## Package 'JOPS'

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Maintainer Paul Eilers [p.eilers@erasmusmc.nl](mailto:p.eilers@erasmusmc.nl)
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Author Paul Eilers [aut, cre], Brian Marx [aut], Bin Li [aut], Jutta Gampe [aut], Maria Xose Rodriguez-Alvarez [aut]
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bbase Compute a B-spline basis matrix

## Description

Compute a B-spline basis matrix using evenly spaced knots.

## Usage

bbase $(x, x l=\min (x), x r=\max (x), n s e g=10, b d e g=3)$

## Arguments

$x \quad a$ vector of argument values, at which the $B$-spline basis functions are to be evaluated.
$x l \quad$ the lower limit of the domain of $x$; default is $\min (x)$.
$x r \quad$ the upper limit of the domain of $x$; default is $\max (x)$.
nseg the number of equally sized segments between xl and xr ; default is 10 .
bdeg the degree of the splines, usually 1,2 , or 3 (default).

## Details

If $x l$ is larger than $\min (x)$, it will be adjusted to $\min (x)$ and a warning wil be given. If $x r$ is smaller than $\max (x)$, it will be adjusted to $\max (x)$ and a warning wil be given. The values of the design parameters $x, x l, x r$, ndeg, bdeg and type $=$ 'bbase' are added to the list of attributes of the matrix.

## Value

A matrix with length $(x)$ rows and nseg + bdeg columns.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

Eilers, P.H.C. and Marx, B.D. (1996). Flexible smoothing with B-splines and penalties (with comments and rejoinder), Statistical Science, 11: 89-121.

Eilers, P.H.C. and B.D. Marx (2010). Splines, knots and penalties. Wiley Interdisciplinary Reviews: Computational Statistics. Wiley: NY. DOI: 10.1002/wics. 125

## Examples

```
# Compute and plot a B-spline basis matrix
x = seq(0, 360, by = 2)
B = bbase(x, 0, 360, nseg = 8, bdeg = 3)
matplot(x, B, type = 'l', lty = 1, lwd = 2, xlab = 'x', ylab = '')
```


## binit

 Translated number vector to bin index.
## Description

Translates number vector to bin index, given lower and upper limits of the domain and number of bins. A support function for (smoothing) histograms.

## Usage

binit(x, xmin $=\min (x), x \max =\max (x), \operatorname{nbin}=100)$

## Arguments

x
$x$ min the lower limit of the domain.
$x \max \quad$ the upper limit of the domain.
nbin the number of bins (default=100).

## Value

A list with components:

| xbin | a vector of length $(x)$ with elements giving the bin index. |
| :--- | :--- |
| xgrid | a vector of length(nbin) with the midpoints of the bins. |
| nbin | the number of bins. |

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

```
bone_data Spinal bone relative mineral density
```


## Description

Relative spinal bone mineral density measurements on 261 North American adolescents. Each value is the difference in spnbmd taken on two consecutive visits, divided by the average. The age is the average age over the two visits.

## Usage

data(bone_data)

## Format

A dataframe with four columns:
idnum ID of the child
age age
gender male or female
spnbmd Relative Spinal bone mineral density.

## Source

https://web.stanford.edu/~hastie/ElemStatLearn/datasets/bone.data

## References

Bachrach, L.K., Hastie, T., Wang, M.-C., Narasimhan, B., Marcus, R. (1999). Bone Mineral Acquisition in Healthy Asian, Hispanic, Black and Caucasian Youth. A Longitudinal Study. J Clin Endocrinol Metab 84, 4702-12.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Description

Computes a circular B-spline basis matrix using evenly spaced knots.

## Usage

$\operatorname{cbase}(x, x l=\min (x), x r=\max (x), n s e g=10, b d e g=3)$

## Arguments

x
$x l \quad$ the lower limit of the domain of $x$; default is $\min (x)$.
$x r \quad$ the upper limit of the domain of $x$; default is $\max (x)$.
nseg the number of B-spline segments (default 10) between xl and xr .
bdeg the degree of the basis, usually 1,2 , or 3 (default).

## Details

If $x l$ is larger than $\min (x)$, it wil be adjusted to $\min (x)$ and a warning wil be given. If $x r$ is smaller than $\max (x)$, it wil be adjusted to $\max (x)$ and a warning wil be given.
The design parameters $x, x l, x r, n d e g$, bdeg and type $=$ 'cbase $'$ are added to the list of attributes.
In a circular basis, the B-splines are wrapped around the boundaries of the domain. Use a circular basis for data like directions or angles. It should be combined with a circular penalty matrix, as computed by cdiff().

## Value

A matrix with length ( $x$ ) rows and nseg columns.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

\# Compute and plot a circular B-spline basis matrix
$x=\operatorname{seq}(0,360$, by $=2)$
$B=$ cbase ( $x, 0,360$, nseg $=8$, bdeg = 3)
matplot(x, B, type = 'l', lty = 1, lwd = 2, xlab = 'x', ylab = '')
title('Note how the ends connect smoothly meet at boundaries' )

$$
\text { cdiff } \quad \text { Compute a second order circular differencing matrix }
$$

## Description

Compute difference matrix used for circular penalities.

## Usage

cdiff( $n$ )

## Arguments

$\mathrm{n} \quad$ number of rows (and columns) of the square differencing matrix.

## Value

A square matrix with n rows and columns.

## Author(s)

Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
# Compare standard and circular differencing matrix
n = 8
D1 = diff(diag(n), diff = 2)
D2 = cdiff(n)
oldpar = par(no.readonly = TRUE)
on.exit(par(oldpar))
par(mfrow = c(1, 2))
image(t(D1))
```

```
title('Linear differencing matrix')
image(t(D2))
title('Circular differencing matrix')
```

    CGHsim Simulation of CGH data
    
## Description

A crude simulation of comparative genomic hybridization (CGH) data.

## Usage

data(CGHsim)

## Format

A data frame with 400 rows and two columns:
y $\log \mathrm{R}$ ratio
$x$ Genomic position (but in fact the row number).

## Source

The simulation program could not be located anymore. But the data have a very simple structure.

```
clone_base Clone a B-spline basis for new x
```


## Description

Extract basis parameters from an existing B-splines basis matrix, and use them for computing a new basis at new values of $x$.

## Usage

clone_base(B, x)

## Arguments

B
x
a B-splines basis matrix, computed with bbase() or cbase(). a vector of new argument values.

## Details

If values in $x$ are outside the domain used for computing $B$, they will be discarded, with a warning.

## Value

A matrix with number of rows=length(xnew).

## Author(s)

Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
x = seq(0, 10, length = 20)
n = length(x)
y = sin(x/2) + rnorm(n) * 0.2
B = bbase(x)
nb = ncol(B)
D = diff(diag(nb), diff = 2)
lambda = 1
a = solve(t(B) %*% B + lambda * t(D)%*% D, t(B) %*% y)
# Clone basis on finer grid
xg = seq(0, 10, length = 200)
Bg = clone_base(B, xg)
yg = Bg %*% a
plot(x, y)
lines(xg, yg, col = 'blue')
```

Complaints Environmental complaints from the Rijnomond area of The Netherlands

## Description

Environmental complaints about odors from the Rijnmond region (near Rotterdam in the Netherlands) in 1988.

## Usage

data(Complaints)

## Format

A dataframe with two columns:
freq The daily number of complaints.
count The number of days the specific complaint frequency occurred.

## Details

In 1988, the Rijnmond Environmental Agency registered approximately 20,000 complaints about odors from regional inhabitants.

## Source

Personal information from Paul Eilers.

## Examples

plot(Complaints\$freq, Complaints\$count, type = 'h', xlab = 'Number of complaints per day', ylab = 'Frequency')
count2d Create a matrix of counts.

## Description

Count the number of occurrences of pairs of positive integers in two vectors, producing a matrix.

## Usage

count $2 \mathrm{~d}(\mathrm{xb}, \mathrm{yb}, \mathrm{nb})$

## Arguments

| xb | a vector of integers. |
| :--- | :--- |
| yb | a vector of integers. |
| nb | a vector of length 2 that provides the number of bins for the 2D histogram on $x$ <br> and $y$. |

## Details

This function builds a two-dimensional histogram, based on two two vectors of bin numbers (obtained with binit). Rows where $x[i]>n b[1]$ or $y[i]>n b[2]$ are discarded without a warning.

## Value

A matrix with nb [1] rows and nb [2] columns with counts. It serves as the input for two-dimensional histogram smoothing.

## Description

Calculates the deviance and returns the ML estimated dispersion parameter for a variety of response distributions for P -spline fitting within the GLM framework.

## Usage

dev_calc $($
family = "gaussian",
$y$,
mu ,
m_binomial $=0 * y+1$, r_gamma $=0 * y+1$
)

## Arguments

family the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" distribution. Quotes are needed; default "family = gaussian".
y
$\mathrm{mu} \quad$ the P -spline estimated mean for the glm response vector of length m .
m_binomial a vector of binomial trials having length ( $y$ ), when family = "binomial". Default is 1 vector.
r_gamma a vector of gamma shape parameters, when family = "Gamma". Default is 1 vector.

## Value

A list with two fields:
dev the estimated deviance.
dispersion_parm
the ML estimated dispersion parameter.

Disks Prices of hard disk drives

## Description

Prices and capacities of hard disk drives, as advertised in a Dutch computer monthly in 1999. Prices are given in Dutch guilders; the Euro did not yet exist.

## Usage

data(Disks)

## Format

A dataframe with six columns:
Year 1999-2000
Month month, 1-12
Size capacity in Gb
Buffer buffer size (Mb)
RPM rotating speed (rpm)
PriceDG in Dutch Guilders, divide by 2.2 for Euro.

## Source

Personal information from Paul Eilers.
ECG A section of an ECG (electrocardiogram)

## Description

The data set includes two signals, respiration and the ECG. Both signals are distorted by strong 60 Hz interference from the mains power.

## Usage

data(ECG)

## Format

A data frame with three columns:
time time in seconds
resp respiration, arbitrary units
ecg ECG, arbitrary units.

## Source

https://physionet.org/content/fantasia/1.0.0/

## References

Iyengar N, Peng C-K, Morin R, Goldberger AL, Lipsitz LA. Age-related alterations in the fractal scaling of cardiac interbeat interval dynamics. Am J Physiol, 1996; 271: 1078-1084.

Standard citation for PhysioNet: Goldberger AL, Amaral LAN, Glass L, Hausdorff JM, Ivanov PCh, Mark RG, Mietus JE, Moody GB, Peng C-K, Stanley HE. PhysioBank, PhysioToolkit, and PhysioNet: Components of a New Research Resource for Complex Physiologic Signals (2003). Circulation. 101(23):e215-e220.

## fitampl Fit amplitude coeffcients in the bundle model for expectiles

## Description

There are two functions for fitting the expectile bundle model, one for estimating asymmetry parameters (fitasy), the other for estimating the amplitude function, fitampl, this function. See the details below.

## Usage

fitampl(y, B, alpha, p, a, pord = 2, lambda)

## Arguments

$y \quad a \quad$ response vector.
B a proper B-spline basis matrix, see bbase().
alpha a vector of B-spline coefficients.
$p \quad a$ vector of asymmetries.
a a vector of asymmetry parameters.
pord the order of the difference penalty, default is 2 .
lambda the positive tuning parameter for the penalty.

## Details

The expectile bundle model determines a set of expectile curves for a point cloud with data vectors x and y , as $\psi_{j} x_{i}=a_{j} g\left(x_{i}\right)$. Here $a_{j}$ is the asymmetry parameter corresponding to a given asymmetry $p_{j}$. A vector of asymmetries with all $0<p_{j}<1$ is specified by the user.
The asymmetric least squares objective function is

$$
\sum_{j} \sum_{i} w_{i j}\left(y_{i}-\sum_{j} a_{j} g_{j}\left(x_{i}\right)\right)^{2} .
$$

The function $g(\cdot)$ is called the amplitude. The weights depend on the residuals:

$$
w_{i j}=p_{j}
$$

if $y_{i}>a_{j} g\left(x_{i}\right)$ and $w_{i j}=1-p_{j}$ otherwise.
The amplitude function is a sum of B -splines with coefficients alpha. There is no direct solution, so alpha and the asymmetry parameters a must be updated alternatingly. See the example.

## Value

a vector of estimated $B$-spline coefficients.

## Note

This is a simplification of the model described in the reference. There is no explict term for the trend.

## Author(s)

Paul Eilers

## References

Schnabel, S.K. and Eilers, P.H.C. (2013) A location-scale model for non-crossing expectile curves. Stat 2: 171-183.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
# Get the data
data(bone_data)
x = bone_data$age
y = bone_data$spnbmd
m <- length(x)
# Set asymmetry levels
p = c(0.005, 0.01, 0.02, 0.05, 0.2, 0.5, 0.8, 0.9, 0.95, 0.98, 0.99, 0.995)
np <- length(p)
# Set P-spline parameters
x0 <- 5
x1 <- 30
ndx <- 20
bdeg <- 3
pord <- 2
# Compute bases
B <- bbase(x, x0, x1, ndx, bdeg)
xg <- seq(from = min(x), to = max(x), length = 100)
Bg <- clone_base(B, xg)
```

