define c-nonmonotonicity as the aggregated nonmonotonicity between any two columns in confs this is one of few c-structuredness indices not between 0 and 1</i>
-------------------	--

---

## Description

c-nonmonotonicity calculates the c-nonmonotonicity based on the maximum asymmetric score We define c-nonmonotonicity as the aggregated nonmonotonicity between any two columns in confs this is one of few c-structuredness indices not between 0 and 1

## Usage

```
c_nonmonotonicity(
  confs,
  aggr = NULL,
  alpha = 1,
  C = 15,
  var.thr = 1e-05,
  zeta = NULL
)
```

## Arguments

confs	a numeric matrix or data frame
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.
alpha	an optional number of cells allowed in the X-by-Y search-grid. Default value is 1
C	an optional number determining the starting point of the X-by-Y search-grid. When trying to partition the x-axis into X columns, the algorithm will start with at most C X clumps. Default value is 15.
var.thr	minimum value allowed for the variance of the input variables, since mine can not be computed in case of variance close to 0. Default value is 1e-5.
zeta	integer in [0,1] (?). If NULL (default) it is set to 1-MIC. It can be set to zero for noiseless functions, but the default choice is the most appropriate parametrization for general cases (as stated in Reshef et al. SOM). It provides robustness.

## Value

a numeric value; nonmonotonicity (aggregated maximal asymmetric score MAS, see [mine](#))

**Examples**

```
x<-seq(-3,3,length.out=200)
y<-sqrt(3^2-x^2)
z<- sin(y-x)
confs<-cbind(x,y,z)
c_nonmonotonicity(confs)
```

---

c\_outlying

*c-outlying*


---

**Description**

Measures the c-outlying structure

**Usage**

```
c_outlying(conf, aggr = NULL)
```

**Arguments**

conf	A numeric matrix.
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; outlying (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf3<-smacof::smacofSym(delts,ndim=3)$conf
c_outlying(conf3)
```

---

c\_regularity

*c-regularity calculates c-regularity as 1 - OPTICS cordillera for k=2.  
The higher the more regular.*


---

**Description**

c-regularity calculates c-regularity as 1 - OPTICS cordillera for k=2. The higher the more regular.

**Usage**

```
c_regularity(
  confs,
  voidarg = NULL,
  q = 1,
  epsilon = 2 * max(dist(confs)),
  distmeth = "euclidean",
  dmax = NULL,
  digits = 10,
  scale = 0,
  ...
)
```

**Arguments**

<code>confs</code>	a numeric matrix or a dist object
<code>voidarg</code>	a placeholder to allow to pass NULL as strucpar and not interfere with the other arguments
<code>q</code>	The norm used for the Cordillera. Defaults to 1 (and should always be 1 imo).
<code>epsilon</code>	The epsilon parameter for OPTICS (called <code>epsilon_max</code> in the paper). Defaults to 2 times the maximum distance between any two points.
<code>distmeth</code>	The distance to be computed if X is not a symmetric matrix or a dist object (otherwise ignored). Defaults to Euclidean distance.
<code>dmax</code>	The winsorization value for the highest allowed reachability. If used for comparisons this should be supplied. If no value is supplied, it is NULL (default), then <code>dmax</code> is taken from the data as minimum of epsilon or the largest reachability.
<code>digits</code>	The precision to round the raw Cordillera and the norm factor. Defaults to 10.
<code>scale</code>	Should X be scaled if it is an asymmetric matrix or data frame? Can take values TRUE or FALSE or a numeric value. If TRUE or 1, standardisation is to mean=0 and sd=1. If 2, no centering is applied and scaling of each column is done with the root mean square of each column. If 3, no centering is applied and scaling of all columns is done as $X/\max(\text{standard deviation}(\text{allcolumns}))$ . If 4, no centering is applied and scaling of all columns is done as $X/\max(\text{rmsq}(\text{allcolumns}))$ . If FALSE, 0 or any other numeric value, no standardisation is applied. Defaults to 0.
<code>...</code>	Additional arguments to be passed to <a href="#">cordillera</a>

**Value**

a numeric value; regularity

**Examples**

```
hpts<-expand.grid(seq(-5,5),seq(-5,5))
c_regularity(hpts)
hpts2<-cbind(jitter(hpts[,1]),jitter(hpts[,2]))
c_regularity(hpts2)
```

---

c_shepardness	<i>c-shepardness calculates the c-shepardness as the correlation between a loess smoother of the transformed distances and the transformed dissimilarities</i>
---------------	--

---

**Description**

c-shepardness calculates the c-shepardness as the correlation between a loess smoother of the transformed distances and the transformed dissimilarities

**Usage**

```
c_shepardness(object, voidarg = NULL)
```

**Arguments**

object	an object of class smacofP
voidarg	empty argument to allow passing NULL as strucpar

**Value**

a numeric value

**Examples**

```
delts<-smacof::kinshipdelta  
res<-smacofx::postmds(delts)  
c_shepardness(res)
```

---

c_skininness	<i>c-skininness</i>
--------------	---------------------

---

**Description**

Measures the c-skininness structure

**Usage**

```
c_skininness(conf, aggr = NULL)
```

**Arguments**

conf	A numeric matrix.
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; skinniness (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacof::smacofSym(delts)$conf
plot(conf,pch=19,asp=1)
c_skininess(conf)
```

---

c\_sparsity

*c-sparsity*


---

**Description**

Measures the c-sparsity structure

**Usage**

```
c_sparsity(conf, aggr = NULL)
```

**Arguments**

conf	A numeric matrix.
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; sparsity (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacof::smacofSym(delts)$conf
plot(conf,pch=19,asp=1)
c_sparsity(conf)
```

---

c_striatedness	<i>c-striatedness</i>
----------------	-----------------------

---

**Description**

Measures the c-striatedness structure

**Usage**

```
c_striatedness(conf, aggr = NULL)
```

**Arguments**

conf	A numeric matrix.
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; striatedness (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacof::smacofSym(delts)$conf
plot(conf,pch=19,asp=1)
c_striatedness(conf)
```

---

c_stringiness	<i>c-stringiness</i>
---------------	----------------------

---

**Description**

Measures the c-stringiness structure

**Usage**

```
c_stringiness(conf, aggr = NULL)
```

**Arguments**

conf	A numeric matrix.
aggr	the aggregation function for configurations of more than two dimensions. Defaults to max.

**Value**

a numeric value; stringiness (see [scagnostics](#))

**Examples**

```
delts<-smacof::kinshipdelta
conf<-smacof::smacofSym(delts)$conf
plot(conf,pch=19,asp=1)
c_stringiness(conf)
```

---

jackmds.stops	<i>MDS Jackknife for stops objects</i>
---------------	--

---

**Description**

These functions perform an MDS Jackknife and plot the corresponding solution.

