

Package: kedd (via r-universe)

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License GPL (>= 2)

Description Smoothing techniques and computing bandwidth selectors of the nth derivative of a probability density for one-dimensional data (described in Arsalane Chouaib Guidoum (2020) <[arXiv:2012.06102](https://arxiv.org/abs/2012.06102)> [stat.CO]).

Title Kernel Estimator and Bandwidth Selection for Density and Its Derivatives

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kedd-package	<i>Kernel Estimator and Bandwidth Selection for Density and Its Derivatives</i>
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Description

Smoothing techniques and computing bandwidth selectors of the r 'th derivative of a probability density for one-dimensional data.

Details

Package: kedd
 Type: Package
 Version: 1.0.4
 Date: 2024-01-27
 License: GPL (≥ 2)

There are four main types of functions in this package:

1. Compute the derivatives and convolutions of a kernel function (1-d).
2. Compute the kernel estimators for density and its derivatives (1-d).
3. Computing the bandwidth selectors (1-d).
4. Displaying kernel estimators.

Main Features

Convolutions and derivatives in kernel function:

In non-parametric statistics, a kernel is a weighting function used in non-parametric estimation techniques. The kernels functions $K(x)$ are used in derivatives of kernel density estimator to estimate $\hat{f}_h^{(r)}(x)$, satisfying the following three requirements:

1. $\int_{\mathbb{R}} K(x)dx = 1$
2. $\int_{\mathbb{R}} xK(x)dx = 0$
3. $\mu_2(K) = \int_{\mathbb{R}} x^2K(x)dx < \infty$

Several types of kernel functions $K(x)$ are commonly used in this package: Gaussian, Epanechnikov, Uniform (rectangular), Triangular, Triweight, Tricube, Biweight (quartic), Cosine.

The function `kernel.fun` for kernel derivative $K^{(r)}(x)$ and `kernel.conv` for kernel convolution $K^{(r)} * K^{(r)}(x)$, where we write formally:

$$K^{(r)}(x) = \frac{d^r}{dx^r} K(x)$$

$$K^{(r)} * K^{(r)}(x) = \int_{-\infty}^{+\infty} K^{(r)}(y)K^{(r)}(x-y)dy$$

for $r = 0, 1, 2, \dots$

Estimators of r'th derivative of a density function:

A *natural estimator* of the r'th derivative of a density function $f(x)$ is:

$$\hat{f}_h^{(r)}(x) = \frac{d^r}{dx^r} \frac{1}{nh} \sum_{i=1}^n K\left(\frac{x-X_i}{h}\right) = \frac{1}{nh^{r+1}} \sum_{i=1}^n K^{(r)}\left(\frac{x-X_i}{h}\right)$$

Here, X_1, X_2, \dots, X_n is an i.i.d. sample of size n from the distribution with density $f(x)$, $K(x)$ is the kernel function which we take to be a symmetric probability density with at least r non zero derivatives when estimating $f^{(r)}(x)$, and h is the bandwidth, this parameter is very important that controls the degree of smoothing applied to the data.

The case ($r = 0$) is the standard kernel density estimator (e.g. Silverman 1986, Wolfgang 1991, Scott 1992, Wand and Jones 1995, Jeffrey 1996, Bowman and Azzalini 1997, Alexandre 2009), properties of such derivative estimators are well known e.g. Sheather and Jones (1991), Jones and Kappenman (1991), Wolfgang (1991). For the case ($r > 0$), is derivative of kernel density estimator (e.g. Bhattacharya 1967, Schuster 1969, Alekseev 1972, Wolfgang et al 1990, Jones 1992, Stoker 1993) and for applications which require the estimation of density derivatives can be found in Singh (1977).

For r'th derivatives of kernel density estimator one-dimensional, the main function is `dkde`. For display, its plot method calls `plot.dkde`, and if to add a plot using `lines.dkde`.

```
R> data(trimodal)
R> dkde(x = trimodal, deriv.order = 0, kernel = "gaussian")
```

```
Data: trimodal (200 obs.);      Kernel: gaussian
Derivative order: 0;    Bandwidth 'h' = 0.1007
      eval.points      est.fx
Min.   :-2.91274   Min.   :0.0000066
1st Qu.: -1.46519   1st Qu.: 0.0669750
Median :-0.01765   Median : 0.1682045
Mean   :-0.01765   Mean    : 0.1723692
3rd Qu.: 1.42989   3rd Qu.: 0.2484626
Max.   : 2.87743   Max.    : 0.4157340
```

```
R> dkde(x = trimodal, deriv.order = 1, kernel = "gaussian")
```

```
Data: trimodal (200 obs.);      Kernel: gaussian
Derivative order: 1;    Bandwidth 'h' = 0.09094
      eval.points      est.fx
Min.   :-2.87358   Min.   :-1.740447
1st Qu.: -1.44562   1st Qu.: -0.343952
Median :-0.01765   Median : 0.009057
Mean   :-0.01765   Mean    : 0.0000000
3rd Qu.: 1.41031   3rd Qu.: 0.415343
Max.   : 2.83828   Max.    : 1.256891
```

Bandwidth selectors:

The most important factor in the r 'th derivative kernel density estimate is a choice of the bandwidth h for one-dimensional observations. Because of its role in controlling both the amount and the direction of smoothing, this choice is particularly important. We present the popular bandwidth selection (for more details see references) methods in this package:

- Optimal Bandwidth (AMISE); with `deriv.order >= 0`, name of this function is `h.amise`. For display, its plot method calls `plot.h.amise`, and to add a plot used `lines.h.amise`.
- Maximum-likelihood cross-validation (MLCV); with `deriv.order = 0`, name of this function is `h.mlcv`. For display, its plot method calls `plot.h.mlcv`, and to add a plot used `lines.h.mlcv`.
- Unbiased cross validation (UCV); with `deriv.order >= 0`, name of this function is `h.ucv`. For display, its plot method calls `plot.h.ucv`, and to add a plot used `lines.h.ucv`.
- Biased cross validation (BCV); with `deriv.order >= 0`, name of this function is `h.bcv`. For display, its plot method calls `plot.h.bcv`, and to add a plot used `lines.h.bcv`.
- Complete cross-validation (CCV); with `deriv.order >= 0`, name of this function is `h.ccv`. For display, its plot method calls `plot.h.ccv`, and to add a plot used `lines.h.ccv`.
- Modified cross-validation (MCV); with `deriv.order >= 0`, name of this function is `h.mcv`. For display, its plot method calls `plot.h.mcv`, and to add a plot used `lines.h.mcv`.
- Trimmed cross-validation (TCV); with `deriv.order >= 0`, name of this function is `h.tcv`. For display, its plot method calls `plot.h.tcv`, and to add a plot used `lines.h.tcv`.

```
R> data(trimodal)
R> h.bcv(x = trimodal, whichbcv = 1, deriv.order = 0, kernel = "gaussian")

Call:      Biased Cross-Validation 1
Derivative order = 0
Data: trimodal (200 obs.);      Kernel: gaussian
Min BCV = 0.004511636; Bandwidth 'h' = 0.4357812

R> h.ccv(x = trimodal, deriv.order = 1, kernel = "gaussian")

Call:      Complete Cross-Validation
Derivative order = 1
Data: trimodal (200 obs.);      Kernel: gaussian
Min CCV = 0.01985078; Bandwidth 'h' = 0.5828336

R> h.tcv(x = trimodal, deriv.order = 2, kernel = "gaussian")

Call:      Trimmed Cross-Validation
Derivative order = 2
Data: trimodal (200 obs.);      Kernel: gaussian
Min TCV = -295.563; Bandwidth 'h' = 0.08908582

R> h.ucv(x = trimodal, deriv.order = 3, kernel = "gaussian")

Call:      Unbiased Cross-Validation
Derivative order = 3
Data: trimodal (200 obs.);      Kernel: gaussian
Min UCV = -63165.18; Bandwidth 'h' = 0.1067236
```

For an overview of this package, see `vignette("kedd")`.

