# Package 'locfit'

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aic

Compute Akaike's Information Criterion.

#### **Description**

The calling sequence for aic matches those for the locfit or locfit.raw functions. The fit is not returned; instead, the returned object contains Akaike's information criterion for the fit.

The definition of AIC used here is -2\*log-likelihood + pen\*(fitted d.f.). For quasi-likelihood, and local regression, this assumes the scale parameter is one. Other scale parameters can effectively be used by changing the penalty.

The AIC score is exact (up to numerical roundoff) if the ev="data" argument is provided. Otherwise, the residual sum-of-squares and degrees of freedom are computed using locfit's standard interpolation based approximations.

## Usage

```
aic(x, ..., pen=2)
```

## **Arguments**

x model formula

... other arguments to locfit

pen penalty for the degrees of freedom term

## See Also

```
locfit, locfit.raw, aicplot
```

aicplot

Compute an AIC plot.

## **Description**

The aicplot function loops through calls to the aic function (and hence to locfit), using a different smoothing parameter for each call. The returned structure contains the AIC statistic for each fit, and can be used to produce an AIC plot.

## Usage

```
aicplot(..., alpha)
```

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## **Arguments**

... arguments to the aic, locfit functions.

alpha Matrix of smoothing parameters. The aicplot function loops through calls to

aic, using each row of alpha as the smoothing parameter in turn. If alpha is provided as a vector, it will be converted to a one-column matrix, thus interpret-

ing each component as a nearest neighbor smoothing parameter.

#### Value

An object with class "gcvplot", containing the smoothing parameters and AIC scores. The actual plot is produced using plot.gcvplot.

#### See Also

```
locfit, locfit.raw, gcv, aic, plot.gcvplot
```

## **Examples**

```
data(morths)
plot(aicplot(deaths~age,weights=n,data=morths,family="binomial",
    alpha=seq(0.2,1.0,by=0.05)))
```

ais

Australian Institute of Sport Dataset

# **Description**

The first two columns are the gender of the athlete and their sport. The remaining 11 columns are various measurements made on the athletes.

# Usage

```
data(ais)
```

#### **Format**

A dataframe.

## Source

Cook and Weisberg (1994).

#### References

Cook and Weisberg (1994). An Introduction to Regression Graphics. Wiley, New York.

6 ang

ang

Angular Term for a Locfit model.

# Description

The ang() function is used in a locfit model formula to specify that a variable should be treated as an angular or periodic term. The scale argument is used to set the period.

```
ang(x) is equivalent to lp(x, style="ang").
```

## Usage

```
ang(x,...)
```

# **Arguments**

x numeric variable to be treated periodically.

... Other arguments to 1p.

## References

Loader, C. (1999). Local Regression and Likelihood. Springer, NY (Section 6.2).

# See Also

locfit.

## **Examples**

```
# generate an x variable, and a response with period 0.2
x <- seq(0,1,length=200)
y <- sin(10*pi*x)+rnorm(200)/5

# compute the periodic local fit. Note the scale argument is period/(2pi)
fit <- locfit(y~ang(x,scale=0.2/(2*pi)))

# plot the fit over a single period
plot(fit)

# plot the fit over the full range of the data
plot(fit,xlim=c(0,1))</pre>
```

bad 7

bad

Example dataset for bandwidth selection

# Description

Example dataset from Loader (1999).

## Usage

data(bad)

#### **Format**

Data Frame with x and y variables.

# References

Loader, C. (1999). Bandwidth Selection: Classical or Plug-in? Annals of Statistics 27.

border

Cricket Batting Dataset

## **Description**

Scores in 265 innings for Australian batsman Allan Border.

#### Usage

data(border)

#### **Format**

A dataframe with day (decimalized); not out indicator and score. The not out indicator should be used as a censoring variable.

## **Source**

Compiled from the Cricinfo archives.

## References

CricInfo: The Home of Cricket on the Internet. https://www.espncricinfo.com/

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chemdiab

Chemical Diabetes Dataset

# Description

Numeric variables are rw, fpg, ga, ina and sspg. Classifier cc is the Diabetic type.

## Usage

data(chemdiab)

#### **Format**

Data frame with five numeric measurements and categroical response.

#### **Source**

Reaven and Miller (1979).

#### References

Reaven, G. M. and Miller, R. G. (1979). An attempt to define the nature of chemical diabetes using a multidimensional analysis. Diabetologia 16, 17-24.

claw54

Claw Dataset

# Description

A random sample of size 54 from the claw density of Marron and Wand (1992), as used in Figure 10.5 of Loader (1999).

# Usage

data(claw54)

## **Format**

Numeric vector with length 54.

## **Source**

Randomly generated.

cldem 9

#### References

Loader, C. (1999). Local Regression and Likelihood. Springer, New York.

Marron, J. S. and Wand, M. P. (1992). Exact mean integrated squared error. Annals of Statistics 20, 712-736.

cldem

Example data set for classification

## **Description**

Observations from Figure 8.7 of Loader (1999).

## Usage

data(cldem)

#### **Format**

Data Frame with x and y variables.

#### References

Loader, C. (1999). Local Regression and Likelihood. Springer, New York.

cltest

Test dataset for classification

## **Description**

200 observations from a 2 population model. Under population 0,  $x_{1,i}$  has a standard normal distribution, and  $x_{2,i}=(2-x_{1,i}^2+z_i)/3$ , where  $z_i$  is also standard normal. Under population 1,  $x_{2,i}=-(2-x_{1,i}^2+z_i)/3$ . The optimal classification regions form a checkerboard pattern, with horizontal boundary at  $x_2=0$ , vertical boundaries at  $x_1=\pm\sqrt{2}$ .

This is the same model as the cltrain dataset.

## Usage

```
data(cltest)
```

# **Format**

Data Frame. Three variables x1, x2 and y. The latter indicates class membership.

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cltrain

Training dataset for classification

## Description

200 observations from a 2 population model. Under population 0,  $x_{1,i}$  has a standard normal distribution, and  $x_{2,i}=(2-x_{1,i}^2+z_i)/3$ , where  $z_i$  is also standard normal. Under population 1,  $x_{2,i}=-(2-x_{1,i}^2+z_i)/3$ . The optimal classification regions form a checkerboard pattern, with horizontal boundary at  $x_2=0$ , vertical boundaries at  $x_1=\pm\sqrt{2}$ .

This is the same model as the cltest dataset.

# Usage

```
data(cltrain)
```

#### **Format**

Data Frame. Three variables x1, x2 and y. The latter indicates class membership.

co2

Carbon Dioxide Dataset

# Description

Monthly time series of carbon dioxide measurements at Mauna Loa, Hawaii from 1959 to 1990.

## Usage

data(co2)

## **Format**

Data frame with year, month and co2 variables.

## **Source**

Boden, Sepanski and Stoss (1992).

#### References

Boden, Sepanski and Stoss (1992). Trends '91: A compedium of data on global change - Highlights. Carbon Dioxide Information Analysis Center, Oak Ridge National Laboratory.

*cp* 11

ср

Compute Mallows' Cp for local regression models.

#### **Description**

The calling sequence for cp matches those for the locfit or locfit.raw functions. The fit is not returned; instead, the returned object contains Cp criterion for the fit.

Cp is usually computed using a variance estimate from the largest model under consideration, rather than  $\sigma^2 = 1$ . This will be done automatically when the cpplot function is used.

The Cp score is exact (up to numerical roundoff) if the ev="data" argument is provided. Otherwise, the residual sum-of-squares and degrees of freedom are computed using locfit's standard interpolation based approximations.

## Usage

```
cp(x, ..., sig2=1)
```

## Arguments

x model formula or numeric vector of the independent variable.

... other arguments to locfit and/or locfit.raw.

sig2 residual variance estimate.

#### See Also

```
locfit, locfit.raw, cpplot
```

cpar

Conditionally parametric term for a Locfit model.

#### **Description**

A term entered in a locfit model formula using cpar will result in a fit that is conditionally parametric. Equivalent to lp(x, style="cpar").

This function is presently almost deprecated. Specifying a conditionally parametric fit as  $y^x1+cpar(x2)$  wil no longer work; instead, the model is specified as  $y^2p(x1,x2,style=c("n","cpar"))$ .

#### Usage

```
cpar(x,...)
```

#### **Arguments**

x numeric variable.

... Other arguments to link{lp}().

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## See Also

```
locfit
```

#### **Examples**

```
data(ethanol, package="locfit")
# fit a conditionally parametric model
fit <- locfit(NOx ~ lp(E, C, style=c("n","cpar")), data=ethanol)</pre>
plot(fit)
# one way to force a parametric fit with locfit
fit <- locfit(NOx ~ cpar(E), data=ethanol)</pre>
```

cpplot

Compute a Cp plot.

## **Description**

The cpplot function loops through calls to the cp function (and hence to link{locfit}), using a different smoothing parameter for each call. The returned structure contains the Cp statistic for each fit, and can be used to produce an AIC plot.

## Usage

```
cpplot(..., alpha, sig2)
```

#### **Arguments**

sig2

arguments to the cp, locfit functions. alpha Matrix of smoothing parameters. The cpplot function loops through calls to cp, using each row of alpha as the smoothing parameter in turn. If alpha is pro-

vided as a vector, it will be converted to a one-column matrix, thus interpreting each component as a nearest neighbor smoothing parameter.

Residual variance. If not specified, the residual variance is computed using the

fitted model with the fewest residual degrees of freedom.

## Value

An object with class "gcvplot", containing the smoothing parameters and CP scores. The actual plot is produced using plot.gcvplot.

#### See Also

```
locfit, locfit.raw, gcv, aic, plot.gcvplot
```

# **Examples**

```
data(ethanol)
plot(cpplot(NOx~E, data=ethanol, alpha=seq(0.2,1.0, by=0.05)))
```

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crit

Compute critical values for confidence intervals.

## **Description**

Every "locfit" object contains a critical value object to be used in computing and ploting confidence intervals. By default, a 95% pointwise confidence level is used. To change the confidence level, the critical value object must be substituted using crit and crit<-.

#### Usage

```
crit(fit, const=c(0, 1), d=1, cov=0.95, rdf=0)
crit(fit) <- value</pre>
```

## **Arguments**

fit	"locfit" object. This is optional; if a fit is provided, defaults for the other arguments are taken from the critical value currently stored on this fit, rather than the usual values above. crit(fit) with no other arguments will just return the current critical value.
const	Tube formula constants for simultaneous bands (the default, $c(0,1)$ , produces pointwise coverage). Usually this is generated by the kappa0 function and should not be provided by the user.
d	Dimension of the fit. Again, users shouldn't usually provide it.
cov	Coverage Probability for critical values.
rdf	Residual degrees of freedom. If non-zero, the critical values are based on the Student's t distribution. When rdf=0, the normal distribution is used.
value	Critical value object generated by crit or kappa0.

