

# Package ‘qkerntool’

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**Title** Q-Kernel-Based and Conditionally Negative Definite Kernel-Based Machine Learning Tools

**Version** 1.18

**Description** Nonlinear machine learning tool for classification, clustering and dimensionality reduction. It integrates 12 q-kernel functions and 14 conditional negative definite kernel functions and includes the q-kernel and conditional negative definite kernel version of density-based spatial clustering of applications with noise, spectral clustering, generalized discriminant analysis, principal component analysis, multidimensional scaling, locally linear embedding, Sammon's mapping and t-Distributed stochastic neighbor embedding.

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## R topics documented:

as.cndkernmatrix . . . . .	2
as.qkernmatrix . . . . .	3
bases . . . . .	4

blkdiag . . . . .	6
cndkernel-class . . . . .	7
cndkernelmatrix . . . . .	8
cnds . . . . .	9
Eucdist . . . . .	11
mfeat_pix . . . . .	12
qkdbscan . . . . .	13
qkdbscan-class . . . . .	16
qkernel-class . . . . .	17
qkernelmatrix . . . . .	19
qkgda . . . . .	20
qkgda-class . . . . .	24
qkIsomap . . . . .	25
qkIsomap-class . . . . .	28
qkLLE . . . . .	30
qkLLE-class . . . . .	33
qkMDS . . . . .	34
qkMDS-class . . . . .	37
qkpca . . . . .	39
qkpca-class . . . . .	42
qkpre-class . . . . .	43
qkspecc . . . . .	44
qkspecc-class . . . . .	47
qkspeclust . . . . .	48
qsammon . . . . .	50
qsammon-class . . . . .	53
qtSNE . . . . .	54
qtSNE-class . . . . .	57

## Index 60

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as.cndkernelmatrix      *Assing cndkernelmatrix class to matrix objects*

---

### Description

as.cndkernelmatrix in package **qkerntool** can be used to create the cndkernelmatrix class to matrix objects representing a CND kernel matrix. These matrices can then be used with the cndkernelmatrix interfaces which most of the functions in **qkerntool** support.

### Usage

```
## S4 method for signature 'matrix'
as.cndkernelmatrix(x, center = FALSE)
```

**Arguments**

x                    matrix to be assigned the cndkernelmatrix class  
 center                center the cndkernel matrix in feature space (default: FALSE)

**Author(s)**

Yusen Zhang  
 <yusenzhang@126.com>

**See Also**

[cndkernelmatrix](#), [qkernelmatrix](#)

**Examples**

```
## Create the data
x <- rbind(matrix(rnorm(10),,2),matrix(rnorm(10,mean=3),,2))
y <- matrix(c(rep(1,5),rep(-1,5)))

### Use as.cndkernelmatrix to label the cov. matrix as a CND kernel matrix
### which is eq. to using a linear kernel

K <- as.cndkernelmatrix(crossprod(t(x)))

K
```

---

as.qkernelmatrix                    *Assing qkernelmatrix class to matrix objects*

---

**Description**

as.qkernelmatrix in package **qkerntool** can be used to create the qkernelmatrix class to matrix objects representing a q kernel matrix. These matrices can then be used with the qkernelmatrix interfaces which most of the functions in **qkerntool** support.

**Usage**

```
## S4 method for signature 'matrix'
as.qkernelmatrix(x, center = FALSE)
```

**Arguments**

x                    matrix to be assigned the qkernelmatrix class  
 center                center the kernel matrix in feature space (default: FALSE)

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**See Also**

[qkernelmatrix](#), [cndkernelmatrix](#)

**Examples**

```
## Create the data
x <- rbind(matrix(rnorm(10),,2),matrix(rnorm(10,mean=3),,2))
y <- matrix(c(rep(1,5),rep(-1,5)))

### Use as.qkernelmatrix to label the cov. matrix as a qkernel matrix
### which is eq. to using a linear kernel

K <- as.qkernelmatrix(crossprod(t(x)))

K
```

---

bases

*qKernel Functions*

---

**Description**

The kernel generating functions provided in `qkerntool`.

The Non Linear Kernel  $k(x, y) = \frac{1}{2(1-q)}(q^{-\alpha\|x\|^2} + q^{-\alpha\|y\|^2} - 2q^{-\alpha x'y})$ .

The Gaussian kernel  $k(x, y) = \frac{1}{1-q}(1 - q^{(\|x-y\|^2/\sigma)})$ .

The Laplacian Kernel  $k(x, y) = \frac{1}{1-q}(1 - q^{(\|x-y\|/\sigma)})$ .

The Rational Quadratic Kernel  $k(x, y) = \frac{1}{1-q}(1 - q^{\frac{\|x-y\|^2}{\|x-y\|^2+c}})$ .

The Multiquadric Kernel  $k(x, y) = \frac{1}{1-q}(q^c - q^{\sqrt{\|x-y\|^2+c}})$ .

The Inverse Multiquadric Kernel  $k(x, y) = \frac{1}{1-q}(q^{-\frac{1}{c}} - q^{-\frac{1}{\sqrt{\|x-y\|^2+c}}})$ .

The Wave Kernel  $k(x, y) = \frac{1}{1-q}(q^{-1} - q^{-\frac{\theta}{\|x-y\|} \sin \frac{\|x-y\|}{\theta}})$ .

The d Kernel  $k(x, y) = \frac{1}{1-q}[1 - q^{(\|x-y\|^d)}]$ .

The Log Kernel  $k(x, y) = \frac{1}{1-q}[1 - q^l n(\|x-y\|^d + 1)]$ .

The Cauchy Kernel  $k(x, y) = \frac{1}{1-q}(q^{-1} - q^{-\frac{1}{1+\|x-y\|^2/\sigma}})$ .

The Chi-Square Kernel  $k(x, y) = \frac{1}{1-q}(1 - q^{\sum 2(x-y)^2/(x+y)\gamma})$ .

The Generalized T-Student Kernel  $k(x, y) = \frac{1}{1-q}(q^{-1} - q^{-\frac{1}{1+\|x-y\|^d}})$ .

**Usage**

```

rbfbase(sigma=1,q=0.8)
nonlbase(alpha = 1,q = 0.8)
laplbase(sigma = 1, q = 0.8)
ratibase(c = 1, q = 0.8)
multibase(c = 1, q = 0.8)
invbase(c = 1, q = 0.8)
wavbase(theta = 1,q = 0.8)
powbase(d = 2, q = 0.8)
logbase(d = 2, q = 0.8)
caubase(sigma = 1, q = 0.8)
chibase(gamma = 1, q = 0.8)
studbase(d = 2, q = 0.8)

```

**Arguments**

q	for all the qkernel function.
sigma	for the Radial Basis qkernel function "rbfbase" , the Laplacian qkernel function "laplbase" and the Cauchy qkernel function "caubase".
alpha	for the Non Linear qkernel function "nonlbase".
c	for the Rational Quadratic qkernel function "ratibase" , the Multiquadric qkernel function "multibase" and the Inverse Multiquadric qkernel function "invbase".
theta	for the Wave qkernel function "wavbase".
d	for the d qkernel function "powbase" , the Log qkernel function "logbase" and the Generalized T-Student qkernel function "studbase".
gamma	for the Chi-Square qkernel function "chibase".

**Details**

The kernel generating functions are used to initialize a kernel function which calculates the kernel function value between two feature vectors in a Hilbert Space. These functions can be passed as a qkernel argument on almost all functions in **qkerntool**(e.g., qkgda, qkpca etc).

**Value**

Return an S4 object of class qkernel which extends the function class. The resulting function implements the given kernel calculating the kernel function value between two vectors.

qpar                    a list containing the kernel parameters (hyperparameters) used.

The kernel parameters can be accessed by the qpar function.

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**See Also**

[qkernelmatrix](#), [cndkernelmatrix](#)

**Examples**

```
qkfunc <- rfbbase(sigma=1,q=0.8)
qkfunc

qpar(qkfunc)

## create two vectors
x <- rnorm(10)
y <- rnorm(10)

## calculate dot product
qkfunc(x,y)
```

---

blkdiag

*Block diagonal concatenation of matrix*

---

**Description**

$Y = \text{BLKDIAG}(A,B,\dots)$  produces  $\text{diag}(A,B,\dots)$

**Usage**

```
blkdiag(x)
```

**Arguments**

x                    a list of matrix

**Value**

E - Block diagonal concatenation of matrix

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

---

cndkernel-class      *Class "cndkernel" "nonlkernel" "polykernel" "rbfkernel" "laplkernel"*

---

## Description

The built-in kernel classes in **qkerntool**

## Objects from the Class

Objects can be created by calls of the form `new("nonlkernel")`, `new{"polykernel"}`, `new{"rbfkernel"}`, `new{"laplkernel"}`, `new{"anokernel"}`, `new{"ratikernel"}`, `new{"multkernel"}`, `new{"invkernel"}`, `new{"wavkernel"}`, `new{"powkernel"}`, `new{"logkernel"}`, `new{"caukernel"}`, `new{"chikernel"}`, `new{"studkernel"}`

or by calling the `nonlcnd`, `polycnd`, `rbfcnd`, `laplcnd`, `anocnd`, `ratcnd`, `multcnd`, `invcnd`, `wavcnd`, `powcnd`, `logcnd`, `caucnd`, `chicnd`, `studcnd` functions etc..

## Slots

`.Data`: Object of class "function" containing the kernel function

`qpar`: Object of class "list" containing the kernel parameters

## Methods

**cndkernmatrix** signature(`kernel = "rbfkernel"`, `x = "matrix"`): computes the kernel matrix

## Author(s)

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## See Also

[qkernmatrix](#), [cndkernmatrix](#)

## Examples

```
cndkfunc <- rbfncnd(gamma = 1)
cndkfunc

qpar(cndkfunc)

## create two vectors
x <- rnorm(10)
y <- rnorm(10)

cndkfunc(x,y)
```

---

cndkernelmatrix                      *CND Kernel Matrix functions*

---

### Description

cndkernelmatrix calculates the kernel matrix  $K_{ij} = k(x_i, x_j)$  or  $K_{ij} = k(x_i, y_j)$ .

### Usage

```
## S4 method for signature 'cndkernel'
cndkernelmatrix(cndkernel, x, y = NULL)
```

### Arguments

cndkernel	the cndkernel function to be used to calculate the CND kernel matrix. This has to be a function of class cndkernel, i.e. which can be generated either one of the build in kernel generating functions (e.g., rbf <sub>cnd</sub> nonl <sub>cnd</sub> etc.) or a user defined function of class cndkernel taking two vector arguments and returning a scalar.
x	a data matrix to be used to calculate the kernel matrix.
y	second data matrix to calculate the kernel matrix.

### Details

Common functions used during kernel based computations.

The cndkernel parameter can be set to any function, of class cndkernel, which computes the kernel function value in feature space between two vector arguments. **qkerneltool** provides more than 10 CND kernel functions which can be initialized by using the following functions:

- nonl<sub>cnd</sub> Non Linear cndkernel function
- poly<sub>cnd</sub> Polynomial cndkernel function
- rbf<sub>cnd</sub> Gaussian cndkernel function
- lapl<sub>cnd</sub> Laplacian cndkernel function
- anoc<sub>cnd</sub> ANOVA cndkernel function
- ratic<sub>cnd</sub> Rational Quadratic cndkernel function
- mult<sub>cnd</sub> Multiquadric cndkernel function
- invc<sub>cnd</sub> Inverse Multiquadric cndkernel function
- wavc<sub>cnd</sub> Wave cndkernel function
- powc<sub>cnd</sub> d cndkernel function
- logc<sub>cnd</sub> Log cndkernel function
- cauc<sub>cnd</sub> Cauchy cndkernel function
- chic<sub>cnd</sub> Chi-Square cndkernel function
- stud<sub>cnd</sub> Generalized T-Student cndkernel function

(see example.)



**Value**

cndkernelmatrix returns a conditionally negative definite matrix with a zero diagonal element.

**Author(s)**

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**See Also**

[nonlbase](#), [rbfbase](#), [laplbase](#), [ratibase](#), [multbase](#), [invbase](#), [wavbase](#), [powbase](#), [logbase](#),  
[caubase](#), [chibase](#), [studbase](#)

**Examples**

```
## use the iris data
data(iris)
dt <- as.matrix(iris[, -5])

## initialize cndkernel function
lapl <- laplcnd(gamma = 1)
lapl

## calculate cndkernel matrix
cndkernelmatrix(lapl, dt)
```

**Description**

The kernel generating functions provided in qkerntool.

The Non Linear Kernel  $k(x, y) = [\exp(\alpha\|x\|^2) + \exp(\alpha\|y\|^2) - 2\exp(\alpha x'y)]/2$ .

The Polynomial kernel  $k(x, y) = [(\alpha\|x\|^2 + c)^d + (\alpha\|y\|^2 + c)^d - 2(\alpha x'y + c)^d]/2$ .

The Gaussian kernel  $k(x, y) = 1 - \exp(-\|x - y\|^2/\gamma)$ .

The Laplacian Kernel  $k(x, y) = 1 - \exp(-\|x - y\|/\gamma)$ .

The ANOVA Kernel  $k(x, y) = n - \sum \exp(-\sigma(x - y)^2)^d$ .

The Rational Quadratic Kernel  $k(x, y) = \|x - y\|^2/(\|x - y\|^2 + c)$ .

The Multiquadric Kernel  $k(x, y) = \sqrt{(\|x - y\|^2 + c^2) - c}$ .

The Inverse Multiquadric Kernel  $k(x, y) = 1/c - 1/\sqrt{\|x - y\|^2 + c^2}$ .

The Wave Kernel  $k(x, y) = 1 - \frac{\theta}{\|x - y\|} \sin \frac{\|x - y\|}{\theta}$ .

The d Kernel  $k(x, y) = \|x - y\|^d$ .

The Log Kernel  $k(x, y) = \log(\|x - y\|^d + 1)$ .

The Cauchy Kernel  $k(x, y) = 1 - 1/(1 + \|x - y\|^2/\gamma)$ .

The Chi-Square Kernel  $k(x, y) = \sum 2(x - y)^2/(x + y)$ .

The Generalized T-Student Kernel  $k(x, y) = 1 - 1/(1 + \|x - y\|^d)$ .

**Usage**

```

nonlcnd(alpha = 1)
polycnd(d = 2, alpha = 1, c = 1)
rbfcnd(gamma = 1)
laplcnd(gamma = 1)
anocnd(d = 2, sigma = 1)
raticnd(c = 1)
multcnd(c = 1)
invcnd(c = 1)
wavcnd(theta = 1)
powcnd(d = 2)
logcnd(d = 2)
caucnd(gamma = 1)
chicnd( )
studcnd(d = 2)

```

**Arguments**

alpha	for the Non Linear cndkernel function "nonlcnd" and the Polynomial cndkernel function "polycnd".
gamma	for the Radial Basis cndkernel function "rbfcnd" and the Laplacian cndkernel function "laplcnd" and the Cauchy cndkernel function "caucnd".
sigma	for the ANOVA cndkernel function "anocnd".
theta	for the Wave cndkernel function "wavcnd".
c	for the Rational Quadratic cndkernel function "raticnd", the Polynomial cndkernel function "polycnd", the Multiquadric cndkernel function "multcnd" and the Inverse Multiquadric cndkernel function "invcnd".
d	for the Polynomial cndkernel function "polycnd", the ANOVA cndkernel function "anocnd", the cndkernel function "powcnd", the Log cndkernel function "logcnd" and the Generalized T-Student cndkernel function "studcnd".