Requirements

R version \geq 2.15.0

Licence

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See Also

ks, **KernSmooth**, **sm**, **np**, **locfit**, **feature**, **GenKern**.

Claw, Bimodal, Kurtotic, Outlier, Trimodal
Datasets

Description

A random sample of size 200 from the claw, bimodal, kurtotic, outlier and trimodal Gaussian density.

Usage

```
data(claw)
data(bimodal)
data(kurtotic)
data(outlier)
data(trimodal)
```

Format

Numeric vector with length 200.

Details

Generate 200 random numbers, distributed according to a normal mixture, using `rnorMix` in package **nor1mix**.

```
## Claw density
claw <- rnorMix(n=200, MW.nm10)
plot(MW.nm10)

## Bimodal density
bimodal <- rnorMix(n=200, MW.nm7)
plot( MW.nm7)

## Kurtotic density
kurtotic <- rnorMix(n=200, MW.nm4)
plot(MW.nm4)

## Outlier density
outlier <- rnorMix(n=200, MW.nm5)
plot( MW.nm5)

## Trimodal density
trimodal <- rnorMix(n=200, MW.nm9)
plot(MW.nm9)
```

Source

Randomly generated a normal mixture with the function `rnorMix` in package **nor1mix**.

References

Martin, M. (2013). **nor1mix**: Normal (1-d) mixture models (S3 classes and methods). R *package version 1.1-4*.

Description

The (S3) generic function `dkde` computes the r 'th derivative of kernel density estimator for one-dimensional data. Its default method does so with the given kernel and bandwidth h for one-dimensional observations.

Usage

```
dkde(x, ...)
## Default S3 method:
dkde(x, y = NULL, deriv.order = 0, h, kernel = c("gaussian",
        "epanechnikov", "uniform", "triangular", "triweight",
        "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	the data from which the estimate is to be computed.
<code>y</code>	the points of the grid at which the density derivative is to be estimated; the defaults are $\tau * h$ outside of $\text{range}(x)$, where $\tau = 4$.
<code>deriv.order</code>	derivative order (scalar).
<code>h</code>	the smoothing bandwidth to be used, can also be a character string giving a rule to choose the bandwidth, see h.bcv . The default h.ucv .
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

A simple estimator for the density derivative can be obtained by taking the derivative of the kernel density estimate. If the kernel $K(x)$ is differentiable r times then the r 'th density derivative estimate can be written as:

$$\hat{f}_h^{(r)}(x) = \frac{1}{nh^{r+1}} \sum_{i=1}^n K^{(r)}\left(\frac{x - X_i}{h}\right)$$

where,

$$K^{(r)}(x) = \frac{d^r}{dx^r} K(x)$$

for $r = 0, 1, 2, \dots$

The following assumptions on the density $f^{(r)}(x)$, the bandwidth h , and the kernel $K(x)$:

1. The $(r + 2)$ derivative $f^{(r+2)}(x)$ is continuous, square integrable and ultimately monotone.
2. $\lim_{n \rightarrow \infty} h = 0$ and $\lim_{n \rightarrow \infty} nh^{2r+1} = \infty$ i.e., as the number of samples n is increased h approaches zero at a rate slower than $1/n^{2r+1}$.

3. $K(x) \geq 0$ and $\int_{\mathbb{R}} K(x) dx = 1$. The kernel function is assumed to be symmetric about the origin i.e., $\int_{\mathbb{R}} x K(x) dx = 0$ for even r and has finite second moment i.e., $\mu_2(K) = \int_{\mathbb{R}} x^2 K(x) dx < \infty$.

Some theoretical properties of the estimator $\hat{f}_h^{(r)}$ have been investigated, among others, by Bhattacharya (1967), Schuster (1969). Let us now turn to the statistical properties of estimator. We are interested in the mean squared error since it combines squared bias and variance.

The **bias** can be written as:

$$E \left[\hat{f}_h^{(r)}(x) \right] - f^{(r)}(x) = \frac{1}{2} h^2 \mu_2(K) f^{(r+2)}(x) + o(h^2)$$

The **variance** of the estimator can be written as:

$$VAR \left[\hat{f}_h^{(r)}(x) \right] = \frac{f(x) R(K^{(r)})}{nh^{2r+1}} + o(1/nh^{2r+1})$$

with, $R(K^{(r)}) = \int_{\mathbb{R}} (K^{(r)}(x))^2 dx$.

The **MSE** (Mean Squared Error) for kernel density derivative estimators can be written as:

$$MSE \left(\hat{f}_h^{(r)}(x), f^{(r)}(x) \right) = \frac{f(x) R(K^{(r)})}{nh^{2r+1}} + \frac{1}{4} h^4 \mu_2^2(K) f^{(r+2)}(x)^2 + o(h^4 + 1/nh^{2r+1})$$

It follows that the MSE-optimal bandwidth for estimating $\hat{f}_h^{(r)} S(x)$, is of order $n^{-1/(2r+5)}$. Therefore, the estimation of $\hat{f}_h^{(1)}(x)$ requires a bandwidth of order $n^{-1/7}$ compared to the optimal $n^{-1/5}$ for estimating $f(x)$ itself. It reveals the increasing difficulty in problems of estimating higher derivatives.

The **MISE** (Mean Integrated Squared Error) can be written as:

$$MISE \left(\hat{f}_h^{(r)}(x), f^{(r)}(x) \right) = AMISE \left(\hat{f}_h^{(r)}(x), f^{(r)}(x) \right) + o(h^4 + 1/nh^{2r+1})$$

where,

$$AMISE \left(\hat{f}_h^{(r)}(x), f^{(r)}(x) \right) = \frac{1}{nh^{2r+1}} R(K^{(r)}) + \frac{1}{4} h^4 \mu_2^2(K) R(f^{(r+2)})$$

with: $R(f^{(r)}(x)) = \int_{\mathbb{R}} (f^{(r)}(x))^2 dx$.

The performance of kernel is measured by **MISE** or **AMISE** (Asymptotic MISE).

If the bandwidth h is missing from `dkde`, then the default bandwidth is `h.ucv(x, deriv.order, kernel)` (Unbiased cross-validation, see `h.ucv`).

For more details see references.

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use.
deriv.order	the derivative order to use.
h	the bandwidth value to use.
eval.points	the coordinates of the points where the density derivative is estimated.
est.fx	the estimated density derivative values.

Note

This function are available in other packages such as **KernSmooth**, **sm**, **np**, **GenKern** and **locfit** if `deriv.order=0`, and in **ks** package for Gaussian kernel only if $0 \leq \text{deriv.order} \leq 10$.

Author(s)

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- Wolfgang, H. (1991). *Smoothing Techniques, With Implementation in S*. Springer-Verlag, New York.

See Also

`plot.dkde`, see `density` in package "stats" if `deriv.order = 0`, and `kdde` in package **ks**.

Examples