## Value

Critical value object.

#### See Also

```
locfit, plot.locfit, kappa0, crit<-.</pre>
```

## **Examples**

```
# compute and plot 99% confidence intervals, with local variance estimate.
data(ethanol)
fit <- locfit(NOx~E,data=ethanol)
crit(fit) <- crit(fit,cov=0.99)
plot(fit,band="local")

# compute and plot 99% simultaneous bands
crit(fit) <- kappa0(NOx~E,data=ethanol,cov=0.99)
plot(fit,band="local")</pre>
```

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dat

Locfit - data evaluation structure.

## **Description**

dat is used to specify evaluation on the given data points for locfit.raw().

## Usage

```
dat(cv=FALSE)
```

#### **Arguments**

С٧

Whether cross-validation should be done.

density.lf

Density estimation using Locfit

## **Description**

This function provides an interface to Locfit, in the syntax of (a now old version of) the S-Plus density function. This can reproduce density results, but allows additional locfit.raw arguments, such as the degree of fit, to be given.

It also works in double precision, whereas density only works in single precision.

## Usage

```
density.lf(x, n = 50, window = "gaussian", width, from, to,
  cut = if(iwindow == 4.) 0.75 else 0.5,
  ev = lfgrid(mg = n, 11 = from, ur = to),
  deg = 0, family = "density", link = "ident", ...)
```

## **Arguments**

Χ	numeric vector of observations whose density is to be estimated.
n	number of evaluation points. Equivalent to the locfit.raw mg argument.
window	Window type to use for estimation. Equivalent to the locfit.raw kern argument. This includes all the density windows except cosine.
width	Window width. Following density, this is the full width; not the half-width usually used by Locfit and many other smoothers.
from	Lower limit for estimation domain.
to	Upper limit for estimation domain.
cut	Controls default expansion of the domain.

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ev	Locfit evaluation structure – default lfgrid().
deg	Fitting degree – default 0 for kernel estimation.
family	Fitting family – default is "density".
link	Link function – default is the "identity".
	Additional arguments to locfit, raw, with standard defaults.

## Value

A list with components x (evaluation points) and y (estimated density).

#### See Also

```
density, locfit, locfit.raw
```

## **Examples**

```
data(geyser)
density.lf(geyser, window="tria")
# the same result with density, except less precision.
density(geyser, window="tria")
```

diab

Exhaust emissions

## **Description**

NOx exhaust emissions from a single cylinder engine. Two predictor variables are E (the engine's equivalence ratio) and C (Compression ratio).

## Usage

```
data(ethanol)
```

#### **Format**

Data frame with NOx, E and C variables.

#### **Source**

Brinkman (1981). Also studied extensively by Cleveland (1993).

## References

Brinkman, N. D. (1981). Ethanol fuel - a single-cylinder engine study of efficiency and exhaust emissions. SAE transactions 90, 1414-1424.

Cleveland, W. S. (1993). Visualizing data. Hobart Press, Summit, NJ.

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ethanol

Exhaust emissions

# Description

NOx exhaust emissions from a single cylinder engine. Two predictor variables are E (the engine's equivalence ratio) and C (Compression ratio).

# Usage

data(ethanol)

#### **Format**

Data frame with NOx, E and C variables.

## **Source**

Brinkman (1981). Also studied extensively by Cleveland (1993).

## References

Brinkman, N. D. (1981). Ethanol fuel - a single-cylinder engine study of efficiency and exhaust emissions. SAE transactions 90, 1414-1424.

Cleveland, W. S. (1993). Visualizing data. Hobart Press, Summit, NJ.

expit

Inverse logistic link function

# Description

Computes  $e^x/(1+e^x)$ . This is the inverse of the logistic link function,  $\log(p/(1-p))$ .

# Usage

expit(x)

# **Arguments**

Х

numeric vector

fitted.locfit 17

fitted.locfit

Fitted values for a "locfit" object.

#### **Description**

Evaluates the fitted values (i.e. evaluates the surface at the original data points) for a Locfit object. This function works by reconstructing the model matrix from the original formula, and predicting at those points. The function may be fooled; for example, if the original data frame has changed since the fit, or if the model formula includes calls to random number generators.

#### Usage

```
## S3 method for class 'locfit'
fitted(object, data=NULL, what="coef", cv=FALSE,
studentize=FALSE, type="fit", tr, ...)
```

## **Arguments**

object "locfit" object.

data The data frame for the original fit. Usually, this shouldn't be needed, especially

when the function is called directly. It may be needed when called inside another

function.

what What to compute fitted values of. The default, what="coef", works with the

fitted curve itself. Other choices include "nlx" for the length of the weight diagram; "infl" for the influence function; "band" for the bandwidth; "degr" for the local polynomial degree; "lik" for the maximized local likelihood; "rdf" for the local residual degrees of freedom and "vari" for the variance function.

The interpolation algorithm for some of these quantities is questionable.

cv If TRUE, leave-one-out cross validated fitted values are approximated. Won't

make much sense, unless what="coef".

studentize If TRUE, residuals are studentized.

type Type of fit or residuals to compute. The default is "fit" for fitted.locfit,

and "dev" for residuals.locfit. Other choices include "pear" for Pearson residuals; "raw" for raw residuals, "ldot" for likelihood derivative; "d2" for the deviance residual squared; lddot for the likelihood second derivative. Gen-

erally, type should only be used when what="coef".

tr Back transformation for likelihood models.

... arguments passed to and from methods.

## Value

A numeric vector of the fitted values.

#### See Also

```
locfit, predict.locfit, residuals.locfit
```

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formula.locfit

Formula from a Locfit object.

# Description

Extract the model formula from a locfit object.

## Usage

```
## S3 method for class 'locfit' formula(x, \ldots)
```

# Arguments

x locfit object.

... Arguments passed to and from other methods.

#### Value

Returns the formula from the locfit object.

#### See Also

locfit

gam.lf

Locfit call for Generalized Additive Models

# Description

This is a locfit calling function used by lf() terms in additive models. It is not normally called directly by users.

## Usage

```
gam.lf(x, y, w, xeval, ...)
```

# **Arguments**

```
x numeric predictor
y numeric response
w prior weights
xeval evaluation points
... other arguments to locfit.raw()
```

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#### See Also

```
locfit, locfit.raw, lf, gam
```

gam.slist

Vector of GAM special terms

## Description

This vector adds "lf" to the default vector of special terms recognized by a gam() model formula. To ensure this is recognized, attach the Locfit library with library(locfit,first=T).

#### **Format**

Character vector.

#### See Also

1f, gam

gcv

Compute generalized cross-validation statistic.

## **Description**

The calling sequence for gcv matches those for the locfit or locfit.raw functions. The fit is not returned; instead, the returned object contains Wahba's generalized cross-validation score for the fit

The GCV score is exact (up to numerical roundoff) if the ev="data" argument is provided. Otherwise, the residual sum-of-squares and degrees of freedom are computed using locfit's standard interpolation based approximations.

For likelihood models, GCV is computed uses the deviance in place of the residual sum of squares. This produces useful results but I do not know of any theory validating this extension.

#### Usage

```
gcv(x, ...)
```

# **Arguments**

x, ... Arguments passed on to locfit or locfit.raw.

#### See Also

```
locfit, locfit.raw, gcvplot
```

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gcvplot

Compute a generalized cross-validation plot.

# Description

The gcvplot function loops through calls to the gcv function (and hence to link{locfit}), using a different smoothing parameter for each call. The returned structure contains the GCV statistic for each fit, and can be used to produce an GCV plot.

## Usage

```
gcvplot(..., alpha, df=2)
```

## **Arguments**

... arguments to the gcv, locfit functions.

alpha Matrix of smoothing parameters. The gcvplot function loops through calls to

gcv, using each row of alpha as the smoothing parameter in turn. If alpha is provided as a vector, it will be converted to a one-column matrix, thus interpret-

ing each component as a nearest neighbor smoothing parameter.

df Degrees of freedom to use as the x-axis. 2=trace(L'L).

#### Value

An object with class "gcvplot", containing the smoothing parameters and GCV scores. The actual plot is produced using plot.gcvplot.

## See Also

```
locfit, locfit.raw, gcv, plot.gcvplot, summary.gcvplot
```

#### **Examples**

```
data(ethanol)
plot(gcvplot(NOx~E,data=ethanol,alpha=seq(0.2,1.0,by=0.05)))
```

geyser

Old Faithful Geyser Dataset

## **Description**

The durations of 107 eruptions of the Old Faithful Geyser.

## Usage

```
data(geyser)
```

geyser.round 21

#### **Format**

A numeric vector of length 107.

#### Source

Scott (1992). Note that several different Old Faithful Geyser datasets (including the faithful dataset in R's base library) have been used in various places in the statistics literature. The version provided here has been used in density estimation and bandwidth selection work.

#### References

Scott, D. W. (1992). Multivariate Density Estimation: Theory, Practice and Visualization. Wiley.

geyser.round

Discrete Old Faithful Geyser Dataset

## **Description**

This is a variant of the geyser dataset, where each observation is rounded to the nearest 0.05 minutes, and the counts tallied.

## Usage

```
data(geyser.round)
```

# **Format**

Data Frame with variables duration and count.

## **Source**

Scott (1992). Note that several different Old Faithful Geyser datasets (including the faithful dataset in R's base library) have been used in various places in the statistics literature. The version provided here has been used in density estimation and bandwidth selection work.

#### References

Scott, D. W. (1992). Multivariate Density Estimation: Theory, Practice and Visualization. Wiley.

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hatmatrix

Weight diagrams and the hat matrix for a local regression model.

# Description

hatmatrix() computes the weight diagrams (also known as equivalent or effective kernels) for a local regression smooth. Essentially, hatmatrix() is a front-end to locfit(), setting a flag to compute and return weight diagrams, rather than the fit.

## Usage

```
hatmatrix(formula, dc=TRUE, ...)
```

## **Arguments**

formula	model formula.
dc	derivative adjustment (see locfit.raw)
	Other arguments to locfit and locfit.raw.

#### Value

A matrix with n rows and p columns; each column being the weight diagram for the corresponding locfit fit point. If ev="data", this is the transpose of the hat matrix.