**Details**

The kernel generating functions are used to initialize a kernel function which calculates the kernel function value between two feature vectors in a Hilbert Space. These functions can be passed as a `kernel` argument on almost all functions in **qkerneltool**.

**Value**

Return an S4 object of class `cndkernel` which extends the function class. The resulting function implements the given kernel calculating the kernel function value between two vectors.

`qpar` a list containing the kernel parameters (hyperparameters) used.

The kernel parameters can be accessed by the `qpar` function.

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**See Also**

[cndkernmatrix](#), [qkernmatrix](#)

**Examples**

```
cndkfunc <- rbfcdnd(gamma = 1)
cndkfunc

qpar(cndkfunc)

## create two vectors
x <- rnorm(10)
y <- rnorm(10)

## calculate dot product
cndkfunc(x,y)
```

---

Euclidist

*Computes the Euclidean(square Euclidean) distance matrix*

---

**Description**

Euclidist Computes the Euclidean(square Euclidean) distance matrix.

**Arguments**

x (Nx $D$ ) matrix (N samples, D features)  
y (Mx $D$ ) matrix (M samples, D features)  
sEuclidean can be TRUE or FALSE, FALSE to Compute the Euclidean distance matrix.

**Value**

E - (MxN) Euclidean (square Euclidean) distances between vectors in x and y

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

### Examples

```
###
data(iris)
testset <- sample(1:150,20)
x <- as.matrix(iris[-testset,-5])
y <- as.matrix(iris[testset,-5])

##
res0 <- Eucdist(x)
res1 <- Eucdist(x, x, sEuclidean = FALSE)
res2 <- Eucdist(x, y = NULL, sEuclidean = FALSE)
res3 <- Eucdist(x, x, sEuclidean = TRUE)
res4 <- Eucdist(x, y = NULL)
res5 <- Eucdist(x, sEuclidean = FALSE)
```

---

mfeat\_pix

*mfeat\_pix dataset*

---

### Description

This dataset consists of features of handwritten numerals ('0'-'9') extracted from a collection of Dutch utility maps. 200 patterns per class (for a total of 2,000 patterns) have been digitized in binary images. This dataset is about 240 pixel averages in 2 x 3 windows

### Usage

```
data("mfeat_pix")
```

### Format

A data frame with 2000 observations on the following 240 variables.

### Source

<https://archive.ics.uci.edu/ml/datasets/Multiple+Features>

### Examples

```
data(mfeat_pix)
```

qkdbscan

*qKernel-DBSCAN density reachability and connectivity clustering***Description**

Similar to the Density-Based Spatial Clustering of Applications with Noise (or DBSCAN) algorithm, qKernel-DBSCAN is a density-based clustering algorithm that can be applied under both linear and non-linear situations.

**Usage**

```
## S4 method for signature 'matrix'
qkdbscan(x, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
eps = 0.25, MinPts = 5, hybrid = TRUE, seeds = TRUE, showplot = FALSE,
countmode = NULL, na.action = na.omit, ...)

## S4 method for signature 'cndkernmatrix'
qkdbscan(x, eps = 0.25, MinPts = 5, seeds = TRUE,
showplot = FALSE, countmode = NULL, ...)

## S4 method for signature 'qkernmatrix'
qkdbscan(x, eps = 0.25, MinPts = 5, seeds = TRUE,
showplot = FALSE, countmode = NULL, ...)

## S4 method for signature 'qkdbscan'
predict(object, data, newdata = NULL, predict.max = 1000, ...)
```

**Arguments**

x	the data matrix indexed by row, or a kernel matrix of <code>cndkernmatrix</code> or <code>qkernmatrix</code> .
kernel	<p>the kernel function used in training and predicting. This parameter can be set to any function, of class <code>kernel</code>, which computes a kernel function value between two vector arguments. <code>qkerntool</code> provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• <code>rbfbase</code> Radial Basis qkernel function "Gaussian"</li> <li>• <code>nonlbase</code> Non Linear qkernel function</li> <li>• <code>laplbase</code> Laplbase qkernel function</li> <li>• <code>ratibase</code> Rational Quadratic qkernel function</li> <li>• <code>multbase</code> Multiquadric qkernel function</li> <li>• <code>invbase</code> Inverse Multiquadric qkernel function</li> <li>• <code>wavbase</code> Wave qkernel function</li> <li>• <code>powbase</code> Power qkernel function</li> <li>• <code>logbase</code> Log qkernel function</li> <li>• <code>caubase</code> Cauchy qkernel function</li> </ul>

- chibase Chi-Square qkernel function
- studbase Generalized T-Student qkernel function
- nonlcnd Non Linear cndkernel function
- polycnd Polynomial cndkernel function
- rbfncnd Radial Basis cndkernel function "Gaussian"
- laplcnd Laplacian cndkernel function
- anocnd ANOVA cndkernel function
- raticnd Rational Quadratic cndkernel function
- multcnd Multiquadric cndkernel function
- invcnd Inverse Multiquadric cndkernel function
- wavcnd Wave cndkernel function
- powcnd Power cndkernel function
- logcnd Log cndkernel function
- caucnd Cauchy cndkernel function
- chicnd Chi-Square cndkernel function
- studcnd Generalized T-Student cndkernel function

The kernel parameter can also be set to a user defined function of class kernel by passing the function name as an argument.

qpar

the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :

- sigma, q for the Radial Basis qkernel function "rbfbase" , the Laplacian qkernel function "laplbase" and the Cauchy qkernel function "caubase".
- alpha, q for the Non Linear qkernel function "nonlbase".
- c, q for the Rational Quadratic qkernel function "ratibase" , the Multi-quadric qkernel function "multbase" and the Inverse Multiquadric qkernel function "invbase".
- theta, q for the Wave qkernel function "wavbase".
- d, q for the Power qkernel function "powbase" , the Log qkernel function "logbase" and the Generalized T-Student qkernel function "studbase".
- alpha for the Non Linear cndkernel function "nonlcnd".
- power, alpha, c for the Polynomial cndkernel function "polycnd".
- gamma for the Radial Basis cndkernel function "rbfncnd" and the Laplacian cndkernel function "laplcnd" and the Cauchy cndkernel function "caucnd".
- power, sigma for the ANOVA cndkernel function "anocnd".
- c for the Rational Quadratic cndkernel function "raticnd" , the Multiquadric cndkernel function "multcnd" and the Inverse Multiquadric cndkernel function "invcnd".
- theta for the Wave cndkernel function "wavcnd".
- power for the Power cndkernel function "powcnd" , the Log cndkernel function "logcnd" and the Generalized T-Student cndkernel function "studcnd".

Hyper-parameters for user defined kernels can be passed through the qpar parameter as well.

eps	reachability distance, see Ester et al. (1996). (default:0.25)
MinPts	reachability minimum number of points, see Ester et al.(1996).(default : 5)
hybrid	whether the algorithm expects raw data but calculates partial distance matrices, can be TRUE or FALSE
seeds	can be TRUE or FALSE, FALSE to not include the <code>issseed</code> -vector in the <code>dbscan</code> -object.
showplot	whether to show the plot or not, can be TRUE or FALSE
na.action	a function to specify the action to be taken if NAs are found. The default action is <code>na.omit</code> , which leads to rejection of cases with missing values on any required variable. An alternative is <code>na.fail</code> , which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)
countmode	NULL or vector of point numbers at which to report progress.
object	object of class <code>dbscan</code> .
data	matrix or <code>data.frame</code> .
newdata	matrix or <code>data.frame</code> with raw data to predict.
predict.max	max. batch size for predictions.
...	Further arguments transferred to plot methods.

### Details

The data can be passed to the `qkdbscan` function in a `matrix`, in addition `qkdbscan` also supports input in the form of a kernel matrix of class `qkernelmatrix` or class `kernelmatrix`.

### Value

`predict(qkdbscan-method)` gives out a vector of predicted clusters for the points in `newdata`.

`qkdbscan` gives out an S4 object which is a LIST with components

<code>clust</code>	integer vector coding cluster membership with noise observations (singletons) coded as 0
<code>eps</code>	parameter <code>eps</code>
<code>MinPts</code>	parameter <code>MinPts</code>
<code>kcall</code>	the function call
<code>kernel</code>	the kernel function used
<code>xmatrix</code>	the original data matrix

all the slots of the object can be accessed by accessor functions.

### Note

The `predict` function can be used to embed new data on the new space.

### Author(s)

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<yusenzhang@126.com>

**References**

Martin Ester, Hans-Peter Kriegel, Joerg Sander, Xiaowei Xu(1996).  
*A Density-Based Algorithm for Discovering Clusters in Large Spatial Databases with Noise*  
 Institute for Computer Science, University of Munich.  
*Proceedings of 2nd International Conference on Knowledge Discovery and Data Mining (KDD-96)*

**See Also**

[qkernelmatrix](#), [cndkernelmatrix](#)

**Examples**

```
# a simple example using the iris
data(iris)
test <- sample(1:150,20)
x<- as.matrix(iris[-test,-5])
ds <- qkdbscan (x,kernel="laplbase",qpar=list(sigma=3.5,q=0.8),eps=0.15,
MinPts=5,hybrid = FALSE)
plot(ds,x)
emb <- predict(ds, x, as.matrix(iris[test,-5]))
points(iris[test,], col= as.integer(1+emb))
```

---

qkdbscan-class	Class "qkdbscan"
----------------	------------------

---

**Description**

The qkernel-DBSCAN class.

**Objects of class "qkdbscan"**

Objects can be created by calls of the form `new("qkdbscan", ...)`. or by calling the `qkdbscan` function.

**Slots**

**clust:** Object of class "vector" containing the cluster membership of the samples

**eps:** Object of class "numeric" containing the reachability distance

**MinPts:** Object of class "numeric" containing the reachability minimum number of points

**isseed:** Object of class "logical" containing the logical vector indicating whether a point is a seed (not border, not noise)



**Methods**

**clust** signature(object = "qkdbscan"): returns the cluster membership  
**kcall** signature(object = "qkdbscan"): returns the performed call  
**cndkernf** signature(object = "qkdbscan"): returns the used kernel function  
**eps** signature(object = "qkdbscan"): returns the reachability distance  
**MinPts** signature(object = "qkdbscan"): returns the reachability minimum number of points  
**predict** signature(object = "qkdbscan"): embeds new data  
**xmatrix** signature(object = "qkdbscan"): returns the used data matrix

**Author(s)**

Yusen Zhang  
 <yusenzhang@126.com>

**See Also**

[qkernel-class](#), [cndkernel-class](#)

**Examples**

```
# a simple example using the iris data
x<- as.matrix(iris[,-5])
ds <- qkdbscan (x,kernel="laplbase",qpar=list(sigma=3.5,q=0.8),eps=0.15,
MinPts=5,hybrid = FALSE)
# print the results
clust(ds)
eps(ds)
MinPts(ds)
cndkernf(ds)
xmatrix(ds)
kcall(ds)
```

---

qkernel-class	<i>Class "qkernel" "rbfqkernel" "nonlqkernel" "laplqkernel" "ratiqkernel"</i>
---------------	---

---

**Description**

The built-in kernel classes in **qkerntool**

## Objects from the Class

Objects can be created by calls of the form `new("rbfqkernel")`, `new{"nonlqkernel"}`, `new{"laplqkernel"}`, `new{"ratiqkernel"}`, `new{"multqkernel"}`, `new{"invqkernel"}`, `new{"wavqkernel"}`, `new{"powqkernel"}`, `new{"logqkernel"}`, `new{"cauqkernel"}`, `new{"chiqkernel"}`, `new{"studqkernel"}`

or by calling the `rbfbase`, `nonlbase`, `laplbase`, `ratibase`, `multbase`, `invbase`, `wavbase`, `powbase`, `logbase`, `caubase`, `chibase`, `studbase` functions etc..

## Slots

`.Data`: Object of class "function" containing the kernel function

`qpar`: Object of class "list" containing the kernel parameters

## Methods

**qkernelmatrix** signature(kernel = "rbfqkernel", x = "matrix"): computes the qkernel matrix

## Author(s)

Yusen Zhang  
<yusenzhang@126.com>

## See Also

[qkernelmatrix](#), [cndkernelmatrix](#)

## Examples

```
qkfunc <- rbfbase(sigma=1,q=0.8)
qkfunc

qpar(qkfunc)

## create two vectors
x <- rnorm(10)
y <- rnorm(10)

## calculate dot product
qkfunc(x,y)
```

---

qkernelmatrix

*qKernel Matrix functions*


---

### Description

qkernelmatrix calculates the qkernel matrix  $K_{ij} = k(x_i, x_j)$  or  $K_{ij} = k(x_i, y_j)$ .

### Usage

```
## S4 method for signature 'qkernel'
qkernelmatrix(qkernel, x, y = NULL)
```

### Arguments

qkernel	the kernel function to be used to calculate the qkernel matrix. This has to be a function of class qkernel, i.e. which can be generated either one of the build in kernel generating functions (e.g., rbfbase etc.) or a user defined function of class qkernel taking two vector arguments and returning a scalar.
x	a data matrix to be used to calculate the kernel matrix
y	second data matrix to calculate the kernel matrix

### Details

Common functions used during kernel based computations.

The qkernel parameter can be set to any function, of class qkernel, which computes the kernel function value in feature space between two vector arguments. **qkerntool** provides more than 10 qkernel functions which can be initialized by using the following functions:

- nonlbase Non Linear qkernel function
- rbfbase Gaussian qkernel function
- laplbase Laplacian qkernel function
- ratibase Rational Quadratic qkernel function
- multbase Multiquadric qkernel function
- invbase Inverse Multiquadric qkernel function
- wavbase Wave qkernel function
- powbase d qkernel function
- logbase Log qkernel function
- caubase Cauchy qkernel function
- chibase Chi-Square qkernel function
- studbase Generalized T-Student qkernel function

(see example.)

**Value**

qkernmatrix returns a conditionally negative definite matrix with a zero diagonal element.

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**See Also**

[nonlcnd](#), [rbfcnd](#), [polycnd](#), [laplcnd](#), [anocnd](#), [raticnd](#), [multcnd](#), [invcnd](#), [wavcnd](#), [powcnd](#), [logcnd](#), [caucnd](#), [chicnd](#), [studcnd](#)

**Examples**

```
data(iris)
dt <- as.matrix(iris[, -5])

## initialize kernel function
rbf <- rfbase(sigma = 1.4, q=0.8)
rbf

## calculate qkernel matrix
qkernmatrix(rbf, dt)
```

---

qkgda

*qKernel Generalized Discriminant Analysis*

---

**Description**

The qkernel Generalized Discriminant Analysis is a method that deals with nonlinear discriminant analysis using kernel function operator.