```
    n <- ncol(B)
    lambda = 1
    alpha <- rep(1,n)
    a = p
    for (it in 1:20){
        alpha <- fitampl(y, B, alpha, p, a, pord, lambda)
        alpha <- alpha / sqrt(mean(alpha ^ 2))
        anew <- fitasy(y, B, alpha, p, a)
        da = max(abs(a - anew))
        a = anew
    cat(it, da, '\n')
        if (da < 1e-6) break
    }
    # Compute bundle on grid
    ampl <- Bg %*% alpha
    Z <- ampl %*% a
    # Plot data and bundle
    plot(x, y, pch = 15, cex = 0.7, col = 'grey', xlab = 'Age', ylab = 'Density')
    cols = colorspace::rainbow_hcl(np, start = 10, end = 350)
    matlines(xg, Z, lty = 1, lwd = 2, col = cols)
```

fitasy Fit asymmetry parameters in the expectile bundle model

## Description

There are two functions for fitting the expectile bundle model, the present one for estimating asymmetry parameters (fitasy), the other for estimating the amplitude function, fitampl. See the details below.

## Usage

fitasy(y, B, b, p, c0)

## Arguments

y
a response vector.
B
a proper B-spline basis matrix, see bbase().
b
a vector of B-spline coefficients.
p
a vector of asymmetries with values between 0 and 1.
c0
a vector.

## Details

The expectile bundle model determines a set of expectile curves for a point cloud with data vectors x and y , as $\psi_{j} x_{i}=a_{j} g\left(x_{i}\right)$. Here $a_{j}$ is the asymmetry parameter corresponding to a given asymmetry $p_{j}$. A vector of asymmetries with all $0<p_{j}<1$ is specified by the user.
The asymmetric least squares objective function is

$$
\sum_{j} \sum_{i} w_{i j}\left(y_{i}-\sum_{j} a_{j} g_{j}\left(x_{i}\right)\right)^{2}
$$

The function $g(\cdot)$ is called the amplitude. The weights depend on the residuals:

$$
w_{i j}=p_{j}
$$

if $y_{i}>a_{j} g\left(x_{i}\right)$ and $w_{i j}=1-p_{j}$ otherwise.
The amplitude function is a sum of B -splines with coefficients alpha. There is no direct solution, so alpha and the asymmetry parameters a must be updated alternatingly. See the example.

## Value

a vector of estimated asymmetry parameters .

## Note

This is a simplification of the model described in the reference. There is no explict term for the trend.

## Author(s)

## Paul Eilers

## References

Schnabel, S.K. and Eilers, P.H.C. (2013) A location-scale model for non-crossing expectile curves. Stat 2: 171-183.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
# Get the data
data(bone_data)
x = bone_data$age
y = bone_data$spnbmd
m <- length(x)
# Set asymmetry levels
p = c(0.005, 0.01, 0.02, 0.05, 0.2, 0.5, 0.8, 0.9, 0.95, 0.98, 0.99, 0.995)
np <- length(p)
# Set P-spline parameters
```

```
x0 <- 5
x1<- 30
ndx <- 20
bdeg <- 3
pord <- 2
# Compute bases
B <- bbase(x, x0, x1, ndx, bdeg)
xg <- seq(from = min(x), to = max(x), length = 100)
Bg <- clone_base(B, xg)
n <- ncol(B)
lambda = 1
alpha <- rep(1,n)
a = p
for (it in 1:20){
    alpha <- fitampl(y, B, alpha, p, a, pord, lambda)
    alpha <- alpha / sqrt(mean(alpha ^ 2))
    anew <- fitasy(y, B, alpha, p, a)
    da = max(abs(a - anew))
    a = anew
    cat(it, da, '\n')
        if (da < 1e-6) break
}
# Compute bundle on grid
ampl <- Bg %*% alpha
Z <- ampl %*% a
# Plot data and bundle
plot(x, y, pch = 15, cex = 0.7, col = 'grey', xlab = 'Age', ylab = 'Density')
cols = colorspace::rainbow_hcl(np, start = 10, end = 350)
matlines(xg, Z, lty = 1, lwd = 2, col = cols)
```

    G519C18 Chromosome G519C18 data
    
## Description

An extract of the data set G519 in the Bioconductor package Vega, for chromosome 18.

## Usage

data(G519C18)

## Format

A dataframe with two columns:
y Probe position
x $\log R$ Ratio.

## References

https://www.bioconductor.org/packages/release/bioc/html/Vega.html

## Examples

plot(G519C18\$x, G519C18\$y, type = 'l', ylab = 'LRR', xlab = 'Position', main = 'Chromosome 18')

Greece_deaths Deaths in Greece in 1960.

## Description

Deaths in Greece in 1960.

## Usage

data(Greece_deaths)

## Format

A dataframe with three columns:
Age 0-85
Male male deaths
Female female deaths.

## Details

All counts for ages above 84 have been grouped to one number for age 85 .

## Source

Personal information from Aris Perperoglou.

Hepatitis Prevalence of Hepatitis among a sample of Bulgarian males.

## Description

Prevalence of Hepatitis among a sample of Bulgarian males.

## Usage

data(Hepatitis)

## Format

A data frame with three columns:
Age years
Infected number of infected persons
Sampled number of sampled persons.

## Source

Table 2 in Keiding (1991).

## References

N. Keiding (1991) Age-Specific Incidence and Prevalence: A Statistical Perspective. JRSS-A 154, 371-396.

## hist2d Compute a $2 D$ histogram

## Description

Compute a two-dimesnional histogram from two vectors (of the same length), x and y .

## Usage

hist2d(x, y, nb = c(100, 100), xlim = range(x), ylim = range(y))

## Arguments

x
$y \quad$ a numeric vector of the same length as $x$.
$\mathrm{nb} \quad$ a vector $\mathrm{c}(\mathrm{nbx}, \mathrm{nby})$, or a scalar nb , providing the number of bins for x , and y ; default is 100 ; see details.
$x \lim \quad$ a vector $c(x \min , x \max )$ containing the limits of the domain of $x$; default range $(x)$.
$y \lim \quad$ a vector $c(y \min , y \max )$ containing the limits of the domain of $y$; default range $(y)$.

## Details

If nb is scalar, it is extended to $\mathrm{c}(\mathrm{nb}, \mathrm{nb})$, so that both dimensions will have the same number of bins.
Elements of $x(y)$ that fall outside the range specified by $x$ lim ( ylim ) are not counted.

## Value

A list with components:
H a matrix of dimension nbx by nby containing bin counts.
xgrid a vector of length $n b x$ representing centers of the bins for $x$.
ygrid a vector of length nby representing centers of the bins for $y$.
$x b$ a vector giving the bin number of each element of $x$.
ybin a vector giving the bin number of each element of $y$.

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

data(faithful)
$x=$ faithful\$eruptions
$y=$ faithful\$waiting
$C=\operatorname{hist} 2 d(x, y, c(50,50))$
image (C\$xgrid, C\$ygrid, C\$H, xlab='Eruption length (min)', ylab='Waiting time (min)')
title('Old Faithful geyser')

```
hist2dsm Smooth a 2D histogram
```


## Description

Fit a 2D smooth P-spline surface to a matrix of counts, assuming Poisson distributed observations.

## Usage

hist2dsm(
$Y$,
nsegx = 10,
nsegy = nsegx,
bdeg $=3$,
lambdax = 10,
lambday $=$ lambdax,

```
    dx = 3,
    dy = dx,
    Mu = Y + 0.01,
    kappa = 1e-04,
    tol = 1e-05
)
```


## Arguments

| Y | a matrix of counts. |
| :---: | :---: |
| nsegx | the number of knots along $\times$ (default=10). |
| nsegy | the number of evenly spaced knots along y for Tensor product B-spline basis (default=10). |
| bdeg | the degree of the basis, default is 3 . |
| lambdax | the positive number for the tuning parameter along x . |
| lambday | the positive number for the tuning parameter along y . |
| dx | the order of the difference penalty along x , default is 3 . |
| dy | the order of the difference penalty along y , default is 3 . |
| Mu | the initialization of the mean (default $Y+0.01$ ). |
| kappa | a (small, positive) number for ridge tuning parameter to stabilize estimation (default 1e-4). |
| tol | the convergence criterion (default 1e-5). |

## Value

A list with elements:
ed the effective dimension of the smooth 2D surface.
$\mathrm{Mu} \quad$ a matrix with the smooth estimates, with dimensions of $\operatorname{dim}(Y)$
pen the numerical value of the penalty.

## Author(s)

Paul Eilers

## References

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
x = faithful$eruptions
y = faithful$waiting
h = hist2d(x, y, c(100, 100))
sm = hist2dsm(h$H, nsegx = 25, nsegy = 25, bdeg = 3, lambdax = 10, lambday = 10)
image(h$xgrid, h$ygrid, sm$Mu, xlab = 'Eruption length (min)',
    ylab = 'Waiting time (min)', main = 'Old Faithful')
```

    indiumoxide
        An X-ray diffractogram.
    
## Description

An X-ray diffractogram.

## Usage

data(indiumoxide)

## Format

A matrix with two columns:
angle the angles (degrees) of diffraction
count corresponding photon counts.

## Details

An X-ray diffractogram of Indium-Tin oxide.
These data have been taken from the source of package Diffractometry, which is no longer available from CRAN in binary form.

## Source

P.L. Davies, U. Gather, M. Meise, D. Mergel, T. Mildenberger (2008). Residual based localization and quantification of peaks in x-ray diffractograms, Annals of Applied Statistics, Vol. 2, No. 3, 861-886.

## Examples

```
angle = indiumoxide[,1]
photon = indiumoxide[,2]
plot(angle, type = 'l', photon, xlab = 'Angle', ylab = 'Photon count')
```

inverse_link Inverse link function, used for GLM fitting.

## Description

Inverse link function, used for GLM fitting.

## Usage

inverse_link(x, link)

## Arguments

$x \quad$ scalar, vector, or matrix input.
link the link function, one of "identity", "log", "sqrt", "logit", "probit", "cloglog", "loglog", "reciprocal"; quotes are needed (default "identity").

## Value

The inverse link function applied to $x$. If link is not in the above list of allowed names, NULL will be returned.

```
JOPS Joys of P-Splines
```


## Description

A package for working with and learning about P -splines. P-splines combine B -splines with discrete penalties to build a very flexible and effective smooth models. They can handle non-normal data in the style of generalized linear models.
This package provides functions for constructing B-spline bases and penalty matrices. It solves the penalized likelihood equations efficiently.
Several methods are provided to determine the values of penalty parameters automatically, using cross-validation, AIC, mixed models or fast Bayesian algorithms.
This package is a companion to the book by Eilers and Marx (2021). The book presents the underlying theory and contains many examples and the code $R$ for each example is available on the website https://psplines.bitbucket.io

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C. and Marx, B.D. (1996). Flexible smoothing with B-splines and penalties (with comments and rejoinder), Statistical Science, 11: 89-121.

JOPS_colors Custom color ramp.

## Description

Custom color ramp.

## Usage

JOPS_colors(n)

## Arguments

n number of steps.

## Value

custom color ramp.

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

JOPS_point Themeing functions used to unify ggplot features

## Description

Custom size and color of points.

## Usage

JOPS_point(s_size = 1.5)

## Arguments

s_size point size parameter for ggplot2 $($ default $=1.5)$.

## Value

themeing function for ggplot2 features.

JOPS_theme Custom theme for ggplot

## Description

Set a ggplot theme in black and white, with centered titles.
Set a ggplot theme in black and white, with centered titles.

## Usage

JOPS_theme(h_just = 0.5)
JOPS_theme(h_just = 0.5)

## Arguments

h_just horizontal justification for ggplot2.

## Value

custom theme for ggplot.
Custom theming function used to unify ggplot features.

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
LAPS_dens Bayesian density estimation

## Description

Bayesian density estimation with P-splines and Laplace approximation.

## Usage

LAPS_dens(B, P, y, loglambdas, tol = 1e-05, mon = FALSE)

## Arguments

B
P
$y \quad$ vector (length $m$ ) of counts, usually a histogram.
loglambdas
tol convergence tolerance (relative change in coefficients), default 1e-5.
mon TRUE or FALSE to monitor the iteration history (default FALSE).

## Details

The B-spline basis should be based on the midpoints of the histogram bins. See the example below. This function is based on the paper of Gressani and Lambert (2018) and code input by Oswaldo Gressani.

## Value

A list with elements:

| alpha | P-spline coefficients of length $n$. |
| :--- | :--- |
| weights | weights from the Laplace approximation, which sum to 1 and are the same <br> length as loglambdas. |
| mu | a vector of length $m$ of expected values. <br> Cov |
| covariance matrix ( m by m ) of $\log (\mathrm{mu})$. |  |
| lambda | the penalty parameter. |

## Author(s)

## Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Gressani, O. and Lambert, P. (2018). Fast Bayesian inference using Laplace approximations in a flexible promotion time cure model based on P-splines. Computational Statistics and Data Analysis 124, 151-167.

## Examples

```
# Smoothing a histogram of Old Faithful eruption durations
data(faithful)
durations = faithful[, 1] # Eruption length
# Histogram with narrow bin widths
bw = 0.05
hst = hist(durations, breaks = seq(1, 6, by = bw), plot = TRUE)
```

```
x = hst$mids
y = hst$counts
# B-spline basis matrices, for fitting and plotting
nseg = 30
B = bbase(x, nseg = nseg)
xg = seq(min(x), max(x), by = 0.01)
Bg = bbase(xg, nseg = nseg)
n = ncol(B)
# Penalty matrix
D2 = diff(diag(n), diff = 2)
P2 = t(D2) %*% D2
# Fit the model
loglambs = seq(-1, 2, by = 0.05)
laps2 = LAPS_dens(B, P2, y, loglambs, mon = FALSE)
fhat2 = exp(Bg %*% laps2$alpha)
lines(xg, fhat2, col = "blue", lwd = 2)
```

Mixture Mixture Data

## Description

The mixture data were obtained in an unpublished experiment in 2001 by Zhenyu Wang at University of Amsterdam, under the supervision of Age Smilde. We are grateful for the permission to use the data.

## Usage

data(Mixture)

## Format

A list consisting of the following:
fractions a $34 \times 3$ matrix of mixure fractions (rows sum to unity): Water (subboiled demi water (self made)), 1, 2ethanediol (99.8\% Sigma-Aldrich Germany), 3amino1propanol (99\% Merk Schuchardt Germany)
xspectra spectra array, 34 (observations) x 401 (wavelenths channels) x 12 (temperatures (C): 30, $35,37.5,40,45,47.5,50,55,60,62.5,65,70$ )
wl wavelengths for the spectra, 700 to $1100(\mathrm{~nm})$, by 1 nm .

## Details

The following instruments and chemicals were used in the experiment: HP 8453 spectrophotometer (Hewlett-Packard, Palo Alto, CA); 2cm closed quartz cuvette with glass thermostatable jacket; Pt100 temperature sensor; Neslab microprocessor EX-111 circulator bath; UV-visible Chemstation software (Rev A.02.04) on a Hewlett-Packard Vectra XM2 PC.

## References

Eilers, P. H. C., and Marx, B. D. (2003). Multivariate calibration with temperature interaction using two-dimensional penalized signal regression. Chemometrics and Intellegent Laboratory Systems, 66, 159-174.
Marx, B. D., Eilers, P. H. C., and Li, B. (2011). Multidimensional single-index signal regression. Chemometrics and Intelligent Laboratory Systems, 109(2), 120-130. [see the Appendix within]
Zhenyou Wang and Age Smilde, Univeristy of Amsterdam, The Netherlands. Personal communication.