#### See Also

```
locfit, plot.locfit.1d, plot.locfit.2d, plot.locfit.3d, lines.locfit, predict.locfit
```

heart

Survival Times of Heart Transplant Recipients

# Description

The survival times of 184 participants in the Stanford heart transplant program.

## Usage

```
data(heart)
```

#### **Format**

Data frame with surv, cens and age variables.

insect 23

#### **Source**

Miller and Halperin (1982). The original dataset includes information on additional patients who never received a transplant. Other authors reported earlier versions of the data.

#### References

Miller, R. G. and Halperin, J. (1982). Regression with censored data. Biometrika 69, 521-531.

insect

Insect Dataset

# Description

An experiment measuring death rates for insects, with 30 insects at each of five treatment levels.

#### Usage

data(insect)

#### **Format**

Data frame with lconc (dosage), deaths (number of deaths) and nins (number of insects) variables.

#### **Source**

Bliss (1935).

## References

Bliss (1935). The calculation of the dosage-mortality curve. Annals of Applied Biology 22, 134-167.

iris

Fisher's Iris Data (subset)

## **Description**

Four measurements on each of fifty flowers of two species of iris (Versicolor and Virginica) – A classification dataset. Fisher's original dataset contained a third species (Setosa) which is trivially seperable.

## Usage

data(iris)

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## **Format**

Data frame with species, petal.wid, petal.len, sepal.wid, sepal.len.

#### **Source**

Fisher (1936). Reproduced in Andrews and Herzberg (1985) Chapter 1.

#### References

Andrews, D. F. and Herzberg, A. M. (1985). Data. Springer-Verlag.

Fisher, R. A. (1936). The Use of Multiple Measurements in Taxonomic Problems. Annals of Eugenics 7, Part II. 179-188.

kangaroo

Kangaroo skull measurements dataset

# Description

Variables are sex (m/f), spec (giganteus, melanops, fuliginosus) and 18 numeric measurements.

## Usage

data(kangaroo)

## **Format**

Data frame with measurements on the skulls of 101 kangaroos. (number of insects) variables.

#### **Source**

Andrews and Herzberg (1985) Chapter 53.

#### References

Andrews, D. F. and Herzberg, A. M. (1985). Data. Springer-Verlag, New York.

kappa0 25

	kappa0	Critical Values for Simultaneous Confidence Bands.
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## **Description**

The geometric constants for simultaneous confidence bands are computed, as described in Sun and Loader (1994) (bias adjustment is not implemented here). These are then passed to the crit function, which computes the critical value for the confidence bands.

The method requires both the weight diagrams l(x), the derivative l'(x) and (in 2 or more dimensions) the second derivatives l''(x). These are implemented exactly for a constant bandwidth. For nearest neighbor bandwidths, the computations are approximate and a warning is produced.

The theoretical justification for the bands uses normality of the random errors  $e_1, \ldots, e_n$  in the regression model, and in particular the spherical symmetry of the error vector. For non-normal distributions, and likelihood models, one relies on central limit and related theorems.

Computation uses the product Simpson's rule to evaluate the multidimensional integrals (The domain of integration, and hence the region of simultaneous coverage, is determined by the flim argument). Expect the integration to be slow in more than one dimension. The mint argument controls the precision.

#### Usage

```
kappa0(formula, cov=0.95, ev=lfgrid(20), ...)
```

# Arguments

formula	Local regression model formula. A "locfit" object can also be provided; in this case the formula and other arguments are extracted from this object.
cov	Coverage Probability for critical values.
ev	Locfit evaluation structure. Should usually be a grid – this specifies the integration rule.
	Other arguments to locfit. Important arguments include flim and alpha.

#### Value

A list with components for the critical value, geometric constants, e.t.c. Can be passed directly to plot.locfit as the crit argument.

## References

Sun, J. and Loader, C. (1994). Simultaneous confidence bands for linear regression and smoothing. Annals of Statistics 22, 1328-1345.

## See Also

```
locfit, plot.locfit, crit, crit<-.</pre>
```

26 kdeb

## **Examples**

```
# compute and plot simultaneous confidence bands
data(ethanol)
fit <- locfit(NOx~E,data=ethanol)
crit(fit) <- kappa0(NOx~E,data=ethanol)
plot(fit,crit=crit,band="local")</pre>
```

kdeb

Bandwidth selectors for kernel density estimation.

## **Description**

Function to compute kernel density estimate bandwidths, as used in the simulation results in Chapter 10 of Loader (1999).

This function is included for comparative purposes only. Plug-in selectors are based on flawed logic, make unreasonable and restrictive assumptions and do not use the full power of the estimates available in Locfit. Any relation between the results produced by this function and desirable estimates are entirely coincidental.

## Usage

```
kdeb(x, h0 = 0.01 * sd, h1 = sd, meth = c("AIC", "LCV", "LSCV", "BCV", "SJPI", "GKK"), kern = "gauss", gf = 2.5)
```

# Arguments

X	One dimensional data vector.
h0	Lower limit for bandwidth selection. Can be fairly small, but h0=0 would cause problems.
h1	Upper limit.
meth	Required selection method(s).
kern	Kernel. Most methods require kern="gauss", the default for this function only.
gf	Standard deviation for the gaussian kernel. Default 2.5, as Locfit's standard. Most papers use 1.

## Value

Vector of selected bandwidths.

#### References

Loader, C. (1999). Local Regression and Likelihood. Springer, New York.

km.mrl 27

km.mrl

Mean Residual Life using Kaplan-Meier estimate

## Description

This function computes the mean residual life for censored data using the Kaplan-Meier estimate of the survival function. If S(t) is the K-M estimate, the MRL for a censored observation is computed as  $(\int_t^\infty S(u)du)/S(t)$ . We take S(t)=0 when t is greater than the largest observation, regardless of whether that observation was censored.

When there are ties between censored and uncensored observations, for definiteness our ordering places the censored observations before uncensored.

This function is used by locfit.censor to compute censored regression estimates.

# Usage

```
km.mrl(times, cens)
```

## **Arguments**

times Observed survival times.

cens Logical variable indicating censoring. The coding is 1 or TRUE for censored; 0

or FALSE for uncensored.

#### Value

A vector of the estimated mean residual life. For uncensored observations, the corresponding estimate is 0.

## References

```
Buckley, J. and James, I. (1979). Linear Regression with censored data. Biometrika 66, 429-436. Loader, C. (1999). Local Regression and Likelihood. Springer, NY (Section 7.2).
```

#### See Also

```
locfit.censor
```

# **Examples**

```
# censored regression using the Kaplan-Meier estimate.
data(heart, package="locfit")
fit <- locfit.censor(log10(surv+0.5)~age, cens=cens, data=heart, km=TRUE)
plotbyfactor(heart$age, 0.5+heart$surv, heart$cens, ylim=c(0.5,16000), log="y")
lines(fit, tr=function(x)10^x)</pre>
```

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lcv

Compute Likelihood Cross Validation Statistic.

## **Description**

The calling sequence for lcv matches those for the locfit or locfit.raw functions. The fit is not returned; instead, the returned object contains likelihood cross validation score for the fit.

The LCV score is exact (up to numerical roundoff) if the ev="cross" argument is provided. Otherwise, the influence and cross validated residuals are computed using locfit's standard interpolation based approximations.

## Usage

```
lcv(x, ...)
```

## Arguments

x model formula

.. other arguments to locfit

#### See Also

```
locfit, locfit.raw, lcvplot
```

lcvplot

Compute the likelihood cross-validation plot.

# Description

The lcvplot function loops through calls to the lcv function (and hence to link{locfit}), using a different smoothing parameter for each call. The returned structure contains the likelihood cross validation statistic for each fit, and can be used to produce an LCV plot.

## Usage

```
lcvplot(..., alpha)
```

#### **Arguments**

... arguments to the lcv, locfit functions.

alpha

Matrix of smoothing parameters. The aicplot function loops through calls to lcv, using each row of alpha as the smoothing parameter in turn. If alpha is provided as a vector, it will be converted to a one-column matrix, thus interpreting each component as a nearest neighbor smoothing parameter.

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#### Value

An object with class "gcvplot", containing the smoothing parameters and LCV scores. The actual plot is produced using plot.gcvplot.

#### See Also

```
locfit, locfit.raw, gcv, lcv, plot.gcvplot
```

#### **Examples**

```
data(ethanol)
plot(lcvplot(NOx~E,data=ethanol,alpha=seq(0.2,1.0,by=0.05)))
```

left

One-sided left smooth for a Locfit model.

## **Description**

The left() function is used in a locfit model formula to specify a one-sided smooth: when fitting at a point x, only data points with  $x_i \leq x$  should be used. This can be useful in estimating points of discontinuity, and in cross-validation for forecasting a time series. left(x) is equivalent to lp(x,style="left").

When using this function, it will usually be necessary to specify an evaluation structure, since the fit is not smooth and locfit's interpolation methods are unreliable. Also, it is usually best to use deg=0 or deg=1, otherwise the fits may be too variable. If nearest neighbor bandwidth specification is used, it does not recognize left().

## Usage

```
left(x,...)
```

#### **Arguments**

```
x numeric variable.
... Other arguments to 1p().
```

#### See Also

```
locfit, lp, right
```

## **Examples**

```
# compute left and right smooths
data(penny)
xev <- (1945:1988)+0.5
fitl <- locfit(thickness~left(year,h=10,deg=1), ev=xev, data=penny)
fitr <- locfit(thickness~right(year,h=10,deg=1),ev=xev, data=penny)
# plot the squared difference, to show the change points.
plot( xev, (predict(fitr,where="ev") - predict(fitl,where="ev"))^2 )</pre>
```

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Locfit term in Additive Model formula

1f

## **Description**

This function is used to specify a smooth term in a gam() model formula.

This function is designed to be used with the S-Plus gam() function. For R users, there are at least two different gam() functions available. Most current distributions of R will include the mgcv library by Simon Wood; 1f() is not compatable with this function.

On CRAN, there is a gam package by Trevor Hastie, similar to the S-Plus version. 1f() should be compatable with this, although it's untested.

