## Package: refund (via r-universe)

September 11, 2024

Type Package

Title Regression with Functional Data

**Version** 0.1-35

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**Depends** R (>= 3.5.0)

**Imports** fda, Matrix, lattice, boot, mgcv (>= 1.9), MASS, magic, nlme, gamm4, lme4, RLRsim, splines, grpreg, ggplot2, stats, pbs, methods

Suggests RColorBrewer, reshape2, testthat

**Description** Methods for regression for functional data, including function-on-scalar, scalar-on-function, and function-on-function regression. Some of the functions are applicable to image data.

License GPL (>= 2)

LazyLoad yes

LazyData true

URL https://github.com/refunders/refund

BugReports https://github.com/refunders/refund/issues

Collate 'Omegas.R' 'af.R' 'af\_old.R' 'amc.R' 'ccb.fpc.R' 
'create.prep.func.R' 'coefficients.pfr.R' 'dt\_basis.R' 
'irreg2mat.R' 'fbps.R' 'fgam.R' 'fosr.R' 'fosr.perm.R' 
'fosr.perm.fit.R' 'fosr.perm.test.R' 'fosr.vs.R' 'fosr2s.R' 
'fpc.R' 'fpca2s.R' 'fpca.sc.R' 'fpca.face.R' 'fpca.ssvd.R' 
'fpcr.R' 'fpcr.setup.R' 'lf.R' 'lf\_old.R' 'lf.vd.R' 'lofocv.R' 
'lpeer.R' 'lpfr.R' 'quadWeights.R' 'lw.test.R' 'osplinepen2d.R' 
'parse.predict.pfr.R' 'peer.R' 'peer\_old.R' 'pffr-ff.R' 
'pffr-ffpc.R' 'pffr-methods.R' 'pffr-pcre.R' 'pffr-robust.R' 
'pffr-sff.R' 'pffr-utilities.R' 'pffr.R' 'pfr.R' 'pfr\_old.R' 
'pi\_basis.R' 'plot.fosr.R' 'plot.fosr.perm.R' 'plot.fosr.vs.R' 
'plot.fpcr.R' 'plot.lpeer.R' 'plot.peer.R' 'plot.pfr.R' 
'poridge.R' 'postprocess.pfr.R' 'predict.fgam.R'

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'preprocess.pfr.R' 'pspline.setting.R' 'pwcv.R' 'summary.pfr.R'
're.R' 'rlrt.pfr.R' 'vis.fgam.R' 'predict.fosr.vs.R'
'CD4-data.R' 'content-data.R' 'COVID19-data.R' 'DTI-data.R'
'DTI2-data.R' 'PEER.Sim-data.R' 'gasoline-data.R' 'vis.pfr.R'
'GLS_CS.R' 'Gibbs_CS_FPCA.R' 'Gibbs_CS_Wish.R'
'Gibbs_Mult_FPCA.R' 'Gibbs_Mult_Wish.R' 'OLS_CS.R'
'VB_CS_FPCA.R' 'VB_CS_Wish.R' 'VB_Mult_FPCA.R' 'VB_Mult_Wish.R'
'XtSiginvX.R' 'bayes_fosr.R' 'f_sum.R' 'f_sum2.R' 'f_sum4.R'
'f_trace.R' 'mfpca.sc.R' 'mfpca.face.R' 'face.Cov.mfpca.R'
'fpca.lfda.R' 'predict.fbps.R' 'select_knots.R'

RoxygenNote 7.2.3

**Encoding** UTF-8 **Repository** https://refunders.r-universe.dev

RemoteUrl https://github.com/refunders/refund

RemoteRef HEAD

**RemoteSha** 3b13697891f489e19b01afbdcb0cb856f22c69ab

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af

Construct an FGAM regression term

## Description

Defines a term  $\int_T F(X_i(t),t) dt$  for inclusion in an mgcv::gam-formula (or bam or gamm or gamm4:::gamm) as constructed by pfr, where F(x,t) is an unknown smooth bivariate function and  $X_i(t)$  is a functional predictor on the closed interval T. See smooth.terms for a list of bivariate basis and penalty options; the default is a tensor product basis with marginal cubic regression splines for estimating F(x,t).

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## Usage

```
af(
   X,
   argvals = NULL,
   xind = NULL,
   basistype = c("te", "t2", "s"),
   integration = c("simpson", "trapezoidal", "riemann"),
   L = NULL,
   presmooth = NULL,
   presmooth.opts = NULL,
   Xrange = range(X, na.rm = T),
   Qtransform = FALSE,
   ...
)
```

# Arguments X

functional predictors, typically expressed as an N by J matrix, where N is the

number of columns and J is the number of evaluation points. May include missing/sparse functions, which are indicated by NA values. Alternatively, can be an

object of class "fd"; see fd.

argvals indices of evaluation of X, i.e.  $(t_{i1}, ..., t_{iJ})$  for subject i. May be entered as either a

length-J vector, or as an N by J matrix. Indices may be unequally spaced. Entering as a matrix allows for different observations times for each subject. If NULL, defaults to an equally-spaced grid between 0 or 1 (or within X\$basis\$rangeval

if X is a fd object.)

xind same as argvals. It will not be supported in the next version of refund.

basistype defaults to "te", i.e. a tensor product spline to represent F(x,t) Alternatively,

use "s" for bivariate basis functions (see s) or "t2" for an alternative parame-

terization of tensor product splines (see t2)

integration method used for numerical integration. Defaults to "simpson"'s rule for calcu-

lating entries in L. Alternatively and for non-equidistant grids, "trapezoidal"

or "riemann".

L an optional N by ncol(argvals) matrix giving the weights for the numerical

integration over t. If present, overrides integration.

presmooth string indicating the method to be used for preprocessing functional predictor

prior to fitting. Options are fpca.sc, fpca.face, fpca.ssvd, fpca.bspline, and fpca.interpolate. Defaults to NULL indicateing no preprocessing. See

create.prep.func.

presmooth.opts list including options passed to preprocessing method create.prep.func.

Xrange numeric; range to use when specifying the marginal basis for the *x*-axis. It may

be desired to increase this slightly over the default of range(X) if concerned about predicting for future observed curves that take values outside of range(X)

Qtransform logical; should the functional be transformed using the empirical cdf and apply-

ing a quantile transformation on each column of X prior to fitting?

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optional arguments for basis and penalization to be passed to the function indicated by basistype. These could include, for example, "bs", "k", "m", etc. See te or s for details.

#### Value

A list with the following entries:

call a "call" to te (or s, t2) using the appropriately constructed covariate and

weight matrices.

argvals the argvals argument supplied to af

L the matrix of weights used for the integration

xindname the name used for the functional predictor variable in the formula used by mgcv

tindname the name used for argvals variable in the formula used by mgcv

Lname the name used for the L variable in the formula used by mgcv

presmooth the presmooth argument supplied to af Xrange the Xrange argument supplied to af

prep. func a function that preprocesses data based on the preprocessing method specified

in presmooth. See create.prep.func

#### Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com>, Fabian Scheipl, and Jonathan Gellar

## References

McLean, M. W., Hooker, G., Staicu, A.-M., Scheipl, F., and Ruppert, D. (2014). Functional generalized additive models. *Journal of Computational and Graphical Statistics*, **23** (1), pp. 249-269. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3982924/.

#### See Also

```
pfr, lf, mgcv's linear.functional.terms, pfr for examples
```

## **Examples**

```
## Not run:
data(DTI)
## only consider first visit and cases (no PASAT scores for controls)
DTI1 <- DTI[DTI$visit==1 & DTI$case==1,]
DTI2 <- DTI1[complete.cases(DTI1),]

## fit FGAM using FA measurements along corpus callosum
## as functional predictor with PASAT as response
## using 8 cubic B-splines for marginal bases with third
## order marginal difference penalties
## specifying gamma > 1 enforces more smoothing when using
## GCV to choose smoothing parameters
fit1 <- pfr(pasat ~ af(cca, k=c(8,8), m=list(c(2,3), c(2,3)),</pre>
```

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```
presmooth="bspline", bs="ps"),
            method="GCV.Cp", gamma=1.2, data=DTI2)
plot(fit1, scheme=2)
vis.pfr(fit1)
## af term for the cca measurements plus an lf term for the rcst measurements
## leave out 10 samples for prediction
test <- sample(nrow(DTI2), 10)</pre>
fit2 \leftarrow pfr(pasat \sim af(cca, k=c(7,7), m=list(c(2,2), c(2,2)), bs="ps",
                       presmooth="fpca.face") +
                    lf(rcst, k=7, m=c(2,2), bs="ps"),
            method="GCV.Cp", gamma=1.2, data=DTI2[-test,])
par(mfrow=c(1,2))
plot(fit2, scheme=2, rug=FALSE)
vis.pfr(fit2, select=1, xval=.6)
pred <- predict(fit2, newdata = DTI2[test,], type='response', PredOutOfRange = TRUE)</pre>
sqrt(mean((DTI2$pasat[test] - pred)^2))
## Try to predict the binary response disease status (case or control)
    using the quantile transformed measurements from the rcst tract
    with a smooth component for a scalar covariate that is pure noise
DTI3 <- DTI[DTI$visit==1,]</pre>
DTI3 <- DTI3[complete.cases(DTI3$rcst),]</pre>
z1 <- rnorm(nrow(DTI3))</pre>
fit3 <- pfr(case \sim af(rcst, k=c(7,7), m = list(c(2, 1), c(2, 1)), bs="ps",
                      presmooth="fpca.face", Qtransform=TRUE) +
                    s(z1, k = 10), family="binomial", select=TRUE, data=DTI3)
par(mfrow=c(1,2))
plot(fit3, scheme=2, rug=FALSE)
abline(h=0, col="green")
# 4 versions: fit with/without Qtransform, plotted with/without Qtransform
fit4 <- pfr(case ~ af(rcst, k=c(7,7), m = list(c(2, 1), c(2, 1)), bs="ps",
                      presmooth="fpca.face", Qtransform=FALSE) +
                    s(z1, k = 10), family="binomial", select=TRUE, data=DTI3)
par(mfrow=c(2,2))
zlms < -c(-7.2,4.3)
plot(fit4, select=1, scheme=2, main="QT=FALSE", zlim=zlms, xlab="t", ylab="rcst")
plot(fit4, select=1, scheme=2, Qtransform=TRUE, main="QT=FALSE", rug=FALSE,
     zlim=zlms, xlab="t", ylab="p(rcst)")
plot(fit3, select=1, scheme=2, main="QT=TRUE", zlim=zlms, xlab="t", ylab="rcst")
plot(fit3, select=1, scheme=2, Qtransform=TRUE, main="QT=TRUE", rug=FALSE,
     zlim=zlms, xlab="t", ylab="p(rcst)")
vis.pfr(fit3, select=1, plot.type="contour")
## End(Not run)
```

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#### **Description**

Defines a term  $\int_T F(X_i(t),t) dt$  for inclusion in an mgcv::gam-formula (or bam or gamm or gamm4:::gamm) as constructed by fgam, where F(x,t)\$ is an unknown smooth bivariate function and  $X_i(t)$  is a functional predictor on the closed interval T. Defaults to a cubic tensor product B-spline with marginal second-order difference penalties for estimating F(x,t). The functional predictor must be fully observed on a regular grid

## Usage

#### **Arguments**

X	an N by J=ncol(argvals) matrix of function evaluations $X_i(t_{i1}),.,X_i(t_{iJ}); i=1,.,N$ .
argvals	matrix (or vector) of indices of evaluations of $X_i(t)$ ; i.e. a matrix with $i$ th row $(t_{i1},.,t_{iJ})$
xind	Same as argvals. It will discard this argument in the next version of refund.
basistype	defaults to "te", i.e. a tensor product spline to represent $F(x,t)$ Alternatively, use "s" for bivariate basis functions (see s) or "t2" for an alternative parameterization of tensor product splines (see t2)
integration	method used for numerical integration. Defaults to "simpson"'s rule for calculating entries in L. Alternatively and for non-equidistant grids, "trapezoidal" or "riemann". "riemann" integration is always used if L is specified
L	optional weight matrix for the linear functional
splinepars	optional arguments specifying options for representing and penalizing the function $F(x,t)$ . Defaults to a cubic tensor product B-spline with marginal second-order difference penalties, i.e. list(bs="ps", m=list(c(2, 2), c(2, 2)), see te or s for details
presmooth	logical; if true, the functional predictor is pre-smoothed prior to fitting; see smooth.basisPar
Xrange	numeric; range to use when specifying the marginal basis for the $x$ -axis. It may be desired to increase this slightly over the default of range(X) if concerned about predicting for future observed curves that take values outside of range(X)

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Qtransform

logical; should the functional be transformed using the empirical cdf and applying a quantile transformation on each column of X prior to fitting? This ensures Xrange=c(0,1). If Qtransform=TRUE and presmooth=TRUE, presmoothing is done prior to transforming the functional predictor

#### Value

A list with the following entries:

- call a "call" to te (or s, t2) using the appropriately constructed covariate and weight matrices.
- 2. argvals the argvals argument supplied to af
- 3. L-the matrix of weights used for the integration
- 4. xindname the name used for the functional predictor variable in the formula used by mgcv.
- 5. tindname the name used for argvals variable in the formula used by mgcv
- 6. Lname the name used for the L variable in the formula used by mgcv
- 7. presmooth the presmooth argument supplied to af
- 8. Qtranform the Qtransform argument supplied to af
- 9. Xrange the Xrange argument supplied to af
- 10. ecdflist a list containing one empirical cdf function from applying ecdf to each (possibly presmoothed) column of X. Only present if Qtransform=TRUE
- 11. Xfd an fd object from presmoothing the functional predictors using smooth.basisPar. Only present if presmooth=TRUE. See fd.

#### Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com> and Fabian Scheipl

#### References

McLean, M. W., Hooker, G., Staicu, A.-M., Scheipl, F., and Ruppert, D. (2014). Functional generalized additive models. *Journal of Computational and Graphical Statistics*, **23** (1), pp. 249-269. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3982924/.

#### See Also

```
fgam, 1f, mgcv's linear.functional.terms, fgam for examples
```

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Bayesian Function-on-scalar regression

#### **Description**

Wrapper function that implements several approaches to Bayesian function- on-scalar regression. Currently handles real-valued response curves; models can include subject-level random effects in a multilevel framework. The residual curve error structure can be estimated using Bayesian FPCA or a Wishart prior. Model parameters can be estimated using a Gibbs sampler or variational Bayes.

## Usage

```
bayes_fosr(formula, data = NULL, est.method = "VB", cov.method = "FPCA", ...)
```

## **Arguments**

formula	a formula indicating the structure of the proposed model. Random intercepts are designated using re().
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
est.method	method used to estimate model parameters. Options are "VB", "Gibbs", and "GLS" with "VB" as default. Variational Bayes is a fast approximation to the full posterior and often provides good point estimates, but may be unreliable for inference. "GLS" doesn't do anything Bayesian – just fits an unpenalized GLS estimator for the specified model.
cov.method	method used to estimate the residual covariance structure. Options are "FPCA" and "Wishart", with default "FPCA"
	additional arguments that are passed to individual fitting functions.

#### Author(s)

```
Jeff Goldsmith <ajg2202@cumc.columbia.edu>
```

#### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

## **Examples**

```
## Not run:
library(reshape2)
library(dplyr)
library(ggplot2)
```

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```
##### Cross-sectional real-data examples #####
## organize data
data(DTI)
DTI = subset(DTI, select = c(cca, case, pasat))
DTI = DTI[complete.cases(DTI),]
DTI$gender = factor(sample(c("male", "female"), dim(DTI)[1], replace = TRUE))
DTI$status = factor(sample(c("RRMS", "SPMS", "PPMS"), dim(DTI)[1], replace = TRUE))
## fit models
default = bayes_fosr(cca ~ pasat, data = DTI)
VB = bayes_fosr(cca ~ pasat, data = DTI, Kp = 4, Kt = 10)
Gibbs = bayes_fosr(cca ~ pasat, data = DTI, Kt = 10, est.method = "Gibbs", cov.method = "Wishart",
                   N.iter = 500, N.burn = 200)
OLS = bayes_fosr(cca ~ pasat, data = DTI, Kt = 10, est.method = "OLS")
GLS = bayes_fosr(cca ~ pasat, data = DTI, Kt = 10, est.method = "GLS")
## plot results
models = c("default", "VB", "Gibbs", "OLS", "GLS")
intercepts = sapply(models, function(u) get(u)$beta.hat[1,])
slopes = sapply(models, function(u) get(u)$beta.hat[2,])
plot.dat = melt(intercepts); colnames(plot.dat) = c("grid", "method", "value")
ggplot(plot.dat, aes(x = grid, y = value, group = method, color = method)) +
   geom_path() + theme_bw()
plot.dat = melt(slopes); colnames(plot.dat) = c("grid", "method", "value")
ggplot(plot.dat, aes(x = grid, y = value, group = method, color = method)) +
   geom_path() + theme_bw()
## fit a model with an interaction
fosr.dti.interaction = bayes_fosr(cca ~ pasat*gender, data = DTI, Kp = 4, Kt = 10)
##### Longitudinal real-data examples #####
data(DTI2)
class(DTI2$cca) = class(DTI2$cca)[-1]
DTI2 = subset(DTI2, select = c(cca, id, pasat))
DTI2 = DTI2[complete.cases(DTI2),]
default = bayes_fosr(cca ~ pasat + re(id), data = DTI2)
VB = bayes_fosr(cca ~ pasat + re(id), data = DTI2, Kt = 10, cov.method = "Wishart")
## End(Not run)
```

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#### **Description**

Uses iterated expectation and variances to obtain corrected estimates and inference for functional expansions.

## Usage

```
ccb.fpc(
   Y,
   argvals = NULL,
   nbasis = 10,
   pve = 0.99,
   n.boot = 100,
   simul = FALSE,
   sim.alpha = 0.95
)
```

#### **Arguments**

Υ	matrix of observed functions for which estimates and covariance matrices are desired.
argvals	numeric; function argument.
nbasis	number of splines used in the estimation of the mean function and the bivariate smoothing of the covariance matrix
pve	proportion of variance explained used to choose the number of principal components to be included in the expansion.
n.boot	number of bootstrap iterations used to estimate the distribution of FPC decomposition objects.
simul	TRUE or FALSE, indicating whether critical values for simultaneous confidence intervals should be estimated
sim.alpha	alpha level of the simultaneous intervals.

#### **Details**

To obtain corrected curve estimates and variances, this function accounts for uncertainty in FPC decomposition objects. Observed curves are resampled, and a FPC decomposition for each sample is constructed. A mixed-model framework is used to estimate curves and variances conditional on each decomposition, and iterated expectation and variances combines both model-based and decomposition-based uncertainty.

#### Value

Yhat	a matrix whose rows are the estimates of the curves in Y.
Yhat.boot	a list containing the estimated curves within each bootstrap iteration.
diag.var	diagonal elements of the covariance matrices for each estimated curve.
VarMats	a list containing the estimated covariance matrices for each curve in Y.
crit.val	estimated critical values for constructing simultaneous confidence intervals.

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#### Author(s)

Jeff Goldsmith < jeff.goldsmith@columbia.edu>

#### References

Goldsmith, J., Greven, S., and Crainiceanu, C. (2013). Corrected confidence bands for functional data using principal components. *Biometrics*, 69(1), 41–51.

### **Examples**

```
## Not run:
data(cd4)
# obtain a subsample of the data with 25 subjects
set.seed(1236)
sample = sample(1:dim(cd4)[1], 25)
Y.sub = cd4[sample,]
# obtain a mixed-model based FPCA decomposition
Fit.MM = fpca.sc(Y.sub, var = TRUE, simul = TRUE)
# use iterated variance to obtain curve estimates and variances
Fit.IV = ccb.fpc(Y.sub, n.boot = 25, simul = TRUE)
# for one subject, examine curve estimates, pointwise and simultaneous itervals
EX = 2
EX.IV = cbind(Fit.IV$Yhat[EX,],
      Fit.IV$Yhat[EX,] + 1.96 * sqrt(Fit.IV$diag.var[EX,]),
      Fit.IV$Yhat[EX,] - 1.96 * sqrt(Fit.IV$diag.var[EX,]),
      Fit.IV$Yhat[EX,] + Fit.IV$crit.val[EX] * sqrt(Fit.IV$diag.var[EX,]),
      Fit.IV$Yhat[EX,] - Fit.IV$crit.val[EX] * sqrt(Fit.IV$diag.var[EX,]))
EX.MM = cbind(Fit.MM$Yhat[EX,],
      Fit.MM$Yhat[EX,] + 1.96 * sqrt(Fit.MM$diag.var[EX,]),
      Fit.MM$Yhat[EX,] - 1.96 * sqrt(Fit.MM$diag.var[EX,]),
      Fit.MM$Yhat[EX,] + Fit.MM$crit.val[EX] * sqrt(Fit.MM$diag.var[EX,]),
      Fit.MM$Yhat[EX,] - Fit.MM$crit.val[EX] * sqrt(Fit.MM$diag.var[EX,]))
# plot data for one subject, with curve and interval estimates
d = as.numeric(colnames(cd4))
plot(d[which(!is.na(Y.sub[EX,]))], Y.sub[EX,which(!is.na(Y.sub[EX,]))], type = 'o',
  pch = 19, cex = .75, ylim = range(0, 3400), xlim = range(d),
    xlab = "Months since seroconversion", lwd = 1.2, ylab = "Total CD4 Cell Count",
      main = "Est. & CI - Sampled Data")
matpoints(d, EX.IV, col = 2, type = 'l', lwd = c(2, 1, 1, 1, 1), lty = c(1,1,1,2,2))
matpoints(d, EX.MM, col = 4, type = 'l', lwd = c(2, 1, 1, 1, 1), lty = c(1,1,1,2,2))
legend("topright", c("IV Est", "IV PW Int", "IV Simul Int",
    expression(paste("MM - ", hat(theta), " Est", sep = "")),
    expression(paste("MM - ", hat(theta), " PW Int", sep = "")),
    expression(paste("MM - ", hat(theta), " Simul Int", sep = ""))),
```

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```
lty=c(1,1,2,1,1,2), lwd = c(2.5,.75,.75,2.5,.75),
col = c("red","red","red","blue","blue","blue"))
## End(Not run)
```

cd4

Observed CD4 cell counts

## Description

CD4 cell counts for 366 subjects between months -18 and 42 since seroconversion. Each subject's observations are contained in a single row.

#### **Format**

A data frame made up of a 366 x 61 matrix of CD4 cell counts

## References

Goldsmith, J., Greven, S., and Crainiceanu, C. (2013). Corrected confidence bands for functional data using principal components. *Biometrics*, 69(1), 41–51.

cmdscale\_lanczos

Faster multi-dimensional scaling

## **Description**

This is a modified version of cmdscale that uses the Lanczos procedure (slanczos) instead of eigen. Called by smooth.construct.pco.smooth.spec.

## Usage

```
cmdscale_lanczos(d, k = 2, eig = FALSE, add = FALSE, x.ret = FALSE)
```

## **Arguments**

d	a distance structure as returned by dist, or a full symmetric matrix of distances or dissimilarities.
k	the maximum dimension of the space which the data are to be represented in; must be in $\{1, 2,, n-1\}$ .
eig	logical indicating whether eigenvalues should be returned.
add	logical indicating if the additive constant of Cailliez (1983) should be computed, and added to the non-diagonal dissimilarities such that the modified dissimilarities are Euclidean.
x.ret	indicates whether the doubly centred symmetric distance matrix should be returned.

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## Value

```
as cmdscale
```

## Author(s)

David L Miller, based on code by R Core.

## References

Cailliez, F. (1983). The analytical solution of the additive constant problem. *Psychometrika*, 48, 343-349.

#### See Also

```
smooth.construct.pco.smooth.spec
```

coef.pffr

Get estimated coefficients from a pffr fit

## Description

Returns estimated coefficient functions/surfaces  $\beta(t)$ ,  $\beta(s,t)$  and estimated smooth effects f(z), f(x,z) or f(x,z,t) and their point-wise estimated standard errors. Not implemented for smooths in more than 3 dimensions.

## Usage

```
## S3 method for class 'pffr'
coef(
  object,
  raw = FALSE,
  se = TRUE,
  freq = FALSE,
  sandwich = FALSE,
  seWithMean = TRUE,
  n1 = 100,
  n2 = 40,
  n3 = 20,
  Ktt = NULL,
  ...
)
```

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#### **Arguments**

object	a fitted pffr-object
raw	$logical, defaults \ to \ FALSE. \ If \ TRUE, the function simply \ returns \ object \$ coefficients$
se	logical, defaults to TRUE. Return estimated standard error of the estimates?
freq	logical, defaults to FALSE. If FALSE, use posterior variance object\$Vp for variability estimates, else use object\$Ve. See gamObject
sandwich	logical, defaults to FALSE. Use a Sandwich-estimator for approximate variances? See Details. THIS IS AN EXPERIMENTAL FEATURE, USE A YOUR OWN RISK.
seWithMean	logical, defaults to TRUE. Include uncertainty about the intercept/overall mean in standard errors returned for smooth components?
n1	see below
n2	see below
n3	n1, n2, n3 give the number of gridpoints for 1-/2-/3-dimensional smooth terms used in the marginal equidistant grids over the range of the covariates at which the estimated effects are evaluated.
Ktt	(optional) an estimate of the covariance operator of the residual process $\epsilon_i(t) \sim N(0,K(t,t'))$ , evaluated on yind of object. If not supplied, this is estimated from the crossproduct matrices of the observed residual vectors. Only relevant for sandwich CIs.
	other arguments, not used.

## **Details**

The seWithMean-option corresponds to the "iterms"-option in predict.gam. The sandwich-options works as follows: Assuming that the residual vectors  $\epsilon_i(t), i=1,\ldots,n$  are i.i.d. realizations of a mean zero Gaussian process with covariance K(t,t'), we can construct an estimator for K(t,t') from the n replicates of the observed residual vectors. The covariance matrix of the stacked observations  $\text{vec}(Y_i(t))$  is then given by a block-diagonal matrix with n copies of the estimated K(t,t') on the diagonal. This block-diagonal matrix is used to construct the "meat" of a sandwich covariance estimator, similar to Chen et al. (2012), see reference below.

#### Value

If raw==FALSE, a list containing

- pterms a matrix containing the parametric / non-functional coefficients (and, optionally, their se's)
- smterms a named list with one entry for each smooth term in the model. Each entry contains
  - coef a matrix giving the grid values over the covariates, the estimated effect (and, optionally, the se's). The first covariate varies the fastest.
  - x, y, z the unique gridpoints used to evaluate the smooth/coefficient function/coefficient surface
  - xlim, ylim, zlim the extent of the x/y/z-axes
  - xlab, ylab, zlab the names of the covariates for the x/y/z-axes

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- dim the dimensionality of the effect
- main the label of the smooth term (a short label, same as the one used in summary.pffr)

#### Author(s)

Fabian Scheipl

#### References

Chen, H., Wang, Y., Paik, M.C., and Choi, A. (2013). A marginal approach to reduced-rank penalized spline smoothing with application to multilevel functional data. *Journal of the American Statistical Association*, 101, 1216–1229.

## See Also

plot.gam, predict.gam which this routine is based on.

coefboot.pffr

Simple bootstrap CIs for pffr

#### **Description**

This function resamples observations in the data set to obtain approximate CIs for different terms and coefficient functions that correct for the effects of dependency and heteroskedasticity of the residuals along the index of the functional response, i.e., it aims for correct inference if the residuals along the index of the functional response are not i.i.d.

## Usage

