

Package ‘gplm’

May 8, 2026

Type Package

Title Generalized Partial Linear Models (GPLM)

Version 0.7-4

Date 2016-08-28

Author Marlene Mueller

Maintainer Marlene Mueller <marlene.mueller@gmx.de>

Description Provides functions for estimating a generalized partial linear model, a semiparametric variant of the generalized linear model (GLM) which replaces the linear predictor by the sum of a linear and a nonparametric function.

Depends AER

License GPL (>= 2)

NeedsCompilation yes

Repository CRAN

Date/Publication 2016-08-29 08:18:41

Contents

bandwidth.scott	2
convol	3
create.grid	4
glm.inverse.link	5
glm.link	6
glm.ll	7
glm.lld	8
kbackfit	9
kde	10
kernel.constants	11
kernel.function	12
kgplm	13
kreg	16
sgplm1	17

bandwidth.scott	<i>Scott's rule of thumb</i>
-----------------	------------------------------

Description

Calculates Scott's rule of thumb bandwidth vector.

Usage

```
bandwidth.scott(x, kernel = "biweight", product = TRUE)
```

Arguments

x	n x d matrix, data
kernel	text string, see kernel.function
product	(if d>1) product or spherical kernel

Details

The default bandwidth vector is computed by Scott's rule of thumb for the Gaussian kernel and adapted to the chosen kernel function.

Value

d x 1 bandwidth vector used for calculation

Author(s)

Marlene Mueller

References

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

See Also

[kernel.function](#), [kde](#)

Examples

```
## two-dimensional data
n <- 1000
u <- runif(n)
thresh <- 0.4
x1 <- rnorm(n)*(u<thresh) +rnorm(n,mean=3)*(u>=thresh)
x2 <- rnorm(n)*(u<thresh) +rnorm(n,mean=9)*(u>=thresh)
bandwidth.scott( cbind(x1,x2) )
```

convol	<i>Kernel convolution</i>
--------	---------------------------

Description

Calculates the convolution of data with a kernel function.

Usage

```
convol(x, h = 1, grid = NULL, y = 1, w = 1, p = 2, q = 2,
       product = TRUE, sort = TRUE)
```

Arguments

x	n x d matrix, data
h	scalar or 1 x d, bandwidth(s)
grid	m x d matrix, where to calculate the convolution (default = x)
y	n x c matrix, optional responses
w	scalar or n x 1 or 1 x m or n x m, optional weights
p	integer or text, see kernel.function
q	integer, see kernel.function
product	(if d>1) product or spherical kernel
sort	logical, TRUE if data need to be sorted

Details

The kernel convolution which is calculated is $\sum_i K_h(x_i - grid_j) y_i w_{ij}$ for $i = 1, \dots, n$ and $j = 1, \dots, m$. The kernel function is determined by the kernel parameters p and q, see [kernel.function](#). The default kernel is the biweight (quartic) kernel function. Note that the DLL requires the data matrix to be sorted by its first column.

Value

m x c matrix

Author(s)

Marlene Mueller

See Also

[kernel.function](#), [kde](#), [kreg](#)

Examples

```
n <- 100
x <- rnorm(n)
convol(x, h=0.8, grid=-3:3)/n ## estimates density of x at points -3:3
```

`create.grid`*Create a grid for kernel estimation*

Description

Helps to define a grid for kernel density or regression estimates (univariate or multivariate).

Usage

```
create.grid(grid.list, sort=TRUE)
```

Arguments

<code>grid.list</code>	list of 1-dimensional vectors containing the grid values for each dimension
<code>sort</code>	sort the vectors (can be set to FALSE if vectors are already sorted in ascending order)

Details

This function allows easily to define grids for the "gplm" package. If the data are d-dimensional and the grid vector lengths are n1, ... nd, then the output is a (n1*...*nd) x d matrix with each row corresponding to one d-dimensional data point at which the function estimate is to be calculated.