```
## EXAMPLE 1: Simple example of a Gaussian density derivative

x <- rnorm(100)
dkde(x,deriv.order=0) ## KDE of f
dkde(x,deriv.order=1) ## KDDE of d/dx f
dkde(x,deriv.order=2) ## KDDE of d^2/x^2 f
dkde(x,deriv.order=3) ## KDDE of d^3/x^3 f
oldpar <- par(no.readonly = TRUE)
dev.new()
par(mfrow=c(2,2))
plot(dkde(x,deriv.order=0))
plot(dkde(x,deriv.order=1))
plot(dkde(x,deriv.order=2))
plot(dkde(x,deriv.order=3))
par(oldpar)

## EXAMPLE 2: Bimodal Gaussian density derivative
## show the kernels in the dkde parametrization

fx <- function(x) 0.5 * dnorm(x,-1.5,0.5) + 0.5 * dnorm(x,1.5,0.5)
fx1 <- function(x) 0.5 *(-4*x-6)* dnorm(x,-1.5,0.5) + 0.5 *(-4*x+6) *
      dnorm(x,1.5,0.5)

## 'h = 0.3' ; 'Derivative order = 0'

kernels <- eval(formals(dkde.default)$kernel)
dev.new()
plot(dkde(bimodal,h=0.3),sub=paste("Derivative order = 0",",",
  "Bandwidth =0.3 "),ylim=c(0,0.5), main = "Bimodal Gaussian Density")
for(i in 2:length(kernels))
  lines(dkde(bimodal, h = 0.3, kernel = kernels[i]), col = i)
curve(fx,add=TRUE,lty=8)
legend("topright", legend = c(TRUE,kernels), col = c("black",seq(kernels)),
  lty = c(8,rep(1,length(kernels))),cex=0.7, inset = .015)

## 'h = 0.6' ; 'Derivative order = 1'

kernels <- eval(formals(dkde.default)$kernel)[-3]
dev.new()
plot(dkde(bimodal,deriv.order=1,h=0.6),main = "Bimodal Gaussian Density Derivative",sub=paste
  ("Derivative order = 1",",", "Bandwidth =0.6"),ylim=c(-0.6,0.6))
for(i in 2:length(kernels))
  lines(dkde(bimodal,deriv.order=1, h = 0.6, kernel = kernels[i]), col = i)
curve(fx1,add=TRUE,lty=8)
legend("topright", legend = c(TRUE,kernels), col = c("black",seq(kernels)),
  lty = c(8,rep(1,length(kernels))),cex=0.7, inset = .015)
```

h.amise

*AMISE for Optimal Bandwidth Selectors***Description**

The (S3) generic function `h.amise` evaluates the asymptotic mean integrated squared error **AMISE** for optimal smoothing parameters h of r 'th derivative of kernel density estimator one-dimensional.

Usage

```
h.amise(x, ...)
## Default S3 method:
h.amise(x, deriv.order = 0, lower = 0.1 * hos, upper = 2 * hos,
        tol = 0.1 * lower, kernel = c("gaussian", "epanechnikov", "triweight",
        "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	vector of data values.
<code>deriv.order</code>	derivative order (scalar).
<code>lower, upper</code>	range over which to minimize. The default is almost always satisfactory. <code>hos</code> (Over-smoothing) is calculated internally from an kernel, see details.
<code>tol</code>	the convergence tolerance for <code>optimize</code> .
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

`h.amise` asymptotic mean integrated squared error implements for choosing the optimal bandwidth h of a r 'th derivative kernel density estimator.

We Consider the following AMISE version of the r 'th derivative of f the r 'th derivative of the kernel estimate (see Scott 1992, pp 131):

$$AMISE(h; r) = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{1}{4}h^4\mu_2^2(K)R(f^{(r+2)})$$

The optimal bandwidth minimizing this function is:

$$h_{(r)}^* = \left[\frac{(2r+1)R(K^{(r)})}{\mu_2^2(K)R(f^{(r+2)})} \right]^{1/(2r+5)} n^{-1/(2r+5)}$$

whereof

$$\inf_{h>0} AMISE(h; r) = \frac{2r+5}{4} R(K^{(r)})^{\frac{4}{(2r+5)}} \left[\frac{\mu_2^2(K)R(f^{(r+2)})}{2r+1} \right]^{\frac{2r+1}{2r+5}} n^{-\frac{4}{2r+5}}$$

which is the smallest possible AMISE for estimation of $f^{(r)}(x)$ using the kernel $K(x)$, where $R(K^{(r)}) = \int_R K^{(r)}(x)^2 dx$ and $\mu_2(K) = \int_R x^2 K(x) dx$.

The range over which to minimize is hos Oversmoothing bandwidth, the default is almost always satisfactory. See George and Scott (1985), George (1990), Scott (1992, pp 165), Wand and Jones (1995, pp 61).

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use
deriv.order	the derivative order to use.
h	value of bandwidth parameter.
amise	the AMISE value.

Author(s)

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References

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- Wand, M. P. and Jones, M. C. (1995). *Kernel Smoothing*. Chapman and Hall, London.

See Also

[plot.h.amise](#), see [nmise](#) in package [sm](#) this function evaluates the mean integrated squared error of a density estimate (`deriv.order = 0`) which is constructed from data which follow a normal distribution.

Examples

```
## Derivative order = 0
h.amise(kurtotic,deriv.order = 0)
```

```
## Derivative order = 1
h.amise(kurtotic,deriv.order = 1)
```

h.bcv

*Biased Cross-Validation for Bandwidth Selection***Description**

The (S3) generic function `h.bcv` computes the biased cross-validation bandwidth selector of r 'th derivative of kernel density estimator one-dimensional.

Usage

```
h.bcv(x, ...)
## Default S3 method:
h.bcv(x, whichbcv = 1, deriv.order = 0, lower = 0.1 * hos, upper = 2 * hos,
      tol = 0.1 * lower, kernel = c("gaussian", "epanechnikov",
      "triweight", "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	vector of data values.
<code>whichbcv</code>	method selected, 1 = BCV1 or 2 = BCV2, see details.
<code>deriv.order</code>	derivative order (scalar).
<code>lower, upper</code>	range over which to minimize. The default is almost always satisfactory. <code>hos</code> (Over-smoothing) is calculated internally from an kernel, see details.
<code>tol</code>	the convergence tolerance for <code>optimize</code> .
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

`h.bcv` biased cross-validation implements for choosing the bandwidth h of a r 'th derivative kernel density estimator. if `whichbcv = 1` then **BCV1** is selected (Scott and George 1987), and if `whichbcv = 2` used **BCV2** (Jones and Kappenman 1991).

Scott and George (1987) suggest a method which has as its immediate target the **AMISE** (e.g. Silverman 1986, section 3.3). We denote $\hat{\theta}_r(h)$ and $\bar{\theta}_r(h)$ (Peter and Marron 1987, Jones and Kappenman 1991) by:

$$\hat{\theta}_r(h) = \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(r)} * K^{(r)} \left(\frac{X_j - X_i}{h} \right)$$

and

$$\bar{\theta}_r(h) = \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(2r)}\left(\frac{X_j - X_i}{h}\right)$$

Scott and George (1987) proposed using $\hat{\theta}_r(h)$ to estimate $f^{(r)}(x)$. Thus, $\hat{h}_{BCV1}^{(r)}$, say, is the h that minimises:

$$BCV1(h; r) = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{1}{4}\mu_2^2(K)h^4\hat{\theta}_{r+2}(h)$$

and we define $\hat{h}_{BCV2}^{(r)}$ as the minimiser of (Jones and Kappenman 1991):

$$BCV2(h; r) = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{1}{4}\mu_2^2(K)h^4\bar{\theta}_{r+2}(h)$$

where $K^{(r)} * K^{(r)}(x)$ is the convolution of the r 'th derivative kernel function $K^{(r)}(x)$ (see [kernel.conv](#) and [kernel.fun](#)); $R(K^{(r)}) = \int_{\mathbb{R}} K^{(r)}(x)^2 dx$ and $\mu_2(K) = \int_{\mathbb{R}} x^2 K(x) dx$.

The range over which to minimize is hos Oversmoothing bandwidth, the default is almost always satisfactory. See George and Scott (1985), George (1990), Scott (1992, pp 165), Wand and Jones (1995, pp 61).

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use
deriv.order	the derivative order to use.
whichbcv	method selected.
h	value of bandwidth parameter.
min.bcv	the minimal BCV value.

Author(s)

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References

- Jones, M. C. and Kappenman, R. F. (1991). On a class of kernel density estimate bandwidth selectors. *Scandinavian Journal of Statistics*, **19**, 337–349.
- Jones, M. C., Marron, J. S. and Sheather, S. J. (1996). A brief survey of bandwidth selection for density estimation. *Journal of the American Statistical Association*, **91**, 401–407.
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- Wand, M. P. and Jones, M. C. (1995). *Kernel Smoothing*. Chapman and Hall, London.
- Wolfgang, H. (1991). *Smoothing Techniques, With Implementation in S*. Springer-Verlag, New York.

See Also

`plot.h.bcv`, see `bw.bcv` in package "stats" and `bcv` in package **MASS** for Gaussian kernel only if `deriv.order = 0`, `Hbcv` for bivariate data in package **ks** for Gaussian kernel only if `deriv.order = 0`, `kdeb` in package **locfit** if `deriv.order = 0`.

Examples