**Usage**

```
## S4 method for signature 'matrix'
qkgda(x, label, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
      features = 0, th = 1e-4, na.action = na.omit, ...)

## S4 method for signature 'cndkernmatrix'
qkgda(x, label, features = 0, th = 1e-4, na.action = na.omit, ...)
## S4 method for signature 'qkernmatrix'
qkgda(x, label, features = 0, th = 1e-4, ...)
```

**Arguments**

x	the data matrix indexed by row, or a kernel matrix of <code>cnkernelmatrix</code> or <code>qkernelmatrix</code> .
label	The original labels of the samples.
kernel	<p>the kernel function used in training and predicting. This parameter can be set to any function, of class <code>kernel</code>, which computes a kernel function value between two vector arguments. <code>qkerntool</code> provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• <code>rbfbase</code> Radial Basis <code>qkernel</code> function "Gaussian"</li> <li>• <code>nonlbase</code> Non Linear <code>qkernel</code> function</li> <li>• <code>laplbase</code> Laplbase <code>qkernel</code> function</li> <li>• <code>ratibase</code> Rational Quadratic <code>qkernel</code> function</li> <li>• <code>multbase</code> Multiquadric <code>qkernel</code> function</li> <li>• <code>invbase</code> Inverse Multiquadric <code>qkernel</code> function</li> <li>• <code>wavbase</code> Wave <code>qkernel</code> function</li> <li>• <code>powbase</code> Power <code>qkernel</code> function</li> <li>• <code>logbase</code> Log <code>qkernel</code> function</li> <li>• <code>caubase</code> Cauchy <code>qkernel</code> function</li> <li>• <code>chibase</code> Chi-Square <code>qkernel</code> function</li> <li>• <code>studbase</code> Generalized T-Student <code>qkernel</code> function</li> <li>• <code>nonlcnd</code> Non Linear <code>cnkernel</code> function</li> <li>• <code>polycnd</code> Polynomial <code>cnkernel</code> function</li> <li>• <code>rbfcnd</code> Radial Basis <code>cnkernel</code> function "Gaussian"</li> <li>• <code>laplcnd</code> Laplacian <code>cnkernel</code> function</li> <li>• <code>anocnd</code> ANOVA <code>cnkernel</code> function</li> <li>• <code>ratıcnd</code> Rational Quadratic <code>cnkernel</code> function</li> <li>• <code>multcnd</code> Multiquadric <code>cnkernel</code> function</li> <li>• <code>invcnd</code> Inverse Multiquadric <code>cnkernel</code> function</li> <li>• <code>wavcnd</code> Wave <code>cnkernel</code> function</li> <li>• <code>powcnd</code> Power <code>cnkernel</code> function</li> <li>• <code>logcnd</code> Log <code>cnkernel</code> function</li> <li>• <code>caucnd</code> Cauchy <code>cnkernel</code> function</li> <li>• <code>chıcnd</code> Chi-Square <code>cnkernel</code> function</li> <li>• <code>studcnd</code> Generalized T-Student <code>cnkernel</code> function</li> </ul> <p>The kernel parameter can also be set to a user defined function of class <code>kernel</code> by passing the function name as an argument.</p>
qpar	<p>the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :</p> <ul style="list-style-type: none"> <li>• <code>sigma</code>, <code>q</code> for the Radial Basis <code>qkernel</code> function "<code>rbfbase</code>" , the Laplacian <code>qkernel</code> function "<code>laplbase</code>" and the Cauchy <code>qkernel</code> function "<code>caubase</code>".</li> <li>• <code>alpha</code>, <code>q</code> for the Non Linear <code>qkernel</code> function "<code>nonlbase</code>".</li> </ul>

- $c$ ,  $q$  for the Rational Quadratic  $q$ kernel function "ratibase" , the Multi-quadric  $q$ kernel function "multibase" and the Inverse Multiquadric  $q$ kernel function "invbase".
- $\theta$ ,  $q$  for the Wave  $q$ kernel function "wavbase".
- $d$ ,  $q$  for the Power  $q$ kernel function "powbase" , the Log  $q$ kernel function "logbase" and the Generalized T-Student  $q$ kernel function "studbase".
- $\alpha$  for the Non Linear  $cn$ dkernel function "nonlcn".
- $d$ ,  $\alpha$ ,  $c$  for the Polynomial  $cn$ dkernel function "polycn".
- $\gamma$  for the Radial Basis  $cn$ dkernel function "rbfcn" and the Laplacian  $cn$ dkernel function "laplcn" and the Cauchy  $cn$ dkernel function "caucn".
- $d$ ,  $\sigma$  for the ANOVA  $cn$ dkernel function "anocn".
- $c$  for the Rational Quadratic  $cn$ dkernel function "raticn" , the Multiquadric  $cn$ dkernel function "multcn" and the Inverse Multiquadric  $cn$ dkernel function "invcn".
- $\theta$  for the Wave  $cn$ dkernel function "wavcn".
- $d$  for the Power  $cn$ dkernel function "powcn" , the Log  $cn$ dkernel function "logcn" and the Generalized T-Student  $cn$ dkernel function "studcn".

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

<code>features</code>	Number of features (principal components) to return. (default: 0 , all)
<code>th</code>	the value of the eigenvalue under which principal components are ignored (only valid when <code>features = 0</code> ). (default : 0.0001)
<code>na.action</code>	A function to specify the action to be taken if NAs are found. The default action is <code>na.omit</code> , which leads to rejection of cases with missing values on any required variable. An alternative is <code>na.fail</code> , which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)
<code>...</code>	additional parameters

## Details

The  $q$ kernel Generalized Discriminant Analysis method provides a mapping of the input vectors into high dimensional feature space, generalizing the classical Linear Discriminant Analysis to non-linear discriminant analysis.

The data can be passed to the `qkgda` function in a `matrix`, in addition `qkgda` also supports input in the form of a kernel matrix of class `qkernelmatrix` or class `cnkernelmatrix`.

## Value

An S4 object containing the eigenvectors and their normalized projections, along with the corresponding eigenvalues and the original function.

<code>prj</code>	The normalized projections on eigenvectors)
<code>eVal</code>	The corresponding eigenvalues
<code>eVec</code>	The corresponding eigenvectors
<code>kcAll</code>	The formula of the function called

cndkernf        The kernel function used  
 xmatrix        The original data matrix

all the slots of the object can be accessed by accessor functions.

### Note

The predict function can be used to embed new data on the new space

### Author(s)

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### References

1. Baudat, G, and F. Anouar:  
*Generalized discriminant analysis using a kernel approach*  
 Neural Computation 12.10(2000),2385
2. Deng Cai, Xiaofei He, and Jiawei Han:  
*Speed Up Kernel Discriminant Analysis*  
 The VLDB Journal, January, 2011, vol.20, no.1, 21-33.

### See Also

[qkernmatrix](#), [cndkernmatrix](#)

### Examples

```

Iris <- data.frame(rbind(iris3[, ,1], iris3[, ,2], iris3[, ,3]), Sp = rep(c("1", "2", "3"), rep(50, 3)))
testset <- sample(1:150, 20)
train <- as.matrix(iris[-testset, -5])
test <- as.matrix(iris[testset, -5])
Sp = rep(c("1", "2", "3"), rep(50, 3))
labels <- as.numeric(Sp)
trainlabel <- labels[-testset]
testlabel <- labels[testset]

kgda1 <- qkgda(train, label=trainlabel, kernel = "ratibase", qpar = list(c=1, q=0.9), features = 2)

prj(kgda1)
eVal(kgda1)
eVec(kgda1)
kcall(kgda1)
# xmatrix(kgda1)

# print the principal component vectors
prj(kgda1)
# plot the data projection on the components
plot(kgda1@prj, col=as.integer(train), xlab="1st Principal Component", ylab="2nd Principal Component")

```

---

qkgda-class

Class "qkgda"

---

### Description

The qkernel Generalized Discriminant Analysis class

### Objects of class "qkgda"

Objects can be created by calls of the form `new("qkgda", ...)`. or by calling the `qkgda` function.

### Slots

**prj**: Object of class "matrix" containing the normalized projections on eigenvectors

**eVal**: Object of class "matrix" containing the corresponding eigenvalues

**eVec**: Object of class "matrix" containing the corresponding eigenvectors

**label**: Object of class "matrix" containing the categorical variables that the categorical data be assigned to one of the categories

### Methods

**prj** signature(object = "qkgda"): returns the normalized projections

**eVal** signature(object = "qkgda"): returns the eigenvalues

**eVec** signature(object = "qkgda"): returns the eigenvectors

**kcall** signature(object = "qkgda"): returns the performed call

**cndkernf** signature(object = "qkgda"): returns the used kernel function

**predict** signature(object = "qkgda"): embeds new data

**xmatrix** signature(object = "qkgda"): returns the used data matrix

### Author(s)

Yusen Zhang  
<yusenzhang@126.com>

### See Also

[qkernel-class](#), [cndkernel-class](#)



**Examples**

```

Iris <- data.frame(rbind(iris3[,1], iris3[,2], iris3[,3]), Sp = rep(c("1","2","3"), rep(50,3)))
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
test <- as.matrix(iris[testset,-5])
Sp = rep(c("1","2","3"), rep(50,3))
labels <-as.numeric(Sp)
trainlabel <- labels[-testset]
testlabel <- labels[testset]

kgda1 <- qkgda(train, label=trainlabel, kernel = "ratibase", qpar = list(c=1,q=0.9),features = 2)

prj(kgda1)
eVal(kgda1)
eVec(kgda1)
cndkernf(kgda1)
kcall(kgda1)

```

---

qkIsomap

*qKernel Isometric Feature Mapping*


---

**Description**

Computes the Isomap embedding as introduced in 2000 by Tenenbaum, de Silva and Langford.

**Usage**

```

## S4 method for signature 'matrix'
qkIsomap(x, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
dims = 2, k, mod = FALSE, plotResiduals = FALSE, verbose = TRUE, na.action = na.omit, ...)

## S4 method for signature 'cndkernmatrix'
qkIsomap(x, dims = 2, k, mod = FALSE, plotResiduals = FALSE,
verbose = TRUE, na.action = na.omit, ...)

## S4 method for signature 'qkernmatrix'
qkIsomap(x, dims = 2, k, mod = FALSE, plotResiduals = FALSE,
verbose = TRUE, na.action = na.omit, ...)

```

**Arguments**

x	N x D matrix (N samples, D features) or a kernel matrix of cndkernmatrix or qkernmatrix.
kernel	the kernel function used in training and predicting. This parameter can be set to any function, of class kernel, which computes a kernel function value between two vector arguments. qkerntool provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:

- rfbbase Radial Basis qkernel function "Gaussian"
- nonlbase Non Linear qkernel function
- laplbase Laplbase qkernel function
- ratibase Rational Quadratic qkernel function
- multbase Multiquadric qkernel function
- invbase Inverse Multiquadric qkernel function
- wavbase Wave qkernel function
- powbase Power qkernel function
- logbase Log qkernel function
- caubase Cauchy qkernel function
- chibase Chi-Square qkernel function
- studbase Generalized T-Student qkernel function
- nonlcnd Non Linear cndkernel function
- polycnd Polynomial cndkernel function
- rbfncnd Radial Basis cndkernel function "Gaussian"
- laplcnd Laplacian cndkernel function
- anocnd ANOVA cndkernel function
- raticnd Rational Quadratic cndkernel function
- multcnd Multiquadric cndkernel function
- invcnd Inverse Multiquadric cndkernel function
- wavgnd Wave cndkernel function
- powcnd Power cndkernel function
- logcnd Log cndkernel function
- caucnd Cauchy cndkernel function
- chicnd Chi-Square cndkernel function
- studcnd Generalized T-Student cndkernel function

The kernel parameter can also be set to a user defined function of class kernel by passing the function name as an argument.

qpar

the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :

- sigma, q for the Radial Basis qkernel function "rbfbase" , the Laplacian qkernel function "laplbase" and the Cauchy qkernel function "caubase".
- alpha, q for the Non Linear qkernel function "nonlbase".
- c, q for the Rational Quadratic qkernel function "ratibase" , the Multi-quadric qkernel function "multbase" and the Inverse Multiquadric qkernel function "invbase".
- theta, q for the Wave qkernel function "wavbase".
- d, q for the Power qkernel function "powbase" , the Log qkernel function "logbase" and the Generalized T-Student qkernel function "studbase".
- alpha for the Non Linear cndkernel function "nonlcnd".
- d, alpha, c for the Polynomial cndkernel function "polycnd".

- `gamma` for the Radial Basis cndkernel function "rbfcnd" and the Laplacian cndkernel function "laplcnd" and the Cauchy cndkernel function "caucnd".
- `d, sigma` for the ANOVA cndkernel function "anocnd".
- `c` for the Rational Quadratic cndkernel function "raticnd", the Multiquadric cndkernel function "multcnd" and the Inverse Multiquadric cndkernel function "invcnd".
- `theta` for the Wave cndkernel function "wavcnd".
- `d` for the Power cndkernel function "powcnd", the Log cndkernel function "logcnd" and the Generalized T-Student cndkernel function "studcnd".

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

<code>dims</code>	vector containing the target space dimension(s)
<code>k</code>	number of neighbours
<code>mod</code>	use modified Isomap algorithm
<code>plotResiduals</code>	show a plot with the residuals between the high and the low dimensional data
<code>verbose</code>	show a summary of the embedding procedure at the end
<code>na.action</code>	A function to specify the action to be taken if NAs are found. The default action is <code>na.omit</code> , which leads to rejection of cases with missing values on any required variable. An alternative is <code>na.fail</code> , which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)
<code>...</code>	additional parameters

## Details

The `qkIsomap` is a nonlinear dimension reduction technique, that preserves global properties of the data. That means, that geodesic distances between all samples are captured best in the low dimensional embedding.

This R version is based on the Matlab implementation by Tenenbaum and uses Floyd's Algorithm to compute the neighbourhood graph of shortest distances, when calculating the geodesic distances. A modified version of the original Isomap algorithm is included. It respects nearest and farthest neighbours.

To estimate the intrinsic dimension of the data, the function can plot the residuals between the high and the low dimensional data for a given range of dimensions.

## Value

`qkIsomap` gives out an S4 object which is a LIST with components

<code>prj</code>	a $N \times \text{dim}$ matrix ( $N$ samples, $\text{dim}$ features) with the reduced input data (list of several matrices if more than one dimension was specified).
<code>dims</code>	the dimension of the target space.
<code>Residuals</code>	the residual variances for all dimensions.
<code>eVal</code>	the corresponding eigenvalues.
<code>eVec</code>	the corresponding eigenvectors.
<code>cndkernf</code>	the kernel function used.

`kcall`            The formula of the function called  
all the slots of the object can be accessed by accessor functions.

### Author(s)

Yusen Zhang  
<yusenzhang@126.com>

### References

Tenenbaum, J. B. and de Silva, V. and Langford, J. C., "A global geometric framework for nonlinear dimensionality reduction.", 2000; Matlab code is available at <http://waldron.stanford.edu/~isomap/>

### Examples

```
# another example using the iris
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
labeltrain<- as.integer(iris[-testset,5])
test <- as.matrix(iris[testset,-5])
# ratibase(c=1,q=0.8)
d_low = qkIsomap(train, kernel = "ratibase", qpar = list(c=1,q=0.8),
                 dims=2, k=5, plotResiduals = TRUE)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")

prj(d_low)
dims(d_low)
Residuals(d_low)
eVal(d_low)
eVec(d_low)
kcall(d_low)
cndkernf(d_low)
```

---

qkIsomap-class

*qKernel Isomap embedding*

---

### Description

The qKernel Isometric Feature Mapping class

### Objects of class "qkIsomap"

Objects can be created by calls of the form `new("qkIsomap", ...)`. or by calling the `qkIsomap` function.