Value

m x d grid matrix

Author(s)

Marlene Mueller

See Also

[expand.grid](#), [kde](#), [kreg](#)

Examples

```
v1 <- 1:5
v2 <- 3:1
grid <- create.grid(list(v1,v2))

x <- matrix(rnorm(60),30,2)
v1 <- seq(min(x[,1]),max(x[,1]),length=10)
v2 <- seq(min(x[,2]),max(x[,2]),length=5)
grid <- create.grid(list(v1,v2))
```

glm.inverse.link	<i>Link function for GLM</i>
------------------	------------------------------

Description

Defines the link function for a GLM.

Usage

```
glm.inverse.link(mu, family="gaussian", link="identity", k=1)
```

Arguments

mu	n x 1, linear predictors
family	text string, family of distributions (e.g. "gaussian" or "bernoulli", see details for glm.ll)
link	text string, link function (depending on family, see details for glm.ll)
k	integer > 0, parameter for the negative binomial

Value

n x 1, vector eta (predictors)

Author(s)

Marlene Mueller

See Also

[glm.ll](#), [glm.lld](#), [glm.link](#)

Examples

```
glm.inverse.link(c(0.25,0.5), family="bernoulli", link="logit")
```

glm.link *(Inverse) Link function for GLM*

Description

Defines the inverse link function for a GLM.

Usage

```
glm.link(eta, family="gaussian", link="identity", k=1)
```

Arguments

eta	n x 1, linear predictors
family	text string, family of distributions (e.g. "gaussian" or "bernoulli", see details for glm.ll)
link	text string, link function (depending on family, see details for glm.ll)
k	integer > 0, parameter for the negative binomial

Value

n x 1, vector mu (responses)

Author(s)

Marlene Mueller

See Also

[glm.ll](#), [glm.lld](#), [glm.inverse.link](#)

Examples

```
glm.link(c(-1,2), family="bernoulli", link="logit")
```

`glm.ll`*Log-likelihood for GLM*

Description

Calculates the log-likelihood function of a GLM. Currently only the gaussian and the bernoulli family are implemented.

Usage

```
glm.ll(mu, y, phi=1, family="gaussian", k=1)
```

Arguments

<code>mu</code>	<code>n x 1</code> , predicted regression function
<code>y</code>	<code>n x 1</code> , responses
<code>phi</code>	scalar, nuisance parameter (σ^2 for the gaussian and inverse gaussian families, ν for the gamma family)
<code>family</code>	text string, family of distributions (e.g. "gaussian" or "bernoulli", see details below)
<code>k</code>	integer > 0 , parameter for the negative binomial

Details

Implemented are the "gaussian" family (with links "identity" and "log"), the "bernoulli" family (with links "logit" and "probit"), the "gamma" family (with link "inverse"), the "poisson" family (with link "log"), the "inverse.gaussian" family (with link "inverse.squared") and the "negative.binomial" (with its canonical "log" type link).

The default value `k=1` leads to the geometric distribution (as a special case of the negative binomial).

Value

log-likelihood value

Author(s)

Marlene Mueller

See Also

[glm.lld](#), [glm.link](#)

Examples

```
glm.ll(rep(0.4,2), c(0,1), family="bernoulli")
```

`glm.lld`*Log-likelihood derivatives for GLM*

Description

Computes first and second derivatives of the individual log-likelihood with respect to the linear predictor. Currently only the gaussian (with identity link) and the bernoulli family (with logit and probit links) are implemented.

Usage

```
glm.lld(eta, y, family="gaussian", link="identity", k=1)
```

Arguments

<code>eta</code>	<code>n x 1</code> , linear predictors
<code>y</code>	<code>n x 1</code> , responses
<code>family</code>	text string, family of distributions (e.g. "gaussian" or "bernoulli", see details for glm.ll)
<code>link</code>	text string, link function (depending on family, see details for glm.ll)
<code>k</code>	integer > 0 , parameter for the negative binomial

Details

See details for [glm.ll](#).