**Usage**

```
## S3 method for class 'stops'
jackmds(object, eps = 1e-06, itmax = 5000, verbose = FALSE)
```

**Arguments**

object	Object of class pcops.
eps	Convergence criterion
itmax	Maximum number of iterations
verbose	If 'TRUE', intermediate stress is printed out.

**Details**

In order to examine the stability solution of an MDS, a Jackknife on the configurations can be performed (see de Leeuw & Meulman, 1986) and plotted. The plot shows the jackknife configurations which are connected to their centroid. In addition, the original configuration (transformed through Procrustes) is plotted. The Jackknife function itself returns also a stability measure (as ratio of between and total variance), a measure for cross validity, and the dispersion around the original smacof solution.

Note that this jackknife only resamples the configuration given the selected hyperparameters, so uncertainty with respect to the hyperparameter selection is not incorporated.



**Value**

An object of class 'smacofJK', see [jackmds](#). With values

- smacof.conf: Original configuration
- jackknife.confboot: An array of n-1 configuration matrices for each Jackknife MDS solution
- comparison.conf: Centroid Jackknife configurations (comparison matrix)
- cross: Cross validity
- stab: Stability coefficient
- disp: Dispersion
- loss: Value of the loss function (just used internally)
- ndim: Number of dimensions
- call: Model call
- niter: Number of iterations
- nobj: Number of objects

**Examples**

```
diso<-kinshipdelta
fit <- stops(diso,loss="stress",lower=1,upper=5)
res.jk <- jackmds(fit)
plot(res.jk)
```

---

knn\_dist

*calculate k nearest neighbours from a distance matrix*


---

**Description**

calculate k nearest neighbours from a distance matrix

**Usage**

```
knn_dist(dis, k)
```

**Arguments**

dis	distance matrix
k	number of nearest neighbours (Note that with a tie, the function returns the alphanumerically first one!)

ljoptim

*(Adaptive) Version of Luus-Jaakola Optimization***Description**

Adaptive means that the search space reduction factors in the number of iterations; makes convergence faster at about 100 iterations

**Usage**

```
ljoptim(
  x,
  fun,
  ...,
  red = ifelse(adaptive, 0.99, 0.95),
  lower,
  upper,
  acc = 1e-06,
  accd = 1e-04,
  itmax = 1000,
  verbose = 0,
  adaptive = TRUE
)
```

**Arguments**

x	optional starting values
fun	function to minimize
...	additional arguments to be passed to the function to be optimized
red	value of the reduction of the search region
lower	The lower constraints of the search region
upper	The upper constraints of the search region
acc	if the numerical accuracy of two successive target function values is below this, stop the optimization; defaults to 1e-6
accd	if the width of the search space is below this, stop the optimization; defaults to 1e-4
itmax	maximum number of iterations
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
adaptive	should the adaptive version be used? defaults to TRUE.

**Value**

A list with the components (`optim`)

- `par` The position of the optimum in the search space (parameters that minimize the function; `argmin fun`)
- `value` The value of the objective function at the optimum (`min fun`)
- `counts` The number of iterations performed at convergence with entries `fnction` for the number of iterations and `gradient` which is always NA at the moment
- `convergence` 0 successful completion by the `accd` or `acc` criterion, 1 indicate iteration limit was reached, 99 is a problem
- `message` is NULL (only for compatibility or future use)

**Examples**

```
fbana <- function(x) {
  x1 <- x[1]
  x2 <- x[2]
  100 * (x2 - x1 * x1)^2 + (1 - x1)^2
}
res1<-ljoptim(c(-1.2,1),fbana,lower=-5,upper=5,accd=1e-16,acc=1e-16)
res1

set.seed(210485)
fwild <- function (x) 10*sin(0.3*x)*sin(1.3*x^2) + 0.00001*x^4 + 0.2*x+80
plot(fwild, -50, 50, n = 1000, main = "ljoptim() minimising 'wild function'")
res2<-ljoptim(50, fwild,lower=-50,upper=50,adaptive=FALSE,accd=1e-16,acc=1e-16)
points(res2$par,res2$value,col="red",pch=19)
res2
```

---

match\_partial\_ignorecase\_nopunct

*function for lookup that partially matched and ignores cases and punctuation*

---

**Description**

function for lookup that partially matched and ignores cases and punctuation

**Usage**

```
match_partial_ignorecase_nopunct(lookup, entry, ...)
```

**Arguments**

<code>lookup</code>	the lookup string
<code>entry</code>	the registry entry
<code>...</code>	additional arguments to <code>pmatch</code>

---

Pendigits500

*Pen digits*


---

### Description

These data are a random sample of 500 of the 10992 pendigits data from Alimoglu (1996). The original data were from 44 writers who handwrote 250 times the digits 0,...,9. The digits were written inside a rectangular box with a resolution of 500 x 500 pixels and the first 10 per writer were ignored for further analysis. This led to 10992 digits. They were recorded in small time intervals by following the trajectory of the pen on the 500 x 500 grid and then normalized. From the normalized trajectory 8 points (x and y axis position) were randomly selected for each handwritten digit, leading to 16 predictors variables. We extracted a random sample of 500 of them.

### Usage

```
data(Pendigits500)
```

### Format

A data frame with 500 rows and 17 variables

### Details

The variables are

- The rownames of Pendigits500 refer to the data point of the 10992 original data
- V1-V16: trajectory points (x, y coordinate) of the grid
- digits: The digit actually written (the label)

### Source

From A. Izenman (2010) Modern multivariate statistical techniques. Springer.

---

plot.stops

*S3 plot method for stops objects*


---

### Description

S3 plot method for stops objects

### Usage

```
## S3 method for class 'stops'
plot(x, plot.type = "confplot", main, asp = 1, ...)
```

**Arguments**

<code>x</code>	an object of class stops
<code>plot.type</code>	String indicating which type of plot to be produced: "confplot", "resplot", "Shepard", "stressplot", "bubbleplot" (see details)
<code>main</code>	the main title of the plot
<code>asp</code>	aspect ratio of x/y axis; defaults to 1; setting to 1 will lead to an accurate representation of the fitted distances.
<code>...</code>	Further plot arguments passed: see 'plot.smacof' and 'plot' for detailed information. Details: See plot.smacofP

**Value**

no return value, just plots

---

<code>print.stops</code>	<i>S3 print method for stops objects</i>
--------------------------	--

---

**Description**

S3 print method for stops objects

**Usage**

```
## S3 method for class 'stops'
print(x, ...)
```

**Arguments**

<code>x</code>	stops object
<code>...</code>	additional arguments

**Value**

no return value, just prints

---

<code>print.summary.stops</code>	<i>S3 print method for summary.stops</i>
----------------------------------	--

