

# Package ‘maotai’

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**Type** Package

**Title** Tools for Matrix Algebra, Optimization and Inference

**Version** 0.2.0

**Description** Matrix is an universal and sometimes primary object/unit in applied mathematics and statistics. We provide a number of algorithms for selected problems in optimization and statistical inference. For general exposition to the topic with focus on statistical context, see the book by Banerjee and Roy (2014, ISBN:9781420095388).

**Encoding** UTF-8

**License** MIT + file LICENSE

**Suggests** igraph

**Imports** Matrix, Rcpp, Rdpack, RSpectra, Rtsne, RANN, cluster, labdsv, shapes, stats, utils, fastcluster, dbscan

**LinkingTo** Rcpp, RcppArmadillo, RcppDist

**RdMacros** Rdpack

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bmds

*Bayesian Multidimensional Scaling*


---

## Description

A Bayesian formulation of classical Multidimensional Scaling is presented. Even though this method is based on MCMC sampling, we only return maximum a posterior (MAP) estimate that maximizes the posterior distribution. Due to its nature without any special tuning, increasing `mc.iter` requires much computation.

## Usage

```
bmds(
  data,
  ndim = 2,
  par.a = 5,
  par.alpha = 0.5,
  par.step = 1,
  mc.iter = 8128,
  verbose = TRUE
)
```

**Arguments**

<code>data</code>	an $(n \times p)$ matrix whose rows are observations.
<code>ndim</code>	an integer-valued target dimension.
<code>par.a</code>	hyperparameter for conjugate prior on variance term, i.e., $\sigma^2 \sim IG(a, b)$ . Note that $b$ is chosen appropriately as in paper.
<code>par.alpha</code>	hyperparameter for conjugate prior on diagonal term, i.e., $\lambda_j \sim IG(\alpha, \beta_j)$ . Note that $\beta_j$ is chosen appropriately as in paper.
<code>par.step</code>	stepsize for random-walk, which is standard deviation of Gaussian proposal.
<code>mc.iter</code>	the number of MCMC iterations.
<code>verbose</code>	a logical; TRUE to show iterations, FALSE otherwise.

**Value**

a named list containing

**embed** an  $(n \times ndim)$  matrix whose rows are embedded observations.

**stress** discrepancy between embedded and original data as a measure of error.

**References**

Oh M, Raftery AE (2001). "Bayesian Multidimensional Scaling and Choice of Dimension." *Journal of the American Statistical Association*, **96**(455), 1031–1044.

**Examples**

```
## use simple example of iris dataset
data(iris)
idata = as.matrix(iris[,1:4])

## run Bayesian MDS
# let's run 10 iterations only.
iris.cmds = cmds(idata, ndim=2)
iris.bmds = bmds(idata, ndim=2, mc.iter=5, par.step=(2.38^2))

## extract coordinates and class information
cx = iris.cmds$embed # embedded coordinates of CMDS
bx = iris.bmds$embed # BMDS
icol = iris[,5] # class information

## visualize
opar <- par(no.readonly=TRUE)
par(mfrow=c(2,1))
mc = paste0("CMDS with STRESS=",round(iris.cmds$stress,4))
mb = paste0("BMDS with STRESS=",round(iris.bmds$stress,4))
plot(cx, col=icol,pch=19,main=mc)
plot(bx, col=icol,pch=19,main=mb)
par(opar)
```

---

boot.mblock	<i>Generate Index for Moving Block Bootstrapping</i>
-------------	--

---

### Description

Assuming data being dependent with cardinality  $N$ , `boot.mblock` returns a vector of index that is used for moving block bootstrapping.

### Usage

```
boot.mblock(N, b = max(2, round(N/10)))
```

### Arguments

N	the number of observations.
b	the size of a block to be drawn.

### Value

a vector of length  $N$  for moving block bootstrap sampling.

### References

Kunsch HR (1989). "The Jackknife and the Bootstrap for General Stationary Observations." *The Annals of Statistics*, **17**(3), 1217–1241.

### Examples

```
## example : bootstrap confidence interval of mean and variances
vec.x = seq(from=0,to=10,length.out=100)
vec.y = sin(1.21*vec.x) + 2*cos(3.14*vec.x) + rnorm(100,sd=1.5)
data.mu = mean(vec.y)
data.var = var(vec.y)

## apply moving block bootstrapping
nreps = 50
vec.mu = rep(0,nreps)
vec.var = rep(0,nreps)
for (i in 1:nreps){
  sample.id = boot.mblock(100, b=10)
  sample.y = vec.y[sample.id]
  vec.mu[i] = mean(sample.y)
  vec.var[i] = var(sample.y)
  print(paste("iteration ",i,"/",nreps," complete.", sep=""))
}

## visualize
opar <- par(no.readonly=TRUE)
```

```
par(mfrow=c(1,3), pty="s")
plot(vec.x, vec.y, type="l", main="1d signal") # 1d signal
hist(vec.mu, main="mean CI", xlab="mu")      # mean
abline(v=data.mu, col="red", lwd=4)
hist(vec.var, main="variance CI", xlab="sigma") # variance
abline(v=data.var, col="blue", lwd=4)
par(opar)
```

---

boot.stationary      *Generate Index for Stationary Bootstrapping*

---

## Description

Assuming data being dependent with cardinality  $N$ , `boot.stationary` returns a vector of index that is used for stationary bootstrapping. To describe, starting points are drawn from uniform distribution over  $1:N$  and the size of each block is determined from geometric distribution with parameter  $p$ .

## Usage

```
boot.stationary(N, p = 0.25)
```

## Arguments

<code>N</code>	the number of observations.
<code>p</code>	parameter for geometric distribution with the size of each block.