**Slots**

**prj**: Object of class "matrix" containing the Nxdim matrix (N samples, dim features) with the reduced input data (list of several matrices if more than one dimension specified)

**dims**: Object of class "numeric" containing the dimension of the target space (default 2)

**connum**: Object of class "numeric" containing the number of connected components in graph

**Residuals**: Object of class "vector" containing the residual variances for all dimensions

**eVal**: Object of class "vector" containing the corresponding eigenvalues

**eVec**: Object of class "vector" containing the corresponding eigenvectors

**Methods**

**prj** signature(object = "qkIsomap"): returns the Nxdim matrix (N samples, dim features)

**dims** signature(object = "qkIsomap"): returns the dimension

**Residuals** signature(object = "qkIsomap"): returns the residual variances

**eVal** signature(object = "qkIsomap"): returns the eigenvalues

**eVec** signature(object = "qkIsomap"): returns the eigenvectors

**xmatrix** signature(object = "qkIsomap"): returns the used data matrix

**kcall** signature(object = "qkIsomap"): returns the performed call

**cndkernf** signature(object = "qkIsomapa"): returns the used kernel function

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**See Also**

[qkernel-class](#), [cndkernel-class](#), [qkIsomap](#)

**Examples**

```
# another example using the iris data
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
labeltrain<- as.integer(iris[-testset,5])
test <- as.matrix(iris[testset,-5])
# ratibase(c=1,q=0.8)
d_low = qkIsomap(train, kernel = "ratibase", qpar = list(c=1,q=0.8),
                dims=2, k=5, plotResiduals = TRUE)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")

prj(d_low)
dims(d_low)
Residuals(d_low)
```

```
eVal(d_low)
eVec(d_low)
kcall(d_low)
cndkernf(d_low)
```

---

qkLLE

*qKernel Locally Linear Embedding*


---

## Description

Computes the qkernel Locally Linear Embedding

## Usage

```
## S4 method for signature 'matrix'
qkLLE(x, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
      dims = 2, k, na.action = na.omit, ...)
## S4 method for signature 'cndkernmatrix'
qkLLE(x, dims = 2, k, na.action = na.omit, ...)
## S4 method for signature 'qkernmatrix'
qkLLE(x, dims = 2, k, na.action = na.omit,...)
```

## Arguments

x	N x D matrix (N samples, D features) or a kernel matrix of cndkernmatrix or qkernmatrix.
kernel	<p>the kernel function used in training and predicting. This parameter can be set to any function, of class kernel, which computes a kernel function value between two vector arguments. qkerntool provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• rbfbase Radial Basis qkernel function "Gaussian"</li> <li>• nonlbase Non Linear qkernel function</li> <li>• laplbase Laplbase qkernel function</li> <li>• ratibase Rational Quadratic qkernel function</li> <li>• multbase Multiquadric qkernel function</li> <li>• invbase Inverse Multiquadric qkernel function</li> <li>• wavbase Wave qkernel function</li> <li>• powbase Power qkernel function</li> <li>• logbase Log qkernel function</li> <li>• caubase Cauchy qkernel function</li> <li>• chibase Chi-Square qkernel function</li> <li>• studbase Generalized T-Student qkernel function</li> <li>• nonlcnd Non Linear cndkernel function</li> <li>• polycnd Polynomial cndkernel function</li> <li>• rbfcd Radial Basis cndkernel function "Gaussian"</li> </ul>

- `laplcnd` Laplacian cndkernel function
- `anocnd` ANOVA cndkernel function
- `raticnd` Rational Quadratic cndkernel function
- `multcnd` Multiquadric cndkernel function
- `invcnd` Inverse Multiquadric cndkernel function
- `wavcnd` Wave cndkernel function
- `powcnd` Power cndkernel function
- `logcnd` Log cndkernel function
- `caucnd` Cauchy cndkernel function
- `chicnd` Chi-Square cndkernel function
- `studcnd` Generalized T-Student cndkernel function

The kernel parameter can also be set to a user defined function of class `kernel` by passing the function name as an argument.

`qpar` the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :

- `sigma`, `q` for the Radial Basis qkernel function "`rbfbase`", the Laplacian qkernel function "`laplbase`" and the Cauchy qkernel function "`caubase`".
- `alpha`, `q` for the Non Linear qkernel function "`nonlbase`".
- `c`, `q` for the Rational Quadratic qkernel function "`ratibase`", the Multiquadric qkernel function "`multbase`" and the Inverse Multiquadric qkernel function "`invbase`".
- `theta`, `q` for the Wave qkernel function "`wavbase`".
- `d`, `q` for the Power qkernel function "`powbase`", the Log qkernel function "`logbase`" and the Generalized T-Student qkernel function "`studbase`".
- `alpha` for the Non Linear cndkernel function "`nonlcnd`".
- `power`, `alpha`, `c` for the Polynomial cndkernel function "`polycnd`".
- `gamma` for the Radial Basis cndkernel function "`rbfcnd`" and the Laplacian cndkernel function "`laplcnd`" and the Cauchy cndkernel function "`caucnd`".
- `power`, `sigma` for the ANOVA cndkernel function "`anocnd`".
- `c` for the Rational Quadratic cndkernel function "`raticnd`", the Multiquadric cndkernel function "`multcnd`" and the Inverse Multiquadric cndkernel function "`invcnd`".
- `theta` for the Wave cndkernel function "`wavcnd`".
- `power` for the Power cndkernel function "`powcnd`", the Log cndkernel function "`logcnd`" and the Generalized T-Student cndkernel function "`studcnd`".

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

`dims` dimension of the target space

`k` the number of nearest neighbours.

`na.action` A function to specify the action to be taken if NAs are found. The default action is `na.omit`, which leads to rejection of cases with missing values on any required variable. An alternative is `na.fail`, which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)

`...` additional parameters

## Details

The qkernel Locally Linear Embedding (qkLLE) preserves local properties of the data by representing each sample in the data by a linear combination of its  $k$  nearest neighbours with each neighbour weighted independently. qkLLE finally chooses the low-dimensional representation that best preserves the weights in the target space. It is an extension of Locally Linear Embedding (LLE) with qkernel method.

## Value

It returns an S4 object containing the principal component vectors along with the corresponding eigenvalues.

prj	a matrix with the reduced input data
dims	dimension of the target space
eVal	The corresponding eigenvalues
eVec	The corresponding eigenvectors
cnDKernf	the kernel function used

all the slots of the object can be accessed by accessor functions.

## Author(s)

Yusen Zhang  
<yusenzhang@126.com>

## References

Roweis, Sam T. and Saul, Lawrence K., "Nonlinear Dimensionality Reduction by Locally Linear Embedding", 2000;

## Examples

```
## S4 method for signature 'matrix'
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
labeltrain<- as.integer(iris[-testset,5])
test <- as.matrix(iris[testset,-5])
plot(train ,col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")
# ratibase(c=1,q=0.8)
d_low <- qkLLE(train, kernel = "ratibase", qpar = list(c=1,q=0.8), dims=2, k=5)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")

## S4 method for signature 'qkernmatrix'
# ratibase(c=0.1,q=0.8)
qkfunc <- ratibase(c=0.1,q=0.8)
ktrain1 <- qkernmatrix(qkfunc,train)
d_low <- qkLLE(ktrain1, dims = 2, k=5)
```



```
#plot the data projection on the components
plot(prj(d_low),col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")
```

---

qkLLE-class

Class "qkLLE"

---

## Description

The qKernel Locally Linear Embedding class

## Objects of class "qkLLE"

Objects can be created by calls of the form `new("qkLLE", ...)`. or by calling the `qkLLE` function.

## Slots

**prj**: Object of class "matrix" containing the reduced input data  
**dims**: Object of class "numeric" containing the dimension of the target space (default 2)  
**eVal**: Object of class "vector" containing the corresponding eigenvalues  
**eVec**: Object of class "matrix" containing the corresponding eigenvectors

## Methods

**prj** signature(object = "qkLLE"): returns the reduced input data  
**dims** signature(object = "qkLLE"): returns the dimension  
**eVal** signature(object = "qkLLE"): returns the eigenvalues  
**eVec** signature(object = "qkLLE"): returns the eigenvectors  
**xmatrix** signature(object = "qkLLE"): returns the used data matrix  
**kcall** signature(object = "qkLLE"): returns the performed call  
**cndkernf** signature(object = "qkLLE"): returns the used kernel function

## Author(s)

Yusen Zhang  
<yusenzhang@126.com>

## See Also

[qkernel-class](#), [cndkernel-class](#)

**Examples**

```

## S4 method for signature 'matrix'
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
labeltrain<- as.integer(iris[-testset,5])
test <- as.matrix(iris[testset,-5])
plot(train ,col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")
# ratibase(c=1,q=0.8)
d_low <- qkLLE(train, kernel = "ratibase", qpar = list(c=1,q=0.8), dims=2, k=5)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain,xlab="1st Principal Component",ylab="2nd Principal Component")

## S4 method for signature 'qkernmatrix'
# ratibase(c=0.1,q=0.8)
qkfunc <- ratibase(c=0.1,q=0.8)
ktrain1 <- qkernmatrix(qkfunc,train)
d_low <- qkLLE(ktrain1, dims = 2, k=5)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain,xlab="1st Principal Component",ylab="2nd Principal Component")

```

---

qkMDS

*qKernel Metric Multi-Dimensional Scaling*


---

**Description**

The qkernel Metric Multi-Dimensional Scaling is a nonlinear form of Metric Multi-Dimensional Scaling

**Usage**

```

## S4 method for signature 'matrix'
qkMDS(x, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
dims = 2, plotResiduals = FALSE, verbose = TRUE, na.action = na.omit, ...)

## S4 method for signature 'cndkernmatrix'
qkMDS(x, dims = 2,plotResiduals = FALSE,
verbose = TRUE, na.action = na.omit, ...)

## S4 method for signature 'qkernmatrix'
qkMDS(x, dims = 2,plotResiduals = FALSE,
verbose = TRUE, na.action = na.omit, ...)

```

**Arguments**

x                    N x D matrix (N samples, D features) or a kernel matrix of cndkernmatrix or qkernmatrix.

kernel	<p>the kernel function used in training and predicting. This parameter can be set to any function, of class kernel, which computes a kernel function value between two vector arguments. qkernntool provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• rbfbase Radial Basis qkernel function "Gaussian"</li> <li>• nonlbase Non Linear qkernel function</li> <li>• laplbase Laplbase qkernel function</li> <li>• ratibase Rational Quadratic qkernel function</li> <li>• multbase Multiquadric qkernel function</li> <li>• invbase Inverse Multiquadric qkernel function</li> <li>• wavbase Wave qkernel function</li> <li>• powbase Power qkernel function</li> <li>• logbase Log qkernel function</li> <li>• caubase Cauchy qkernel function</li> <li>• chibase Chi-Square qkernel function</li> <li>• studbase Generalized T-Student qkernel function</li> <li>• nonlcnd Non Linear cndkernel function</li> <li>• polycnd Polynomial cndkernel function</li> <li>• rbfncnd Radial Basis cndkernel function "Gaussian"</li> <li>• laplcnd Laplacian cndkernel function</li> <li>• anocnd ANOVA cndkernel function</li> <li>• raticnd Rational Quadratic cndkernel function</li> <li>• multcnd Multiquadric cndkernel function</li> <li>• invcnd Inverse Multiquadric cndkernel function</li> <li>• wavgnd Wave cndkernel function</li> <li>• powcnd Power cndkernel function</li> <li>• logcnd Log cndkernel function</li> <li>• caucnd Cauchy cndkernel function</li> <li>• chicnd Chi-Square cndkernel function</li> <li>• studcnd Generalized T-Student cndkernel function</li> </ul> <p>The kernel parameter can also be set to a user defined function of class kernel by passing the function name as an argument.</p>
qpar	<p>the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :</p> <ul style="list-style-type: none"> <li>• sigma, q for the Radial Basis qkernel function "rbfbase" , the Laplacian qkernel function "laplbase" and the Cauchy qkernel function "caubase".</li> <li>• alpha, q for the Non Linear qkernel function "nonlbase".</li> <li>• c, q for the Rational Quadratic qkernel function "ratibase" , the Multi-quadric qkernel function "multbase" and the Inverse Multiquadric qkernel function "invbase".</li> <li>• theta, q for the Wave qkernel function "wavbase".</li> </ul>

- `d`, `q` for the Power qkernel function "powbase", the Log qkernel function "logbase" and the Generalized T-Student qkernel function "studbase".
- `alpha` for the Non Linear cndkernel function "nonlcnd".
- `d`, `alpha`, `c` for the Polynomial cndkernel function "polycnd".
- `gamma` for the Radial Basis cndkernel function "rbfcnd" and the Laplacian cndkernel function "laplcnd" and the Cauchy cndkernel function "caucnd".
- `d`, `sigma` for the ANOVA cndkernel function "anocnd".
- `c` for the Rational Quadratic cndkernel function "raticnd", the Multiquadric cndkernel function "multcnd" and the Inverse Multiquadric cndkernel function "invcnd".
- `theta` for the Wave cndkernel function "wavcnd".
- `d` for the Power cndkernel function "powcnd", the Log cndkernel function "logcnd" and the Generalized T-Student cndkernel function "studcnd".

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

<code>dims</code>	vector containing the target space dimension(s)
<code>plotResiduals</code>	show a plot with the residuals between the high and the low dimensional data
<code>verbose</code>	show a summary of the embedding procedure at the end
<code>na.action</code>	A function to specify the action to be taken if NAs are found. The default action is <code>na.omit</code> , which leads to rejection of cases with missing values on any required variable. An alternative is <code>na.fail</code> , which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)
<code>...</code>	additional parameters

## Details

There are several versions of non-metric multidimensional scaling in R, but **qkerntool** offers the following unique combination of using qKernel methods

## Value

qkMDS gives out an S4 object which is a LIST with components

<code>prj</code>	a $N \times \text{dim}$ matrix ( $N$ samples, $\text{dim}$ features) with the reduced input data (list of several matrices if more than one dimension was specified).
<code>dims</code>	the dimension of the target space.
<code>Residuals</code>	the residual variances for all dimensions.
<code>eVal</code>	the corresponding eigenvalues.
<code>eVec</code>	the corresponding eigenvectors.
<code>cndkernf</code>	the kernel function used.
<code>kcall</code>	The formula of the function called

all the slots of the object can be accessed by accessor functions.

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**References**

Kruskal, J.B. 1964a. Multidimensional scaling by optimizing goodness-of-fit to a nonmetric hypothesis. *Psychometrika* 29, 1–28.