```
pclm Fit a composite link model
```


## Description

Fit a smooth latent distribution using the penalized composite link model (PCLM).

## Usage

$\operatorname{pclm}(y, C, B, \operatorname{lambda}=1, \operatorname{pord}=2$, itmax $=50$, show $=$ FALSE $)$

## Arguments

$y \quad a$ vector of counts, length $m$.
C a composition matrix, $m$ by $q$.
B a B-spline basis matrix, q by $n$.
lambda the penalty parameter.
pord the the order of the difference penalty (default $=2$ ).
itmax the maximum number of iterations (default $=50$ ).
show $\quad$ Set to TRUE or FALSE to display iteration history $($ default $=$ FALSE $)$.

## Details

The composite link model assumes that $E(y)=\mu=C \exp (B \alpha)$, where $\exp (B \alpha)$ is a latent discrete distribution, usually on a finer grid than that for $y$.
Note that sum(gamma) $==\operatorname{sum}(\mathrm{mu})$.

## Value

A list with the following items:

| alpha | the estimated B-spline coefficients, length n. |
| :--- | :--- |
| gamma | the estimated latent distribution, length q. |
| mu | estimated values of y, length m. |
| dev | the deviance of the model. |
| ed | the effective model dimension. |
| aic | Akaike's Information Criterion. |

## Author(s)

Paul Eilers and Jutta Gampe

## References

Eilers, P. H. C. (2007). III-posed problems with counts, the composite link model and penalized likelihood. Statistical Modelling, 7(3), 239-254.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

\# Left and right boundaries, and counts, of wide intervals of the data
$\mathrm{cb}<-\mathrm{c}(0,20,30,40,50,60)$
ce <- c $20,30,40,50,60,70)$
$\mathrm{y}<-\mathrm{c}(79,54,19,1,1,0)$
\# Construct the composition matrix
m <- length $(\mathrm{y})$
$\mathrm{n}<-\max (\mathrm{ce})$
C <- matrix(0, m, n)
for (i in 1:m) C[i, cb[i]:ce[i]] <- 1
mids $=(c b+c e) / 2-0.5$
widths = ce - cb + 1
dens = y / widths / sum(y)
$x=(1: n)-0.5$
$B=$ bbase ( $x$ )
fit $=\operatorname{pclm}(y, C, B, \operatorname{lambda}=2$, pord $=2$, show $=$ TRUE $)$
gamma = fit\$gamma / sum(fit\$gamma)
\# Plot density estimate and data
plot(x, gamma, type = 'l', lwd = 2, xlab = "Lead Concentration", ylab = "Density")
$\operatorname{rect}(c b, 0, c e, d e n s$, density $=\operatorname{rep}(10,6)$, angle $=\operatorname{rep}(45,6)$ )
plot.ps2dglm
Plotting function for ps2DGLM

## Description

Plotting function for 2D P-spline (GLM) smooothing (using ps2DGLM with class ps2dglm).

## Usage

```
## S3 method for class 'ps2dglm'
plot(x, ..., xlab = " ", ylab = " ", Resol = 100, se = 2)
```


## Arguments

x
... other parameters.
$x$ lab label for the x -axis, e.g. "my x " (quotes required).
$y l a b \quad$ label for the $y$-axis, e.g. "my y" (quotes required).
Resol resolution for plotting, default Resol $=100$.
se a scalar, e.g. $s e=2$ to produce twice se surfaces, set se $>0$ (or set se $=0$ to supress).

## Value

Plot a plot of the mean (inverse link) 2D P-spline (GLM) smooth surface.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
library(fields)
library(JOPS)
# Extract data
library(rpart)
Kyphosis <- kyphosis$Kyphosis
Age <- kyphosis$Age
Start <- kyphosis$Start
y <- 1 * (Kyphosis == "present") # make y 0/1
fit <- ps2DGLM(
    Data = cbind(Start, Age, y),
    Pars = rbind(c(1, 18, 10, 3, .1, 2), c(1, 206, 10, 3, .1, 2)),
    family = "binomial"
)
plot(fit, xlab = "Start", ylab = "Age")
#title(main = "Probability of Kyphosis")
```

```
plot.ps2dnormal Plotting function for ps2DNormal
```


## Description

Plotting function for 2D P-spline smooothing (using ps2DNormal with class ps2dnormal).

## Usage

```
## S3 method for class 'ps2dnormal'
plot(x, ..., xlab = " ", ylab = " ", Resol = 100)
```


## Arguments

x
... other parameters.
$x l a b \quad$ label for the $x$-axis, e.g. "my $x "$ (quotes required).
$y l a b \quad$ label for the $y$-axis, e.g. "my y" (quotes required).
Resol resolution for plotting, default Resol $=100$.

## Value

Plot a plot of the smooth 2D P-spline smooth surface.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
library(SemiPar)
library(fields)
library(spam)
library(JOPS)
# Get the data
data(ethanol)
x <- ethanol$C
y <- ethanol$E
z <- ethanol$NOx
```

```
# Set parameters for domain
xlo <- 7
xhi <- 19
ylo <- 0.5
yhi <- 1.25
# Set P-spline parameters, fit and compute surface
xpars <- c(xlo, xhi, 10, 3, 3, 1)
ypars <- c(ylo, yhi, 10, 3, 3, 1)
Pars1 <- rbind(xpars, ypars)
fit <- ps2DNormal(cbind(x, y, z), Pars = Pars1)
plot(fit, xlab = "C", ylab = "E")
```

plot.ps2dsignal Plotting function for ps2DSignal

## Description

Plotting function for 2D P-spline signal regression coefficients (using ps2DSignal with class ps2dsignal). Although standard error surface bands can be comuputed they are intentially left out as they are not interpretable, and there is generally little data to steer such a high-dimensional parameterization.

## Usage

```
## S3 method for class 'ps2dsignal'
plot(x, ..., xlab = " ", ylab = " ", Resol = 200)
```


## Arguments

| x | the P-spline object, usually from ps2DSignal. |
| :--- | :--- |
| $\ldots$ | other parameters. |
| xlab | label for the x-axis, e.g. "my x" (quotes required). |
| ylab | label for the y-axis, e.g. "my y" (quotes required). |
| Resol | Resolution of bgrid (default Resol $=200$ ). |

## Value

Plot a plot of the 2D P-spline signal coefficent surface.

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B.D. and Eilers, P.H.C. (2005). Multidimensional penalized signal regression, Technometrics, 47: 13-22.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(fields)
library(JOPS)
# Get the data
x0 <- Sugar$X
x0 <- x0 - apply(x0, 1, mean) # center Signal
y <- as.vector(Sugar$y[, 3]) # Response is Ash
# Inputs for two-dimensional signal regression
nseg <- c(7, 37)
pord <- c(3, 3)
min_ <- c(230, 275)
max_ <- c(340, 560)
M1_index <- rev(c(340, 325, 305, 290, 255, 240, 230))
M2_index <- seq(from = 275, to = 560, by = .5)
p1 <- length(M1_index)
p2 <- length(M2_index)
# Fit optimal model based on LOOCV
opt_lam <- c(8858.6679, 428.1332) # Found via svcm
Pars_opt <- rbind(
    c(min_[1], max_[1], nseg[1], 3, opt_lam[1], pord[1]),
    c(min_[2], max_[2], nseg[2], 3, opt_lam[2], pord[2]))
fit <- ps2DSignal(y, x0, p1, p2, "unfolded", M1_index, M2_index,
            Pars_opt, int = FALSE, ridge_adj = 1e-4 )
# Plotting coefficient image
plot(fit)
```

plot.pspfit
Plotting function for psNormal, psPoisson, psBinomial

## Description

Plotting function for P -spline smooth with normal, Poisson, or binomial responses (class pspfit), with or without standard error bands.

## Usage

\#\# S3 method for class 'pspfit'
plot (x, ..., se = 2, xlab = "", ylab = "", col = "black", pch = 1)

## Arguments

x
...
se
xlab
ylab
col
pch
the P-spline object, usually from psNormal, psPoisson, psBinomial. other parameters.
a scalar, e.g. $s e=2$ to produce twice se bands, set se $>0$ (or set se=0 to supress). label for the x -axis.
label for the $y$-axis.
color for points.
point character.

## Value

Plot a plot of the mean (inverse link) smoothed normal, Poisson, or binomial responses, with or without se bands.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
library(JOPS)
#Extract data
library(MASS)
# Get the data
data(mcycle)
x = mcycle$times
y = mcycle$accel
fit1 = psNormal(x, y, nseg = 20, bdeg = 3, pord = 2, lambda = .8)
plot(fit1, se = 2, xlab = "time (ms)", ylab = "accel")
library(JOPS)
library(boot)
# Extract the data
Count = hist(coal$date, breaks=c(1851:1963), plot = FALSE)$counts
Year = c(1851:1962)
xl = min(Year)
xr = max(Year)
# Poisson smoothing
nseg = 20
bdeg = 3
```

```
fit1=psPoisson(Year, Count, xl, xr, nseg, bdeg, pord = 2,
lambda = 1)
names(fit1)
plot(fit1, xlab = "Year", ylab = "Count", se = 2)
library(JOPS)
#Extract data
library(rpart)
Kyphosis = kyphosis$Kyphosis
Age =kyphosis$Age
y = 1 * (Kyphosis == "present") # make y 0/1
# Binomial smoothing
fit1 = psBinomial(Age, y, xl = min(Age), xr = max(Age), nseg = 20,
    bdeg = 3, pord = 2, lambda = 1)
names(fit1)
plot(fit1, xlab = "Age", ylab = '0/1', se = 2)
```

plot.pssignal
Plotting function for psSignal

## Description

Plotting function for signal regression P-spline smooth coefficients (using psSignal with class pssignal), with or without standard error bands.

## Usage

\#\# S3 method for class 'pssignal'
plot (x, ..., se = 2, xlab = "", ylab = "", col = "black", lty = 1)

## Arguments

x
...
se
xlab
ylab
col
lty

## Value

Plot
a plot of the smooth P-spline signal coefficent vector, with or without standard error bands.

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B.D. and Eilers, P.H.C. (1999). Generalized linear regression for sampled signals and curves: A P-spline approach. Technometrics, 41(1): 1-13.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
# Get the data
library(fds)
data(nirc)
iindex=nirc$x
x=nirc$y
sel= 50:650 #1200 <= x & x<= 2400
X=X[sel, ]
iindex=iindex[sel]
dX=diff(X)
diindex=iindex[-1]
y=as.vector(labc[1,1:40])
oout = 23
dX=t(dX[,-oout])
y=y[-oout]
fit2 = psSignal(y, dX, diindex, nseg = 25,lambda = 0.0001)
plot(fit2, se = 2, xlab = 'Coefficient Index', ylab= "ps Smooth Coeff")
title(main='25 B-spline segments with tuning=0.0001')
names(fit2)
```

plot.psvcsignal Plotting function for psVCSignal

## Description

Plotting function for varying-coefficent signal regression P-spline smooth coefficients (using psVCSignal with class psvcsignal). Although se surface bands can be comuputed they are intentially left out as they are not interpretable, and there is generally little data to steer such a high-dimensional parameterization.

## Usage

```
## S3 method for class 'psvcsignal'
plot(x, ..., xlab = " ", ylab = " ", Resol = 100)
```


## Arguments

X
... other parameters.
$x l a b \quad$ label for the $x$-axis, e.g. "my $x$ " (quotes required).
$y l a b \quad l a b e l$ for the $y$-axis, e.g. "my y" (quotes required).
Resol resolution for plotting, default Resol $=100$.

## Value

Plot a two panel plot, one of the 2D P-spline signal coefficient surface and another that displays several slices of the smooth coefficient vectors at fixed levels of the varying index.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P. H. C. and Marx, B. D. (2003). Multivariate calibration with temperature interaction using two-dimensional penalized signal regression. Chemometrics and Intellegent Laboratory Systems, 66, 159-174.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(fds)
data(nirc)
iindex <- nirc$x
x <- nirc$y
sel <- 50:650 # 1200 <= x & x<= 2400
X <- X[sel, ]
iindex <- iindex[sel]
dX <- diff(X)
diindex <- iindex[-1]
y <- as.vector(labc[1, 1:40]) # percent fat
t_var <- as.vector(labc[4, 1:40]) # percent flour
oout <- 23
dX <- t(dX[, -oout])
y <- y[-oout]
t_var = t_var[-oout]
Pars = rbind(c(min(diindex), max(diindex), 25, 3, 1e-7, 2),
c(min(t_var), max(t_var), 20, 3, 0.0001, 2))
fit1 <- psVCSignal(y, dX, diindex, t_var, Pars = Pars,
family = "gaussian", link = "identity", int = TRUE)
plot(fit1, xlab = "Coefficient Index", ylab = "VC: % Flour")
names(fit1)
```

plot.simpsr Plotting function for sim_psr

## Description

Plotting function for single-index signal regression with tensor product P -splines (using sim_psr with class simpsr).

## Usage

\#\# S3 method for class 'simpsr'
plot(x, ..., xlab = " ", ylab = " ", Resol = 100)

## Arguments

x
... other parameters.
$x l a b \quad l a b e l$ for the $x$-axis, e.g. "my $x "$ (quotes required).
$y l a b \quad l a b e l$ for the $y$-axis, e.g. "my y" (quotes required).
Resol resolution for plotting, default Resol $=100$.

## Value

Plot a two panel plot, one for the estimated P-spline signal coefficent vector, and another for the estimated (unkown) P-spline smooth link function.

## Author(s)

Paul Eilers, Brian Marx, and Bin Li

## References

Eilers, P.H.C., B. Li, B.D. Marx (2009). Multivariate calibration with single-index signal regression, Chemometrics and Intellegent Laboratory Systems, 96(2), 196-202.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
# Get the data
library(fds)
data(nirc)
iindex <- nirc$x
x <- nirc$y
sel <- 50:650 # 1200 <= x & x<= 2400
X <- X[sel, ]
```

```
iindex <- iindex[sel]
dX <- diff(X)
diindex <- iindex[-1]
y <- as.vector(labc[1, 1:40])
oout <- 23
dX <- t(dX[, -oout])
y <- y[-oout]
pords <- c(2, 2)
nsegs <- c(27, 7)
bdegs = c(3, 3)
lambdas <- c(1e-6, .1)
max_iter <- 100
# Single-index model
fit <- sim_psr(y, dX, diindex, nsegs, bdegs, lambdas, pords,
    max_iter)
plot(fit, xlab = "Wavelength (nm)", ylab = " ")
```

plot.simvcpsr Plotting function for sim_vcpsr

## Description

Plotting function for varying-coefficent single-index signal regression using tensor P -splines (using sim_vcpsr with class simvcpsr).

## Usage

```
## S3 method for class 'simvcpsr'
plot(x, ..., xlab = " ", ylab = " ", Resol = 100)
```


## Arguments

| $x$ | the P-spline object, usually from sim_vcpsr. |
| :--- | :--- |
| $\ldots$ | other parameters. |
| xlab | label for the x-axis, e.g. "my x" (quotes required). |
| ylab | label for the y-axis, e.g. "my y" (quotes required). |
| Resol | resolution for plotting, default Resol $=100$. |

## Value

Plot a plot of the estimated 2D P-spline signal coefficent surface along with the companion plot of the estimated 2D P-spline varying link function surface. Slices of these plots, at fixed levels of the indexing covariate, are also provided.