---

**Description**

S3 print method for summary.stops

**Usage**

```
## S3 method for class 'summary.stops'  
print(x, ...)
```

**Arguments**

<code>x</code>	object of class summary.stops
<code>...</code>	additional arguments

**Value**

no return value, just prints

---

<code>residuals.stops</code>	<i>S3 residuals method for stops</i>
------------------------------	--------------------------------------

---

**Description**

S3 residuals method for stops

**Usage**

```
## S3 method for class 'stops'  
residuals(object, ...)
```

**Arguments**

<code>object</code>	object of class stops
<code>...</code>	addditional arguments

**Value**

a vector of residuals (observed minus fitted distances)

stoploss

*Calculate the weighted multiobjective loss function used in STOPS***Description**

Calculate the weighted multiobjective loss function used in STOPS

**Usage**

```
stoploss(
  obj,
  stressweight = 1,
  structures,
  strucweight = rep(-1/length(structures), length(structures)),
  strucpars,
  stoptype = c("additive", "multiplicative"),
  verbose = 0,
  registry = struc_reg
)
```

**Arguments**

obj	object returned inside a stop_* function. Uses the stress.m slot for getting the stress.
stressweight	weight to be used for the fit measure; defaults to 1
structures	which c-structuredness indices to be included in the loss
strucweight	the weights of the structuredness indices; defaults to -1/#number of structures
strucpars	a list of parameters to be passed to the c-structuredness indices in the same order as the values in structures. If the index has no parameters or you want to use the defaults, supply NULL. (alternatively a named list that has the structure name as the element name).
stoptype	what type of weighted combination should be used? Can be 'additive' or 'multiplicative'.
verbose	verbose output
registry	an object of class registry. This can be used to add additional c-structuredness indices. Defaults to the registry created via .onLoad in zzz.R

**Value**

a list with calculated stoploss (`$stoploss`), structuredness indices (`$strucinidices`) and hyperparameters (`$parameters` and `$theta`)

---

 stops

*High Level STOPS Function*


---

## Description

This allows to fit STOPS models as described in Rusch, Mair, Hornik (2023).

## Usage

```
stops(
  dis,
  loss = "stress",
  theta = 1,
  type = "ratio",
  structures,
  ndim = 2,
  weightmat = NULL,
  init = NULL,
  stressweight = 1,
  strucweight,
  strucpars,
  optimmethod = c("SANN", "ALJ", "pso", "Kriging", "tgp", "direct", "stogo", "cobyla",
    "crs2lm", "isres", "mlsl", "neldermead", "sbplx", "hjk", "cmaes"),
  lower,
  upper,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  initpoints = 10,
  itmax = 50,
  itmaxps = 10000,
  model,
  control,
  registry = struc_reg,
  ...
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
loss	which loss function to be used for fitting, defaults to stress.
theta	hyperparameter vector starting values for the transformation functions. If the length is smaller than the number of hyperparameters for the MDS version the vector gets recycled (see the corresponding stop_XXX function or the vignette for how theta must look like exactly for each loss). If larger than the number of hyperparameters for the MDS method, an error is thrown. If completely missing theta is set to 1 and recycled.



type	type of MDS optimal scaling (implicit transformation). One of "ratio", "interval" or "ordinal". Default is "ratio". Not every type can be used with every loss, only ratio works with all.
structures	character vector of which c-structuredness indices should be considered; if missing no structure is considered.
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights; defaults to 1 for all off diagonals
init	(optional) initial configuration
stressweight	weight to be used for the fit measure; defaults to 1
strucweight	vector of weights to be used for the c-structuredness indices (in the same order as in structures); defaults to $-1/\text{length}(\text{structures})$ for each index
strucpars	(possibly named with the structure). Metaparameters for the structuredness indices (gamma in the article). It's safest for it be a list of lists with the named arguments for the structuredness indices and the order of the lists must be like the order of structures. So something like this <code>list(list(par1Struc1=par1Struc1, par2Struc1=par2Struc1, ...), ...)</code> where <code>parYStrucX</code> are the named arguments for the metaparameter Y of the structure X the list elements corresponds to. For a structure without parameters, set NULL. Parameters in different list elements <code>parYStrucX</code> can have the same name. For example, say we want to use <code>cclusteredness</code> with metaparameters <code>epsilon=10</code> and <code>k=4</code> (and the default for the other parameters), <code>cdependence</code> with no metaparameters and <code>cfaithfulness</code> with metaparameter <code>k=7</code> one would <code>list(list(epsilon=10, k=4), list(NULL), list(dis=obdiss, k=6))</code> for structures vector <code>("cclusteredness", "cdependence", "cfaitfulness")</code> . The parameter lists must be in the same ordering as the indices in structures. If missing it is set to NULL and defaults are used. It is also possible to supply a structure's metaparameters as a list of vectors with named elements if the metaparameters are scalars, so like <code>list(c(par1Struc1=parStruc1, par2Struc1=par1Struc1, ...), c(par1Struc2=parStruc2, ...))</code> . That can have unintended consequences if the metaparameter is a vector or matrix.
optimmethod	What solver to use. Currently supported are Bayesian optimization with Gaussian Process priors and Kriging ("Kriging", see <a href="#">EG0.nsteps</a> ), Bayesian optimization with treed Gaussian processes with jump to linear models ("tgp", see <a href="#">dopt.gp</a> ), Adaptive LJ Search ("ALJ"), Particle Swarm optimization ("pso", see <a href="#">psoptim</a> ), simulated annealing ("SANN", <a href="#">optim</a> ), "direct ( <a href="#">direct</a> )", Stochastic Global Optimization ("stogo", <a href="#">stogo</a> ), COBYLA ("cobyla", <a href="#">cobyla</a> ), Controlled Random Search 2 with local mutation ("crs2lm", <a href="#">crs2lm</a> ), Improved Stochastic Ranking Evolution Strategy ("isres", <a href="#">isres</a> ), Multi-Level Single-Linkage ("mlsl", <a href="#">mlsl</a> ), Nelder-Mead ("neldermead", <a href="#">neldermead</a> ), Subplex ("sbplx", <a href="#">sbplx</a> ), Hooke-Jeeves Pattern Search ("hjk", <a href="#">hjk</a> ), CMA-ES ("cmaes", <a href="#">cma_es</a> ). Defaults to "ALJ" version. "tgp", "ALJ", "Kriging" and "pso" usually work well for relatively low values of 'itmax'.
lower	The lower constraints of the search region. Needs to be a numeric vector of the same length as the parameter vector theta.
upper	The upper constraints of the search region. Needs to be a numeric vector of the same length as the parameter vector theta.

verbose	numeric value that prints information on the fitting process; >2 is very verbose.
stoptype	which aggregation for the multi objective target function? Either 'additive' (default) or 'multiplicative'
initpoints	number of initial points to fit the surrogate model for Bayesian optimization; default is 10.
itmax	maximum number of iterations of the outer optimization (for theta) or number of steps of Bayesian optimization; default is 50. We recommend a higher number for ALJ (around 150). Note that due to the inner workings of some solvers, this may or may not correspond to the actual number of function evaluations performed (or PS models fitted). E.g., with tgp the actual number of function evaluation of the PS method is between itmax and 6*itmax as tgp samples 1-6 candidates from the posterior and uses the best candidate. For pso it is the number of particles $s$ times itmax. For cmaes it is usually a bit higher than itmax. This currently may get overruled by a control argument if it is used (and then set to either ewhat is supplied by control or to the default of the method).
itmaxps	maximum number of iterations of the inner optimization (to obtain the PS configuration)
model	a character specifying the surrogate model to use. For Kriging it specifies the covariance kernel for the GP prior; see <a href="#">covTensorProduct-class</a> defaults to "powerexp". For tgp it specifies the non stationary process used see <a href="#">bgp</a> , defaults to "btgp1lm"
control	a control argument passed to the outer optimization procedure. Will override any other control arguments passed, especially verbose and itmax. For the effect of control, see the functions <code>pomp::sannbox</code> for SANN and <code>pso::psoptim</code> for pso, <code>cmaes::cma_es</code> for cmaes, <code>dfoptim::hjk</code> for hjk and the <code>nloptr</code> docs for the algorithms <code>direct</code> , <code>stogo</code> , <code>cobyqa</code> , <code>crs2lm</code> , <code>isres</code> , <code>mlsl</code> , <code>neldermead</code> , <code>sbplx</code> .
registry	an object of class <code>registry</code> containing the c-structuredness indices. Defaults to the what is created <code>.onLoad</code> .
...	additional arguments passed to the outer optimization procedures (not fully tested).

## Details

The combination of c-structuredness indices and stress uses the `stress.m` values, which are the explicitly normalized stresses. Reported however is the stress-1 value which is `sqrt(stress.m)`.

## Value

A list with the components

- `stoploss`: the stoploss value
- `optim`: the object returned from the optimization procedure
- `stressweight`: the stressweight
- `strucweight`: the vector of structure weights
- `call`: the call
- `optimmethod`: The solver selected

- loss: The PS badness-of-fit function
- nobj: the number of objects in the configuration
- type: The type of stoploss scalacrisation (additive or multiplicative)
- fit: The fitted PS object (most importantly \$fit\$conf the fitted configuration)
- stoptype: Type of stoploss combinatio

### Examples