**Examples**

```
# another example using the iris
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
labeltrain<- as.integer(iris[-testset,5])
test <- as.matrix(iris[testset,-5])
# ratibase(c=1,q=0.8)
d_low = qkMDS(train, kernel = "ratibase", qpar = list(c=1,q=0.9),dims = 2,
              plotResiduals = TRUE)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")

prj(d_low)
dims(d_low)
Residuals(d_low)
eVal(d_low)
eVec(d_low)
kcall(d_low)
cndkernf(d_low)
```

---

qkMDS-class

*qKernel Metric Multi-Dimensional Scaling*


---

**Description**

The qkernel Metric Multi-Dimensional Scaling class

**Objects of class "qkMDS"**

Objects can be created by calls of the form `new("qkMDS", ...)`. or by calling the `qkMDS` function.

**Slots**

`prj`: Object of class "matrix" containing the Nxdim matrix (N samples, dim features) with the reduced input data (list of several matrices if more than one dimension specified)

`dims`: Object of class "numeric" containing the dimension of the target space (default 2)

`connum`: Object of class "numeric" containing the number of connected components in graph

**Residuals:** Object of class "vector" containing the residual variances for all dimensions

**eVal:** Object of class "vector" containing the corresponding eigenvalues

**eVec:** Object of class "vector" containing the corresponding eigenvectors

## Methods

**prj** signature(object = "qkMDS"): returns the Nxdim matrix (N samples, dim features)

**dims** signature(object = "qkMDS"): returns the dimension

**Residuals** signature(object = "qkMDS"): returns the residual variances

**eVal** signature(object = "qkMDS"): returns the eigenvalues

**eVec** signature(object = "qkMDS"): returns the eigenvectors

**xmatrix** signature(object = "qkMDS"): returns the used data matrix

**kcall** signature(object = "qkMDS"): returns the performed call

**cndkernf** signature(object = "qkMDS"): returns the used kernel function

## Author(s)

Yusen Zhang

<yusenzhang@126.com>

## See Also

[qkernel-class](#), [cndkernel-class](#), [qkMDS](#)

## Examples

```
# another example using the iris
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[-testset,-5])
labeltrain<- as.integer(iris[-testset,5])
test <- as.matrix(iris[testset,-5])
# ratibase(c=1,q=0.8)
d_low = qkMDS(train, kernel = "ratibase", qpar = list(c=1,q=0.8),
              dims=2, plotResiduals = TRUE)
#plot the data projection on the components
plot(prj(d_low),col=labeltrain, xlab="1st Principal Component",ylab="2nd Principal Component")

prj(d_low)
dims(d_low)
Residuals(d_low)
eVal(d_low)
eVec(d_low)
kcall(d_low)
cndkernf(d_low)
```

**Description**

The qkernel Principal Components Analysis is a nonlinear form of principal component analysis.

**Usage**

```
## S4 method for signature 'formula'
qkpca(x, data = NULL, na.action, ...)
## S4 method for signature 'matrix'
qkpca(x, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
      features = 0, th = 1e-4, na.action = na.omit, ...)
## S4 method for signature 'cndkernmatrix'
qkpca(x, features = 0, th = 1e-4, ...)
## S4 method for signature 'qkernmatrix'
qkpca(x, features = 0, th = 1e-4, ...)
```

**Arguments**

- |        |   |
|--------|---|
| x      | the data matrix indexed by row, a formula describing the model or a kernel matrix of <code>cndkernmatrix</code> or <code>qkernmatrix</code> .   |
| data   | an optional data frame containing the variables in the model (when using a formula).  |
| kernel | <p>the kernel function used in training and predicting. This parameter can be set to any function, of class <code>kernel</code>, which computes a kernel function value between two vector arguments. <code>qkerntool</code> provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• <code>rbfbase</code> Radial Basis qkernel function "Gaussian"</li> <li>• <code>nonlbase</code> Non Linear qkernel function</li> <li>• <code>laplbase</code> Laplbase qkernel function</li> <li>• <code>ratibase</code> Rational Quadratic qkernel function</li> <li>• <code>multbase</code> Multiquadric qkernel function</li> <li>• <code>invbase</code> Inverse Multiquadric qkernel function</li> <li>• <code>wavbase</code> Wave qkernel function</li> <li>• <code>powbase</code> d qkernel function</li> <li>• <code>logbase</code> Log qkernel function</li> <li>• <code>caubase</code> Cauchy qkernel function</li> <li>• <code>chibase</code> Chi-Square qkernel function</li> <li>• <code>studbase</code> Generalized T-Student qkernel function</li> <li>• <code>nonlcnd</code> Non Linear cndkernel function</li> <li>• <code>polycnd</code> Polynomial cndkernel function</li> <li>• <code>rbfcnd</code> Radial Basis cndkernel function "Gaussian"</li> </ul> |

- `laplcnd` Laplacian cndkernel function
- `anocnd` ANOVA cndkernel function
- `raticnd` Rational Quadratic cndkernel function
- `multcnd` Multiquadric cndkernel function
- `invcnd` Inverse Multiquadric cndkernel function
- `wavcnd` Wave cndkernel function
- `powcnd` power cndkernel function
- `logcnd` Log cndkernel function
- `caucnd` Cauchy cndkernel function
- `chicnd` Chi-Square cndkernel function
- `studcnd` Generalized T-Student cndkernel function

The kernel parameter can also be set to a user defined function of class kernel by passing the function name as an argument.

`qpar`

the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :

- `sigma`, `q` for the Radial Basis qkernel function "`rbfbase`", the Laplacian qkernel function "`laplbase`" and the Cauchy qkernel function "`caubase`".
- `alpha`, `q` for the Non Linear qkernel function "`nonlbase`".
- `c`, `q` for the Rational Quadratic qkernel function "`ratibase`", the Multiquadric qkernel function "`multbase`" and the Inverse Multiquadric qkernel function "`invbase`".
- `theta`, `q` for the Wave qkernel function "`wavbase`".
- `d`, `q` for the d qkernel function "`powbase`", the Log qkernel function "`logbase`" and the Generalized T-Student qkernel function "`studbase`".
- `alpha` for the Non Linear cndkernel function "`nonlcnd`".
- `d`, `alpha`, `c` for the Polynomial cndkernel function "`polycnd`".
- `gamma` for the Radial Basis cndkernel function "`rbfcnd`" and the Laplacian cndkernel function "`laplcnd`" and the Cauchy cndkernel function "`caucnd`".
- `d`, `sigma` for the ANOVA cndkernel function "`anocnd`".
- `c` for the Rational Quadratic cndkernel function "`raticnd`", the Multiquadric cndkernel function "`multcnd`" and the Inverse Multiquadric cndkernel function "`invcnd`".
- `theta` for the Wave cndkernel function "`wavcnd`".
- `d` for the power cndkernel function "`powcnd`", the Log cndkernel function "`logcnd`" and the Generalized T-Student cndkernel function "`studcnd`".

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

`features`

Number of features (principal components) to return. (default: 0 , all)

`th`

the value of the eigenvalue under which principal components are ignored (only valid when `features = 0`). (default : 0.0001)

`na.action`

A function to specify the action to be taken if NAs are found. The default action is `na.omit`, which leads to rejection of cases with missing values on any required variable. An alternative is `na.fail`, which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)



... additional parameters

### Details

Using kernel functions one can efficiently compute principal components in high-dimensional feature spaces, related to input space by some non-linear map.

The data can be passed to the qkPCA function in a matrix, in addition qkPCA also supports input in the form of a kernel matrix of class qkernmatrix or class cndkernmatrix.

### Value

An S4 object containing the principal component vectors along with the corresponding eigenvalues.

pcv	a matrix containing the principal component vectors (column wise)
eVal	The corresponding eigenvalues
rotated	The original data projected (rotated) on the principal components
cndkernf	the kernel function used
xmatrix	The original data matrix

all the slots of the object can be accessed by accessor functions.

### Note

The predict function can be used to embed new data on the new space

### Author(s)

Yusen Zhang  
<yusenzhang@126.com>

### References

Schoelkopf B., A. Smola, K.-R. Mueller :  
*Nonlinear component analysis as a kernel eigenvalue problem*  
Neural Computation 10, 1299-1319  
<http://citeseerx.ist.psu.edu/viewdoc/summary?doi=10.1.1.29.1366>

### See Also

[qkernmatrix](#), [cndkernmatrix](#)

### Examples

```
# another example using the iris data
data(iris)
test <- sample(1:150,20)
qkpc <- qkPCA(~.,data=iris[-test,-5],kernel="rbfbase",
             qpar=list(sigma=50,q=0.8),features=2)

# print the principal component vectors
```

```
pcv(qkpc)
#plot the data projection on the components
plot(rotated(qkpc),col=as.integer(iris[-test,5]),
      xlab="1st Principal Component",ylab="2nd Principal Component")

# embed remaining points
emb <- predict(qkpc,iris[test,-5])
points(emb,col=as.integer(iris[test,5]))
```

---

qkpc-class

*Class "qkpc"*


---

### Description

The qkernel Principal Components Analysis class

### Objects of class "qkpc"

Objects can be created by calls of the form `new("qkpc", ...)`. or by calling the `qkpc` function.

### Slots

**pcv**: Object of class "matrix" containing the principal component vectors

**eVal**: Object of class "vector" containing the corresponding eigenvalues

**rotated**: Object of class "matrix" containing the projection of the data on the principal components

### Methods

**eVal** signature(object = "qkpc"): returns the eigenvalues

**pcv** signature(object = "qkpc"): returns the principal component vectors

**predict** signature(object = "qkpc"): embeds new data

**rotated** signature(object = "qkpc"): returns the projected data

**xmatrix** signature(object = "qkpc"): returns the used data matrix

**kcall** signature(object = "qkpc"): returns the performed call

**cndkernf** signature(object = "qkpc"): returns the used kernel function

### Author(s)

Yusen Zhang

<yusenzhang@126.com>

### See Also

[qkernel-class](#), [cndkernel-class](#)

**Examples**

```
# another example using the iris data
data(iris)
test <- sample(1:150,20)
qkpc <- qkpca(~.,iris[-test,-5], kernel = "rbfbase",
             qpar = list(sigma = 50, q = 0.8), features = 2)

# print the principal component vectors
pcv(qkpc)
rotated(qkpc)
cndkernf(qkpc)
eVal(qkpc)
xmatrix(qkpc)
names(eVal(qkpc))
```

---

qkprc-class

*Class "qkprc"*


---

**Description**

The qKernel Prehead class

**Objects of class "qkprc"**

Objects from the class cannot be created directly but only contained in other classes.

**Slots**

**cndkernf**: Object of class "kfunction" containing the kernel function used

**qpar**: Object of class "list" containing the kernel parameters used

**xmatrix**: Object of class "input" containing the data matrix used

**ymatrix**: Object of class "input" containing the data matrix used

**kcall**: Object of class "ANY" containing the function call

**terms**: Object of class "ANY" containing the function terms

**n.action**: Object of class "ANY" containing the action performed on NA

**Methods**

**cndkernf** signature(object = "qkprc"): returns the used kernel function

**xmatrix** signature(object = "qkprc"): returns the used data matrix

**ymatrix** signature(object = "qkprc"): returns the used data matrix

**kcall** signature(object = "qkprc"): returns the performed call

**Author(s)**

Yusen Zhang  
<yusenzhang@126.com>

**See Also**

[qkernel-class](#), [cndkernel-class](#)

---

qkspecc

*qkernel spectral Clustering*

---

**Description**

A qkernel spectral clustering algorithm. Clustering is performed by embedding the data into the subspace of the eigenvectors of a graph Laplacian matrix.

**Usage**

```
## S4 method for signature 'matrix'
qkspecc(x, kernel = "rbfbase", qpar = list(sigma = 2, q = 0.9),
        Nocent=NA, normalize="symmetric", maxk=20, iterations=200,
        na.action = na.omit, ...)

## S4 method for signature 'cndkernmatrix'
qkspecc(x, Nocent=NA, normalize="symmetric",
        maxk=20, iterations=200, ...)

## S4 method for signature 'qkernmatrix'
qkspecc(x, Nocent=NA, normalize="symmetric",
        maxk=20, iterations=200, ...)
```

**Arguments**

x	the matrix of data to be clustered or a kernel Matrix of class <code>qkernmatrix</code> or <code>cndkernmatrix</code> .
kernel	the kernel function used in computing the affinity matrix. This parameter can be set to any function, of class <code>kernel</code> , which computes a kernel function value between two vector arguments. <code>kernlab</code> provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings: <ul style="list-style-type: none"> <li>• <code>rbfbase</code> Radial Basis qkernel function "Gaussian"</li> <li>• <code>nonlbase</code> Non Linear qkernel function</li> <li>• <code>laplbase</code> Laplbase qkernel function</li> <li>• <code>ratibase</code> Rational Quadratic qkernel function</li> <li>• <code>multbase</code> Multiquadric qkernel function</li> <li>• <code>invbase</code> Inverse Multiquadric qkernel function</li> </ul>

- wavbase Wave qkernel function
- powbase d qkernel function
- logbase Log qkernel function
- caubase Cauchy qkernel function
- chibase Chi-Square qkernel function
- studbase Generalized T-Student qkernel function
- nonlcnd Non Linear cndkernel function
- polycnd Polynomial cndkernel function
- rbfcd Radial Basis cndkernel function "Gaussian"
- laplcnd Laplacian cndkernel function
- anocnd ANOVA cndkernel function
- raticnd Rational Quadratic cndkernel function
- multcnd Multiquadric cndkernel function
- invcnd Inverse Multiquadric cndkernel function
- wvcnd Wave cndkernel function
- powcnd d cndkernel function
- logcnd Log cndkernel function
- caucnd Cauchy cndkernel function
- chicnd Chi-Square cndkernel function
- studcnd Generalized T-Student cndkernel function

The kernel parameter can also be set to a user defined function of class kernel by passing the function name as an argument.

qpar

a character string or the list of hyper-parameters (kernel parameters). The default character string `list(sigma = 2, q = 0.9)` uses a heuristic to determine a suitable value for the width parameter of the RBF kernel. The second option "local" (local scaling) uses a more advanced heuristic and sets a width parameter for every point in the data set. This is particularly useful when the data incorporates multiple scales. A list can also be used containing the parameters to be used with the kernel function. Valid parameters for existing kernels are :

- sigma, q for the Radial Basis qkernel function "rbfbase" , the Laplacian qkernel function "laplbase" and the Cauchy qkernel function "caubase".
- alpha, q for the Non Linear qkernel function "nonlbase".
- c, q for the Rational Quadratic qkernel function "ratibase" , the Multiquadric qkernel function "multbase" and the Inverse Multiquadric qkernel function "invbase".
- theta, q for the Wave qkernel function "wavbase".
- d, q for the d qkernel function "powbase" , the Log qkernel function "logbase" and the Generalized T-Student qkernel function "studbase".
- alpha for the Non Linear cndkernel function "nonlcnd".
- d, alpha, c for the Polynomial cndkernel function "polycnd".
- gamma for the Radial Basis cndkernel function "rbfcd" and the Laplacian cndkernel function "laplcnd" and the Cauchy cndkernel function "caucnd".
- d, sigma for the ANOVA cndkernel function "anocnd".

