Package 'cops'

November 1, 2019

Title Cluster Optimized Proximity Scaling

Version 1.0-2

Date 2019-10-28

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

Description Cluster optimized proximity scaling (COPS) refers to multidimensional scaling (MDS) methods that aim at pronouncing the clustered appearance of the configuration. They achieve this by transforming proximities/distances with power functions and augment the fitting criterion with a clusteredness index, the OP-TICS Cordillera (Rusch, Hornik & Mair, 2018, <doi:10.1080/10618600.2017.1349664>). There are two variants: One for finding the configuration directly for given parameters (COPS-C) for ratio, interval and non-metric MDS (Borg & Groenen, 2005, ISBN:978-0-387-28981-6), and one for using the augmented fitting criterion to find optimal parameters (P-COPS). The package contains various functions, wrappers, methods and classes for fitting, plotting and displaying different MDS models in a COPS framework like ratio, interval and non-metric MDS for COPS-C and P-COPS with Torgerson scaling (Torgerson, 1958, ISBN:978-0471879459), scaling by majorizing a complex function (SMA-COF; de Leeuw, 1977, <https://escholarship.org/uc/item/4ps3b5mj>), Sammon mapping (Sammon, 1969, <doi:10.1109/T-C.1969.222678>), elastic scaling (McGee, 1966, <doi:10.1111/j.2044-8317.1966.tb00367.x>), sstress (Takane, Young & de Leeuw, 1977, <doi:10.1007/BF02293745>, rstress (de Leeuw, Groenen & Mair, 2016, <https://rpubs.com/deleeuw/142619>), powerstress (Buja & Swayne, 2002 <doi:10.1007/s00357-001-0031-0>) and power elastic scaling, power Sammon mapping and approximated power stress (Rusch, Mair & Hornik, 2015, <https://bachs59.wu.ac.at/4888/>). All of these models can also solely be fit as MDS with power transformations. The package further contains a function for pattern search optimization, the ``Adaptive Luus-Jakola Algorithm" (Rusch, Mair & Hornik, 2015, https://bach-s59.wu.ac.at/4888/>).

Depends R (>= 3.1.2), cordillera (>= 0.7-2), smacof (>= 1.10-4)

Imports MASS, minqa, pso, scatterplot3d, NlcOptim, Rsolnp, dfoptim, subplex, cmaes, crs, nloptr, rgl, rgenoud, GenSA

Enhances stats

Suggests testthat

License GPL-2 | GPL-3

LazyData true

URL http://r-forge.r-project.org/projects/stops/

RoxygenNote 6.1.1

Encoding UTF-8

NeedsCompilation no

Author Thomas Rusch [aut, cre] (<https://orcid.org/0000-0002-7773-2096>), Jan de Leeuw [aut], Patrick Mair [aut]

Repository CRAN

Date/Publication 2019-11-01 09:30:02 UTC

R topics documented:

BankingCrisesDistances														
cmdscale	 	•										•		 3
conf_adjust	 	•										•		
cops	 	•												 5
copstress	 	•												 7
copstressMin	 	•												 8
cop_apstress	 	•										•		 11
cop_cmdscale	 	•						 •				•		 13
cop_elastic	 	•												 14
cop_powerelastic	 	•						 •				•		 15
cop_powermds	 	•										•		 17
cop_powersammon														
cop_powerstress	 	•										•	•	 19
cop_rstress	 	•										•		 21
cop_sammon	 	•										•	•	 22
cop_sammon2														
cop_smacofSphere														
cop_smacofSym	 	•	 •	•••		 •		 •			•	•	•	 26
cop_sstress														
doubleCenter														
enorm	 	•	 •	•••		 •		 •			•	•	•	 29
ljoptim														
mkBmat														
mkPower														
mkPowerALTERN														
pcops														
pdist	 	•	 •	•••		 •		 •			•	•	•	
plot.cops	 	•	 •	•••		 •		 •			•	•	•	 35
plot.pcops	 	•										•	•	 36
plot.smacofP														
plot3d.cmdscale														
plot3dstatic	 	•						 •				•		 40

2

plot3dstatic.cmdscale	 40
powerStressFast	 41
powerStressMin	 43
procruster	 44
sammon	 45
scale_adjust	 45
secularEq	 46
sqdist	 46
torgerson	 47
	48

Index

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)

cmdscale

Wrapper to cmdscale for S3 class

Description

Wrapper to cmdscale for S3 class

Usage

cmdscale(d, k = 2, eig = TRUE, ...)

Arguments

d	a distance structure such as that returned by 'dist' or a full symmetric matrix containing the dissimilarities
k	the maximum dimension of the space which the data are to be represented in
eig	indicates whether eigenvalues should be returned.
	additional parameters passed to cmdscale. See cmdscale

Value

See cmdscale. This wrapper only adds an extra slot to the list with the call, adds column labels to the \$points and assigns S3 class 'cmdscale'.

Examples

```
dis<-as.matrix(smacof::kinshipdelta)
res<-cmdscale(dis)</pre>
```

conf_adjust

conf_adjust: a function to procrustes adjust two matrices

Description

conf_adjust: a function to procrustes adjust two matrices

Usage

```
conf_adjust(conf1, conf2, verbose = FALSE, eps = 1e-12, itmax = 100)
```

Arguments

conf1	reference configuration, a numeric matrix
conf2	another configuration, a numeric matrix
verbose	should adjustment be output; default to FALSE
eps	numerical accuracy
itmax	maximum number of iterations

Value

a list with ref.conf being the reference configuration, other.conf the adjusted coniguration and comparison.conf the comparison configuration

Description

Cluster optimized proximity scaling (COPS) refers to multidimensional scaling methods that aim at pronouncing the clustered appearance of the configuration. They achieve this by transforming proximities/distances with power functions and augment the fitting criterion with a clusteredness index, the OPTICS Cordillera (Rusch, Hornik & Mair 2017). There are two variants: One for finding the configuration directly for given parameters (COPS-C), and one for using the augmented fitting criterion to find optimal parameters for the power transformations (P-COPS). The package contains various functions, wrappers, methods and classes for fitting, plotting and displaying different MDS models in a COPS framework like Torgerson scaling, SMACOF, Sammon mapping, elastic scaling, symmetric SMACOF, spherical SMACOF, sstress, rstress, powernds, power elastic scaling, power sammon mapping, powerstress. All of these models can also solely be fit as MDS with power transformations. The package further contains functions for optimization (Adaptive LJ Algorithmus).

Minimizing copstress for a clustered MDS configuration. Allows to choose COPS-C (finding a configuration from copstress with cordillera penalty) and profile COPS (finding hyperparameters for MDS models with power transformations). It is wrapper for copstressMin and pcops.