## Value

a vector of length  $N$  for moving block bootstrap sampling.

## References

Politis DN, Romano JP (1994). "The Stationary Bootstrap." *Journal of the American Statistical Association*, **89**(428), 1303. ISSN 01621459, doi: [10.2307/2290993](https://doi.org/10.2307/2290993).

## Examples

```
## example : bootstrap confidence interval of mean and variances
vec.x = seq(from=0, to=10, length.out=100)
vec.y = sin(1.21*vec.x) + 2*cos(3.14*vec.x) + rnorm(100, sd=1.5)
data.mu = mean(vec.y)
data.var = var(vec.y)

## apply stationary bootstrapping
nreps = 50
vec.mu = rep(0, nreps)
```

```

vec.var = rep(0,nreps)
for (i in 1:nreps){
  sample.id = boot.stationary(100)
  sample.y = vec.y[sample.id]
  vec.mu[i] = mean(sample.y)
  vec.var[i] = var(sample.y)
  print(paste("iteration ",i,"/",nreps," complete.", sep=""))
}

## visualize
opar <- par(no.readonly=TRUE)
par(mfrow=c(1,3), pty="s")
plot(vec.x, vec.y, type="l", main="1d signal") # 1d signal
hist(vec.mu, main="mean CI", xlab="mu") # mean
abline(v=data.mu, col="red", lwd=4)
hist(vec.var, main="variance CI", xlab="sigma") # variance
abline(v=data.var, col="blue", lwd=4)
par(opar)

```

---

cayleymenger

*Cayley-Menger Determinant*


---

## Description

Cayley-Menger determinant is a formula of a  $n$ -dimensional simplex with respect to the squares of all pairwise distances of its vertices.

## Usage

```
cayleymenger(data)
```

## Arguments

`data` an  $(n \times p)$  matrix of row-stacked observations.

## Value

a list containing

**det** determinant value.

**vol** volume attained from the determinant.

**Examples**

```
## USE 'IRIS' DATASET
data(iris)
X = as.matrix(iris[,1:4])

## COMPUTE CAYLEY-MENGER DETERMINANT
# since k=4 < n=149, it should be zero.
cayleymenger(X)
```

---

checkdist

*Check for Distance Matrix*

---

**Description**

This function checks whether the distance matrix  $D := d_{ij} = d(x_i, x_j)$  satisfies three axioms to make itself a semimetric, which are (1)  $d_{ii} = 0$ , (2)  $d_{ij} > 0$  for  $i \neq j$ , and (3)  $d_{ij} = d_{ji}$ .

**Usage**

```
checkdist(d)
```

**Arguments**

d "dist" object or  $(N \times N)$  matrix of pairwise distances.

**Value**

a logical; TRUE if it satisfies metric property, FALSE otherwise.

**See Also**

[checkmetric](#)

**Examples**

```
## Let's use L2 distance matrix of iris dataset
data(iris)
dx = as.matrix(stats::dist(iris[,1:4]))

# perturb d(i,j)
dy = dx
dy[1,2] <- dy[2,1] <- 10

# run the algorithm
checkdist(dx)
checkdist(dy)
```

---

`checkmetric`*Check for Metric Matrix*

---

### Description

This function checks whether the distance matrix  $D := d_{ij} = d(x_i, x_j)$  satisfies four axioms to make itself a semimetric, which are (1)  $d_{ii} = 0$ , (2)  $d_{ij} > 0$  for  $i \neq j$ , (3)  $d_{ij} = d_{ji}$ , and (4)  $d_{ij} \leq d_{ik} + d_{kj}$ .

### Usage

```
checkmetric(d)
```

### Arguments

`d` "dist" object or  $(N \times N)$  matrix of pairwise distances.

### Value

a logical; TRUE if it satisfies metric property, FALSE otherwise.

### See Also

[checkdist](#)

### Examples

```
## Let's use L2 distance matrix of iris dataset
data(iris)
dx = as.matrix(stats::dist(iris[,1:4]))

# perturb d(i,j)
dy = dx
dy[1,2] <- dy[2,1] <- 10

# run the algorithm
checkmetric(dx)
checkmetric(dy)
```



**Description**

Classical multidimensional scaling aims at finding low-dimensional structure by preserving pairwise distances of data.

**Usage**

```
cmds(data, ndim = 2)
```

**Arguments**

**data** an  $(n \times p)$  matrix whose rows are observations.  
**ndim** an integer-valued target dimension.

**Value**

a named list containing

**embed** an  $(n \times ndim)$  matrix whose rows are embedded observations.

**stress** discrepancy between embedded and original data as a measure of error.

**References**

Torgerson WS (1952). "Multidimensional Scaling: I. Theory and Method." *Psychometrika*, **17**(4), 401–419. ISSN 0033-3123, 1860-0980, doi: [10.1007/BF02288916](https://doi.org/10.1007/BF02288916).

**Examples**

```
## use simple example of iris dataset
data(iris)
idata = as.matrix(iris[,1:4])
icol = as.factor(iris[,5]) # class information

## run Classical MDS
iris.cmds = cmds(idata, ndim=2)

## visualize
opar <- par(no.readonly=TRUE)
plot(iris.cmds$embed, col=icol,
     main=paste0("STRESS=", round(iris.cmds$stress, 4)))
par(opar)
```

---

`dpmeans`*DP-means Algorithm for Clustering Euclidean Data*

---

### Description

DP-means is a nonparametric clustering method motivated by DP mixture model in that the number of clusters is determined by a parameter  $\lambda$ . The larger the  $\lambda$  value is, the smaller the number of clusters is attained. In addition to the original paper, we added an option to randomly permute an order of updating for each observation's membership as a common heuristic in the literature of cluster analysis.