```
coefboot.pffr(
  object,
  n1 = 100,
  n2 = 40,
  n3 = 20,
  B = 100,
  ncpus = getOption("boot.ncpus", 1),
  parallel = c("no", "multicore", "snow"),
  c1 = NULL,
  conf = c(0.9, 0.95),
  type = "percent",
  method = c("resample", "residual", "residual.c"),
  showProgress = TRUE,
  ...
)
```

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## **Arguments**

object	a fitted pffr-model
n1	see coef.pffr
n2	see coef.pffr
n3	see coef.pffr
В	number of bootstrap replicates, defaults to (a measly) 100
ncpus	see boot. Defaults to getOption("boot.ncpus", 1L) (like boot).
parallel	see boot
cl	see boot
conf	desired levels of bootstrap CIs, defaults to 0.90 and 0.95
type	type of bootstrap interval, see boot.ci. Defaults to "percent" for percentile-based CIs.
method	either "resample" (default) to resample response trajectories, or "residual" to resample responses as fitted values plus residual trajectories or "residual.c" to resample responses as fitted values plus residual trajectories that are centered at zero for each gridpoint.
showProgress	TRUE/FALSE
	not used

## Value

a list with similar structure as the return value of coef.pffr, containing the original point estimates of the various terms along with their bootstrap CIs.

## Author(s)

Fabian Scheipl

coefficients.pfr Extract coefficient functions from a fitted pfr-object

## Description

This function is used to extract a coefficient from a fitted 'pfr' model, in particular smooth functions resulting from including functional terms specified with 1f, af, etc. It can also be used to extract smooths genereated using mgcv's s, te, or t2.

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## Usage

```
## S3 method for class 'pfr'
coefficients(
  object,
  select = 1,
  coords = NULL,
  n = NULL,
  se = ifelse(length(object$smooth) & select, TRUE, FALSE),
  seWithMean = FALSE,
  useVc = TRUE,
  Qtransform = FALSE,
)
## S3 method for class 'pfr'
coef(
  object,
  select = 1,
  coords = NULL,
  n = NULL,
  se = ifelse(length(object$smooth) & select, TRUE, FALSE),
  seWithMean = FALSE,
  useVc = TRUE,
  Qtransform = FALSE,
)
```

## Arguments

object	return object from pfr
select	integer indicating the index of the desired smooth term in object\$smooth. Enter 0 to request the raw coefficients (i.e., object\$coefficients) and standard errors (if se==TRUE).
coords	named list indicating the desired coordinates where the coefficient function is to be evaluated. Names must match the argument names in object\$smooth[[select]]\$term. If NULL, uses n to generate equally-spaced coordinates.
n	integer vector indicating the number of equally spaced coordinates for each argument. If length 1, the same number is used for each argument. Otherwise, the length must match object\$smooth[[select]]\$dim.
se	if TRUE, returns pointwise standard error estimates. Defaults to FALSE if raw coefficients are being returned; otherwise TRUE.
seWithMean	if TRUE the standard errors include uncertainty about the overall mean; if FALSE, they relate purely to the centered smooth itself. Marra and Wood (2012) suggests that TRUE results in better coverage performance for GAMs.
useVc	if TRUE, standard errors are calculated using a covariance matrix that has been corrected for smoothing parameter uncertainty. This matrix will only be available under ML or REML smoothing.

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Qtransform

For additive functional terms, TRUE indicates the coefficient should be extracted on the quantile-transformed scale, whereas FALSE indicates the scale of the original data. Note this is different from the Qtransform argument of af, which specifies the scale on which the term is fit.

... these arguments are ignored

#### Value

a data frame containing the evaluation points, coefficient function values and optionally the SE's for the term indicated by select.

#### Author(s)

Jonathan Gellar and Fabian Scheipl

#### References

Marra, G and S.N. Wood (2012) Coverage Properties of Confidence Intervals for Generalized Additive Model Components. Scandinavian Journal of Statistics.

content

The CONTENT child growth study

#### **Description**

The CONTENT child growth study was funded by the Sixth Framework Programme of the European Union, Project CONTENT (INCO-DEV-3-032136) and was led by Dr. William Checkley. The study was conducted between May 2007 and February 2011 in Las Pampas de San Juan Miraflores and Nuevo Paraiso, two peri-urban shanty towns with high population density located on the southern edge of Lima city in Peru.

#### Usage

data(content)

#### **Format**

A list made up of

id Numeric vector of subject ID numbers;

**malfe0** Numeric vector of the sex of the child, 1 for male and 0 for female;

weightkg Numeric vector of the weight of the child measured in kilograms(kg);

**height** Numeric vector of the height of the child measured in centimeters;

agedays Numeric vector of the age of the child measured in days;

**cbmi** Numeric vector of the BMI of the child;

**zlen** Numeric vector of the height-for-age z-scores;

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- zwei Numeric vector of the weight-for-age z-scores;
- **zwfl** Numeric vector of the weight-for-height z-scores;
- zbmi Numeric vector of the BMI-for-age z-scores;

#### References

Crainiceanu, C., Goldsmith, J., Leroux, A., Cui, E. (2023). Functional Data Analysis with R. *Chapman & Hall/CRC Statistics* 

COVID19 The US weekly all-cause mortality and COVID19-associated deaths 2020	is in
--	-------

#### **Description**

The COVID19 mortality data used in the "Functional Data Analysis with R" book

#### **Usage**

data(COVID19)

#### **Format**

A list made up of

- **US\_weekly\_mort** A numeric vector of length 207, which contains the total number of weekly all-cause deaths in the US from January 14, 2017 to December 26, 2020;
- **US\_weekly\_mort\_dates** A vector of dates of length 207, which contains the weeks corresponding to the US\_weekly\_mort vector;
- US\_weekly\_mort\_CV19 A numeric vector of length 52, which contains the total number of weekly COVID 19 deaths in the US from January 4, 2020 to December 26, 2020;
- US\_weekly\_mort\_CV19\_dates A vector of dates of length 52, which contains the weeks corresponding to the US\_weekly\_mort\_CV19 vector;
- US\_weekly\_excess\_mort\_2020 A numeric vector of length 52, which contains the US weekly excess mortality (total mortality in one week in 2020 minus total mortality in the corresponding week of 2019) from January 4, 2020 to December 26, 2020;
- US\_weekly\_excess\_mort\_2020\_dates A vector dates of length 52, which contains the weeks corresponding to the US\_weekly\_excess\_mort\_2020 vector.;
- US\_states\_names A vector of strings containing the names of 52 US states and territories in alphabetic order. These are the states for which all-cause and Covid-19 data are available in this data set;
- US\_states\_population A numeric vector containing the population of the 52 states in the vector US\_states\_names estimated as of July 1, 2020. The order of the vector US\_states\_population is the same as that of US\_states\_names;

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**States\_excess\_mortality** A numeric 52 x 52 dimensional matrix that contains the weekly US excess mortality in 52 states and territories. Each row corresponds to one state in the same order as the vector US\_states\_names. Each column corresponds to a week in 2020 corresponding to the order in the vector US\_weekly\_excess\_mort\_2020\_dates. The (i,j)th entry of the matrix is the difference in all-cause mortality during the week j of 2020 and 2019 for state i;

- **States\_excess\_mortality\_per\_million** A numeric 52 x 52 dimensional matrix that contains the weekly US excess mortality in 52 states and territories per one million individuals. This is obtained by dividing every row (corresponding to a state) of States\_excess\_mortality by the population of that state stored in US\_states\_population and multiplying by one million;
- **States\_CV19\_mortality** A numeric 52 x 52 dimensional matrix that contains the weekly US Covid-19 mortality in 52 states and territories. Each row corresponds to one state in the same order as the vector US\_states\_names. Each column corresponds to a week in 2020 corresponding to the order in the vector US\_weekly\_excess\_mort\_2020\_dates;
- **States\_CV19\_mortality\_per\_million** A numeric 52 x 52 dimensional matrix that contains the weekly US Covid-19 mortality in 52 states and territories per one million individuals. This is obtained by dividing every row (corresponding to a state) of States\_CV19\_mortality by the population of that state stored in US\_states\_population and multiplying by one million.

#### References

Crainiceanu, C., Goldsmith, J., Leroux, A., Cui, E. (2023). Functional Data Analysis with R. *Chapman & Hall/CRC Statistics* 

create.prep.func

Construct a function for preprocessing functional predictors

#### **Description**

Prior to using functions X as predictors in a scalar-on-function regression, it is often necessary to presmooth curves to remove measurement error or interpolate to a common grid. This function creates a function to do this preprocessing depending on the method specified.

#### Usage

```
create.prep.func(
   X,
   argvals = seq(0, 1, length = ncol(X)),
   method = c("fpca.sc", "fpca.face", "fpca.ssvd", "bspline", "interpolate"),
   options = NULL
)
```

#### **Arguments**

Χ

an N by J=ncol(argvals) matrix of function evaluations  $X_i(t_{i1}),..,X_i(t_{iJ}); i=1,..,N$ . For FPCA-based processing methods, these functions are used to define the eigen decomposition used to preprocess current and future data (for example, in predict.pfr)

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argvals matrix (or vector) of indices of evaluations of  $X_i(t)$ ; i.e. a matrix with *i*th row

 $(t_{i1},.,t_{iJ})$ 

method character string indicating the preprocessing method. Options are "fpca.sc",

"fpca.face", "fpca.ssvd", "bspline", and "interpolate". The first three use the corresponding existing function; "bspline" uses an (unpenalized) cubic bspline smoother with nbasis basis functions; "interpolate" uses linear

interpolation.

options list of options passed to the preprocessing method; as an example, options for

fpca.sc include pve, nbasis, and npc.

#### Value

a function that returns the preprocessed functional predictors, with arguments

newX The functional predictors to process

argvals. Indices of evaluation of newX

options. Any options needed to preprocess the predictor functions

#### Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

#### See Also

```
pfr, fpca.sc, fpca.face, fpca.ssvd
```

DTI

Diffusion Tensor Imaging: tract profiles and outcomes

#### **Description**

Fractional anisotropy (FA) tract profiles for the corpus callosum (cca) and the right corticospinal tract (rcst). Accompanying the tract profiles are the subject ID numbers, visit number, total number of scans, multiple sclerosis case status and Paced Auditory Serial Addition Test (pasat) score.

#### **Format**

A data frame made up of

cca A 382 x 93 matrix of fractional anisotropy tract profiles from the corpus callosum;

rest A 382 x 55 matrix of fractional anisotropy tract profiles from the right corticospinal tract;

**ID** Numeric vector of subject ID numbers;

visit Numeric vector of the subject-specific visit numbers;

visit.time Numeric vector of the subject-specific visit time, measured in days since first visit;

Nscans Numeric vector indicating the total number of visits for each subject;

case Numeric vector of multiple sclerosis case status: 0 - healthy control, 1 - MS case;

sex factor variable indicated subject's sex;

pasat Numeric vector containing the PASAT score at each visit.

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#### **Details**

If you use this data as an example in written work, please include the following acknowledgment: "The MRI/DTI data were collected at Johns Hopkins University and the Kennedy-Krieger Institute"

DTI2 uses mean diffusivity of the the corpus callosum rather than FA, and parallel diffusivity of the rcst rather than FA. Please see the documentation for DTI2.

#### References

Goldsmith, J., Bobb, J., Crainiceanu, C., Caffo, B., and Reich, D. (2011). Penalized Functional Regression. *Journal of Computational and Graphical Statistics*, 20, 830 - 851.

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2010). Longitudinal Penalized Functional Regression for Cognitive Outcomes on Neuronal Tract Measurements. *Journal of the Royal Statistical Society: Series C*, 61, 453 - 469.

DTI2

Diffusion Tensor Imaging: more fractional anisotropy profiles and outcomes

### **Description**

A diffusion tensor imaging dataset used in Swihart et al. (2012). Mean diffusivity profiles for the corpus callosum (cca) and parallel diffusivity for the right corticospinal tract (rcst). Accompanying the profiles are the subject ID numbers, visit number, and Paced Auditory Serial Addition Test (pasat) score. We thank Dr. Daniel Reich for making this dataset available.

#### **Format**

A data frame made up of

cca a 340 x 93 matrix of fractional anisotropy profiles from the corpus callosum;

rcst a 340 x 55 matrix of fractional anisotropy profiles from the right corticospinal tract;

id numeric vector of subject ID numbers;

visit numeric vector of the subject-specific visit numbers;

**pasat** numeric vector containing the PASAT score at each visit.

#### **Details**

If you use this data as an example in written work, please include the following acknowledgment: "The MRI/DTI data were collected at Johns Hopkins University and the Kennedy-Krieger Institute"

Note: DTI2 uses mean diffusivity of the the corpus callosum rather than fractional anisotropy (FA), and parallel diffusivity of the rest rather than FA. Please see the documentation for DTI for more about the DTI dataset.

expand.call 25

#### References

Goldsmith, J., Bobb, J., Crainiceanu, C., Caffo, B., and Reich, D. (2011). Penalized functional regression. *Journal of Computational and Graphical Statistics*, 20(4), 830–851.

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2012). Longitudinal penalized functional regression for cognitive outcomes on neuronal tract measurements. *Journal of the Royal Statistical Society: Series C*, 61(3), 453–469.

Swihart, B. J., Goldsmith, J., and Crainiceanu, C. M. (2014). Restricted Likelihood Ratio Tests for Functional Effects in the Functional Linear Model. *Technometrics*, 56, 483–493.

expand.call

Return call with all possible arguments

#### **Description**

Return a call in which all of the arguments which were supplied or have presets are specified by their full names and their supplied or default values.

#### Usage

```
expand.call(
  definition = NULL,
  call = sys.call(sys.parent(1)),
  expand.dots = TRUE
)
```

#### **Arguments**

definition a function. See match.call.

call an unevaluated call to the function specified by definition. See match.call.

expand.dots logical. Should arguments matching ... in the call be included or left as a ... argument? See match.call.

#### Value

An object of mode "call".

## Author(s)

Fabian Scheipl

#### See Also

```
match.call
```

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fbps

Sandwich smoother for matrix data

## Description

A fast bivariate *P*-spline method for smoothing matrix data.

## Usage

```
fbps(
  data,
  subj = NULL,
  covariates = NULL,
 knots = 35,
 knots.option = "equally-spaced",
 periodicity = c(FALSE, FALSE),
 p = 3,
 m = 2,
 lambda = NULL,
  selection = "GCV",
  search.grid = T,
  search.length = 100,
 method = "L-BFGS-B",
 lower = -20,
 upper = 20,
  control = NULL
)
```

## Arguments

data	n1 by n2 data matrix without missing data
subj	vector of subject id (corresponding to the columns of data); defaults to NULL
covariates	list of two vectors of covariates of lengths n1 and n2; if NULL, then generates equidistant covariates
knots	list of two vectors of knots or number of equidistant knots for all dimensions; defaults to 35
knots.option	knot selection method; defaults to "equally-spaced"
periodicity	vector of two logical, indicating periodicity in the direction of row and column; defaults to c(FALSE, FALSE)
р	degrees of B-splines; defaults to 3
m	order of differencing penalty; defaults to 2
lambda	user-specified smoothing parameters; defaults to NULL
selection	selection of smoothing parameter; defaults to "GCV"
search.grid	logical; defaults to TRUE, if FALSE, uses optim

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search.length number of equidistant (log scale) smoothing parameter; defaults to 100

method see optim; defaults to L-BFGS-B

lower, upper bounds for log smoothing parameter, passed to optim; defaults are -20 and 20.

control see optim

#### **Details**

The smoothing parameter can be user-specified; otherwise, the function uses grid searching method or optim for selecting the smoothing parameter.

#### Value

A list with components

lambda vector of length 2 of selected smoothing parameters

Yhat fitted data

trace trace of the overall smoothing matrix gcv value of generalized cross validation
Theta matrix of estimated coefficients

#### Author(s)

Luo Xiao <lxiao@jhsph.edu>

#### References

Xiao, L., Li, Y., and Ruppert, D. (2013). Fast bivariate *P*-splines: the sandwich smoother. *Journal of the Royal Statistical Society: Series B*, 75(3), 577–599.

## **Examples**

```
#############################
#### True function #####
n1 <- 60
n2 <- 80
x < -(1:n1)/n1-1/2/n1
z <- (1:n2)/n2-1/2/n2
MY <- array(0,c(length(x),length(z)))
sigx <- .3
sigz <- .4
for(i in 1:length(x))
for(j in 1:length(z))
\#MY[i,j] < .75/(pi*sigx*sigz) *exp(-(x[i]-.2)^2/sigx^2-(z[j]-.3)^2/sigz^2)
\#MY[i,j] <- MY[i,j] + .45/(pi*sigx*sigz) *exp(-(x[i]-.7)^2/sigx^2-(z[j]-.8)^2/sigz^2)
MY[i,j] = sin(2*pi*(x[i]-.5)^3)*cos(4*pi*z[j])
}
```

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```
#############################
#### Observed data #####
#############################
sigma <- 1
Y \leftarrow MY + sigma*rnorm(n1*n2,0,1)
##############################
     Estimation
                  #####
#############################
est <- fbps(Y,list(x=x,z=z))</pre>
mse <- mean((est$Yhat-MY)^2)</pre>
cat("mse of fbps is",mse,"\n")
cat("The smoothing parameters are:",est$lambda,"\n")
######### Compare the estimated surface with the true surface ########
par(mfrow=c(1,2))
persp(x,z,MY,zlab="f(x,z)",zlim=c(-1,2.5), phi=30,theta=45,expand=0.8,r=4,
     col="blue",main="True surface")
persp(x,z,est\$Yhat,zlab="f(x,z)",zlim=c(-1,2.5),phi=30,theta=45,
     expand=0.8,r=4,col="red",main="Estimated surface")
```

Construct a function-on-function regression term

#### **Description**

ff

Defines a term  $\int_{s_{lo,i}}^{s_{hi,i}} X_i(s)\beta(t,s)ds$  for inclusion in an mgcv::gam-formula (or bam or gamm or gamm4:::gamm4) as constructed by pffr.

Defaults to a cubic tensor product B-spline with marginal first order differences penalties for  $\beta(t,s)$  and numerical integration over the entire range  $[s_{lo,i},s_{hi,i}]=[\min(s_i),\max(s_i)]$  by using Simpson weights. Can't deal with any missing X(s), unequal lengths of  $X_i(s)$  not (yet?) possible. Unequal integration ranges for different  $X_i(s)$  should work.  $X_i(s)$  is assumed to be numeric (duh...).

#### Usage

```
ff(
    X,
    yind = NULL,
    xind = seq(0, 1, 1 = ncol(X)),
    basistype = c("te", "t2", "ti", "s", "tes"),
    integration = c("simpson", "trapezoidal", "riemann"),
    L = NULL,
    limits = NULL,
    splinepars = if (basistype != "s") {
        list(bs = "ps", m = list(c(2, 1), c(2, 1)), k
        = c(5, 5))
    } else {
```

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```
list(bs = "tp", m = NA)
},
  check.ident = TRUE
)
```

#### **Arguments**

X	an n by ncol(xind) matrix of function evaluations $X_i(s_{i1}),\dots,X_i(s_{iS}); i=1,\dots,n.$
yind	$\ensuremath{\textit{DEPRECATED}}$ used to supply matrix (or vector) of indices of evaluations of $Y_i(t),$ no longer used.
xind	vector of indices of evaluations of $X_i(s)$ , i.e., $(s_1, \ldots, s_S)$
basistype	defaults to "te", i.e. a tensor product spline to represent $\beta(t,s)$ . Alternatively, use "s" for bivariate basis functions (see mgcv's s) or "t2" for an alternative parameterization of tensor product splines (see mgcv's t2).
integration	method used for numerical integration. Defaults to "simpson"'s rule for calculating entries in L. Alternatively and for non-equidistant grids, "trapezoidal" or "riemann". "riemann" integration is always used if limits is specified
L	optional: an n by $ncol(xind)$ matrix giving the weights for the numerical integration over $s$ .
limits	defaults to NULL for integration across the entire range of $X(s)$ , otherwise specifies the integration limits $s_{hi}(t), s_{lo}(t)$ : either one of "s <t" "s<='t"' <math="" for="" or="">(s_{hi}(t), s_{lo}(t)) = (t, 0] or <math>[t, 0]</math>, respectively, or a function that takes s as the first and t as the second argument and returns TRUE for combinations of values (s,t) if s falls into the integration range for the given t. This is an experimental feature and not well tested yet; use at your own risk.</t">
splinepars	optional arguments supplied to the basistype-term. Defaults to a cubic tensor product B-spline with marginal first difference penalties, i.e. list(bs="ps", $m=list(c(2, 1), c(2, 1))$ ). See te or s in $mgcv$ for details
check.ident	check identifiability of the model spec. See Details and References. Defaults to $TRUE$ .

#### **Details**

If check ident==TRUE and basistype!="s" (the default), the routine checks conditions for non-identifiability of the effect. This occurs if a) the marginal basis for the functional covariate is rank-deficient (typically because the functional covariate has lower rank than the spline basis along its index) and simultaneously b) the kernel of  $\mathrm{Cov}(X(s))$  is not disjunct from the kernel of the marginal penalty over s. In practice, a) occurs quite frequently, and b) occurs usually because curve-wise mean centering has removed all constant components from the functional covariate. If there is kernel overlap,  $\beta(t,s)$  is constrained to be orthogonal to functions in that overlap space (e.g., if the overlap contains constant functions, constraints " $\int \beta(t,s)ds = 0$  for all t" are enforced). See reference for details.

A warning is always given if the effective rank of Cov(X(s)) (defined as the number of eigenvalues accounting for at least 0.995 of the total variance in  $X_i(s)$ ) is lower than 4. If  $X_i(s)$  is of very low rank, ffpc-term may be preferable.

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#### Value

A list containing

call a "call" to te (or s or t2) using the appropriately constructed covariate and

weight matrices

data a list containing the necessary covariate and weight matrices

#### Author(s)

Fabian Scheipl, Sonja Greven

#### References

For background on check.ident:

```
Scheipl, F., Greven, S. (2016). Identifiability in penalized function-on-function regression models. Electronic Journal of Statistics, 10(1), 495–526. https://projecteuclid.org/journals/electronic-journal-of-statistics/volume-10/issue-1/Identifiability-in-penalized-function-on-functi 10.1214/16-EJS1123.full
```

## See Also

```
mgcv's linear.functional.terms
```

ffpc

Construct a PC-based function-on-function regression term

## **Description**

Defines a term  $\int X_i(s)\beta(t,s)ds$  for inclusion in an mgcv::gam-formula (or bam or gamm or gamm4:::gamm4) as constructed by pffr.

## Usage

```
ffpc(
   X,
   yind = NULL,
   xind = seq(0, 1, length = ncol(X)),
   splinepars = list(bs = "ps", m = c(2, 1), k = 8),
   decomppars = list(pve = 0.99, useSymm = TRUE),
   npc.max = 15
)
```

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#### **Arguments**

	by ncol(xind) matrix of function evaluations $X_i(s_{i1}),\ldots,X_i(s_{iS}); i=\ldots,n.$
•	PRECATED used to supply matrix (or vector) of indices of evaluations of ), no longer used.
	ix (or vector) of indices of evaluations of $X_i(t)$ , defaults to seq(0, 1, gth=ncol(X)).
	onal arguments supplied to the basistype-term. Defaults to a cubic Base with first difference penalties and 8 basis functions for each $\tilde{\beta}_k(t)$ .
decomppars para	meters for the FPCA performed with fpca.sc.
npc.max max	imal number $K$ of FPCs to use, regardless of decomppars; defaults to 15

#### **Details**

In contrast to ff, ffpc does an FPCA decomposition  $X(s) \approx \sum_{k=1}^K \xi_{ik} \Phi_k(s)$  using fpca.sc and represents  $\beta(t,s)$  in the function space spanned by these  $\Phi_k(s)$ . That is, since

$$\int X_i(s)\beta(t,s)ds = \sum_{k=1}^K \xi_{ik} \int \Phi_k(s)\beta(s,t)ds = \sum_{k=1}^K \xi_{ik}\tilde{\beta}_k(t),$$

the function-on-function term can be represented as a sum of K univariate functions  $\hat{\beta}_k(t)$  in t each multiplied by the FPC scores  $\xi_{ik}$ . The truncation parameter K is chosen as described in fpca.sc. Using this instead of ff() can be beneficial if the covariance operator of the  $X_i(s)$  has low effective rank (i.e., if K is small). If the covariance operator of the  $X_i(s)$  is of (very) high rank, i.e., if K is large, ffpc() will not be very efficient.

To reduce model complexity, the  $\tilde{\beta}_k(t)$  all have a single joint smoothing parameter (in mgcv, they get the same id, see s).

Please see pffr for details on model specification and implementation.

#### Value

A list containing the necessary information to construct a term to be included in a mgcv::gam-formula.

#### Author(s)

Fabian Scheipl

#### **Examples**

```
## Not run:
set.seed(1122)
n <- 55
S <- 60
T <- 50
s <- seq(0,1, 1=S)</pre>
```

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```
t < - seq(0,1, l=T)
#generate X from a polynomial FPC-basis:
rankX <- 5
Phi <- cbind(1/sqrt(S), poly(s, degree=rankX-1))
lambda <- rankX:1
Xi <- sapply(lambda, function(l)</pre>
            scale(rnorm(n, sd=sqrt(l)), scale=FALSE))
X <- Xi %*% t(Phi)
beta.st <- outer(s, t, function(s, t) cos(2 * pi * s * t))
y \leftarrow (1/S*X) \%*\% beta.st + 0.1 * matrix(rnorm(n * T), nrow=n, ncol=T)
data <- list(y=y, X=X)</pre>
# set number of FPCs to true rank of process for this example:
m.pc <- pffr(y \sim c(1) + 0 + ffpc(X, yind=t, decomppars=list(npc=rankX)),
        data=data, yind=t)
summary(m.pc)
m.ff <- pffr(y \sim c(1) + 0 + ff(X, yind=t), data=data, yind=t)
summary(m.ff)
# fits are very similar:
all.equal(fitted(m.pc), fitted(m.ff))
# plot implied coefficient surfaces:
layout(t(1:3))
persp(t, s, t(beta.st), theta=50, phi=40, main="Truth",
    ticktype="detailed")
plot(m.ff, select=1, zlim=range(beta.st), theta=50, phi=40,
    ticktype="detailed")
title(main="ff()")
ffpcplot(m.pc, type="surf", auto.layout=FALSE, theta = 50, phi = 40)
title(main="ffpc()")
# show default ffpcplot:
ffpcplot(m.pc)
## End(Not run)
```

ffpcplot

Plot PC-based function-on-function regression terms

#### **Description**

Convenience function for graphical summaries of ffpc-terms from a pffr fit.