Usage

```
cops(dis, variant = c("1", "2", "Variant1", "Variant2", "v1", "v2",
    "COPS-C", "P-COPS", "configuration-c", "profile", "copstress-c",
    "p-copstress", "COPS-P", "copstress-p", "cops-c", "p-cops"), ...)
```

Arguments

dis	a dissimilarity matrix or a dist object
variant	a character string specifying which variant of COPS to fit. Allowed is any of the following "1", "2", "Variant1", "Variant2", "v1", "v2", "COPS-C", "P-COPS", "configuration-c", "profile", "copstress-c", "p-copstress". Defaults to "COPS-C".
	arguments to be passed to copstressMin (for Variant 1) or pcops (for Variant 2).

Details

The cops package provides five categories of important functions:

Models & Algorithms:

- cops() ... high level interface to fit COPS models as described in Rusch et al. (2015). By setting cordweight to zero they can also be used to fit metric MDS for many different models, see below.
- copstressMin()... The workhorse for fitting a COPS-C model. Can also be called directly.
- pcops()... The workhorse for fitting a P-COPS model. Can also be called directly.

cops

 powerStressMin()... a workhorse for fitting s-stress, r-stress (de Leeuw, 2014), p-stress (e.g., Rusch et al., 2015), Sammon mapping with power transformations (powersammon) and elastic scaling with power transformation (powerelastic). They can conveniently also be fitted via the cops functions and setting stressweight=1 and cordweight or by the dedicated functions starting with cops_XXX where XXX is the method and setting stressweight=1 and cordweight=0. It uses the nested majorization algorithm for r-stress of De Leeuw (2014).

Optimization functions:

• ljoptim() ... An (adaptive) version of the Luus-Jakola random search

Wrappers and convenience functions:

- conf_adjust(): procrustes adjustment of configurations
- cmdscale(), sammon(): wrappers for that return S3 objects to be used with cops
- copstress() ... a function to calculate copstress (Rusch et al., 2015)
- cop_smacofSym(), cop_sammon(), cop_cmdscale(), cop_rstress(), cop_powerstress(), cop_smacofSphere(), cop_sammon2(), cop_elastic(), cop_sstress(), cop_powerelastic(), cop_powersammon(): cop versions of these MDS models.

Methods: For most of the objects returned by the high-level functions S3 classes and methods for standard generics were implemented, including print, summary, plot, plot3d, plot3dstatic.

References:

- Rusch, T., Mair, P. \& Hornik, K. (2015) COPS: Cluster optimized proximity scaling, Report 2015/1, Discussion Paper Series, Center for Empirical Research Methods, WU Vienna University of Economics and Business.
- Rusch, T., Mair, P. & Hornik, K. (2017) Assessing and quantifying clusteredness: The OP-TICS Cordillera. Journal of Computational and Graphical Statistics. Forthcoming. doi = 10.1080/10618600.2017.1349664, http://dx.doi.org/10.1080/10618600.2017.1349664

Authors: Thomas Rusch, Jan de Leeuw, Patrick Mair

Maintainer: Thomas Rusch

Value

For COPS-C Variant 1 see copstressMin, for P-COPS Variant 2 see pcops

Examples

```
library(cordillera)
data(BankingCrisesDistances)
```

```
#shorthand function for COPS-C (finding configuration with copstress)
res<-cops(BankingCrisesDistances[,1:69],variant="COPS-C",stressweight=0.98,cordweight=0.02)
res
summary(res)
plot(res)
plot(res, "reachplot")</pre>
```

copstress

```
plot(res,"transplot")
plot(res,"Shepard")
#shorthand function for P-COPS (hyperparameter search for powerstress)
res<-cops(BankingCrisesDistances[,1:69],variant="P-COPS")</pre>
res
summary(res)
plot(res)
plot(res,"reachplot")
plot(res,"transplot")
plot(res,"Shepard")
dis<-as.matrix(smacof::kinshipdelta)</pre>
#COPS-C with equal weight to stress and cordillera
res1<-cops(dis,variant="COPS-C",stressweight=0.5,cordweight=0.5,</pre>
minpts=2,itmax=1000) #use higher itmax in real
res1
summary(res1)
plot(res1)
plot(res1,"reachplot")
#s-stress type copstress (i.e. kappa=2, lambda=2)
res3<-cops(dis,variant="COPS-C",kappa=2,lambda=2,stressweight=0.5,cordweight=0.5)</pre>
res3
summary(res3)
plot(res3)
#power-stress type profile copstress
# search for optimal kappa and lambda between kappa=0.5,lambda=0.5 and kappa=2,lambda=5
# nu is fixed on -1
ws<-1/dis
diag(ws)<-1
res5<-cops(dis,variant="P-COPS",loss="powerstress",</pre>
theta=c(1.4,3,-1),lower=c(1,0.5,-1),upper=c(3,5,-1),weightmat=ws,
stressweight=0.9,cordweight=0.1)
res5
summary(res5)
plot(res5)
```

```
copstress
```

Calculates copstress for given MDS object

Description

Calculates copstress for given MDS object

Usage

```
copstress(obj, stressweight = 1, cordweight = 5, q = 1, minpts = 2,
epsilon = 10, rang = NULL, verbose = 0, normed = TRUE,
scale = c("std", "sd", "proc", "none"), init, ...)
```

Arguments

obj	MDS object (supported are sammon, cmdscale, smacof, rstress, powermds)
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to 2
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is very verbose (copstress level), >3 is extremely (up to MDS optimization level)
normed	should the cordillera be normed; defaults to TRUE
scale	should the configuration be scale adjusted.
init	a reference configuration when doing procrustes adjustment
	additional arguments to be passed to the cordillera function

Value

A list with the components

- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- cordillera: the cordillera object

copstressMin

Fitting a COPS-C Model (COPS Variant 1).