### Usage

```
dpmeans(  
  data,  
  lambda = 1,  
  maxiter = 1234,  
  abstol = 1e-06,  
  permute.order = FALSE  
)
```

### Arguments

<code>data</code>	an $(n \times p)$ data matrix for each row being an observation.
<code>lambda</code>	a threshold to define a new cluster.
<code>maxiter</code>	maximum number of iterations.
<code>abstol</code>	stopping criterion
<code>permute.order</code>	a logical; TRUE if random order for permutation is used, FALSE otherwise.

### Value

a named list containing

**cluster** an  $(n \times ndim)$  matrix whose rows are embedded observations.

**centers** a list containing information for out-of-sample prediction.

### References

Kulis B, Jordan MI (2012). "Revisiting K-Means: New Algorithms via Bayesian Nonparametrics." In *Proceedings of the 29th International Conference on International Conference on Machine Learning*, ICML'12, 1131–1138. ISBN 978-1-4503-1285-1.

**Examples**

```

## define data matrix of two clusters
x1 = matrix(rnorm(50*3,mean= 2), ncol=3)
x2 = matrix(rnorm(50*3,mean=-2), ncol=3)
X = rbind(x1,x2)
lab = c(rep(1,50),rep(2,50))

## run dpmeans with several lambda values
solA <- dpmeans(X, lambda= 5)$cluster
solB <- dpmeans(X, lambda=10)$cluster
solC <- dpmeans(X, lambda=20)$cluster

## visualize the results
opar <- par(no.readonly=TRUE)
par(mfrow=c(1,4), pty="s")
plot(X,col=lab, pch=19, cex=.8, main="True", xlab="x", ylab="y")
plot(X,col=solA, pch=19, cex=.8, main="dpmeans lbd=5", xlab="x", ylab="y")
plot(X,col=solB, pch=19, cex=.8, main="dpmeans lbd=10", xlab="x", ylab="y")
plot(X,col=solC, pch=19, cex=.8, main="dpmeans lbd=20", xlab="x", ylab="y")
par(opar)

## let's find variations by permuting orders of update
## used setting : lambda=20, we will 8 runs
sol8 <- list()
for (i in 1:8){
  sol8[[i]] = dpmeans(X, lambda=20, permute.order=TRUE)$cluster
}

## let's visualize
vpar <- par(no.readonly=TRUE)
par(mfrow=c(2,4), pty="s")
for (i in 1:8){
  pm = paste("permute no.",i,sep="")
  plot(X,col=sol8[[i]], pch=19, cex=.8, main=pm, xlab="x", ylab="y")
}
par(vpar)

```

---

ecdfdist

*Distance Measures between Multiple Empirical Cumulative Distribution Functions*


---

**Description**

We measure distance between two empirical cumulative distribution functions (ECDF). For simplicity, we only take an input of [ecdf](#) objects from [stats](#) package.

**Usage**

```
ecdfdist(elist, method = c("KS", "Lp", "Wasserstein"), p = 2, as.dist = FALSE)
```

**Arguments**

<code>elist</code>	a length $N$ list of ecdf objects.
<code>method</code>	name of the distance/dissimilarity measure. Case insensitive.
<code>p</code>	exponent for Lp or Wasserstein distance.
<code>as.dist</code>	a logical; TRUE to return dist object, FALSE to return an $(N \times N)$ symmetric matrix of pairwise distances.

**Value**

either dist object of an  $(N \times N)$  symmetric matrix of pairwise distances by `as.dist` argument.

**See Also**

[ecdf](#)

**Examples**

```
## toy example : 10 of random and uniform distributions
mylist = list()
for (i in 1:10){
  mylist[[i]] = stats::ecdf(stats::rnorm(50, sd=2))
}
for (i in 11:20){
  mylist[[i]] = stats::ecdf(stats::runif(50, min=-5))
}

## compute Kolmogorov-Smirnov distance
dm = ecdfdist(mylist, method="KS")

## visualize
mks = "KS distances of 2 Types"
opar = par(no.readonly=TRUE)
par(pty="s")
image(dm[,nrow(dm):1], axes=FALSE, main=mks)
par(opar)
```

**Description**

We measure distance between two sets of empirical cumulative distribution functions (ECDF). For simplicity, we only take an input of [ecdf](#) objects from **stats** package.

**Usage**

```
ecdfdist2(elist1, elist2, method = c("KS", "Lp", "Wasserstein"), p = 2)
```

**Arguments**

<code>elist1</code>	a length $M$ list of <code>ecdf</code> objects.
<code>elist2</code>	a length $N$ list of <code>ecdf</code> objects.
<code>method</code>	name of the distance/dissimilarity measure. Case insensitive.
<code>p</code>	exponent for <code>Lp</code> or Wasserstein distance.

**Value**

an  $(M \times N)$  matrix of pairwise distances.