```
data(kinshipdelta,package="smacof")
strucpar<-list(NULL,NULL) #parameters for indices
res1<-stops(kinshipdelta,loss="stress",
structures=c("cclumpiness","cassociation"),strucpars=strucpar,
lower=0,upper=10,itmax=10)
res1

#use higher itmax in general, we use 5 just to shorten the tests
data(BankingCrisesDistances)
strucpar<-list(c(epsilon=10,minpts=2),NULL) #parameters for indices
res1<-stops(BankingCrisesDistances[,1:69],loss="stress",verbose=0,
structures=c("cclusteredness","clinearity"),strucpars=strucpar,
lower=0,upper=10,itmax=5)
res1

strucpar<-list(list(alpha=0.6,C=15,var.thr=1e-5,zeta=NULL),
list(alpha=0.6,C=15,var.thr=1e-5,zeta=NULL))
res1<-stops(BankingCrisesDistances[,1:69],loss="stress",verbose=0,
structures=c("cfunctionality","ccomplexity"),strucpars=strucpar,
lower=0,upper=10,itmax=5)
res1
```

---

stop\_apstress

*STOPS version of approximated power stress models.*


---

### Description

This uses an approximation to power stress that can make use of smacof as workhorse. Free parameters are kappa, lambda and nu.

### Usage

```
stop_apstress(
  dis,
  theta = c(1, 1, 1),
  type = "ratio",
  ndim = 2,
```

```

weightmat = 1 - diag(nrow(dis)),
init = NULL,
itmaxi = 1000,
...,
stressweight = 1,
structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
  "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
  "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
  "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
strucweight = rep(1/length(structures), length(structures)),
strucpars,
verbose = 0,
stoptype = c("additive", "multiplicative"),
registry = struc_reg
)

```

### Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of parameters to optimize over. Must be of length three, with the first the kappa argument, the second the lambda argument and the third the nu argument. One cannot supply epsilon and tau as of yet. Defaults to 1 1 1.
<code>type</code>	MDS type.
<code>ndim</code>	number of dimensions of the target space
<code>weightmat</code>	(optional) a binary matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>itmaxi</code>	number of iterations. default is 1000.
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures vector. See examples.
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value (sqrt stress.m)
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda, nu)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop\_bcmds

*STOPS version of Box Cox Stress*


---

## Description

STOPS version of Box Cox Stress

## Usage

```
stop_bcmds(
  dis,
  theta = c(1, 1, 0),
  type = "ratio",
  weightmat = NULL,
  init = NULL,
  ndim = 2,
  itmaxi = 5000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; the first is mu (for the fitted distances), the second lambda (for the proximities), the third nu (for the weights). If a scalar is given it is recycled. Defaults to 1 1 0.
type	MDS type. Is ignored here.

weightmat	(not used)
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	which structures to look for
strucweight	weight to be used for the structures; defaults to 0.5
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_clca	<i>STOPS version of CLCA.</i>
-----------	-------------------------------

---

### Description

CLCA with free lambda0 and 20 epochs. Should we add alpha0?

**Usage**

```

stop_clca(
  dis,
  theta = 3 * max(sd(dis)),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)

```

**Arguments**

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of explicit parameters; lambda0 for the maximal neighbourhood. Defaults to 100.
<code>type</code>	MDS type.
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value hat prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

**Value**

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (tau)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop\_cldae

*STOPS version of CLDA with free epsilon.*


---

**Description**

CLDA with free lambda0 and epsilon and 20 epochs. Should we add alpha0?

**Usage**

```
stop_cldae(
  dis,
  theta = rep(3 * max(sd(dis)), 2),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```



**Arguments**

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of explicit parameters; first is $\lambda_0$ for the maximal neighbourhood and second is $k$ for the number of neighbours for the geodesic distance.
<code>type</code>	MDS type.
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

**Value**

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `struc`: the structuredness indices
- `parameters`: the parameters used for fitting ( $\tau$ )
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object

---

stop\_cldak

*STOPS version of CLDA with free k.*


---

### Description

CLDA with free lambda0 and k and 20 epochs. Should we add alpha0?

### Usage

```
stop_cldak(
  dis,
  theta = c(3 * max(sd(dis)), nrow(dis)/4),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskinniness", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

### Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of explicit parameters; first is lambda0 for the maximal neighbourhood and second is k for the number of neighbours for the geodesic distance.
type	MDS type.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.

strucweight	weight to be used for the structures; defaults to 1/number of structures
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (tau)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_cmdscale	<i>STOPS version of strain</i>
---------------	--------------------------------

---

### Description

The free parameter is lambda for power transformations of the observed proximities.