Description

Minimizing Copstress to obtain a clustered MDS configuration with given hyperparameters theta.

copstressMin

Usage

```
copstressMin(delta, kappa = 1, lambda = 1, nu = 1, theta = c(kappa,
lambda, nu), type = c("ratio", "interval", "ordinal"),
ties = "primary", weightmat = 1 - diag(nrow(delta)), ndim = 2,
init = NULL, stressweight = 0.975, cordweight = 0.025, q = 1,
minpts = ndim + 1, epsilon = 10, dmax = NULL, rang,
optimmethod = c("NelderMead", "Newuoa", "BFGS", "SANN", "hjk", "solnl",
"solnp", "subplex", "snomadr", "hjk-Newuoa", "hjk-BFGS", "BFGS-hjk",
"Newuoa-hjk", "cmaes", "direct", "direct-Newuoa", "direct-BFGS",
"genoud", "gensa"), verbose = 0, scale = c("sd", "rmsq", "std",
"proc", "none"), normed = TRUE, accuracy = 1e-07, itmax = 5000,
stresstype = c("stress-1", "stress"), ...)
```

Arguments

delta	numeric matrix or dist object of a matrix of proximities
kappa	power transformation for fitted distances
lambda	power transformation for proximities
nu	power transformation for weights
theta	the theta vector of powers; the first is kappa (for the fitted distances if it exists), the second lambda (for the observed proximities if it exist), the third is nu (for the weights if it exists). If less than three elements are is given as argument, it will be recycled. Defaults to 1 1 1. Will override any kappa, lmabda, nu parameters if they are given and do not match
type	what type of MDS to fit. Currently one of "ratio", "interval" or "ordinal". Default is "ratio".
ties	the handling of ties for ordinal (nonmetric) MDS. Possible are "primary" (de-fault), "secondary" or "tertiary".
weightmat	(optional) a matrix of nonnegative weights; defaults to 1 for all off diagonals
ndim	number of dimensions of the target space
init	(optional) initial configuration
stressweight	weight to be used for the fit measure; defaults to 0.975
cordweight	weight to be used for the cordillera; defaults to 0.025
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
dmax	The winsorization limit of reachability distances in the OPTICS Cordillera. If supplied, it should be either a numeric value that matches max(rang) or NULL; if NULL it is found as 1.5 times (for kappa >1) or 1 times (for kappa <=1) the maximum reachbility value of the power torgerson model with the same lambda. If dmax and rang are supplied and dmax is not max(rang), a warning is given and rang takes precedence.

rang	range of the reachabilities to be considered. If missing it is found from the initial configuration by taking 0 as the lower boundary and dmax (see above) as upper boundary. See also cordillera
optimmethod	What optimizer to use? Choose one string of 'Newuoa' (from package minqa), 'NelderMead', 'hjk' (Hooke-Jeeves algorithm from dfoptim), 'solnl' (from nl-cOptim), 'solnp' (from Rsolnp), 'subplex' (from subplex), 'SANN' (simulated annealing), 'BFGS', 'snomadr' (from crs), 'genoud' (from rgenoud), 'gensa' (from GenSA), 'cmaes' (from cmaes) and 'direct' (from nloptr). See the according R packages for details on these solvers. There are also combinations that proved to work well good, like 'hjk-Newuoa', 'hjk-BFGS', 'BFGS-hjk', 'Newuoa-hjk', 'direct-Newuoa' and 'direct-BFGS'. Usually hjk, BFGS, newuoa, subplex and solnl work rather well in an acceptable time frame (depending on the smoothness of copstress). Default is 'hjk-Newuoa'.
verbose	numeric value hat prints information on the fitting process; >2 is very verbose
scale	Allows to scale the configuration for the OC (the scaled configuration is also returned as \$conf). One of none (so no scaling), sd (configuration divided by the maximum standard deviation of the columns), std (standardize all columns !NOTE: This does not preserve the relative distances of the optimal config), proc (procrustes adjustment to the initial fit) and rmsq (configuration divided by the maximum root mean square of the columns). Default is sd.
normed	should the cordillera be normed; defaults to TRUE
accuracy	numerical accuracy, defaults to 1e-7
itmax	maximum number of iterations. Defaults to 5000.
stresstype	which stress to use in the copstress. Defaults to stress-1. If anything else is set, explicitly normed stress which is (stress-1)^2. Using stress-1 puts more weight on MDS fit.
	additional arguments to be passed to the optimization procedure

Value

A list with the components

- delta: the original transformed dissimilarities
- obsdiss: the explicitly normed transformed dissimilarities (which are approximated by the fit)
- confdist: the fitted distances
- conf: the configuration to which the scaling of argument scale was applied
- confo: the unscaled but explicitly normed configuration returned from the fitting procedure. Scaling applied to confo gives conf.
- par, pars : the theta vector of powers tranformations (kappa,lambda,nu)
- niter: number of iterations of the optimizer.
- stress: the square root of explicitly normalized stress (calculated for confo).
- spp: stress per point
- ndim: number of dimensions
- model: Fitted model name with optimizer

- call: the call
- nobj: the number of objects
- type, loss, losstype: stresstype
- stress.m: The stress used for copstress. If stresstype="stress-1" this is like \$stress else it is stress^2
- · stress.en: another ways to calculate the stress
- · deltaorig: the original untransformed dissimilarities
- copstress: the copstress loss value
- resmat: the matrix of residuals
- weightmat: the matrix of untransformed weights
- OC: the (normed) OPTICS Cordillera object (calculated for scaled conf)
- OCv: the (normed) OPTICS Cordillera value alone (calculated for scaled conf)
- optim: the object returned from the optimization procedure
- stressweight, cordweight: the weights of the stress and OC respectively (v_1 and v_2)
- optimmethod: The solver used
- type: the type of MDS fitted

Examples

dis<-as.matrix(smacof::kinshipdelta)</pre>

```
#Copstress with equal weight to stress and cordillera
res1<-copstressMin(dis,stressweight=0.5,cordweight=0.5,itmax=1000) #use higher itmax about 10000
res1
summary(res1)
plot(res1) #super clustered
```

cop_apstress	PCOPS versions of approximated power stress models. This uses an
	approximation and makes use of smacof.

Description

PCOPS versions of approximated power stress models. This uses an approximation and makes use of smacof.

Usage

```
cop_apstress(dis, theta = c(1, 1, 1), ndim = 2, weightmat = NULL,
init = NULL, itmaxi = 1000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, normed = TRUE, scale,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the tau and upsilon trans- formation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second the tau argument and the third the upsilon argument. Defaults to 1 1 1
ndim	number of dimensions of the target space
weightmat	(optional) a binary matrix of nonnegative weights
init	(optional) initial configuration
itmaxi	number of iterations. default is 1000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
normed	should the cordillera be normed; defaults to TRUE
scale	should the configuration be scale adjusted
stresstype	which stress to report. Only takes smacofs default stress currrently.

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure (which has all smacofB elements and some more
- cordillera: the cordillera object

cop_cmdscale

Description

PCOPS version of strain

Usage

```
cop_cmdscale(dis, theta = c(1, 1, 1), weightmat = NULL, ndim = 2,
init = NULL, itmaxi = 1000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, scale = 3, normed = TRUE,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second and third the lambda and the nu argument (the latter is fixed to 1). Defaults to 1 1 1
weightmat	(optional) a matrix of nonnegative weights
ndim	number of dimensions of the target space
init	(optional) initial configuration
itmaxi	number of iterations. No effect here.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the corrdillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report. Only takes cmdscales default stress currrently.