**See Also**

[ecdf](#) [ecdfdist](#)

**Examples**

```
## toy example
# first list : 10 of random and uniform distributions
mylist1 = list()
for (i in 1:10){ mylist1[[i]] = stats::ecdf(stats::rnorm(50, sd=2))}
for (i in 11:20){mylist1[[i]] = stats::ecdf(stats::runif(50, min=-5))}

# second list : 15 uniform and random distributions
mylist2 = list()
for (i in 1:15){ mylist2[[i]] = stats::ecdf(stats::runif(50, min=-5))}
for (i in 16:30){mylist2[[i]] = stats::ecdf(stats::rnorm(50, sd=2))}

## compute Kolmogorov-Smirnov distance
dm2ks = ecdfdist2(mylist1, mylist2, method="KS")
dm2lp = ecdfdist2(mylist1, mylist2, method="lp")
dm2wa = ecdfdist2(mylist1, mylist2, method="wasserstein")
nrs = nrow(dm2ks)

## visualize
```

```

opar = par(no.readonly=TRUE)
par(mfrow=c(1,3), pty="s")
image(dm2ks[,nrs:1], axes=FALSE, main="Kolmogorov-Smirnov")
image(dm2lp[,nrs:1], axes=FALSE, main="L2")
image(dm2wa[,nrs:1], axes=FALSE, main="Wasserstein")
par(opar)

```

---

epmeans

*EP-means Algorithm for Clustering Empirical Distributions*


---

### Description

EP-means is a variant of k-means algorithm adapted to cluster multiple empirical cumulative distribution functions under metric structure induced by Earth Mover's Distance.

### Usage

```
epmeans(elist, k = 2)
```

### Arguments

`elist` a length  $N$  list of either vector or ecdf objects.  
`k` the number of clusters.

### Value

a named list containing

**cluster** an integer vector indicating the cluster to which each ecdf is allocated.

**centers** a length  $k$  list of centroid ecdf objects.

### References

Henderson K, Gallagher B, Eliassi-Rad T (2015). "EP-MEANS: An Efficient Nonparametric Clustering of Empirical Probability Distributions." In *Proceedings of the 30th Annual ACM Symposium on Applied Computing - SAC '15*, 893–900. ISBN 978-1-4503-3196-8, doi: [10.1145/2695664.2695860](https://doi.org/10.1145/2695664.2695860).

### Examples

```

## two sets of 1d samples, 10 each and add some noise
# set 1 : mixture of two gaussians
# set 2 : single gamma distribution

# generate data
elist = list()

```

```

for (i in 1:10){
  elist[[i]] = stats::ecdf(c(rnorm(100, mean=-2), rnorm(50, mean=2)))
}
for (j in 11:20){
  elist[[j]] = stats::ecdf(rgamma(100,1) + rnorm(100, sd=sqrt(0.5)))
}

# run EP-means with k clusters
# change the value below to see different settings
myk = 2
epout = epmeans(elist, k=myk)

# visualize
opar = par(no.readonly=TRUE)
par(mfrow=c(1,myk))
for (k in 1:myk){
  idk = which(epout$cluster==k)
  for (i in 1:length(idk)){
    if (i<2){
      pm = paste("class ",k," (size=",length(idk),")",sep="")
      plot(elist[[idk[i]]], verticals=TRUE, lwd=0.25, do.points=FALSE, main=pm)
    } else {
      plot(elist[[idk[i]]], add=TRUE, verticals=TRUE, lwd=0.25, do.points=FALSE)
    }
    plot(epout$centers[[k]], add=TRUE, verticals=TRUE, lwd=2, col="red", do.points=FALSE)
  }
}
par(opar)

```

---

kmeanspp

*K-Means++ Clustering Algorithm*


---

## Description

*k*-means++ algorithm is known to be a smart, careful initialization technique. It is originally intended to return a set of *k* points as initial centers though it can still be used as a rough clustering algorithm by assigning points to the nearest points.

## Usage

```
kmeanspp(data, k = 2)
```

## Arguments

**data** an ( $n \times p$ ) matrix whose rows are observations.  
**k** the number of clusters.

**Value**

a length- $n$  vector of class labels.

**References**

Arthur D, Vassilvitskii S (2007). “K-Means++: The Advantages of Careful Seeding.” In *In Proceedings of the 18th Annual ACM-SIAM Symposium on Discrete Algorithms*.

**Examples**

```
## use simple example of iris dataset
data(iris)
mydata = as.matrix(iris[,1:4])
mycol = as.factor(iris[,5])

## find the low-dimensional embedding for visualization
my2d = cmds(mydata, ndim=2)$embed

## apply 'kmeanspp' with different numbers of k's.
k2 = kmeanspp(mydata, k=2)
k3 = kmeanspp(mydata, k=3)
k4 = kmeanspp(mydata, k=4)
k5 = kmeanspp(mydata, k=5)
k6 = kmeanspp(mydata, k=6)

## visualize
opar <- par(no.readonly=TRUE)
par(mfrow=c(2,3))
plot(my2d, col=k2, main="k=2", pch=19, cex=0.5)
plot(my2d, col=k3, main="k=3", pch=19, cex=0.5)
plot(my2d, col=k4, main="k=4", pch=19, cex=0.5)
plot(my2d, col=k5, main="k=5", pch=19, cex=0.5)
plot(my2d, col=k6, main="k=6", pch=19, cex=0.5)
plot(my2d, col=mycol, main="true cluster", pch=19, cex=0.5)
par(opar)
```

**Description**

We modify generalized Procrustes analysis for large-scale data by first setting a subset of anchor points and applying the attained transformation to the rest data. If `sub.id` is a vector  $1:\dim(x)[1]$ , it uses all observations as anchor points, reducing to the conventional generalized Procrustes analysis.

**Usage**

```
lgpa(x, sub.id = 1:(dim(x)[1]), scale = TRUE, reflect = FALSE)
```



**Arguments**

x	a $(k \times m \times n)$ 3d array, where $k$ is the number of points, $m$ the number of dimensions, and $n$ the number of samples.
sub.id	a vector of indices for defining anchor points.
scale	a logical; TRUE if scaling is applied, FALSE otherwise.
reflect	a logical; TRUE if reflection is required, FALSE otherwise.

**Value**

a  $(k \times m \times n)$  3d array of aligned samples.