#### Usage

```
ffpcplot(
```

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```
object,
type = c("fpc+surf", "surf", "fpc"),
pages = 1,
se.mult = 2,
ticktype = "detailed",
theta = 30,
phi = 30,
plot = TRUE,
auto.layout = TRUE
)
```

#### **Arguments**

object a fitted pffr-model

type one of "fpc+surf", "surf" or "fpc": "surf" shows a perspective plot of the coefficient surface implied by the estimated effect functions of the FPC scores, "fpc"

shows three plots: 1) a scree-type plot of the estimated eigenvalues of the functional covariate, 2) the estimated eigenfunctions, and 3) the estimated coefficient

functions associated with the FPC scores. Defaults to showing both.

pages the number of pages over which to spread the output. Defaults to 1. (Irrelevant

if auto.layout=FALSE.)

se.mult display estimated coefficient functions associated with the FPC scores with plus/minus

this number time the estimated standard error. Defaults to 2.

ticktype see persp.
theta see persp.
phi see persp.

plot produce plots or only return plotting data? Defaults to TRUE.

auto.layout should the the function set a suitable layout automatically? Defaults to TRUE

#### Value

primarily produces plots, invisibly returns a list containing the data used for the plots.

#### Author(s)

Fabian Scheipl

#### **Examples**

```
## Not run:
    #see ?ffpc
## End(Not run)
```

34 fgam

fgam

Functional Generalized Additive Models

#### **Description**

Implements functional generalized additive models for functional and scalar covariates and scalar responses. Additionally implements functional linear models. This function is a wrapper for mgcv's gam and its siblings to fit models of the general form

$$g(E(Y_i)) = \beta_0 + \int_{T_1} F(X_{i1}, t)dt + \int_{T_2} \beta(t)X_{i2}dt + f(z_{i1}) + f(z_{i2}, z_{i3}) + \dots$$

with a scalar (but not necessarily continuous) response Y, and link function g

#### Usage

```
fgam(formula, fitter = NA, tensortype = c("te", "t2"), ...)
```

## Arguments

formula	a formula with special terms as for gam, with additional special terms $af()$ , $1f()$ , $re()$ .
fitter	the name of the function used to estimate the model. Defaults to gam if the matrix of functional responses has less than 2e5 data points and to bam if not. "gamm" (see gamm) and "gamm4" (see gamm4) are valid options as well.
tensortype	defaults to te, other valid option is t2
•••	additional arguments that are valid for gam or bam; for example, specify a gamma > 1 to increase amount of smoothing when using GCV to choose smoothing parameters or method="REML" to change to REML for estimation of smoothing parameters (default is GCV).

## Value

a fitted fgam-object, which is a gam-object with some additional information in a fgam-entry. If fitter is "gamm" or "gamm4", only the \$gam part of the returned list is modified in this way.

#### Warning

Binomial responses should be specified as a numeric vector rather than as a matrix or a factor.

#### Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com> and Fabian Scheipl

#### References

McLean, M. W., Hooker, G., Staicu, A.-M., Scheipl, F., and Ruppert, D. (2014). Functional generalized additive models. *Journal of Computational and Graphical Statistics*, **23** (1), pp. 249-269. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3982924/.

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#### See Also

```
af, lf, predict.fgam, vis.fgam
```

## **Examples**

```
data(DTI)
## only consider first visit and cases (no PASAT scores for controls)
y <- DTI$pasat[DTI$visit==1 & DTI$case==1]</pre>
X <- DTI$cca[DTI$visit==1 & DTI$case==1, ]</pre>
X_2 <- DTI$rcst[DTI$visit==1 & DTI$case==1, ]</pre>
## remove samples containing missing data
ind <- rowSums(is.na(X)) > 0
ind2 < - rowSums(is.na(X_2)) > 0
y \leftarrow y[!(ind \mid ind2)]
X \leftarrow X[!(ind \mid ind2),]
X_2 \leftarrow X_2[!(ind | ind2), ]
N <- length(y)
## fit fgam using FA measurements along corpus callosum
## as functional predictor with PASAT as response
## using 8 cubic B-splines for marginal bases with third
## order marginal difference penalties
## specifying gamma > 1 enforces more smoothing when using
## GCV to choose smoothing parameters
#fit <- fgam(y ~ af(X, k = c(8, 8), m = list(c(2, 3), c(2, 3))), gamma = 1.2)
## fgam term for the cca measurements plus an flm term for the rcst measurements
## leave out 10 samples for prediction
test <- sample(N, 10)
#fit <- fgam(y ~ af(X, k = c(7, 7), m = list(c(2, 2), c(2, 2))) +
        lf(X_2, k=7, m = c(2, 2)), subset=(1:N)[-test])
#plot(fit)
## predict the ten left outs samples
#pred <- predict(fit, newdata = list(X=X[test, ], X_2 = X_2[test, ]), type='response',</pre>
 #
                  PredOutOfRange = TRUE)
#sqrt(mean((y[test] - pred)^2))
## Try to predict the binary response disease status (case or control)
## using the quantile transformed measurements from the rcst tract
## with a smooth component for a scalar covariate that is pure noise
y <- DTI$case[DTI$visit==1]</pre>
X <- DTI$cca[DTI$visit==1, ]</pre>
X_2 <- DTI$rcst[DTI$visit==1, ]</pre>
ind <- rowSums(is.na(X)) > 0
ind2 <- rowSums(is.na(X_2)) > 0
y \leftarrow y[!(ind \mid ind2)]
X \leftarrow X[!(ind \mid ind2),]
```

36 fosr

fosr

Function-on-scalar regression

#### **Description**

Fit linear regression with functional responses and scalar predictors, with efficient selection of optimal smoothing parameters.

## Usage

```
fosr(
  formula = NULL,
  Y = NULL
  fdobj = NULL,
  data = NULL,
  Χ,
  con = NULL,
  argvals = NULL,
 method = c("OLS", "GLS", "mix"),
  \label{eq:gam.method} \texttt{gam.method} \; \texttt{=} \; \texttt{c("REML", "ML", "GCV.Cp", "GACV.Cp", "P-REML", "P-ML")},
  cov.method = c("naive", "mod.chol"),
  lambda = NULL,
  nbasis = 15,
  norder = 4,
  pen.order = 2,
 multi.sp = ifelse(method == "OLS", FALSE, TRUE),
  pve = 0.99,
 max.iter = 1,
 maxlam = NULL
  cv1 = FALSE,
  scale = FALSE
)
```

## **Arguments**

formula Formula for fitting fosr. If used, data argument must not be null.

Y, fdobj the functional responses, given as either an  $n \times d$  matrix Y or a functional data object (class "fd") as in the **fda** package.

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data	data frame containing the predictors and responses.
X	the model matrix, whose columns represent scalar predictors. Should ordinarily include a column of 1s.
con	a row vector or matrix of linear contrasts of the coefficient functions, to be constrained to equal zero.
argvals	the $d$ argument values at which the coefficient functions will be evaluated.
method	estimation method: "OLS" for penalized ordinary least squares, "GLS" for penalized generalized least squares, "mix" for mixed effect models.
gam.method	smoothing parameter selection method, to be passed to gam: "REML" for restricted maximum likelihood, "GCV.Cp" for generalized cross-validation.
cov.method	covariance estimation method: the current options are naive or modified Cholesky. See Details.
lambda	smoothing parameter value. If NULL, the smoothing parameter(s) will be estimated. See Details.
nbasis, norder	number of basis functions, and order of splines (the default, 4, gives cubic splines), for the B-spline basis used to represent the coefficient functions. When the functional responses are supplied using fdobj, these arguments are ignored in favor of the values pertaining to the supplied object.
pen.order	order of derivative penalty.
multi.sp	a logical value indicating whether separate smoothing parameters should be estimated for each coefficient function. Currently must be FALSE if method = "OLS".
pve	if method = 'mix', the percentage of variance explained by the principal components; defaults to 0.99.
max.iter	maximum number of iterations if method = "GLS".
maxlam	maximum smoothing parameter value to consider (when lamvec=NULL; see lofocv).
cv1	logical value indicating whether a cross-validation score should be computed even if a single fixed lambda is specified (when method = "OLS").
scale	logical value or vector determining scaling of the matrix X (see scale, to which the value of this argument is passed).

## **Details**

The GLS method requires estimating the residual covariance matrix, which has dimension  $d \times d$  when the responses are given by Y, or  $nbasis \times nbasis$  when they are given by fdobj. When cov.method = "naive", the ordinary sample covariance is used. But this will be singular, or non-singular but unstable, in high-dimensional settings, which are typical. cov.method = "mod.chol" implements the modified Cholesky method of Pourahmadi (1999) for estimation of covariance matrices whose inverse is banded. The number of bands is chosen to maximize the p-value for a sphericity test (Ledoit and Wolf, 2002) applied to the "prewhitened" residuals. Note, however, that the banded inverse covariance assumption is sometimes inappropriate, e.g., for periodic functional responses.

There are three types of values for argument lambda:

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1. if NULL, the smoothing parameter is estimated by gam (package **mgcv**) if method = "GLS", or by optimize if method = "OLS";

- 2. if a scalar, this value is used as the smoothing parameter (but only for the initial model, if method = "GLS");
- 3. if a vector, this is used as a grid of values for optimizing the cross-validation score (provided method = "OLS"; otherwise an error message is issued).

Please note that currently, if multi.sp = TRUE, then lambda must be NULL and method must be "GLS".

## Value

An object of class fosr, which is a list with the following elements:

fd	object of class "fd" representing the estimated coefficient functions. Its main components are a basis and a matrix of coefficients with respect to that basis.
pca.resid	if method = "mix", an object representing a functional PCA of the residuals, performed by fpca.sc if the responses are in raw form or by pca.fd if in functional-data-object form.
U	if method = "mix", an $n \times m$ matrix of random effects, where $m$ is the number of functional PC's needed to explain proportion pve of the residual variance. These random effects can be interpreted as shrunken FPC scores.
yhat, resid	objects of the same form as the functional responses (see arguments Y and fdobj), giving the fitted values and residuals.
est.func	matrix of values of the coefficient function estimates at the points given by argvals.
se.func	matrix of values of the standard error estimates for the coefficient functions, at the points given by argvals.
argvals	points at which the coefficient functions are evaluated.
fit	fit object outputted by amc.
edf	effective degrees of freedom of the fit.
lambda	smoothing parameter, or vector of smoothing parameters.
cv	cross-validated integrated squared error if method="OLS", otherwise NULL.
roughness	value of the roughness penalty.
resp.type	"raw" or "fd", indicating whether the responses were supplied in raw or functional-data-object form.

## Author(s)

Philip Reiss <phil.reiss@nyumc.org>, Lan Huo, and Fabian Scheipl

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#### References

Ledoit, O., and Wolf, M. (2002). Some hypothesis tests for the covariance matrix when the dimension is large compared to the sample size. *Annals of Statistics*, 30(4), 1081–1102.

Pourahmadi, M. (1999). Joint mean-covariance models with applications to longitudinal data: unconstrained parameterisation. *Biometrika*, 86(3), 677–690.

Ramsay, J. O., and Silverman, B. W. (2005). *Functional Data Analysis*, 2nd ed., Chapter 13. New York: Springer.

Reiss, P. T., Huang, L., and Mennes, M. (2010). Fast function-on-scalar regression with penalized basis expansions. *International Journal of Biostatistics*, 6(1), article 28. Available at https://pubmed.ncbi.nlm.nih.gov/21969982/

#### See Also

```
plot.fosr
```

```
## Not run:
require(fda)
# The first two lines, adapted from help(fRegress) in package fda,
# set up a functional data object representing daily average
# temperatures at 35 sites in Canada
daybasis25 <- create.fourier.basis(rangeval=c(0, 365), nbasis=25,</pre>
                  axes=list('axesIntervals'))
Temp.fd <- with(CanadianWeather, smooth.basisPar(day.5,</pre>
                dailyAv[,,'Temperature.C'], daybasis25)$fd)
modmat = cbind(1, model.matrix(~ factor(CanadianWeather$region) - 1))
constraints = matrix(c(0,1,1,1,1), 1)
# Penalized OLS with smoothing parameter chosen by grid search
olsmod = fosr(fdobj = Temp.fd, X = modmat, con = constraints, method="OLS", lambda=100*10:30)
plot(olsmod, 1)
# Test use formula to fit fosr
set.seed(2121)
data1 <- pffrSim(scenario="ff", n=40)</pre>
formod = fosr(Y~xlin+xsmoo, data=data1)
plot(formod, 1)
# Penalized GLS
glsmod = fosr(fdobj = Temp.fd, X = modmat, con = constraints, method="GLS")
plot(glsmod, 1)
## End(Not run)
```

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fosr.perm

Permutation testing for function-on-scalar regression

## **Description**

fosr.perm() is a wrapper function calling fosr.perm.fit(), which fits models to permuted data, followed by fosr.perm.test(), which performs the actual simultaneous hypothesis test. Calling the latter two functions separately may be useful for performing tests at different significance levels. By default, fosr.perm() produces a plot using the plot function for class fosr.perm.

# Usage

```
fosr.perm(
 Y = NULL
  fdobj = NULL,
 Χ,
  con = NULL,
 X0 = NULL,
  con0 = NULL,
  argvals = NULL,
  lambda = NULL,
  lambda0 = NULL,
 multi.sp = FALSE,
 nperm,
  level = 0.05,
 plot = TRUE,
  xlabel = "",
  title = NULL,
 prelim = if (multi.sp) 0 else 15,
)
fosr.perm.fit(
 Y = NULL,
  fdobj = NULL,
 Χ,
  con = NULL,
 X0 = NULL,
  con0 = NULL,
  argvals = NULL,
  lambda = NULL,
  lambda0 = NULL,
 multi.sp = FALSE,
 nperm,
 prelim,
)
```

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```
fosr.perm.test(x, level = 0.05)
## S3 method for class 'fosr.perm'
plot(x, level = 0.05, xlabel = "", title = NULL, ...)
```

# Arguments

Y, fdobj	the functional responses, given as either an $n \times d$ matrix Y or a functional data object (class "fd") as in the <b>fda</b> package.
Χ	the design matrix, whose columns represent scalar predictors.
con	a row vector or matrix of linear contrasts of the coefficient functions, to be restricted to equal zero.
X0	design matrix for the null-hypothesis model. If NULL, the null hypothesis is the intercept-only model.
con0	linear constraints for the null-hypothesis model.
argvals	the $d$ argument values at which the coefficient functions will be evaluated.
lambda	smoothing parameter value. If NULL, the smoothing parameter(s) will be estimated. See four for details.
lambda0	smoothing parameter for null-hypothesis model.
multi.sp	a logical value indicating whether separate smoothing parameters should be estimated for each coefficient function. Currently must be FALSE if method = "OLS".
nperm	number of permutations.
level	significance level for the simultaneous test.
plot	logical value indicating whether to plot the real- and permuted-data pointwise F-type statistics.
xlabel	x-axis label for plots.
title	title for plot.
prelim	number of preliminary permutations. The smoothing parameter in the main permutations will be fixed to the median value from these preliminary permutations. If prelim=0, this is not done. Preliminary permutations are not available when multi.sp = TRUE (hence the complicated default).
	for fosr.perm and fosr.perm.fit, additional arguments passed to fosr. These arguments may include max.iter, method, gam.method, and scale. For plot.fosr.perm, graphical parameters (see par) for the plot.
X	object of class fosr.perm, outputted by fosr.perm, fosr.perm.fit, or fosr.perm.test.

## Value

fosr.perm or fosr.perm.test produces an object of class fosr.perm, which is a list with the elements below. fosr.perm.fit also outputs an object of this class, but without the last five elements.

F pointwise F-type statistics at each of the points given by argvals.

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F. perm a matrix, each of whose rows gives the pointwise F-type statistics for a permuted

data set.

argvals points at which F-type statistics are computed.

lambda.real smoothing parameter(s) for the real-data fit.

lambda.prelim smoothing parameter(s) for preliminary permuted-data fits.

lambda.perm smoothing parameter(s) for main permuted-data fits.

lambda0.real, lambda0.prelim, lambda0.perm

as above, but for null hypothesis models.

level significance level of the test. critval critical value for the test.

signif vector of logical values indicating whether significance is attained at each of the

points argvals.

n2s subset of 1, ..., length(argvals) identifying the points at which the test statistic

changes from non-significant to significant.

s2n points at which the test statistic changes from significant to non-significant.

### Author(s)

Philip Reiss <phil.reiss@nyumc.org> and Lan Huo

### References

Reiss, P. T., Huang, L., and Mennes, M. (2010). Fast function-on-scalar regression with penalized basis expansions. *International Journal of Biostatistics*, 6(1), article 28. Available at https://pubmed.ncbi.nlm.nih.gov/21969982/

#### See Also

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```
box()
axis(2)
axisIntervals(1)
## End(Not run)
```

fosr.vs

Function-on Scalar Regression with variable selection

## **Description**

Implements an iterative algorithm for function-on-scalar regression with variable selection by alternatively updating the coefficients and covariance structure.

## Usage

```
fosr.vs(
  formula,
  data,
  nbasis = 10,
  method = c("ls", "grLasso", "grMCP", "grSCAD"),
  epsilon = 1e-05,
  max.iter_num = 100
)
```

# Arguments

formula an object of class "formula": an expression of the model to be fitted.

data a data frame that contains the variables in the model.

nbasis number of B-spline basis functions used.

method group variable selection method to be used ("grLasso", "grMCP", "grSCAD"

refer to group Lasso, group MCP and group SCAD, respectively) or "1s" for

least squares estimation.

epsilon the convergence criterion.

max.iter\_num maximum number of iterations.

## Value

A fitted fosr.vs-object, which is a list with the following elements:

formula an object of class "formula": an expression of the model to be fitted.

coefficients the estimated coefficient functions.

fitted.values the fitted curves. residuals the residual curves.

vcov the estimated variance-covariance matrix when convergence is achieved.

method group variable selection method to be used or "1s" for least squares estimation.

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## Author(s)

Yakuan Chen <yc2641@cumc.columbia.edu>

#### References

Chen, Y., Goldsmith, J., and Ogden, T. (2016). Variable selection in function-on-scalar regression. *Stat* 5 88-101

## See Also

```
grpreg
```

```
## Not run:
set.seed(100)
I = 100
p = 20
D = 50
grid = seq(0, 1, length = D)
beta.true = matrix(0, p, D)
beta.true[1,] = sin(2*grid*pi)
beta.true[2,] = cos(2*grid*pi)
beta.true[3,] = 2
psi.true = matrix(NA, 2, D)
psi.true[1,] = sin(4*grid*pi)
psi.true[2,] = cos(4*grid*pi)
lambda = c(3,1)
set.seed(100)
X = matrix(rnorm(I*p), I, p)
C = cbind(rnorm(I, mean = 0, sd = lambda[1]), rnorm(I, mean = 0, sd = lambda[2]))
fixef = X%*%beta.true
pcaef = C %*% psi.true
error = matrix(rnorm(I*D), I, D)
Yi.true = fixef
Yi.pca = fixef + pcaef
Yi.obs = fixef + pcaef + error
data = as.data.frame(X)
data$Y = Yi.obs
fit.fosr.vs = fosr.vs(Y~., data = data, method="grMCP")
plot(fit.fosr.vs)
## End(Not run)
```

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fosr2s

Two-step function-on-scalar regression

# Description

This function performs linear regression with functional responses and scalar predictors by (1) fitting a separate linear model at each point along the function, and then (2) smoothing the resulting coefficients to obtain coefficient functions.

## Usage

```
fosr2s(
   Y,
   X,
   argvals = seq(0, 1, , ncol(Y)),
   nbasis = 15,
   norder = 4,
   pen.order = norder - 2,
   basistype = "bspline"
)
```

## **Arguments**

Υ	the functional responses, given as an $n \times d$ matrix.
X	$n\times p$ model matrix, whose columns represent scalar predictors. Should ordinarily include a column of 1s.
argvals	the $d$ argument values at which the functional responses are evaluated, and at which the coefficient functions will be evaluated.
nbasis	number of basis functions used to represent the coefficient functions.
norder	norder of the spline basis, when basistype="bspline" (the default, 4, gives cubic splines).
pen.order	order of derivative penalty.
basistype	type of basis used. The basis is created by an appropriate constructor function from the <b>fda</b> package; see basisfd. Only "bspline" and "fourier" are supported.

## **Details**

Unlike fosr and pffr, which obtain smooth coefficient functions by minimizing a penalized criterion, this function introduces smoothing only as a second step. The idea was proposed by Fan and Zhang (2000), who employed local polynomials rather than roughness penalization for the smoothing step.

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## Value

An object of class fosr, which is a list with the following elements:

fd	object of class "fd" representing the estimated coefficient functions. Its main components are a basis and a matrix of coefficients with respect to that basis.
raw.coef	$d\times p$ matrix of coefficient estimates from regressing on X separately at each point along the function.
raw.se	$d \times p$ matrix of standard errors of the raw coefficient estimates.
yhat	$n \times d$ matrix of fitted values.
est.func	$d\times p$ matrix of coefficient function estimates, obtained by smoothing the columns of raw.coef.
se.func	$d \times p$ matrix of coefficient function standard errors.
argvals	points at which the coefficient functions are evaluated.
lambda	smoothing parameters (chosen by REML) used to smooth the $p$ coefficient functions with respect to the supplied basis.

## Author(s)

Philip Reiss <phil.reiss@nyumc.org> and Lan Huo

### References

Fan, J., and Zhang, J.-T. (2000). Two-step estimation of functional linear models with applications to longitudinal data. *Journal of the Royal Statistical Society, Series B*, 62(2), 303–322.

## See Also

fosr, pffr

fpc	Construct a FPC regression term	

# Description

Constructs a functional principal component regression (Reiss and Ogden, 2007, 2010) term for inclusion in an mgcv::gam-formula (or bam or gamm or gamm4:::gamm) as constructed by pfr. Currently only one-dimensional functions are allowed.

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### Usage

```
fpc(
   X,
   argvals = NULL,
   method = c("svd", "fpca.sc", "fpca.face", "fpca.ssvd"),
   ncomp = NULL,
   pve = 0.99,
   penalize = (method == "svd"),
   bs = "ps",
   k = 40,
   ...
)
```

## **Arguments**

Χ	functional predictors, typically expressed as an N by J matrix, where N is the
	number of columns and J is the number of evaluation points. May include miss-
	ing/sparse functions, which are indicated by NA values. Alternatively, can be an
	object of class "fd"; see fd.

argvals indices of evaluation of X, i.e.  $(t_{i1},..,t_{iJ})$  for subject i. May be entered as either a length-J vector, or as an N by J matrix. Indices may be unequally spaced. Entering as a matrix allows for different observations times for each subject. If NULL, default to an equally-spaced grid between 0 or 1 (or within X\$basis\$rangeval

if X is a fd object.)

method the method used for finding principal components. The default is an uncon-

strained SVD of the XB matrix. Alternatives include constrained (functional)

principal components approaches

ncomp number of principal components. if NULL, chosen by pve

pve proportion of variance explained; used to choose the number of principal com-

ponents

penalize if TRUE, a roughness penalty is applied to the functional estimate. Defaults to

TRUE if method=="svd" (corresponding to the FPCR\_R method of Reiss and

Ogden (2007)), and FALSE if method!="svd" (corresponding to FPCR\_C).

bs two letter character string indicating the mgcv-style basis to use for pre-smoothing

Χ

k the dimension of the pre-smoothing basis

additional options to be passed to lf. These include arguals, integration,

and any additional options for the pre-smoothing basis (as constructed by mgcv::s),

such as m.

#### **Details**

fpc is a wrapper for 1f, which defines linear functional predictors for any type of basis for inclusion in a pfr formula. fpc simply calls 1f with the appropriate options for the fpc basis and penalty construction.

This function implements both the FPCR-R and FPCR-C methods of Reiss and Ogden (2007). Both methods consist of the following steps:

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- 1. project X onto a spline basis B
- 2. perform a principal components decomposition of XB
- 3. use those PC's as the basis in fitting a (generalized) functional linear model

This implementation provides options for each of these steps. The basis for in step 1 can be specified using the arguments bs and k, as well as other options via . . .; see s for these options. The type of PC-decomposition is specified with method. And the FLM can be fit either penalized or unpenalized via penalize.

The default is FPCR-R, which uses a b-spline basis, an unconstrained principal components decomposition using svd, and the FLM fit with a second-order difference penalty. FPCR-C can be selected by using a different option for method, indicating a constrained ("functional") PC decomposition, and by default an unpenalized fit of the FLM.

FPCR-R is also implemented in fpcr; here we implement the method for inclusion in a pfr formula.

#### Value

The result of a call to 1f.

#### NOTE

Unlike fpcr, fpc within a pfr formula does not automatically decorrelate the functional predictors from additional scalar covariates.

## Author(s)

Jonathan Gellar <JGellar@mathematica-mpr.com>, Phil Reiss <phil.reiss@nyumc.org>, Lan Huo <lan.huo@nyumc.org>, and Lei Huang <huangracer@gmail.com>

#### References

Reiss, P. T. (2006). Regression with signals and images as predictors. Ph.D. dissertation, Department of Biostatistics, Columbia University. Available at http://works.bepress.com/phil\_reiss/11/.

Reiss, P. T., and Ogden, R. T. (2007). Functional principal component regression and functional partial least squares. *Journal of the American Statistical Association*, 102, 984-996.

Reiss, P. T., and Ogden, R. T. (2010). Functional generalized linear models with images as predictors. *Biometrics*, 66, 61-69.

## See Also

```
1f, smooth.construct.fpc.smooth.spec
```

```
data(gasoline)
par(mfrow=c(3,1))

# Fit PFCR_R
gasmod1 <- pfr(octane ~ fpc(NIR, ncomp=30), data=gasoline)
plot(gasmod1, rug=FALSE)</pre>
```

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```
est1 <- coef(gasmod1)
# Fit FPCR_C with fpca.sc
gasmod2 <- pfr(octane ~ fpc(NIR, method="fpca.sc", ncomp=6), data=gasoline)
plot(gasmod2, se=FALSE)
est2 <- coef(gasmod2)

# Fit penalized model with fpca.face
gasmod3 <- pfr(octane ~ fpc(NIR, method="fpca.face", penalize=TRUE), data=gasoline)
plot(gasmod3, rug=FALSE)
est3 <- coef(gasmod3)

par(mfrow=c(1,1))
ylm <- range(est1$value)*1.35
plot(value ~ X.argvals, type="l", data=est1, ylim=ylm)
lines(value ~ X.argvals, col=2, data=est2)
lines(value ~ X.argvals, col=3, data=est3)</pre>
```

fpca.face

Functional principal component analysis with fast covariance estimation

# **Description**

A fast implementation of the sandwich smoother (Xiao et al., 2013) for covariance matrix smoothing. Pooled generalized cross validation at the data level is used for selecting the smoothing parameter.