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_elastic *PCOPS versions of elastic scaling models (via smacofSym)*

Description

PCOPS versions of elastic scaling models (via smacofSym)

Usage

```
cop_elastic(dis, theta = c(1, 1, -2), ndim = 2, weightmat = 1,
init = NULL, itmaxi = 1000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, normed = TRUE, scale = 3,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second the lambda argument and the third the nu argument (here internally fixed to -2). Defaults to $1 \ 1 \ -2$
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. default is 1000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the corrdillera; defaults to 1

14

minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
normed	should the cordillera be normed; defaults to TRUE
scale	should the configuration be scale adjusted
stresstype	which stress to report. Only takes smacofs default stress currrently.

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_powerelastic *PCOPS version of elastic scaling with powers*

Description

PCOPS version of elastic scaling with powers

Usage

```
cop_powerelastic(dis, theta = c(1, 1, -2), weightmat = 1 -
diag(nrow(dis)), init = NULL, ndim = 2, itmaxi = 10000, ...,
stressweight = 1, cordweight = 0.5, q = 1, minpts = ndim + 1,
epsilon = 10, rang = NULL, verbose = 0, scale = 3,
normed = TRUE, stresstype = c("default", "stress1", "rawstress",
"normstress", "enormstress", "enormstress1"))
```

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) and nu as the third (fixed to -2). If a scalar for the free parameters is given it is recycled. Defaults to 1 1 -2.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations. default is 10000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report? Defaults to explicitly normed stress

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_powermds

Description

PCOPS version of powermds

Usage

```
cop_powermds(dis, theta = c(1, 1, 1), weightmat = 1 - diag(nrow(dis)),
init = NULL, ndim = 2, itmaxi = itmaxi, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, scale = 3, normed = TRUE,
stresstype = c("default", "stress1", "rawstress", "normstress",
"enormstress", "enormstress1"))
```

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), nu is fixed to 1. If a scalar is given it is recycled. Defaults to 1 1 1.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations. default is 10000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely ver- bose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report? Defaults to whatever whim is my default (currently explicitly normed stress)

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_powersammon *PCOPS version of sammon with powers*

Description

PCOPS version of sammon with powers

Usage

```
cop_powersammon(dis, theta = c(1, 1, -1), weightmat = 1 -
diag(nrow(dis)), init = NULL, ndim = 2, itmaxi = 10000, ...,
stressweight = 1, cordweight = 0.5, q = 1, minpts = ndim + 1,
epsilon = 10, rang = NULL, verbose = 0, scale = 3,
normed = TRUE, stresstype = c("default", "stress1", "rawstress",
"normstress", "enormstress", "enormstress1"))
```

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 (fixed to -1). If a scalar is given it is recycled for the free parameters. Defaults to 1 1 -1.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations. default is 10000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1

18

cop_powerstress

epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report? Defaults to explicitly normed stress

Value

A list with the components

- stress: the stress
- · stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_powerstress

COPS version of powerstress

Description

COPS version of powerstress

Usage

```
cop_powerstress(dis, theta = c(1, 1, 1), weightmat = 1 -
diag(nrow(dis)), init = NULL, ndim = 2, itmaxi = 10000, ...,
stressweight = 1, cordweight = 0.5, q = 1, minpts = ndim + 1,
epsilon = 10, rang = NULL, verbose = 0, scale = "sd",
normed = TRUE, stresstype = c("default", "stress1", "rawstress",
"normstress", "enormstress", "enormstress1"))
```

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.
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations. default is 10000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report? Defaults to explicitly normed stress

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_rstress

Description

PCOPS version of rstress

Usage

```
cop_rstress(dis, theta = c(1, 1, 1), weightmat = 1 - diag(nrow(dis)),
init = NULL, ndim = 2, itmaxi = 10000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, scale = 3, normed = TRUE,
stresstype = c("default", "stress1", "rawstress", "normstress",
"enormstress", "enormstress1"))
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the kappa transformation for the fitted distances proximities, or a vector where the first is the kappa argument for the fitted distances and the second the lambda argument, the third the nu argument (here internally fixed to 1). Defaults to 1 1 1
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations. default is 10000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
epsilon rang	· · ·
·	to 10 range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an
rang	to 10 range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness. numeric value hat prints information on the fitting process; >2 is extremely ver-
rang verbose	to 10 range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness. numeric value hat prints information on the fitting process; >2 is extremely ver- bose

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_sammon *PCOPS version of sammon mapping*

Description

PCOPS version of sammon mapping

Usage

```
cop_sammon(dis, theta = c(1, 1, -1), ndim = 2, init = NULL,
weightmat = NULL, itmaxi = 100, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, scale = 3, normed = TRUE,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second the lambda argument and the third the nu argument (here internally fixed to -1). Defaults to $1 \ 1 \ -1$
ndim	number of dimensions of the target space
init	(optional) initial configuration
weightmat	(optional) a matrix of nonnegative weights
itmaxi	number of iterations. default is 1000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the corrdillera; defaults to 1

22

minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report. Only takes smacofs default stress currently.

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_sammon2

COPS versions of Sammon mapping models (via smacofSym)

Description

COPS versions of Sammon mapping models (via smacofSym)

Usage

```
cop_sammon2(dis, theta = c(1, 1, -1), ndim = 2, weightmat = NULL,
init = NULL, itmaxi = 1000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, normed = TRUE, scale = 3,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second the lambda argument, the thrid the nu argument (fixed to -1). Defaults to 1 1 -1.
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights (NOT the sammon weights)
init	(optional) initial configuration
itmaxi	number of iterations. default is 1000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the corrdillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
normed	should the cordillera be normed; defaults to TRUE
scale	should the configuration be scale adjusted
stresstype	which stress to report. Only takes smacofs default stress currrently.