Value

List with components:

<code>l11</code>	<code>n x 1</code> , vector of first derivatives
<code>l12</code>	<code>n x 1</code> , vector of second derivatives
<code>l11.2</code>	<code>n x 1</code> , ratio <code>l11/l12</code>

Author(s)

Marlene Mueller

See Also

[glm.ll](#), [glm.link](#)

Examples

```
glm.lld(c(-1,2), c(0,1), family="bernoulli", link="logit")
```

kbackfit

*Backfitting for an additive model using kernel regression***Description**

Implements kernel-based backfitting in an additive model, optional with a partial linear term.

Usage

```
kbackfit(t, y, h, x = NULL, grid = NULL, weights.conv = 1,
         offset = 0, method = "generic",
         max.iter = 50, eps.conv = 1e-04, m.start = NULL,
         kernel = "biweight")
```

Arguments

y	n x 1 vector, responses
t	n x q matrix, data for nonparametric part
h	scalar or 1 x q, bandwidth(s)
x	optional, n x p matrix, data for linear part
grid	m x q matrix, where to calculate the nonparametric function (default = t)
weights.conv	weights for convergence criterion
offset	offset
method	one of "generic", "linit" or "modified"
max.iter	maximal number of iterations
eps.conv	convergence criterion
m.start	n x q matrix, start values for m
kernel	text string, see kernel.function

Value

List with components:

c	constant
b	p x 1 vector, linear coefficients
m	n x q matrix, nonparametric marginal function estimates
m.grid	m x q matrix, nonparametric marginal function estimates on grid
rss	residual sum of squares

Author(s)

Marlene Mueller

See Also

[kernel.function](#), [kreg](#)

kde *Kernel density estimation*

Description

Calculates a kernel density estimate (univariate or multivariate).

Usage

```
kde(x, bandwidth = NULL, grid = TRUE, kernel = "biweight",
    product = TRUE, sort = TRUE)
```

Arguments

x	n x d matrix, data
bandwidth	scalar or 1 x d, bandwidth(s)
grid	logical or m x d matrix (where to calculate the density)
kernel	text string, see kernel.function
product	(if d>1) product or spherical kernel
sort	logical, TRUE if data need to be sorted

Details

The kernel density estimator is calculated as $\frac{1}{n} \sum_i K_h(x_i - grid_j)$ for $i = 1, \dots, n$ and $j = 1, \dots, m$. The default bandwidth vector is computed by Scott's rule of thumb (adapted to the chosen kernel function).

Value

List with components:

x	m x d matrix, where density has been calculated
y	m x 1 vector, density estimates
bandwidth	bandwidth vector used for calculation
rearrange	if sort=TRUE, index to rearrange x and y to its original order.

Author(s)

Marlene Mueller

See Also

[kernel.function](#), [convol](#), [kreg](#)

Examples

```

n <- 1000
x <- rnorm(n)
plot(kde(x), type="l")

## mixed normal data
n <- 1000
u <- runif(n)
thresh <- 0.4
x <- rnorm(n)*(u<thresh) +rnorm(n,mean=3)*(u>=thresh)
h <- 1
fh <- kde(x,bandwidth=h)
plot(kde(x,bandwidth=h),type="l",lwd=2); rug(x)
lines(kde(x,bandwidth=h*1.2),col="red")
lines(kde(x,bandwidth=h*1.4),col="orange")
lines(kde(x,bandwidth=h/1.2),col="blue")
lines(kde(x,bandwidth=h/1.4),col="cyan")

## two-dimensional data
n <- 1000
u <- runif(n)
thresh <- 0.4
x1 <- rnorm(n)*(u<thresh) +rnorm(n,mean=3)*(u>=thresh)
x2 <- rnorm(n)*(u<thresh) +rnorm(n,mean=9)*(u>=thresh)

grid1 <- seq(min(x1),max(x1),length=20) ## grid for x1
grid2 <- seq(min(x2),max(x2),length=25) ## grid for x2

fh <- kde( cbind(x1,x2), grid=create.grid(list(grid1,grid2)) )
o <- order(fh$x[,2],fh$x[,1])
density <- (matrix(fh$y[o],length(grid1),length(grid2)))

par(mfrow=c(2,2))
plot(kde(x1),type="l",main="x1"); rug(x1)
plot(kde(x2),type="l",main="x2"); rug(x2)
persp(grid1,grid2,density,main="KDE",
       theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
contour(grid1,grid2,density, main="KDE Contours")
points(x1,x2,col="red",pch=18,cex=0.5)
par(mfrow=c(1,1))