**Author(s)**

Kisung You

**References**

Goodall C (1991). "Procrustes Methods in the Statistical Analysis of Shape." *Journal of the Royal Statistical Society. Series B (Methodological)*, **53**(2), 285–339. ISSN 00359246.

**Examples**

```
## Not run:
## This should be run if you have 'shapes' package installed.
library(shapes)
data(gorf.dat)

## apply anchor-based method and original procGPA
out.proc = shapes::procGPA(gorf.dat, scale=TRUE)$rotated # procGPA from shapes package
out.anc4 = lgpa(gorf.dat, sub.id=c(1,4,9,7), scale=TRUE) # use 4 points
out.anc7 = lgpa(gorf.dat, sub.id=1:7, scale=TRUE) # use all but 1 point as anchors

## visualize
opar = par(no.readonly=TRUE)
par(mfrow=c(3,4), pty="s")
plot(out.proc[, ,1], main="procGPA No.1", pch=18)
plot(out.proc[, ,2], main="procGPA No.2", pch=18)
plot(out.proc[, ,3], main="procGPA No.3", pch=18)
plot(out.proc[, ,4], main="procGPA No.4", pch=18)
plot(out.anc4[, ,1], main="4 Anchors No.1", pch=18, col="blue")
plot(out.anc4[, ,2], main="4 Anchors No.2", pch=18, col="blue")
plot(out.anc4[, ,3], main="4 Anchors No.3", pch=18, col="blue")
plot(out.anc4[, ,4], main="4 Anchors No.4", pch=18, col="blue")
plot(out.anc7[, ,1], main="7 Anchors No.1", pch=18, col="red")
plot(out.anc7[, ,2], main="7 Anchors No.2", pch=18, col="red")
plot(out.anc7[, ,3], main="7 Anchors No.3", pch=18, col="red")
plot(out.anc7[, ,4], main="7 Anchors No.4", pch=18, col="red")
par(opar)

## End(Not run)
```

lyapunov

*Solve Lyapunov Equation***Description**

The Lyapunov equation is of form

$$AX + XA^T = Q$$

where  $A$  and  $Q$  are square matrices of same size. Above form is also known as *continuous* form. This is a wrapper of `armadillo`'s `sylvester` function.

**Usage**

```
lyapunov(A, Q)
```

**Arguments**

$A$	a $(p \times p)$ matrix as above.
$Q$	a $(p \times p)$ matrix as above.

**Value**

a solution matrix  $X$  of size  $(p \times p)$ .

**References**

Sanderson C, Curtin R (2016). "Armadillo: A Template-Based C++ Library for Linear Algebra." *The Journal of Open Source Software*, **1**(2), 26.

Eddelbuettel D, Sanderson C (2014). "RcppArmadillo: Accelerating R with High-Performance C++ Linear Algebra." *Computational Statistics and Data Analysis*, **71**, 1054–1063.

**Examples**

```
## simulated example
# generate square matrices
A = matrix(rnorm(25),nrow=5)
X = matrix(rnorm(25),nrow=5)
Q = A%%X + X%%t(A)

# solve using 'lyapunov' function
solX = lyapunov(A,Q)
## Not run:
pm1 = "* Experiment with Lyapunov Solver"
pm2 = paste("* Absolute Error : ",norm(solX-X,"f"),sep="")
pm3 = paste("* Relative Error : ",norm(solX-X,"f")/norm(X,"f"),sep="")
cat(paste(pm1,"\n",pm2,"\n",pm3,sep=""))

## End(Not run)
```

---

matderiv	<i>Numerical Approximation to Gradient of a Function with Matrix Argument</i>
----------	---

---

### Description

For a given function  $f : \mathbf{R}^{n \times p} \rightarrow \mathbf{R}$ , we use finite difference scheme that approximates a gradient at a given point  $x$ . In Riemannian optimization, this can be used as a proxy for ambient gradient. Use with care since it may accumulate numerical error.

### Usage

```
matderiv(fn, x, h = 0.001)
```

### Arguments

fn	a function that takes a matrix of size $(n \times p)$ and returns a scalar value.
x	an $(n \times p)$ matrix where the gradient is to be computed.
h	step size for centered difference scheme.

### Value

an approximate numerical gradient matrix of size  $(n \times p)$ .

### References

Kincaid D, Cheney EW (2009). *Numerical Analysis: Mathematics of Scientific Computing*, number 2 in Pure and Applied Undergraduate Texts, 3. ed edition. American Mathematical Society, Providence, RI.

### Examples

```
## function f(X) = <a,Xb> for two vectors 'a' and 'b'
# derivative w.r.t X is ab'
# take an example of (5x5) symmetric positive definite matrix

# problem settings
a <- rnorm(5)
b <- rnorm(5)
ftn <- function(X){
  return(sum(as.vector(X%*%b)*a))
} # function to be taken derivative
myX <- matrix(rnorm(25),nrow=5) # point where derivative is evaluated
myX <- myX%*%t(myX)

# main computation
sol.true <- base::outer(a,b)
sol.num1 <- matderiv(ftn, myX, h=1e-1) # step size : 1e-1
```

```

sol.num2 <- matderiv(ftn, myX, h=1e-5) #           1e-3
sol.num3 <- matderiv(ftn, myX, h=1e-9) #           1e-5

## visualize/print the results
expar = par(no.readonly=TRUE)
par(mfrow=c(2,2),pty="s")
image(sol.true, main="true solution")
image(sol.num1, main="h=1e-1")
image(sol.num2, main="h=1e-5")
image(sol.num3, main="h=1e-9")
par(expar)

ntrue = norm(sol.true,"f")
cat('* Relative Errors in Frobenius Norm ')
cat(paste("* h=1e-1 : ",norm(sol.true-sol.num1,"f")/ntrue,sep=""))
cat(paste("* h=1e-5 : ",norm(sol.true-sol.num2,"f")/ntrue,sep=""))
cat(paste("* h=1e-9 : ",norm(sol.true-sol.num3,"f")/ntrue,sep=""))

```

---

mmd2test

*Kernel Two-sample Test with Maximum Mean Discrepancy*


---

## Description

Maximum Mean Discrepancy (MMD) as a measure of discrepancy between samples is employed as a test statistic for two-sample hypothesis test of equal distributions. Kernel matrix  $K$  is a symmetric square matrix that is positive semidefinite.