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_smacofSphere PCOPS versions of smacofSphere models

Description

PCOPS versions of smacofSphere models

Usage

```
cop_smacofSphere(dis, theta = c(1, 1, 1), ndim = 2, weightmat = NULL,
init = NULL, itmaxi = 1000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, normed = TRUE, scale = 3,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second the lambda argument and the third the nu argument. Defaults to 1 1 1
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
itmaxi	number of iterations. default is 1000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the corrdillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
normed	should the cordillera be normed; defaults to TRUE
scale	should the configuration be scale adjusted
stresstype	which stress to report. Only takes smacofs default stress currently.

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

cop_smacofSym PCOPS versions of smacofSym models

Description

PCOPS versions of smacofSym models

Usage

```
cop_smacofSym(dis, theta = c(1, 1, 1), ndim = 2, weightmat = NULL,
init = NULL, itmaxi = 1000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, normed = TRUE, scale,
stresstype = "default")
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 1) and the second the lambda argument and the third the nu argument (here internally fixed to 1). Defaults to $1 \ 1 \ 1$
ndim	number of dimensions of the target space
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
itmaxi	number of iterations. default is 1000
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1

26

cop_sstress

minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
normed	should the cordillera be normed; defaults to TRUE
scale	should the configuration be scale adjusted
stresstype	which stress to report. Only takes smacofs default stress currrently.

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure (which has all smacofB elements and some more
- cordillera: the cordillera object

cop_sstress

PCOPS version of sstress

Description

PCOPS version of sstress

Usage

```
cop_sstress(dis, theta = c(2, 1, 1), weightmat = 1 - diag(nrow(dis)),
init = NULL, ndim = 2, itmaxi = 10000, ..., stressweight = 1,
cordweight = 0.5, q = 1, minpts = ndim + 1, epsilon = 10,
rang = NULL, verbose = 0, scale = 3, normed = TRUE,
stresstype = c("default", "stress1", "rawstress", "normstress",
"enormstress", "enormstress1"))
```

Arguments

dis	numeric matrix or dist object of a matrix of proximities
theta	the theta vector of powers; this is either a scalar of the lambda transformation for the observed proximities, or a vector where the first is the kappa argument for the fitted distances (here internally fixed to 2) and the second the lambda argument and the third the nu argument (internally fixed to 1). Defaults to $2\ 1\ 1$
weightmat	(optional) a matrix of nonnegative weights
init	(optional) initial configuration
ndim	number of dimensions of the target space
itmaxi	number of iterations. default is 10000.
	additional arguments to be passed to the fitting procedure
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; defaults to 0.5
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the distances (min distance minus max distance). If NULL (default) the cordillera will be normed to each configuration's maximum distance, so an absolute value of goodness-of-clusteredness.
verbose	numeric value hat prints information on the fitting process; >2 is extremely verbose
scale	should the configuration be scale adjusted
normed	should the cordillera be normed; defaults to TRUE
stresstype	which stress to report? Defaults to explicitly normed stress

Value

A list with the components

- stress: the stress
- stress.m: default normalized stress
- copstress: the weighted loss value
- OC: the Optics cordillera value
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

doubleCenter

Description

Double centering of a matrix

Usage

doubleCenter(x)

Arguments

x numeric matrix

Value

the double centered matrix

enorm

Explicit Normalization Normalizes distances

Description

Explicit Normalization Normalizes distances

Usage

enorm(x, w = 1)

Arguments

х	numeric matrix
w	weight

Value

a constant

```
ljoptim
```

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

х	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 contraints of the search region
upper	The upper contraints 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 hat prints information on the fitting process; >2 is extremely ver- bose
adaptive	should the adaptive version be used? defaults to TRUE.

Value

A list with the components (see also optim)

- par The position of the optimimum 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)

mkBmat

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</pre>
```

```
mkBmat
```

Auxfunction1 only used internally

Description

Auxfunction1 only used internally

Usage

mkBmat(x)

Arguments

x matrix

Value

a matrix

mkPower

Take matrix to a power

Description

Take matrix to a power

Usage

mkPower(x, r)

Arguments

х	matrix
r	numeric (power)

Value

a matrix

mkPowerALTERN Take matrix to a power

Description

Take matrix to a power

Usage

mkPowerALTERN(x, r)

Arguments

Х	matrix
r	numeric (power)

Value

a matrix

```
pcops
```

Profile COPS Function (aka COPS Variant 2)

Description

Metaparameter selection for MDS models baseed on the Profile COPS approach (COPS Variant 2). It uses copstress for hyperparameter selection. It is a special case of a STOPS model.

Usage

```
pcops(dis, loss = c("stress", "smacofSym", "smacofSphere", "strain",
  "sammon", "rstress", "powermds", "sstress", "elastic", "powersammon",
  "powerelastic", "powerstress", "sammon2", "powerstrain", "apstress"),
  weightmat = NULL, ndim = 2, init = NULL, theta = c(1, 1, 1),
  stressweight = 1, cordweight, q = 2, minpts = ndim + 1,
  epsilon = 100, rang, optimmethod = c("ALJ", "pso", "SANN", "DIRECT",
  "DIRECTL", "stogo", "MADS", "hjk"), lower = c(1, 1, 0.5),
  upper = c(5, 5, 2), verbose = 0, scale = c("proc", "sd", "none",
  "std"), normed = TRUE, s = 4, stresstype = "default",
  acc = 1e-07, itmaxo = 200, itmaxi = 10000, ...)
```

32

pcops

pcops

Arguments	
dis	numeric matrix or dist object of a matrix of proximities
loss	which loss function to be used for fitting, defaults to strain. Currently allows for the following models:
	• Power transformations of observed proximities only: Strain loss or classi- cal scaling (strain, workhorse is cmdscale), Kruskall's stress for symmet- ric matrices (smacofSym or stress and smacofSphere for scaling onto a sphere; workhorse is smacof), Sammon mapping (sammon or sammon2; for the earlier the workhorse is sammon from MASS for the latter it is smacof), elastic scaling (elastic, the workhorse is smacof), Takane et al's S-Stress sstress (workhorse is powerstressMin)
	• Power transformations of fitted distances only: De Leeuw's r-stress rstress (workhorse is powerstressMin)
	 Power transformations of fitted distances and observed proximities: Power- mds powermds, Sammon mapping and elastic scaling with powers (powersammon, powerelastic, powerstress; workhorse is powerstressMin) Approximation to power stress: Approximated power stress (apstress;
	workhorse is smacof)
weightmat	(optional) a matrix of nonnegative weights; defaults to 1 for all off diagonals
ndim	number of dimensions of the target space
init	(optional) initial configuration. If not supplied, the Torgerson scaling result of the dissimilarity matrix dis^theta[2]/enorm(dis^theta[2],weightmat) is used.
theta	the theta vector of powers; the first is kappa (for the fitted distances if it exists), the second lambda (for the observed proximities if it exist), the third is nu (for the weights if it exists). If a scalar is given as argument, it will take the role designated by the loss argument. Defaults to 1 1 1
stressweight	weight to be used for the fit measure; defaults to 1
cordweight	weight to be used for the cordillera; if missing gets estimated from the initial configuration so that copstress = 0 for theta= $c(1,1)$
q	the norm of the cordillera; defaults to 1
minpts	the minimum points to make up a cluster in OPTICS; defaults to ndim+1
epsilon	the epsilon parameter of OPTICS, the neighbourhood that is checked; defaults to 10
rang	range of the minimum reachabilities to be considered. If missing it is found from the initial configuration by taking 1.5 times the maximal minimum reachability of the model with theta= $c(1,1)$. If NULL it will be normed to each configuration's minimum and maximum distance, so an absolute value of goodness-of-clusteredness. Note that the latter is not necessarily desirable when comparing configurations for their relative clusteredness. See also cordillera
optimmethod	What general purpose optimizer to use? Defaults to our adaptive LJ version (ALJ). Also allows particle swarm optimization with s particles ("pso") and simulated annealing ("SANN"), "DIRECT" and "DIRECTL", Hooke-Jeeves ("hjk"), StoGo ("stogo"), and "MADS". We recommend not using SANN and pso with the rstress, sstress and the power stress models. We amde good experiences with ALJ, stogo, DIRECT and DIRECTL and also MADS.

lower	The lower contraints of the search region
upper	The upper contraints of the search region
verbose	numeric value hat prints information on the fitting process; >2 is extremely ver- bose
scale	should the configuration be scaled and/or centered for calculating the cordillera? "std" standardizes each column of the configurations to mean=0 and sd=1, "sd" scales the configuration by the maximum standard devation of any column, "proc" adjusts the fitted configuration to the init configuration (or the Togerson scaling solution if init=NULL). This parameter only has an effect for calculating the cordillera, the fitted and returned configuration is NOT scaled.
normed	should the cordillera be normed; defaults to TRUE
S	number of particles if pso is used
stresstype	what stress to be used for comparisons between solutions. Currently not imple- mented and pcops uses explicitly normalized stress for copstress (not stress-1). Stress-1 is reported by the print function though.
асс	termination threshold difference of two successive outer minimization steps.
itmaxo	iterations of the outer step (optimization over the hyperparmeters; if solver allows it). Defaults to 200.
itmaxi	iterations of the inner step (optimization of the MDS). Defaults to 10000 (which is huge).
	additional arguments to be passed to the optimization procedure

Value

A list with the components

- copstress: the weighted loss value
- OC: the Optics cordillera
- optim: the object returned from the optimization procedure
- stress: the stress (square root of stress.m)
- stress.m: default normalized stress
- parameters: the parameters used for fitting (kappa, lambda)
- fit: the returned object of the fitting procedure
- cordillera: the cordillera object

Examples