## Usage

```
fpca.face(
  Y = NULL
 ydata = NULL,
 Y.pred = NULL,
  argvals = NULL,
 pve = 0.99,
  npc = NULL,
  var = FALSE,
  simul = FALSE,
  sim.alpha = 0.95,
  center = TRUE,
  knots = 35,
  p = 3,
 m = 2,
  lambda = NULL,
  alpha = 1,
  search.grid = TRUE,
  search.length = 100,
```

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```
method = "L-BFGS-B",
lower = -20,
upper = 20,
control = NULL,
periodicity = FALSE
)
```

## **Arguments**

Y, ydata the user must supply either Y, a matrix of functions observed on a regular grid,

or a data frame ydata representing irregularly observed functions. See Details.

Y. pred if desired, a matrix of functions to be approximated using the FPC decomposi-

tion.

argvals numeric; function argument.

pve proportion of variance explained: used to choose the number of principal com-

ponents.

npc how many smooth SVs to try to extract, if NA (the default) the hard thresholding

rule of Gavish and Donoho (2014) is used (see Details, References).

var logical; should an estimate of standard error be returned?

simul logical; if TRUE curves will we simulated using Monte Carlo to obtain an esti-

mate of the sim. alpha quantile at each argval; ignored if var == FALSE

sim.alpha numeric; if simul==TRUE, quantile to estimate at each argval; ignored if var

== FALSE

center logical; center Y so that its column-means are 0? Defaults to TRUE

knots number of knots to use or the vectors of knots; defaults to 35

p integer; the degree of B-splines functions to usem integer; the order of difference penalty to use

lambda smoothing parameter; if not specified smoothing parameter is chosen using

optim or a grid search

alpha numeric; tuning parameter for GCV; see parameter gamma in gam

search.grid logical; should a grid search be used to find lambda? Otherwise, optim is used

search.length integer; length of grid to use for grid search for lambda; ignored if search.grid

is FALSE

method method to use; see optim

lower see optim
upper see optim
control see optim

periodicity Option for a periodic spline basis. Defaults to FALSE.

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#### Value

A list with components

- 1. Yhat If Y. pred is specified, the smooth version of Y. pred. Otherwise, if Y. pred=NULL, the smooth version of Y.
- 2. scores matrix of scores
- 3. mu mean function
- 4. npc number of principal components
- 5. efunctions matrix of eigenvectors
- 6. evalues vector of eigenvalues
- 7. pve The percent variance explained by the returned number of PCs

if var == TRUE additional components are returned

- 1. sigma2 estimate of the error variance
- 2. VarMats list of covariance function estimate for each subject
- 3. diag.var matrix containing the diagonals of each matrix in
- 4. crit.val list of estimated quantiles; only returned if simul == TRUE

## Author(s)

Luo Xiao

#### References

Xiao, L., Li, Y., and Ruppert, D. (2013). Fast bivariate *P*-splines: the sandwich smoother, *Journal of the Royal Statistical Society: Series B*, 75(3), 577-599.

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C. (2016). Fast covariance estimation for high-dimensional functional data. *Statistics and Computing*, 26, 409-421. DOI: 10.1007/s11222-014-9485-x.

## See Also

fpca.sc for another covariance-estimate based smoothing of Y; fpca2s and fpca.ssvd for two SVD-based smoothings.

```
#### settings
I <- 50 # number of subjects
J <- 3000 # dimension of the data
t <- (1:J)/J # a regular grid on [0,1]
N <- 4 #number of eigenfunctions
sigma <- 2 ##standard deviation of random noises
lambdaTrue <- c(1,0.5,0.5^2,0.5^3) # True eigenvalues

case = 1
### True Eigenfunctions</pre>
```

```
if(case==1) phi <- sqrt(2)*cbind(sin(2*pi*t),cos(2*pi*t),</pre>
                           sin(4*pi*t),cos(4*pi*t))
if(case==2) phi <- cbind(rep(1,J), sqrt(3)*(2*t-1),
                      sqrt(5)*(6*t^2-6*t+1),
                     sqrt(7)*(20*t^3-30*t^2+12*t-1))
#############
           Generate Data
xi <- matrix(rnorm(I*N),I,N);</pre>
xi <- xi %*% diag(sqrt(lambdaTrue))</pre>
X \leftarrow xi \%\% t(phi); \# of size I by J
Y <- X + sigma*matrix(rnorm(I*J),I,J)
results <- fpca.face(Y,center = TRUE, argvals=t,knots=100,pve=0.99)
# calculate percent variance explained by each PC
evalues = results$evalues
pve_vec = evalues * results$npc/sum(evalues)
########
                FACE
Phi <- results$efunctions
eigenvalues <- results$evalues
for(k in 1:N){
 if(Phi[,k] %*% phi[,k]< 0)
   Phi[,k] \leftarrow - Phi[,k]
}
### plot eigenfunctions
par(mfrow=c(N/2,2))
seq <- (1:(J/10))*10
for(k in 1:N){
 plot(t[seq], Phi[seq, k]*sqrt(J), type="1", lwd = 3,
     ylim = c(-2,2),col = "red",
     ylab = paste("Eigenfunction ",k,sep=""),
     xlab="t",main="FACE")
 lines(t[seq],phi[seq,k],lwd = 2, col = "black")
```

fpca.lfda

Longitudinal Functional Data Analysis using FPCA

### **Description**

Implements longitudinal functional data analysis (Park and Staicu, 2015). It decomposes longitudinally-observed functional observations in two steps. It first applies FPCA on a properly defined marginal

covariance function and obtain estimated scores (mFPCA step). Then it further models the underlying process dynamics by applying another FPCA on a covariance of the estimated scores obtained in the mFPCA step. The function also allows to use a random effects model to study the underlying process dynamics instead of a KL expansion model in the second step. Scores in mFPCA step are estimated using numerical integration. Scores in sFPCA step are estimated under a mixed model framework.

## Usage

```
fpca.lfda(
 Υ,
  subject.index,
  visit.index,
  obsT = NULL,
  funcArg = NULL,
  numTEvalPoints = 41,
 newdata = NULL,
  fbps.knots = c(5, 10),
  fbps.p = 3,
  fbps.m = 2,
 mFPCA.pve = 0.95,
 mFPCA.knots = 35,
 mFPCA.p = 3,
 mFPCA.m = 2,
 mFPCA.npc = NULL,
 LongiModel.method = c("fpca.sc", "lme"),
  sFPCA.pve = 0.95,
  sFPCA.nbasis = 10,
  sFPCA.npc = NULL,
  gam.method = "REML",
 gam.kT = 10
)
```

#### **Arguments**

Y	a matrix of which each row corresponds to one curve observed on a regular and dense grid (dimension of N by m; $N = total$ number of observed functions; $m = total$ number of grid points)
subject.index	subject id; vector of length N with each element corresponding a row of Y
visit.index	index for visits (repeated measures); vector of length N with each element corresponding a row of Y
obsT	actual time of visits at which a function is observed; vector of length $N$ with each element corresponding a row of $\Upsilon$
funcArg	numeric; function argument
numTEvalPoints	total number of evaluation time points for visits; used for pre-binning in sFPCA step; defaults to 41
newdata	an optional data frame providing predictors (i for subject id / Ltime for visit time) with which prediction is desired; defaults to NULL

fbps.knots list of two vectors of knots or number of equidistanct knots for all dimensions for a fast bivariate P-spline smoothing (fbps) method used to estimate a bivariate, smooth mean function; defaults to c(5,10); see fbps fbps.p integer; degrees of B-spline functions to use for a fbps method; defaults to 3; see fbps fbps.m integer; order of differencing penalty to use for a fbps method; defaults to 2; see proportion of variance explained for a mFPCA step; used to choose the number mFPCA.pve of principal components (PCs); defaults to 0.95; see fpca. face mFPCA.knots number of knots to use or the vectors of knots in a mFPCA step; used for obtain a smooth estimate of a covariance function; defaults to 35; see fpca. face integer; the degree of B-spline functions to use in a mFPCA step; defaults to 3; mFPCA.p see fpca.face mFPCA.m integer; order of differencing penalty to use in a mFPCA step; defaults to 2; see fpca.face pre-specified value for the number of principal components; if given, it overrides mFPCA.npc pve; defaults to NULL; see fpca. face LongiModel.method model and estimation method for estimating covariance of estimated scores from a mFPCA step; either KL expansion model or random effects model; defaults to fpca.sc proportion of variance explained for sFPCA step; used to choose the number of sFPCA.pve principal components; defaults to 0.95; see fpca.sc number of B-spline basis functions used in sFPCA step for estimation of the sFPCA.nbasis mean function and bivariate smoothing of the covariance surface; defaults to 10; see fpca.sc pre-specified value for the number of principal components; if given, it overrides sFPCA.npc pve; defaults to NULL; see fpca.sc gam.method smoothing parameter estimation method when gam is used for predicting score functions at unobserved visit time, T; defaults to REML; see gam gam.kT dimension of basis functions to use; see gam

#### Value

# A list with components

obsData observed data (input)

i subject id

funcArg function argument

visitTime visit times

 $\label{eq:fitted_values} \textbf{fitted values (in-sample); of the same dimension as } Y$ 

fitted.values.all

a list of which each component consists of a subject's fitted values at all pairs of evaluation points (s and T)

```
predicted.values
                  predicted values for variables provided in newdata
bivariateSmoothMeanFunc
                  estimated bivariate smooth mean function
mFPCA.efunctions
                  estimated eigenfunction in a mFPCA step
                 estimated eigenvalues in a mFPCA step
mFPCA.evalues
mFPCA.npc
                  number of principal components selected with pre-specified pve in a mFPCA
mFPCA.scree.eval
                  estimated eigenvalues obtained with pre-specified pve = 0.9999; for scree plot
sFPCA.xiHat.bySubj
                  a list of which each component consists of a subject's predicted score functions
                  evaluated at equidistanced grid in direction of visit time, T
sFPCA.npc
                  a vector of numbers of principal components selected in a sFPCA step with
                  pre-specified pve; length of mFPCA.npc
mFPCA.covar
                  estimated marginal covariance
sFPCA.longDynCov.k
                  a list of estimated covariance of score function; length of mFPCA.npc
```

#### **Details**

A random effects model is recommended when a set of visit times for all subjects and visits is not dense in its range.

### Author(s)

So Young Park <spark13@ncsu.edu>, Ana-Maria Staicu

### References

Park, S.Y. and Staicu, A.M. (2015). Longitudinal functional data analysis. Stat 4 212-226.

```
lfpca.dti <- fpca.lfda(Y = Y, subject.index = id,</pre>
                     visit.index = visit.index, obsT = visit.time,
                     LongiModel.method = 'lme',
                     mFPCA.pve = 0.95)
TT \leftarrow seq(0,1,length.out=41); ss = seq(0,1,length.out=93)
# estimated mean function
persp(x = ss, y = TT, z = t(lfpca.dti$bivariateSmoothMeanFunc),
     xlab="s", ylab="visit times", zlab="estimated mean fn", col='light blue')
# first three estimated marginal eigenfunctions
matplot(ss, lfpca.dti$mFPCA.efunctions[,1:3], type='l', xlab='s', ylab='estimated eigen fn')
# predicted scores function corresponding to first two marginal PCs
matplot(TT, do.call(cbind, lapply(lfpca.dti$sFPCA.xiHat.bySubj, function(a) a[,1])),
       xlab="visit time (T)", ylab="xi_hat(T)", main = "k = 1", type='l')
matplot(TT, do.call(cbind, lapply(lfpca.dti$sFPCA.xiHat.bySubj, function(a) a[,2])),
       xlab="visit time (T)", ylab="xi_hat(T)", main = "k = 2", type='1')
# prediction of cca of first two subjects at T = 0, 0.5 and 1 (black, red, green)
matplot(ss, t(lfpca.dti$fitted.values.all[[1]][c(1,21,41),]),
      type='1', lty = 1, ylab="", xlab="s", main = "Subject = 1")
matplot(ss, t(lfpca.dti$fitted.values.all[[2]][c(1,21,41),]),
      type='1', lty = 1, ylab="", xlab="s", main = "Subject = 2")
### Illustration with simulated data ###
# data generation
set.seed(1)
n <- 100 # number of subjects
ss \leftarrow seq(0,1,length.out=101)
TT <- seq(0, 1, length.out=41)
mi <- runif(n, min=6, max=15)</pre>
ij <- sapply(mi, function(a) sort(sample(1:41, size=a, replace=FALSE)))</pre>
# error variances
sigma <- 0.1
sigma_wn <- 0.2
lambdaTrue <- c(1,0.5) # True eigenvalues
eta1True <- c(0.5, 0.5^2, 0.5^3) # True eigenvalues
eta2True < c(0.5<sup>2</sup>, 0.5<sup>3</sup>) # True eigenvalues
phi <- sqrt(2)*cbind(sin(2*pi*ss),cos(2*pi*ss))</pre>
psi1 \leftarrow cbind(rep(1, length(TT)), sqrt(3)*(2*TT-1), sqrt(5)*(6*TT^2-6*TT+1))
psi2 <- sqrt(2)*cbind(sin(2*pi*TT),cos(2*pi*TT))</pre>
zeta1 <- sapply(eta1True, function(a) rnorm(n = n, mean = 0, sd = a))</pre>
```

```
zeta2 <- sapply(eta2True, function(a) rnorm(n = n, mean = 0, sd = a))</pre>
xi1 <- unlist(lapply(1:n, function(a) (zeta1 %*% t(psi1))[a,ij[[a]]] ))</pre>
xi2 <- unlist(lapply(1:n, function(a) (zeta2 %*% t(psi2))[a,ij[[a]]] ))</pre>
xi <- cbind(xi1, xi2)</pre>
Tij <- unlist(lapply(1:n, function(i) TT[ij[[i]]] ))</pre>
i <- unlist(lapply(1:n, function(i) rep(i, length(ij[[i]]))))</pre>
j <- unlist(lapply(1:n, function(i) 1:length(ij[[i]])))</pre>
X <- xi %*% t(phi)</pre>
meanFn <- function(s,t){ 0.5*t + 1.5*s + 1.3*s*t}
mu <- matrix(meanFn(s = rep(ss, each=length(Tij)), t=rep(Tij, length(ss))), nrow=nrow(X))</pre>
Y \leftarrow mu + X +
 matrix(rnorm(nrow(X)*ncol(phi), 0, sigma), nrow=nrow(X)) %*% t(phi) + #correlated error
  matrix(rnorm(length(X), 0, sigma\_wn), nrow=nrow(X)) # white noise
matplot(ss, t(Y[which(i==2),]), type='l', ylab="", xlab="functional argument",
      main="observations from subject i = 2")
# END: data generation
\# Illustration I : when covariance of scores from a mFPCA step is estimated using fpca.sc
est <- fpca.lfda(Y = Y,
                subject.index = i, visit.index = j, obsT = Tij,
                funcArg = ss, numTEvalPoints = length(TT),
                newdata = data.frame(i = c(1:3), Ltime = c(Tij[1], 0.2, 0.5)),
                fbps.knots = 35, fbps.p = 3, fbps.m = 2,
                LongiModel.method='fpca.sc',
                mFPCA.pve = 0.95, mFPCA.knots = 35, mFPCA.p = 3, mFPCA.m = 2,
                sFPCA.pve = 0.95, sFPCA.nbasis = 10, sFPCA.npc = NULL,
                gam.method = 'REML', gam.kT = 10)
# mean function (true vs. estimated)
par(mfrow=c(1,2))
persp(x=TT, y = ss, z = t(sapply(TT, function(a) meanFn(s=ss, t = a))),
       xlab="visit times", ylab="s", zlab="true mean fn")
persp(x = TT, y = ss, est\$bivariateSmoothMeanFunc,
 xlab="visit times", ylab="s", zlab="estimated mean fn", col='light blue')
par(mfrow=c(1,2))
# marginal covariance fn (true vs. estimated)
image(phi%*%diag(lambdaTrue)%*%t(phi))
image(est$mFPCA.covar)
# eigenfunctions (true vs. estimated)
matplot(ss, phi, type='l')
matlines(ss, cbind(est$mFPCA.efunctions[,1], est$mFPCA.efunctions[,2]), type='1', lwd=2)
```

```
# scree plot
plot(cumsum(est$mFPCA.scree.eval)/sum(est$mFPCA.scree.eval), type='l',
     ylab = "Percentage of variance explained")
points(cumsum(est$mFPCA.scree.eval)/sum(est$mFPCA.scree.eval), pch=16)
################
                 sFPCA step ################
par(mfrow=c(1,2))
print(est$mFPCA.npc) # k = 2
\# covariance of score functions for k = 1 (true vs. estimated)
image(psi1%*%diag(eta1True)%*%t(psi1), main='TRUE')
image(est$sFPCA.longDynCov.k[[1]], main='ESTIMATED')
# covariance of score functions for k = 2 (true vs. estimated)
image(psi2%*%diag(eta2True)%*%t(psi2))
image(est$sFPCA.longDynCov.k[[2]])
# estimated scores functions
matplot(TT, do.call(cbind,lapply(est$sFPCA.xiHat.bySubj, function(a) a[,1])),
       xlab="visit time", main="k=1", type='l', ylab="", col=rainbow(100, alpha = 1),
        lwd=1, lty=1)
matplot(TT, do.call(cbind,lapply(est$sFPCA.xiHat.bySubj, function(a) a[,2])),
        xlab="visit time", main="k=2",type='l', ylab="", col=rainbow(100, alpha = 1),
        lwd=1, lty=1)
###############
                par(mfrow=c(1,2))
# fitted
matplot(ss, t(Y[which(i==1),]), type='l', ylab="", xlab="functional argument")
matlines(ss, t(est$fitted.values[which(i==1),]), type='l', lwd=2)
# sanity check : expect fitted and predicted (obtained using info from newdata)
               values to be the same
plot(ss, est$fitted.values[1,], type='p', xlab="", ylab="", pch = 1, cex=1)
lines(ss, est$predicted.values[1,], type='1', lwd=2, col='blue')
all.equal(est$predicted.values[1,], est$fitted.values[1,])
# Illustration II : when covariance of scores from a mFPCA step is estimated using lmer
est.lme <- fpca.lfda(Y = Y,
                   subject.index = i, visit.index = j, obsT = Tij,
                   funcArg = ss, numTEvalPoints = length(TT),
                   newdata = data.frame(i = c(1:3), Ltime = c(Tij[1], 0.2, 0.5)),
                   fbps.knots = 35, fbps.p = 3, fbps.m = 2,
                   LongiModel.method='lme',
                   mFPCA.pve = 0.95, mFPCA.knots = 35, mFPCA.p = 3, mFPCA.m = 2,
                   gam.method = 'REML', gam.kT = 10)
par(mfrow=c(2,2))
```

fpca.sc

Functional principal components analysis by smoothed covariance

## **Description**

Decomposes functional observations using functional principal components analysis. A mixed model framework is used to estimate scores and obtain variance estimates.

## Usage

```
fpca.sc(
  Y = NULL
 ydata = NULL,
 Y.pred = NULL,
  argvals = NULL,
  random.int = FALSE,
 nbasis = 10,
 pve = 0.99,
  npc = NULL,
  var = FALSE,
  simul = FALSE,
  sim.alpha = 0.95,
  useSymm = FALSE,
 makePD = FALSE,
  center = TRUE,
  cov.est.method = 2,
  integration = "trapezoidal"
)
```

### **Arguments**

Y, ydata the user must supply either Y, a matrix of functions observed on a regular grid, or a data frame ydata representing irregularly observed functions. See Details. Y.pred if desired, a matrix of functions to be approximated using the FPC decomposiargvals the argument values of the function evaluations in Y, defaults to a equidistant grid from 0 to 1. If TRUE, the mean is estimated by gamm4 with random intercepts. If FALSE (the random.int default), the mean is estimated by gam treating all the data as independent. number of B-spline basis functions used for estimation of the mean function and nbasis bivariate smoothing of the covariance surface. pve proportion of variance explained: used to choose the number of principal components. prespecified value for the number of principal components (if given, this overnpc rides pve). TRUE or FALSE indicating whether model-based estimates for the variance of var FPCA expansions should be computed. simul logical: should critical values be estimated for simultaneous confidence intervals? sim.alpha 1 - coverage probability of the simultaneous intervals. logical, indicating whether to smooth only the upper triangular part of the naive useSymm covariance (when cov.est.method==2). This can save computation time for large data sets, and allows for covariance surfaces that are very peaked on the diagonal. makePD logical: should positive definiteness be enforced for the covariance surface estimate? center logical: should an estimated mean function be subtracted from Y? Set to FALSE if you have already demeaned the data using your favorite mean function estimate. covariance estimation method. If set to 1, a one-step method that applies a cov.est.method bivariate smooth to the  $y(s_1)y(s_2)$  values. This can be very slow. If set to 2 (the default), a two-step method that obtains a naive covariance estimate which is then smoothed. integration quadrature method for numerical integration; only 'trapezoidal' is currently supported.

### Details

This function computes a FPC decomposition for a set of observed curves, which may be sparsely observed and/or measured with error. A mixed model framework is used to estimate curve-specific scores and variances.

FPCA via kernel smoothing of the covariance function, with the diagonal treated separately, was proposed in Staniswalis and Lee (1998) and much extended by Yao et al. (2005), who introduced the 'PACE' method. fpca.sc uses penalized splines to smooth the covariance function, as developed by Di et al. (2009) and Goldsmith et al. (2013).

The functional data must be supplied as either

• an  $n \times d$  matrix Y, each row of which is one functional observation, with missing values allowed; or

• a data frame ydata, with columns '.id' (which curve the point belongs to, say i), '.index' (function argument such as time point t), and '.value' (observed function value  $Y_i(t)$ ).

#### Value

An object of class fpca containing:

Yhat FPC approximation (projection onto leading components) of Y. pred if speci-

fied, or else of Y.

Y the observed data

scores  $n \times npc$  matrix of estimated FPC scores.

mu estimated mean function (or a vector of zeroes if center==FALSE).

efunctions  $d \times npc$  matrix of estimated eigenfunctions of the functional covariance, i.e., the

FPC basis functions.

evalues estimated eigenvalues of the covariance operator, i.e., variances of FPC scores.

npc number of FPCs: either the supplied npc, or the minimum number of basis func-

tions needed to explain proportion pve of the variance in the observed curves.

argument values of eigenfunction evaluations

pve The percent variance explained by the returned number of PCs

sigma2 estimated measurement error variance.

diag.var diagonal elements of the covariance matrices for each estimated curve.

VarMats a list containing the estimated covariance matrices for each curve in Y.

crit.val estimated critical values for constructing simultaneous confidence intervals.

### Author(s)

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#### References

Di, C., Crainiceanu, C., Caffo, B., and Punjabi, N. (2009). Multilevel functional principal component analysis. *Annals of Applied Statistics*, 3, 458–488.

Goldsmith, J., Greven, S., and Crainiceanu, C. (2013). Corrected confidence bands for functional data using principal components. *Biometrics*, 69(1), 41–51.

Staniswalis, J. G., and Lee, J. J. (1998). Nonparametric regression analysis of longitudinal data. *Journal of the American Statistical Association*, 93, 1403–1418.

Yao, F., Mueller, H.-G., and Wang, J.-L. (2005). Functional data analysis for sparse longitudinal data. *Journal of the American Statistical Association*, 100, 577–590.