# Usage

```
lf(..., alpha=0.7, deg=2, scale=1, kern="tcub", ev=rbox(), maxk=100)
```

## **Arguments**

```
... numeric predictor variable(s) alpha, deg, scale, kern, ev, maxk these are as in locfit.raw.
```

## See Also

```
locfit, locfit.raw, gam.lf, gam
```

#### **Examples**

```
## Not run:
# fit an additive semiparametric model to the ethanol data.
stopifnot(require(gam))
# The `gam' package must be attached _before_ `locfit', otherwise
# the following will not work.
data(ethanol, package = "lattice")
fit <- gam(NOx ~ lf(E) + C, data=ethanol)
op <- par(mfrow=c(2, 1))
plot(fit)
par(op)
## End(Not run)</pre>
```

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lfeval

Extract Locfit Evaluation Structure.

## **Description**

Extracts the evaluation structure from a "locfit" object. This object has the class "lfeval", and has its own set of methods for plotting e.t.c.

## Usage

```
lfeval(object)
```

## **Arguments**

object

"locfit" object

## Value

"lfeval" object.

#### See Also

```
locfit, plot.lfeval, print.lfeval
```

lfgrid

Locfit - grid evaluation structure.

# Description

lfgrid() is used to specify evaluation on a grid of points for locfit.raw(). The structure computes a bounding box for the data, and divides that into a grid with specified margins.

# Usage

```
lfgrid(mg=10, ll, ur)
```

# Arguments

mg	Number of grid points along each margin. Can be a single number (which is applied in each dimension), or a vector specifying a value for each dimension.
11	Lower left limits for the grid. Length should be the number of dimensions of the data provided to locfit.raw().
ur	Upper right limits for the grid. By default, 11 and ur are generated as the bounding box for the data.

32 Iflim

## **Examples**

```
data(ethanol, package="locfit")
plot.eval(locfit(NOx ~ lp(E, C, scale=TRUE), data=ethanol, ev=lfgrid()))
```

1fknots

Extraction of fit-point information from a Locfit object.

#### **Description**

Extracts information, such as fitted values, influence functions from a "locfit" object.

# Usage

```
lfknots(x, tr, what = c("x", "coef", "h", "nlx"), delete.pv = TRUE)
```

### **Arguments**

x Fitted object from locfit().

tr Back transformation. Default is the invers link function from the Locfit object.

what What to return; default is c("x", "coef", "h", "nlx"). Allowed fields are x (fit points); coef (fitted values); f1 (local slope); nlx (length of the weight

diagram); nlx1 (estimated derivative of nlx); se (standard errors); infl (influence function); infla (slope of influence function); lik (maximixed local log-likelihood and local degrees of freedom); h (bandwidth) and deg (degree of

fit).

delete.pv If T, pseudo-vertices are deleted.

#### Value

A matrix with one row for each fit point. Columns correspond to the specified what vector; some fields contribute multiple columns.

1flim

Construct Limit Vectors for Locfit fits.

## **Description**

This function is used internally to interpret xlim and flim arguments. It should not be called directly.

## Usage

```
lflim(limits, nm, ret)
```

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## **Arguments**

limits Limit argument.nm Variable names.ret Initial return vector.

## Value

Vector with length 2\*dim.

#### See Also

locfit

1fmarg

Generate grid margins.

# Description

This function is usually called by plot.locfit.

## Usage

```
lfmarg(xlim, m = 40)
```

## **Arguments**

valim Vector of limits for the grid. Should be of length 2\*d; the first d components

represent the lower left corner, and the next d components the upper right corner.

Can also be a "locfit" object.

m Number of points for each grid margin. Can be a vector of length d.

## Value

A list, whose components are the d grid margins.

#### See Also

```
locfit, plot.locfit
```

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lines.locfit

Add locfit line to existing plot

## **Description**

Adds a Locfit line to an existing plot. 11ines is for use within a panel function for Lattice.

## Usage

```
## $3 method for class 'locfit'
lines(x, m=100, tr=x$trans, ...)
## $3 method for class 'locfit'
llines(x, m=100, tr=x$trans, ...)
```

## **Arguments**

x locfit object. Should be a model with one predictor.

m Number of points to evaluate the line at.

tr Transformation function to use for plotting. Default is the inverse link function,

or the identity function if derivatives are required.

. . . Other arguments to the default lines function.

#### See Also

```
locfit, plot.locfit, lines
```

livmet

liver Metastases dataset

## **Description**

Survival times for 622 patients diagnosed with Liver Metastases.

Beware, the censoring variable is coded as 1 = uncensored, so use cens=1-z in locfit() calls.

# Usage

```
data(livmet)
```

#### **Format**

Data frame with survival times (t), censoring indicator (z) and a number of covariates.

## Source

Haupt and Mansmann (1995)

locfit 35

#### References

Haupt, G. and Mansmann, U. (1995) CART for Survival Data. Statlib Archive.

locfit

Local Regression, Likelihood and Density Estimation.

# Description

locfit is the model formula-based interface to the Locfit library for fitting local regression and likelihood models.

locfit is implemented as a front-end to locfit.raw. See that function for options to control smoothing parameters, fitting family and other aspects of the fit.

#### Usage

## **Arguments**

formula	Model Formula; e.g. $y\sim lp(x)$ for a regression model; $\sim lp(x)$ for a density estimation model. Use of $lp()$ on the RHS is recommended, especially when non-default smoothing parameters are used.
data	Data Frame.
weights	Prior weights (or sample sizes) for individual observations. This is typically used where observations have unequal variance.
cens	Censoring indicator. 1 (or TRUE) denotes a censored observation. 0 (or FALSE) denotes uncensored.
base	Baseline for local fitting. For local regression models, specifying a base is equivalent to using y-base as the reponse. But base also works for local likelihood.
subset	Subset observations in the data frame.
geth	Don't use.
• • •	Other arguments to locfit.raw() (or the lfproc).
lfproc	A processing function to compute the local fit. Default is locfit.raw(). Other choices include locfit.robust(), locfit.censor() and locfit.quasi().

#### Value

An object with class "locfit". A standard set of methods for printing, ploting, etc. these objects is provided.

## References

Loader, C. (1999). Local Regression and Likelihood. Springer, New York.

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## See Also

```
locfit.raw
```

#### **Examples**

```
# fit and plot a univariate local regression
data(ethanol, package="locfit")
fit <- locfit(NOx ~ E, data=ethanol)
plot(fit, get.data=TRUE)

# a bivariate local regression with smaller smoothing parameter
fit <- locfit(NOx~lp(E,C,nn=0.5,scale=0), data=ethanol)
plot(fit)

# density estimation
data(geyser, package="locfit")
fit <- locfit( ~ lp(geyser, nn=0.1, h=0.8))
plot(fit,get.data=TRUE)</pre>
```

locfit.censor

Censored Local Regression

## **Description**

locfit.censor produces local regression estimates for censored data. The basic idea is to use an EM style algorithm, where one alternates between estimating the regression and the true values of censored observations.

locfit.censor is designed as a front end to locfit.raw with data vectors, or as an intermediary between locfit and locfit.raw with a model formula. If you can stand the syntax, the second calling sequence above will be slightly more efficient than the third.

## Usage

```
locfit.censor(x, y, cens, ..., iter=3, km=FALSE)
```

#### **Arguments**

X	Either a locfit model formula or a numeric vector of the predictor variable.
У	If x is numeric, y gives the response variable.
cens	Logical variable indicating censoring. The coding is 1 or TRUE for censored; 0 or FALSE for uncensored.
• • •	Other arguments to locfit.raw
iter	Number of EM iterations to perform
km	If km=TRUE, the estimation of censored observations uses the Kaplan-Meier estimate, leading to a local version of the Buckley-James estimate. If km=F, the estimation is based on a normal model (Schmee and Hahn). Beware of claims that B-J is nonparametric; it makes stronger assumptions on the upper tail of

survival distributions than most authors care to admit.

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#### Value

locfit object.

#### References

Buckley, J. and James, I. (1979). Linear Regression with censored data. Biometrika 66, 429-436.

Loader, C. (1999). Local Regression and Likelihood. Springer, NY (Section 7.2).

Schmee, J. and Hahn, G. J. (1979). A simple method for linear regression analysis with censored data (with discussion). Technometrics 21, 417-434.

#### See Also

```
km.mrl, locfit, locfit.raw
```

# **Examples**

```
data(heart, package="locfit") fit <- locfit.censor(log10(surv+0.5) ^{\sim} age, cens=cens, data=heart) ## Can also be written as: ## Not run: fit <- locfit(log10(surv + 0.5) ^{\sim} age, cens=cens, data=heart, lfproc=locfit.censor) with(heart, plotbyfactor(age, 0.5 + surv, cens, ylim=c(0.5, 16000), log="y")) lines(fit, tr=function(x) 10^{\wedge}x)
```

locfit.matrix

Reconstruct a Locfit model matrix.

### **Description**

Reconstructs the model matrix, and associated variables such as the response, prior weights and censoring indicators, from a locfit object. This is used by functions such as fitted.locfit; it is not normally called directly. The function will only work properly if the data frame has not been changed since the fit was constructed.

# Usage

```
locfit.matrix(fit, data)
```

# **Arguments**

```
fit Locfit object data Data Frame.
```

#### Value

A list with variables x (the model matrix); y (the response); w (prior weights); sc (scales); ce (censoring indicator) and base (baseline fit).

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### See Also

```
locfit, fitted.locfit, residuals.locfit
```

locfit.quasi

Local Quasi-Likelihood with global reweighting.

# Description

locfit.quasi assumes a specified mean-variance relation, and performs iterartive reweighted local regression under this assumption. This is appropriate for local quasi-likelihood models, and is an alternative to specifying a family such as "qpoisson".

locfit.quasi is designed as a front end to locfit.raw with data vectors, or as an intermediary between locfit and locfit.raw with a model formula. If you can stand the syntax, the second calling sequence above will be slightly more efficient than the third.

# Usage

```
locfit.quasi(x, y, weights, ..., iter=3, var=abs)
```

# **Arguments**

x Either a locfit model formula or a numeric vector of the predictor variable.

y If x is numeric, y gives the response variable.

weights Case weights to use in the fitting.

Other arguments to locfit.raw

iter Number of EM iterations to perform

var Function specifying the assumed relation between the mean and variance.

### Value

```
"locfit" object.
```

#### See Also

```
locfit, locfit.raw
```

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locfit.raw

Local Regression, Likelihood and Density Estimation.

#### **Description**

locfit.raw is an interface to Locfit using numeric vectors (for a model-formula based interface, use locfit). Although this function has a large number of arguments, most users are likely to need only a small subset.

The first set of arguments (x, y, weights, cens, and base) specify the regression variables and associated quantities.