```
## EXAMPLE 1:

x <- rnorm(100)
h.bcv(x, whichbcv = 1, deriv.order = 0)
h.bcv(x, whichbcv = 2, deriv.order = 0)

## EXAMPLE 2:

## Derivative order = 0

h.bcv(kurtotic, deriv.order = 0)

## Derivative order = 1

h.bcv(kurtotic, deriv.order = 1)
```

h.ccv

Complete Cross-Validation for Bandwidth Selection

Description

The (S3) generic function `h.ccv` computes the complete cross-validation bandwidth selector of r 'th derivative of kernel density estimator one-dimensional.

Usage

```
h.ccv(x, ...)
## Default S3 method:
h.ccv(x, deriv.order = 0, lower = 0.1 * hos, upper = hos,
      tol = 0.1 * lower, kernel = c("gaussian", "triweight",
      "tricube", "biweight", "cosine"), ...)
```

Arguments

x	vector of data values.
deriv.order	derivative order (scalar).
lower, upper	range over which to minimize. The default is almost always satisfactory. hos (Over-smoothing) is calculated internally from an kernel, see details.
tol	the convergence tolerance for <code>optimize</code> .
kernel	a character string giving the smoothing kernel to be used, with default "gaussian".
...	further arguments for (non-default) methods.

Details

h.ccv complete cross-validation implements for choosing the bandwidth h of a r 'th derivative kernel density estimator.

Jones and Kappenman (1991) proposed a so-called complete cross-validation (CCV) in kernel density estimator. This method can be extended to the estimation of derivative of the density, basing our estimate of integrated squared density derivative (Peter and Marron 1987) on the $\bar{\theta}_r(h)$'s, we get the following, start from $R(\hat{f}_h^{(r)}) - \bar{\theta}_r(h)$ as an estimate of MISE. Thus, $\hat{h}_{CCV}^{(r)}$, say, is the h that minimises:

$$CCV(h; r) = R(\hat{f}_h^{(r)}) - \bar{\theta}_r(h) + \frac{1}{2}\mu_2(K)h^2\bar{\theta}_{r+1}(h) + \frac{1}{24}(6\mu_2^2(K) - \delta(K))h^4\bar{\theta}_{r+2}(h)$$

with

$$R(\hat{f}_h^{(r)}) = \int (\hat{f}_h^{(r)}(x))^2 dx = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(r)} * K^{(r)} \left(\frac{X_j - X_i}{h} \right)$$

and

$$\bar{\theta}_r(h) = \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(2r)} \left(\frac{X_j - X_i}{h} \right)$$

and $K^{(r)} * K^{(r)}(x)$ is the convolution of the r 'th derivative kernel function $K^{(r)}(x)$ (see [kernel.conv](#) and [kernel.fun](#)); $R(K^{(r)}) = \int_{\mathbb{R}} K^{(r)}(x)^2 dx$ and $\mu_2(K) = \int_{\mathbb{R}} x^2 K(x) dx$, $\delta(K) = \int_{\mathbb{R}} x^4 K(x) dx$.

The range over which to minimize is hos Oversmoothing bandwidth, the default is almost always satisfactory. See George and Scott (1985), George (1990), Scott (1992, pp 165), Wand and Jones (1995, pp 61).

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use
deriv.order	the derivative order to use.
h	value of bandwidth parameter.
min.ccv	the minimal CCV value.

Author(s)

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References

Jones, M. C. and Kappenman, R. F. (1991). On a class of kernel density estimate bandwidth selectors. *Scandinavian Journal of Statistics*, **19**, 337–349.

Peter, H. and Marron, J.S. (1987). Estimation of integrated squared density derivatives. *Statistics and Probability Letters*, **6**, 109–115.

See Also

[plot.h.ccv.](#)

Examples

```
## Derivative order = 0
h.ccv(kurtotic,deriv.order = 0)

## Derivative order = 1
h.ccv(kurtotic,deriv.order = 1)
```

h.mcv

Modified Cross-Validation for Bandwidth Selection

Description

The (S3) generic function `h.mcv` computes the modified cross-validation bandwidth selector of r 'th derivative of kernel density estimator one-dimensional.

Usage

```
h.mcv(x, ...)
## Default S3 method:
h.mcv(x, deriv.order = 0, lower = 0.1 * hos, upper = 2 * hos,
      tol = 0.1 * lower, kernel = c("gaussian", "epanechnikov", "triweight",
      "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	vector of data values.
<code>deriv.order</code>	derivative order (scalar).
<code>lower, upper</code>	range over which to minimize. The default is almost always satisfactory. <code>hos</code> (Over-smoothing) is calculated internally from an kernel, see details.
<code>tol</code>	the convergence tolerance for optimize .

kernel a character string giving the smoothing kernel to be used, with default "gaussian".
 ... further arguments for (non-default) methods.

Details

h.mcv modified cross-validation implements for choosing the bandwidth h of a r 'th derivative kernel density estimator.

Stute (1992) proposed a so-called modified cross-validation (MCV) in kernel density estimator. This method can be extended to the estimation of derivative of a density, the essential idea based on approximated the problematic term by the aid of the Hajek projection (see Stute 1992). The minimization criterion is defined by:

$$MCV(h; r) = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n \varphi^{(r)}\left(\frac{X_j - X_i}{h}\right)$$

whit

$$\varphi^{(r)}(c) = \left(K^{(r)} * K^{(r)} - K^{(2r)} - \frac{\mu_2(K)}{2} K^{(2r+2)} \right) (c)$$

and $K^{(r)} * K^{(r)}(x)$ is the convolution of the r 'th derivative kernel function $K^{(r)}(x)$ (see [kernel.conv](#) and [kernel.fun](#)); $R(K^{(r)}) = \int_{\mathbb{R}} K^{(r)}(x)^2 dx$ and $\mu_2(K) = \int_{\mathbb{R}} x^2 K(x) dx$.