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B. D. (2015). Varying-coefficient single-index signal regression. Chemometrics and Intellegent Laboratory Systems, 143, 111-121.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
\#' @ examples \# Load libraries library(fields) \# Needed for plotting
\# Get the data Dat <- Mixture
\# Dimensions: observations, temperature index, signal m <-34 p1 <- $401 \mathrm{p} 2<-12$
\# Stacking mixture data, each mixture has 12 signals stacked \# The first differenced spectra are also computed. mixture_data <- matrix $(0$, nrow $=\mathrm{p} 2 * \mathrm{~m}, \mathrm{ncol}=\mathrm{p} 1)$ for (ii in 1:m)
mixture_data[((ii - 1) * p2 + 1):(ii * p2), 1:p1] <- t(as.matrix (Dat\$xspectra[ii, , ])) d_mixture_data $<-\mathrm{t}(\operatorname{diff}(\mathrm{t}($ mixture_data $)))$
\# Response (typo fixed) and index for signal y_mixture <- Dat\$fractions y_mixture[17, 3] <- 0.1501 index_mixture <- Dat\$wl
\# Select response and replicated for the 12 temps \# Column 1: water; 2: ethanediol; 3: amino-1propanol $\mathrm{y}<-\operatorname{as.vector}\left(\mathrm{y} \_\right.$mixture[, 2]) y $<-\mathrm{rep}(\mathrm{y}$, each $=\mathrm{p} 2$ )
bdegs $=c(3,3,3,3)$ pords $<-\mathrm{c}(2,2,2,2)$ nsegs $<-\mathrm{c}(12,5,5,5) \#$ Set to $\mathrm{c}(27,7,7,7)$ for given lambdas mins $<-\mathrm{c}(700,30)$ maxs $<-\mathrm{c}(1100,70)$ lambdas $<-\mathrm{c}(1 \mathrm{e}-11,100,0.5,1)$ \# based on svcm search x _index $<-\operatorname{seq}($ from $=701$, to $=1100$, by $=1)$ \# for dX t_var_sub $<-\mathrm{c}(30,35,37.5,40,45$, $47.5,50,55,60,62.5,65,70)$ t_var <- rep(t_var_sub, m) max_iter <- 2 \# Set higher in practice, e.g. 100 int <- TRUE
\# Defining $x$ as first differenced spectra, number of channels. $x<-d \_m i x t u r e \_d a t a$
\# Single-index VC model using optimal tuning fit <- sim_vcpsr(y, x, t_var, x_index, nsegs, bdegs, lambdas, pords, max_iter = max_iter, $\operatorname{mins}=\operatorname{mins}, \operatorname{maxs}=\operatorname{maxs}$ )
plot(fit, xlab = "Wavelength (nm)", ylab = "Temp C")
predict.ps2dglm Predict function for ps2DGLM

## Description

Prediction function which returns both linear predictor and inverse link predictions at arbitrary ( $x$, y) data locations (using ps2DGLM with class ps2dglm).

## Usage

\#\# S3 method for class 'ps2dglm'
predict(object, ..., XY, type = "mu")

## Arguments

object an object using ps2DGLM.
... other parameters.
$X Y \quad$ a matrix of arbitrary $(x, y)$ locations for desired prediction.
type the mean value type = "mu" (default) or linear predictor type = "eta".

## Value

pred the estimated mean (inverse link function) (default) or the linear predictor prediction with type $=$ "eta", for arbitary $(x, y)$ locations in XY.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(fields)
library(JOPS)
# Extract data
library(rpart)
Kyphosis <- kyphosis$Kyphosis
Age <- kyphosis$Age
Start <- kyphosis$Start
y <- 1 * (Kyphosis == "present") # make y 0/1
fit <- ps2DGLM(
        Data = cbind(Start, Age, y),
        Pars = rbind(c(1, 18, 10, 3, .1, 2), c(1, 206, 10, 3, .1, 2)),
        family = "binomial", link = "logit")
predict(fit, XY = cbind(Start, Age)[1:5,])
```

predict.ps2dnormal Predict function for ps2DNormal

## Description

Prediction function which returns linear predictions at arbitrary ( $\mathrm{x}, \mathrm{y}$ ) data locations (using ps2DNormal with class ps2dnormal).

## Usage

\#\# S3 method for class 'ps2dnormal'
predict(object, ..., XY)

## Arguments

object an object using ps2DNormal.
... other parameters.
$X Y \quad$ a matrix of arbitrary $(x, y)$ locations for desired prediction.

## Value

pred the estimated mean at $(x, y)$ locations, in $X Y$.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(SemiPar)
library(fields)
library(spam)
library(JOPS)
# Get the data
data(ethanol)
x <- ethanol$C
y <- ethanol$E
z <- ethanol$NOx
# Set parameters for domain
xlo <- 7
xhi <- 19
ylo <- 0.5
yhi <- 1.25
# Set P-spline parameters, fit and compute surface
xpars <- c(xlo, xhi, 10, 3, 0.01, 1)
ypars <- c(ylo, yhi, 10, 3, 0.1, 1)
Pars1 <- rbind(xpars, ypars)
fit <- ps2DNormal(cbind(x, y, z), Pars = Pars1)
predict(fit, XY = cbind(x, y)[1:5, ])
```

predict.ps2dsignal Predict function for ps2DSignal

## Description

Prediction function which returns both linear predictor and inverse link predictions for arbitrary 2D signals (using ps2DSignal with class ps2dsignal).

## Usage

\#\# S3 method for class 'ps2dsignal'
predict(object, ..., M_pred, M_type = "unfolded", type = "mu")

## Arguments

object an object using ps2DSignal.
... other parameters.
M_pred a matrix of q arbitrary "stacked" or "unfolded" signal matrices of dimension (q by p 1 ) by p 2 or $q$ by ( $p 1$ by p 2 , respectively, for desired prediction (default "unfolded").
M_type $\quad$ stacked" or "unfolded" (default).
type the mean value type = "mu" $($ default $)$ or linear predictor type = "eta".

## Value

pred the estimated mean (inverse link function) or the linear predictor prediction with type = "eta", for arbitary 2D signals in M_pred.

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B.D. and Eilers, P.H.C. (2005). Multidimensional penalized signal regression, Technometrics, 47: 13-22.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(fields)
library(JOPS)
# Get the data
x0 <- Sugar$X
x0 <- x0 - apply(x0, 1, mean) # center Signal
```

```
y <- as.vector(Sugar$y[, 3]) # Response is Ash
# Inputs for two-dimensional signal regression
nseg <- c(7, 37)
pord <- c(3, 3)
min_ <- c(230, 275)
max_ <- c(340, 560)
M1_index <- rev(c(340, 325, 305, 290, 255, 240, 230))
M2_index <- seq(from = 275, to = 560, by = .5)
p1 <- length(M1_index)
p2 <- length(M2_index)
# Fit optimal model based on LOOCV
opt_lam <- c(8858.6679, 428.1332) # Found via svcm
Pars_opt <- rbind(
    c(min_[1], max_[1], nseg[1], 3, opt_lam[1], pord[1]),
    c(min_[2], max_[2], nseg[2], 3, opt_lam[2], pord[2])
)
fit <- ps2DSignal(y, x0, p1, p2, "unfolded", M1_index, M2_index,
    Pars_opt,int = TRUE, ridge_adj = 0.0001,
    M_pred = x0 )
predict(fit, M_pred= x0, type = "mu", M_type = "unfolded")
```

predict.pspfit

## Description

Prediction function which returns both linear predictor and inverse link predictions at arbitrary data locations (using psNormal, psBinomial, psPoisson with class pspfit).

## Usage

\#\# S3 method for class 'pspfit'
predict(object, ..., x, type = "mu")

## Arguments

object an object using psNormal, psBinomial, or psPoisson .
... other parameters.
$x \quad$ a scalar or vector of arbitrary $x$ locations for desired prediction.
type the mean value type = "mu" (default) or linear predictor type = "eta".

## Value

pred the estimated mean (inverse link function) (default) or the linear predictor prediction with type $=$ "eta", at arbitary $\times$ locations.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
library(boot)
# Extract the data
Count <- hist(coal$date, breaks = c(1851:1963), plot = FALSE)$counts
Year <- c(1851:1962)
xl <- min(Year)
xr <- max(Year)
# Poisson smoothing
nseg <- 20
bdeg <- 3
fit1 <- psPoisson(Year, Count, xl, xr, nseg, bdeg, pord = 2, lambda = 1)
names(fit1)
plot(fit1, xlab = "Year", ylab = "Count", se = 2)
predict(fit1, x = fit1$x[1:5])
predict(fit1, x = fit1$x[1:5], type = "eta")
```

```
predict.pssignal Predict function for psSignal
```


## Description

Prediction function which returns both linear predictor and inverse link predictions, for an arbitrary matrix of signals (using psSignal with class pssignal).

## Usage

\#\# S3 method for class 'pssignal'
predict (object, ..., X_pred, type = "mu")

## Arguments

object an object using psSignal.
... other parameters.
X_pred a matrix of arbitrary signals with $\operatorname{ncol}(X)==$ length(x_index) locations for desired prediction.
type the mean value type = "mu" (default) or linear predictor type = "eta".

## Value

pred the estimated mean (inverse link function) (default) or the linear predictor prediction with type $=$ "eta", for a matrix of signals in X_pred.

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B.D. and Eilers, P.H.C. (1999). Generalized linear regression for sampled signals and curves: A P-spline approach. Technometrics, 41(1): 1-13.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
# Get the data
library(fds)
data(nirc)
iindex=nirc$x
X=nirc$y
sel= 50:650 #1200<= x & x<= 2400
X=X[sel,]
iindex=iindex[sel]
dX=diff(X)
diindex=iindex[-1]
y=as.vector(labc[1,1:40])
oout=23
dX=t(dX[,-oout])
y=y[-oout]
fit1 = psSignal(y, dX, diindex, nseg = 25,lambda = 0.0001)
predict(fit1, X_pred = dX[1:5, ])
predict(fit1, X_pred = dX[1:5, ], type = 'eta')
```


## Description

Prediction function which returns both linear predictor and inverse link predictions for an arbitrary matrix of signals with their vector of companion indexing covariates (using psVCSignal with class psvcsignal).

## Usage

\#\# S3 method for class 'psvcsignal'
predict(object, ..., X_pred, t_pred, type = "mu")

## Arguments

object an object using psVCSignal.
... other parameters.
X_pred a matrix of $q$ arbitrary signal vectors of dimension $q$ by $p 1$ for desired prediction.
t_pred a q vector for the varying index variable associated with X_pred.
type the mean value type ="mu" (default) or linear predictor type = "eta".

## Value

pred the estimated mean (inverse link function) (default) or the linear predictor prediction with type $=$ "eta", at signals in matrix X_pred and covariates in vector t_pred.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P. H. C. and Marx, B. D. (2003). Multivariate calibration with temperature interaction using two-dimensional penalized signal regression. Chemometrics and Intellegent Laboratory Systems, 66, 159-174.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
    library(fds)
    data(nirc)
    iindex <- nirc$x
    x <- nirc$y
    sel <- 50:650 # 1200 <= x & x<= 2400
    X <- X[sel, ]
    iindex <- iindex[sel]
    dX <- diff(X)
    diindex <- iindex[-1]
    y <- as.vector(labc[1, 1:40]) # percent fat
    t_var <- as.vector(labc[4, 1:40]) # percent flour
    oout <- 23
    dX <- t(dX[, -oout])
    y <- y[-oout]
    t_var = t_var[-oout]
    Pars = rbind(c(min(diindex), max(diindex), 25, 3, 1e-7, 2),
    c(min(t_var), max(t_var), 20, 3, 0.0001, 2))
    fit1 <- psVCSignal(y, dX, diindex, t_var, Pars = Pars,
    family = "gaussian", link = "identity", int = TRUE)
    predict(fit1, X_pred = dX[1:5,], t_pred = t_var[1:5])
```

predict.simpsr Predict function for sim_psr

## Description

Prediction function which returns single-index inverse link linear predictions at arbitrary data locations (using sim_psr with class simpsr).

## Usage

\#\# S3 method for class 'simpsr'
predict(object, ..., X_pred)

## Arguments

$$
\begin{array}{ll}
\text { object } & \text { an object using sim_psr. } \\
\ldots & \text { other parameters. } \\
\text { X_pred } & \begin{array}{l}
\text { a matrix of arbitrary signals with ncol (X_pred) }=\text { length (x_index) locations } \\
\text { for desired prediction. }
\end{array}
\end{array}
$$

## Value

pred the estimated (inverse single-index) mean for the signals in X_pred.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C., B. Li, B.D. Marx (2009). Multivariate calibration with single-index signal regression, Chemometrics and Intellegent Laboratory Systems, 96(2), 196-202.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

library(JOPS)
\# Get the data
library(fds)
data(nirc)
iindex <- nirc\$x
x <- nirc\$y
sel <- 50:650 \# $1200<=x \& x<=2400$
X <- X[sel, ]
iindex <- iindex[sel]
dX <- diff( $X$ )
diindex <- iindex[-1]
y <- as.vector(labc[1, 1:40])
oout <- 23
$d X<-t(d X[$, -oout $])$
y <- y[-oout]
pords <- c(2, 2)
nsegs <- c(27, 7)
bdegs $=c(3,3)$
lambdas <- c(1e-6, .1)
max_iter <- 100
\# Single-index model
fit <- sim_psr(y, dX, diindex, nsegs, bdegs, lambdas, pords, max_iter)
predict(fit, X_pred = dX)

```
predict.simvcpsr Predict function for sim_vcpsr
```


## Description

Prediction function which returns varying-coefficient single-index inverse link linear predictions at arbitrary data locations (using sim_vcpsr with class simvepsr).

## Usage

\#\# S3 method for class 'simvcpsr'
predict(object, ..., X_pred, t_pred)

## Arguments

object an object using sim_vcpsr.
... other parameters.
X_pred a matrix of arbitrary signals with ncol (X_pred) = length(x_index) locations for desired prediction.
t_pred a q vector for the VC index variable associated with X_pred.

## Value

pred the estimated (inverse single-index) mean for the signals in the matrix X_pred, with the companion vector of indexing covariates in t_pred.