```
## Not run:
library(ggplot2)
library(reshape2)
data(cd4)
Fit.MM = fpca.sc(cd4, var = TRUE, simul = TRUE)
Fit.mu = data.frame(mu = Fit.MM$mu,
                    d = as.numeric(colnames(cd4)))
Fit.basis = data.frame(phi = Fit.MM$efunctions,
                       d = as.numeric(colnames(cd4)))
## for one subject, examine curve estimate, pointwise and simultaneous itervals
EX.MM = data.frame(fitted = Fit.MM$Yhat[EX,],
           ptwise.UB = Fit.MM$Yhat[EX,] + 1.96 * sqrt(Fit.MM$diag.var[EX,]),
           ptwise.LB = Fit.MM$Yhat[EX,] - 1.96 * sqrt(Fit.MM$diag.var[EX,]),
          simul.UB = Fit.MM$Yhat[EX,] + Fit.MM$crit.val[EX] * sqrt(Fit.MM$diag.var[EX,]),
          simul.LB = Fit.MM$Yhat[EX,] - Fit.MM$crit.val[EX] * sqrt(Fit.MM$diag.var[EX,]),
           d = as.numeric(colnames(cd4)))
## plot data for one subject, with curve and interval estimates
EX.MM.m = melt(EX.MM, id = 'd')
ggplot(EX.MM.m, aes(x = d, y = value, group = variable, color = variable, linetype = variable)) +
  geom_path() +
  scale_linetype_manual(values = c(fitted = 1, ptwise.UB = 2,
                        ptwise.LB = 2, simul.UB = 3, simul.LB = 3) +
  scale_color_manual(values = c(fitted = 1, ptwise.UB = 2,
                     ptwise.LB = 2, simul.UB = 3, simul.LB = 3)) +
  labs(x = 'Months since seroconversion', y = 'Total CD4 Cell Count')
## plot estimated mean function
ggplot(Fit.mu, aes(x = d, y = mu)) + geom_path() +
  labs(x = 'Months since seroconversion', y = 'Total CD4 Cell Count')
## plot the first two estimated basis functions
Fit.basis.m = melt(Fit.basis, id = 'd')
ggplot(subset(Fit.basis.m, variable %in% c('phi.1', 'phi.2')), aes(x = d,
y = value, group = variable, color = variable)) + geom_path()
## input a dataframe instead of a matrix
nid <- 20
nobs <- sample(10:20, nid, rep=TRUE)</pre>
ydata <- data.frame(</pre>
    .id = rep(1:nid, nobs),
    .index = round(runif(sum(nobs), 0, 1), 3))
ydata$.value <- unlist(tapply(ydata$.index,</pre>
                              ydata$.id,
                              function(x)
                                   runif(1, -.5, .5) +
                                   dbeta(x, runif(1, 6, 8), runif(1, 3, 5))
```

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```
)
Fit.MM = fpca.sc(ydata=ydata, var = TRUE, simul = FALSE)
## End(Not run)
```

fpca.ssvd

Smoothed FPCA via iterative penalized rank one SVDs.

## **Description**

Implements the algorithm of Huang, Shen, Buja (2008) for finding smooth right singular vectors of a matrix X containing (contaminated) evaluations of functional random variables on a regular, equidistant grid. If the number of smooth SVs to extract is not specified, the function hazards a guess for the appropriate number based on the asymptotically optimal truncation threshold under the assumption of a low rank matrix contaminated with i.i.d. Gaussian noise with unknown variance derived in Donoho, Gavish (2013). Please note that Donoho, Gavish (2013) should be regarded as experimental for functional PCA, and will typically not work well if you have more observations than grid points.

## Usage

```
fpca.ssvd(
  Y = NULL
 ydata = NULL,
  argvals = NULL,
  npc = NA,
  center = TRUE,
 maxiter = 15,
  tol = 1e-04,
  diffpen = 3,
  gridsearch = TRUE,
  alphagrid = 1.5^{-20:40},
  lower.alpha = 1e-05,
  upper.alpha = 1e+07,
  verbose = FALSE,
  integration = "trapezoidal"
)
```

# Arguments

Y data matrix (rows: observations; columns: grid of eval. points)
ydata a data frame ydata representing irregularly observed functions. NO

a data frame ydata representing irregularly observed functions. NOT IMPLE-MENTED for this method.

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argvals the argument values of the function evaluations in Y, defaults to a equidistant grid from 0 to 1. See Details. how many smooth SVs to try to extract, if NA (the default) the hard thresholding npc rule of Donoho, Gavish (2013) is used (see Details, References). center center Y so that its column-means are 0? Defaults to TRUE how many iterations of the power algorithm to perform at most (defaults to 15) maxiter tol convergence tolerance for power algorithm (defaults to 1e-4) diffpen difference penalty order controlling the desired smoothness of the right singular vectors, defaults to 3 (i.e., deviations from local quadratic polynomials). gridsearch use optimize or a grid search to find GCV-optimal smoothing parameters? defaults to TRUE. alphagrid grid of smoothing parameter values for grid search lower limit for for smoothing parameter if !gridsearch lower.alpha upper.alpha upper limit for smoothing parameter if !gridsearch generate graphical summary of progress and diagnostic messages? defaults to verbose **FALSE** 

#### **Details**

integration

ignored, see Details.

Note that fpca.ssvd computes smoothed orthonormal eigenvectors of the supplied function evaluations (and associated scores), not (!) evaluations of the smoothed orthonormal eigenfunctions. The smoothed orthonormal eigenvectors are then rescaled by the length of the domain defined by argvals to have a quadratic integral approximately equal to one (instead of crossproduct equal to one), so they approximate the behavior of smooth eigenfunctions. If argvals is not equidistant, fpca.ssvd will simply return the smoothed eigenvectors without rescaling, with a warning.

#### Value

an fpca object like that returned from fpca.sc, with entries Yhat, the smoothed trajectories, Y, the observed data, scores, the estimated FPC loadings, mu, the column means of Y (or a vector of zeroes if !center), efunctions, the estimated smooth FPCs (note that these are orthonormal vectors, not evaluations of orthonormal functions if argvals is not equidistant), evalues, their associated eigenvalues, and npc, the number of smooth components that were extracted.

## Author(s)

Fabian Scheipl

### References

Huang, J. Z., Shen, H., and Buja, A. (2008). Functional principal components analysis via penalized rank one approximation. *Electronic Journal of Statistics*, 2, 678-695

Donoho, D.L., and Gavish, M. (2013). The Optimal Hard Threshold for Singular Values is 4/sqrt(3). eprint arXiv:1305.5870. Available from https://arxiv.org/abs/1305.5870.

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## See Also

fpca.sc and fpca.face for FPCA based on smoothing a covariance estimate; fpca2s for a faster SVD-based approach.

## **Examples**

```
## as in Sec. 6.2 of Huang, Shen, Buja (2008):
set.seed(2678695)
n <- 101
m < -101
s1 <- 20
s2 <- 10
s <- 4
t <- seq(-1, 1, 1=m)
v1 \leftarrow t + sin(pi*t)
v2 <- cos(3*pi*t)
V <- cbind(v1/sqrt(sum(v1^2)), v2/sqrt(sum(v2^2)))</pre>
U <- matrix(rnorm(n*2), n, 2)
D \leftarrow diag(c(s1^2, s2^2))
eps <- matrix(rnorm(m*n, sd=s), n, m)</pre>
Y \leftarrow U%*D%*%t(V) + eps
smoothSV <- fpca.ssvd(Y, verbose=TRUE)</pre>
layout(t(matrix(1:4, nr=2)))
clrs <- sapply(rainbow(n), function(c)</pre>
           do.call(rgb, as.list(c(col2rgb(c)/255, .1))))
matplot(V, type="1", lty=1, col=1:2, xlab="",
         main="FPCs: true", bty="n")
matplot(smoothSV$efunctions, type="l", lty=1, col=1:5, xlab="",
         main="FPCs: estimate", bty="n")
matplot(1:m, t(U%*%D%*%t(V)), type="1", lty=1, col=clrs, xlab="", ylab="",
         main="true smooth Y", bty="n")
matplot(1:m, t(smoothSV$Yhat), xlab="", ylab="",
         type="l", lty=1,col=clrs, main="estimated smooth Y", bty="n")
```

fpca2s

Functional principal component analysis by a two-stage method

# Description

This function performs functional PCA by performing an ordinary singular value decomposition on the functional data matrix, then smoothing the right singular vectors by smoothing splines.

## Usage

```
fpca2s(
   Y = NULL,
   ydata = NULL,
```

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```
argvals = NULL,
npc = NA,
center = TRUE,
smooth = TRUE
```

### Arguments

Υ	data matrix (rows: observations; columns: grid of eval. points)
ydata	a data frame ydata representing irregularly observed functions. NOT IMPLE-MENTED for this method.
argvals	the argument values of the function evaluations in Y, defaults to a equidistant grid from 0 to 1. See Details.
npc	how many smooth SVs to try to extract, if NA (the default) the hard thresholding rule of Donoho, Gavish (2013) is used (see Details, References).
center	center Y so that its column-means are 0? Defaults to TRUE
smooth	logical; defaults to TRUE, if NULL, no smoothing of eigenvectors.

#### **Details**

Note that fpca2s computes smoothed orthonormal eigenvectors of the supplied function evaluations (and associated scores), not (!) evaluations of the smoothed orthonormal eigenfunctions. The smoothed orthonormal eigenvectors are then rescaled by the length of the domain defined by argvals to have a quadratic integral approximately equal to one (instead of crossproduct equal to one), so they approximate the behavior of smooth eigenfunctions. If argvals is not equidistant, fpca2s will simply return the smoothed eigenvectors without rescaling, with a warning.

# Value

an fpca object like that returned from fpca.sc, with entries Yhat, the smoothed trajectories, Y, the observed data, scores, the estimated FPC loadings, mu, the column means of Y (or a vector of zeroes if !center), efunctions, the estimated smooth FPCs (note that these are orthonormal vectors, not evaluations of orthonormal functions if argvals is not equidistant), evalues, their associated eigenvalues, and npc, the number of smooth components that were extracted.

## Author(s)

Luo Xiao <lxiao@jhsph.edu>, Fabian Scheipl

### References

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C., (2013), Fast covariance estimation for high-dimensional functional data. (submitted) https://arxiv.org/abs/1306.5718.

Gavish, M., and Donoho, D. L. (2014). The optimal hard threshold for singular values is 4/sqrt(3). *IEEE Transactions on Information Theory*, 60(8), 5040–5053.

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## See Also

fpca.sc and fpca.face for FPCA based on smoothing a covariance estimate; fpca.ssvd for another SVD-based approach.

```
#### settings
I <- 50 # number of subjects
J <- 3000 # dimension of the data
t \leftarrow (1:J)/J \# a regular grid on [0,1]
N <- 4 #number of eigenfunctions
sigma <- 2 ##standard deviation of random noises
lambdaTrue \leftarrow c(1,0.5,0.5<sup>2</sup>,0.5<sup>3</sup>) # True eigenvalues
case = 1
### True Eigenfunctions
if(case==1) phi <- sqrt(2)*cbind(sin(2*pi*t),cos(2*pi*t),</pre>
                               sin(4*pi*t), cos(4*pi*t))
if(case==2) phi <- cbind(rep(1,J),sqrt(3)*(2*t-1),
                       sqrt(5)*(6*t^2-6*t+1),
                       sqrt(7)*(20*t^3-30*t^2+12*t-1))
########
                                   ##############
            Generate Data
xi <- matrix(rnorm(I*N),I,N);</pre>
xi <- xi%*%diag(sqrt(lambdaTrue))</pre>
X \leftarrow xi\%\%(phi); \# of size I by J
Y <- X + sigma*matrix(rnorm(I*J),I,J)
results <- fpca2s(Y,npc=4,argvals=t)</pre>
####
                  SVDS
Phi <- results$efunctions
eigenvalues <- results$evalues
for(k in 1:N){
  if(Phi[,k]%*%phi[,k]< 0)
    Phi[,k] \leftarrow - Phi[,k]
}
### plot eigenfunctions
par(mfrow=c(N/2,2))
seq <- (1:(J/10))*10
for(k in 1:N){
    plot(t[seq], Phi[seq, k]*sqrt(J), type='l', lwd = 3,
         ylim = c(-2,2), col = 'red',
         ylab = paste('Eigenfunction ',k,sep=''),
         xlab='t',main='SVDS')
```

```
lines(t[seq],phi[seq,k],lwd = 2, col = 'black')
}
```

fpcr

Functional principal component regression

## **Description**

Implements functional principal component regression (Reiss and Ogden, 2007, 2010) for generalized linear models with scalar responses and functional predictors.

# Usage

```
fpcr(
 у,
 xfuncs = NULL,
 fdobj = NULL,
 ncomp = NULL,
 pve = 0.99,
 nbasis = NULL,
 basismat = NULL,
  penmat = NULL,
 argvals = NULL,
  covt = NULL,
 mean.signal.term = FALSE,
  spline.order = NULL,
 family = "gaussian",
 method = "REML",
  sp = NULL,
 pen.order = 2,
 cv1 = FALSE,
 nfold = 5,
  store.cv = FALSE,
  store.gam = TRUE,
)
```

# Arguments

У	scalar outcome vector.
xfuncs	for 1D predictors, an $n \times d$ matrix of signals/functional predictors, where $n$ is the length of y and $d$ is the number of sites at which each signal is defined. For 2D predictors, an $n \times d1 \times d2$ array representing $n$ images of dimension $d1 \times d2$ .
fdobj	functional data object (class "fd") giving the functional predictors. Allowed only for 1D functional predictors.
ncomp	number of principal components. If NULL, this is chosen by pve.

pve proportion of variance explained: used to choose the number of principal com-

ponents.

number(s) of B-spline basis functions: either a scalar, or a vector of values to be

compared. For 2D predictors, tensor product B-splines are used, with the given basis dimension(s) in each direction; alternatively, nbasis can be given in the form list(v1, v2), in which case cross-validation will be performed for each combination of the first-dimension basis sizes in v1 and the second-dimension basis sizes in v2. Ignored if fdobj is supplied. If fdobj is *not* supplied, this defaults to 40 (i.e., 40 B-spline basis functions) for 1D predictors, and 15 (i.e., tensor product B-splines with 15 functions per dimension) for 2D predictors.

basismat a  $d \times K$  matrix of values of K basis functions at the d sites.

penmat a  $K \times K$  matrix defining a penalty on the basis coefficients.

argvals points at which the functional predictors and the coefficient function are eval-

uated. By default, if 1D functional predictors are given by the  $n \times d$  matrix xfuncs, argvals is set to d equally spaced points from 0 to 1; if they are given by fdobj, argvals is set to 401 equally spaced points spanning the domain of the given functions. For 2D (image) predictors supplied as an  $n \times d1 \times d2$  array, argvals defaults to a list of (1) d1 equally spaced points from 0 to 1; (2) d2

equally spaced points from 0 to 1.

covt covariates: an n-row matrix, or a vector of length n.

mean.signal.term

logical: should the mean of each subject's signal be included as a covariate?

spline.order order of B-splines used, if fdobj is not supplied; defaults to 4, i.e., cubic B-

splines.

family generalized linear model family. Current version supports "gaussian" (the de-

fault) and "binomial".

method smoothing parameter selection method, passed to function gam; see the gam doc-

umentation for details.

sp a fixed smoothing parameter; if NULL, an optimal value is chosen (see method).

pen.order order of derivative penalty applied when estimating the coefficient function; de-

faults to 2.

cv1 logical: should cross-validation be performed to select the best model if only

one set of tuning parameters provided? By default, FALSE. Note that, if there are

multiple sets of tuning parameters provided, cv is always performed.

nfold the number of validation sets ("folds") into which the data are divided; by de-

fault, 5.

store.cv logical: should a CV result table be in the output? By default, FALSE.

store.gam logical: should the gam object be included in the output? Defaults to TRUE.

... other arguments passed to function gam.

#### **Details**

One-dimensional functional predictors can be given either in functional data object form, using argument fdobj (see the **fda** package of Ramsay, Hooker and Graves, 2009, and Method 1 in the

example below), or explicitly, using xfuncs (see Method 2 in the example). In the latter case, arguments basismat and penmat can also be used to specify the basis and/or penalty matrices (see Method 3).

For two-dimensional predictors, functional data object form is not supported. Instead of radial B-splines as in Reiss and Ogden (2010), this implementation employs tensor product cubic B-splines, sometimes known as bivariate O-splines (Ormerod, 2008).

For purposes of interpreting the fitted coefficients, please note that the functional predictors are decorrelated from the scalar predictors before fitting the model (when there are no scalar predictors other than the intercept, this just means the columns of the functional predictor matrix are demeaned); see section 3.2 of Reiss (2006) for details.

#### Value

A list with components

gamObject if store.gam = TRUE, an object of class gam (see gamObject in the mgcv pack-

age documentation).

fhat coefficient function estimate.

se pointwise Bayesian standard error.

undecor.coef undecorrelated coefficient for covariates.

argvals the supplied value of argvals.

cv. table a table giving the CV criterion for each combination of nbasis and ncomp, if

store.cv = TRUE; otherwise, the CV criterion only for the optimized combina-

tion of these parameters. Set to NULL if CV is not performed.

nbasis, ncomp when CV is performed, the values of nbasis and comp that minimize the CV

criterion.

## Author(s)

Philip Reiss <phil.reiss@nyumc.org>, Lan Huo <lan.huo@nyumc.org>, and Lei Huang <huangracer@gmail.com>

#### References

Ormerod, J. T. (2008). On semiparametric regression and data mining. Ph.D. dissertation, School of Mathematics and Statistics, University of New South Wales.

Ramsay, J. O., Hooker, G., and Graves, S. (2009). Functional Data Analysis with R and MATLAB. New York: Springer.

Reiss, P. T. (2006). Regression with signals and images as predictors. Ph.D. dissertation, Department of Biostatistics, Columbia University.

Reiss, P. T., and Ogden, R. T. (2007). Functional principal component regression and functional partial least squares. *Journal of the American Statistical Association*, 102, 984–996.

Reiss, P. T., and Ogden, R. T. (2010). Functional generalized linear models with images as predictors. *Biometrics*, 66, 61–69.

Wood, S. N. (2006). Generalized Additive Models: An Introduction with R. Boca Raton, FL: Chapman & Hall.

```
require(fda)
### 1D functional predictor example ###
####### Octane data example ########
data(gasoline)
# Create the requisite functional data objects
bbasis = create.bspline.basis(c(900, 1700), 40)
wavelengths = 2*450:850
nir <- t(gasoline$NIR)</pre>
gas.fd = smooth.basisPar(wavelengths, nir, bbasis)$fd
# Method 1: Call fpcr with fdobj argument
gasmod1 = fpcr(gasoline$octane, fdobj = gas.fd, ncomp = 30)
plot(gasmod1, xlab="Wavelength")
## Not run:
# Method 2: Call fpcr with explicit signal matrix
gasmod2 = fpcr(gasoline$octane, xfuncs = gasoline$NIR, ncomp = 30)
# Method 3: Call fpcr with explicit signal, basis, and penalty matrices
gasmod3 = fpcr(gasoline$octane, xfuncs = gasoline$NIR,
               basismat = eval.basis(wavelengths, bbasis),
               penmat = getbasispenalty(bbasis), ncomp = 30)
# Check that all 3 calls yield essentially identical estimates
all.equal(gasmod1$fhat, gasmod2$fhat, gasmod3$fhat)
# But note that, in general, you'd have to specify argvals in Method 1
# to get the same coefficient function values as with Methods 2 & 3.
## End(Not run)
### 2D functional predictor example ###
n = 200; d = 70
# Create true coefficient function
ftrue = matrix(0,d,d)
ftrue[40:46,34:38] = 1
# Generate random functional predictors, and scalar responses
ii = array(rnorm(n*d^2), dim=c(n,d,d))
iimat = ii; dim(iimat) = c(n,d^2)
yy = iimat %*% as.vector(ftrue) + rnorm(n, sd=.3)
mm = fpcr(yy, ii, ncomp=40)
image(ftrue)
contour(mm$fhat, add=TRUE)
## Not run:
### Cross-validation ###
cv.gas = fpcr(gasoline$octane, xfuncs = gasoline$NIR,
```

f\_sum

Sum computation 1

## **Description**

Internal function used compute a sum in FPCA-based covariance updates

# Usage

```
f_sum(mu.q.c, sig.q.c, theta, obspts.mat)
```

## **Arguments**

mu.q.c current value of mu.q.c sig.q.c current value of sig.q.c

theta spline basis

obspts.mat matrix indicating the points on which data is observed

## Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

 $f\_sum2$ 

Sum computation 2

# Description

Internal function used compute a sum in FPCA-based covariance updates

# Usage

```
f_sum2(y, fixef, mu.q.c, kt, theta)
```

## **Arguments**

y outcome matrix

fixef current estimate of fixed effects

mu.q.c current value of mu.q.c kt number of basis functions

theta spline basis

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### Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

f\_sum4

Sum computation 2

### **Description**

Internal function used compute a sum in FPCA-based covariance updates

## Usage

```
f_sum4(mu.q.c, sig.q.c, mu.q.bpsi, sig.q.bpsi, theta, obspts.mat)
```

### **Arguments**

mu.q.c current value of mu.q.c
sig.q.c current value of sig.q.c
mu.q.bpsi current value of mu.q.bpsi
sig.q.bpsi current value of sig.q.bpsi
theta current value of theta

obspts.mat matrix indicating where curves are observed

## Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

f\_trace

Trace computation

# Description

Internal function used compute a trace in FPCA-based covariance updates

### **Usage**

```
f_trace(Theta_i, Sig_q_Bpsi, Kp, Kt)
```

# Arguments

Theta\_i basis functions on observed grid points
Sig\_q\_Bpsi variance of FPC basis coefficients

Kp number of FPCs

Kt number of spline basis functions

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#### Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

gasoline

Octane numbers and NIR spectra of gasoline

# **Description**

Near-infrared reflectance spectra and octane numbers of 60 gasoline samples. Each NIR spectrum consists of log(1/reflectance) measurements at 401 wavelengths, in 2-nm intervals from 900 nm to 1700 nm. We thank Prof. John Kalivas for making this data set available.

#### **Format**

A data frame comprising

octane a numeric vector of octane numbers for the 60 samples.

**NIR** a 60 x 401 matrix of NIR spectra.

### **Source**

Kalivas, John H. (1997). Two data sets of near infrared spectra. *Chemometrics and Intelligent Laboratory Systems*, 37, 255–259.

#### References

For applications of functional principal component regression to this data set:

Reiss, P. T., and Ogden, R. T. (2007). Functional principal component regression and functional partial least squares. *Journal of the American Statistical Association*, 102, 984–996.

Reiss, P. T., and Ogden, R. T. (2009). Smoothing parameter selection for a class of semiparametric linear models. *Journal of the Royal Statistical Society, Series B*, 71(2), 505–523.

#### See Also

fpcr

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gibbs\_cs\_fpca Cross-sectional FoSR using a Gibbs sampler and FPCA

# Description

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using a Gibbs sampler and estimates the residual covariance surface using FPCA.

# Usage

```
gibbs_cs_fpca(
  formula,
 Kt = 5,
 Kp = 2,
  data = NULL,
  verbose = TRUE,
 N.iter = 5000,
 N.burn = 1000,
  SEED = NULL,
  sig2.me = 0.01,
  alpha = 0.1,
 Aw = NULL,
 Bw = NULL,
 Apsi = NULL,
 Bpsi = NULL
)
```

## **Arguments**

formula	a formula indicating the structure of the proposed model.
Kt	number of spline basis functions used to estimate coefficient functions
Кр	number of FPCA basis functions to be estimated
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
verbose	logical defaulting to TRUE – should updates on progress be printed?
N.iter	number of iterations used in the Gibbs sampler
N.burn	number of iterations discarded as burn-in
SEED	seed value to start the sampler; ensures reproducibility
sig2.me	starting value for measurement error variance
alpha	tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha = 0 is all second-derivative penalty)
Aw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects

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Bw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
Apsi	hyperparameter for inverse gamma controlling variance of spline terms for FPC effects
Bpsi	hyperparameter for inverse gamma controlling variance of spline terms for FPC effects

## Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

gibbs\_cs\_wish

Cross-sectional FoSR using a Gibbs sampler and Wishart prior

# **Description**

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using a Gibbs sampler and estimates the residual covariance surface using a Wishart prior.

# Usage

```
gibbs_cs_wish(
formula,
Kt = 5,
data = NULL,
verbose = TRUE,
N.iter = 5000,
N.burn = 1000,
alpha = 0.1,
min.iter = 10,
max.iter = 50,
Aw = NULL,
Bw = NULL,
v = NULL,
SEED = NULL
)
```

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# Arguments

formula	a formula indicating the structure of the proposed model.
Kt	number of spline basis functions used to estimate coefficient functions
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
verbose	logical defaulting to TRUE – should updates on progress be printed?
N.iter	number of iterations used in the Gibbs sampler
N.burn	number of iterations discarded as burn-in
alpha	tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha = $0$ is all second-derivative penalty)
min.iter	minimum number of iterations
max.iter	maximum number of iterations
Aw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
Bw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
V	hyperparameter for inverse Wishart prior on residual covariance
SEED	seed value to start the sampler; ensures reproducibility

# Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

## References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

gibbs_mult_fpca	Multilevel FoSR using a Gibbs sampler and FPCA	
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# Description

Fitting function for function-on-scalar regression for longitudinal data. This function estimates model parameters using a Gibbs sampler and estimates the residual covariance surface using FPCA.

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### Usage

```
gibbs_mult_fpca(
   formula,
   Kt = 5,
   Kp = 2,
   data = NULL,
   verbose = TRUE,
   N.iter = 5000,
   N.burn = 1000,
   sig2.me = 0.01,
   alpha = 0.1,
   SEED = NULL
)
```

# Arguments

formula	a f	formula ir	ndicating	the structure	of the	proposed	model.
---------	-----	------------	-----------	---------------	--------	----------	--------

Kt number of spline basis functions used to estimate coefficient functions

Kp number of FPCA basis functions to be estimated

data an optional data frame, list or environment containing the variables in the model.

If not found in data, the variables are taken from environment(formula), typi-

cally the environment from which the function is called.

verbose logical defaulting to TRUE – should updates on progress be printed?

N.iter number of iterations used in the Gibbs sampler

N. burn number of iterations discarded as burn-in

sig2.me starting value for measurement error variance

alpha tuning parameter balancing second-derivative penalty and zeroth-derivative penalty

(alpha = 0 is all second-derivative penalty)

SEED seed value to start the sampler; ensures reproducibility

## Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

#### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

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Multilevel FoSR using a Gibbs sampler and Wishart prior

## **Description**

Fitting function for function-on-scalar regression for multilevel data. This function estimates model parameters using a Gibbs sampler and estimates the residual covariance surface using a Wishart prior.

# Usage

```
gibbs_mult_wish(
formula,
Kt = 5,
data = NULL,
verbose = TRUE,
N.iter = 5000,
N.burn = 1000,
alpha = 0.1,
Az = NULL,
Bz = NULL,
Aw = NULL,
Bw = NULL,
v = NULL,
SEED = NULL
```

# Arguments

a formula indicating the structure of the proposed model.
number of spline basis functions used to estimate coefficient functions
an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
logical defaulting to TRUE – should updates on progress be printed?
number of iterations used in the Gibbs sampler
number of iterations discarded as burn-in
tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha = 0 is all second-derivative penalty)
hyperparameter for inverse gamma controlling variance of spline terms for subject-level effects
hyperparameter for inverse gamma controlling variance of spline terms for subject-level effects
hyperparameter for inverse gamma controlling variance of spline terms for population-level effects

gls\_cs

Bw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
V	hyperparameter for inverse Wishart prior on residual covariance
SEED	seed value to start the sampler; ensures reproducibility

# Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

#### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

gls\_cs

Cross-sectional FoSR using GLS

# Description

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using GLS: first, an OLS estimate of spline coefficients is estimated; second, the residual covariance is estimated using an FPC decomposition of the OLS residual curves; finally, a GLS estimate of spline coefficients is estimated. Although this is in the 'BayesFoSR' package, there is nothing Bayesian about this FoSR.

## Usage