The range over which to minimize is hos Oversmoothing bandwidth, the default is almost always satisfactory. See George and Scott (1985), George (1990), Scott (1992, pp 165), Wand and Jones (1995, pp 61).

Value

x data points - same as input.
 data.name the deparsed name of the x argument.
 n the sample size after elimination of missing values.
 kernel name of kernel to use
 deriv.order the derivative order to use.
 h value of bandwidth parameter.
 min.mcv the minimal MCV value.

Author(s)

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References

Heidenreich, N. B., Schindler, A. and Sperlich, S. (2013). Bandwidth selection for kernel density estimation: a review of fully automatic selectors. *Advances in Statistical Analysis*.
 Stute, W. (1992). Modified cross validation in density estimation. *Journal of Statistical Planning and Inference*, **30**, 293–305.

See Also[plot.h.mcv.](#)**Examples**

```
## Derivative order = 0

h.mcv(kurtotic,deriv.order = 0)

## Derivative order = 1

h.mcv(kurtotic,deriv.order = 1)
```

h.mlcv

Maximum-Likelihood Cross-validation for Bandwidth Selection

Description

The (S3) generic function `h.mlcv` computes the maximum likelihood cross-validation (Kullback-Leibler information) bandwidth selector of a one-dimensional kernel density estimate.

Usage

```
h.mlcv(x, ...)
## Default S3 method:
h.mlcv(x, lower = 0.1, upper = 5, tol = 0.1 * lower,
       kernel = c("gaussian", "epanechnikov", "uniform", "triangular",
                  "triweight", "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	vector of data values.
<code>lower, upper</code>	range over which to maximize. The default is almost always satisfactory.
<code>tol</code>	the convergence tolerance for optimize .
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

`h.mlcv` maximum-likelihood cross-validation implements for choosing the optimal bandwidth h of kernel density estimator.

This method was proposed by Habbema, Hermans, and Van den Broeck (1971) and by Duin (1976). The maximum-likelihood cross-validation (MLCV) function is defined by:

$$MLCV(h) = n^{-1} \sum_{i=1}^n \log [\hat{f}_{h,i}(x)]$$

the estimate $\hat{f}_{h,i}(x)$ on the subset $\{X_j\}_{j \neq i}$ denoting the leave-one-out estimator, can be written:

$$\hat{f}_{h,i}(X_i) = \frac{1}{(n-1)h} \sum_{j \neq i} K\left(\frac{X_j - X_i}{h}\right)$$

Define that h_{mlcv} as good which approaches the finite maximum of $MLCV(h)$:

$$h_{mlcv} = \arg \max_h MLCV(h) = \arg \max_h \left(n^{-1} \sum_{i=1}^n \log \left[\sum_{j \neq i} K\left(\frac{X_j - X_i}{h}\right) \right] - \log[(n-1)h] \right)$$

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use
h	value of bandwidth parameter.
mlcv	the maximal likelihood CV value.

Author(s)

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References

Habbema, J. D. F., Hermans, J., and Van den Broek, K. (1974) A stepwise discrimination analysis program using density estimation. *Compstat 1974: Proceedings in Computational Statistics*. Physica Verlag, Vienna.

Duin, R. P. W. (1976). On the choice of smoothing parameters of Parzen estimators of probability density functions. *IEEE Transactions on Computers*, **C-25**, 1175–1179.

See Also

[plot.h.mlcv](#), see [lcv](#) in package [locfit](#).

Examples

```
h.mlcv(bimodal)
h.mlcv(bimodal, kernel = "epanechnikov")
```

Description

The (S3) generic function `h.tcv` computes the trimmed cross-validation bandwidth selector of r 'th derivative of kernel density estimator one-dimensional.

Usage

```
h.tcv(x, ...)
## Default S3 method:
h.tcv(x, deriv.order = 0, lower = 0.1 * hos, upper = 2 * hos,
      tol = 0.1 * lower, kernel = c("gaussian", "epanechnikov", "uniform",
      "triangular", "triweight", "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	vector of data values.
<code>deriv.order</code>	derivative order (scalar).
<code>lower, upper</code>	range over which to minimize. The default is almost always satisfactory. <code>hos</code> (Over-smoothing) is calculated internally from an kernel, see details.
<code>tol</code>	the convergence tolerance for <code>optimize</code> .
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

`h.tcv` trimmed cross-validation implements for choosing the bandwidth h of a r 'th derivative kernel density estimator.

Feluch and Koronacki (1992) proposed a so-called trimmed cross-validation (TCV) in kernel density estimator, a simple modification of the unbiased (least-squares) cross-validation criterion. We consider the following "trimmed" version of "unbiased", to be minimized with respect to h :

$$\int \left(\hat{f}_h^{(r)}(x) \right)^2 - 2 \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(2r)} \left(\frac{X_j - X_i}{h} \right) \chi(|X_i - X_j| > c_n)$$

where $\chi(\cdot)$ denotes the indicator function and c_n is a sequence of positive constants, $c_n/h^{2r+1} \rightarrow 0$ as $n \rightarrow \infty$, and

$$\int \left(\hat{f}_h^{(r)}(x) \right)^2 = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(r)} * K^{(r)} \left(\frac{X_j - X_i}{h} \right)$$

the trimmed cross-validation function is defined by:

$$TCV(h; r) = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n \varphi^{(r)}\left(\frac{X_j - X_i}{h}\right)$$

whit

$$\varphi^{(r)}(c) = \left(K^{(r)} * K^{(r)} - 2K^{(2r)} \chi(|c| > c_n/h^{2r+1}) \right) (c)$$

here we take $c_n = 1/n$, for assure the convergence. Where $K^{(r)} * K^{(r)}(x)$ is the convolution of the r 'th derivative kernel function $K^{(r)}(x)$ (see [kernel.conv](#) and [kernel.fun](#)).

The range over which to minimize is hos Oversmoothing bandwidth, the default is almost always satisfactory. See George and Scott (1985), George (1990), Scott (1992, pp 165), Wand and Jones (1995, pp 61).

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use
deriv.order	the derivative order to use.
h	value of bandwidth parameter.
min.tcv	the minimal TCV value.

Author(s)

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References

Feluch, W. and Koronacki, J. (1992). A note on modified cross-validation in density estimation. *Computational Statistics and Data Analysis*, **13**, 143–151.

See Also

[plot.h.tcv](#).

Examples

```
## Derivative order = 0
h.tcv(kurtotic,deriv.order = 0)

## Derivative order = 1
h.tcv(kurtotic,deriv.order = 1)
```


Description

The (S3) generic function `h.ucv` computes the unbiased (least-squares) cross-validation bandwidth selector of r 'th derivative of kernel density estimator one-dimensional.

Usage

```
h.ucv(x, ...)
## Default S3 method:
h.ucv(x, deriv.order = 0, lower = 0.1 * hos, upper = 2 * hos,
      tol = 0.1 * lower, kernel = c("gaussian", "epanechnikov", "uniform",
      "triangular", "triweight", "tricube", "biweight", "cosine"), ...)
```

Arguments

<code>x</code>	vector of data values.
<code>deriv.order</code>	derivative order (scalar).
<code>lower, upper</code>	range over which to minimize. The default is almost always satisfactory. <code>hos</code> (Over-smoothing) is calculated internally from an kernel, see details.
<code>tol</code>	the convergence tolerance for optimize .
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

`h.ucv` unbiased (least-squares) cross-validation implements for choosing the bandwidth h of a r 'th derivative kernel density estimator.