### Usage

```
stop_cmdscale(
  dis,
  theta = 1,
  type = "ratio",
  weightmat = NULL,
  ndim = 2,
  init = NULL,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
```

```

    strucpars,
    verbose = 0,
    stoptype = c("additive", "multiplicative"),
    itmaxi = 1000,
    add = TRUE,
    registry = struc_reg
  )

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; this must be a scalar of the lambda transformation for the observed proximities.
<code>type</code>	MDS type. Ignored here.
<code>weightmat</code>	(optional) a matrix of nonnegative weights. Not used.
<code>ndim</code>	number of dimensions of the target space
<code>init</code>	(optional) initial configuration
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structuredness indices to be included in the loss
<code>strucweight</code>	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
<code>strucpars</code>	the parameters for the structuredness indices
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
<code>itmaxi</code>	number of iterations. No effect here.
<code>add</code>	if TRUE <code>dis</code> is made to Euclidean distances
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the badness-of-fit value (this isn't stress here but  $1 - (\text{sum\_ndim}(\text{max}(\text{eigenvalues}, 0)) / \text{sum\_n}(\text{max}(\text{eigenvalues}, 0)))$ , 1-GOF[2])
- `stress.m`: explicitly normalized stress (manually calculated)
- `stoploss`: the weighted loss value
- `indices`: the values of the structuredness indices
- `parameters`: the parameters used for fitting (lambda)
- `fit`: the returned object of the fitting procedure, which is `cmdscalex` object with some extra slots for the parameters and stresses
- `stopobj`: the `stopobj` object

stop\_elastic

*STOPS versions of elastic scaling models (via smacofSym)***Description**

The free parameter is lambda for power transformations the observed proximities. The fitted distances power is internally fixed to 1 and the power for the weights=delta is -2. Allows for a weight matrix because of smacof.

**Usage**

```
stop_elastic(
  dis,
  theta = 1,
  type = "ratio",
  ndim = 2,
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  itmaxi = 1000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

**Arguments**

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this must be a scalar of the lambda transformation for the observed proximities. Defaults to 1.
type	MDS type. Defaults to 'ratio'.
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights (NOT the elscal weights)
init	(optional) initial configuration
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1

structures	which structuredness indices to be included in the loss
strucweight	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
strucpars	the parameters for the structuredness indices
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
stoptype	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 ( $\sqrt{\text{stress.m}}$ )
- stress.m: default normalized stress (used for STOPS)
- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting ( $\lambda$ )
- fit: the returned object of the fitting procedure
- stopobj: the stopobj objects

---

stop_isomap1	<i>STOPS version of isomap to optimize over integer <math>k</math>.</i>
--------------	---

---

### Description

Free parameter is  $k$ .

### Usage

```
stop_isomap1(
  dis,
  theta = 3,
  type = "ratio",
  weightmat = NULL,
  ndim = 2,
  init = NULL,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
```

```

    strucpars,
    verbose = 0,
    stoctype = c("additive", "multiplicative"),
    itmaxi = NULL,
    registry = struc_reg
  )

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the number of shortest dissimilarities retained for a point (nearest neighbours), the isomap parameter. Must be a numeric scalar. Defaults to 3.
<code>type</code>	MDS type. Is "ratio".
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>ndim</code>	number of dimensions of the target space
<code>init</code>	(optional) initial configuration
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structuredness indices to be included in the loss
<code>strucweight</code>	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
<code>strucpars</code>	the parameters for the structuredness indices
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoctype</code>	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
<code>itmaxi</code>	placeholder for compatibility in stops call; not used
<code>registry</code>	registry object with c-structuredness indices.

## Details

Currently this version is a bit less flexible than the vegan one, as the only allowed parameter for isomap is the theta (k in isomap, no epsilon) and the shortest path is always estimated with argument "shortest". Also note that fragmentedOK is always set to TRUE which means that for theta that is too small only the largest connected group will be analyzed. If that's not wanted just set the theta higher.

## Value

A list with the components

- `stress`: Not really stress but 1-GOF[2] where GOF is the second element returned from `sma-cofx::cmdscale` (the sum of the first `ndim` eigenvalues divided by the sum of all absolute eigenvalues).
- `stress.m`: default normalized stress (sqrt explicitly normalized stress; really the stress this time)

- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop\_isomap2

*STOPS version of isomap over real epsilon.*


---

### Description

Free parameter is eps.

### Usage

```
stop_isomap2(
  dis,
  theta = stats::quantile(dis, 0.1),
  type = "ratio",
  weightmat = NULL,
  ndim = 2,
  init = NULL,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskinniness", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  itmaxi = NULL,
  registry = struc_reg
)
```

### Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the number of shortest dissimilarities retained for a point (neighbourhood region), the isomap parameter. Defaults to the 0.1 quantile of the empirical distribution of dis.
type	MDS type. Is "ratio".
weightmat	(optional) a matrix of nonnegative weights
ndim	number of dimensions of the target space



init	(optional) initial configuration
stressweight	weight to be used for the fit measure; defaults to 1
structures	which structuredness indices to be included in the loss
strucweight	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
strucpars	the parameters for the structuredness indices
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
stoptype	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
itmaxi	placeholder for compatibility in stops call; not used
registry	registry object with c-structuredness indices.

## Details

Currently this version is a bit less flexible than the vegan one, as the only allowed parameter for isomap is the theta (epsilon in isomap) and the shortest path is always estimated with argument "shortest". Also note that fragmentedOK is always set to TRUE which means that for theta that is too small only the largest conected group will be analyzed. If that's not wanted just set the theta higher.

## Value

A list with the components

- stress: Not really stress but 1-GOF[2] where GOF is the second element returned from cmd-scale (the sum of the first ndim absolute eigenvalues divided by the sum of all absolute eigenvalues).
- stress.m: default normalized stress (sqrt explicitly normalized stress; really the stress this time)
- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

stop\_lmds

*STOPS version of lmds***Description**

STOPS version of lmds

**Usage**

```
stop_lmds(
  dis,
  theta = c(2, 0.5),
  type = "ratio",
  weightmat = NULL,
  init = NULL,
  ndim = 2,
  itmaxi = 5000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

**Arguments**

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; the first is k (for the neighbourhood), the second tau (for the penalty) . If a scalar is given it is recycled. Defaults to 2 and 0.5.
type	MDS type. Ignored.
weightmat	(not used)
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	which structures to look for
strucweight	weight to be used for the structures; defaults to 0.5

strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structure
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

## Value

A list with the components

- stress: the stress-1
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_powerelastic	<i>STOPS version of elastic scaling with powers for proximities and distances</i>
-------------------	---

---

## Description

This is power stress with free kappa and lambda but rho is fixed to -2 and the weights are delta.