```
gls_cs(
  formula,
  data = NULL,
  Kt = 5,
  basis = "bs",
  sigma = NULL,
  verbose = TRUE,
  CI.type = "pointwise"
)
```

# Arguments

formula	a formula indicating the structure of the proposed model.
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
Kt	number of spline basis functions used to estimate coefficient functions
basis	basis type; options are "bs" for b-splines and "pbs" for periodic b-splines

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sigma	optional covariance matrix used in GLS; if NULL, OLS will be used to estimated fixed effects, and the covariance matrix will be estimated from the residuals.
verbose	logical defaulting to TRUE – should updates on progress be printed?
CI.type	Indicates CI type for coefficient functions; options are "pointwise" and "simultaneous"

## Author(s)

Jeff Goldsmith <a jg 2202@cumc.columbia.edu>

#### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

1f

Construct an FLM regression term

## **Description**

Defines a term  $\int_T \beta(t) X_i(t) dt$  for inclusion in an mgcv: :gam-formula (or bam or gamm or gamm4: ::gamm) as constructed by pfr, where  $\beta(t)$  is an unknown coefficient function and  $X_i(t)$  is a functional predictor on the closed interval T. See smooth. terms for a list of basis and penalty options; the default is thin-plate regression splines, as this is the default option for s.

# Usage

```
lf(
    X,
    argvals = NULL,
    xind = NULL,
    integration = c("simpson", "trapezoidal", "riemann"),
    L = NULL,
    presmooth = NULL,
    presmooth.opts = NULL,
    ...
)
```

## **Arguments**

Χ

functional predictors, typically expressed as an N by J matrix, where N is the number of columns and J is the number of evaluation points. May include missing/sparse functions, which are indicated by NA values. Alternatively, can be an object of class "fd"; see fd.

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argvals indices of evaluation of X, i.e.  $(t_{i1}, ., t_{iJ})$  for subject i. May be entered as either a

length-J vector, or as an N by J matrix. Indices may be unequally spaced. Entering as a matrix allows for different observations times for each subject. If NULL, defaults to an equally-spaced grid between 0 or 1 (or within X\$basis\$rangeval

if X is a fd object.)

xind same as argvals. It will not be supported in the next version of refund.

integration method used for numerical integration. Defaults to "simpson"'s rule for calcu-

lating entries in L. Alternatively and for non-equidistant grids, "trapezoidal"

or "riemann".

L an optional N by ncol(argvals) matrix giving the weights for the numerical

integration over t. If present, overrides integration.

presmooth string indicating the method to be used for preprocessing functional predictor

prior to fitting. Options are fpca.sc, fpca.face, fpca.ssvd, fpca.bspline, and fpca.interpolate. Defaults to NULL indicating no preprocessing. See

create.prep.func.

presmooth.opts list including options passed to preprocessing method create.prep.func.

... optional arguments for basis and penalization to be passed to mgcv::s. These

could include, for example, "bs", "k", "m", etc. See s for details.

#### Value

a list with the following entries

call a call to te (or s, t2) using the appropriately constructed covariate and weight

matrices

argvals the argvals argument supplied to 1f

L the matrix of weights used for the integration

xindname the name used for the functional predictor variable in the formula used by mgcv

tindname the name used for argvals variable in the formula used by mgcv

LXname the name used for the L variable in the formula used by mgcv

presmooth the presmooth argument supplied to 1f

prep. func a function that preprocesses data based on the preprocessing method specified

in presmooth. See create.prep.func

# Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com>, Fabian Scheipl, and Jonathan Gellar

#### References

Goldsmith, J., Bobb, J., Crainiceanu, C., Caffo, B., and Reich, D. (2011). Penalized functional regression. *Journal of Computational and Graphical Statistics*, 20(4), 830-851.

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2012). Longitudinal penalized functional regression for cognitive outcomes on neuronal tract measurements. *Journal of the Royal Statistical Society: Series C*, 61(3), 453-469.

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### See Also

pfr, af, mgcv's smooth.terms and linear.functional.terms; pfr for additional examples

### **Examples**

```
data(DTI)
DTI1 <- DTI[DTI$visit==1 & complete.cases(DTI),]</pre>
# We can apply various preprocessing options to the DTI data
fit1 <- pfr(pasat ~ lf(cca, k=30), data=DTI1)</pre>
fit2 <- pfr(pasat ~ lf(cca, k=30, presmooth="fpca.sc",
                        presmooth.opts=list(nbasis=8, pve=.975)), data=DTI1)
fit3 <- pfr(pasat ~ lf(cca, k=30, presmooth="fpca.face",
                        presmooth.opts=list(m=3, npc=9)), data=DTI1)
fit4 <- pfr(pasat ~ lf(cca, k=30, presmooth="fpca.ssvd"), data=DTI1)</pre>
fit5 <- pfr(pasat ~ lf(cca, k=30, presmooth="bspline",
                        presmooth.opts=list(nbasis=8)), data=DTI1)
fit6 <- pfr(pasat ~ lf(cca, k=30, presmooth="interpolate"), data=DTI1)
# All models should result in similar fits
fits <- as.data.frame(lapply(1:6, function(i)</pre>
 get(paste0("fit",i))$fitted.values))
names(fits) <- c("none", "fpca.sc", "fpca.face", "fpca.ssvd", "bspline", "interpolate")</pre>
pairs(fits)
```

lf.vd

Construct a VDFR regression term

## Description

This function defines the a variable-domain functional regression term for inclusion in an gamformula (or bam or gamm or gamm4::gamm as constructed by pfr. These are functional predictors for which each function is observed over a domain of different width. The default is the term  $1/T_i \int_0^{T_i} X_i(t) \beta(t,T_i) dt$ , where  $X_i(t)$  is a functional predictor of length  $T_i$  and  $\beta(t,T_i)$  is an unknown bivariate coefficient function. Various domain transformations are available, such as lagging or domain-standardizing the coordinates, or parameterizing the interactions; these often result in improved model fit. Basis choice is fully customizable using the options of s and te.

# Usage

```
lf.vd(
   X,
   argvals = seq(0, 1, 1 = ncol(X)),
   vd = NULL,
   integration = c("simpson", "trapezoidal", "riemann"),
   L = NULL,
   basistype = c("s", "te", "t2"),
```

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```
transform = NULL,
mp = TRUE,
...
)
```

### **Arguments**

Х	matrix containing variable-domain functions. Should be $NxJ$ , where $N$ is the number of subjects and $J$ is the maximum number of time points per subject. Most rows will have NA values in the right-most columns, corresponding to unobserved time points.
argvals	indices of evaluation of X, i.e. $(t_{i1},.,t_{iJ})$ for subject $i$ . May be entered as either a length-J vector, or as an N by J matrix. Indices may be unequally spaced. Entering as a matrix allows for different observations times for each subject.
vd	vector of values of containing the variable-domain width ( $T_i$ above). Defaults to the argvals value corresponding to the last non-NA element of $X_i(t)$ .
integration	method used for numerical integration. Defaults to "simpson"'s rule for calculating entries in L. Alternatively and for non-equidistant grids, "trapezoidal" or "riemann".
L	an optional N by ncol(argvals) matrix giving the weights for the numerical integration over t. If present, overrides integration.
basistype	character string indicating type of bivariate basis used. Options include "s" (the default), "te", and "t2", which correspond to mgcv::s, mgcv::te, and mgcv::t2.
transform	character string indicating an optional basis transformation; see Details for options.
тр	for transform=="linear" or transform=="quadratic", TRUE to use multiple penalties for the smooth (one for each marginal basis). If FALSE, penalties are concatonated into a single block-diagonal penalty matrix (with one smoothing parameter).
	optional arguments for basis and penalization to be passed to the function indicated by basistype. These could include, for example, "bs", "k", "m", etc. See te or s for details.

## **Details**

The variable-domain functional regression model uses the term  $\frac{1}{T_i}\int_0^{T_i}X_i(t)\beta(t,T_i)dt$  to incorporate a functional predictor with subject-specific domain width. This term imposes a smooth (non-parametric) interaction between t and  $T_i$ . The domain of the coefficient function is the triangular (or trapezoidal) surface defined by  $t,T_i:0\leq t\leq T_i$ . The default basis uses bivariate thin-plate regression splines.

Different basis transformations can result in different properties; see Gellar, et al. (2014) for a more complete description. We make five basis transformations easily accessible using the transform argument. This argument is a character string that can take one of the following values:

1. "lagged": transforms argvals to argvals - vd

2. "standardized": transforms argvals to argvals/vd, and then rescales vd linearly so it ranges from  $0\ \text{to}\ 1$ 

- 3. "linear": first transforms the domain as in "standardized", then parameterizes the interaction with "vd" to be linear
- 4. "quadratic": first transforms the domain as in "standardized", then parameterizes the interaction with "vd" to be quadratic
- 5. "noInteraction": first transforms the domain as in "standardized", then reduces the bivariate basis to univariate with no effect of vd. This would be equivalent to using 1f on the domain-standardized predictor functions.

The practical effect of using the "lagged" basis is to increase smoothness along the right (diagonal) edge of the resultant estimate. The practical effect of using a "standardized" basis is to allow for greater smoothness at high values of  $T_i$  compared to lower values.

These basis transformations rely on the basis constructors available in the mgcvTrans package. For more specific control over the transformations, you can use bs="dt" and/or bs="pi"; see smooth.construct.dt.smooth.spec or smooth.construct.pi.smooth.spec for an explanation of the options (entered through the xt argument of lf.vd/s).

Note that tensor product bases are only recommended when a standardized transformation is used. Without this transformation, just under half of the "knots" used to define the basis will fall outside the range of the data and have no data available to estimate them. The penalty allows the corresponding coefficients to be estimated, but results may be unstable.

#### Value

a list with the following entries

call a call to s or te, using the appropriately constructed weight matrices
data data used by the call, which includes the matrices indicated by argname, Tindname,

and LXname

L the matrix of weights used for the integration

the name used for the argvals variable in the formula used by mgcv::gam
the name used for the Tind variable in the formula used by mgcv::gam

LXname the name of the by variable used by s or te in the formula for mgcv::gam

## Author(s)

Jonathan E. Gellar < JGellar@mathematica-mpr.com>

#### References

Gellar, Jonathan E., Elizabeth Colantuoni, Dale M. Needham, and Ciprian M. Crainiceanu. Variable-Domain Functional Regression for Modeling ICU Data. Journal of the American Statistical Association, 109(508):1425-1439, 2014.

#### See Also

```
pfr, lf, mgcv's linear.functional.terms.
```

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### **Examples**

```
## Not run:
 data(sofa)
 fit.vd1 <- pfr(death ~ lf.vd(SOFA) + age + los,
                 family="binomial", data=sofa)
 fit.vd2 <- pfr(death ~ lf.vd(SOFA, transform="lagged") + age + los,
                 family="binomial", data=sofa)
 fit.vd3 <- pfr(death ~ lf.vd(SOFA, transform="standardized") + age + los,</pre>
                 family="binomial", data=sofa)
 fit.vd4 <- pfr(death ~ lf.vd(SOFA, transform="standardized",</pre>
                                basistype="te") + age + los,
                 family="binomial", data=sofa)
 fit.vd5 <- pfr(death ~ lf.vd(SOFA, transform="linear", bs="ps") + age + los,</pre>
                 family="binomial", data=sofa)
 fit.vd6 <- pfr(death ~ lf.vd(SOFA, transform="quadratic", bs="ps") + age + los,</pre>
                 family="binomial", data=sofa)
 fit.vd7 <- pfr(death ~ lf.vd(SOFA, transform="noInteraction", bs="ps") + age + los,</pre>
                 family="binomial", data=sofa)
 ests <- lapply(1:7, function(i) {
    c.i <- coef(get(paste0("fit.vd", i)), n=173, n2=173)</pre>
    c.i[(c.i$SOFA.arg <= c.i$SOFA.vd),]</pre>
 })
 # Try plotting for each i
 i <- 1
 lims <- c(-2,8)
 if (requireNamespace("ggplot2", quietly = TRUE) &
      requireNamespace("RColorBrewer", quietly = TRUE)) {
        est <- ests[[i]]</pre>
        est$value[est$value<lims[1]] <- lims[1]</pre>
        est$value[est$value>lims[2]] <- lims[2]
        ggplot2::ggplot(est, ggplot2::aes(SOFA.arg, SOFA.vd)) +
          ggplot2::geom_tile(ggplot2::aes(colour=value, fill=value)) +
          ggplot2::scale_fill_gradientn( name="", limits=lims,
                     colours=rev(RColorBrewer::brewer.pal(11, "Spectral"))) +
          ggplot2::scale_colour_gradientn(name="", limits=lims,
                    colours=rev(RColorBrewer::brewer.pal(11, "Spectral"))) +
          ggplot2::scale_y\_continuous(expand = c(0,0)) +
          ggplot2::scale_x_continuous(expand = c(0,0)) +
          ggplot2::theme_bw()
 }
## End(Not run)
```

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## **Description**

Defines a term  $\int_T \beta(t) X_i(t) dt$  for inclusion in an gam-formula (or bam or gamm or gamm4) as constructed by fgam, where  $\beta(t)$  is an unknown coefficient function and  $X_i(t)$  is a functional predictor on the closed interval T. Defaults to a cubic B-spline with second-order difference penalties for estimating  $\beta(t)$ . The functional predictor must be fully observed on a regular grid.

## Usage

```
lf_old(
    X,
    argvals = seq(0, 1, l = ncol(X)),
    xind = NULL,
    integration = c("simpson", "trapezoidal", "riemann"),
    L = NULL,
    splinepars = list(bs = "ps", k = min(ceiling(n/4), 40), m = c(2, 2)),
    presmooth = TRUE
)
```

### **Arguments**

X	an N by J=ncol(argvals) matrix of function evaluations $X_i(t_{i1}),.,X_i(t_{iJ}); i=1,.,N.$
argvals	matrix (or vector) of indices of evaluations of $X_i(t)$ ; i.e. a matrix with $i$ th row $(t_{i1},.,t_{iJ})$
xind	same as argvals. It will not be supported in the next version of refund.
integration	method used for numerical integration. Defaults to "simpson"'s rule for calculating entries in L. Alternatively and for non-equidistant grids, "trapezoidal" or "riemann". "riemann" integration is always used if L is specified
L	an optional N by $ncol(argvals)$ matrix giving the weights for the numerical integration over t
splinepars	optional arguments specifying options for representing and penalizing the functional coefficient $\beta(t)$ . Defaults to a cubic B-spline with second-order difference penalties, i.e. list(bs="ps", m=c(2, 1)) See te or s for details
presmooth	logical; if true, the functional predictor is pre-smoothed prior to fitting. See smooth.basisPar

## Value

a list with the following entries

- 1. call a call to te (or s, t2) using the appropriately constructed covariate and weight matrices
- 2. argvals the argvals argument supplied to 1f
- 3. L the matrix of weights used for the integration
- 4. xindname the name used for the functional predictor variable in the formula used by mgcv
- 5. tindname the name used for argvals variable in the formula used by mgcv

lpeer lpeer

- 6. LXname the name used for the L variable in the formula used by mgcv
- 7. presmooth the presmooth argument supplied to 1f
- 8. Xfd an fd object from presmoothing the functional predictors using smooth.basisPar. Only present if presmooth=TRUE. See fd

### Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com> and Fabian Scheipl

#### See Also

```
fgam, af, mgcv's linear.functional.terms, fgam for examples
```

lpeer

Longitudinal Functional Models with Structured Penalties

## **Description**

Implements longitudinal functional model with structured penalties (Kundu et al., 2012) with scalar outcome, single functional predictor, one or more scalar covariates and subject-specific random intercepts through mixed model equivalence.

# Usage

```
lpeer(
   Y,
   subj,
   t,
   funcs,
   argvals = NULL,
   covariates = NULL,
   comm.pen = TRUE,
   pentype = "Ridge",
   L.user = NULL,
   f_t = NULL,
   Q = NULL,
   phia = 10^3,
   se = FALSE,
   ...
)
```

## **Arguments**

Y vector of all outcomes over all visits or timepoints
subj vector containing the subject number for each observation
t vector containing the time information when the observation are taken

funcs matrix containing observed functional predictors as rows. Rows with NA and Inf values will be deleted. argvals matrix (or vector) of indices of evaluations of  $X_i(t)$ ; i.e. a matrix with ith row  $(t_{i1},.,t_{iJ})$ matrix of scalar covariates. covariates logical value indicating whether common penalty for all the components of recomm.pen gression function. Default is TRUE. type of penalty: either decomposition based penalty ('DECOMP') or ridge ('RIDGE') pentype or second-order difference penalty ('D2') or any user defined penalty ('USER'). For decomposition based penalty user need to specify Q matrix in Q argument (see details). For user defined penalty user need to specify L matrix in L argument (see details). For Ridge and second-order difference penalty, specification for arguments L and Q will be ignored. Default is 'RIDGE'. L.user penalty matrix. Need to be specified with pentype='USER'. When comm.pen=TRUE, Number of columns need to be equal with number of columns of matrix specified to funcs. When comm.pen=FALSE, Number of columns need to be equal with the number of columns of matrix specified to funcs times the number of components of regression function. Each row represents a constraint on functional predictor. This argument will be ignored when value of pentype is other than 'USER'. f\_t vector or matrix with number of rows equal to number of total observations and number of columns equal to d (see details). If matrix then each column pertains to single function of time and the value in the column represents the realization corresponding to time vector t. The column with intercept or multiple of intercept will be dropped. A NULL value refers to time-invariant regression function. Default value is NULL. Q Q matrix to derive decomposition based penalty. Need to be specified with pentype='DECOMP'. When comm.pen=TRUE, number of columns must equal number of columns of matrix specified to funcs. When comm.pen=FALSE, Number of columns need to be equal with the number of columns of matrix specified to funcs times the number of components of regression function. Each row represents a basis function where functional predictor is expected lie according to prior belief. This argument will be ignored when value of pentype is other than 'DECOMP'. phia scalar value of a in decomposition based penalty. Needs to be specified with pentype='DECOMP'.

#### Details

se

If there are any missing or infinite values in Y, subj, t, covariates, funcs and f\_t, the corresponding row (or observation) will be dropped, and infinite values are not allowed for these arguments. Neither Q nor L may contain missing or infinite values. 1peer() fits the following model:

logical; calculate standard error when TRUE.

additional arguments passed to 1me.

$$y_{i(t)} = X_{i(t)}^T \beta + \int W_{i(t)}(s) \gamma(t, s) ds + Z_{i(t)} u_i + \epsilon_{i(t)}$$

where  $\epsilon_{i(t)}$   $N(0, \sigma^2)$  and  $u_i$   $N(0, \sigma_u^2)$ . For all the observations, predictor function  $W_{i(t)}(s)$  is evaluated at K sampling points. Here, regression function  $\gamma(t, s)$  is represented in terms of (d+1) component functions  $\gamma_0(s), ..., \gamma_d(s)$  as follows

$$\gamma(t,s) = \gamma_0(s) + f_1(t)\gamma_1(s) + f_d(t)\gamma_d(s)$$

Values of  $y_{i(t)}, X_{i(t)}$  and  $W_{i(t)}(s)$  are passed through argument Y, covariates and funcs, respectively. Number of elements or rows in Y, t, subj, covariates (if not NULL) and funcs need to be equal.

Values of  $f_1(t), ..., f_d(t)$  are passed through  $f_t$  argument. The matrix passed through  $f_t$  argument should have d columns where each column represents one and only one of  $f_1(t), ..., f_d(t)$ .

The estimate of (d+1) component functions  $\gamma_0(s),...,\gamma_d(s)$  is obtained as penalized estimated. The following 3 types of penalties can be used for a component function:

- i. Ridge:  $I_K$
- ii. Second-order difference:  $[d_{i,j}]$  with  $d_{i,i} = d_{i,i+2} = 1, d_{i,i+1} = -2$ , otherwise  $d_{i,j} = 0$
- iii. Decomposition based penalty:  $bP_Q + a(I P_Q)$  where  $P_Q = Q^T(QQ^T)^{-1}Q$

For Decomposition based penalty the user must specify pentype= 'DECOMP' and the associated Q matrix must be passed through the Q argument. Alternatively, one can directly specify the penalty matrix by setting pentype= 'USER' and using the L argument to supply the associated L matrix.

If Q (or L) matrix is similar for all the component functions then argument comm. pen should have value TRUE and in that case specified matrix to argument Q (or L) should have K columns. When Q (or L) matrix is different for all the component functions then argument comm. pen should have value FALSE and in that case specified matrix to argument Q (or L) should have K(d+1) columns. Here first K columns pertains to first component function, second K columns pertains to second component functions, and so on.

Default penalty is Ridge penalty for all the component functions and user needs to specify 'RIDGE'. For second-order difference penalty, user needs to specify 'D2'. When pentype is 'RIDGE' or 'D2' the value of comm.pen is always TRUE and comm.pen=FALSE will be ignored.

## Value

## A list containing:

fit result of the call to lme fitted.vals predicted outcomes

BetaHat parameter estimates for scalar covariates including intercept

se.Beta standard error of parameter estimates for scalar covariates including intercept

Beta parameter estimates with standard error for scalar covariates including intercept

GammaHat estimates of components of regression functions. Each column represents one

estimates of components of regression functions. Each column represents on

component function.

Se.Gamma standard error associated with GammaHat

AIC AIC value of fit (smaller is better)

BIC BIC value of fit (smaller is better)

logLik (restricted) log-likelihood at convergence

lambda list of estimated smoothing parameters associated with each component function

V	conditional variance of Y treating only random intercept as random one.
V1	unconditional variance of Y
N	number of subjects
K	number of Sampling points in functional predictor
TotalObs	total number of observations over all subjects
Sigma.u	estimated sd of random intercept.
sigma	estimated within-group error standard deviation.

#### Author(s)

Madan Gopal Kundu <mgkundu@iupui.edu>

### References

Kundu, M. G., Harezlak, J., and Randolph, T. W. (2012). Longitudinal functional models with structured penalties (arXiv:1211.4763 [stat.AP]).

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323–353.

#### See Also

```
peer, plot.lpeer
```

# **Examples**

```
## Not run:
# Example 1: Estimation with Ridge penalty
#-----
##Load Data
data(DTI)
## Extract values for arguments for lpeer() from given data
cca = DTI$cca[which(DTI$case == 1),]
DTI = DTI[which(DTI$case == 1),]
##1.1 Fit the model with single component function
     gamma(t,s)=gamm0(s)
t<- DTI$visit
fit.cca.lpeer1 = lpeer(Y=DTI$pasat, t=t, subj=DTI$ID, funcs = cca)
plot(fit.cca.lpeer1)
##1.2 Fit the model with two component function
     gamma(t,s)=gamm0(s) + t*gamma1(s)
fit.cca.lpeer2 = lpeer(Y=DTI$pasat, t=t, subj=DTI$ID, funcs = cca,
                   f_t=t, se=TRUE)
plot(fit.cca.lpeer2)
```

```
# Example 2: Estimation with structured penalty (need structural
        information about regression function or predictor function)
##Load Data
data(PEER.Sim)
## Extract values for arguments for lpeer() from given data
W<- PEER.Sim[,c(3:(K+2))]
Y<- PEER.Sim[,K+3]
t<- PEER.Sim[,2]
id<- PEER.Sim[,1]</pre>
##Load Q matrix containing structural information
data(Q)
##2.1 Fit the model with two component function
    gamma(t,s)=gamm0(s) + t*gamma1(s)
Fit1<- lpeer(Y=Y, subj=id, t=t, covariates=cbind(t), funcs=W,
     pentype='DECOMP', f_t=cbind(1,t), Q=Q, se=TRUE)
Fit1$Beta
plot(Fit1)
##2.2 Fit the model with three component function
      gamma(t,s)=gamm0(s) + t*gamma1(s) + t^2*gamma1(s)
Fit2<- lpeer(Y=Y, subj=id, t=t, covariates=cbind(t), funcs=W,
       pentype='DECOMP', f_t=cbind(1,t, t^2), Q=Q, se=TRUE)
Fit2$Beta
plot(Fit2)
##2.3 Fit the model with two component function with different penalties
      gamma(t,s)=gamm0(s) + t*gamma1(s)
Q1 \leftarrow cbind(Q, Q)
Fit3<- lpeer(Y=Y, subj=id, t=t, covariates=cbind(t), comm.pen=FALSE, funcs=W,
       pentype='DECOMP', f_t=cbind(1,t), Q=Q1, se=TRUE)
##2.4 Fit the model with two component function with user defined penalties
      gamma(t,s)=gamm0(s) + t*gamma1(s)
phia<- 10<sup>3</sup>
P_Q \leftarrow t(Q)%*solve(Q%*%t(Q))%*% Q
L<- phia*(diag(K)- P_Q) + 1*P_Q
Fit4<- lpeer(Y=Y, subj=id, t=t, covariates=cbind(t), funcs=W,
       pentype='USER', f_t=cbind(1,t), L=L, se=TRUE)
L1<- adiag(L, L)
Fit5<- lpeer(Y=Y, subj=id, t=t, covariates=cbind(t), comm.pen=FALSE, funcs=W,
       pentype='USER', f_t=cbind(1,t), L=L1, se=TRUE)
## End(Not run)
```

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lpfr

Longitudinal penalized functional regression

### **Description**

Implements longitudinal penalized functional regression (Goldsmith et al., 2012) for generalized linear functional models with scalar outcomes and subject-specific random intercepts.

# Usage

```
lpfr(
   Y,
   subj,
   covariates = NULL,
   funcs,
   kz = 30,
   kb = 30,
   smooth.cov = FALSE,
   family = "gaussian",
   method = "REML",
   ...
)
```

# Arguments

Υ	vector of all outcomes over all visits
subj	vector containing the subject number for each observation
covariates	matrix of scalar covariates
funcs	matrix or list of matrices containing observed functional predictors as rows. NA values are allowed.
kz	dimension of principal components basis for the observed functional predictors
kb	dimension of the truncated power series spline basis for the coefficient function
smooth.cov	logical; do you wish to smooth the covariance matrix of observed functions? Increases computation time, but results in smooth principal components
family	generalized linear model family
method	method for estimating the smoothing parameters; defaults to REML
	additional arguments passed to gam to fit the regression model.