```

kernel.constants

Kernel constants

Description

Calculates several constants of a (product) kernel function.

Usage

```
kernel.constants(kernel = "biweight", d = 1, product = TRUE)
```

Arguments

kernel	text string, see kernel.function
d	integer (dimension of the kernel)
product	(if d>1) product or spherical kernel

Details

The constants which are calculated are the second moment, the square norm and the canonical bandwidth of the kernel (only the two latter terms depend on the dimension d).

Value

List with components:

m2	second moment
c2	square norm
d0	canonical bandwidth

Author(s)

Marlene Mueller

See Also

[kernel.function](#)

Examples

```
kernel.constants()           ## default (biweight), d=1
kernel.constants("epanechnikov",1) ## epanechnikov, d=1
kernel.constants("epanechnikov",2) ## product epanechnikov, d=2
```

kernel.function *Kernel function*

Description

Calculates several kernel functions (uniform, triangle, epanechnikov, biweight, triweight, gaussian).

Usage

```
kernel.function(u, kernel = "biweight", product = TRUE)
```

Arguments

u	n x d matrix
kernel	text string
product	(if d>1) product or spherical kernel

Details

The kernel parameter is a text string specifying the univariate kernel function which is either the gaussian pdf or proportional to $(1 - |u|^p)^q$. Possible text strings are "triangle" (p=q=1), "uniform" (p=1, q=0), "epanechnikov" (p=2, q=1), "biweight" or "quartic" (p=q=2), "triweight" (p=2, q=3), "gaussian" or "normal" (gaussian pdf).

The multivariate kernels are obtained by a product of univariate kernels $K(u_1)...K(u_d)$ or by a spherical (radially symmetric) kernel proportional to $K(||u||)$. (The resulting kernel is a density, i.e. integrates to 1.)

Value

n x 1 vector of kernel weights

Author(s)

Marlene Mueller

Examples

```
kernel.function(0)                ## default (biweight)
kernel.function(0, kernel="epanechnikov") ## epanechnikov
kernel.function(0, kernel="gaussian")   ## equals dnorm(0)
```

kgplm

Generalized partial linear model

Description

Fits a generalized partial linear model (kernel-based) using the (generalized) Speckman estimator or backfitting (in the generalized case combined with local scoring) for two additive component functions.

Usage

```
kgplm(x, t, y, h, family, link,
      b.start=NULL, m.start=NULL, grid = NULL,
      offset = 0, method = "speckman", sort = TRUE, weights = 1,
      weights.trim = 1, weights.conv = 1, max.iter = 25, eps.conv = 1e-8,
      kernel = "biweight", kernel.product = TRUE, verbose = FALSE)
```