- `c` for the Rational Quadratic cndkernel function "raticnd", the Multiquadric cndkernel function "multcnd" and the Inverse Multiquadric cndkernel function "invcnd".
- `theta` for the Wave cndkernel function "wavgnd".
- `d` for the d cndkernel function "powcnd", the Log cndkernel function "logcnd" and the Generalized T-Student cndkernel function "studcnd". where `length` is the length of the strings considered, `lambda` the decay factor and `normalized` a logical parameter determining if the kernel evaluations should be normalized.

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

<code>Nocent</code>	the number of clusters.
<code>normalize</code>	Normalisation of the Laplacian ("none", "symmetric" or "random-walk").
<code>maxk</code>	If <code>k</code> is NA, an upper bound for the automatic estimation. Defaults to 20.
<code>iterations</code>	the maximum number of iterations allowed.
<code>na.action</code>	the action to perform on NA.
<code>...</code>	additional parameters.

## Details

The `qkernel` spectral clustering works by embedding the data points of the partitioning problem into the subspace of the eigenvectors corresponding to the  $k$  smallest eigenvalues of the graph Laplacian matrix. Using a simple clustering method like `kmeans` on the embedded points usually leads to good performance. It can be shown that `qkernel` spectral clustering methods boil down to graph partitioning.

The data can be passed to the `qkspecc` function in a `matrix`, in addition `qkspecc` also supports input in the form of a kernel matrix of class `qkernelmatrix` or `cndkernelmatrix`.

## Value

An S4 object of class `qkspecc` which extends the class `vector` containing integers indicating the cluster to which each point is allocated. The following slots contain useful information

<code>clust</code>	The cluster assignments
<code>eVec</code>	The corresponding eigenvector
<code>eVal</code>	The corresponding eigenvalues
<code>ymatrix</code>	The eigenvectors corresponding to the $k$ smallest eigenvalues of the graph Laplacian matrix.

## Author(s)

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<yusenzhang@126.com>

**References**

Andrew Y. Ng, Michael I. Jordan, Yair Weiss  
*On Spectral Clustering: Analysis and an Algorithm*  
 Neural Information Processing Symposium 2001

**See Also**

[qkernelmatrix](#), [cndkernelmatrix](#), [qkpca](#)

**Examples**

```
data("iris")
x=as.matrix(iris[,-5])

qspe <- qkspecc(x, kernel = "rbfbase", qpar = list(sigma = 10, q = 0.9),
               Nocent=3, normalize="symmetric", maxk=15, iterations=1200)
plot(x, col = clust(qspe))

qkfunc <- nonlbase(alpha=1/15, q=0.8)
Ktrain <- qkernelmatrix(qkfunc, x)
qspe <- qkspecc(Ktrain, Nocent=3, normalize="symmetric", maxk=20)
plot(x, col = clust(qspe))
```

---

qkspecc-class

*Class "qkspecc"*


---

**Description**

The qKernel Spectral Clustering Class

**Objects from the Class**

Objects can be created by calls of the form `new("qkspecc", ...)`. or by calling the function `qkspecc`.

**Slots**

`clust`: Object of class "vector" containing the cluster assignments

`eVec`: Object of class "matrix" containing the corresponding eigenvector in each cluster

`eVal`: Object of class "vector" containing the corresponding eigenvalue for each cluster

`withinss`: Object of class "vector" containing the within-cluster sum of squares for each cluster

**Methods**

**clust** signature(object = "qkspecc"): returns the cluster assignments  
**eVec** signature(object = "qkspecc"): returns the corresponding eigenvector in each cluster  
**eVal** signature(object = "qkspecc"): returns the corresponding eigenvalue for each cluster  
**xmatrix** signature(object = "qkspecc"): returns the original data matrix or a kernel Matrix  
**ymatrix** signature(object = "qkspecc"): returns The eigenvectors corresponding to the  $k$  smallest eigenvalues of the graph Laplacian matrix.  
**cndkernf** signature(object = "qkspecc"): returns the used kernel function  
**kcall** signature(object = "qkspecc"): returns the performed call

**Author(s)**

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**See Also**

[qkspecc](#), [qkernel-class](#), [cndkernel-class](#)

**Examples**

```
## Cluster the iris data set.
data("iris")
x=as.matrix(iris[,-5])

qspe <- qkspecc(x,kernel = "rbfbase", qpar = list(sigma = 10, q = 0.9),
               Nocent=3, normalize="symmetric", maxk=15, iterations=1200)

clust(qspe)
eVec(qspe)
eVal(qspe)
xmatrix(qspe)
ymatrix(qspe)
cndkernf(qspe)
```

---

qkspeclust

*qkernel spectral Clustering*


---

**Description**

This is also a qkernel spectral clustering algorithm which uses three ways to assign labels after the laplacian embedding: kmeans, hclust and dbscan.

**Usage**

```
## S4 method for signature 'qkspecc'
qkspeclust(x, clustmethod = "kmeans",
           Nocent=NULL,iterations=NULL, hmethod=NULL,eps = NULL, MinPts = NULL)
```



**Arguments**

x	object of class qkspecc.
clustmethod	the strategy to use to assign labels in the embedding space. There are three ways to assign labels after the laplacian embedding: kmeans, hclust and dbscan.
Nocent	the number of clusters
iterations	the maximum number of iterations allowed for "kmeans".
hmethod	the agglomeration method for "hclust". This should be (an unambiguous abbreviation of) one of "ward.D", "ward.D2", "single", "complete", "average" (= UPGMA), "mcquitty" (= WPGMA), "median" (= WPGMC) or "centroid" (= UPGMC).
eps	Reachability distance for "dbscan".
MinPts	Reachability minimum no. of points for "dbscan".

**Details**

The qkernel spectral clustering works by embedding the data points of the partitioning problem into the subspace of the eigenvectors corresponding to the  $k$  smallest eigenvalues of the graph Laplacian matrix. Using the simple clustering methods like kmeans, hclust and dbscan on the embedded points usually leads to good performance. It can be shown that qkernel spectral clustering methods boil down to graph partitioning.

**Value**

An S4 object of class qkspecc which extends the class vector containing integers indicating the cluster to which each point is allocated. The following slots contain useful information

clust	The cluster assignments
eVec	The corresponding eigenvector
eVal	The corresponding eigenvalues
xmatrix	The original data matrix
ymatrix	The real valued matrix of eigenvectors corresponding to the $k$ smallest eigenvalues of the graph Laplacian matrix
cnkernf	The kernel function used

**Author(s)**

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<yusenzhang@126.com>

**References**

Andrew Y. Ng, Michael I. Jordan, Yair Weiss  
*On Spectral Clustering: Analysis and an Algorithm*  
Neural Information Processing Symposium 2001

**See Also**

[qkernelmatrix](#), [cndkernelmatrix](#), [qkspecc-class](#), [qkspecc](#)

**Examples**

```
data("iris")
x=as.matrix(iris[ , -5])

qspe <- qkspecc(x, kernel = "rbfbase", qpar = list(sigma = 90, q = 0.9),
               Nocent=3, normalize="symmetric", maxk=15, iterations=1200)
plot(x, col = clust(qspe))

qspec <- qkspeclust(qspe, clustmethod = "hclust", Nocent=3, hmethod="ward.D2")
plot(x, col = clust(qspec))
plot(qspec)
```

---

 qsammon

*qKernel Sammon Mapping*


---

**Description**

The qkernel Sammon Mapping is an implementation for Sammon mapping, one of the earliest dimension reduction techniques that aims to find low-dimensional embedding that preserves pairwise distance structure in high-dimensional data space. qsammon is a nonlinear form of Sammon Mapping.

**Usage**

```
## S4 method for signature 'matrix'
qsammon(x, kernel = "rbfbase", qpar = list(sigma = 0.5, q = 0.9),
        dims = 2, Initialisation = 'random', MaxHalves = 20,
        MaxIter = 500, TolFun = 1e-7, na.action = na.omit, ...)

## S4 method for signature 'cndkernelmatrix'
qsammon(cndkernel, x, k, dims = 2, Initialisation = 'random',
        MaxHalves = 20, MaxIter = 500, TolFun = 1e-7, ...)

## S4 method for signature 'qkernelmatrix'
qsammon(qkernel, x, k, dims = 2, Initialisation = 'random',
        MaxHalves = 20, MaxIter = 500, TolFun = 1e-7, ...)
```

**Arguments**

x	the data matrix indexed by row or a kernel matrix of <code>condkernelmatrix</code> or <code>qkernelmatrix</code> .
kernel	<p>the kernel function used in training and predicting. This parameter can be set to any function, of class <code>kernel</code>, which computes a kernel function value between two vector arguments. <code>qkerneltool</code> provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• <code>rbfbase</code> Radial Basis <code>qkernel</code> function "Gaussian"</li> <li>• <code>nonlbase</code> Non Linear <code>qkernel</code> function</li> <li>• <code>laplbase</code> Laplbase <code>qkernel</code> function</li> <li>• <code>ratibase</code> Rational Quadratic <code>qkernel</code> function</li> <li>• <code>multbase</code> Multiquadric <code>qkernel</code> function</li> <li>• <code>invbase</code> Inverse Multiquadric <code>qkernel</code> function</li> <li>• <code>wavbase</code> Wave <code>qkernel</code> function</li> <li>• <code>powbase d</code> <code>qkernel</code> function</li> <li>• <code>logbase</code> Log <code>qkernel</code> function</li> <li>• <code>caubase</code> Cauchy <code>qkernel</code> function</li> <li>• <code>chibase</code> Chi-Square <code>qkernel</code> function</li> <li>• <code>studbase</code> Generalized T-Student <code>qkernel</code> function</li> <li>• <code>nonlcnd</code> Non Linear <code>condkernel</code> function</li> <li>• <code>polycnd</code> Polynomial <code>condkernel</code> function</li> <li>• <code>rbfcnd</code> Radial Basis <code>condkernel</code> function "Gaussian"</li> <li>• <code>laplcnd</code> Laplacian <code>condkernel</code> function</li> <li>• <code>anocnd</code> ANOVA <code>condkernel</code> function</li> <li>• <code>ratıcnd</code> Rational Quadratic <code>condkernel</code> function</li> <li>• <code>multcnd</code> Multiquadric <code>condkernel</code> function</li> <li>• <code>invcnd</code> Inverse Multiquadric <code>condkernel</code> function</li> <li>• <code>wavcnd</code> Wave <code>condkernel</code> function</li> <li>• <code>powcnd d</code> <code>condkernel</code> function</li> <li>• <code>logcnd</code> Log <code>condkernel</code> function</li> <li>• <code>caucnd</code> Cauchy <code>condkernel</code> function</li> <li>• <code>chıcnd</code> Chi-Square <code>condkernel</code> function</li> <li>• <code>studcnd</code> Generalized T-Student <code>condkernel</code> function</li> </ul> <p>The kernel parameter can also be set to a user defined function of class <code>kernel</code> by passing the function name as an argument.</p>
qpar	<p>the list of hyper-parameters (kernel parameters). This is a list which contains the parameters to be used with the kernel function. Valid parameters for existing kernels are :</p> <ul style="list-style-type: none"> <li>• <code>sigma</code>, <code>q</code> for the Radial Basis <code>qkernel</code> function "<code>rbfbase</code>" , the Laplacian <code>qkernel</code> function "<code>laplbase</code>" and the Cauchy <code>qkernel</code> function "<code>caubase</code>".</li> <li>• <code>alpha</code>, <code>q</code> for the Non Linear <code>qkernel</code> function "<code>nonlbase</code>".</li> <li>• <code>c</code>, <code>q</code> for the Rational Quadratic <code>qkernel</code> function "<code>ratibase</code>" , the Multi-quadric <code>qkernel</code> function "<code>multbase</code>" and the Inverse Multiquadric <code>qkernel</code> function "<code>invbase</code>".</li> </ul>

- `theta`, `q` for the Wave `qkernel` function "wavbase".
- `d`, `q` for the `d kernel` function "powbase", the Log `kernel` function "logbase" and the Generalized T-Student `kernel` function "studbase".
- `alpha` for the Non Linear `cndkernel` function "nonlnd".
- `d`, `alpha`, `c` for the Polynomial `cndkernel` function "polycnd".
- `gamma` for the Radial Basis `cndkernel` function "rbfcnd" and the Laplacian `cndkernel` function "laplnd" and the Cauchy `cndkernel` function "caucnd".
- `d`, `sigma` for the ANOVA `cndkernel` function "anocnd".
- `c` for the Rational Quadratic `cndkernel` function "raticnd", the Multiquadric `cndkernel` function "multcnd" and the Inverse Multiquadric `cndkernel` function "invcnd".
- `theta` for the Wave `cndkernel` function "wavcnd".
- `d` for the `d cndkernel` function "powcnd", the Log `cndkernel` function "logcnd" and the Generalized T-Student `cndkernel` function "studcnd".

Hyper-parameters for user defined kernels can be passed through the `qpar` parameter as well.

<code>kernel</code>	the kernel function to be used to calculate the <code>kernel</code> matrix.
<code>cndkernel</code>	the <code>cndkernel</code> function to be used to calculate the <code>CND</code> kernel matrix.
<code>k</code>	the dimension of the original data.
<code>dims</code>	Number of features to return. (default: 2)
<code>Initialisation</code>	"random" or "pca"; the former performs fast random projection and the latter performs standard PCA (default : "random")
<code>MaxHalves</code>	maximum number of step halvings. (default : 20)
<code>MaxIter</code>	the maximum number of iterations allowed. (default : 500)
<code>TolFun</code>	relative tolerance on objective function. (default : 1e-7)
<code>na.action</code>	A function to specify the action to be taken if NAs are found. The default action is <code>na.omit</code> , which leads to rejection of cases with missing values on any required variable. An alternative is <code>na.fail</code> , which causes an error if NA cases are found. (NOTE: If given, this argument must be named.)
<code>...</code>	additional parameters

## Details

Using kernel functions one can efficiently compute principal components in high-dimensional feature spaces, related to input space by some non-linear map.

The data can be passed to the `qsammon` function in a `matrix`, in addition `qsammon` also supports input in the form of a kernel matrix of class `qkernelmatrix` or class `cndkernelmatrix`.

## Value

<code>dimRed</code>	The matrix whose rows are embedded observations.
<code>kcall</code>	The function call contained
<code>cndkernelf</code>	The kernel function used
all the slots of the object can be accessed by accessor functions.	

**Author(s)**

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**References**

Sammon, J.W. (1969) *A Nonlinear Mapping for Data Structure Analysis*. IEEE Transactions on Computers, C-18 5:401-409.

**See Also**

[qkernelmatrix](#), [cndkernelmatrix](#)

**Examples**

```
data(iris)
train <- as.matrix(iris[,1:4])
labeltrain <- as.integer(iris[,5])
## S4 method for signature 'matrix'
kpc2 <- qsammon(train, kernel = "rbfbase", qpar = list(sigma = 2, q = 0.9), dims = 2,
               Initialisation = 'pca', TolFun = 1e-5)
plot(dimRed(kpc2), col = as.integer(labeltrain))
cndkernel(kpc2)
```

---

 qsammon-class

 Class "qsammon"
 

---

**Description**

The qKernel Sammon Mapping class

**Objects of class "qsammon"**

Objects can be created by calls of the form `new("qsammon", ...)`. or by calling the `qsammon` function.