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B. D. (2015). Varying-coefficient single-index signal regression. Chemometrics and Intellegent Laboratory Systems, 143, 111-121.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
# Load libraries
library(fields) # Needed for plotting
# Get the data
Dat <- Mixture
# Dimensions: observations, temperature index, signal
m <- 34
p1 <- 401
p2 <- 12
# Stacking mixture data, each mixture has 12 signals stacked
# The first differenced spectra are also computed.
mixture_data <- matrix(0, nrow = p2 * m, ncol = p1)
for (ii in 1:m)
{
    mixture_data[((ii - 1) * p2 + 1):(ii * p2), 1:p1] <-
        t(as.matrix(Dat$xspectra[ii, , ]))
    d_mixture_data <- t(diff(t(mixture_data)))
}
# Response (typo fixed) and index for signal
y_mixture <- Dat$fractions
y_mixture[17, 3] <- 0.1501
index_mixture <- Dat$wl
```

```
    # Select response and replicated for the 12 temps
    # Column 1: water; 2: ethanediol; 3: amino-1-propanol
    y <- as.vector(y_mixture[, 2])
    y <- rep(y, each = p2)
    bdegs = c(3, 3, 3, 3)
    pords <- c(2, 2, 2, 2)
    nsegs <- c(12, 5, 5, 5) # Set to c(27, 7, 7 , 7) for given lambdas
    mins <- c(700, 30)
    maxs <- c(1100, 70)
    lambdas <- c(1e-11, 100, 0.5, 1) # based on svcm search
    x_index <- seq(from = 701, to = 1100, by = 1) # for dX
    t_var_sub <- c(30, 35, 37.5, 40, 45, 47.5, 50, 55, 60, 62.5, 65, 70)
    t_var <- rep(t_var_sub, m)
    max_iter <- 2 # Set higher in practice, e.g. 100
    int <- TRUE
    # Defining x as first differenced spectra, number of channels.
    x <- d_mixture_data
    # Single-index VC model using optimal tuning
fit <- sim_vcpsr(y, x, t_var, x_index, nsegs, bdegs, lambdas, pords,
    max_iter = max_iter, mins = mins, maxs = maxs)
predict(fit, X_pred = x, t_pred = t_var)
```

ps2DGLM Two-dimensional smoothing of scattered normal or non-normal (GLM) responses using tensor product $P$-splines.

## Description

ps2DGLM is used to smooth scattered normal or non-normal responses, with aniosotripic penalization of tensor product P -splines.

## Usage

```
ps2DGLM(
    Data,
    Pars = rbind(c(min(Data[, 1]), max(Data[, 1]), 10, 3, 1, 2), c(min(Data[, 2]),
        max(Data[, 2]), 10, 3, 1, 2)),
    ridge_adj = 0,
    XYpred = Data[, 1:2],
    z_predicted = NULL,
    se_pred = 2,
    family = "gaussian",
    link = "default",
```

```
        m_binomial = rep(1, nrow(Data)),
        wts = rep(1, nrow(Data)),
        r_gamma = rep(1, nrow(Data))
)
```


## Arguments

| Data | a matrix of 3 columns $x, y, z$ of equal length; the response is $z$. <br> a matrix of 2 rows, where the first and second row sets the P-spline paramters <br> for $x$ and $y$, respectively. Each row consists of: min max nseg bdeg lambda <br> pord. The min and max set the ranges, nseg (default 10) is the number of evenly <br> spaced segments between min and max, bdeg is the degree of the basis (default <br> 3 for cubic), lambda is the (positive) tuning parameter for the penalty (default <br> 1), pord is the number for the order of the difference penalty (default 2). <br> a ridge penalty tuning parameter, usually set to small value, e.g. 1e-8 to stabilize <br> estimation (default 0). |
| :--- | :--- |
| ridge_adj |  |

## Details

Support functions needed: pspline_fitter, bbase, and pspline_2dchecker.
Value

| pcoef | a vector of length (Pars $[1,3]+\operatorname{Pars}[1,4]) *(\operatorname{Pars}[2,3]+\operatorname{Pars}[2,4])$ of (un- <br> folded) estimated P-spline coefficients. |
| :--- | :--- |
| mu | a vector of length $(z)$ of smooth estimated means (at the $x, y$ locations). |
| devf_df | the deviance of fit. |
| aic | the approximate effective dimension of fit. |
| df_resid | AIC. |

```
cv leave-one-out standard error prediction, when family = 'gaussian'.
cv_predicted standard error prediction for y_predict, when family = 'gaussian'.
avediff_pred mean absolute difference prediction, when family = 'gaussian'.
Pars the design and tuning parameters (see arguments above).
dispersion_parm
estimate of dispersion, dev/df_resid.
summary_predicted
    inverse link prediction vectors, and se_pred bands.
eta_predicted estimated linear predictor of length(z).
press_mu leave-one-out prediction of mean, when family = 'gaussian'.
bin_percent_correct
    percent correct classification based on 0.5 cut-off (when family = "binomial").
Data a matrix of 3 columns x,y,z of equal length; the response is z.
Q the tensor product B-spline basis.
qr the Q-R of the model.
```


## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## See Also

ps2DNormal

## Examples

```
library(fields)
library(JOPS)
# Extract data
library(rpart)
Kyphosis <- kyphosis$Kyphosis
Age <- kyphosis$Age
Start <- kyphosis$Start
y <- 1 * (Kyphosis == "present") # make y 0/1
fit <- ps2DGLM(
    Data = cbind(Start, Age, y),
    Pars = rbind(c(1, 18, 10, 3, .1, 2), c(1, 206, 10, 3, .1, 2)),
    family = "binomial", link = "logit")
plot(fit, xlab = "Start", ylab = "Age")
#title(main = "Probability of Kyphosis")
```

ps2DNormal Two-dimensional smoothing scattered (normal) data using P-splines.

## Description

ps2DNormal is used to smooth scattered (normal) data, with aniosotripic penalization of tensor product P -splines.

## Usage

```
    ps2DNormal(
        Data,
        Pars = rbind(c(min(Data[, 1]), max(Data[, 1]), 10, 3, 1, 2), c(min(Data[, 2]),
            max(Data[, 2]), 10, 3, 1, 2)),
        XYpred = expand.grid(Data[, 1], Data[, 2])
    )
```


## Arguments

Data a matrix of 3 columns $x, y, z$ of equal length; the response is $z$.
Pars a matrix of 2 rows, where the first and second row sets the P -spline paramters for x and y , respectively. Each row consists of: min max nseg bdeg lambda pord. The min and max set the ranges, nseg (default 10) is the number of evenly spaced segments between min and max, bdeg is the degree of the basis (default 3 for cubic), lambda is the (positive) tuning parameter for the penalty (default $1)$, pord is the number for the order of the difference penalty (default 2 ),
XYpred a matrix with two columns $(x, y)$ that give the coordinates of (future) prediction; the default is the data locations.

## Details

Support functions needed: pspline_fitter, bbase, and pspline_2dchecker.

## Value

coef a vector of length (Pars[1,3]+Pars[1,4])*(Pars[2,3]+Pars[2,4]) of (unfolded) estimated P -spline coefficients.
fit a vector of length ( $y$ ) of smooth estimated means (at the $x, y$ locations).
pred a vector of length nrow(XYpred) of (future) predictions.
Pars the design and tuning parameters (see arguments above).
cv leave-one-out standard error of prediction or root average PRESS.
h "hat" diagonals of tensor P-spline fit.
B tensor product B-spline basis used for fitting.

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## See Also

ps2DGLM

## Examples

```
library(SemiPar)
library(fields)
library(spam)
library(JOPS)
# Get the data
data(ethanol)
x <- ethanol$C
y <- ethanol$E
z <- ethanol$NOx
# Set parameters for domain
xlo <- 7
xhi <- 19
ylo <- 0.5
yhi <- 1.25
# Set P-spline parameters, fit and compute surface
xpars <- c(xlo, xhi, 10, 3, 3, 1)
ypars <- c(ylo, yhi, 10, 3, 3, 1)
Pars1 <- rbind(xpars, ypars)
fit <- ps2DNormal(cbind(x, y, z), Pars = Pars1)
plot(fit, xlab = "C", ylab = "E")
```

ps2DSignal Two-dimensional penalized signal regression using $P$-splines.

## Description

ps2DSignal is a function used to regress a (glm) response onto a two-dimensional signal or image, with aniosotripic penalization of tensor product P -splines.

```
Usage
    ps2DSignal(
        y ,
        M,
        p1,
        p2,
        M_type = "stacked",
        M1_index \(=c(1: p 1)\),
        M2_index \(=c(1: p 2)\),
        Pars \(=\operatorname{rbind}(c(1, p 1,10,3,1,2), c(1, p 2,10,3,1,2))\),
        ridge_adj = 1e-06,
        M_pred = M,
        y_predicted = NULL,
        family = "gaussian",
        link = "default",
        m_binomial \(=1+0\) * \(y\),
        wts = \(1+0\) * \(y\),
        r_gamma \(=1+0\) * \(y\),
        int = TRUE,
        se_pred \(=2\)
    )
```


## Arguments

y
M
p1 the row dimension of the image.
p2 the column dimension of the image.
M_type $\quad$ stacked" (signal as matrix) or "unfolded" (signal as vector).
M1_index an index of length p 1 for rows of regressor matrix (default is a simple sequence).
M2_index an index of length p2 for columns of regressor matrix (default is a simple sequence).
Pars a matrix of 2 rows, where the first and second row sets the P -spline paramters for $x$ and $y$, respectively. Each row consists of: min max nseg bdeg lambda pord. The min and max set the ranges, nseg (default 10) is the number of evenly spaced segments between min and max, bdeg is the degree of the basis (default 3 for cubic), lambda is the (positive) tuning parameter for the penalty (default 1 ), pord is the number for the order of the difference penalty (default 2 ).
ridge_adj A ridge penalty tuning parameter (usually set to small value, default $1 e-6$, to stabilize estimation).
M_pred (e.g. stacked (q*p1) by p2 signal inputs or (unfolded) q by (p1 * p2) signal inputs for $q$ new predictions.

| y_predicted | a vector of responses from a cv data set (assoc. with M_pred), when family = "gaussian". |
| :---: | :---: |
| family | the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" distribution. Quotes are needed. Default is "gaussian". |
| link | the link function, one of "identity", "log", "sqrt", "logit", "probit", "cloglog", "loglog", "reciprocal"; quotes are needed (default "identity"). |
| m_binomial | a vector of binomial trials having length(y). Default is 1 vector for family $=$ "binomial", NULL otherwise. |
| wts | the weight vector of length ( y ). Default is 1 . |
| r_gamma | a vector of gamma shape parameters. Default is 1 vector for for family $=$ "Gamma", NULL otherwise. |
| int | set to TRUE or FALSE to include intercept term in linear predictor (default TRUE). |
| se_pred | a scalar, e.g. se $=2$ (default) to produce twice se surfaces, set se $>0$. Used for CIs at XYpred locations. |

## Details

Support functions needed: pspline_fitter, bbase, and pspline_2dchecker.

## Value

pcoef a vector of length (Pars[1,3]+Pars[1,4])*(Pars[2,3]+Pars[2, 4]) of (unfolded) estimated P -spline coefficients for tensor surface.
summary_predicted
inverse link prediction vectors, and standard error surfaces.
dev deviance of fit.
eff_df the approximate effective dimension of fit.
aic
df_resid approximate df residual.
cv leave-one-out standard error prediction, when family = "gaussian".
cv_predicted standard error prediction for y_predict, when family = "gaussian".
avediff_pred mean absolute difference prediction, when family = 'gaussian'.
Pars design and tuning parameters (see above arguments).
Dispersion_parm
estimate of dispersion, dev/df_resid.
summary_predicted
inverse link prediction vectors at M_pred, and standard error bands.
eta_predicted estimated linear predictor of length(y).
press_mu leave-one-out prediction of mean, when family = "gaussian".
bin_percent_correct
percent correct classification based on 0.5 cut-off, when family = "binomial",
NULL otherwise.

| B | Tensor basis ( $\mathrm{p} 1 \times \mathrm{p} 2$ ) by ( $\mathrm{n} 1 \times \mathrm{n} 2$ ) for 2D signal regression. |
| :---: | :---: |
| Q | Effective regressors (mby $n 1 * n 2$ ) for 2D signal regression. |
| Ahat | smooth P-spline coefficient vector of length p1 x p2, constructed by B \%*\% pcoef. |
| M | the signal/image regressors. |
| y | the response vector. |
| M1 index | index of length p 1 for rows of regressor matrix. |
| M2index | index of length p 2 for columns of regressor matrix. |
| M_type | "stacked" or "unfolded". |
| w | GLM weight vector of length $m$. |
| h | "hat" diagonals. |
| ridge_adj | additional ridge tuning parameter to stabilize estimation. |

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B.D. and Eilers, P.H.C. (2005). Multidimensional penalized signal regression, Technometrics, 47: 13-22.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(fields)
library(JOPS)
# Get the data
x0 <- Sugar$X
x0 <- x0 - apply(x0, 1, mean) # center Signal
y <- as.vector(Sugar$y[, 3]) # Response is Ash
# Inputs for two-dimensional signal regression
nseg <- c(7, 37)
pord <- c(3, 3)
min_ <- c(230, 275)
max_ <- c(340, 560)
M1_index <- rev(c(340, 325, 305, 290, 255, 240, 230))
M2_index <- seq(from = 275, to = 560, by = .5)
p1 <- length(M1_index)
p2 <- length(M2_index)
# Fit optimal model based on LOOCV
opt_lam <- c(8858.6679, 428.1332) # Found via svcm
Pars_opt <- rbind(
    c(min_[1], max_[1], nseg[1], 3, opt_lam[1], pord[1]),
```

```
    c(min_[2], max_[2], nseg[2], 3, opt_lam[2], pord[2])
)
fit <- ps2DSignal(y, x0, p1, p2, "unfolded", M1_index, M2_index,
    Pars_opt,int = TRUE, ridge_adj = 0.0001,
    M_pred = x0 )
# Plotting coefficient image
    plot(fit)
```

ps2D_PartialDeriv Partial derivative two-dimensional smoothing scattered (normal) data using $P$-splines.

## Description

ps2D_PartialDeriv provides the partial derivative P-spline surface along x, with aniosotripic penalization of tensor product B-splines.

## Usage

```
ps2D_PartialDeriv(
        Data,
        Pars = rbind(c(min(Data[, 1]), max(Data[, 1]), 10, 3, 1, 2), c(min(Data[, 2]),
            max(Data[, 2]), 10, 3, 1, 2)),
        XYpred = cbind(Data[, 1], Data[, 2])
    )
```


## Arguments

Data a matrix of 3 columns $x, y, z$ of equal length; the response is $z$.
Pars a matrix of 2 rows, where the first and second row sets the P-spline paramters for $x$ and $y$, respectively. Each row consists of: min max nseg bdeg lambda pord. The min and max set the ranges, nseg (default 10 ) is the number of evenly spaced segments between min and max, bdeg is the degree of the basis (default 3 for cubic), lambda is the (positive) tuning parameter for the penalty (default 1 ), pord is the number for the order of the difference penalty (default 2 ).

XYpred a matrix with two columns $(x, y)$ that give the coordinates of (future) prediction; the default is the data locations.

## Details

This is support function for sim_vcpsr.

| Value |  |
| :---: | :---: |
| coef | a vector of length (Pars[1, 3] + Pars[1, 4]) * (Pars[1, 3] + Pars[1, 4]). of (unfolded) estimated P -spline coefficients. |
| B | the tensor product B-spline matrix of dimensions m by length(coef). |
| fit | a vector of length ( y ) of smooth estimated means (at the $\mathrm{x}, \mathrm{y}$ locations). |
| pred | a vector of length nrow (XYpred) of (future) predictions. |
| d_coef | a vector of length (Pars[1, 3] + Pars[1, 4]-1) * (Pars[1, 3]+Pars[1, 4]). of (unfolded) partial derivative estimated P -spline coefficients. |
| B_d | the tensor product B-spline matrix of dimensions $m$ by lengh(d_coef), associated with the partial derivative of the tensor basis. |
| d_fit | a vector of length ( $y$ ) of partial derivative (along $x$ ) of the smooth estimated means (at the $x, y$ locations). |
| d_pred | a vector of length nrow(XYpred) of partial derivative (future) predictions. |
| Pars | a matrix of 2 rows, where each the first (second) row sets the P -spline paramters for $x(y)$ : min max nseg bdeg lambda pord. See the argument above. |
| cV | root leave-one-out CV or root average PRESS. |
| XYpred | a matrix with two columns $(x, y)$ that give the coordinates of (future) prediction; the default is the data locations. |

## Author(s)

Brian Marx

## References

Marx, B. D. (2015). Varying-coefficient single-index signal regression. Chemometrics and Intelligent Laboratory Systems, 143, 111-121.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

```
psBinomial
```

Smoothing scattered binomial data using P-splines.