Arguments

x	n x p matrix, data for linear part
y	n x 1 vector, responses
t	n x q matrix, data for nonparametric part
h	scalar or 1 x q, bandwidth(s)

family	text string, family of distributions (e.g. "gaussian" or "bernoulli", see details for glm.l1)
link	text string, link function (depending on family, see details for glm.l1)
b.start	p x 1 vector, start values for linear part
m.start	n x 1 vector, start values for nonparametric part
grid	m x q matrix, where to calculate the nonparametric function (default = t)
offset	offset
method	"speckman" or "backfit"
sort	logical, TRUE if data need to be sorted
weights	binomial weights
weights.trim	trimming weights for fitting the linear part
weights.conv	weights for convergence criterion
max.iter	maximal number of iterations
eps.conv	convergence criterion
kernel	text string, see kernel.function
kernel.product	(if p>1) product or spherical kernel
verbose	print additional convergence information

Value

List with components:

b	p x 1 vector, linear coefficients
b.cov	p x p matrix, linear coefficients
m	n x 1 vector, nonparametric function estimate
m.grid	m x 1 vector, nonparametric function estimate on grid
it	number of iterations
deviance	deviance
df.residual	approximate degrees of freedom (residuals)
aic	Akaike's information criterion

Author(s)

Marlene Mueller

References

- Mueller, M. (2001). Estimation and testing in generalized partial linear models – A comparative study. *Statistics and Computing*, 11:299–309.
- Hastie, T. and Tibshirani, R. (1990). *Generalized Additive Models*. London: Chapman and Hall.

See Also

[kernel.function](#), [kreg](#)

Examples

```
## data
n <- 1000; b <- c(1,-1); rho <- 0.7
m <- function(t){ 1.5*sin(pi*t) }
x1 <- runif(n,min=-1,max=1); u <- runif(n,min=-1,max=1)
t <- runif(n,min=-1,max=1); x2 <- round(m(rho*t + (1-rho)*u))
x <- cbind(x1,x2)
y <- x %*% b + m(t) + rnorm(n)

## partial linear model (PLM)
gh <- kgplm(x,t,y,h=0.25,family="gaussian",link="identity")
o <- order(t)
plot(t[o],m(t[o]),type="l",col="green")
lines(t[o],gh$m[o]); rug(t)

## partial linear probit model (GPLM)
y <- (y>0)
gh <- kgplm(x,t,y,h=0.25,family="bernoulli",link="probit")

o <- order(t)
plot(t[o],m(t[o]),type="l",col="green")
lines(t[o],gh$m[o]); rug(t)

## data with two-dimensional m-function
n <- 1000; b <- c(1,-1); rho <- 0.7
m <- function(t1,t2){ 1.5*sin(pi*t1)+t2 }
x1 <- runif(n,min=-1,max=1); u <- runif(n,min=-1,max=1)
t1 <- runif(n,min=-1,max=1); t2 <- runif(n,min=-1,max=1)
x2 <- round( m( rho*t1 + (1-rho)*u , t2 ) )
x <- cbind(x1,x2); t <- cbind(t1,t2)
y <- x %*% b + m(t1,t2) + rnorm(n)

## partial linear model (PLM)
grid1 <- seq(min(t[,1]),max(t[,1]),length=20)
grid2 <- seq(min(t[,2]),max(t[,2]),length=25)
grid <- create.grid(list(grid1,grid2))

gh <- kgplm(x,t,y,h=0.5,grid=grid,family="gaussian",link="identity")

o <- order(grid[,2],grid[,1])
est.m <- (matrix(gh$m.grid[o],length(grid1),length(grid2)))
orig.m <- outer(grid1,grid2,m)
par(mfrow=c(1,2))
persp(grid1,grid2,orig.m,main="Original Function",
       theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
persp(grid1,grid2,est.m,main="Estimated Function",
       theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
par(mfrow=c(1,1))
```

kreg *Kernel regression*

Description

Calculates a kernel regression estimate (univariate or multivariate).