## Usage

```
stop_powerelastic(
  dis,
  theta = c(1, 1),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 1e+05,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
```

```

    strucpars,
    verbose = 0,
    stoptype = c("additive", "multiplicative"),
    registry = struc_reg
  )

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; a vector of length two where the first element is kappa (for the fitted distances), the second lambda (for the observed proximities). If a scalar for the free parameters is given it is recycled. Defaults to 1 1.
<code>type</code>	MDS type. Defaults to "ratio".
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structures to look for
<code>strucweight</code>	weight to be used for the structures; defaults to 0.5
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `struc`: the structuredness indices
- `parameters`: the parameters used for fitting (kappa, lambda)
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object

---

stop\_powermds

*STOPS version of powermds*


---

## Description

This is power stress with free kappa and lambda but rho is fixed to 1, so no weight transformation.

## Usage

```
stop_powermds(
  dis,
  theta = c(1, 1),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; a vector of length 2 where the first element is kappa (for the fitted distances), the second lambda (for the observed proximities). If a scalar is given it is recycled. Defaults to 1,1.
type	MDS type. Defaults to "ratio".
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	which structures to look for

strucweight	weight to be used for the structures; defaults to 0.5
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop\_powersammon

*STOPS version of sammon with powers*


---

### Description

This is power stress with free kappa and lambda but rho is fixed to -1 and the weights are delta.

### Usage

```
stop_powersammon(
  dis,
  theta = c(1, 1),
  type = "ratio",
  weightmat = NULL,
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
```

```

    strucweight = rep(1/length(structures), length(structures)),
    strucpars,
    verbose = 0,
    stoptype = c("additive", "multiplicative"),
    registry = struc_reg
  )

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; a vector of length two where the first element is kappa (for the fitted distances), the second lambda (for the observed proximities). If a scalar is given it is recycled for the free parameters. Defaults to 1 1.
<code>type</code>	MDS type. Defaults to "ratio".
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structures to look for
<code>strucweight</code>	weight to be used for the structures; defaults to 0.5
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the stress
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `struc`: the structuredness indices
- `parameters`: the parameters used for fitting (kappa, lambda)
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object

---

stop_powerstress	<i>STOPS version of powerstress</i>
------------------	-------------------------------------

---

## Description

Power stress with free kappa and lambda and rho.

## Usage

```
stop_powerstress(
  dis,
  theta = c(1, 1, 1),
  type = "ratio",
  weightmat = NULL,
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; the first is kappa (for the fitted distances), the second lambda (for the observed proximities), the third nu (for the weights). If a scalar is given it is recycled. Defaults to 1 1 1.
type	MDS type.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.



strucweight	weight to be used for the structures; defaults to 1/number of structures
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda, nu)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_rpowerstress	<i>STOPS version of restricted powerstress</i>
-------------------	--

---

### Description

STOPS version of restricted powerstress

### Usage

```
stop_rpowerstress(
  dis,
  theta = c(1, 1, 1),
  type = "ratio",
  weightmat = NULL,
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
```

```

    strucweight = rep(1/length(structures), length(structures)),
    strucpars,
    verbose = 0,
    stoptype = c("additive", "multiplicative"),
    registry = struc_reg
  )

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; the first two arguments are for kappa and lambda and should be equal (for the fitted distances and observed proximities), the third nu (for the weights). Internally the kappa and lambda are equated. If a scalar is given it is recycled (so all elements of theta are equal); if a vector of length 2 is given, it gets expanded to <code>c(theta[1],theta[1],theta[2])</code> . Defaults to 1 1 1.
<code>type</code>	MDS type. Defaults to "ratio".
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations. default is 10000.
<code>...</code>	additional arguments to be passed to the fitting procedure <code>powerStressMin</code>
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures vector. See examples.
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `struc`: the structuredness indices
- `parameters`: the parameters used for fitting (kappa=lambda, nu)
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object

---

stop_rstress	<i>STOPS version of rstress</i>
--------------	---------------------------------

---

## Description

Free parameter is  $\kappa=2r$  for the fitted distances.

## Usage

```
stop_rstress(
  dis,
  theta = 1,
  type = "ratio",
  weightmat = NULL,
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this must be a scalar of the $\kappa=2*r$ transformation for the fitted distances proximities. Defaults to 1. Note that what is returned is $r$ , not $\kappa$ .
type	MDS type. Default is "ratio"
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations.
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	which structuredness indices to be included in the loss

strucweight	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
strucpars	the parameters for the structuredness indices
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop\_sammon

*STOPS version of Sammon mapping*


---

### Description

Uses smacofx::sammon. The free parameter is lambda for power transformations of the observed proximities.

### Usage

```
stop_sammon(
  dis,
  theta = 1,
  type = "ratio",
  ndim = 2,
  init = NULL,
  weightmat = NULL,
  itmaxi = 1000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "chierarchy", "cconvexity", "cstriatedness", "coutlying", "cskininess", "csparsity",
```

```

    "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; this must be a scalar of the lambda transformation for the observed proximities. Defaults to 1.
<code>type</code>	MDS type. Ignored here.
<code>ndim</code>	number of dimensions of the target space
<code>init</code>	(optional) initial configuration
<code>weightmat</code>	a matrix of nonnegative weights. Has no effect here.
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structuredness indices to be included in the loss
<code>strucweight</code>	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
<code>strucpars</code>	the parameters for the structuredness indices
<code>verbose</code>	numeric value hat prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the stress/1 \*sqrt stress(
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `indices`: the values of the structuredness indices
- `parameters`: the parameters used for fitting
- `fit`: the returned object of the fitting procedure `smacofx::sammon`
- `stopobj`: the stopobj object

---

stop\_sammon2

*Another STOPS version of Sammon mapping models (via smacofSym)*


---

## Description

Uses Smacof, so it can deal with a weight matrix too. The free parameter is lambda for power transformations of the observed proximities. The fitted distances power is internally fixed to 1 and the power for the weights=delta is -1.

## Usage

```
stop_sammon2(
  dis,
  theta = 1,
  type = "ratio",
  ndim = 2,
  weightmat = NULL,
  init = NULL,
  itmaxi = 1000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this must be a scalar of the lambda transformation for the observed proximities. Defaults to 1.
type	MDS type
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1

structures	which structuredness indices to be included in the loss
strucweight	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
strucpars	the parameters for the structuredness indices
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
stoptype	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 ( $\sqrt{\text{stress.m}}$ )
- stress.m: default normalized stress (used for STOPS)
- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting ( $\lambda$ )
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_smacofSphere	<i>STOPS versions of smacofSphere models</i>
-------------------	--

---

### Description

The free parameter is  $\lambda$  for power transformations the observed proximities. The fitted distances power is internally fixed to 1 and the power for the weights is 1.