## **Details**

Functional predictors are entered as a matrix or, in the case of multiple functional predictors, as a list of matrices using the funcs argument. Missing values are allowed in the functional predictors, but it is assumed that they are observed over the same grid. Functional coefficients and confidence bounds are returned as lists in the same order as provided in the funcs argument, as are principal component and spline bases.

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#### Value

fit result of the call to gam fitted.vals predicted outcomes

betaHat list of estimated coefficient functions

beta.covariates

parameter estimates for scalar covariates

ranef vector of subject-specific random intercepts

X design matrix used in the model fit

phi list of truncated power series spline bases for the coefficient functions

psi list of principal components basis for the functional predictors

varBetaHat list containing covariance matrices for the estimated coefficient functions

Bounds list of bounds of a 95% confidence interval for the estimated coefficient func-

tions

## Author(s)

Jeff Goldsmith <jeff.goldsmith@columbia.edu>

#### References

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2012). Longitudinal penalized functional regression for cognitive outcomes on neuronal tract measurements. *Journal of the Royal Statistical Society: Series C*, 61(3), 453–469.

## **Examples**

```
## Not run:
# use longitudinal data to regress continuous outcomes on
# functional predictors (continuous outcomes only recorded for
# case == 1)
data(DTI)
# subset data as needed for this example
cca = DTI$cca[which(DTI$case == 1),]
rcst = DTI$rcst[which(DTI$case == 1),]
DTI = DTI[which(DTI$case == 1),]
# note there is missingness in the functional predictors
apply(is.na(cca), 2, mean)
apply(is.na(rcst), 2, mean)
# fit two models with single functional predictors and plot the results
fit.cca = lpfr(Y=DTI$pasat, subj=DTI$ID, funcs = cca, smooth.cov=FALSE)
```

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```
fit.rcst = lpfr(Y=DTI$pasat, subj=DTI$ID, funcs = rcst, smooth.cov=FALSE)
par(mfrow = c(1,2))
matplot(cbind(fit.cca$BetaHat[[1]], fit.cca$Bounds[[1]]),
  type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
  main = "CCA")
matplot(cbind(fit.rcst$BetaHat[[1]], fit.rcst$Bounds[[1]]),
  type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
  main = "RCST")
# fit a model with two functional predictors and plot the results
fit.cca.rcst = lpfr(Y=DTI$pasat, subj=DTI$ID, funcs = list(cca,rcst),
  smooth.cov=FALSE)
par(mfrow = c(1,2))
matplot(cbind(fit.cca.rcst$BetaHat[[1]], fit.cca.rcst$Bounds[[1]]),
  type = '1', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
  main = "CCA")
matplot(cbind(fit.cca.rcst$BetaHat[[2]], fit.cca.rcst$Bounds[[2]]),
  type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
  main = "RCST")
## End(Not run)
```

mfpca.face

Multilevel functional principal components analysis with fast covariance estimation

### Description

Decompose dense or sparse multilevel functional observations using multilevel functional principal component analysis with the fast covariance estimation approach.

## Usage

```
mfpca.face(
   Y,
   id,
   visit = NULL,
   twoway = TRUE,
   weight = "obs",
   argvals = NULL,
   pve = 0.99,
   npc = NULL,
   p = 3,
   m = 2,
   knots = 35,
   silent = TRUE
)
```

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## **Arguments**

Υ	A multilevel functional dataset on a regular grid stored in a matrix. Each row of the data is the functional observations at one visit for one subject. Missingness is allowed and need to be labeled as NA. The data must be specified.
id	A vector containing the id information to identify the subjects. The data must be specified.
visit	A vector containing information used to identify the visits. If not provided, assume the visit id are 1,2, for each subject.
twoway	Logical, indicating whether to carry out twoway ANOVA and calculate visit-specific means. Defaults to TRUE.
weight	The way of calculating covariance. weight = "obs" indicates that the sample covariance is weighted by observations. weight = "subj" indicates that the sample covariance is weighted equally by subjects. Defaults to "obs".
argvals	A vector containing observed locations on the functional domain.
pve	Proportion of variance explained. This value is used to choose the number of principal components for both levels.
npc	Pre-specified value for the number of principal components. If given, this over-rides pve.
p	The degree of B-splines functions to use. Defaults to 3.
m	The order of difference penalty to use. Defaults to 2.
knots	Number of knots to use or the vectors of knots. Defaults to 35.
silent	Logical, indicating whether to not display the name of each step. Defaults to TRUE.

## **Details**

The fast MFPCA approach (Cui et al., 2023) uses FACE (Xiao et al., 2016) to estimate covariance functions and mixed model equations (MME) to predict scores for each level. As a result, it has lower computational complexity than MFPCA (Di et al., 2009) implemented in the mfpca.sc function, and can be applied to decompose data sets with over 10000 subjects and over 10000 dimensions.

# Value

## A list containing:

Yhat FPC approximation (projection onto leading components) of Y, estimated curves

for all subjects and visits

Yhat. subject Estimated subject specific curves for all subjects

Y. df The observed data

mu estimated mean function (or a vector of zeroes if center==FALSE).

eta The estimated visit specific shifts from overall mean. scores A matrix of estimated FPC scores for level1 and level2. mfpca.sc 97

efunctions	A matrix of estimated eigenfunctions of the functional covariance, i.e., the FPC basis functions for levels 1 and 2.
evalues	Estimated eigenvalues of the covariance operator, i.e., variances of FPC scores for levels 1 and 2.
pve	The percent variance explained by the returned number of PCs.
npc	Number of FPCs: either the supplied npc, or the minimum number of basis functions needed to explain proportion pve of the variance in the observed curves for levels 1 and 2.
sigma2	Estimated measurement error variance.

### Author(s)

Ruonan Li <rli20@ncsu.edu>, Erjia Cui <ecui@umn.edu>

#### References

Cui, E., Li, R., Crainiceanu, C., and Xiao, L. (2023). Fast multilevel functional principal component analysis. *Journal of Computational and Graphical Statistics*, 32(3), 366-377.

Di, C., Crainiceanu, C., Caffo, B., and Punjabi, N. (2009). Multilevel functional principal component analysis. *Annals of Applied Statistics*, 3, 458-488.

Xiao, L., Ruppert, D., Zipunnikov, V., and Crainiceanu, C. (2016). Fast covariance estimation for high-dimensional functional data. *Statistics and Computing*, 26, 409-421.

## **Examples**

## **Description**

Decomposes functional observations using functional principal components analysis. A mixed model framework is used to estimate scores and obtain variance estimates.

## Usage

```
mfpca.sc(
   Y = NULL,
   id = NULL,
   visit = NULL,
   twoway = FALSE,
   argvals = NULL,
   nbasis = 10,
```

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```
pve = 0.99,
npc = NULL,
makePD = FALSE,
center = TRUE,
cov.est.method = 2,
integration = "trapezoidal"
)
```

### **Arguments**

Υ	The user must supply a matrix of functions on a regular grid
id	Must be supplied, a vector containing the id information used to identify clusters
visit	A vector containing information used to identify visits. Defaults to NULL.
twoway	logical, indicating whether to carry out twoway ANOVA and calculate visit-specific means. Defaults to FALSE.
argvals	function argument.
nbasis	number of B-spline basis functions used for estimation of the mean function and bivariate smoothing of the covariance surface.
pve	proportion of variance explained: used to choose the number of principal components.
npc	prespecified value for the number of principal components (if given, this overrides pve).
makePD	logical: should positive definiteness be enforced for the covariance surface estimate? Defaults to FALSE Only FALSE is currently supported.
center	logical: should an estimated mean function be subtracted from Y? Set to FALSE if you have already demeaned the data using your favorite mean function estimate.
cov.est.method	covariance estimation method. If set to 1, a one-step method that applies a bivariate smooth to the $y(s_1)y(s_2)$ values. This can be very slow. If set to 2 (the default), a two-step method that obtains a naive covariance estimate which is then smoothed. 2 is currently supported.
integration	quadrature method for numerical integration; only "trapezoidal" is currently supported.

## **Details**

This function computes a multilevel FPC decomposition for a set of observed curves, which may be sparsely observed and/or measured with error. A mixed model framework is used to estimate level 1 and level 2 scores.

MFPCA was proposed in Di et al. (2009), with variations for MFPCA with sparse data in Di et al. (2014). mfpca.sc uses penalized splines to smooth the covariance functions, as Described in Di et al. (2009) and Goldsmith et al. (2013).

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#### Value

An object of class mfpca containing:

Yhat FPC approximation (projection onto leading components) of Y, estimated curves

for all subjects and visits

Yhat. subject estimated subject specific curves for all subjects

Y the observed data

scores  $n \times npc$  matrix of estimated FPC scores for level1 and level2. mu estimated mean function (or a vector of zeroes if center==FALSE).

efunctions  $d \times npc$  matrix of estimated eigenfunctions of the functional covariance, i.e., the

FPC basis functions for levels 1 and 2.

evalues estimated eigenvalues of the covariance operator, i.e., variances of FPC scores

for levels 1 and 2.

npc number of FPCs: either the supplied npc, or the minimum number of basis func-

tions needed to explain proportion pve of the variance in the observed curves for

levels 1 and 2.

sigma2 estimated measurement error variance.

eta the estimated visit specific shifts from overall mean.

### Author(s)

Julia Wrobel <jw3134@cumc.columbia.edu>, Jeff Goldsmith <jeff.goldsmith@columbia.edu>, and Chongzhi Di

#### References

Di, C., Crainiceanu, C., Caffo, B., and Punjabi, N. (2009). Multilevel functional principal component analysis. *Annals of Applied Statistics*, 3, 458–488.

Di, C., Crainiceanu, C., Caffo, B., and Punjabi, N. (2014). Multilevel sparse functional principal component analysis. *Stat*, 3, 126–143.

Goldsmith, J., Greven, S., and Crainiceanu, C. (2013). Corrected confidence bands for functional data using principal components. *Biometrics*, 69(1), 41–51.

# Examples

```
## Not run:
data(DTI)
DTI = subset(DTI, Nscans < 6) ## example where all subjects have 6 or fewer visits
id = DTI$ID
Y = DTI$cca
mfpca.DTI = mfpca.sc(Y=Y, id = id, twoway = TRUE)
## End(Not run)</pre>
```

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model.matrix.pffr

Obtain model matrix for a pffr fit

# Description

Obtain model matrix for a pffr fit

## Usage

```
## S3 method for class 'pffr'
model.matrix(object, ...)
```

## **Arguments**

object a fitted pffr-object

... other arguments, passed to predict.gam.

#### Value

A model matrix

# Author(s)

Fabian Scheipl

ols\_cs

Cross-sectional FoSR using GLS

## **Description**

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using GLS: first, an OLS estimate of spline coefficients is estimated; second, the residual covariance is estimated using an FPC decomposition of the OLS residual curves; finally, a GLS estimate of spline coefficients is estimated. Although this is in the 'BayesFoSR' package, there is nothing Bayesian about this FoSR.

## Usage

```
ols_cs(formula, data = NULL, Kt = 5, basis = "bs", verbose = TRUE)
```

#### **Arguments**

Tormula a formula indicating the structure of the proposed model	formula	a formula indicating the structure of the proposed model
--	---------	--

data an optional data frame, list or environment containing the variables in the model.

If not found in data, the variables are taken from environment(formula), typi-

cally the environment from which the function is called.

kt number of spline basis functions used to estimate coefficient functions basis basis type; options are "bs" for b-splines and "pbs" for periodic b-splines verbose logical defaulting to TRUE – should updates on progress be printed?

### Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

#### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

pco\_predict\_preprocess

Make predictions using pco basis terms

## **Description**

This function performs the necessary preprocessing for making predictions with gam models that include pco basis terms. The function pco\_predict\_preprocess builds a data.frame (or augments an existing one) to be used with the usual predict function.

### Usage

```
pco_predict_preprocess(model, newdata = NULL, dist_list)
```

## Arguments

model a fitted gam model with at least one term of class "pco. smooth".

newdata data frame including the new values for any non-pco terms in the original fit. If

there were none, this can be left as NULL.

dist\_list a list of  $n \times n*$  matrices, one per pco term in the model, giving the distances

from the n\* prediction points to the n design points (original observations). List entry names should correspond to the names of the terms in the model (e.g., if the model includes a s(x) term, dist\_list must include an element named

"x").

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#### **Details**

Models with pco basis terms are fitted by inputting distances among the observations and then regressing (with a ridge penalty) on leading principal coordinates arising from these distances. To perform prediction, we must input the distances from the new data points to the original points, and then "insert" the former into the principal coordinate space by the interpolation method of Gower (1968) (see also Miller, 2012).

An example of how to use this function in practice is shown in smooth.construct.pco.smooth.spec.

#### Value

a data.frame with the coordinates for the new data inserted into principal coordinate space, in addition to the supplied newdata if this was non-NULL. This can be used as the newdata argument in a call to predict.gam.

### Author(s)

David L Miller

#### References

Gower, J. C. (1968). Adding a point to vector diagrams in multivariate analysis. Biometrika, 55(3), 582-585.

Miller, D. L. (2012). On smooth models for complex domains and distances. PhD dissertation, Department of Mathematical Sciences, University of Bath.

# See Also

```
smooth.construct.pco.smooth.spec
```

pcre

pffr-constructor for functional principal component-based functional random intercepts.

# **Description**

pffr-constructor for functional principal component-based functional random intercepts.

### Usage

```
pcre(id, efunctions, evalues, yind, ...)
```

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## Arguments

id grouping variable a factor

efunctions matrix of eigenfunction evaluations on gridpoints yind (<length of yind> x <no. of used eigenfunctions>)

evalues eigenvalues associated with efunctions

yind vector of gridpoints on which efunctions are evaluated.

... not used

#### Value

a list used internally for constructing an appropriate call to mgcv::gam

#### **Details**

Fits functional random intercepts  $B_i(t)$  for a grouping variable id using as a basis the functions  $\phi_m(t)$  in efunctions with variances  $\lambda_m$  in evalues:  $B_i(t) \approx \sum_m^M \phi_m(t) \delta_{im}$  with independent  $\delta_{im} \sim N(0,\sigma^2\lambda_m)$ , where  $\sigma^2$  is (usually) estimated and controls the overall contribution of the  $B_i(t)$  while the relative importance of the M basisfunctions is controlled by the supplied variances lambda\_m. Can be used to model smooth residuals if id is simply an index of observations. Differing from scalar random effects in mgcv, these effects are estimated under a "sum-to-zero-foreach-t"-constraint – specifically  $\sum_i \hat{b}_i(t) = 0$  (not  $\sum_i n_i \hat{b}_i(t) = 0$ ) where \$n\_i\$ is the number of observed curves for subject i, so the intercept curve for models with unbalanced group sizes no longer corresponds to the global mean function.

efunctions and evalues are typically eigenfunctions and eigenvalues of an estimated covariance operator for the functional process to be modeled, i.e., they are a functional principal components basis.

#### Author(s)

Fabian Scheipl

# Examples

```
## Not run:
residualfunction <- function(t){
#generate quintic polynomial error functions
          drop(poly(t, 5)%*%rnorm(5, sd=sqrt(2:6)))
}
# generate data Y(t) = mu(t) + E(t) + white noise
set.seed(1122)
n <- 50
T <- 30
t <- seq(0,1, l=T)
# E(t): smooth residual functions
E <- t(replicate(n, residualfunction(t)))
int <- matrix(scale(3*dnorm(t, m=.5, sd=.5) - dbeta(t, 5, 2)), byrow=T, n, T)
Y <- int + E + matrix(.2*rnorm(n*T), n, T)
data <- data.frame(Y=I(Y))</pre>
```

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```
# fit model under independence assumption:
summary(m0 \leftarrow pffr(Y \sim 1, yind=t, data=data))
# get first 5 eigenfunctions of residual covariance
# (i.e. first 5 functional PCs of empirical residual process)
Ehat <- resid(m0)</pre>
fpcE <- fpca.sc(Ehat, npc=5)</pre>
efunctions <- fpcE$efunctions
evalues <- fpcE$evalues</pre>
data$id <- factor(1:nrow(data))</pre>
# refit model with fpc-based residuals
m1 <- pffr(Y ~ 1 + pcre(id=id, efunctions=efunctions, evalues=evalues, yind=t), yind=t, data=data)
t1 <- predict(m1, type="terms")</pre>
summary(m1)
#compare squared errors
mean((int-fitted(m0))^2)
mean((int-t1[[1]])^2)
mean((E-t1[[2]])^2)
# compare fitted & true smooth residuals and fitted intercept functions:
layout(t(matrix(1:4,2,2)))
matplot(t(E), lty=1, type="l", ylim=range(E, t1[[2]]))
matplot(t(t1[[2]]), lty=1, type="l", ylim=range(E, t1[[2]]))
plot(m1, select=1, main="m1", ylim=range(Y))
lines(t, int[1,], col=rgb(1,0,0,.5))
plot(m0, select=1, main="m0", ylim=range(Y))
lines(t, int[1,], col=rgb(1,0,0,.5))
## End(Not run)
```

peer

Construct a PEER regression term in a pfr formula

### **Description**

Defines a term  $\int_T \beta(t) X_i(t) dt$  for inclusion in a pfr formula, where  $\beta(t)$  is estimated with structured penalties (Randolph et al., 2012).

# Usage

```
peer(
   X,
   argvals = NULL,
   pentype = "RIDGE",
   Q = NULL,
   phia = 10^3,
   L = NULL,
   ...
)
```

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#### **Arguments**

argvals

pentype

. . .

functional predictors, typically expressed as an N by J matrix, where N is the number of columns and J is the number of evaluation points. May include missing/sparse functions, which are indicated by NA values. Alternatively, can be an object of class "fd"; see fd.

indices of evaluation of X, i.e.  $(t_{i1},..,t_{iJ})$  for subject i. May be entered as either a length-J vector, or as an N by J matrix. Indices may be unequally spaced. Entering as a matrix allows for different observations times for each subject. If NULL, defaults to an equally-spaced grid between 0 or 1 (or within X\$basis\$rangeval if X is a fd object.)

the type of penalty to apply, one of "RIDGE", "D", "DECOMP", or "USER"; see

Details.

Q matrix Q used for pentype="DECOMP"; see Details.

phia scalar a used for pentype="DECOMP"; see Details.

L user-supplied penalty matrix for pentype="USER"; see Details.

additional arguments to be passed to 1f (and then possibly s). Arguments processed by 1f include, for example, integration for specifying the method of numerical integration. Arguments processed by s include information related to basis and penalization, such as m for specifying the order of the difference penalty; See Details. xt-argument is not allowed for peer-terms and will cause an error.

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#### **Details**

peer is a wrapper for 1f, which defines linear functional predictors for any type of basis. It simply calls 1f with the appropriate options for the peer basis and penalty construction. The type of penalty is determined by the pentype argument. There are four types of penalties available:

- 1. pentype=="RIDGE" for a ridge penalty, the default
- 2. pentype=="D" for a difference penalty. The order of the difference penalty may be specified by supplying an m argument (default is 2).
- 3. pentype=="DECOMP" for a decomposition-based penalty,  $bP_Q + a(I P_Q)$ , where  $P_Q = Q^t(QQ^t)^{-1}Q$ . The Q matrix must be specified by Q, and the scalar a by phia. The number of columns of Q must be equal to the length of the data. Each row represents a basis function where the functional predictor is expected to lie, according to prior belief.
- 4. pentype=="USER" for a user-specified penalty matrix, supplied by the L argument.

The original stand-alone implementation by Madan Gopal Kundu is available in peer\_old.

#### Author(s)

Jonathan Gellar < JGellar@mathematica-mpr.com> and Madan Gopal Kundu < mgkundu@iupui.edu>

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### References

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323-353.

Kundu, M. G., Harezlak, J., and Randolph, T. W. (2012). Longitudinal functional models with structured penalties (arXiv:1211.4763 [stat.AP]).

#### See Also

```
pfr, smooth.construct.peer.smooth.spec
```

## **Examples**

```
# Example 1: Estimation with D2 penalty
#-----
data(DTI)
DTI = DTI[which(DTI$case == 1),]
fit.D2 = pfr(pasat ~ peer(cca, pentype="D"), data=DTI)
plot(fit.D2)
#-----
# Example 2: Estimation with structured penalty (need structural
         information about regression function or predictor function)
data(PEER.Sim)
data(Q)
PEER.Sim1<- subset(PEER.Sim, t==0)
# Setting k to max possible value
fit.decomp <- pfr(Y ~ peer(W, pentype="Decomp", Q=Q, k=99), data=PEER.Sim1)</pre>
plot(fit.decomp)
## End(Not run)
```

PEER.Sim

Simulated longitudinal data with functional predictor and scalar response, and structural information associated with predictor function

# Description

PEER.Sim contains simulated observations from 100 subjects, each observed at 4 distinct time-points. At each timepoint bumpy predictor profile is generated randomly and the scalar response variable is generated considering a time-varying regression function and subject intercept. Accompanying the functional predictor and scalar response are the subject ID numbers and time of measurements.

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### **Format**

The data frame PEER.Sim is made up of subject ID number(id), subject-specific time of measurement (t), functional predictor profile (W.1-W.100) and scalar response (Y)

## **Details**

Q represents the  $7 \times 100$  matrix where each row provides structural information about the functional predictor profile for data PEER. Sim. For specific details about the simulation and Q matrix, please refer to Kundu et. al. (2012).

#### References

Kundu, M. G., Harezlak, J., and Randolph, T. W. (2012). Longitudinal functional models with structured penalties. (please contact J. Harezlak at <harezlak@iupui.edu>)

peer\_old

Functional Models with Structured Penalties

# Description

Implements functional model with structured penalties (Randolph et al., 2012) with scalar outcome and single functional predictor through mixed model equivalence.

### Usage

```
peer_old(
   Y,
   funcs,
   argvals = NULL,
   pentype = "Ridge",
   L.user = NULL,
   Q = NULL,
   phia = 10^3,
   se = FALSE,
   ...
)
```

## **Arguments**

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pentype	type of penalty. It can be either decomposition based penalty (DECOMP) or ridge (RIDGE) or second-order difference penalty (D2) or any user defined penalty (USER). For decomposition based penalty user need to specify Q matrix in Q argument (see details). For user defined penalty user need to specify L matrix in L argument (see details). For Ridge and second-order difference penalty, specification for arguments L and Q will be ignored. Default is RIDGE.
L.user	penalty matrix. Need to be specified with pentype='USER'. Number of columns need to be equal with number of columns of matrix specified to funcs. Each row represents a constraint on functional predictor. This argument will be ignored when value of pentype is other than USER.
Q	Q matrix to derive decomposition based penalty. Need to be specified with pentype='DECOMP'. Number of columns need to be equal with number of columns of matrix specified to funcs. Each row represents a basis function where functional predictor is expected lie according to prior belief. This argument will be ignored when value of pentype is other than DECOMP.
phia	Scalar value of a in decomposition based penalty. Need to be specified with pentype='DECOMP'.
se	logical; calculate standard error when TRUE.
• • •	additional arguments passed to the 1me function.

### **Details**

If there are any missing or infinite values in Y, and funcs, the corresponding row (or observation) will be dropped. Neither Q nor L may contain missing or infinite values.

peer\_old() fits the following model:

$$y_i = \int W_i(s)\gamma(s)ds + \epsilon_i$$

where  $\epsilon_i N(0, \sigma^2)$ . For all the observations, predictor function  $W_i(s)$  is evaluated at K sampling points. Here,  $\gamma(s)$  denotes the regression function.

Values of  $y_i$  and  $W_i(s)$  are passed through argument Y and funcs, respectively. Number of elements or rows in Y and funcs need to be equal.

The estimate of regression functions  $\gamma(s)$  is obtained as penalized estimated. Following 3 types of penalties can be used:

i. Ridge:  $I_K$ 

ii. Second-order difference:  $[d_{i,j}]$  with  $d_{i,i} = d_{i,i+2} = 1, d_{i,i+1} = -2$ , otherwise  $d_{i,i} = 0$ 

iii. Decomposition based penalty:  $bP_Q + a(I - P_Q)$  where  $P_Q = Q^T(QQ^T)^{-1}Q$ 

For Decomposition based penalty user need to specify pentype='DECOMP' and associated Q matrix need to be passed through Q argument.

Alternatively, user can pass directly penalty matrix through argument L. For this user need to specify pentype='USER' and associated L matrix need to be passed through L argument.

Default penalty is Ridge penalty and user needs to specify RIDGE. For second-order difference penalty, user needs to specify D2.

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#### Value

a list containing:

fit result of the call to lme fitted.vals predicted outcomes

Gamma estimates with standard error for regression function

GammaHat estimates of regression function

se.Gamma standard error associated with GammaHat

AIC AIC value of fit (smaller is better)

BIC BIC value of fit (smaller is better)

logLik (restricted) log-likelihood at convergence

lambda estimates of smoothing parameter

N number of subjects

K number of Sampling points in functional predictor sigma estimated within-group error standard deviation.

### Author(s)

Madan Gopal Kundu <mgkundu@iupui.edu>

### References

Kundu, M. G., Harezlak, J., and Randolph, T. W. (2012). Longitudinal functional models with structured penalties (arXiv:1211.4763 [stat.AP]).

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323–353.

## See Also

```
lpeer, plot.peer
```

```
## Not run:
#------
# Example 1: Estimation with D2 penalty
#-------
## Load Data
data(DTI)

## Extract values for arguments for peer() from given data
cca = DTI$cca[which(DTI$case == 1),]
DTI = DTI[which(DTI$case == 1),]

##1.1 Fit the model
fit.cca.peer1 = peer(Y=DTI$pasat, funcs = cca, pentype='D2', se=TRUE)
```

pffr

```
plot(fit.cca.peer1)

#-----
# Example 2: Estimation with structured penalty (need structural
# information about regression function or predictor function)
#------

## Load Data
data(PEER.Sim)

## Extract values for arguments for peer() from given data
PEER.Sim1<- subset(PEER.Sim, t==0)
W<- PEER.Sim1$W
Y<- PEER.Sim1$Y

##Load Q matrix containing structural information
data(Q)

##2.1 Fit the model
Fit1<- peer(Y=Y, funcs=W, pentype='Decomp', Q=Q, se=TRUE)
plot(Fit1)