## Usage

```
mmd2test(K, label, method = c("b", "u"), mc.iter = 999)
```

## Arguments

K	kernel matrix or an object of kernelMatrix class from <b>kernlab</b> package.
label	label vector of class indices.
method	type of estimator to be used. "b" for biased and "u" for unbiased estimator of MMD.
mc.iter	the number of Monte Carlo resampling iterations.

## Value

a (list) object of S3 class htest containing:

**statistic** a test statistic.

**p.value**  $p$ -value under  $H_0$ .

**alternative** alternative hypothesis.  
**method** name of the test.  
**data.name** name(s) of provided kernel matrix.

## References

Gretton A, Borgwardt KM, Rasch MJ, Schölkopf B, Smola A (2012). "A Kernel Two-Sample Test." *J. Mach. Learn. Res.*, **13**, 723–773. ISSN 1532-4435.

## Examples

```
## small test for CRAN submission
dat1 <- matrix(rnorm(60, mean= 1), ncol=2) # group 1 : 30 obs of mean 1
dat2 <- matrix(rnorm(50, mean=-1), ncol=2) # group 2 : 25 obs of mean -1

dmat <- as.matrix(dist(rbind(dat1, dat2))) # Euclidean distance matrix
kmat <- exp(-(dmat^2)) # build a gaussian kernel matrix
lab <- c(rep(1,30), rep(2,25)) # corresponding label

mmd2test(kmat, lab) # run the code !

## Not run:
## WARNING: computationally heavy.
# Let's compute empirical Type 1 error at alpha=0.05
niter = 496
pvals1 = rep(0,niter)
pvals2 = rep(0,niter)
for (i in 1:niter){
  dat = matrix(rnorm(200),ncol=2)
  lab = c(rep(1,50), rep(2,50))
  lbd = 0.1
  kmat = exp(-lbd*(as.matrix(dist(dat))^2))
  pvals1[i] = mmd2test(kmat, lab, method="b")$p.value
  pvals2[i] = mmd2test(kmat, lab, method="u")$p.value
  print(paste("iteration ",i," complete..",sep=""))
}

# Visualize the above at multiple significance levels
alphas = seq(from=0.001, to=0.999, length.out=100)
errors1 = rep(0,100)
errors2 = rep(0,100)
for (i in 1:100){
  errors1[i] = sum(pvals1<=alphas[i])/niter
  errors2[i] = sum(pvals2<=alphas[i])/niter
}

opar <- par(no.readonly=TRUE)
par(mfrow=c(1,2), pty="s")
plot(alphas, errors1, "b", main="Biased Estimator Error",
      xlab="alpha", ylab="error", cex=0.5)
abline(a=0,b=1, lwd=1.5, col="red")
plot(alphas, errors2, "b", main="Unbiased Estimator Error",
```

```
      xlab="alpha", ylab="error", cex=0.5)
abline(a=0,b=1, lwd=1.5, col="blue")
par(opar)

## End(Not run)
```

---

nef

*Negative Eigenfraction*

---

### Description

Negative Eigenfraction (NEF) is a measure of distortion for the data whether they are lying in Euclidean manner or not. When the value is exactly 0, it means the data is Euclidean. On the other hand, when NEF is far away from 0, it means not Euclidean. The concept of NEF is closely related to the definiteness of a Gram matrix.

### Usage

```
nef(data)
```

### Arguments

data            an  $(n \times p)$  matrix whose rows are observations.

### Value

a nonnegative NEF value.

### References

Pełkalska E, Harol A, Duin RPW, Spillmann B, Bunke H (2006). “Non-Euclidean or Non-Metric Measures Can Be Informative.” In Yeung D, Kwok JT, Fred A, Roli F, de Ridder D (eds.), *Structural, Syntactic, and Statistical Pattern Recognition*, 871–880. ISBN 978-3-540-37241-7.

### Examples

```
## use simple example of iris dataset
data(iris)
mydat = as.matrix(iris[,1:4])

## calculate NEF
nef(mydat)
```

---

nem	<i>Negative Eigenvalue Magnitude</i>
-----	--------------------------------------

---

**Description**

Negative Eigenvalue Magnitude (NEM) is a measure of distortion for the data whether they are lying in Euclidean manner or not. When the value is exactly 0, it means the data is Euclidean. On the other hand, when NEM is far away from 0, it means not Euclidean. The concept of NEM is closely related to the definiteness of a Gram matrix.

**Usage**

```
nem(data)
```

**Arguments**

data            an  $(n \times p)$  matrix whose rows are observations.

**Value**

a nonnegative NEM value.

**References**

Pełkalska E, Harol A, Duin RPW, Spillmann B, Bunke H (2006). “Non-Euclidean or Non-Metric Measures Can Be Informative.” In Yeung D, Kwok JT, Fred A, Roli F, de Ridder D (eds.), *Structural, Syntactic, and Statistical Pattern Recognition*, 871–880. ISBN 978-3-540-37241-7.