### Usage

```
stop_smacofSphere(
  dis,
  theta = 1,
  type = "ratio",
  ndim = 2,
  weightmat = NULL,
  init = NULL,
  itmaxi = 1000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
```

```

    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)

```

### Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; this must be a scalar of the lambda transformation for the observed proximities. Defaults to 1.
<code>type</code>	MDS type.
<code>ndim</code>	number of dimensions of the target space
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structuredness indices to be included in the loss
<code>strucweight</code>	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
<code>strucpars</code>	the parameters for the structuredness indices
<code>verbose</code>	numeric value hat prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
<code>registry</code>	registry object with c-structuredness indices.

### Value

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `indices`: the values of the structuredness indices
- `parameters`: the parameters used for fitting
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object



---

stop_smacofSym	<i>STOPS version of smacofSym models</i>
----------------	--

---

## Description

The free parameter is lambda for power transformations the observed proximities. The fitted distances power is internally fixed to 1 and the power for the weights is 1.

## Usage

```
stop_smacofSym(
  dis,
  theta = 1,
  type = "ratio",
  ndim = 2,
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  itmaxi = 1000,
  ...,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "chierarchy", "cconvexity", "cstriatedness", "coutlying", "cskininess", "csparsity",
    "cstringiness", "cclumpiness", "cinequality"),
  stressweight = 1,
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector; must be a scalar for the lambda (proximity) transformation. Defaults to 1.
type	MDS type. Defaults to 'ratio'.
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
itmaxi	number of iterations
...	additional arguments to be passed to the fitting
structures	which structuredness indices to be included in the loss
stressweight	weight to be used for the fit measure; defaults to 1

strucweight	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
strucpars	the parameters for the structuredness indices
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
stoptype	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 ( $\sqrt{\text{stress.m}}$ )
- stress.m: default normalized stress (used for STOPS)
- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting (lambda)
- fit: the returned object of the fitting procedure
- stopobj: the stops object

---

stop_smddae	<i>STOPS version of sparsified multidimensional distance analysis for fixed eps and tau</i>
-------------	---

---

### Description

smdda with free parameters tau and epsilon.

### Usage

```
stop_smddae(
  dis,
  theta = c(100, 100),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
```

```

    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of explicit parameters; first is tau for the neighbourhood, second is epsilon for isomapdist. Defaults to 100, 100.
<code>type</code>	MDS type.
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `struc`: the structuredness indices
- `parameters`: the parameters used for fitting (tau)
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object

---

stop_smddak	<i>STOPS version of sparsified multidimensional distance analysis for fixed k and tau</i>
-------------	---

---

## Description

smdda with free parameters tau and k.

## Usage

```
stop_smddak(
  dis,
  theta = c(100, 10),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of explicit parameters; first is tau for the neighbourhood, second is k. Defaults to 100, 10.
type	MDS type.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.

strucweight	weight to be used for the structures; defaults to 1/number of structures
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

## Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (tau)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_smds	<i>STOPS version of sparsified MDS.</i>
-----------	---

---

## Description

smds with free tau.

## Usage

```
stop_smds(
  dis,
  theta = c(100),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
```

```

    strucweight = rep(1/length(structures), length(structures)),
    strucpars,
    verbose = 0,
    stoptype = c("additive", "multiplicative"),
    registry = struc_reg
  )

```

## Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of explicit parameters; tau for the neighbourhood. Defaults to 100.
<code>type</code>	MDS type.
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value that prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

## Value

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value
- `struc`: the structuredness indices
- `parameters`: the parameters used for fitting (tau)
- `fit`: the returned object of the fitting procedure
- `stopobj`: the stopobj object

---

stop_spmddae	<i>STOPS version of sparsified post multidimensional distance analysis for fixed tau and epsilon.</i>
--------------	---

---

## Description

Sparsified POST MDDA with free kappa, lambda, rho, tau and epsilon. Phew.

## Usage

```
stop_spmddae(
  dis,
  theta = c(1, 1, 1, 100, 100),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of explicit parameters; the first is kappa (for the fitted distances), the second lambda (for the observed proximities), the third nu (for the weights), the fourth tau (for the neighbourhood), the fifth the epsilon for the geodesic distances. If a scalar or vector shorter than 5 is given it is recycled. Defaults to 1 1 1 100 10.
type	MDS type.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure

stressweight	weight to be used for the fit measure; defaults to 1
structures	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
strucweight	weight to be used for the structures; defaults to 1/number of structures
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda, nu, tau)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_spmddak	<i>STOPS version of sparsified post multidimensional distance analysis for fixed tau and k.</i>
--------------	---

---

### Description

Sparsified Post MDDA with free kappa, lambda, rho, tau and k. Phew.

### Usage

```
stop_spmddak(
  dis,
  theta = c(1, 1, 1, 100, 10),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
```



```

structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
  "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
  "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
  "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
strucweight = rep(1/length(structures), length(structures)),
strucpars,
verbose = 0,
stoptype = c("additive", "multiplicative"),
registry = struc_reg
)

```

### Arguments

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of explicit parameters; the first is kappa (for the fitted distances), the second lambda (for the observed proximities), the third nu (for the weights), the fourth tau (for the neighbourhood), the fifth the k for the geodesic distances. If a scalar or vector shorter than 5 is given it is recycled. Defaults to 1 1 1 100 10.
<code>type</code>	MDS type.
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
<code>strucweight</code>	weight to be used for the structures; defaults to 1/number of structures
<code>strucpars</code>	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
<code>verbose</code>	numeric value hat prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
<code>registry</code>	registry object with c-structuredness indices.

### Value

A list with the components

- `stress`: the stress-1 value
- `stress.m`: default normalized stress
- `stoploss`: the weighted loss value

- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda, nu, tau)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop\_spmds

*STOPS version of sparsified POST-MDS for fixed tau*


---

## Description

Sparsified power stress with free kappa, lambda, rho and tau.