Rudemo (1982) and Bowman (1984) proposed a so-called unbiased (least-squares) cross-validation (UCV) in kernel density estimator. An adaptation of unbiased cross-validation is proposed by Wolfgang et al. (1990) for bandwidth choice in the r 'th derivative of kernel density estimator. The essential idea of this methods, for the estimation of $f^{(r)}(x)$ (r is derivative order), is to use the bandwidth h which minimizes the function:

$$UCV(h; r) = \int \left(\hat{f}_h^{(r)}(x) \right)^2 - 2n^{-1}(-1)^r \sum_{i=1}^n \hat{f}_{h,i}^{(2r)}(X_i)$$

The bandwidth minimizing this function is:

$$\hat{h}_{ucv}^{(r)} = \arg \min_{h^{(r)}} UCV(h; r)$$

for $r = 0, 1, 2, \dots$
 where

$$\int \left(\hat{f}_h^{(r)}(x) \right)^2 = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n K^{(r)} * K^{(r)} \left(\frac{X_j - X_i}{h} \right)$$

and $K^{(r)} * K^{(r)}(x)$ is the convolution of the r 'th derivative kernel function $K^{(r)}(x)$ (see [kernel.conv](#) and [kernel.fun](#)).

The estimate $\hat{f}_{h,i}^{(2r)}(x)$ on the subset $\{X_j\}_{j \neq i}$ denoting the leave-one-out estimator, can be written:

$$\hat{f}_{h,i}^{(2r)}(X_i) = \frac{1}{(n-1)h^{2r+1}} \sum_{j \neq i} K^{(2r)} \left(\frac{X_j - X_i}{h} \right)$$

The function $UCV(h; r)$ is unbiased cross-validation in the sense that $E[UCV] = MISE[\hat{f}_h^{(r)}(x)] - R(f^{(r)}(x))$ (see, Scott and George 1987). Can be simplified to give the computationally:

$$UCV(h; r) = \frac{R(K^{(r)})}{nh^{2r+1}} + \frac{(-1)^r}{n(n-1)h^{2r+1}} \sum_{i=1}^n \sum_{j=1; j \neq i}^n \left(K^{(r)} * K^{(r)} - 2K^{(2r)} \right) \left(\frac{X_j - X_i}{h} \right)$$

where $R(K^{(r)}) = \int_R K^{(r)}(x)^2 dx$.

The range over which to minimize is hos Oversmoothing bandwidth, the default is almost always satisfactory. See George and Scott (1985), George (1990), Scott (1992, pp 165), Wand and Jones (1995, pp 61).

Value

x	data points - same as input.
data.name	the deparsed name of the x argument.
n	the sample size after elimination of missing values.
kernel	name of kernel to use
deriv.order	the derivative order to use.
h	value of bandwidth parameter.
min.ucv	the minimal UCV value.

Author(s)

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References

- Bowman, A. (1984). An alternative method of cross-validation for the smoothing of kernel density estimates. *Biometrika*, **71**, 353–360.
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See Also

`plot.h.ucv`, see `bw.ucv` in package "stats" and `ucv` in package **MASS** for Gaussian kernel only if `deriv.order = 0`, `hlscv` in package **ks** for Gaussian kernel only if $0 \leq \text{deriv.order} \leq 5$, `kdeb` in package **locfit** if `deriv.order = 0`.

Examples

```
## Derivative order = 0

h.ucv(kurtotic,deriv.order = 0)

## Derivative order = 1

h.ucv(kurtotic,deriv.order = 1)
```

kernel.conv

Convolutions of r'th Derivative for Kernel Function

Description

The (S3) generic function `kernel.conv` computes the convolution of r 'th derivative for kernel function.

Usage

```
kernel.conv(x, ...)
## Default S3 method:
kernel.conv(x = NULL, deriv.order = 0, kernel = c("gaussian", "epanechnikov",
"uniform", "triangular", "triweight", "tricube",
"biweight", "cosine", "silverman"), ...)
```

Arguments

`x` points at which the convolution of kernel derivative is to be evaluated.

`deriv.order` derivative order (scalar).

`kernel` a character string giving the smoothing kernel to be used, with default "gaussian".

`...` further arguments for (non-default) methods.

Details

The convolution of r 'th derivative for kernel function is written $K^{(r)} * K^{(r)}$. It is defined as the integral of the product of the derivative for kernel. As such, it is a particular kind of integral transform:

$$K^{(r)} * K^{(r)}(x) = \int_{-\infty}^{+\infty} K^{(r)}(y)K^{(r)}(x - y)dy$$

where:

$$K^{(r)}(x) = \frac{d^r}{dx^r}K(x)$$

for $r = 0, 1, 2, \dots$

Value

`kernel` name of kernel to use.

`deriv.order` the derivative order to use.

`x` the n coordinates of the points where the convolution of kernel derivative is evaluated.

`kx` the convolution of kernel derivative values.

Author(s)

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References

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Wolfgang, H. (1991). *Smoothing Techniques, With Implementation in S*. Springer-Verlag, New York.

See Also

[plot.kernel.conv](#), [kernapply](#) in package "stats" for computes the convolution between an input sequence, and [convolve](#) use the Fast Fourier Transform ([fft](#)) to compute the several kinds of convolutions of two sequences.

Examples

```

kernels <- eval(formals(kernel.conv.default)$kernel)
kernels

## gaussian
kernel.conv(x = 0, kernel=kernels[1], deriv.order=0)
kernel.conv(x = 0, kernel=kernels[1], deriv.order=1)

## silverman
kernel.conv(x = 0, kernel=kernels[9], deriv.order=0)
kernel.conv(x = 0, kernel=kernels[9], deriv.order=1)

```

kernel.fun

Derivatives of Kernel Function

Description

The (S3) generic function `kernel.fun` computes the r 'th derivative for kernel density.

Usage

```

kernel.fun(x, ...)
## Default S3 method:
kernel.fun(x = NULL, deriv.order = 0, kernel = c("gaussian", "epanechnikov",
"uniform", "triangular", "triweight", "tricube",
"biweight", "cosine", "silverman"), ...)

```

Arguments

<code>x</code>	points at which the derivative of kernel function is to be evaluated.
<code>deriv.order</code>	derivative order (scalar).
<code>kernel</code>	a character string giving the smoothing kernel to be used, with default "gaussian".
<code>...</code>	further arguments for (non-default) methods.

Details

We give a short survey of some kernels functions $K(x; r)$; where r is derivative order,

- Gaussian: $K(x; \infty) = \frac{1}{\sqrt{2\pi}} \exp\left(-\frac{x^2}{2}\right) 1_{]-\infty, +\infty[}$
- Epanechnikov: $K(x; 2) = \frac{3}{4}(1 - x^2) 1_{(|x| \leq 1)}$
- uniform (rectangular): $K(x; 0) = \frac{1}{2} 1_{(|x| \leq 1)}$
- triangular: $K(x; 1) = (1 - |x|) 1_{(|x| \leq 1)}$
- triweight: $K(x; 6) = \frac{35}{32}(1 - x^2)^3 1_{(|x| \leq 1)}$
- tricube: $K(x; 9) = \frac{70}{81}(1 - |x|^3)^3 1_{(|x| \leq 1)}$
- biweight: $K(x; 4) = \frac{15}{16}(1 - x^2)^2 1_{(|x| \leq 1)}$
- cosine: $K(x; \infty) = \frac{\pi}{4} \cos\left(\frac{\pi}{2}x\right) 1_{(|x| \leq 1)}$
- Silverman: $K(x; r \bmod 8) = \frac{1}{2} \exp\left(-\frac{|x|}{\sqrt{2}}\right) \sin\left(\frac{|x|}{\sqrt{2}} + \frac{\pi}{4}\right) 1_{]-\infty, +\infty[}$

The r 'th derivative for kernel function $K(x)$ is written:

$$K^{(r)}(x) = \frac{d^r}{dx^r} K(x)$$

for $r = 0, 1, 2, \dots$

The r 'th derivative of the **Gaussian kernel** $K(x)$ is given by:

$$K^{(r)}(x) = (-1)^r H_r(x) K(x)$$

where $H_r(x)$ is the r 'th **Hermite polynomial**. This polynomials are set of orthogonal polynomials, for more details see, [hermite.h.polynomials](#) in package **orthopolynom**.