**Examples**

```
## use simple example of iris dataset
data(iris)
mydat = as.matrix(iris[,1:4])

## calculate NEM
nem(mydat)
```

---

pdeterminant	<i>Calculate the Pseudo-Determinant of a Matrix</i>
--------------	---

---

**Description**

When a given square matrix  $A$  is rank deficient, determinant is zero. Still, we can compute the pseudo-determinant by multiplying all non-zero eigenvalues. Since thresholding to determine near-zero eigenvalues is subjective, we implemented the function as of original limit problem. When matrix is non-singular, it coincides with traditional determinant.

**Usage**

```
pdeterminant(A)
```

**Arguments**

A a square matrix whose pseudo-determinant be computed.

**Value**

a scalar value for computed pseudo-determinant.

**References**

Holbrook A (2018). “Differentiating the Pseudo Determinant.” *Linear Algebra and its Applications*, **548**, 293–304.

**Examples**

```
## show the convergence of pseudo-determinant
# settings
n = 10
A = cov(matrix(rnorm(5*n),ncol=n)) # (n x n) matrix
k = as.double(Matrix::rankMatrix(A)) # rank of A

# iterative computation
ntry = 11
del.vec = exp(-(1:ntry))
det.vec = rep(0,ntry)
for (i in 1:ntry){
  del = del.vec[i]
  det.vec[i] = det(A+del*diag(n))/(del^(n-k))
}

# visualize the results
opar <- par(no.readonly=TRUE)
plot(1:ntry, det.vec, main=paste("true rank is ",k," out of ",n,sep=""),"b", xlab="iterations")
abline(h=pdeterminant(A),col="red",lwd=1.2)
par(opar)
```

---

shortestpath

*Find Shortest Path using Floyd-Warshall Algorithm*


---

**Description**

This is a fast implementation of Floyd-Warshall algorithm to find the shortest path in a pairwise sense using RcppArmadillo. A logical input is also accepted. The given matrix should contain pairwise distance values  $d_{i,j}$  where 0 means there exists no path for node  $i$  and  $j$ .



**Usage**

```
shortestpath(dist)
```

**Arguments**

`dist` either an  $(n \times n)$  matrix or a `dist` class object.

**Value**

an  $(n \times n)$  matrix containing pairwise shortest path length.

**References**

Floyd RW (1962). "Algorithm 97: Shortest Path." *Communications of the ACM*, **5**(6), 345.

Warshall S (1962). "A Theorem on Boolean Matrices." *Journal of the ACM*, **9**(1), 11–12.

**Examples**

```
## simple example : a ring graph
# edges exist for pairs
A = array(0,c(10,10))
for (i in 1:9){
  A[i,i+1] = 1
  A[i+1,i] = 1
}
A[10,1] <- A[1,10] <- 1

# compute shortest-path and show the matrix
sdA <- shortestpath(A)

# visualize
opar <- par(no.readonly=TRUE)
par(pty="s")
image(sdA, main="shortest path length for a ring graph")
par(opar)
```

---

sylvester

*Solve Sylvester Equation*


---

**Description**

The Sylvester equation is of form

$$AX + XB = C$$

where  $X$  is the unknown and others are given. Though it's possible to have non-square  $A$  and  $B$  matrices, we currently support square matrices only. This is a wrapper of `armadillo`'s `sylvester` function.

**Usage**

```
sylvester(A, B, C)
```

**Arguments**

A	a $(p \times p)$ matrix as above.
B	a $(p \times p)$ matrix as above.
C	a $(p \times p)$ matrix as above.

**Value**

a solution matrix  $X$  of size  $(p \times p)$ .

**References**

Sanderson C, Curtin R (2016). "Armadillo: A Template-Based C++ Library for Linear Algebra." *The Journal of Open Source Software*, **1**(2), 26.

Eddelbuettel D, Sanderson C (2014). "RcppArmadillo: Accelerating R with High-Performance C++ Linear Algebra." *Computational Statistics and Data Analysis*, **71**, 1054–1063.

**Examples**

```
## simulated example
# generate square matrices
A = matrix(rnorm(25),nrow=5)
X = matrix(rnorm(25),nrow=5)
B = matrix(rnorm(25),nrow=5)
C = A**X + X**B

# solve using 'sylvester' function
solX = sylvester(A,B,C)
pm1 = "* Experiment with Sylvester Solver"
pm2 = paste("Absolute Error : ",norm(solX-X,"f"),sep="")
pm3 = paste("Relative Error : ",norm(solX-X,"f")/norm(X,"f"),sep="")
cat(paste(pm1,"\n",pm2,"\n",pm3,sep=""))
```

## Description

This function provides several algorithms to solve the following problem

$$\max \frac{\text{tr}(V^T AV)}{\text{tr}(V^T BV)} \text{ such that } V^T CV = I$$

where  $V$  is a projection matrix, i.e.,  $V^T V = I$ . Trace ratio optimization is pertained to various linear dimension reduction methods. It should be noted that when  $C = I$ , the above problem is often reformulated as a generalized eigenvalue problem since it's an easier proxy with faster computation.

## Usage

```
trio(
  A,
  B,
  C,
  dim = 2,
  method = c("2003Guo", "2007Wang", "2009Jia", "2012Ngo"),
  maxiter = 1000,
  eps = 1e-10
)
```

## Arguments

A	a $(p \times p)$ symmetric matrix in the numerator term.
B	a $(p \times p)$ symmetric matrix in the denominator term.
C	a $(p \times p)$ symmetric constraint matrix. If not provided, it is set as identical matrix automatically.
dim	an integer for target dimension. It can be considered as the number of loadings.
method	the name of algorithm to be used. Default is 2003Guo.
maxiter	maximum number of iterations to be performed.
eps	stopping criterion for iterative algorithms.