```
dis<-as.matrix(smacof::kinshipdelta)
set.seed(210485)
#configuration is scaled with highest column sd for calculating cordilera
res1<-pcops(dis,loss="strain",lower=0.1,upper=5,minpts=2)
res1
summary(res1)
plot(res1)</pre>
```

pdist

Description

Squared p-distances

Usage

pdist(x, p)

Arguments

Х	numeric matrix
р	p>0 the Minkoswki distance

Value

squared Minkowski distance matrix

plot.cops	S3 plot method for cops objects	
p=00100p0	Se provincentou joi cops cojecis	

Description

S3 plot method for cops objects

Usage

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

Arguments

х	an object of class cops
plot.type	String indicating which type of plot to be produced: "confplot", "reachplot", "resplot", "transplot", "Shepard", "stressplot" (see details)
main	the main title of the plot
asp	aspect ratio of x/y axis; defaults to NA; setting to 1 will lead to an accurate representation of the fitted distances.
	Further plot arguments passed: see 'plot.smacof' and 'plot' for detailed infor- mation.
	Details:
	• Configuration plot (plot type - "configlet"); Plots the MDS configurations

- Reachability plot (plot.type = "confplot"): Plots the OPTICS reachability plot and the OPTICS cordillera
- Residual plot (plot.type = "resplot"): Plots the dissimilarities against the fitted distances.
- Linearized Shepard diagram (plot.type = "Shepard"): Diagram with the transformed observed dissimilarities against the transformed fitted distance as well as loess smooth and a least squares line.
- Transformation Plot (plot.type = "transplot"): Diagram with the observed dissimilarities (lighter) and the transformed observed dissimilarities (darker) against the fitted distances together with loess smoothing lines
- Stress decomposition plot (plot.type = "stressplot", only for SMACOF objects in \$fit): Plots the stress contribution in of each observation. Note that it rescales the stress-per-point (SPP) from the corresponding smacof function to percentages (sum is 100). The higher the contribution, the worse the fit.
- Bubble plot (plot.type = "bubbleplot", only available for SMACOF objects \$fit): Combines the configuration plot with the point stress contribution. The larger the bubbles, the better the fit.

Examples

```
dis<-as.matrix(smacof::kinshipdelta)
resl<-copstressMin(dis,itmax=20)
plot(resl)</pre>
```

plot.pcops S3 plot method for p-cops objects

Description

S3 plot method for p-cops objects

Usage

```
## S3 method for class 'pcops'
plot(x, plot.type = c("confplot"), main, asp = NA, ...)
```

Arguments

х	an object of class cops
plot.type	String indicating which type of plot to be produced: "confplot", "reachplot", "resplot", "transplot", "Shepard", "stressplot" (see details)
main	the main title of the plot
asp	aspect ratio of x/y axis; defaults to NA; setting to 1 will lead to an accurate representation of the fitted distances.

. . .

Further plot arguments passed: see 'plot.smacof' and 'plot' for detailed information.

Details:

- Configuration plot (plot.type = "confplot"): Plots the MDS configurations.
- Reachability plot (plot.type = "confplot"): Plots the OPTICS reachability plot and the OPTICS cordillera
- Residual plot (plot.type = "resplot"): Plots the dissimilarities against the fitted distances.
- Linearized Shepard diagram (plot.type = "Shepard"): Diagram with the transformed observed dissimilarities against the transformed fitted distance as well as loess smooth and a least squares line.
- Transformation Plot (plot.type = "transplot"): Diagram with the observed dissimilarities (lighter) and the transformed observed dissimilarities (darker) against the fitted distances together with loess smoothing lines
- Stress decomposition plot (plot.type = "stressplot", only for SMACOF objects in \$fit): Plots the stress contribution in of each observation. Note that it rescales the stress-per-point (SPP) from the corresponding smacof function to percentages (sum is 100). The higher the contribution, the worse the fit.
- Bubble plot (plot.type = "bubbleplot", only available for SMACOF objects \$fit): Combines the configuration plot with the point stress contribution. The larger the bubbles, the better the fit.

Examples