Another set (scale, alpha, deg, kern, kt, acri and basis) control the amount of smoothing: bandwidth, smoothing weights and the local model. Most of these arguments are deprecated - they'll currently still work, but should be provided through the lp() model term instead.

deriv and dc relate to derivative (or local slope) estimation.

family and link specify the likelihood family.

xlim and renorm may be used in density estimation.

ev specifies the evaluation structure or set of evaluation points.

maxk, itype, mint, maxit and debug control the Locfit algorithms, and will be rarely used.

geth and sty are used by other functions calling locfit.raw, and should not be used directly.

#### Usage

```
locfit.raw(x, y, weights=1, cens=0, base=0,
    scale=FALSE, alpha=0.7, deg=2, kern="tricube", kt="sph",
    acri="none", basis=list(NULL),
    deriv=numeric(0), dc=FALSE,
    family, link="default",
    xlim, renorm=FALSE,
    ev=rbox(),
    maxk=100, itype="default", mint=20, maxit=20, debug=0,
    geth=FALSE, sty="none")
```

### **Arguments**

х	Vector (or matrix) of the independent variable(s). Can be constructed using the $lp()$ function.
у	Response variable for regression models. For density families, y can be omitted.
weights	Prior weights for observations (reciprocal of variance, or sample size).
cens	Censoring indicators for hazard rate or censored regression. The coding is 1 (or TRUE) for a censored observation, and $\emptyset$ (or FALSE) for uncensored observations.
base	Baseline parameter estimate. If provided, the local regression model is fitted as $Y_i = b_i + m(x_i) + \epsilon_i$ , with Locfit estimating the $m(x)$ term. For regression models, this effectively subtracts $b_i$ from $Y_i$ . The advantage of the base formulation is that it extends to likelihood regression models.

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scale Deprecated - see lp().

alpha Deprecated - see lp(). A single number (e.g. alpha=0.7) is interpreted as a

> nearest neighbor fraction. With two componentes (e.g. alpha=c(0.7,1.2)), the first component is a nearest neighbor fraction, and the second component is a fixed component. A third component is the penalty term in locally adaptive

smoothing.

deg Degree of local polynomial. Deprecated - see lp().

Weight function, default = "tcub". Other choices are "rect", "trwt", "tria", kern

"epan", "bisq" and "gauss". Choices may be restricted when derivatives are

required; e.g. for confidence bands and some bandwidth selectors.

Kernel type, "sph" (default); "prod". In multivariate problems, "prod" uses a kt

simplified product model which speeds up computations.

Deprecated - see lp(). acri

basis User-specified basis functions.

deriv Derivative estimation. If deriv=1, the returned fit will be estimating the deriva-

tive (or more correctly, an estimate of the local slope). If deriv=c(1,1) the second order derivative is estimated. deriv=2 is for the partial derivative, with

respect to the second variable, in multivariate settings.

Derivative adjustment. dc

Local likelihood family; "gaussian"; "binomial"; "poisson"; "gamma" and

"geom". Density and rate estimation families are "dens", "rate" and "hazard" (hazard rate). If the family is preceded by a 'q' (for example, family="qbinomial"), quasi-likelihood variance estimates are used. Otherwise, the residual variance (rv) is fixed at 1. The default family is "qgauss" if a response y is provided;

"density" if no response is provided.

link Link function for local likelihood fitting. Depending on the family, choices may

be "ident", "log", "logit", "inverse", "sqrt" and "arcsin".

For density estimation, Locfit allows the density to be supported on a bounded interval (or rectangle, in more than one dimension). The format should be

c(11,u1) where 11 is a vector of the lower bounds and ur the upper bounds. Bounds such as  $[0, \infty)$  are not supported, but can be effectively implemented by

specifying a very large upper bound.

Local likelihood density estimates may not integrate exactly to 1. If renorm=T, renorm

the integral will be estimated numerically and the estimate rescaled. Presently

this is implemented only in one dimension.

The evaluation structure, rbox() for tree structures; lfgrid() for grids; dat() for data points; none() for none. A vector or matrix of evaluation points can also be provided, although in this case you may prefer to use the smooth.lf()

interface to Locfit. Note that arguments flim, mg and cut are now given as arguments to the evaluation structure function, rather than to locfit.raw()

directly (change effective 12/2001).

Controls space assignment for evaluation structures. For the adaptive evaluation structures, it is impossible to be sure in advance how many vertices will be generated. If you get warnings about 'Insufficient vertex space', Locfit's default

assignment can be increased by increasing maxk. The default is maxk=100.

maxk

family

xlim

ev

locfit.robust 41

itype	Integration type for density estimation. Available methods include "prod", "mult" and "mlin"; and "haz" for hazard rate estimation problems. The available in the state of the
	able integration methods depend on model specification (e.g. dimension, degree of fit). By default, the best available method is used.
mint	Points for numerical integration rules. Default 20.
maxit	Maximum iterations for local likelihood estimation. Default 20.
debug	If $> 0$ ; prints out some debugging information.
geth	Don't use!
sty	Deprecated - see lp().

### Value

An object with class "locfit". A standard set of methods for printing, ploting, etc. these objects is provided.

### References

Loader, C., (1999) Local Regression and Likelihood.

	locfit.robust	Robust Local Regression	
--	---------------	-------------------------	--

# **Description**

locfit.robust implements a robust local regression where outliers are iteratively identified and downweighted, similarly to the lowess method (Cleveland, 1979). The iterations and scale estimation are performed on a global basis.

The scale estimate is 6 times the median absolute residual, while the robust downweighting uses the bisquare function. These are performed in the S code so easily changed.

This can be interpreted as an extension of M estimation to local regression. An alternative extension (implemented in locfit via family="qrgauss") performs the iteration and scale estimation on a local basis.

# Usage

```
locfit.robust(x, y, weights, ..., iter=3)
```

# Arguments

X	Either a locfit model formula or a numeric vector of the predictor variable.
У	If x is numeric, y gives the response variable.
weights	weights to use in the fitting.
	Other arguments to locfit.raw.
iter	Number of iterations to perform

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# Value

```
"locfit" object.
```

#### References

Cleveland, W. S. (1979). Robust locally weighted regression and smoothing scatterplots. J. Amer. Statist. Assn. 74, 829-836.

# See Also

```
locfit, locfit.raw
```

1p

Local Polynomial Model Term

# Description

1p is a local polynomial model term for Locfit models. Usually, it will be the only term on the RHS of the model formula.

Smoothing parameters should be provided as arguments to lp(), rather than to locfit().

# Usage

```
lp(..., nn, h, adpen, deg, acri, scale, style)
```

# Arguments

	Predictor variables for the local regression model.
nn	Nearest neighbor component of the smoothing parameter. Default value is 0.7, unless either h or adpen are provided, in which case the default is 0.
h	The constant component of the smoothing parameter. Default: 0.
adpen	Penalty parameter for adaptive fitting.
deg	Degree of polynomial to use.
acri	Criterion for adaptive bandwidth selection.
style	Style for special terms (left, ang e.t.c.). Do not try to set this directly; call locfit instead.
scale	A scale to apply to each variable. This is especially important for multivariate fitting, where variables may be measured in non-comparable units. It is also used to specify the frequency for ang terms. If scale=F (the default) no scaling is performed. If scale=T, marginal standard deviations are used. Alternatively,

a numeric vector can provide scales for the individual variables.

### See Also

```
locfit, locfit.raw
```

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### **Examples**

```
data(ethanol, package="locfit")
# fit with 50% nearest neighbor bandwidth.
fit <- locfit(NOx~lp(E,nn=0.5),data=ethanol)
# bivariate fit.
fit <- locfit(NOx~lp(E,C,scale=TRUE),data=ethanol)
# density estimation
data(geyser, package="locfit")
fit <- locfit.raw(lp(geyser,nn=0.1,h=0.8))</pre>
```

lscv

Least Squares Cross Validation Statistic.

### **Description**

The calling sequence for lscv matches those for the locfit or locfit.raw functions. Note that this function is only designed for density estimation in one dimension. The returned object contains the least squares cross validation score for the fit.

The computation of  $\int \hat{f}(x)^2 dx$  is performed numerically. For kernel density estimation, this is unlikely to agree exactly with other LSCV routines, which may perform the integration analytically.

# Usage

```
lscv(x, ..., exact=FALSE)
```

# **Arguments**

```
    x model formula (or numeric vector, if exact=T)
    ... other arguments to locfit or lscv.exact
    exact
    By default, the computation is approximate. If exact=TRUE, exact computation using lscv.exact is performed. This uses kernel density estimation with a constant bandwidth.
```

#### Value

A vector consisting of the LSCV statistic and fitted degrees of freedom.

#### See Also

```
locfit, locfit.raw, lscv.exact lscvplot
```

### **Examples**

```
# approximate calculation for a kernel density estimate
data(geyser, package="locfit")
lscv(~lp(geyser,h=1,deg=0), ev=lfgrid(100,ll=1,ur=6), kern="gauss")
# same computation, exact
lscv(lp(geyser,h=1),exact=TRUE)
```

44 lscvplot

lscv.exact

Exact LSCV Calculation

### Description

This function performs the exact computation of the least squares cross validation statistic for onedimensional kernel density estimation and a constant bandwidth.

At the time of writing, it is implemented only for the Gaussian kernel (with the standard deviation of 0.4; Locfit's standard).

### Usage

```
lscv.exact(x, h=0)
```

# **Arguments**

x Numeric data vector.

h The bandwidth. If x is constructed with lp(), the bandwidth should be given there instead.

### Value

A vector of the LSCV statistic and the fitted degrees of freedom.

### See Also

```
lscv, lscvplot
```

# **Examples**

```
data(geyser, package="locfit")
lscv.exact(lp(geyser,h=0.25))
# equivalent form using lscv
lscv(lp(geyser, h=0.25), exact=TRUE)
```

lscvplot

Compute the LSCV plot.

# Description

The 1scvplot function loops through calls to the 1scv function (and hence to link{locfit}), using a different smoothing parameter for each call. The returned structure contains the LSCV statistic for each density estimate, and can be used to produce an LSCV plot.

### Usage

```
lscvplot(..., alpha)
```

mcyc 45

# **Arguments**

... arguments to the lscv, locfit functions.

alpha Matrix of smoothing parameters. The lscvplot function loops through calls to

lscv, using each row of alpha as the smoothing parameter in turn. If alpha is provided as a vector, it will be converted to a one-column matrix, thus interpret-

ing each component as a nearest neighbor smoothing parameter.