## Value

a named list containing

**V** a  $(p \times \text{dim})$  projection matrix.

**tr.val** an attained maximum scalar value.

## References

Guo Y, Li S, Yang J, Shu T, Wu L (2003). "A Generalized Foley–Sammon Transform Based on Generalized Fisher Discriminant Criterion and Its Application to Face Recognition." *Pattern Recognition Letters*, **24**(1-3), 147–158.

Wang H, Yan S, Xu D, Tang X, Huang T (2007). “Trace Ratio vs. Ratio Trace for Dimensionality Reduction.” In *2007 IEEE Conference on Computer Vision and Pattern Recognition*, 1–8.

Yangqing Jia, Feiping Nie, Changshui Zhang (2009). “Trace Ratio Problem Revisited.” *IEEE Transactions on Neural Networks*, **20**(4), 729–735.

Ngo TT, Bellalij M, Saad Y (2012). “The Trace Ratio Optimization Problem.” *SIAM Review*, **54**(3), 545–569.

## Examples

```
## simple test
# problem setting
p = 5
mydim = 2
A = matrix(rnorm(p^2),nrow=p); A=A%%t(A)
B = matrix(runif(p^2),nrow=p); B=B%%t(B)
C = diag(p)

# approximate solution via determinant ratio problem formulation
eigAB = eigen(solve(B,A))
V      = eigAB$vectors[,1:mydim]
eigval = sum(diag(t(V)%*%A%*%V))/sum(diag(t(V)%*%B%*%V))

# solve using 4 algorithms
m12 = trio(A,B,dim=mydim, method="2012Ngo")
m09 = trio(A,B,dim=mydim, method="2009Jia")
m07 = trio(A,B,dim=mydim, method="2007Wang")
m03 = trio(A,B,dim=mydim, method="2003Guo")

# print the results
line1 = '* Evaluation of the cost function'
line2 = paste("* approx. via determinant : ",eigval,sep="")
line3 = paste("* trio by 2012Ngo          : ",m12$str.val, sep="")
line4 = paste("* trio by 2009Jia           : ",m09$str.val, sep="")
line5 = paste("* trio by 2007Wang            : ",m07$str.val, sep="")
line6 = paste("* trio by 2003Guo             : ",m03$str.val, sep="")
cat(line1,"\n",line2,"\n",line3,"\n",line4,"\n",line5,"\n",line6)
```

---

tsne

*t-SNE Embedding*

---

## Description

This function is a simple wrapper of [Rtsne](#) function for t-Stochastic Neighbor Embedding for finding low-dimensional structure of the data embedded in the high-dimensional space.

## Usage

```
tsne(data, ndim = 2, ...)
```

**Arguments**

**data** an  $(n \times p)$  matrix whose rows are observations.  
**ndim** an integer-valued target dimension.  
... extra parameters to be used in `Rtsne` function.

**Value**

a named list containing

**embed** an  $(n \times ndim)$  matrix whose rows are embedded observations.

**stress** discrepancy between embedded and original data as a measure of error.

**Examples**

```
## use simple example of iris dataset
data(iris)
mydat = as.matrix(iris[,1:4])
mylab = as.factor(iris[,5])

## run t-SNE and MDS for comparison
iris.cmds = cmds(mydat, ndim=2)
iris.tsne = tsne(mydat, ndim=2)

## extract coordinates and class information
cx = iris.cmds$embed # embedded coordinates of CMDS
tx = iris.tsne$embed # t-SNE

## visualize
# main title
mc = paste("CMDS with STRESS=",round(iris.cmds$stress,4),sep="")
mt = paste("tSNE with STRESS=",round(iris.tsne$stress,4),sep="")

# draw a figure
opar <- par(no.readonly=TRUE)
par(mfrow=c(1,2))
plot(cx, col=mylab, pch=19, main=mc)
plot(tx, col=mylab, pch=19, main=mt)
par(opar)
```

**Description**

Geometric median, also known as L1-median, is a solution to the following problem

$$\operatorname{argmin} \sum_{i=1}^n \|x_i - y\|_2$$

for a given data  $x_1, x_2, \dots, x_n \in R^p$ .

**Usage**

```
weiszfeld(X, weights = NULL, maxiter = 496, abstol = 1e-06)
```

**Arguments**

<code>X</code>	an $(n \times p)$ matrix for $p$ -dimensional signal. If vector is given, it is assumed that $p = 1$ .
<code>weights</code>	NULL for equal weight $\text{rep}(1/n, n)$ ; otherwise, it has to be a vector of length $n$ .
<code>maxiter</code>	maximum number of iterations.
<code>abstol</code>	stopping criterion

**Examples**

```
## generate sin(x) data with noise for 100 replicates
set.seed(496)
t = seq(from=0, to=10, length.out=20)
X = array(0, c(100, 20))
for (i in 1:100){
  X[i,] = sin(t) + stats::rnorm(20, sd=0.5)
}

## compute L1-median and L2-mean
vecL2 = base::colMeans(X)
vecL1 = weiszfeld(X)

## visualize
opar <- par(no.readonly=TRUE)
par(mfrow=c(1,3), pty="s")
matplot(t(X[1:5,]), type="l", main="5 generated data", ylim=c(-2,2))
plot(t, vecL2, type="l", col="blue", main="L2-mean", ylim=c(-2,2))
plot(t, vecL1, type="l", col="red", main="L1-median", ylim=c(-2,2))
par(opar)
```

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