Usage

```
kreg(x, y, bandwidth = NULL, grid = TRUE, kernel = "biweight",
     product = TRUE, sort = TRUE)
```

Arguments

x	n x d matrix, data
y	n x 1 vector, responses
bandwidth	scalar or 1 x d, bandwidth(s)
grid	logical or m x d matrix (where to calculate the regression)
kernel	text string, see kernel.function
product	(if d>1) product or spherical kernel
sort	logical, TRUE if data need to be sorted

Details

The estimator is calculated by Nadaraya-Watson kernel regression. Future extension to local linear (d>1) or polynomial (d=1) estimates is planned. The default bandwidth is computed by Scott's rule of thumb for kde (adapted to the chosen kernel function).

Value

List with components:

x	m x d matrix, where regression has been calculated
y	m x 1 vector, regression estimates
bandwidth	bandwidth used for calculation
df.residual	approximate degrees of freedom (residuals)
rearrange	if sort=TRUE, index to rearrange x and y to its original order.

Author(s)

Marlene Mueller

See Also

[kernel.function](#), [convol](#), [kde](#)

Examples

```

n <- 1000
x <- rnorm(n)
m <- sin(x)
y <- m + rnorm(n)
plot(x,y,col="gray")
o <- order(x); lines(x[o],m[o],col="green")
lines(kreg(x,y),lwd=2)

## two-dimensional
n <- 100
x <- 6*cbind(runif(n), runif(n))-3
m <- function(x1,x2){ 4*sin(x1) + x2 }
y <- m(x[,1],x[,2]) + rnorm(n)
mh <- kreg(x,y)##,bandwidth=1

grid1 <- unique(mh$x[,1])
grid2 <- unique(mh$x[,2])
est.m <- t(matrix(mh$y,length(grid1),length(grid2)))
orig.m <- outer(grid1,grid2,m)
par(mfrow=c(1,2))
persp(grid1,grid2,orig.m,main="Original Function",
      theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
persp(grid1,grid2,est.m,main="Estimated Function",
      theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
par(mfrow=c(1,1))

## now with normal x, note the boundary problem,
## which can be somewhat reduced by a gaussian kernel
n <- 1000
x <- cbind(rnorm(n), rnorm(n))
m <- function(x1,x2){ 4*sin(x1) + x2 }
y <- m(x[,1],x[,2]) + rnorm(n)
mh <- kreg(x,y)##,p="gaussian")

grid1 <- unique(mh$x[,1])
grid2 <- unique(mh$x[,2])
est.m <- t(matrix(mh$y,length(grid1),length(grid2)))
orig.m <- outer(grid1,grid2,m)
par(mfrow=c(1,2))
persp(grid1,grid2,orig.m,main="Original Function",
      theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
persp(grid1,grid2,est.m,main="Estimated Function",
      theta=30,phi=30,expand=0.5,col="lightblue",shade=0.5)
par(mfrow=c(1,1))

```

Description

Fits a generalized partial linear model (based on smoothing spline) using the (generalized) Speckman estimator or backfitting (in the generalized case combined with local scoring) for two additive component functions. In contrast to [kgplm](#), this function can be used only for a 1-dimensional nonparametric function.

Usage

```
sgplm1(x, t, y, spar, df=4, family, link,
       b.start=NULL, m.start=NULL, grid = NULL, offset = 0,
       method = "speckman", weights = 1, weights.trim = 1,
       weights.conv = 1, max.iter = 25, eps.conv = 1e-8,
       verbose = FALSE, ...)
```

Arguments

x	n x p matrix, data for linear part
y	n x 1 vector, responses
t	n x 1 matrix, data for nonparametric part
spar	scalar smoothing parameter, as in smooth.spline
df	scalar equivalent number of degrees of freedom (trace of the smoother matrix), as in smooth.spline
family	text string, family of distributions (e.g. "gaussian" or "bernoulli", see details for glm.l1)
link	text string, link function (depending on family, see details for glm.l1)
b.start	p x 1 vector, start values for linear part
m.start	n x 1 vector, start values for nonparametric part
grid	m x q matrix, where to calculate the nonparametric function (default = t)
offset	offset
method	"speckman" or "backfit"
weights	binomial weights
weights.trim	trimming weights for fitting the linear part
weights.conv	weights for convergence criterion
max.iter	maximal number of iterations
eps.conv	convergence criterion
verbose	print additional convergence information
...	further parameters to be passed to smooth.spline