```
dis<-as.matrix(smacof::kinshipdelta)
resl<-pcops(dis,loss="strain",lower=0.1,upper=5,minpts=2)
plot(resl)
plot(resl,plot.type="Shepard")</pre>
```

plot.smacofP S3 plot method for smacofP objects

Description

S3 plot method for smacofP objects

Usage

```
## S3 method for class 'smacofP'
plot(x, plot.type = "confplot", plot.dim = c(1, 2),
    bubscale = 5, col, label.conf = list(label = TRUE, pos = 3, col = 1,
    cex = 0.8), identify = FALSE, type = "p", pch = 20, asp = 1,
    main, xlab, ylab, xlim, ylim, legend = TRUE, legpos, loess = TRUE,
    ...)
```

Arguments

х	an object of class smacofP	
plot.type	String indicating which type of plot to be produced: "confplot", "resplot", "Shep- ard", "stressplot", "transplot", "bubbleplot" (see details)	
plot.dim	dimensions to be plotted in confplot; defaults to $c(1, 2)$	
bubscale	Scaling factor (size) for the bubble plot	
col	vector of colors for the points	
label.conf	List with arguments for plotting the labels of the configurations in a configura- tion plot (logical value whether to plot labels or not, label position, label color)	
identify	If 'TRUE', the 'identify()' function is called internally that allows to add con- figuration labels by mouse click	
type	What type of plot should be drawn (see also 'plot')	
pch	Plot symbol	
asp	Aspect ratio; defaults to 1 so distances between x and y are represented accurately; can lead to slighlty weird looking plots if the variance on one axis is much smaller than on the other axis; use NA if the standard type of R plot is wanted where the ylim and xlim arguments define the aspect ratio - but then the distances seen are no longer accurate	
main	plot title	
xlab	label of x axis	
ylab	label of y axis	
xlim	scale of x axis	
ylim	scale of y axis	
legend	Flag whether legends should be drawn for plots that have legends	
legpos	Position of legend in plots with legends	
loess	should loess fit be added to Shepard plot	
	Further plot arguments passed: see 'plot.smacof' and 'plot' for detailed infor- mation. Details:	
	 Configuration plot (plot.type = "confplot"): Plots the MDS configurations. Residual plot (plot.type = "resplot"): Plots the dissimilarities against the fitted distances. 	
	• Linearized Shepard diagram (plot.type = "Shepard"): Diagram with the transformed observed dissimilarities against the transformed fitted distance as well as loess curve and a least squares line.	
	• Transformation Plot (plot.type = "transplot"): Diagram with the observed dissimilarities (lighter) and the transformed observed dissimilarities (darker) against the fitted distances together with the nonlinear regression curve	
	• Stress decomposition plot (plot.type = "stressplot"): Plots the stress con- tribution in of each observation. Note that it rescales the stress-per-point (SPP) from the corresponding smacof function to percentages (sum is 100). The higher the contribution, the worse the fit.	
	• Bubble plot (plot.type = "bubbleplot"): Combines the configuration plot with the point stress contribution. The larger the bubbles, the better the fit.	

plot3d.cmdscale

Examples

```
dis<-as.matrix(smacof::kinshipdelta)
res<-powerStressMin(dis)
plot(res)</pre>
```

plot3d.cmdscale 3D plots: plot3d method for class cmdscale

Description

This methods produces a dynamic 3D configuration plot.

Usage

```
## S3 method for class 'cmdscale'
plot3d(x, plot.dim = c(1, 2, 3), xlab, ylab, zlab,
    col, main, bgpng = NULL, ax.grid = TRUE, sphere.rgl = FALSE, ...)
```

Arguments

х	object of class cmdscale
plot.dim	vector of length 3 with dimensions to be plotted
xlab	label of x axis
ylab	label of y axis
zlab	label of z axis
col	color of the text labels
main	plot title
bgpng	Background image from rgl library; 'NULL' for white background
ax.grid	If 'TRUE', axes grid is plotted.
sphere.rgl	If 'TRUE', rgl sphere (background) is plotted.
	Further plot arguments passed: see 'plot3d' in package 'rgl' for detailed infor- mation.

plot3dstatic

Description

A static 3d plot S3 generic

Usage

```
plot3dstatic(x, plot.dim = c(1, 2, 3), main, xlab, ylab, zlab, col, ...)
```

Arguments

х	object
plot.dim	dimensions to plot
main	main title
xlab	label for x axis
ylab	label for y axis
zlab	label for z axis
col	color
	other arguments

Details

A static 3d plot

plot3dstatic.cmdscale 3D plots: plot3dstatic method for class cmdscale

Description

This methods produces a static 3D configuration plot.

Usage

```
## S3 method for class 'cmdscale'
plot3dstatic(x, plot.dim = c(1, 2, 3), main, xlab,
    ylab, zlab, col, ...)
```

powerStressFast

Arguments

x	object of class cmdscale
plot.dim	vector of length 3 with dimensions to be plotted
main	plot title
xlab	label of x axis
ylab	label of y axis
zlab	label of z axis
col	color of the text labels
	Further plot arguments passed: see 'scatterplot3d' in package 'scatterplot3d' for detailed information.

powerStressFast	Power stress minimization by NEWUOA	
-----------------	-------------------------------------	--

Description

An implementation to minimize power stress by a derivative-free trust region optimization algorithm (NEWUOA). Much faster than majorizing as used in powerStressMin but perhaps less accurate.

Usage

```
powerStressFast(delta, kappa = 1, lambda = 1, nu = 1, weightmat = 1
  - diag(nrow(delta)), init = NULL, ndim = 2, acc = 1e-12,
  itmax = 50000, verbose = FALSE)
```

Arguments

delta	dist object or a symmetric, numeric data.frame or matrix of distances
kappa	power of the transformation of the fitted distances; defaults to 1
lambda	the power of the transformation of the proximities; defaults to 1
nu	the power of the transformation for weightmat; defaults to 1
weightmat	a matrix of finite weights
init	starting configuration
ndim	dimension of the configuration; defaults to 2
асс	The smallest value of the trust region radius that is allowed. If not defined, then 1e-10 will be used.
itmax	maximum number of iterations. Default is 50000.
verbose	should iteration output be printed; if > 1 then yes

Value

a smacofP object (inheriting form smacofB, see smacofSym). It is a list with the components

- · delta: Observed dissimilarities, not normalized
- · obsdiss: Observed dissimilarities, normalized
- · confdist: Configuration dissimilarities, NOT normalized
- conf: Matrix of fitted configuration, NOT normalized
- stress: Default stress (stress 1, square root of the explicitly normalized stress on the normalized, transformed dissimilarities)
- spp: Stress per point (based on stress.en)
- ndim: Number of dimensions
- model: Name of smacof model
- niter: Number of iterations
- nobj: Number of objects
- type: Type of MDS model

and some additional components

- gamma: Empty
- stress.m: default stress for the COPS and STOP. Defaults to the explicitly normalized stress on the normalized, transformed dissimilarities
- stress.en: explicitly stress on the normalized, transformed dissimilarities and normalized transformed distances
- · deltaorig: observed, untransformed dissimilarities
- weightmat: weighting matrix

See Also

smacofSym

Examples