#### Value

An object with class "gcvplot", containing the smoothing parameters and LSCV scores. The actual plot is produced using plot.gcvplot.

### See Also

```
locfit, locfit.raw, gcv, lscv, plot.gcvplot
```

mcyc

Acc(De?)celeration of a Motorcycle Hitting a Wall

# **Description**

Measurements of the acceleration of a motorcycle as it hits a wall. Actually, rumored to be a concatenation of several such datasets.

### Usage

```
data(mcyc)
```

### **Format**

Data frame with time and accel variables.

#### **Source**

```
H\"ardle (1990).
```

### References

H\"ardle, W. (1990). Applied Nonparametric Regression. Cambridge University Press.

46 mmsamp

mine

Fracture Counts in Coal Mines

# Description

The number of fractures in the upper seam of coal mines, and four predictor variables. This dataset can be modeled using Poisson regression.

# Usage

data(mine)

#### **Format**

A dataframe with the response frac, and predictor variables extrp, time, seamh and inb.

### **Source**

Myers (1990).

### References

Myers, R. H. (1990). Classical and Modern Regression with Applications (Second edition). PWS-Kent Publishing, Boston.

mmsamp

Test dataset for minimax Local Regression

# **Description**

50 observations, as used in Figure 13.1 of Loader (1999).

# Usage

data(cltest)

#### **Format**

Data Frame with x and y variables.

# References

Loader, C. (1999). Local Regression and Likelihood. Springer, New York.

morths 47

 ${\tt morths}$ 

Henderson and Sheppard Mortality Dataset

# **Description**

Observed mortality for 55 to 99.

### Usage

```
data(morths)
```

### **Format**

Data frame with age, n and number of deaths.

#### **Source**

Henderson and Sheppard (1919).

### References

Henderson, R. and Sheppard, H. N. (1919). Graduation of mortality and other tables. Actuarial Society of America, New York.

none

Locfit Evaluation Structure

# Description

none() is an evaluation structure for locfit.raw(), specifying no evaluation points. Only the initial parametric fit is computed - this is the easiest and most efficient way to coerce Locfit into producing a parametric regression fit.

# Usage

none()

### **Examples**

```
data(ethanol, package="locfit")
# fit a fourth degree polynomial using locfit
fit <- locfit(NOx~E,data=ethanol,deg=4,ev=none())
plot(fit,get.data=TRUE)</pre>
```

48 plot.eval

penny

Penny Thickness Dataset

# **Description**

For each year, 1945 to 1989, the thickness of two U.S. pennies was recorded.

# Usage

data(penny)

### **Format**

A dataframe.

### **Source**

Scott (1992).

#### References

Scott (1992). Multivariate Density Estimation. Wiley, New York.

plot.eval

Plot evaluation points from a 2-d locfit object.

# Description

This function is used to plot the evaluation structure generated by Locfit for a two dimensional fit. Vertices of the tree structure are displayed as 0; pseudo-vertices as \*.

#### Usage

```
plot.eval(x, add=FALSE, text=FALSE, ...)
```

# Arguments

x "locfit" object.

add If TRUE, add to existing plot.

text If TRUE, numbers will be added indicating the order points were added.

... Arguments passed to and from other methods.

# See Also

locfit.

plot.gcvplot 49

# **Examples**

```
data(ethanol, package="locfit")
fit <- locfit(NOx ~ E + C, data=ethanol, scale=0)
plot.eval(fit)</pre>
```

plot.gcvplot

Produce a cross-validation plot.

# **Description**

Plots the value of the GCV (or other statistic) in a gcvplot object against the degrees of freedom of the fit.

### Usage

```
## S3 method for class 'gcvplot'
plot(x, xlab = "Fitted DF", ylab = x$cri, ...)
```

### **Arguments**

```
x A gcvplot object, produced by gcvplot, aicplot etc.
xlab Text label for the x axis.
ylab Text label for the y axis.
... Other arguments to plot .
```

# See Also

```
locfit, locfit.raw, gcv, aicplot, cpplot, gcvplot, lcvplot
```

# **Examples**

```
data(ethanol)
plot(gcvplot(NOx~E,data=ethanol,alpha=seq(0.2,1.0,by=0.05)))
```

50 plot.locfit

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			-vai

Plot a Locfit Evaluation Structure.

# **Description**

Plots the evaluation points from a locfit or lfeval structure, for one- or two-dimensional fits.

### Usage

```
## S3 method for class 'lfeval'
plot(x, add=FALSE, txt=FALSE, ...)
```

# Arguments

X	A Ifeval or locfit object
add	If TRUE, the points will be added to the existing plot. Otherwise, a new plot is created.
txt	If TRUE, the points are annotated with numbers in the order they were entered into the fit.
	Additional graphical parameters.

### Value

```
"lfeval" object.
```

#### See Also

```
lfeval, locfit, print.lfeval
```

plot.locfit

Plot an object of class locfit.

# **Description**

The plot.locfit function generates grids of ploting points, followed by a call to preplot.locfit. The returned object is then passed to plot.locfit.1d, plot.locfit.2d or plot.locfit.3d as appropriate.

# Usage

```
## S3 method for class 'locfit'
plot(x, xlim, pv, tv, m, mtv=6, band="none", tr=NULL,
  what = "coef", get.data=FALSE, f3d=(d == 2) && (length(tv) > 0), ...)
```

plot.locfit 51

# Arguments

X	locfit object.
xlim	Plotting limits. Eg. $xlim=c(0,0,1,1)$ plots over the unit square in two dimensions. Default is bounding box of the data.
pv	Panel variables, to be varied within each panel of a plot. May be specified as a character vector, or variable numbers. There must be one or two panel variables; default is all variables in one or two dimensions; Variable 1 in three or more dimensions. May by specified using either variable numbers or names.
tv	Trellis variables, to be varied from panel to panel of the plot.
m	Controls the plot resolution (within panels, for trellis displays). Default is 100 points in one dimension; 40 points (per dimension) in two or more dimensions.
mtv	Number of points for trellis variables; default 6.
band	Type of confidence bands to add to the plot. Default is "none". Other choices include "global" for bands using a global variance estimate; "local" for bands using a local variance estimate and "pred" for prediction bands (at present, using a global variance estimate). To obtain the global variance estimate for a fit, use rv. This can be changed with rv< Confidence bands, by default, are 95%, based on normal approximations and neglecting bias. To change the critical value or confidence level, or to obtain simultaneous instead of pointwise confidence, the critical value stored on the fit must be changed. See the kappa0 and crit functions.
tr	Transformation function to use for plotting. Default is the inverse link function, or the identity function if derivatives are requested.
what	What to plot. See predict.locfit.
get.data	If TRUE, original data is added to the plot. Default: FALSE.
f3d	Force the locfit.3d class on the prediction object, thereby generating a trellis style plot. Default: FALSE, unless a $tv$ argument is' provided. Not available in R.
	Other arguments to plot.locfit.1d, plot.locfit.2d or plot.locfit.3d as appropriate.

# See Also

```
locfit, plot.locfit.1d, plot.locfit.2d, plot.locfit.3d, lines.locfit, predict.locfit, preplot.locfit\\
```

# Examples

```
x <- rnorm(100)
y <- dnorm(x) + rnorm(100) / 5
plot(locfit(y~x), band="global")
x <- cbind(rnorm(100), rnorm(100))
plot(locfit(~x), type="persp")</pre>
```

52 plot.locfit.2d

plot.locfit.1d Plo

Plot a one dimensional preplot.locfit object.

### **Description**

This function is not usually called directly. It will be called automatically when plotting a one-dimensional locfit or preplot.locfit object.

# Usage

```
## S3 method for class 'locfit.1d'
plot(x, add=FALSE, main="", xlab="default", ylab=x$yname,
   type="l", ylim, lty=1, col=1, ...)
```

### **Arguments**

```
    x One dimensional preplot.locfit object.
    add If TRUE, the plot will be added to the existing plot.
    main, xlab, ylab, type, ylim, lty, col
    Graphical parameters passed on to plot (only if add=FALSE).
    ... Additional graphical parameters to the plot function (only if add=FALSE).
```

#### See Also

```
locfit, plot.locfit, preplot.locfit
```

plot.locfit.2d

Plot a two-dimensional "preplot.locfit" object.

# **Description**

This function is not usually called directly. It will be called automatically when plotting onedimensional locfit or preplot.locfit objects.

### Usage

```
## S3 method for class 'locfit.2d'
plot(x, type="contour", main, xlab, ylab, zlab=x$yname, ...)
```

### **Arguments**

```
x Two dimensional preplot.locfit object.
type one of "contour", "persp", or "image".
main title for the plot.
xlab, ylab text labels for the x- and y-axes.
zlab if type="persp", the label for the z-axis.
... Additional arguments to the contour, persp or image functions.
```

plot.locfit.3d 53

### See Also

```
locfit, plot.locfit, preplot.locfit
```

plot.locfit.3d

Plot a high-dimensional "preplot.locfit" object using trellis displays.

# Description

This function plots cross-sections of a Locfit model (usually in three or more dimensions) using trellis displays. It is not usually called directly, but is invoked by plot.locfit.

The R libraries lattice and grid provide a partial (at time of writing) implementation of trellis. Currently, this works with one panel variable.

### Usage

### **Arguments**

X	"preplot.locfit" object.
main	title for the plot.
pv	Panel variables. These are the variables (either one or two) that are varied within each panel of the display.
tv	Trellis variables. These are varied from panel to panel of the display.
type	Type of display. When there are two panel variables, the choices are "contour", "level" and "persp".
pred.lab	label for the predictor variable.
resp.lab	label for the response variable.
crit	critical value for the confidence level.
	graphical parameters passed to xyplot or contourplot.

### See Also

```
plot.locfit, preplot.locfit
```

54 plot.scb

```
plot.preplot.locfit Plot a "preplot.locfit" object.
```

# Description

The plot.locfit() function is implemented, roughly, as a call to preplot.locfit(), followed by a call to plot.locfitpred(). For most users, there will be little need to call plot.locfitpred() directly.

# Usage

```
## S3 method for class 'preplot.locfit'
plot(x, pv, tv, ...)
```

### **Arguments**

```
x A preplot.locfit object, produced by preplot.locfit().
pv, tv, ... Other arguments to plot.locfit.1d, plot.locfit.2d or plot.locfit.3d as appropriate.
```

### See Also

```
locfit, plot.locfit, preplot.locfit, plot.locfit.1d, plot.locfit.2d, plot.locfit.3d.
```

plot.scb

Plot method for simultaneous confidence bands

# **Description**

Plot method for simultaneous confidence bands created by the scb function.