## Description

psBinomial is used to smooth scattered binomial data using P-splines using a logit link function.

## Usage

```
    psBinomial(
```

        x ,
        y ,
        \(x l=\min (x)\),
        \(x r=\max (x)\),
        nseg \(=10\),
        bdeg \(=3\),
        pord \(=2\),
        lambda \(=1\),
        ntrials \(=0 * y+1\),
        wts = NULL,
        show = FALSE,
        iter = 100,
        xgrid \(=100\)
    )

## Arguments

x
y
xl
xr
nseg

## bdeg

pord
lambda
ntrials
wts
show
iter
xgrid
the vector for the continuous regressor of length ( y ) and the abcissae, on which the B -spline basis is constructed.
the response vector, usually $0 / 1$ or binomial counts.
the lower limit for the domain of $x($ default is $\min (x))$.
the upper limit for the domain of $x$ (default is $\max (x)$ ).
the number of evenly spaced segments between xl and xr .
the number of the degree of the basis, usually 1,2 (default), or 3 .
the number of the order of the difference penalty, usually 1,2 , or 3 (defalult).
the (positive) number for the tuning parameter for the penalty.
the vector for the number of binomial trials (default $=1$ ).
the vector of weights, default is 1 , zeros allowed.
Set to TRUE or FALSE to display iteration history.
a scalar to set the maximum number of iterations, default iter $=100$.
a scalar or a vector that gives the x locations for prediction, useful for plotting. If a scalar (default 100) is used then a uniform grid of this size along ( $\mathrm{xl}, \mathrm{xr}$ ).

## Value

pcoe
p
muha
dev
effdim effective dimension of the smooth.
aic
a vector of length $n$ of estimated $P$-spline coefficients.
a vector of length $m$ of estimated probabilities.
uhat
a vector of length $m$ of estimated means (ntrials*p).
deviance

AIC

| wts | a vector of preset weights (default $=1$ ). |
| :--- | :--- |
| nseg | the number of B-spline segments. |
| bdeg | the degree of the B-spline basis. |
| pord | the order of the difference penalty. |
| family | the GLM family (repsonse distribution). |
| link | the link function. |
| y | the binomial response. |
| x | the regressor on which the basis is constructed. |
| P | "half" of the penalty matrix, P'P = lambda*D'D. |
| B | the B-spline basis. |
| lambda | the positive tuning parameter. |
| dispersion | dispersion parameter estimated dev/ (m-effdim). |
| xgrid | gridded x values,useful for plotting. |
| ygrid | gridded fitted linear predictor values, useful for plotting. |
| pgrid | gridded (inverse link) fitted probability values, useful for plotting. |
| se_eta | gridded standard errors for the linear predictor. |

## Author(s)

## Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
library(JOPS)
# Extract data
library(rpart)
Kyphosis <- kyphosis$Kyphosis
Age <- kyphosis$Age
y <- 1 * (Kyphosis == "present") # make y 0/1
fit1 <- psBinomial(Age, y,
    xl = min(Age), xr = max(Age), nseg = 20,
    bdeg = 3, pord = 2, lambda = 10
)
names(fit1)
plot(fit1, xlab = "Age", ylab = "0/1", se = 2)
```


## Description

psNormal is used to smooth scattered (normal) data using P-splines (with identity link function).

## Usage

psNormal (
x ,
$y$,
$x l=\min (x)$,
$x r=\max (x)$,
nseg $=10$,
bdeg $=3$, pord $=2$,
lambda $=1$, wts = NULL, xgrid $=100$
)

## Arguments

$x \quad$ the vector for the continuous regressor of length $(y)$ and the abcissae used to build the B-spline basis.
$y \quad$ the response vector, usually continuous data.
$x 1 \quad$ the number for the min along $x($ default is $\min (x))$.
$x r \quad$ the number for the max along $x$ (default is $\max (x)$ ).
nseg the number of evenly spaced segments between $x l$ and $x r$.
bdeg the number of the degree of the basis, usually 1,2 (default), or 3.
pord the number of the order of the difference penalty, usually 1,2 , or 3 (defalult).
lambda the (positive) number for the tuning parameter for the penalty (default 1 ).
wts the vector of general weights, default is 1 ; zero allowed.
xgrid
a scalar or a vector that gives the x locations for prediction, useful for plotting. If a scalar (default 100) is used then a uniform grid of this size along ( $x 1, x r$ ).

## Value

pcoeff a vector of length $n$ of estimated $P$-spline coefficients.

B
wts
muhat a vector of length $m$ of smooth estimated means.
a matrix of dimension $m$ by $n$ for the $B$-spline basis matrix.
a vector of length $m$ of weights.

| effdim | estimated effective dimension. |
| :--- | :--- |
| ed_resid | approximate df residual. |
| sigma | square root of MSE. |
| cv | standard error of leave-one-out prediction or root average PRESS. |
| nseg | the number of B-spline segments. |
| bdeg | the degree of the B-spline basis. |
| pord | the order of the difference penalty. |
| lambda | the positive tuning parameter. |
| xgrid | gridded x values, useful for plotting. |
| ygrid | gridded fitted mean values, useful for plotting. |
| se_eta | gridded standard errors for the fitted mean values, useful for plotting. |
| P | "half" of the penalty, such that P'P= lambda D'D. |

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
library(JOPS)
library(MASS)
data(mcycle)
x <- mcycle$times
y <- mcycle$accel
fit1 <- psNormal(x, y, nseg = 20, bdeg = 3, pord = 2, lambda = .8)
plot(fit1, se = 2, xlab = "Time (ms)", ylab = "Acceleration")
```

psNormal_Deriv Derivative for a P-spline fit of scattered (normal) data.

## Description

psNormal_Deriv provides the derivative P -spline fit along x .

## Usage

```
psNormal_Deriv(
        x ,
        \(y\),
        \(x l=\min (x)\),
        \(x r=\max (x)\),
        nseg \(=10\),
        bdeg \(=3\),
        pord \(=2\),
        lambda \(=1\),
        wts \(=\operatorname{rep}(1\), length(y)),
        xgrid \(=x\)
    )
```


## Arguments

x $y \quad$ the response vector, usually continuous data.
$x 1 \quad$ the number for the min along $x($ default is $\min (x))$.
$\mathrm{xr} \quad$ the number for the max along $x$ (default is $\max (x)$ ).
nseg the number of evenly spaced segments between $x l$ and $x r$.
bdeg the number of the degree of the basis, usually 1,2 , or 3 (defalult).
pord the number of the order of the difference penalty, usually 1,2 (defalult), or 3 .
lambda the positive tuning parameter (default 1 ).
wts the vector of weights, default is $1 ; 0 / 1$ allowed.
xgrid a scalar or a vector that gives the x locations for prediction, useful for plotting. If a scalar (default 100) is used then a uniform grid of this size along ( $\mathrm{xl}, \mathrm{xr}$ ).

## Details

This is also a support function needed for sim_psr and sim_vcpsr. SISR (Eilers, Li, Marx, 2009).

## Value

coef a vector of length (nsegs + bdeg) of estimated P-spline coefficients.
B
fit
The B-spline matrix of dimensions $m$ by length (coef).
pred
a vector of length ( $y$ ) of smooth estimated means (at the $x$ locations).
d_coef a vector of length (nsegs + bdeg -1) of differenced (derivative) estimated Pspline coefficients.
B_d The first derivative B-spline matrix of dimensions $m$ by lengh (d_coef).
d_fit a vector of length ( $y$ ) of partial derivative (along $x$ ) of the smooth estimated means (at the $x$ locations).
d_pred a vector of length lenght (xgrid) of partial derivative (future) predictions.

| xl | the number for the min along $x($ default is $\min (x))$. |
| :--- | :--- |
| xr | the number for the max along $x($ default is $\max (x))$. |
| nseg | the number of evenly spaced segments between $x l$ and $x r$. |
| bdeg | the number of the degree of the basis, usually 1,2, or 3 (default). |
| pord | the number of the order of the difference penalty, usually 1,2 (default), or 3. |
| lambda | the positive tuning parameter (default 1$).$ |

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B. D. (2015). Varying-coefficient single-index signal regression. Chemometrics and Intelligent Laboratory Systems, 143, 111-121.
Eilers, P.H.C., B. Li, B.D. Marx (2009). Multivariate calibration with single-index signal regression, Chemometrics and Intellegent Laboratory Systems, 96(2), 196-202.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## See Also

sim_psr sim_vcpsr
pspline2d_checker $\quad P$-spline $2 D$ tensor product checking algorithm for the GLM.

## Description

pspline_2dchecker checks to see if all the 2D tensor inputs associated for P-spines are properly defined.

```
Usage
    pspline2d_checker(
        family,
    link,
    bdeg1,
    bdeg2,
    pord1,
    pord2,
    nseg1,
    nseg2,
    lambda1,
    lambda2,
    ridge_adj,
    wts
)
```


## Arguments

| family | the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" <br> distribution. Quotes are needed. |
| :--- | :--- |
| link | the link function, one of "identity", "log", "sqrt", "logit", "probit", <br> "cloglog", "loglog", "reciprocal"; quotes are needed. |
| bdeg1 | the degree of B-splines. |
| bdeg2 | the degree of B-splines. |
| pord1 | the order of the penalty. |
| pord2 | the order of the penalty. |
| nseg1 | the number of evenly spaced B-spline segmements. |
| nseg2 | the number of evenly spaced B-spline segmements. |
| lambda1 | the positive tuning parameter for the difference penalty. |
| lambda2 | the positive tuning parameter for the difference penalty. |
| ridge_adj | the positive tuning parameter for the ridge penalty. |
| wts | the weight vector, separate from GLM weights. |

## Value

list same as inputs, with warnings if required.

```
pspline_checker P-spline checking algorithm for the GLM.
```


## Description

pspline_checker checks to see if all the inputs associated for P-spines are properly defined.

## Usage

pspline_checker(family, link, bdeg, pord, nseg, lambda, ridge_adj, wts)

## Arguments

| family | the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" <br> distribution. Quotes are needed. |
| :--- | :--- |
| link | the link function, one of "identity", "log", "sqrt", "logit", "probit", <br> "cloglog", "loglog", "reciprocal"; |
| bdeg | the degree of B-splines. |
| pord | the order of the penalty. |
| nseg | the number of evenly-spaced B-spline segmements. |
| lambda | the positive tuning parameter for the difference penalty. |
| ridge_adj | the positive tuning parameter for the ridge penalty. |
| wts | the weight vector, separate from GLM weights. |

## Value

list
same as inputs, with warnings if required.

```
pspline_fitter P-spline fitting algorithm for the GLM.
```


## Description

pspline_fitter appies the method of scoring to a variety of response distributions and link functions within for P -spline fitting within the GLM framework.

## Usage

pspline_fitter(
$y$,
B,
family = "gaussian",
link = "identity",
P ,
P_ridge $=0 * \operatorname{diag}(n c o l(B))$,
wts $=0$ * $y+1$,
m_binomial $=0$ * y + 1,
r_gamma $=0$ * y + 1
)

## Arguments

y
B

## family

link

P
wts
m_binomial
r_gamma

P_ridge ridge ("half") penalty for data augmentation, usually sqrt(lambda_r)*I (default 0).
the glm response vector of length m .
The effective P-spline regressors, e.g. B for B-splines, $\mathrm{Q}=\mathrm{X} \% * \% \mathrm{~B}$ for PSR.
the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" distribution; quotes are needed (default family = "gaussian".)
the link function, one of "identity", "log", "sqrt", "logit", "probit", "cloglog", "loglog", "reciprocal"; quotes are needed (default link = "identity").
P-spline ("half") penalty matrix for data augmentation, such that $P^{\prime} P=1$ ambda D'D.
the weight vector of length $(y)$, separate from GLM weights.
a vector of binomial trials having length $(y)$, when family = "binomial". Default is 1 vector.
a vector of gamma shape parameters, when family = "Gamma". Default is 1 vector.

## Value

coef the estimated P-spline coefficient regressor, using the effective regressors.
w $w t s * w$, GLM weight vector times input weights of length $m$.
$f \quad$ the lsfit object using data augmentation to get $P$-spline coefficient estimates. eta the linear predictor from $f$.

## Description

psPoisson is used to smooth scattered Poisson data using P-splines with a log link function.

## Usage

psPoisson(
x ,
$y$,
$x l=\min (x)$,
$x r=\max (x)$,
nseg $=10$,
bdeg $=3$,
pord $=2$,
lambda $=1$,
wts = NULL,
show $=$ FALSE,
iter = 100,
xgrid $=100$
)

## Arguments

pord the number of the order of the difference penalty, usually 1,2 (default), or 3 .
lambda the (positive) number for the tuning parameter for the penalty (default 1 ).
x
y
xl
xr
nseg

## bdeg

wts
show
iter
xgrid
the vector for the continuous regressor of length $(y)$ and the abcissae used to build the B -spline basis.
the response vector, usually count data.
$1 \quad$ the number for the min along $x($ default is $\min (x))$.
$r \quad$ the number for the max along $x$ (default is $\max (x)$ ).
seg the number of evenly spaced segments between $x l$ and $x r$ (default 10).
the vector of general weights, zeros are allowed (default 1).
Set to TRUE or FALSE to display iteration history (default FALSE).
a scalar to set the maximum number of iterations, default iter=100.
a scalar or a vector that gives the $x$ locations for prediction, useful for plotting. If a scalar (default 100) is used then a uniform grid of this size along ( $\mathrm{xl}, \mathrm{xr}$ ).

## Value

| pcoef | a vector of length $n$ of estimated P-spline coefficients. |
| :--- | :--- |
| muhat | a vector of length $m$ of estimated means. |
| B | the $m$ by $n$ B-spline basis. |
| deffdim | deviance of fit. |
| aic | effective dimension of fit. |
| wts | AIC. |
| nseg | the vector of given prior weights. |
| bdeg | the degree of the B-spline basis. |
| pord | the order of the difference penalty. |
| lambda | the positive tuning parameter. |
| family | the family of the response ("Poisson"). |
| link | the link function used ("log"). |
| xgrid | gridded $x$ values, useful for plotting. |
| ygrid | gridded fitted linear predictor values, useful for plotting. |
| mugrid | gridded (inverse link) fitted mean values, useful for plotting. |
| se_eta | gridded standard errors for the linear predictor. |
| dispersion | Dispersion parameter estimated dev/(m-effdim). |

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

Eilers, P.H.C., Marx, B.D., and Durban, M. (2015). Twenty years of P-splines, SORT, 39(2): 149186.