Value

kernel	name of kernel to use.
deriv.order	the derivative order to use.
x	the n coordinates of the points where the derivative of kernel function is evaluated.
kx	the kernel derivative values.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

References

- Jones, M. C. (1992). Differences and derivatives in kernel estimation. *Metrika*, **39**, 335–340.
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- Silverman, B. W. (1986). *Density Estimation for Statistics and Data Analysis*. Chapman & Hall/CRC. London.

See Also

[plot.kernel.fun](#), [deriv](#) and [D](#) in package "stats" for symbolic and algorithmic derivatives of simple expressions.

Examples

```

kernels <- eval(formals(kernel.fun.default)$kernel)
kernels

## gaussian
kernel.fun(x = 0, kernel=kernels[1], deriv.order=0)
kernel.fun(x = 0, kernel=kernels[1], deriv.order=1)

## silverman
kernel.fun(x = 0, kernel=kernels[9], deriv.order=0)
kernel.fun(x = 0, kernel=kernels[9], deriv.order=1)

```

plot.dkde

Plot for Kernel Density Derivative Estimate

Description

The [plot.dkde](#) function loops through calls to the [dkde](#) function. Plot for kernel density derivative estimate for 1-dimensional data.

Usage

```

## S3 method for class 'dkde'
plot(x, fx = NULL, ...)
## S3 method for class 'dkde'
lines(x, ...)

```

Arguments

x	object of class dkde (output from dkde).
fx	add to graphics the true density derivative (class :function), to compare it by the density derivative to estimate.
...	other graphics parameters, see par in package "graphics".

Details

The 1-d plot is a standard plot of a 1-d curve. If `!is.null(fx)` then a true density derivative is added.

Value

Plot of 1-d kernel density derivative estimates are sent to graphics window.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

See Also

[dkde](#), [plot.density](#) in package "stats" if `deriv.order = 0`.

Examples

```
plot(dkde(kurtotic,deriv.order=0,kernel="gaussian"),sub="")
lines(dkde(kurtotic,deriv.order=0,kernel="biweight"),col="red")
```

plot.h.amise

Plot for Asymptotic Mean Integrated Squared Error

Description

The [plot.h.amise](#) function loops through calls to the [h.amise](#) function. Plot for asymptotic mean integrated squared error function for 1-dimensional data.

Usage

```
## S3 method for class 'h.amise'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.amise'
lines(x,seq.bws=NULL, ...)
```

Arguments

<code>x</code>	object of class <code>h.amise</code> (output from h.amise).
<code>seq.bws</code>	the sequence of bandwidths in which to compute the AMISE function. By default, the procedure defines a sequence of 50 points, from $0.15 \cdot h_{os}$ to $2 \cdot h_{os}$ (Over-smoothing).
<code>...</code>	other graphics parameters, see par in package "graphics".

Value

Plot of 1-d AMISE function are sent to graphics window.

<code>kernel</code>	name of kernel to use.
<code>deriv.order</code>	the derivative order to use.
<code>seq.bws</code>	the sequence of bandwidths.
<code>amise</code>	the values of the AMISE function in the bandwidths grid.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

See Also

[h.amise.](#)

Examples

```
plot(h.amise(bimodal,deriv.order=0))
```

plot.h.bcv

Plot for Biased Cross-Validation

Description

The `plot.h.bcv` function loops through calls to the `h.bcv` function. Plot for biased cross-validation function for 1-dimensional data.

Usage

```
## S3 method for class 'h.bcv'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.bcv'
lines(x,seq.bws=NULL, ...)
```

Arguments

<code>x</code>	object of class <code>h.bcv</code> (output from <code>h.bcv</code>).
<code>seq.bws</code>	the sequence of bandwidths in which to compute the biased cross-validation function. By default, the procedure defines a sequence of 50 points, from $0.15 \times \text{hos}$ to $2 \times \text{hos}$ (Over-smoothing).
<code>...</code>	other graphics parameters, see <code>par</code> in package "graphics".

Value

Plot of 1-d biased cross-validation function are sent to graphics window.

<code>kernel</code>	name of kernel to use.
<code>deriv.order</code>	the derivative order to use.
<code>seq.bws</code>	the sequence of bandwidths.
<code>bcv</code>	the values of the biased cross-validation function in the bandwidths grid.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

See Also[h.bcv.](#)**Examples**

```
## EXAMPLE 1:

plot(h.bcv(trimodal, whichbcv = 1, deriv.order = 0),main="",sub="")
lines(h.bcv(trimodal, whichbcv = 2, deriv.order = 0),col="red")
legend("topright", c("BCV1", "BCV2"),lty=1,col=c("black", "red"),inset = .015)

## EXAMPLE 2:

plot(h.bcv(trimodal, whichbcv = 1, deriv.order = 1),main="",sub="")
lines(h.bcv(trimodal, whichbcv = 2, deriv.order = 1),col="red")
legend("topright", c("BCV1", "BCV2"),lty=1,col=c("black", "red"),inset = .015)
```

plot.h.ccv

*Plot for Complete Cross-Validation***Description**

The [plot.h.ccv](#) function loops through calls to the [h.ccv](#) function. Plot for complete cross-validation function for 1-dimensional data.

Usage

```
## S3 method for class 'h.ccv'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.ccv'
lines(x, seq.bws=NULL, ...)
```

Arguments

x	object of class <code>h.ccv</code> (output from h.ccv).
seq.bws	the sequence of bandwidths in which to compute the complete cross-validation function. By default, the procedure defines a sequence of 50 points, from $0.15 \cdot h_{os}$ to $2 \cdot h_{os}$ (Over-smoothing).
...	other graphics parameters, see par in package "graphics".

Value

Plot of 1-d complete cross-validation function are sent to graphics window.

kernel	name of kernel to use.
deriv.order	the derivative order to use.
seq.bws	the sequence of bandwidths.
ccv	the values of the complete cross-validation function in the bandwidths grid.

Author(s)

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See Also

[h.ccv.](#)

Examples

```
oldpar <- par(no.readonly = TRUE)
par(mfrow=c(2,1))
plot(h.ccv(trimodal,deriv.order=0),main="")
plot(h.ccv(trimodal,deriv.order=1),main="")
par(oldpar)
```

plot.h.mcv

Plot for Modified Cross-Validation

Description

The [plot.h.mcv](#) function loops through calls to the [h.mcv](#) function. Plot for modified cross-validation function for 1-dimensional data.

Usage

```
## S3 method for class 'h.mcv'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.mcv'
lines(x,seq.bws=NULL, ...)
```

Arguments

x	object of class <code>h.mcv</code> (output from h.mcv).
seq.bws	the sequence of bandwidths in which to compute the modified cross-validation function. By default, the procedure defines a sequence of 50 points, from $0.15 \cdot h_{os}$ to $2 \cdot h_{os}$ (Over-smoothing).
...	other graphics parameters, see par in package "graphics".