Value

List with components:

b	p x 1 vector, linear coefficients
b.cov	p x p matrix, linear coefficients
m	n x 1 vector, nonparametric function estimate
m.grid	m x 1 vector, nonparametric function estimate on grid
it	number of iterations
deviance	deviance
df.residual	approximate degrees of freedom (residuals)
aic	Akaike's information criterion

Note

This function is mainly implemented for comparison. It is not really optimized for performance, however since it is spline-based, it should be sufficiently fast. Nevertheless, there might be several possibilities to improve for speed, in particular I guess that the sorting that `smooth.spline` performs in every iteration is slowing down the procedure quite a bit.

Author(s)

Marlene Mueller

References

Mueller, M. (2001) Estimation and testing in generalized partial linear models – A comparative study. *Statistics and Computing*, 11:299–309.

Hastie, T. and Tibshirani, R. (1990) *Generalized Additive Models*. London: Chapman and Hall.

See Also

[kgplm](#)

Examples

```
## generate data
n <- 1000; b <- c(1,-1); rho <- 0.7
mm <- function(t){ 1.5*sin(pi*t) }
x1 <- runif(n,min=-1,max=1); u <- runif(n,min=-1,max=1)
t <- runif(n,min=-1,max=1); x2 <- round(mm(rho*t + (1-rho)*u))
x <- cbind(x1,x2)
y <- x %*% b + mm(t) + rnorm(n)

## fit partial linear model (PLM)
k.plm <- kgplm(x,t,y,h=0.35,family="gaussian",link="identity")
s.plm <- sgplm1(x,t,y,spar=0.95,family="gaussian",link="identity")

o <- order(t)
```

```
ylim <- range(c(mm(t[o]),k.plm$m,s.plm$m),na.rm=TRUE)
plot(t[o],mm(t[o]),type="l",ylim=ylim)
lines(t[o],k.plm$m[o], col="green")
lines(t[o],s.plm$m[o], col="blue")
rug(t); title("Kernel PLM vs. Spline PLM")

## fit partial linear probit model (GPLM)
y <- (y>0)
k.gplm <- kgplm(x,t,y,h=0.35,family="bernoulli",link="probit")
s.gplm <- sgplm1(x,t,y,spar=0.95,family="bernoulli",link="probit")

o <- order(t)
ylim <- range(c(mm(t[o]),k.gplm$m,s.gplm$m),na.rm=TRUE)
plot(t[o],mm(t[o]),type="l",ylim=ylim)
lines(t[o],k.gplm$m[o], col="green")
lines(t[o],s.gplm$m[o], col="blue")
rug(t); title("Kernel GPLM vs. Spline GPLM (Probit)")
```

Index

* **smooth**

- bandwidth.scott, 2
- convol, 3
- create.grid, 4
- glm.inverse.link, 5
- glm.link, 6
- glm.ll, 7
- glm.lld, 8
- kbackfit, 9
- kde, 10
- kernel.constants, 11
- kernel.function, 12
- kgplm, 13
- kreg, 16
- sgplm1, 17

bandwidth.scott, 2

convol, 3, 10, 16

create.grid, 4

expand.grid, 4

glm.inverse.link, 5, 6

glm.link, 5, 6, 7, 8

glm.ll, 5, 6, 7, 8, 14, 18

glm.lld, 5–7, 8

kbackfit, 9

kde, 2–4, 10, 16

kernel.constants, 11

kernel.function, 2, 3, 9, 10, 12, 12, 14–16

kgplm, 13, 18, 19

kreg, 3, 4, 9, 10, 15, 16

sgplm1, 17

smooth.spline, 18, 19