## Examples

```
library(JOPS)
library(boot)
# Extract the data
Count <- hist(coal$date, breaks = c(1851:1963), plot = FALSE)$counts
Year <- c(1851:1962)
xl <- min(Year)
xr <- max(Year)
# Poisson smoothing
nseg <- 20
```

```
    bdeg <- 3
    fit1 <- psPoisson(Year, Count, xl, xr, nseg, bdeg, pord = 2, lambda = 1)
    plot(fit1, xlab = "Year", ylab = "Count", se = 2)
```

psSignal Smooth signal (multivariate calibration) regression using $P$-splines.

## Description

Smooth signal (multivariate calibration) regression using P-splines.

## Usage

```
    psSignal(
```

        \(y\),
        x_signal,
        x_index = c(1:ncol(x_signal)),
        nseg \(=10\),
        bdeg \(=3\),
        pord \(=3\),
        lambda \(=1\),
        wts = \(1+0\) * \(y\),
        family = "gaussian",
        link = "default",
        m_binomial \(=1+0 * y\),
        r_gamma = wts,
        y_predicted \(=\) NULL,
        x_predicted = x_signal,
        ridge_adj = 0,
        int = TRUE
    )

## Arguments

$y$

x_index
nseg the number of evenly spaced segments between xl and xr (default 10).

## bdeg

pord
lambda
wts
$\mathrm{a}(\mathrm{glm})$ response vector, usually continuous, binomial or count data.
a matrix of continuous regressor with nrow(x_signal) == length(y), often a discrete digitization of a signal or histogram or time series. a vector to of length ncol ( $x$ _signal) $==p$, associated with the ordering index of the signal. Default is $1: n c o l\left(x_{-}\right.$signal). the degree of the basis, usually 1,2 , or 3 (defalult). the (positive) tuning parameter for the penalty (default 1 ). the weight vector of length $(\mathrm{y})$; default is 1 .
$\left.\begin{array}{ll}\text { family } & \begin{array}{l}\text { the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" } \\ \text { distribution; quotes are needed. Default is "gaussian". }\end{array} \\ \text { link } & \begin{array}{l}\text { the link function, one of "identity", "log", "sqrt", "logit", "probit", } \\ \text { "cloglog", "loglog", "reciprocal"; quotes are needed (default "identity"). }\end{array} \\ \text { m_binomial } & \begin{array}{l}\text { a vector of binomial trials having length(y); default is } 1 \text { vector for family }= \\ \text { "binomial", NULL otherwise. }\end{array} \\ \text { y_predicted } & \begin{array}{l}\text { a vector of gamma shape parameters. Default is } 1 \text { vector for family = "Gamma", } \\ \text { NULL otherwise. }\end{array} \\ \text { x_predicted } & \begin{array}{l}\text { standard error of external prediction. Default is NULL. }\end{array} \\ \text { a matrix of external signals to yield external prediction. }\end{array}\right]$

## Details

Support functions needed: pspline_fitter, bbase and pspline_checker.


```
y_intercept the estimated y-intercept (when int = TRUE.)
int a logical variable related to use of y-intercept in model.
dispersion_param
    estimate of dispersion, Dev/df_resid.
summary_predicted
inverse link prediction vectors, and twice se bands.
eta_predicted estimated linear predictor of length ( \(y\) ).
press_mu leave-one-out prediction of mean, when family = "gaussian", NULL other-
    wise.
bin_percent_correct
                                    percent correct classification based on 0.5 cut-off, when family = binomial,
                                    NULL otherwise.
\(x\) index a vector to of length ncol (x_signal) \(==\mathrm{p}\), associated with the ordering of the
    signal.
```


## Author(s)

Brian Marx

## References

Marx, B.D. and Eilers, P.H.C. (1999). Generalized linear regression for sampled signals and curves: A P-spline approach. Technometrics, 41(1): 1-13.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
# Get the data
library(fds)
data(nirc)
iindex <- nirc$x
x <- nirc$y
sel <- 50:650 # 1200 <= x & x<= 2400
X <- X[sel, ]
iindex <- iindex[sel]
dX <- diff(X)
diindex <- iindex[-1]
y <- as.vector(labc[1, 1:40]) # percent fat
oout <- 23
dX <- t(dX[, -oout])
y <- y[-oout]
fit1 <- psSignal(y, dX, diindex, nseg = 25, bdeg = 3, lambda = 0.0001,
pord = 2, family = "gaussian", link = "identity", x_predicted = dX, int = TRUE)
plot(fit1, xlab = "Coefficient Index", ylab = "ps Smooth Coeff")
title(main = "25 B-spline segments with tuning = 0.0001")
names(fit1)
```


## Description

psVCSignal is used to regress a (glm) response onto a signal such that the signal coefficients can vary over another covariate $t$. Anisotripic penalization of tensor product B-splines produces a 2 D coefficient surface that can be sliced at $t$.
@details Support functions needed: pspline_fitter, pspline_2dchecker, and bbase.
@import stats

```
Usage
    psVCSignal(
        y,
        X,
        x_index,
        t_var,
        Pars = rbind(c(min(x_index), max(x_index), 10, 3, 1, 2), c(min(t_var), max(t_var),
            10, 3, 1, 2)),
    family = "gaussian",
    link = "default",
    m_binomial = 1 + 0 * y,
    wts = 1 + 0 * y,
    r_gamma = 1 + 0 * y,
    X_pred = X,
    t_pred = t_var,
    y_predicted = NULL,
    ridge_adj = 1e-08,
    int = TRUE
    )
```


## Arguments

| $y$ | a glm response vector of length m , usually continuous, binary/bimomial or counts. |
| :---: | :---: |
| X | a m by p1 Signal matrix of regressors. |
| x_index | p1-vector for index of Signal (e.g. wavelength). |
| t_var | p2-vector with other (indexing) variable in coefficient surface (e.g. temperature, depth, time). |
| Pars | a matrix with 2 rows, each with P-spline parameters: min max nseg bdeg lambda pord, for row and columns of tensor product surface; defaults are min and max for $x_{\text {_index }}$ and $t_{\text {_var }}$ (resp.), nseg $=10$, bdeg $=3$, lambda $=1$, pord $=2$. |
| family | the response distribution, e.g. "gaussian", "binomial", "poisson", "Gamma" distribution; quotes are needed (default "gaussian". |


| link | the link function, one of "identity", "log", "sqrt", "logit", "probit", "cloglog", "loglog", "reciprocal"); quotes are needed (default "identity". |
| :---: | :---: |
| m_binomial | a vector of binomial trials having length ( $y$ ). Default is 1 vector for family $=$ "binomial", NULL otherwise. |
| wts | a m vector of weights (default 1 ). |
| r_gamma | a vector of gamma shape parameters. Default is 1 vector for family = "Gamma", NULL otherwise. |
| X_pred | a matrix of signals with $n \operatorname{col}(X)$ columns for prediction, default is $X$. |
| t_pred | a vector for the VC indexing variable with length nrow(X_pred), default is t_var. |
| y_predicted | a vector for the responses associated with X_pred with length nrow(X_pred) useful for CV when family = "binomial", default is NULL. |
| ridge_adj | a small ridge penalty tuning parameter to regularize estimation (default $1 \mathrm{e}-8$ ). |
| int | intercept set to TRUE or FALSE for intercept term. |

## Value

pcoef a vector of length (Pars[1,3]+Pars[1,4])*(Pars[2,3]+Pars[2,4]) of estimated P -spline coefficients for tensor surface.
summary_predicted
inverse link prediction vectors, and twice se bands.
dev the deviance of fit.
eff_dim the approximate effective dimension of fit.
family the family of the response.
link the link function.
aic AIC.
df_resid approximate df residual.
cv leave-one-out standard error prediction when family = "gaussian", NULL otherwise.
cv_predicted standard error prediction for y_predict when family = "gaussian", NULL otherwise.

Pars design and tuning parameters; see arguments above.
dispersion_parm
estimate of dispersion, Dev/df_resid.
summary_predicted
inverse link prediction vectors, and twice se bands.
eta_predicted estimated linear predictor of length ( y ).
press_mu leave-one-out prediction of mean when family = "gaussian", NULL otherwise.
bin_percent_correct
percent correct classification based on 0.5 cut-off when family = "binomial",
NULL otherwise.

| Bx | B-spline basis matrix of dimension p1 by $n$ 1, along $x \_i n d e x$. |
| :--- | :--- |
| By | B-spline basis matrix of dimension p2 by $n 2$, along t_var. |
| $Q$ | Modified tensor basis $(m$ by $(n 1 * n 2))$ for VC signal regression. |
| yint | the estimated y-intercept $($ when int $=$ TRUE.) |
| int | a logical variable related to use of $y$-intercept in model. |

## Author(s)

Paul Eilers and Brian Marx

## References

Eilers, P.H.C. and Marx, B.D. (2003). Multivariate calibration with temperature interaction using two-dimensional penalized signal regression. Chemometrics and Intellegent Laboratory Systems, 66, 159-174.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(fds)
data(nirc)
iindex <- nirc$x
X <- nirc$y
sel <- 50:650 # 1200 <= x & x<= 2400
X <- X[sel, ]
iindex <- iindex[sel]
dX <- diff(X)
diindex <- iindex[-1]
y <- as.vector(labc[1, 1:40]) # percent fat
t_var <- as.vector(labc[4, 1:40]) # percent flour
oout <- 23
dX <- t(dX[, -oout])
y <- y[-oout]
t_var = t_var[-oout]
Pars = rbind(c(min(diindex), max(diindex), 25, 3, 1e-7, 2),
c(min(t_var), max(t_var), 20, 3, 0.0001, 2))
fit1 <- psVCSignal(y, dX, diindex, t_var, Pars = Pars,
family = "gaussian", link = "identity", int = TRUE)
plot(fit1, xlab = "Coefficient Index", ylab = "VC: % Flour")
names(fit1)
```

rdw

Observations on the widths of red blood cell distributions (RDW).

## Description

Observations on the widths of red blood cell distributions (RDW).

## Usage

data(rdw)

## Format

A vector.

## Source

Erasmus University Medical Centre, Rotterdam, The Netherlands

## Examples

```
data(rdw)
hist(rdw, breaks = 20)
```

```
    rowtens Compute the row tensor product of two matrices
```


## Description

Compute the row tensor product of two matrices with identical numbers of rows.

## Usage

rowtens $(X, Y=X)$

## Arguments

| $X$ | a numeric matrix. |
| :--- | :--- |
| $Y$ | a numeric matrix (if missing, $Y=x)$. |

## Details

The input matrices must have the same number of rows, say $m$. If their numbers of columns are n 1 and n 2 , the result is a matrix with $m$ rows and $n 1 * n 2$ columns. Each row of the result is the Kronecker product of the corresponding rows of $X$ and $Y$.

## Value

The row-wise tensor product of the two matrices.

## Author(s)

Paul Eilers

## References

Eilers, P. H. C. and Currie, I. D. and Durban, M. (2006) Fast and compact smoothing on large multidimensional grids CSDA 50, 61-76.

```
save_PDF Save a plot as a PDF file.
```


## Description

Save a plot as a PDF file in a (default) folder. The present default is determined by the folder structure for the production of the book.

## Usage

```
    save_PDF(
```

        fname = "scratch",
        folder = "../../Graphs",
        show \(=\mathrm{T}\),
        width = 6,
        height \(=4.5\)
    )
    
## Arguments

fname the file name without the extension PDF (default: scratch).
folder the folder for saving PDF plots (default . . / . ./Graphs).
show a logical parameter; if TRUE the full file name will be displayed.
width figure width in inches (default $=6$ ).
height $\quad$ figure height in inches $($ default $=4.5)$.

## Value

save a plot as a PDF file.

## Author(s)

Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

```
set_panels Prepare graphics layout for multiple panels
```


## Description

Adapt margins and axes layout for multiple panels.

## Usage

set_panels(rows = 1, cols = 1)

## Arguments

| rows | number of rows. |
| :--- | :--- |
| cols | number of columns. |

## Value

Prepare graphics layout for multiple panels

## Author(s)

Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

```
set_window Open a graphics window.
```


## Description

Open a a window for graphics, with specified width and height.

## Usage

set_window(width $=6$, height $=4.5$, kill $=$ TRUE, noRStudioGD $=$ TRUE)

## Arguments

width figure width in inches $($ default $=6)$.
height $\quad$ figure height in inches $($ default $=4.5)$.
kill if TRUE (default) closes all graphics windows. Works only for Windows.
noRStudioGD if TRUE: do not use the RStudio device (which does not accept width and height).

## Value

open a graphics window.

## Note

Currently only works for Windows!

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.
sim_psr Single-Index signal regression using $P$-splines

## Description

sim_psr is a single-index signal regression model that estimates both the signal coefficients vector and the unknown link function using P -splines.

## Usage

sim_psr(
$y$,
X ,
x_index = c(1:ncol(X)),
nsegs $=\operatorname{rep}(10,2)$,
bdegs $=\operatorname{rep}(3,3)$,
lambdas $=\operatorname{rep}(1,2)$,
pords $=\operatorname{rep}(2,2)$,
max_iter = 100
)

## Arguments

y
$X \quad$ The signal regressors with dimension $m$ by $p$.
x_index
nsegs a vector of length 2 containing the number of evenly spaced segments between min and max, for each the coefficient vector and the (unknown) link function, resp. (default c $(10,10))$.
bdegs a vector of length 2 containing the degree of B-splines, for the coefficient vector and the (unknown) link function, resp. (default cubic or $\mathrm{c}(3,3)$ ).
lambdas a vector of length 2 containing the positive tuning parameters, for each the coefficient vector and the (unknown) link function, resp. (default $c(1,1)$ ).

| pords | a vector of length 2 containing the difference penalty order, for each the coeffi- <br> cient vector and the (unknown) link function, resp. (defaultc $(2,2)$ ). |
| :--- | :--- |
| max_iter | a scalar for the maximum number of iterations (default 100). |

## Value

$y \quad$ the response vector of length $m$.
alpha the P-spline coefficient vector of length (nsegs[1]+bdeg[1]).
iter the number of iterations used for the single-index fit.
yint the estimated y-intercept for the single-index model.
B
the B-spline matrix built along the signal index, using nsegs[1], used for the coefficient vector.

Q the effective regressors from the psVCSignal portion of the single-index fit with dimension $m$ by length (alpha).
nsegs a vector of length 2 containing the number of evenly spaced segments between min and max, for each the coefficient vector and the link function, resp.
bdegs a vector of length 2 containing the degree of B-splines, for each the coefficient vector and the link function, resp.
lambdas a vector of length 2 containing the positive tuning parameters, for each the coefficient vector and the link function, resp.
pords a vector of length 2 containing the difference penalty order, for each the coefficient vector and the link function, resp.
eta the estimated linear predictor for the single-index fit.
cv the leave-one-out cross-validation statistic or the standard error of prediction for the single-index fit.
delta_alpha change measure in signal-coefficent parameters at convervence.
$x$ index the index of length $p$ for columns of signal matrix.
f_fit the psNormal object, fitting link function $f(e t a)$.
f_eta the predicted values of the link function estimated with f_fit or estimated $\mathrm{f}(\mathrm{eta})$, at $\mathrm{x}=\mathrm{e}$ ta.