Value

Plot of 1-d modified cross-validation function are sent to graphics window.

kernel	name of kernel to use.
deriv.order	the derivative order to use.
seq.bws	the sequence of bandwidths.
mcv	the values of the modified cross-validation function in the bandwidths grid.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

See Also

[h.mcv](#).

Examples

```
oldpar <- par(no.readonly = TRUE)
par(mfrow=c(2,1))
plot(h.mcv(trimodal,deriv.order=0),main="")
plot(h.mcv(trimodal,deriv.order=1),main="")
par(oldpar)
```

plot.h.mlcv

Plot for Maximum-Likelihood Cross-validation

Description

The `plot.h.mlcv` function loops through calls to the `h.mlcv` function. Plot for maximum-likelihood cross-validation function for 1-dimensional data.

Usage

```
## S3 method for class 'h.mlcv'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.mlcv'
lines(x,seq.bws=NULL, ...)
```

Arguments

<code>x</code>	object of class <code>h.mlcv</code> (output from <code>h.mlcv</code>).
<code>seq.bws</code>	the sequence of bandwidths in which to compute the maximum-likelihood cross-validation function. By default, the procedure defines a sequence of 50 points, from $0.15 \cdot h_{os}$ to $2 \cdot h_{os}$ (Over-smoothing).
<code>...</code>	other graphics parameters, see <code>par</code> in package "graphics".

Value

Plot of 1-d maximum-likelihood cross-validation function are sent to graphics window.

<code>kernel</code>	name of kernel to use.
<code>seq.bws</code>	the sequence of bandwidths.
<code>mlcv</code>	the values of the maximum-likelihood cross-validation function in the bandwidths grid.

Author(s)

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See Also

[h.mlcv](#).

Examples

```
plot(h.mlcv(bimodal))
```

plot.h.tcv

Plot for Trimmed Cross-Validation

Description

The [plot.h.tcv](#) function loops through calls to the [h.tcv](#) function. Plot for trimmed cross-validation function for 1-dimensional data.

Usage

```
## S3 method for class 'h.tcv'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.tcv'
lines(x, seq.bws=NULL, ...)
```

Arguments

x	object of class <code>h.tcv</code> (output from h.tcv).
seq.bws	the sequence of bandwidths in which to compute the trimmed cross-validation function. By default, the procedure defines a sequence of 50 points, from $0.15 \cdot h_{os}$ to $2 \cdot h_{os}$ (Over-smoothing).
...	other graphics parameters, see par in package "graphics".

Value

Plot of 1-d trimmed cross-validation function are sent to graphics window.

kernel	name of kernel to use.
deriv.order	the derivative order to use.
seq.bws	the sequence of bandwidths.
tcv	the values of the trimmed cross-validation function in the bandwidths grid.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

See Also

[h.tcv](#).

Examples

```
oldpar <- par(no.readonly = TRUE)
par(mfrow=c(2,1))
plot(h.tcv(trimodal,deriv.order=0),main="")
plot(h.tcv(trimodal,deriv.order=1),seq.bws=seq(0.1,0.5,length.out=50),main="")
par(oldpar)
```

plot.h.ucv

Plot for Unbiased Cross-Validation

Description

The `plot.h.ucv` function loops through calls to the `h.ucv` function. Plot for unbiased cross-validation function for 1-dimensional data.

Usage

```
## S3 method for class 'h.ucv'
plot(x, seq.bws=NULL, ...)
## S3 method for class 'h.ucv'
lines(x,seq.bws=NULL, ...)
```

Arguments

<code>x</code>	object of class <code>h.ucv</code> (output from <code>h.ucv</code>).
<code>seq.bws</code>	the sequence of bandwidths in which to compute the unbiased cross-validation function. By default, the procedure defines a sequence of 50 points, from $0.15 \times h_{os}$ to $2 \times h_{os}$ (Over-smoothing).
<code>...</code>	other graphics parameters, see <code>par</code> in package "graphics".

Value

Plot of 1-d unbiased cross-validation function are sent to graphics window.

<code>kernel</code>	name of kernel to use.
<code>deriv.order</code>	the derivative order to use.
<code>seq.bws</code>	the sequence of bandwidths.
<code>ucv</code>	the values of the unbiased cross-validation function in the bandwidths grid.

Author(s)

Arsalane Chouaib Guidoum <acguidoum@usthb.dz>

See Also

[h.ucv](#).

Examples

```
oldpar <- par(no.readonly = TRUE)
par(mfrow=c(2,1))
plot(h.ucv(trimodal,deriv.order=0),seq.bws=seq(0.06,0.2,length=50))
plot(h.ucv(trimodal,deriv.order=1),seq.bws=seq(0.06,0.2,length=50))
par(oldpar)
```

plot.kernel.conv

Plot for Convolutions of r'th Derivative Kernel Function

Description

The [plot.kernel.conv](#) function loops through calls to the [kernel.conv](#) function. Plot for convolutions of r'th derivative kernel function one-dimensional.

Usage

```
## S3 method for class 'kernel.conv'
plot(x, ...)
```

Arguments

x object of class `kernel.conv` (output from [kernel.conv](#)).

... other graphics parameters, see [par](#) in package "graphics".

Value

Plot of 1-d for convolution of r'th derivative kernel function are sent to graphics window.

Author(s)

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See Also

[kernel.conv](#).

Examples

```
## Gaussian kernel
oldpar <- par(no.readonly = TRUE)
dev.new()
par(mfrow=c(2,2))
plot(kernel.conv(kernel="gaussian",deriv.order=0))
plot(kernel.conv(kernel="gaussian",deriv.order=1))
plot(kernel.conv(kernel="gaussian",deriv.order=2))
plot(kernel.conv(kernel="gaussian",deriv.order=3))

## Silverman kernel

dev.new()
par(mfrow=c(2,2))
plot(kernel.conv(kernel="silverman",deriv.order=0))
plot(kernel.conv(kernel="silverman",deriv.order=1))
plot(kernel.conv(kernel="silverman",deriv.order=2))
plot(kernel.conv(kernel="silverman",deriv.order=3))

par(oldpar)
```

plot.kernel.fun

Plot of r'th Derivative Kernel Function

Description

The `plot.kernel.fun` function loops through calls to the `kernel.fun` function. Plot for r'th derivative kernel function one-dimensional.

Usage

```
## S3 method for class 'kernel.fun'
plot(x, ...)
```

Arguments

x object of class `kernel.fun` (output from `kernel.fun`).

... other graphics parameters, see `par` in package "graphics".

Value

Plot of 1-d for r'th derivative kernel function are sent to graphics window.

Author(s)

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See Also

[kernel.fun.](#)

Examples

```
## Gaussian kernel
oldpar <- par(no.readonly = TRUE)
dev.new()
par(mfrow=c(2,2))
plot(kernel.fun(kernel="gaussian",deriv.order=0))
plot(kernel.fun(kernel="gaussian",deriv.order=1))
plot(kernel.fun(kernel="gaussian",deriv.order=2))
plot(kernel.fun(kernel="gaussian",deriv.order=3))

## Silverman kernel

dev.new()
par(mfrow=c(2,2))
plot(kernel.fun(kernel="silverman",deriv.order=0))
plot(kernel.fun(kernel="silverman",deriv.order=1))
plot(kernel.fun(kernel="silverman",deriv.order=2))
plot(kernel.fun(kernel="silverman",deriv.order=3))

par(oldpar)
```

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