**Slots**

`dimRed`: Object of class "matrix" containing the matrix whose rows are embedded observations

`cndkernel`: Object of class "function" containing the kernel function used

`kcall`: Object of class "ANY" containing the function call

**Methods**

**dimRed** signature(object = "qsammon"): returns the matrix whose rows are embedded observations

**kcall** signature(object = "qsammon"): returns the performed call

**cndkernf** signature(object = "qsammon"): returns the used kernel function

**Author(s)**

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**See Also**

[qsammon](#)

**Examples**

```
data(iris)
train <- as.matrix(iris[,1:4])
labeltrain<- as.integer(iris[,5])
## S4 method for signature 'matrix'
qkpc <- qsammon(train, kernel = "rbfbase", qpar = list(sigma = 0.5, q = 0.9),
               dims = 2, Initialisation = 'pca', MaxHalves = 50)

cndkernf(qkpc)
dimRed(qkpc)
kcall(qkpc)
```

---

qtSNE

*qKernel t-Distributed Stochastic Neighbor Embedding*

---

**Description**

Wrapper for the qkernel t-distributed stochastic neighbor embeddingg. qtSNE is a method for constructing a low dimensional embedding of high-dimensional data, distances or similarities.

**Usage**

```
## S4 method for signature 'matrix'
qtSNE(x, kernel = "rbfbase", qpar = list(sigma = 0.1, q = 0.9),
      initial_config = NULL, no_dims=2, initial_dims=30, perplexity=30, max_iter= 1300,
      min_cost=0, epoch_callback=NULL, epoch=100, na.action = na.omit, ...)
## S4 method for signature 'cndkernmatrix'
qtSNE(x, initial_config = NULL, no_dims=2, initial_dims=30,
      perplexity=30, max_iter = 1000, min_cost=0, epoch_callback=NULL, epoch=100)
## S4 method for signature 'qkernmatrix'
qtSNE(x, initial_config = NULL, no_dims=2, initial_dims=30,
      perplexity=30, max_iter = 1000, min_cost=0, epoch_callback=NULL, epoch=100)
```

**Arguments**

x	the matrix of data to be clustered or a kernel Matrix of class <code>qkernelmatrix</code> or <code>cnkernelmatrix</code> .
kernel	<p>the kernel function used in computing the affinity matrix. This parameter can be set to any function, of class <code>kernel</code>, which computes a kernel function value between two vector arguments. <code>kernlab</code> provides the most popular kernel functions which can be used by setting the kernel parameter to the following strings:</p> <ul style="list-style-type: none"> <li>• <code>rbfbase</code> Radial Basis <code>qkernel</code> function "Gaussian"</li> <li>• <code>nonlbase</code> Non Linear <code>qkernel</code> function</li> <li>• <code>laplbase</code> Laplbase <code>qkernel</code> function</li> <li>• <code>ratibase</code> Rational Quadratic <code>qkernel</code> function</li> <li>• <code>multbase</code> Multiquadric <code>qkernel</code> function</li> <li>• <code>invbase</code> Inverse Multiquadric <code>qkernel</code> function</li> <li>• <code>wavbase</code> Wave <code>qkernel</code> function</li> <li>• <code>powbase</code> Power <code>qkernel</code> function</li> <li>• <code>logbase</code> Log <code>qkernel</code> function</li> <li>• <code>caubase</code> Cauchy <code>qkernel</code> function</li> <li>• <code>chibase</code> Chi-Square <code>qkernel</code> function</li> <li>• <code>studbase</code> Generalized T-Student <code>qkernel</code> function</li> <li>• <code>nonlcnd</code> Non Linear <code>cnkernel</code> function</li> <li>• <code>polycnd</code> Polynomial <code>cnkernel</code> function</li> <li>• <code>rbfcnd</code> Radial Basis <code>cnkernel</code> function "Gaussian"</li> <li>• <code>laplcnd</code> Laplacian <code>cnkernel</code> function</li> <li>• <code>anocnd</code> ANOVA <code>cnkernel</code> function</li> <li>• <code>ratıcnd</code> Rational Quadratic <code>cnkernel</code> function</li> <li>• <code>multcnd</code> Multiquadric <code>cnkernel</code> function</li> <li>• <code>invcnd</code> Inverse Multiquadric <code>cnkernel</code> function</li> <li>• <code>wavcnd</code> Wave <code>cnkernel</code> function</li> <li>• <code>powcnd</code> Power <code>cnkernel</code> function</li> <li>• <code>logcnd</code> Log <code>cnkernel</code> function</li> <li>• <code>caucnd</code> Cauchy <code>cnkernel</code> function</li> <li>• <code>chıcnd</code> Chi-Square <code>cnkernel</code> function</li> <li>• <code>studcnd</code> Generalized T-Student <code>cnkernel</code> function</li> </ul> <p>The kernel parameter can also be set to a user defined function of class <code>kernel</code> by passing the function name as an argument.</p>
qpar	a character string or the list of hyper-parameters (kernel parameters). The default character string <code>list(sigma = 2, q = 0.9)</code> uses a heuristic to determine a suitable value for the width parameter of the RBF kernel. The second option "local" (local scaling) uses a more advanced heuristic and sets a width parameter for every point in the data set. This is particularly useful when the data incorporates multiple scales. A list can also be used containing the parameters to be used with the kernel function. Valid parameters for existing kernels are :

- `sigma` for the Radial Basis qkernel function "rbfbase", the Laplacian qkernel function "laplbase" the Cauchy qkernel function "caubase" and for the ANOVA cndkernel function "anocnd".
- `alpha` for the Non Linear qkernel function "nonlbase",for the Non Linear cndkernel function "nonlcnd",and for the Polynomial cndkernel function "polycnd".
- `c` for the Rational Quadratic qkernel function "ratibase", the Multiquadric qkernel function "multibase", the Inverse Multiquadric qkernel function "invbase",for the Polynomial cndkernel function "polycnd",for the Rational Quadratic cndkernel function "raticnd", the Multiquadric cndkernel function "multcnd" and the Inverse Multiquadric cndkernel function "invcnd".
- `d` for qkernel function "powbase", the Log qkernel function "logbase", the Generalized T-Student qkernel function "studbase", for the Polynomial cndkernel function "polycnd", for the ANOVA cndkernel function "anocnd",for the d cndkernel function "powcnd", the Log cndkernel function "logcnd" and the Generalized T-Student cndkernel function "studcnd".
- `theta` for the Wave qkernel function "wavbase" and for the Wave cndkernel function "wavcnd".
- `gamma` for the Chi-Square qkernel function "chibase",for the Radial Basis cndkernel function "rbfcnd" and the Laplacian cndkernel function "laplcnd" and the Cauchy cndkernel function "caucnd".
- `q` For all qkernel Function. where length is the length of the strings considered, lambda the decay factor and normalized a logical parameter determining if the kernel evaluations should be normalized.

Hyper-parameters for user defined kernels can be passed through the `qkpar` parameter as well.

<code>initial_config</code>	An intitial configure about x (default: NULL)
<code>no_dims</code>	the dimension of the resulting embedding. (default: 2)
<code>initial_dims</code>	The number of dimensions to use in reduction method. (default: 30)
<code>perplexity</code>	Perplexity parameter
<code>max_iter</code>	Number of iterations (default: 1300)
<code>min_cost</code>	The minimum cost for every object after the final iteration
<code>epoch_callback</code>	A callback function used after each epoch (an epoch here means a set number of iterations)
<code>epoch</code>	The interval of the number of iterations displayed (default: 100)
<code>na.action</code>	the action to perform on NA
<code>...</code>	Other arguments that can be passed to qtSNE

## Details

When the `initial_config` argument is specified, the algorithm will automatically enter the final momentum stage. This stage has less large scale adjustment to the embedding, and is intended for small scale tweaking of positioning. This can greatly speed up the generation of embeddings for various similar X datasets, while also preserving overall embedding orientation.



**Value**

qtSNE gives out an S4 object which is a LIST with components

dimRed            Matrix containing the new representations for the objects after qtSNE  
cndkernf          The kernel function used

**Author(s)**

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**References**

Maaten, L. Van Der, 2014. Accelerating t-SNE using Tree-Based Algorithms. Journal of Machine Learning Research, 15, p.3221-3245.

van der Maaten, L.J.P. & Hinton, G.E., 2008. Visualizing High-Dimensional Data Using t-SNE. Journal of Machine Learning Research, 9, pp.2579-2605.

**Examples**

```
## Not run:
#use iris data set
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[,1:4])

colors = rainbow(length(unique(iris$Species)))
names(colors) = unique(iris$Species)
#for matrix
ecb = function(x,y){
  plot(x,t='n');
  text(x,labels=iris$Species, col=colors[iris$Species])
}
kpc2 <- qtSNE(train, kernel = "rbfbase", qpar = list(sigma=1,q=0.8),
  epoch_callback = ecb, perplexity=10, max_iter = 500)

## End(Not run)
```

---

qtSNE-class

*Class "qtSNE"*

---

**Description**

An S4 Class for qtSNE.

## Details

The qtSNE is a method that uses Qkernel t-Distributed Stochastic Neighborhood Embedding between the distance matrices in high and low-dimensional space to embed the data. The method is very well suited to visualize complex structures in low dimensions.

## Objects from the Class

Objects can be created by calls of the form `new("qtSNE", ...)`. or by calling the function `qtSNE`.

## Slots

`dimRed` Matrix containing the new representations for the objects after qtSNE

`cndkernf` The kernel function used

## Method

`dimRed` signature(object="qtSNE"): return a new representation matrix

`cndkernf` signature(object="qtSNE"): return the kernel used

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## References

Maaten, L. van der, 2014. Accelerating t-SNE using Tree-Based Algorithms. Journal of Machine Learning Research 15, 3221-3245.

van der Maaten, L., Hinton, G., 2008. Visualizing Data using t-SNE. J. Mach. Learn. Res. 9, 2579-2605.

## See Also

[qtSNE](#)

## Examples

```
## Not run:
#use iris data set
data(iris)
testset <- sample(1:150,20)
train <- as.matrix(iris[,1:4])

colors = rainbow(length(unique(iris$Species)))
names(colors) = unique(iris$Species)
#for matrix
ecb = function(x,y){
  plot(x,t='n');
  text(x,labels=iris$Species, col=colors[iris$Species])
}
```

```
kpc2 <- qtSNE(train, kernel = "rbfbase", qpar = list(sigma=1,q=0.8),
              epoch_callback = ecb, perplexity=10, max_iter = 500)

#cndernf
cndkernf(kpc2)

#dimRed
plot(dimRed(kpc2),col=train)

## End(Not run)
```

# Index

- \*Topic **algebra**
  - [cndkernmatrix](#), [8](#)
  - [qkernmatrix](#), [19](#)
- \*Topic **array**
  - [cndkernmatrix](#), [8](#)
  - [qkernmatrix](#), [19](#)
- \*Topic **classes**
  - [cndkernel-class](#), [7](#)
  - [qkdbscan-class](#), [16](#)
  - [qkernel-class](#), [17](#)
  - [qkgda-class](#), [24](#)
  - [qkIsomap-class](#), [28](#)
  - [qkLLE-class](#), [33](#)
  - [qkMDS-class](#), [37](#)
  - [qkpca-class](#), [42](#)
  - [qkprc-class](#), [43](#)
  - [qkspecc-class](#), [47](#)
  - [qsammon-class](#), [53](#)
- \*Topic **classif**
  - [qkgda](#), [20](#)
- \*Topic **cluster**
  - [qkdbscan](#), [13](#)
  - [qkpca](#), [39](#)
  - [qkspecc](#), [44](#)
  - [qkspeclust](#), [48](#)
  - [qsammon](#), [50](#)
- \*Topic **datasets**
  - [mfeat\\_pix](#), [12](#)
- \*Topic **methods**
  - [as.cndkernmatrix](#), [2](#)
  - [as.qkernmatrix](#), [3](#)
- \*Topic **symbolmath**
  - [bases](#), [4](#)
  - [cnds](#), [9](#)
- [anocnd](#), [20](#)
- [anocnd \(cnds\)](#), [9](#)
- [anokernel-class \(cndkernel-class\)](#), [7](#)
- [as.cndkernmatrix](#), [2](#)
- [as.cndkernmatrix, matrix-method \(as.cndkernmatrix\)](#), [2](#)
- [as.cndkernmatrix-methods \(as.cndkernmatrix\)](#), [2](#)
- [as.qkernmatrix](#), [3](#)
- [as.qkernmatrix, matrix-method \(as.qkernmatrix\)](#), [3](#)
- [as.qkernmatrix-methods \(as.qkernmatrix\)](#), [3](#)
- [bases](#), [4](#)
- [blkdiag](#), [6](#)
- [caubase](#), [9](#)
- [caubase \(bases\)](#), [4](#)
- [caucnd](#), [20](#)
- [caucnd \(cnds\)](#), [9](#)
- [caukernel-class \(cndkernel-class\)](#), [7](#)
- [cauqkernel-class \(qkernel-class\)](#), [17](#)
- [chibase](#), [9](#)
- [chibase \(bases\)](#), [4](#)
- [chicnd](#), [20](#)
- [chicnd \(cnds\)](#), [9](#)
- [chikernel-class \(cndkernel-class\)](#), [7](#)
- [chiqkernel-class \(qkernel-class\)](#), [17](#)
- [clust \(qkdbscan-class\)](#), [16](#)
- [clust, qkdbscan-method \(qkdbscan-class\)](#), [16](#)
- [clust, qkspecc-method \(qkspecc-class\)](#), [47](#)
- [clust<- \(qkdbscan-class\)](#), [16](#)
- [clust<-, qkdbscan-method \(qkdbscan-class\)](#), [16](#)
- [clust<-, qkspecc-method \(qkspecc-class\)](#), [47](#)
- [cndkernel-class](#), [7](#)
- [cndkernf \(qkprc-class\)](#), [43](#)
- [cndkernf, qkprc-method \(qkprc-class\)](#), [43](#)
- [cndkernf<- \(qkprc-class\)](#), [43](#)
- [cndkernf<-, qkprc-method \(qkprc-class\)](#), [43](#)