### Usage

```
## S3 method for class 'scb'
plot(x, add=FALSE, ...)
```

# Arguments

```
x scb object created by scb.add If TRUE, bands will be added to the existing plot.. . . Arguments passed to and from other methods.
```

# See Also

scb

plotbyfactor 55

### **Examples**

plotbyfactor

*x-y scatterplot, colored by levels of a factor.* 

# Description

Produces a scatter plot of x-y data, with different classes given by a factor f. The different classes are identified by different colours and/or symbols.

# Usage

```
plotbyfactor(x, y, f, data, col = 1:10, pch = "0", add = FALSE, lg,
    xlab = deparse(substitute(x)), ylab = deparse(substitute(y)),
    log = "", ...)
```

### **Arguments**

x	Variable for x axis.
У	Variable for y axis.
f	Factor (or variable for which as.factor() works).
data	data frame for variables x, y, f. Default: sys.parent().
col	Color numbers to use in plot. Will be replicated if shorter than the number of levels of the factor f. Default: 1:10.
pch	Vector of plot characters. Replicated if necessary. Default: "O".
add	If TRUE, add to existing plot. Otherwise, create new plot.
lg	Coordinates to place a legend. Default: Missing (no legend).
xlab, ylab	Axes labels.
log	Should the axes be in log scale? Use "x", "y", or "xy" to specify which axis to be in log scale.
	Other graphical parameters, labels, titles e.t.c.

# **Examples**

```
data(iris)
plotbyfactor(petal.wid, petal.len, species, data=iris)
```

56 predict.locfit

points.locfit

Add 'locfit' points to existing plot

### Description

This function shows the points at which the local fit was computed directly, rather than being interpolated. This can be useful if one is unsure of the validity of interpolation.

#### Usage

```
## S3 method for class 'locfit'
points(x, tr, ...)
```

### **Arguments**

x "locfit" object. Should be a model with one predictor.

tr Back transformation.

... Other arguments to the default points function.

#### See Also

```
locfit, plot.locfit, points
```

predict.locfit

Prediction from a Locfit object.

# Description

The locfit function computes a local fit at a selected set of points (as defined by the ev argument). The predict.locfit function is used to interpolate from these points to any other points. The method is based on cubic hermite polynomial interpolation, using the estimates and local slopes at each fit point.

The motivation for this two-step procedure is computational speed. Depending on the sample size, dimension and fitting procedure, the local fitting method can be expensive, and it is desirable to keep the number of points at which the direct fit is computed to a minimum. The interpolation method used by predict.locfit() is usually much faster, and can be computed at larger numbers of points.

### Usage

preplot.locfit 57

### **Arguments**

object Fitted object from locfit().

newdata Points to predict at. Can be given in several forms: vector/matrix; list, data frame.

se.fit If TRUE, standard errors are computed along with the fitted values.

where, what, band

arguments passed on to preplot.locfit.

Additional arguments to preplot.locfit.

### Value

If se.fit=T, a numeric vector of predictors. If se.fit=T, a list with components fit, se.fit and residual.scale.

# **Examples**

```
data(ethanol, package="locfit")
fit <- locfit(NOx ~ E, data=ethanol)
predict(fit,c(0.6,0.8,1.0))</pre>
```

preplot.locfit

Prediction from a Locfit object.

### **Description**

preplot.locfit can be called directly, although it is more usual to call plot.locfit or predict.locfit. The advantage of preplot.locfit is in S-Plus 5, where arithmetic and transformations can be performed on the "preplot.locfit" object.

plot(preplot(fit)) is essentially synonymous with plot(fit).

# Usage

```
## S3 method for class 'locfit'
preplot(object, newdata=NULL, where, tr=NULL, what="coef",
   band="none", get.data=FALSE, f3d=FALSE, ...)
```

#### **Arguments**

object Fitted object from locfit().

newdata Points to predict at. Can be given in several forms: vector/matrix; list, data

frame.

where An alternative to newdata. Choices include "grid" for the grid 1fmarg(object);

"data" for the original data points and "fitp" for the direct fitting points (ie.

no interpolation).

tr Transformation for likelihood models. Default is the inverse of the link function.

58 preplot.locfit.raw

what

What to compute predicted values of. The default, what="coef", works with the fitted curve itself. Other choices include "nlx" for the length of the weight diagram; "infl" for the influence function; "band" for the bandwidth; "degr" for the local polynomial degree; "lik" for the maximized local likelihood; "rdf" for the local residual degrees of freedom and "vari" for the variance function. The interpolation algorithm for some of these quantities is questionable.

band

Compute standard errors for the fit and include confidence bands on the returned object. Default is "none". Other choices include "global" for bands using a global variance estimate; "local" for bands using a local variance estimate and "pred" for prediction bands (at present, using a global variance estimate). To obtain the global variance estimate for a fit, use rv. This can be changed with rv<-. Confidence bands, by default, are 95%, based on normal approximations and neglecting bias. To change the critical value or confidence level, or to obtain simultaneous instead of pointwise confidence, the critical value stored on the fit must be changed. See the kappa0 and crit functions.

get.data

If TRUE, the original data is attached to the returned object, and added to the plot.

f3d

If TRUE, sets a flag that forces ploting using the trellis style. Not available in R.

. . .

arguments passed to and from other methods.

#### Value

An object with class "preplot.locfit", containing the predicted values and additional information used to construct the plot.

### See Also

```
locfit, predict.locfit, plot.locfit.
```

preplot.locfit.raw

Prediction from a Locfit object.

# **Description**

preplot.locfit.raw is an internal function used by predict.locfit and preplot.locfit. It should not normally be called directly.

#### **Usage**

```
## S3 method for class 'locfit.raw'
preplot(object, newdata, where, what, band, ...)
```

print.gcvplot 59

### **Arguments**

object Fitted object from locfit().

newdata New data points.

where Type of data provided in newdata.

what What to compute predicted values of.

band Compute standard errors for the fit and include confidence bands on the returned

object.

... Arguments passed to and from other methods.

# Value

A list containing raw output from the internal prediction routines.

### See Also

```
locfit, predict.locfit, preplot.locfit.
```

print.gcvplot

Print method for gcvplot objects

# **Description**

Print method for "gcvplot" objects. Actually, equivalent to plot.gcvplot(). scb function.

# Usage

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

# Arguments

x gcvplot object.

... Arguments passed to and from other methods.

#### See Also

```
gcvplot, plot.gcvplot summary.gcvplot
```

print.locfit

print.lfeval

Print the Locfit Evaluation Points.

# Description

Prints a matrix of the evaluation points from a locfit or lfeval structure.

# Usage

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

# **Arguments**

x A lfeval or locfit object

... Arguments passed to and from other methods.

### Value

Matrix of the fit points.

# See Also

```
lfeval, locfit, plot.lfeval
```

print.locfit

Print method for "locfit" object.

### **Description**

Prints a short summary of a "locfit" object.

# Usage

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

# **Arguments**

x locfit object.

... Arguments passed to and from other methods.

### See Also

locfit

print.preplot.locfit 61

```
print.preplot.locfit Print method for preplot.locfit objects.
```

# Description

Print method for objects created by the preplot.locfit function.

# Usage

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

# Arguments

```
x "preplot.locfit" object.
```

... Arguments passed to and from other methods.

#### See Also

```
preplot.locfit, predict.locfit
```

print.scb

Print method for simultaneous confidence bands

# Description

Print method for simultaneous confidence bands created by the scb function.

# Usage

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

# Arguments

x "scb" object created by scb.

... Arguments passed to and from other methods.

# See Also

scb

62 rbox

```
print.summary.locfit Print a Locfit summary object.
```

# Description

Print method for "summary.locfit" objects.

# Usage

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

# **Arguments**

x Object from summary.locfit.

... Arguments passed to and from methods.

### See Also

```
summary.locfit()
```

rbox

Local Regression, Likelihood and Density Estimation.

# Description

rbox() is used to specify a rectangular box evaluation structure for locfit.raw(). The structure begins by generating a bounding box for the data, then recursively divides the box to a desired precision.

# Usage

```
rbox(cut=0.8, type="tree", 11, ur)
```

# **Arguments**

type	If type="tree", the cells are recursively divided according to the bandwidths at each corner of the cell; see Chapter 11 of Loader (1999). If type="kdtree", the K-D tree structure used in Loess (Cleveland and Grosse, 1991) is used.
cut	Precision of the tree; a smaller value of cut results in a larger tree with more nodes being generated.
11	Lower left corner of the initial cell. Length should be the number of dimensions of the data provided to locfit.raw().
ur	Upper right corner of the initial cell. By default, 11 and ur are generated as the bounding box for the data.

regband 63

### References

Loader, C. (1999). Local Regression and Likelihood. Springer, New York.

Cleveland, W. and Grosse, E. (1991). Computational Methods for Local Regression. Statistics and Computing 1.

# **Examples**

```
data(ethanol, package="locfit")
plot.eval(locfit(NOx~E+C,data=ethanol,scale=0,ev=rbox(cut=0.8)))
plot.eval(locfit(NOx~E+C,data=ethanol,scale=0,ev=rbox(cut=0.3)))
```

regband

Bandwidth selectors for local regression.

# **Description**

Function to compute local regression bandwidths for local linear regression, implemented as a front end to locfit().

This function is included for comparative purposes only. Plug-in selectors are based on flawed logic, make unreasonable and restrictive assumptions and do not use the full power of the estimates available in Locfit. Any relation between the results produced by this function and desirable estimates are entirely coincidental.

### Usage

```
regband(formula, what = c("CP", "GCV", "GKK", "RSW"), deg=1, ...)
```

### **Arguments**

formula Model Formula (one predictor).

what Methods to use.

deg Degree of fit.

... Other Locfit options.

# Value

Vector of selected bandwidths.

64 right

residua	als	Inctit

Fitted values and residuals for a Locfit object.

### **Description**

residuals.locfit is implemented as a front-end to fitted.locfit, with the type argument set.

### Usage

```
## S3 method for class 'locfit'
residuals(object, data=NULL, type="deviance", ...)
```

# Arguments

object locfit object.

data The data frame for the original fit. Usually, shouldn't be needed.

type Type of fit or residuals to compute. The default is "fit" for fitted.locfit,

and "dev" for residuals.locfit. Other choices include "pear" for Pearson residuals; "raw" for raw residuals, "ldot" for likelihood derivative; "d2" for the deviance residual squared; lddot for the likelihood second derivative. Gen-

erally, type should only be used when what="coef".