```
dis<-smacof::kinshipdelta
res<-powerStressFast(as.matrix(dis),kappa=2,lambda=1.5)
res
summary(res)
plot(res)</pre>
```

powerStressMin

Description

An implementation to minimize power stress by minimization-majorization. Usually more accurate but slower than powerStressFast. Uses a repeat loop.

Usage

```
powerStressMin(delta, kappa = 1, lambda = 1, nu = 1, weightmat = 1
  - diag(nrow(delta)), init = NULL, ndim = 2, acc = 1e-10,
  itmax = 50000, verbose = FALSE)
```

Arguments

delta	dist object or a symmetric, numeric data.frame or matrix of distances
kappa	power of the transformation of the fitted distances; defaults to 1
lambda	the power of the transformation of the proximities; defaults to 1
nu	the power of the transformation for weightmat; defaults to 1
weightmat	a matrix of finite weights
init	starting configuration
ndim	dimension of the configuration; defaults to 2
acc	numeric accuracy of the iteration
itmax	maximum number of iterations. Default is 50000.
verbose	should iteration output be printed; if > 1 then yes

Value

a smacofP object (inheriting form smacofB, see smacofSym). It is a list with the components

- · delta: Observed dissimilarities, not normalized
- · obsdiss: Observed dissimilarities, normalized
- confdist: Configuration dissimilarities, NOT normalized
- conf: Matrix of fitted configuration, NOT normalized
- stress: Default stress (stress 1; sqrt of explicitly normalized stress)
- spp: Stress per point (based on stress.en)
- ndim: Number of dimensions
- model: Name of smacof model
- niter: Number of iterations
- nobj: Number of objects
- type: Type of MDS model

and some additional components

- stress.m: default stress for the COPS and STOP defaults to the explicitly normalized stress on the normalized, transformed dissimilarities
- stress.en: a manually calculated stress on the normalized, transformed dissimilarities and normalized transformed distances which is not correct
- deltaorig: observed, untransformed dissimilarities
- weightmat: weighting matrix

See Also

smacofSym

Examples

```
dis<-smacof::kinshipdelta
res<-powerStressMin(as.matrix(dis),kappa=2,lambda=1.5,itmax=1000)
res
summary(res)
plot(res)</pre>
```

procruster

procruster: a procrustes function

Description

procruster: a procrustes function

Usage

procruster(x)

Arguments

x numeric matrix

Value

a matrix

sammon

Description

Wrapper to sammon for S3 class

Usage

sammon(d, y = NULL, k = 2, ...)

Arguments

d	a distance structure such as that returned by 'dist' or a full symmetric matrix. Data are assumed to be dissimilarities or relative distances, but must be positive except for self-distance. This can contain missing values.
У	An initial configuration. If NULL, 'cmdscale' is used to provide the classical solution. (If there are missing values in 'd', an initial configuration must be provided.) This must not have duplicates.
k	The dimension of the configuration
	Additional parameters passed to sammon, see sammon

Value

See sammon. This wrapper only adds an extra slot to the list with the call, adds column labels to the \$points and assigns S3 classes 'sammon', 'cmdscale'. It also adds a slot obsdiss with normalized dissimilarities.

Examples

dis<-as.matrix(smacof::kinshipdelta)
res<-sammon(dis)</pre>

scale_adjust Adjusts a configuration

Description

Adjusts a configuration

Usage

```
scale_adjust(conf, ref, scale = c("sd", "std", "proc", "none"))
```

Arguments

conf	a configuration
ref	a reference configuration (only for scale="proc")
scale	Scale adjustment. "std" standardizes each column of the configurations to mean=0 and sd=1, "sd" scales the configuration by the maximum standard devation of any column, "proc" adjusts the fitted configuration to the reference

Value

The scale adjusted configuration.

secularEq Secular Equation
Description
Secular Equation
Usage

secularEq(a, b)

Arguments

а	matrix
b	matrix

Value

a matrix

sqdist

Squared distances

Description

Squared distances

Usage

sqdist(x)

Arguments

x numeric matrix

torgerson

Value

squared distance matrix

torgerson Torgerson scaling

Description

Torgerson scaling

Usage

torgerson(delta, p = 2)

Arguments

delta	symmetric, numeric matrix of distances
р	target space dimensions

Value

a n x p matrix (the configuration)

Examples

dis<-as.matrix(smacof::kinshipdelta)
res<-torgerson(dis)</pre>

Index

*Topic clustering cops, 5 copstressMin, 8 pcops, 32 *Topic **multivariate** cop_apstress, 11 cop_cmdscale, 13 cop_elastic, 14 cop_powerelastic, 15 cop_powermds, 17 cop_powersammon, 18 cop_powerstress, 19 cop_rstress, 21 cop_sammon, 22 cop_sammon2, 23 cop_smacofSphere, 25 cop_smacofSym, 26 cop_sstress, 27 cops, 5 copstress, 7 copstressMin, 8 pcops, 32 BankingCrisesDistances, 3 cmdscale, 3, 4 conf_adjust, 4

```
conn_adjust, 4
cop_apstress, 11
cop_cmdscale, 13
cop_elastic, 14
cop_powerelastic, 15
cop_powersdmmon, 18
cop_powerstress, 19
cop_rstress, 21
cop_sammon, 22
cop_sammon2, 23
cop_smacofSphere, 25
cop_smacofSym, 26
cop_sstress, 27
```

cops, 5 cops-package (cops), 5 copstress, 7 copstressMin, 5, 6, 8 cordillera, 10, 33 doubleCenter, 29 enorm, 29 ljoptim, 30 mkBmat, 31mkPower, 31 mkPowerALTERN, 32 optim, *30* pcops, 5, 6, 32 pdist, 35 plot.cops, 35 plot.pcops, 36 plot.smacofP, 37 plot3d.cmdscale, 39 plot3dstatic, 40 plot3dstatic.cmdscale, 40 powerStressFast, 41 powerStressMin, 43 procruster, 44 sammon, 45, 45 scale_adjust, 45 secularEq, 46 smacofSym, 42-44 sqdist, 46

torgerson, 47