- [3](#), [4](#), [6](#), [7](#), [8](#), [11](#), [18](#), [23](#), [41](#), [47](#), [50](#), [53](#)
  - anokernel-method (cndkernmatrix), [8](#)
  - caukernel-method (cndkernmatrix), [8](#)
  - chikernel-method (cndkernmatrix), [8](#)
  - cndkernel-method (cndkernmatrix), [8](#)
  - invkernel-method (cndkernmatrix), [8](#)
  - laplkernel-method (cndkernmatrix), [8](#)
  - logkernel-method (cndkernmatrix), [8](#)
  - multkernel-method (cndkernmatrix), [8](#)
  - nonlkernel-method (cndkernmatrix), [8](#)
  - polykernel-method (cndkernmatrix), [8](#)
  - powkernel-method (cndkernmatrix), [8](#)
  - ratikernel-method (cndkernmatrix), [8](#)
  - rbfkernel-method (cndkernmatrix), [8](#)
  - studkernel-method (cndkernmatrix), [8](#)
  - wavkernel-method (cndkernmatrix), [8](#)
- cndkernmatrix-class (as.cndkernmatrix), [2](#)
- anokernel (cndkernmatrix), [8](#)
  - caukernel (cndkernmatrix), [8](#)
  - chikernel (cndkernmatrix), [8](#)
  - invkernel (cndkernmatrix), [8](#)
  - laplkernel (cndkernmatrix), [8](#)
  - logkernel (cndkernmatrix), [8](#)
  - multkernel (cndkernmatrix), [8](#)
  - nonlkernel (cndkernmatrix), [8](#)
  - polykernel (cndkernmatrix), [8](#)
  - powkernel (cndkernmatrix), [8](#)
  - ratikernel (cndkernmatrix), [8](#)
  - rbfkernel (cndkernmatrix), [8](#)
  - studkernel (cndkernmatrix), [8](#)
  - wavkernel (cndkernmatrix), [8](#)
- nonlkernel (cndkernmatrix), [8](#)
  - polykernel (cndkernmatrix), [8](#)
  - powkernel (cndkernmatrix), [8](#)
  - ratikernel (cndkernmatrix), [8](#)
  - rbfkernel (cndkernmatrix), [8](#)
  - studkernel (cndkernmatrix), [8](#)
  - wavkernel (cndkernmatrix), [8](#)
- cnds, [9](#)
- connum (qkIsomap-class), [28](#)
- connum, qkIsomap-method (qkIsomap-class), [28](#)
- connum, qkMDS-method (qkMDS-class), [37](#)
- connum<- (qkIsomap-class), [28](#)
- connum<- , qkIsomap-method (qkIsomap-class), [28](#)
- connum<- , qkMDS-method (qkMDS-class), [37](#)
- dimRed (qsammon-class), [53](#)
- dimRed, qsammon-method (qsammon-class), [53](#)
- dimRed, qtSNE-method (qtSNE-class), [57](#)
- dimRed<- (qsammon-class), [53](#)
- dimRed<- , qsammon-method (qsammon-class), [53](#)
- dimRed<- , qtSNE-method (qtSNE-class), [57](#)
- dims (qkIsomap-class), [28](#)
- dims, qkIsomap-method (qkIsomap-class), [28](#)
- dims, qkLLE-method (qkLLE-class), [33](#)
- dims, qkMDS-method (qkMDS-class), [37](#)
- dims<- (qkIsomap-class), [28](#)
- dims<- , qkIsomap-method (qkIsomap-class), [28](#)
- dims<- , qkLLE-method (qkLLE-class), [33](#)
- dims<- , qkMDS-method (qkMDS-class), [37](#)
- eps (qkdbscan-class), [16](#)
- eps, qkdbscan-method (qkdbscan-class), [16](#)
- eps<- (qkdbscan-class), [16](#)
- eps<- , qkdbscan-method (qkdbscan-class), [16](#)
- Euclidist, [11](#)

- Eucdist, matrix-method (Eucdist), 11
- eVal (qkgda-class), 24
- eVal, qkgda-method (qkgda-class), 24
- eVal, qkIsomap-method (qkIsomap-class), 28
- eVal, qkLLE-method (qkLLE-class), 33
- eVal, qkMDS-method (qkMDS-class), 37
- eVal, qkpca-method (qkpca-class), 42
- eVal, qkspecc-method (qkspecc-class), 47
- eVal<- (qkgda-class), 24
- eVal<-, qkgda-method (qkgda-class), 24
- eVal<-, qkIsomap-method (qkIsomap-class), 28
- eVal<-, qkLLE-method (qkLLE-class), 33
- eVal<-, qkMDS-method (qkMDS-class), 37
- eVal<-, qkpca-method (qkpca-class), 42
- eVal<-, qkspecc-method (qkspecc-class), 47
- eVec (qkgda-class), 24
- eVec, qkgda-method (qkgda-class), 24
- eVec, qkIsomap-method (qkIsomap-class), 28
- eVec, qkLLE-method (qkLLE-class), 33
- eVec, qkMDS-method (qkMDS-class), 37
- eVec, qkspecc-method (qkspecc-class), 47
- eVec<- (qkgda-class), 24
- eVec<-, qkgda-method (qkgda-class), 24
- eVec<-, qkIsomap-method (qkIsomap-class), 28
- eVec<-, qkLLE-method (qkLLE-class), 33
- eVec<-, qkMDS-method (qkMDS-class), 37
- eVec<-, qkspecc-method (qkspecc-class), 47
  
- fun (qsammon-class), 53
  
- input-class (qkernel-class), 17
- invbase, 9
- invbase (bases), 4
- invcnd, 20
- invcnd (cnnds), 9
- invkernel-class (cndkernel-class), 7
- invqkernel-class (qkernel-class), 17
- isseed (qkdbscan-class), 16
- isseed, qkdbscan-method (qkdbscan-class), 16
- isseed<- (qkdbscan-class), 16
- isseed<-, qkdbscan-method (qkdbscan-class), 16
  
- kcall (qkprc-class), 43
- kcall, qkprc-method (qkprc-class), 43
- kcall<- (qkprc-class), 43
- kcall<-, qkprc-method (qkprc-class), 43
- kfunction-class (qkernel-class), 17
  
- label (qkgda-class), 24
- label, qkgda-method (qkgda-class), 24
- label<- (qkgda-class), 24
- label<-, qkgda-method (qkgda-class), 24
- laplbase, 9
- laplbase (bases), 4
- laplcnd, 20
- laplcnd (cnnds), 9
- laplkernel-class (cndkernel-class), 7
- laplqkernel-class (qkernel-class), 17
- logbase, 9
- logbase (bases), 4
- logcnd, 20
- logcnd (cnnds), 9
- logkernel-class (cndkernel-class), 7
- logqkernel-class (qkernel-class), 17
  
- mfeat\_pix, 12
- MinPts (qkdbscan-class), 16
- MinPts, qkdbscan-method (qkdbscan-class), 16
- MinPts<- (qkdbscan-class), 16
- MinPts<-, qkdbscan-method (qkdbscan-class), 16
  
- multbase, 9
- multbase (bases), 4
- multcnd, 20
- multcnd (cnnds), 9
- multkernel-class (cndkernel-class), 7
- multqkernel-class (qkernel-class), 17
  
- n.action (qkprc-class), 43
- n.action, qkprc-method (qkprc-class), 43
- n.action<- (qkprc-class), 43
- n.action<-, qkprc-method (qkprc-class), 43
- nonlbase, 9
- nonlbase (bases), 4
- nonlcnd, 20
- nonlcnd (cnnds), 9
- nonlkernel-class (cndkernel-class), 7
- nonlqkernel-class (qkernel-class), 17
  
- pcv (qkpca-class), 42

- pcv, qkpca-method (qkpca-class), 42
- pcv<- (qkpca-class), 42
- pcv<- , qkpca-method (qkpca-class), 42
- plot (qkdbscan-class), 16
- plot, qkdbscan-method (qkdbscan-class), 16
- plot, qkspecc-method (qkspecc-class), 47
- polycnd, 20
- polycnd (cnds), 9
- polykernel-class (cndkernel-class), 7
- powbase, 9
- powbase (bases), 4
- powcnd, 20
- powcnd (cnds), 9
- powkernel-class (cndkernel-class), 7
- powqkernel-class (qkernel-class), 17
- predict, qkdbscan-method (qkdbscan), 13
- predict, qkgda-method (qkgda), 20
- predict, qkpca-method (qkpca), 39
- print, qkdbscan-method (qkdbscan), 13
- prj (qkgda-class), 24
- prj, qkgda-method (qkgda-class), 24
- prj, qkIsomap-method (qkIsomap-class), 28
- prj, qkLLE-method (qkLLE-class), 33
- prj, qkMDS-method (qkMDS-class), 37
- prj<- (qkgda-class), 24
- prj<- , qkgda-method (qkgda-class), 24
- prj<- , qkIsomap-method (qkIsomap-class), 28
- prj<- , qkLLE-method (qkLLE-class), 33
- prj<- , qkMDS-method (qkMDS-class), 37
  
- qkdbscan, 13
- qkdbscan, cndkernelmatrix-method (qkdbscan), 13
- qkdbscan, matrix-method (qkdbscan), 13
- qkdbscan, qkernelmatrix-method (qkdbscan), 13
- qkdbscan-class, 16
- qkernel-class, 17
- qkernelmatrix, 3, 4, 6, 7, 11, 16, 18, 19, 23, 41, 47, 50, 53
- qkernelmatrix, cauqkernel-method (qkernelmatrix), 19
- qkernelmatrix, chiqkernel-method (qkernelmatrix), 19
- qkernelmatrix, invqkernel-method (qkernelmatrix), 19
- qkernelmatrix, laplqkernel-method (qkernelmatrix), 19
- qkernelmatrix, logqkernel-method (qkernelmatrix), 19
- qkernelmatrix, multqkernel-method (qkernelmatrix), 19
- qkernelmatrix, nonlqkernel-method (qkernelmatrix), 19
- qkernelmatrix, powqkernel-method (qkernelmatrix), 19
- qkernelmatrix, qkernel-method (qkernelmatrix), 19
- qkernelmatrix, ratiqkernel-method (qkernelmatrix), 19
- qkernelmatrix, rbfqkernel-method (qkernelmatrix), 19
- qkernelmatrix, studqkernel-method (qkernelmatrix), 19
- qkernelmatrix, wavqkernel-method (qkernelmatrix), 19
- qkernelmatrix-class (as. qkernelmatrix), 3
- qkernelmatrix.cauqkernel (qkernelmatrix), 19
- qkernelmatrix.chiqkernel (qkernelmatrix), 19
- qkernelmatrix.invqkernel (qkernelmatrix), 19
- qkernelmatrix.laplqkernel (qkernelmatrix), 19
- qkernelmatrix.logqkernel (qkernelmatrix), 19
- qkernelmatrix.multqkernel (qkernelmatrix), 19
- qkernelmatrix.nonlqkernel (qkernelmatrix), 19
- qkernelmatrix.powqkernel (qkernelmatrix), 19
- qkernelmatrix.ratiqkernel (qkernelmatrix), 19
- qkernelmatrix.rbfqkernel (qkernelmatrix), 19
- qkernelmatrix.studqkernel (qkernelmatrix), 19
- qkernelmatrix.wavqkernel (qkernelmatrix), 19
- qkgda, 20
- qkgda, cndkernelmatrix-method (qkgda), 20
- qkgda, matrix-method (qkgda), 20
- qkgda, qkernelmatrix-method (qkgda), 20
- qkgda-class, 24
- qkIsomap, 25, 29
- qkIsomap, cndkernelmatrix-method (qkIsomap), 25
- qkIsomap, matrix-method (qkIsomap), 25
- qkIsomap, qkernelmatrix-method (qkIsomap),

- 25
- qkIsomap-class, 28
- qkLLE, 30
- qkLLE, cndkernmatrix-method (qkLLE), 30
- qkLLE, matrix-method (qkLLE), 30
- qkLLE, qkernmatrix-method (qkLLE), 30
- qkLLE-class, 33
- qkMDS, 34, 38
- qkMDS, cndkernmatrix-method (qkMDS), 34
- qkMDS, matrix-method (qkMDS), 34
- qkMDS, qkernmatrix-method (qkMDS), 34
- qkMDS-class, 37
- qkpca, 39, 47
- qkpca, cndkernmatrix-method (qkpca), 39
- qkpca, formula-method (qkpca), 39
- qkpca, matrix-method (qkpca), 39
- qkpca, qkernmatrix-method (qkpca), 39
- qkpca-class, 42
- qkprc-class, 43
- qkspecc, 44, 48, 50
- qkspecc, cndkernmatrix-method (qkspecc), 44
- qkspecc, matrix-method (qkspecc), 44
- qkspecc, qkernmatrix-method (qkspecc), 44
- qkspecc-class, 47
- qkspeclust, 48
- qkspeclust, qkspecc-method (qkspeclust), 48
- qpar (qkprc-class), 43
- qpar, cndkernel-method (cndkernel-class), 7
- qpar, qkernel-method (qkernel-class), 17
- qpar, qkprc-method (qkprc-class), 43
- qpar<- (qkprc-class), 43
- qpar<-, qkprc-method (qkprc-class), 43
- qsammon, 50, 54
- qsammon, cndkernmatrix-method (qsammon), 50
- qsammon, matrix-method (qsammon), 50
- qsammon, qkernmatrix-method (qsammon), 50
- qsammon-class, 53
- qtSNE, 54, 58
- qtSNE, cndkernmatrix-method (qtSNE), 54
- qtSNE, matrix-method (qtSNE), 54
- qtSNE, qkernmatrix-method (qtSNE), 54
- qtSNE-class, 57
- ratibase, 9
- ratibase (bases), 4
- raticnd, 20
- raticnd (cnds), 9
- ratickernel-class (cndkernel-class), 7
- ratiqkernel-class (qkernel-class), 17
- rbfbase, 9
- rbfbase (bases), 4
- rbfcnd, 20
- rbfcnd (cnds), 9
- rbfkernel-class (cndkernel-class), 7
- rbfqkernel-class (qkernel-class), 17
- Residuals (qkIsomap-class), 28
- Residuals, qkIsomap-method (qkIsomap-class), 28
- Residuals, qkMDS-method (qkMDS-class), 37
- Residuals<- (qkIsomap-class), 28
- Residuals<-, qkIsomap-method (qkIsomap-class), 28
- Residuals<-, qkMDS-method (qkMDS-class), 37
- rotated (qkpca-class), 42
- rotated, qkpca-method (qkpca-class), 42
- rotated<- (qkpca-class), 42
- rotated<-, qkpca-method (qkpca-class), 42
- show, cndkernel-method (cndkernel-class), 7
- show, qkernel-method (qkernel-class), 17
- show, qkspecc-method (qkspecc), 44
- studbase, 9
- studbase (bases), 4
- studcnd, 20
- studcnd (cnds), 9
- studkernel-class (cndkernel-class), 7
- studqkernel-class (qkernel-class), 17
- terms (qkprc-class), 43
- terms, qkprc-method (qkprc-class), 43
- terms<- (qkprc-class), 43
- terms<-, qkprc-method (qkprc-class), 43
- wavbase, 9
- wavbase (bases), 4
- wavcnd, 20
- wavcnd (cnds), 9
- wavkernel-class (cndkernel-class), 7
- wavqkernel-class (qkernel-class), 17
- withinss (qkspecc-class), 47
- withinss, qkspecc-method (qkspecc-class), 47



`withinss<-` (qkspecc-class), 47  
`withinss<-`, qkspecc-method  
    (qkspecc-class), 47

`xmatrix` (qkprc-class), 43  
`xmatrix`, qkprc-method (qkprc-class), 43  
`xmatrix<-` (qkprc-class), 43  
`xmatrix<-`, qkprc-method (qkprc-class), 43

`ymatrix` (qkprc-class), 43  
`ymatrix`, qkprc-method (qkprc-class), 43  
`ymatrix<-` (qkprc-class), 43  
`ymatrix<-`, qkprc-method (qkprc-class), 43