## Usage

```
stop_spmds(
  dis,
  theta = c(1, 1, 1, 100),
  type = "ratio",
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 10000,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)
```

## Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of explicit parameters; the first is kappa (for the fitted distances), the second lambda (for the observed proximities), the third nu (for the weights), the fourth tau (for the neighbourhood). If a scalar or vector shorter than 4 is given it is recycled. Defaults to 1 1 1 100.
type	MDS type.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration

ndim	number of dimensions of the target space
itmaxi	number of iterations
...	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
structures	a character vector listing the structure indices to use. They always are called "cfoo" with foo being the structure.
strucweight	weight to be used for the structures; defaults to 1/number of structures
strucpars	a list of parameters for the structuredness indices; each list element corresponds to one index in the order of the appearance in structures
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
stoptype	which weighting to be used in the multi-objective optimization? Either 'additive' (default) or 'multiplicative'.
registry	registry object with c-structuredness indices.

### Value

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- struc: the structuredness indices
- parameters: the parameters used for fitting (kappa, lambda, nu, tau)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

stop_sstress	<i>STOPS version of sstress</i>
--------------	---------------------------------

---

### Description

Free parameter is lambda for the observed proximities. Fitted distances are transformed with power 2, weights have exponent of 1. Note that the lambda here works as a multiplier of 2 (as sstress has  $f(\delta^2)$ ).

**Usage**

```

stop_sstress(
  dis,
  theta = 1,
  type = type,
  weightmat = 1 - diag(nrow(dis)),
  init = NULL,
  ndim = 2,
  itmaxi = 1e+05,
  ...,
  stressweight = 1,
  structures = c("cclusteredness", "clinearity", "cdependence", "cmanifoldness",
    "cassociation", "cnonmonotonicity", "cfunctionality", "ccomplexity", "cfaithfulness",
    "cregularity", "chierarchy", "cconvexity", "cstriatedness", "coutlying",
    "cskininess", "csparsity", "cstringiness", "cclumpiness", "cinequality"),
  strucweight = rep(1/length(structures), length(structures)),
  strucpars,
  verbose = 0,
  stoptype = c("additive", "multiplicative"),
  registry = struc_reg
)

```

**Arguments**

<code>dis</code>	numeric matrix or dist object of a matrix of proximities
<code>theta</code>	the theta vector of powers; this must be a scalar of the lambda transformation for the observed proximities. Defaults to 1. Note that the lambda here works as a multiplier of 2 (as sstress has $f(\delta^2)$ ).
<code>type</code>	MDS type.
<code>weightmat</code>	(optional) a matrix of nonnegative weights
<code>init</code>	(optional) initial configuration
<code>ndim</code>	the number of dimensions of the target space
<code>itmaxi</code>	number of iterations
<code>...</code>	additional arguments to be passed to the fitting procedure
<code>stressweight</code>	weight to be used for the fit measure; defaults to 1
<code>structures</code>	which structuredness indices to be included in the loss
<code>strucweight</code>	weight to be used for the structuredness indices; ; defaults to 1/#number of structures
<code>strucpars</code>	the parameters for the structuredness indices
<code>verbose</code>	numeric value hat prints information on the fitting process; >2 is extremely verbose
<code>stoptype</code>	How to construct the target function for the multi objective optimization? Either 'additive' (default) or 'multiplicative'
<code>registry</code>	registry object with c-structuredness indices.

**Value**

A list with the components

- stress: the stress-1 value
- stress.m: default normalized stress
- stoploss: the weighted loss value
- indices: the values of the structuredness indices
- parameters: the parameters used for fitting (lambda)
- fit: the returned object of the fitting procedure
- stopobj: the stopobj object

---

summary.stops

*S3 summary method for stops*

---

**Description**

S3 summary method for stops

**Usage**

```
## S3 method for class 'stops'  
summary(object, ...)
```

**Arguments**

object	object of class stops
...	additional arguments

**Value**

object of class 'summary.stops'

---

Swissroll

*Swiss roll*


---

### Description

A swiss roll data example where 150 data points are arranged on a swiss roll embedded in a 3D space.

### Usage

```
data(Swissroll)
```

### Format

A data frame with 150 rows and 4 columns

### Details

A data frame with the variables (columns)

- x The x axis coordinate for each point
- y The y axis coordinate for each point
- z The z axis coordinate for each point
- col a color code for each point with points along the y axis having the same color (based on the viridis palette)

---

tgpoptim

*Bayesian Optimization by a (treed) Bayesian Gaussian Process Prior (with jumps to linear models) surrogate model Essentially a wrapper for the functionality in tgp that has the same slots as optim with defaults for STOPS models.*

---

### Description

Bayesian Optimization by a (treed) Bayesian Gaussian Process Prior (with jumps to linear models) surrogate model Essentially a wrapper for the functionality in tgp that has the same slots as optim with defaults for STOPS models.

**Usage**

```
tgpoptim(
  x,
  fun,
  ...,
  initpoints = 10,
  lower,
  upper,
  acc = 1e-08,
  itmax = 10,
  verbose = 0,
  model = "bgp"
)
```

**Arguments**

x	optional starting values
fun	function to minimize
...	additional arguments to be passed to the function to be optimized
initpoints	the number of points to sample initially to fit the surrogate model
lower	The lower constraints of the search region
upper	The upper constraints of the search region
acc	if the numerical accuracy of two successive target function values is below this, stop the optimization; defaults to 1e-8
itmax	maximum number of iterations
verbose	numeric value that prints information on the fitting process; >2 is extremely verbose
model	which surrogate model class to use (currently uses defaults only, will extend this to tweak the model)

**Value**

A list with the components (for compatibility with `optim`)

- `par` The position of the optimum in the search space (parameters that minimize the function; `argmin fun`).
- `value` The value of the objective function at the optimum (`min fun`). Note we do not use the last value in the candidate list but the best candidate (which can but need not coincide).
- `svalue` The value of the surrogate objective function at the optimal parameters
- `counts` The number of iterations performed at convergence with entries `fnction` for the number of iterations and `gradient` which is always NA at the moment
- `convergence` 0 successful completion by the `accd` or `acc` criterion, 1 indicate iteration limit was reached, 99 is a problem
- `message` is NULL (only for compatibility or future use)
- `history` the improvement history
- `tgput` the output of the `tg` model

**Examples**

```
fbana <- function(x) {  
  x1 <- x[1]  
  x2 <- x[2]  
  100 * (x2 - x1 * x1)^2 + (1 - x1)^2  
}  
res1<-tgpoptim(c(-1.2,1),fbana,lower=c(-5,-5),upper=c(5,5),acc=1e-16,itmax=20)  
res1  
  
fwild <- function (x) 10*sin(0.3*x)*sin(1.3*x^2) + 0.00001*x^4 + 0.2*x+80  
plot(fwild, -50, 50, n = 1000, main = "Bayesian GP Optimization minimizing 'wild function'")  
set.seed(210485)  
res2<-tgpoptim(50, fwild,lower=-50,upper=50,acc=1e-16,itmax=20,model="btgpllm")  
points(res2$par,res2$value,col="red",pch=19)  
res2
```



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