## Author(s)

Paul Eilers, Brian Marx, and Bin Li

## References

Eilers, P.H.C., B. Li, B.D. Marx (2009). Multivariate calibration with single-index signal regression, Chemometrics and Intellegent Laboratory Systems, 96(2), 196-202.

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
# Get the data
library(fds)
data(nirc)
iindex <- nirc$x
x <- nirc$y
sel <- 50:650 # 1200 <= x & x<= 2400
X <- X[sel, ]
iindex <- iindex[sel]
dX <- diff(X)
diindex <- iindex[-1]
y <- as.vector(labc[1, 1:40])
oout <- 23
dX <- t(dX[, -oout])
y <- y[-oout]
pords <- c(2, 2)
nsegs <- c(27, 7)
bdegs = c(3, 3)
lambdas <- c(1e-6, .1)
max_iter <- 100
# Single-index model
fit <- sim_psr(y, dX, diindex, nsegs, bdegs, lambdas, pords,
    max_iter)
plot(fit, xlab = "Wavelength (nm)", ylab = " ")
```

sim_vcpsr Varying-coefficient single-index signal regression using tensor $P$ -
splines.

## Description

sim_vcpsr is a varying-coefficient single-index signal regression approach that allows both the signal coefficients and the unknown link function to vary with an indexing variable $t$, e.g. temperature. Two surfaces are estimated (coefficent and link) that can be sliced at arbitary t. Anisotripic penalization with P -splines is used on both.

## Usage

sim_vcpsr(
y ,
X,
t_var,
x_index $=c(1: \operatorname{ncol}(X))$,
nsegs $=\operatorname{rep}(10,4)$,

```
    bdegs = rep(3, 4),
    lambdas = rep(1, 4),
    pords = rep(2, 4),
    max_iter = 100,
    mins = c(min(x_index), min(t_var)),
    maxs = c(max(x_index), max(t_var))
)
```


## Arguments

y
X
t_var
x_index
nsegs
bdegs a vector of length 4 containing the degree of $B$-splines, for each the coefficient surface (row and col) and link surface (row and col), resp. (default cubic $\operatorname{rep}(3,4)$ ).
lambdas a vector of length 4 containing the positive tuning parameters, for each the coefficient surface (row and col) and link surface (row and col), resp. (default $\operatorname{rep}(1,4))$.
pords a vector of length 4 containing the difference penalty order, for each the coefficient surface (row and col) and link surface (row and col), resp. (default $\operatorname{rep}(2,4)$ ).
max_iter a scalar for the maximum number of iterations (default 100)
mins A vector length 2, containing min for signal index and $t$ _var, default associated with $x$ _index and $t$ _var minimums; default is respective minimums.
A vector length 2, containing max for signal index and t_var, default associated with $x$ _index and $t \_v a r$ maximums; default is respective maximums.

## Value

y
alpha
iter
yint
Bx

By the B-spline matrix built along the $t_{\text {_ }}$ var index, using nsegs[2], used for the coefficient surface.
Q the effective regressors from the psVCSignal portion of the single-index fit with dimension $m$ by length(alpha).

| t_var | the VC indexing variable of length m . |
| :---: | :---: |
| nsegs | a vector of length 4 containing the number of evenly spaced segments between min and max, for each the coefficient surface (row and col) and link surface (row and col). |
| bdegs | a vector of length 4 containing the degree of B-splines, for each the coefficient surface (row and col) and link surface (row and col). |
| lambdas | a vector of length 4 containing the positive tuning parameters, for each the coefficient surface (row and col) and link surface (row and col). |
| pords | a vector of length 4 containing the difference penalty order, for each the coefficient surface (row and col) and link surface (row and col). |
| mins | a vector length 2, containing min for signal index and t_var. |
| maxs | a vector length 2, containing max for signal index and t_var. |
| eta | the estimated linear predictor for the single-index fit. |
| Pars | a matrix of 2 rows associated with the signal coefficient surface design parameters, each row: $\mathrm{c}(\mathrm{min}, \max , \mathrm{nseg}, \mathrm{bdeg}$, lambda, pord) for linear predictor x_index and t_var, resp. |
| pPars | a matrix of 2 rows associated with the link function design parameters, each row: $c$ (min, max, nseg, bdeg, lambda, pord) for linear predictor eta and t_var, resp. |
| cv | the leave-one-out cross-validation statistic or the standard error of prediction for the single-index fit. |
| delta_alpha | change measure in signal-coefficent parameters at convergence. |
| fit2D | ps2DNormal object, fitting f(eta, $\mathrm{t}_{-}$var). |

## Author(s)

Paul Eilers and Brian Marx

## References

Marx, B. D. (2015). Varying-coefficient single-index signal regression. Chemometrics and Intelligent Laboratory Systems, 143, 111-121.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
# Load libraries
library(fields) # Needed for plotting
# Get the data
Dat <- Mixture
# Dimensions: observations, temperature index, signal
m <- 34
p1 <- 401
p2 <- 12
```

```
# Stacking mixture data, each mixture has 12 signals stacked
# The first differenced spectra are also computed.
mixture_data <- matrix(0, nrow = p2 * m, ncol = p1)
for (ii in 1:m)
{
    mixture_data[((ii - 1) * p2 + 1):(ii * p2), 1:p1] <-
        t(as.matrix(Dat$xspectra[ii, , ]))
    d_mixture_data <- t(diff(t(mixture_data)))
}
# Response (typo fixed) and index for signal
y_mixture <- Dat$fractions
y_mixture[17, 3] <- 0.1501
index_mixture <- Dat$wl
# Select response and replicated for the 12 temps
# Column 1: water; 2: ethanediol; 3: amino-1-propanol
y <- as.vector(y_mixture[, 2])
y <- rep(y, each = p2)
bdegs = c(3, 3, 3, 3)
pords <- c(2, 2, 2, 2)
nsegs <- c(12, 5, 5, 5) # Set to c(27, 7, 7 , 7) for given lambdas
mins <- c(700, 30)
maxs <- c(1100, 70)
lambdas <- c(1e-11, 100, 0.5, 1) # based on svcm search
x_index <- seq(from = 701, to = 1100, by = 1) # for dX
t_var_sub <- c(30, 35, 37.5, 40, 45, 47.5, 50, 55, 60, 62.5, 65, 70)
t_var <- rep(t_var_sub, m)
max_iter <- 2 # Set higher in practice, e.g. 100
int <- TRUE
# Defining x as first differenced spectra, number of channels.
x <- d_mixture_data
# Single-index VC model using optimal tuning
fit <- sim_vcpsr(y, x, t_var, x_index, nsegs, bdegs, lambdas, pords,
    max_iter = max_iter, mins = mins, maxs = maxs)
plot(fit, xlab = "Wavelength (nm)", ylab = "Temp C")
```


## Description

Two-dimensional smoothing of scattered data points with tensor product P -splines.

## Usage

```
SpATS.nogeno(
        response,
        spatial,
        fixed = NULL,
        random \(=\) NULL,
        data,
        family = gaussian(),
        offset = 0,
        weights = NULL,
        control = list(maxit = 100)
    )
```


## Arguments

| response | a character string with the name of the variable that contains the response vari- <br> able of interest. |
| :--- | :--- |
| spatial | a right hand formula object specifying the spatial P-Spline model. See SAP and <br> PSANOVA for more details about how to specify the spatial trend. <br> an optional right hand formula object specifying the fixed effects. |
| fixed | an optional right hand formula object specifying the random effects. Currently, <br> only sets of independent and identically distributed random effects can be incor- <br> porated. |
| data | a data frame containing the variables. |
| family | object of class family specifying the distribution and link function. <br> an optional numerical vector containing an a priori known component to be <br> included in the linear predictor during fitting. |
| weights | an optional numerical vector of weights to be used in the fitting process. By <br> default, the weights are considered to be one. |
| control | a list of control values. |

## Details

This function is a modified version of the function SpATS in the package SpATS. The difference is that genotypes have been removed.

## Value

A list with the following components:

| call | the matched call. |
| :--- | :--- |
| data | the original supplied data argument with a new column with the weights used <br> during the fitting process. |
| model | a list with the model components: response, spatial, fixed and/or random. |
| fitted | a numeric vector with the fitted values. |


| residuals | a numeric vector with deviance residuals. |
| :--- | :--- |
| a two-length vector with the values of the dispersion parameters at convergence. |  |
| For Gaussian responses both elements coincide, being the (REML) estimate of |  |
| dispersion parameter. For non-Gaussian responses, the result depends on the ar- |  |
| gument update.psi of the controlSpATS function. If this argument was spec- |  |
| ified to FALSE (the default), the first component of the vector corresponds to the |  |
| default value used for the dispersion parameter (usually 1). The second element, |  |
| correspond to the (REML) estimate of the dispersion parameter at convergence. |  |
| If the argument update.psi was specified to TRUE, both components coincide |  |
| (as in the Gaussian case). |  |
| a numeric vector with the (REML) variance component estimates. This vector |  |
| contains the variance components associated with the spatial trend, as well as |  |
| those related with the random model terms. |  |

## Author(s)

Maria-Xose Rodriguez-Alvarez and Paul Eilers

## References

Rodriguez-Alvarez, M.X, Boer, M.P., van Eeuwijk, F.A., and Eilers, P.H.C. (2018). Correcting for spatial heterogeneity in plant breeding experiments with P-splines. Spatial Statistics, 23, 52-71. https://doi.org/10.1016/j.spasta.2017.10.003.

## Examples

```
# Get the data
library(SemiPar)
data(ethanol)
# Fit the PS-ANOVA model
ps2d <- SpATS.nogeno(response = "NOx",
    spatial = ~PSANOVA(E, C, nseg = c(20, 20), nest.div = c(2, 2)),
    data = ethanol,
    control = list(maxit = 100, tolerance = 1e-05,
    monitoring = 0, update.psi = FALSE))
# Report effective dimensions, if desired
# print(summary(ps2d))
# Compute component surface and their sum on a fine grid
Tr = obtain.spatialtrend(ps2d, grid = c(100, 100))
# Plot surface and contours
image(Tr$row.p, Tr$col.p, Tr$fit, col = terrain.colors(100), xlab = 'C', ylab = 'E')
contour(Tr$row.p, Tr$col.p, Tr$fit, add = TRUE, col = 'blue')
points(ethanol$C, ethanol$E, pch = '+')
```

spbase

## Compute a sparse B-spline basis on evenly spaced knots

## Description

Constructs a sparse B-spline basis on evenly spaced knots.

## Usage

$\operatorname{spbase}(x, x l=\min (x), x r=\max (x), n s e g=10, b d e g=3)$

## Arguments

$x \quad a$ vector of argument values, at which the B-spline basis functions are to be evaluated.
$x l \quad$ the lower limit of the domain of $x($ default $\min (x))$.
$x r \quad$ the upper limit of the domain of $x($ default $\max (x))$.
nseg the number of evenly spaced segments between $x l$ and $x r$ (default 10).
bdeg the degree of the basis, usually 1,2 , or 3 (default).

## Value

A sparse matrix (in spam format) with length( $x$ ) of rows= and nseg + bdeg columns.

## Author(s)

Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (1996). Flexible smoothing with B-splines and penalties (with comments and rejoinder), Statistical Science, 11: 89-121.
Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

library(JOPS)
\# Basis on grid
$x=\operatorname{seq}(0,4$, length $=1000)$
$B=\operatorname{spbase}(x, 0,4$, nseg $=50$, bdeg = 3)
$n b 1=\operatorname{ncol}(B)$
matplot(x, B, type = 'l', lty = 1, lwd = 1, xlab = 'x', ylab = '')
cat('Dimensions of $B:$ ', nrow(B), 'by', ncol(B), 'with', length(B@entries), 'non-zero elements' )
Sugar Sugar Processing Data

## Description

Sugar was sampled continuously during eight hours to make a mean sample representative for one "shift" (eight hour period). Samples were taken during the three months of operation (the so-called campaign) in late autumn from a sugar plant in Scandinavia giving a total of 268 samples. The sugar was sampled directly from the final unit operation (centrifuge) of the process.

## Usage

data(Sugar)

## Format

A list consisting of the following:
y a $268 \times 3$ matrix of quality parameters: date, color, ash*1000
X fluoresence array, 268 (observations) x [571 (emission channels) x 7 (excitation channels)]
Lab Lab information
$\operatorname{DimX}$ array dimension for $X$
Yidx names (id) for $y$
EmAx Emmission levels for axis (nm)
ExAx Excitation levels for axis (nm)
time
readmetime
Lname
LabNumber
ProcNumber
Proc
DimLab
DimProc

## Source

http://www.models.kvl.dk/Sugar_Process

## References

R. Bro, Exploratory study of sugar production using fluorescence spectroscopy and multi-way analysis, Chemom. Intell. Lab. Syst., 1999, (46), 133-147.

## Suicide Suicide Data Set

## Description

The dataset comprises lengths (in days) of psychiatric treatment spells for patients used as controls in a study of suicide risks.

## Usage

data(Suicide)

## Format

A dataframe with one column: $y$.

## Source

Silverman, B. (1986). Density Estimation for Statistics and Data Analysis. Chapman \& Hall.

## References

Silverman, B. (1986). Density Estimation for Statistics and Data Analysis. Chapman \& Hall.
tpower Compute a truncated power function.

## Description

Compute a truncated power function.

## Usage

tpower (x, knot, p)

## Arguments

$x \quad a \quad$ vector on which the basis is calculated.
knot a scalar giving the truncation point.
$p \quad a \operatorname{scalar}$ power for the basis, e.g. $p=3$ for cubic TPF.

## Value

a vector with the truncated power function.

## Author(s)

Paul Eilers

## References

Eilers, P.H.C. and Marx, B.D. (2021). Practical Smoothing, The Joys of P-splines. Cambridge University Press.

## Examples

```
library(JOPS)
# Basis on grid
x = seq(0, 4, length = 500)
knots = 0:3
Y = outer(x, knots, tpower, 1)
matplot(x, Y, type ='l', lwd = 2, xlab = 'x', ylab = '',
main ='Linear TPF basis')
```


## Description

Brightness of a variable star.

## Usage

data(Varstar)

## Format

A dataframe with eleven columns (V1-V11):
V1 day index
V2 brightness
V3-V11 Paul Eilers, personal communication.

## References

Paul Eilers (personal communication).

Woodsurf Profile of a sanded piece of wood.

## Description

Profile of a sanded piece of wood.

## Usage

data(Woodsurf)

## Format

A data frame with one column: $y$.

## Source

Pandit, S.M. and Wu, S.M. (1993). Time Series and System Analysis with Applications. Krieger Publishing Company.

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