... arguments passed to and from other methods.

### Value

A numeric vector of the residuals.

right

One-sided right smooth for a Locfit model.

### Description

The right() function is used in a locfit model formula to specify a one-sided smooth: when fitting at a point x, only data points with  $x_i \le x$  should be used. This can be useful in estimating points of discontinuity, and in cross-validation for forecasting a time series. right(x) is equivalent to lp(x,style="right").

When using this function, it will usually be necessary to specify an evaluation structure, since the fit is not smooth and locfit's interpolation methods are unreliable. Also, it is usually best to use deg=0 or deg=1, otherwise the fits may be too variable. If nearest neighbor bandwidth specification is used, it does not recognize right().

### Usage

```
right(x,...)
```

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# **Arguments**

x numeric variable.

... Other arguments to lp().

# See Also

```
locfit, left
```

# **Examples**

```
# compute left and right smooths
data(penny)
xev <- (1945:1988)+0.5
fitl <- locfit(thickness~left(year,h=10,deg=1), ev=xev, data=penny)
fitr <- locfit(thickness~right(year, h=10, deg=1), ev=xev, data=penny)
# plot the squared difference, to show the change points.
plot( xev, (predict(fitr, where="ev") - predict(fitl, where="ev"))^2 )</pre>
```

rν

Residual variance from a locfit object.

# **Description**

As part of the locfit fitting procedure, an estimate of the residual variance is computed; the rv function extracts the variance from the "locfit" object. The estimate used is the residual sum of squares (or residual deviance, for quasi-likelihood models), divided by the residual degrees of freedom.

For likelihood (not quasi-likelihood) models, the estimate is 1.0.

### Usage

```
rv(fit)
```

### Arguments

```
fit "locfit" object.
```

#### Value

Returns the residual variance estimate from the "locfit" object.

#### See Also

```
locfit, rv<-
```

66 scb

### **Examples**

```
data(ethanol)
fit <- locfit(NOx~E,data=ethanol)
rv(fit)</pre>
```

rva

Substitute variance estimate on a locfit object.

# Description

By default, Locfit uses the normalized residual sum of squares as the variance estimate when constructing confidence intervals. In some cases, the user may like to use alternative variance estimates; this function allows the default value to be changed.

# Usage

```
rv(fit) <- value</pre>
```

# **Arguments**

```
fit "locfit" object.
value numeric replacement value.
```

### See Also

```
locfit(), rv(), plot.locfit()
```

scb

Simultaneous Confidence Bands

# Description

scb is implemented as a front-end to locfit, to compute simultaneous confidence bands using the tube formula method and extensions, based on Sun and Loader (1994).

# Usage

```
scb(x, ..., ev = lfgrid(20), simul = TRUE, type = 1)
```

sjpi 67

### Arguments

x	A numeric vector or matrix of predictors (as in locfit.raw), or a model formula (as in locfit).
	Additional arguments to locfit.raw.
ev	The evaluation structure to use. See locfit.raw.
simul	Should the coverage be simultaneous or pointwise?
type	Type of confidence bands. type=0 computes pointwise 95% bands. type=1 computes basic simultaneous bands with no corrections. type=2,3,4 are the centered and corrected bands for parametric regression models listed in Table 3 of Sun, Loader and McCormick (2000).

### Value

A list containing the evaluation points, fit, standard deviations and upper and lower confidence bounds. The class is "scb"; methods for printing and ploting are provided.

#### References

Sun J. and Loader, C. (1994). Simultaneous confidence bands in linear regression and smoothing. *The Annals of Statistics* 22, 1328-1345.

Sun, J., Loader, C. and McCormick, W. (2000). Confidence bands in generalized linear models. *The Annals of Statistics* 28, 429-460.

# See Also

```
locfit, print.scb, plot.scb.
```

### **Examples**

sjpi

Sheather-Jones Plug-in bandwidth criterion.

### **Description**

Given a dataset and set of pilot bandwidths, this function computes a bandwidth via the plug-in method, and the assumed 'pilot' relationship of Sheather and Jones (1991). The S-J method chooses the bandwidth at which the two intersect.

The purpose of this function is to demonstrate the sensitivity of plug-in methods to pilot bandwidths and assumptions. This function does not provide a reliable method of bandwidth selection.

68 smooth.If

#### Usage

```
sjpi(x, a)
```

#### **Arguments**

- x data vector
- a vector of pilot bandwidths

### Value

A matrix with four columns; the number of rows equals the length of a. The first column is the plug-in selected bandwidth. The second column is the pilot bandwidths a. The third column is the pilot bandwidth according to the assumed relationship of Sheather and Jones. The fourth column is an intermediate calculation.

#### References

Sheather, S. J. and Jones, M. C. (1991). A reliable data-based bandwidth selection method for kernel density estimation. JRSS-B 53, 683-690.

# See Also

```
locfit, locfit.raw, lcvplot
```

#### **Examples**

smooth.1f

Local Regression, Likelihood and Density Estimation.

### **Description**

smooth.1f is a simple interface to the Locfit library. The input consists of a predictor vector (or matrix) and response. The output is a list with vectors of fitting points and fitted values. Most locfit.raw options are valid.

spence.15

### Usage

```
smooth.lf(x, y, xev=x, direct=FALSE, ...)
```

### **Arguments**

X	Vector (or matrix) of the independent variable(s).
у	Response variable. If omitted, $x$ is treated as the response and the predictor variable is $1:n$ .
xev	Fitting Points. Default is the data vector x.
direct	Logical variable. If T, local regression is performed directly at each fitting point. If F, the standard Locfit method combining fitting and interpolation is used.

... Other arguments to locfit.raw().

# Value

A list with components x (fitting points) and y (fitted values). Also has a call component, so update() will work.

#### See Also

```
locfit(), locfit.raw(), density.lf().
```

### **Examples**

spence.15

Spencer's 15 point graduation rule.

### **Description**

Spencer's 15 point rule is a weighted moving average operation for a sequence of observations equally spaced in time. The average at time t depends on the observations at times t-7,...,t+7.

Except for boundary effects, the function will reproduce polynomials up to degree 3.

### Usage

```
spence.15(y)
```

70 spence.21

#### **Arguments**

У

Data vector of observations at equally spaced points.

#### Value

A vector with the same length as the input vector, representing the graduated (smoothed) values.

#### References

Spencer, J. (1904). On the graduation of rates of sickness and mortality. Journal of the Institute of Actuaries 38, 334-343.

### See Also

```
spence. 21, spencer,
```

### **Examples**

```
data(spencer)
yy <- spence.15(spencer$mortality)
plot(spencer$age, spencer$mortality)
lines(spencer$age, yy)</pre>
```

spence.21

Spencer's 21 point graduation rule.

### Description

Spencer's 21 point rule is a weighted moving average operation for a sequence of observations equally spaced in time. The average at time t depends on the observations at times t-11,...,t+11.

Except for boundary effects, the function will reproduce polynomials up to degree 3.

# Usage

```
spence.21(y)
```

# **Arguments**

У

Data vector of observations at equally spaced points.

#### Value

A vector with the same length as the input vector, representing the graduated (smoothed) values.

#### References

Spencer, J. (1904). On the graduation of rates of sickness and mortality. Journal of the Institute of Actuaries 38, 334-343.

spencer 71

### See Also

```
spence.15, spencer,
```

# **Examples**

```
data(spencer)
yy <- spence.21(spencer$mortality)
plot(spencer$age, spencer$mortality)
lines(spencer$age, yy)</pre>
```

spencer

Spencer's Mortality Dataset

# Description

Observed mortality rates for ages 20 to 45.

# Usage

```
data(spencer)
```

# **Format**

Data frame with age and mortality variables.

# Source

Spencer (1904).

# References

Spencer, J. (1904). On the graduation of rates of sickness and mortality. Journal of the Institute of Actuaries 38, 334-343.

stamp

Stamp Thickness Dataset

# Description

Thicknesses of 482 postage stamps of the 1872 Hidalgo issue of Mexico.

# Usage

```
data(stamp)
```

72 summary.gcvplot

### **Format**

Data frame with thick (stamp thickness) and count (number of stamps) variables.

#### Source

Izenman and Sommer (1988).

### References

Izenman, A. J. and Sommer, C. J. (1988). Philatelic mixtures and multimodal densities. Journal of the American Statistical Association 73, 602-606.

store

Save S functions.

# **Description**

I've gotta keep track of this mess somehow!

# Usage

```
store(data=FALSE, grand=FALSE)
```

# **Arguments**

data whether data objects are to be saved.
grand whether everything is to be saved.

summary.gcvplot

Summary method for a gcvplot structure.

# **Description**

Computes a short summary for a generalized cross-validation plot structure

# Usage

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

# Arguments

object A gcvplot structure produced by a call to gcvplot, cpplot e.t.c. arugments to and from other methods.

summary.locfit 73

# Value

A matrix with two columns; one row for each fit computed in the gcvplot call. The first column is the fitted degrees of freedom; the second is the GCV or other criterion computed.

# See Also

```
locfit, gcv, gcvplot
```

# **Examples**

```
data(ethanol)
summary(gcvplot(NOx~E,data=ethanol,alpha=seq(0.2,1.0,by=0.05)))
```

summary.locfit

Print method for a locfit object.

# Description

Prints a short summary of a "locfit" object.

# Usage

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

# Arguments

object locfit object.

... arguments passed to and from methods.

# Value

A summary.locfit object, containg a short summary of the locfit object.

74 trimod

```
summary.preplot.locfit
```

Summary method for a preplot.locfit object.

### **Description**

Prints a short summary of a "preplot.locfit" object.

# Usage

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

### **Arguments**

```
object preplot.locfit object.
```

... arguments passed to and from methods.

#### Value

The fitted values from a preplot.locfit object.

trimod

Generated sample from a bivariate trimodal normal mixture

# **Description**

This is a random sample from a mixture of three bivariate standard normal components; the sample was used for the examples in Loader (1996).

# **Format**

Data frame with 225 observations and variables x0, x1.

### **Source**

Randomly generated in S.

### References

Loader, C. R. (1996). Local Likelihood Density Estimation. Annals of Statistics 24, 1602-1618.

xbar 75

xbar

 $Loc {\it fit\ Evaluation\ Structure}$ 

# Description

xbar() is an evaluation structure for locfit.raw(), evaluating the fit at a single point, namely, the average of each predictor variable.

# Usage

xbar()

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