## End(Not run)
```

pffr

Penalized flexible functional regression

## **Description**

Implements additive regression for functional and scalar covariates and functional responses. This function is a wrapper for mgcv's gam and its siblings to fit models of the general form  $E(Y_i(t)) = g(\mu(t) + \int X_i(s)\beta(s,t)ds + f(z_{1i},t) + f(z_{2i}) + z_{3i}\beta_3(t) + \dots)$  with a functional (but not necessarily continuous) response Y(t), response function g, (optional) smooth intercept  $\mu(t)$ , (multiple) functional covariates X(t) and scalar covariates  $z_1, z_2$ , etc.

## Usage

```
pffr(
  formula,
  yind,
  data = NULL,
  ydata = NULL,
  algorithm = NA,
  method = "REML",
  tensortype = c("ti", "t2"),
  bs.yindex = list(bs = "ps", k = 5, m = c(2, 1)),
  bs.int = list(bs = "ps", k = 20, m = c(2, 1)),
  ...
)
```

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### **Arguments**

formula	a formula with special terms as for gam, with additional special terms ff(), sff(),ffpc(), pcre() and c().
yind	a vector with length equal to the number of columns of the matrix of functional responses giving the vector of evaluation points $(t_1, \ldots, t_G)$ . If not supplied, yind is set to 1:ncol( <response>).</response>
data	an (optional) data.frame containing the data. Can also be a named list for regular data. Functional covariates have to be supplied as <no. observations="" of=""> by <no. evaluations="" of=""> matrices, i.e. each row is one functional observation.</no.></no.>
ydata	an (optional) data.frame supplying functional responses that are not observed on a regular grid. See Details.
algorithm	the name of the function used to estimate the model. Defaults to gam if the matrix of functional responses has less than 2e5 data points and to bam if not. 'gamm', 'gamm4' and 'jagam' are valid options as well. See Details for 'gamm4' and 'jagam'.
method	Defaults to "REML"-estimation, including of unknown scale. If algorithm="bam", the default is switched to "fREML". See gam and bam for details.
tensortype	which typ of tensor product splines to use. One of "ti" or "t2", defaults to ti. t2-type terms do not enforce the more suitable special constraints for functional regression, see Details.
bs.yindex	a named (!) list giving the parameters for spline bases on the index of the functional response. Defaults to list(bs="ps", k=5,m=c(2, 1)), i.e. 5 cubic B-splines bases with first order difference penalty.
bs.int	a named (!) list giving the parameters for the spline basis for the global functional intercept. Defaults to list(bs="ps", k=20,m=c(2, 1)), i.e. 20 cubic B-splines bases with first order difference penalty.
	additional arguments that are valid for gam, bam, 'gamm4' or 'jagam'. subset is not implemented.

### Value

A fitted pffr-object, which is a gam-object with some additional information in an pffr-entry. If algorithm is "gamm" or "gamm4", only the \$gam part of the returned list is modified in this way. Available methods/functions to postprocess fitted models: summary.pffr, plot.pffr, coef.pffr, fitted.pffr, residuals.pffr, predict.pffr, model.matrix.pffr, qq.pffr, pffr.check. If algorithm is "jagam", only the location of the model file and the usual jagam-object are returned, you have to run the sampler yourself.

## **Details**

The routine can estimate

1. linear functional effects of scalar (numeric or factor) covariates that vary smoothly over t (e.g.  $z_{1i}\beta_1(t)$ , specified as ~z1),

2. nonlinear, and possibly multivariate functional effects of (one or multiple) scalar covariates z that vary smoothly over the index t of Y(t) (e.g.  $f(z_{2i},t)$ , specified in the formula simply as  $\sim$ s(z2))

- 3. (nonlinear) effects of scalar covariates that are constant over t (e.g.  $f(z_{3i})$ , specified as  $\sim c(s(z_3))$ , or  $\beta_3 z_{3i}$ , specified as  $\sim c(z_3)$ ),
- 4. function-on-function regression terms (e.g.  $\int X_i(s)\beta(s,t)ds$ , specified as ~ff(X, yindex=t, xindex=s), see ff). Terms given by sff and ffpc provide nonlinear and FPC-based effects of functional covariates, respectively.
- 5. concurrent effects of functional covariates X measured on the same grid as the response are specified as follows:  $\sim s(x)$  for a smooth, index-varying effect f(X(t),t),  $\sim x$  for a linear index-varying effect  $X(t)\beta(t)$ ,  $\sim c(s(x))$  for a constant nonlinear effect f(X(t)),  $\sim c(x)$  for a constant linear effect  $X(t)\beta$ .
- 6. Smooth functional random intercepts  $b_{0g(i)}(t)$  for a grouping variable g with levels g(i) can be specified via  $\sim$ s(g, bs="re")), functional random slopes  $u_i b_{1g(i)}(t)$  in a numeric variable u via  $\sim$ s(g, u, bs="re")). Scheipl, Staicu, Greven (2013) contains code examples for modeling correlated functional random intercepts using mrf-terms.

Use the c()-notation to denote model terms that are constant over the index of the functional response.

Internally, univariate smooth terms without a c()-wrapper are expanded into bivariate smooth terms in the original covariate and the index of the functional response. Bivariate smooth terms (s(), te() or t2()) without a c()-wrapper are expanded into trivariate smooth terms in the original covariates and the index of the functional response. Linear terms for scalar covariates or categorical covariates are expanded into varying coefficient terms, varying smoothly over the index of the functional response. For factor variables, a separate smooth function with its own smoothing parameter is estimated for each level of the factor.

The marginal spline basis used for the index of the functional response is specified via the global argument bs.yindex. If necessary, this can be overriden for any specific term by supplying a bs.yindex-argument to that term in the formula, e.g.  $\sim s(x, bs.yindex=list(bs="tp", k=7))$  would yield a tensor product spline over x and the index of the response in which the marginal basis for the index of the response are 7 cubic thin-plate spline functions (overriding the global default for the basis and penalty on the index of the response given by the global bs.yindex-argument). Use  $\sim -1 + c(1) + \ldots$  to specify a model with only a constant and no functional intercept.

The functional covariates have to be supplied as a n by <no. of evaluations> matrices, i.e. each row is one functional observation. For data on a regular grid, the functional response is supplied in the same format, i.e. as a matrix-valued entry in data, which can contain missing values.

If the functional responses are *sparse or irregular* (i.e., not evaluated on the same evaluation points across all observations), the ydata-argument can be used to specify the responses: ydata must be a data.frame with 3 columns called '.obs', '.index', '.value' which specify which curve the point belongs to ('.obs'=i), at which t it was observed ('.index'=t), and the observed value ('.value'= $Y_i(t)$ ). Note that the vector of unique sorted entries in ydata\$.obs must be equal to rownames (data) to ensure the correct association of entries in ydata to the corresponding rows of data. For both regular and irregular functional responses, the model is then fitted with the data in

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long format, i.e., for data on a grid the rows of the matrix of the functional response evaluations  $Y_i(t)$  are stacked into one long vector and the covariates are expanded/repeated correspondingly. This means the models get quite big fairly fast, since the effective number of rows in the design matrix is number of observations times number of evaluations of Y(t) per observation.

Note that pffr does not use mgcv's default identifiability constraints (i.e.,  $\sum_{i,t} \hat{f}(z_i,x_i,t) = 0$  or  $\sum_{i,t} \hat{f}(x_i,t) = 0$ ) for tensor product terms whose marginals include the index t of the functional response. Instead,  $\sum_i \hat{f}(z_i,x_i,t) = 0$  for all t is enforced, so that effects varying over t can be interpreted as local deviations from the global functional intercept. This is achieved by using titerms with a suitably modified mc-argument. Note that this is not possible if algorithm='gamm4' since only t2-type terms can then be used and these modified constraints are not available for t2. We recommend using centered scalar covariates for terms like  $z\beta(t)$  (~z) and centered functional covariates with  $\sum_i X_i(t) = 0$  for all t in ff-terms so that the global functional intercept can be interpreted as the global mean function.

The family-argument can be used to specify all of the response distributions and link functions described in family.mgcv. Note that family = "gaulss" is treated in a special way: Users can supply the formula for the variance by supplying a special argument varformula, but this is not modified in the way that the formula-argument is but handed over to the fitter directly, so this is for expert use only. If varformula is not given, pffr will use the parameters from argument bs.int to define a spline basis along the index of the response, i.e., a smooth variance function over t for responses Y(t).

#### Author(s)

Fabian Scheipl, Sonja Greven

#### References

Ivanescu, A., Staicu, A.-M., Scheipl, F. and Greven, S. (2015). Penalized function-on-function regression. Computational Statistics, 30(2):539–568. https://biostats.bepress.com/jhubiostat/paper254/

Scheipl, F., Staicu, A.-M. and Greven, S. (2015). Functional Additive Mixed Models. Journal of Computational & Graphical Statistics, 24(2): 477–501. https://arxiv.org/abs/1207.5947

F. Scheipl, J. Gertheiss, S. Greven (2016): Generalized Functional Additive Mixed Models, Electronic Journal of Statistics, 10(1), 1455–1492. https://projecteuclid.org/journals/electronic-journal-of-stativolume-10/issue-1/Generalized-functional-additive-mixed-models/10.1214/16-EJS1145. full

### See Also

smooth.terms for details of mgcv syntax and available spline bases and penalties.

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```
data1 <- pffrSim(scenario="ff", n=40)</pre>
t <- attr(data1, "yindex")
s <- attr(data1, "xindex")</pre>
m1 <- pffr(Y ~ ff(X1, xind=s), yind=t, data=data1)
summary(m1)
plot(m1, pages=1)
## Not run:
# multivariate model:
\# E(Y(t)) = \beta(t) + \inf X1(s)\beta(s+1)ds + x lin \beta(t) +
        f_1(xte1, xte2) + f_2(xsmoo, t) + \beta_4(xsmoo, t)
data2 <- pffrSim(scenario="all", n=200)</pre>
t <- attr(data2, "yindex")
s <- attr(data2, "xindex")</pre>
m2 \leftarrow pffr(Y \sim ff(X1, xind=s) + \#linear function-on-function
               xlin + #varying coefficient term
            c(te(xte1, xte2)) + #bivariate smooth term in xte1 & xte2, const. over Y-index
               s(xsmoo) + #smooth effect of xsmoo varying over Y-index
               c(xconst), # linear effect of xconst constant over Y-index
       yind=t,
       data=data2)
summary(m2)
plot(m2)
str(coef(m2))
# convenience functions:
preddata <- pffrSim(scenario="all", n=20)</pre>
str(predict(m2, newdata=preddata))
str(predict(m2, type="terms"))
cm2 <- coef(m2)
cm2$pterms
str(cm2$smterms, 2)
str(cm2$smterms[["s(xsmoo)"]]$coef)
# sparse data (80% missing on a regular grid):
set.seed(88182004)
data3 <- pffrSim(scenario=c("int", "smoo"), n=100, propmissing=0.8)</pre>
t <- attr(data3, "yindex")
m3.sparse <- pffr(Y ~ s(xsmoo), data=data3$data, ydata=data3$ydata, yind=t)</pre>
summary(m3.sparse)
plot(m3.sparse,pages=1)
## End(Not run)
```

pffr.check

Some diagnostics for a fitted pffr model

### **Description**

This is simply a wrapper for gam.check().

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### Usage

```
pffr.check(
   b,
   old.style = FALSE,
   type = c("deviance", "pearson", "response"),
   k.sample = 5000,
   k.rep = 200,
   rep = 0,
   level = 0.9,
   rl.col = 2,
   rep.col = "gray80",
   ...
)
```

### Arguments

```
b
                  a fitted pffr-object
old.style
                  If you want old fashioned plots, exactly as in Wood, 2006, set to TRUE.
type
                  type of residuals, see residuals.gam, used in all plots.
k.sample
                  Above this k testing uses a random sub-sample of data.
                  how many re-shuffles to do to get p-value for k testing.
k.rep
                  passed to qq.gam when old.style is FALSE.
rep
level
                  passed to qq.gam when old.style is FALSE.
                  passed to qq.gam when old.style is FALSE.
rl.col
rep.col
                  passed to qq.gam when old.style is FALSE.
                  extra graphics parameters to pass to plotting functions.
```

pffrGLS

Penalized function-on-function regression with non-i.i.d. residuals

### **Description**

Implements additive regression for functional and scalar covariates and functional responses. This function is a wrapper for mgcv's gam and its siblings to fit models of the general form  $Y_i(t) = \mu(t) + \int X_i(s)\beta(s,t)ds + f(z_{1i},t) + f(z_{2i}) + z_{3i}\beta_3(t) + \dots + E_i(t))$  with a functional (but not necessarily continuous) response Y(t), (optional) smooth intercept  $\mu(t)$ , (multiple) functional covariates X(t) and scalar covariates  $z_1, z_2$ , etc. The residual functions  $E_i(t) \sim GP(0,K(t,t'))$  are assumed to be i.i.d. realizations of a Gaussian process. An estimate of the covariance operator K(t,t') evaluated on yind has to be supplied in the hatSigma-argument.

pffrGLS

## Usage

```
pffrGLS(
   formula,
   yind,
   hatSigma,
   algorithm = NA,
   method = "REML",
   tensortype = c("te", "t2"),
   bs.yindex = list(bs = "ps", k = 5, m = c(2, 1)),
   bs.int = list(bs = "ps", k = 20, m = c(2, 1)),
   cond.cutoff = 500,
   ...
)
```

## **Arguments**

formula	a formula with special terms as for gam, with additional special terms $ff()$ and $c()$ . See $pffr$ .
yind	a vector with length equal to the number of columns of the matrix of functional responses giving the vector of evaluation points $(t_1, \ldots, t_G)$ . see pffr
hatSigma	(an estimate of) the within-observation covariance (along the responses' index), evaluated at yind. See Details.
algorithm	the name of the function used to estimate the model. Defaults to gam if the matrix of functional responses has less than 2e5 data points and to bam if not. "gamm" (see gamm) and "gamm4" (see gamm4) are valid options as well.
method	See pffr
tensortype	See pffr
bs.yindex	See pffr
bs.int	See pffr
cond.cutoff	if the condition number of hatSigma is greater than this, hatSigma is made "more" positive-definite via nearPD to ensure a condition number equal to cond.cutoff. Defaults to 500.
	additional arguments that are valid for gam or bam. See pffr.

## Value

```
a fitted pffr-object, see pffr.
```

### **Details**

Note that hatSigma has to be positive definite. If hatSigma is close to positive *semi*-definite or badly conditioned, estimated standard errors become unstable (typically much too small). pffrGLS will try to diagnose this and issue a warning. The danger is especially big if the number of functional observations is smaller than the number of gridpoints (i.e, length(yind)), since the raw covariance estimate will not have full rank.

Please see pffr for details on model specification and implementation.

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THIS IS AN EXPERIMENTAL VERSION AND NOT WELL TESTED YET – USE AT YOUR OWN RISK.

#### Author(s)

Fabian Scheipl

#### See Also

```
pffr, fpca.sc
```

pffrSim

Simulate example data for pffr

### Description

Simulates example data for pffr from a variety of terms. Scenario "all" generates data from a complex multivariate model

$$Y_i(t) = \mu(t) + \int X_{1i}(s)\beta_1(s,t)ds + xlin\beta_3(t) + f(xte1,xte2) + f(xsmoo,t) + \beta_4xconst + f(xfactor,t) + \epsilon_i(t)$$

. Scenarios "int", "ff", "lin", "te", "smoo", "const", "factor", generate data from simpler models containing only the respective term(s) in the model equation given above. Specifying a vector-valued scenario will generate data from a combination of the respective terms. Sparse/irregular response trajectories can be generated by setting propmissing to something greater than 0 (and smaller than 1). The return object then also includes a ydata-item with the sparsified data.

### Usage

```
pffrSim(
    scenario = "all",
    n = 100,
    nxgrid = 40,
    nygrid = 60,
    SNR = 10,
    propmissing = 0,
    limits = NULL
)
```

### **Arguments**

scenario see Description

n number of observations

nxgrid number of evaluation points of functional covariates

nygrid number of evaluation points of the functional response

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SNR the signal-to-noise ratio for the generated data: empirical variance of the additive

predictor divided by variance of the errors.

propmissing proportion of missing data in the response, default = 0. See Details.

limits a function that defines an integration range, see ff

#### **Details**

See source code for details.

### Value

a named list with the simulated data, and the true components of the predictor etc as attributes.

pfr

Penalized Functional Regression

# Description

Implements various approaches to penalized scalar-on-function regression. These techniques include Penalized Functional Regression (Goldsmith et al., 2011), Longitudinal Penalized Functional Regression (Goldsmith, et al., 2012), Functional Principal Component Regression (Reiss and Ogden, 2007), Partially Empirical Eigenvectors for Regression (Randolph et al., 2012), Functional Generalized Additive Models (McLean et al., 2013), and Variable-Domain Functional Regression (Gellar et al., 2014). This function is a wrapper for mgcv's gam and its siblings to fit models with a scalar (but not necessarily continuous) response.

### Usage

```
pfr(formula = NULL, fitter = NA, method = "REML", ...)
```

control the estimation.

### **Arguments**

formula	a formula that could contain any of the following special terms: lf(), af(), lf.vd(), peer(), fpc(), or re(); also mgcv's s(), te(), or t2().
fitter	the name of the function used to estimate the model. Defaults to gam if the matrix of functional responses has less than 2e5 data points and to bam if not. "gamm" (see gamm) and "gamm4" (see gamm4) are valid options as well.
method	The smoothing parameter estimation method. Default is "REML". For options, see gam.
	additional arguments that are valid for gam or bam. These include data and family to specify the input data and outcome family, as well as many options to

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#### Value

A fitted pfr-object, which is a gam-object with some additional information in a \$pfr-element. If fitter is "gamm" or "gamm4", only the \$gam part of the returned list is modified in this way.

### Warning

Binomial responses should be specified as a numeric vector rather than as a matrix or a factor.

#### Author(s)

Jonathan Gellar <JGellar@mathematica-mpr.com>, Mathew W. McLean, Jeff Goldsmith, and Fabian Scheipl

#### References

Goldsmith, J., Bobb, J., Crainiceanu, C., Caffo, B., and Reich, D. (2011). Penalized functional regression. *Journal of Computational and Graphical Statistics*, 20(4), 830-851.

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2012). Longitudinal penalized functional regression for cognitive outcomes on neuronal tract measurements. *Journal of the Royal Statistical Society: Series C*, 61(3), 453-469.

Reiss, P. T., and Ogden, R. T. (2007). Functional principal component regression and functional partial least squares. *Journal of the American Statistical Association*, 102, 984-996.

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323-353.

McLean, M. W., Hooker, G., Staicu, A.-M., Scheipl, F., and Ruppert, D. (2014). Functional generalized additive models. *Journal of Computational and Graphical Statistics*, **23** (1), pp. 249-269. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3982924/.

Gellar, J. E., Colantuoni, E., Needham, D. M., and Crainiceanu, C. M. (2014). Variable-Domain Functional Regression for Modeling ICU Data. Journal of the American Statistical Association, 109(508): 1425-1439.

#### See Also

```
af, lf, lf.vd, fpc, peer, re.
```

```
# See lf(), lf.vd(), af(), fpc(), and peer() for additional examples
data(DTI)
DTI1 <- DTI[DTI$visit==1 & complete.cases(DTI),]
par(mfrow=c(1,2))

# Fit model with linear functional term for CCA
fit.lf <- pfr(pasat ~ lf(cca, k=30, bs="ps"), data=DTI1)
plot(fit.lf, ylab=expression(paste(beta(t))), xlab="t")
## Not run:
# Alternative way to plot</pre>
```

```
bhat.lf <- coef(fit.lf, n=101)</pre>
bhat.lf$upper <- bhat.lf$value + 1.96*bhat.lf$se</pre>
bhat.lf$lower <- bhat.lf$value - 1.96*bhat.lf$se
matplot(bhat.lf$cca.argvals, bhat.lf[,c("value", "upper", "lower")],
        type="1", lty=c(1,2,2), col=1,
        ylab=expression(paste(beta(t))), xlab="t")
# Fit model with additive functional term for CCA, using tensor product basis
fit.af <- pfr(pasat ~ af(cca, Qtransform=TRUE, k=c(7,7)), data=DTI1)</pre>
plot(fit.af, scheme=2, xlab="t", ylab="cca(t)", main="Tensor Product")
plot(fit.af, scheme=2, Qtransform=TRUE,
     xlab="t", ylab="cca(t)", main="Tensor Product")
# Change basistype to thin-plate regression splines
fit.af.s <- pfr(pasat ~ af(cca, basistype="s", Qtransform=TRUE, k=50),</pre>
                 data=DTI1)
plot(fit.af.s, scheme=2, xlab="t", ylab="cca(t)", main="TPRS", rug=FALSE)
plot(fit.af.s, scheme=2, Qtransform=TRUE,
     xlab="t", ylab="cca(t)", main="TPRS", rug=FALSE)
# Visualize bivariate function at various values of x
par(mfrow=c(2,2))
vis.pfr(fit.af, xval=.2)
vis.pfr(fit.af, xval=.4)
vis.pfr(fit.af, xval=.6)
vis.pfr(fit.af, xval=.8)
# Include random intercept for subject
DTI.re <- DTI[complete.cases(DTI$cca),]</pre>
DTI.re$ID <- factor(DTI.re$ID)</pre>
fit.re <- pfr(pasat ~ lf(cca, k=30) + re(ID), data=DTI.re)</pre>
coef.re <- coef(fit.re)</pre>
par(mfrow=c(1,2))
plot(fit.re)
# FPCR R Model
fit.fpc <- pfr(pasat ~ fpc(cca), data=DTI.re)</pre>
plot(fit.fpc)
# PEER Model with second order difference penalty
DTI.use <- DTI[DTI$case==1,]</pre>
DTI.use <- DTI.use[complete.cases(DTI.use$cca),]</pre>
fit.peer <- pfr(pasat ~ peer(cca, argvals=seq(0,1,length=93),</pre>
                              integration="riemann", pentype="D"), data=DTI.use)
plot(fit.peer)
## End(Not run)
```

#### **Description**

This code implements the function pfr() available in refund 0.1-11. It is included to maintain backwards compatibility.

Functional predictors are entered as a matrix or, in the case of multiple functional predictors, as a list of matrices using the funcs argument. Missing values are allowed in the functional predictors, but it is assumed that they are observed over the same grid. Functional coefficients and confidence bounds are returned as lists in the same order as provided in the funcs argument, as are principal component and spline bases. Increasing values of nbasis will increase computational time and the values of nbasis, kz, and kb in relation to each other may need to be adjusted in application-specific ways.

## Usage

```
pfr_old(
    Y,
    subj = NULL,
    covariates = NULL,
    funcs,
    kz = 10,
    kb = 30,
    nbasis = 10,
    family = "gaussian",
    method = "REML",
    smooth.option = "fpca.sc",
    pve = 0.99,
    ...
)
```

# Arguments Y

subj	vector containing the subject number for each observation
covariates	matrix of scalar covariates
funcs	matrix, or list of matrices, containing observed functional predictor

vector of all outcomes over all visits

Funcs matrix, or list of matrices, containing observed functional predictors as rows.

NA values are allowed.

kz can be NULL; can be a scalar, in which case this will be the dimension of principal components basis for each and every observed functional predictors; can

be a vector of length equal to the number of functional predictors, in which case each element will correspond to the dimension of principal components basis

for the corresponding observed functional predictors

kb dimension of the B-spline basis for the coefficient function (note: this is a

change from versions 0.1-7 and previous)

nbasis passed to refund::fpca.sc (note: using fpca.sc is a change from versions 0.1-7

and previous)

family generalized linear model family

method method for estimating the smoothing parameters; defaults to REML

smooth.option method to do FPC decomposition on the predictors. Two options available –

"fpca.sc" or "fpca.face". If using "fpca.sc", a number less than 35 for nbasis

should be used while if using "fpca.face",35 or more is recommended.

pve proportion of variance explained used to choose the number of principal com-

ponents to be included in the expansion.

... additional arguments passed to gam to fit the regression model.

### Value

fit result of the call to gam

fitted.vals predicted outcomes

fitted.vals.level.0

predicted outcomes at population level

fitted.vals.level.1

predicted outcomes at subject-specific level (if applicable)

betaHat list of estimated coefficient functions

beta.covariates

parameter estimates for scalar covariates

varBetaHat list containing covariance matrices for the estimated coefficient functions

Bounds list of bounds of a pointwise 95% confidence interval for the estimated coeffi-

cient functions

X design matrix used in the model fit
D penalty matrix used in the model fit

phi list of B-spline bases for the coefficient functions

psi list of principal components basis for the functional predictors

C stacked row-specific principal component scores

J transpose of psi matrix multiplied by phi

CJ C matrix multiplied J

Z1 design matrix of random intercepts subj subject identifiers as specified by user

fixed.mat the fixed effects design matrix of the pfr as a mixed model rand.mat the fixed effects design matrix of the pfr as a mixed model

N\_subj the number of unique subjects, if subj is specified

p number of scalar covariatesN. pred number of functional covariates

kz as specified

kz.adj For smooth.option="fpca.sc", will be same as kz (or a vector of repeated values

of the specified scalar kz). For smooth.option="fpca.face", will be the corresponding number of principal components for each functional predictor as determined by fpca.face; will be less than or equal to kz on an elemental-wise

level.

kb as specified nbasis as specified

totD number of penalty matrices created for mgcv::gam

funcs as specified covariates as specified smooth.option as specified

#### Warning

Binomial responses should be specified as a numeric vector rather than as a matrix or a factor.

### Author(s)

Bruce Swihart Swihart com> and Jeff Goldsmith jeff.goldsmith@columbia.edu>

#### References

Goldsmith, J., Bobb, J., Crainiceanu, C., Caffo, B., and Reich, D. (2011). Penalized functional regression. *Journal of Computational and Graphical Statistics*, 20(4), 830-851.

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2012). Longitudinal penalized functional regression for cognitive outcomes on neuronal tract measurements. *Journal of the Royal Statistical Society: Series C*, 61(3), 453-469.

Swihart, Bruce J., Goldsmith, Jeff; and Crainiceanu, Ciprian M. (July 2012). Testing for functional effects. Johns Hopkins University Dept. of Biostatistics Working Paper 247, available at https://biostats.bepress.com/jhubiostat/paper247/ American Statistical Association, 109(508): 1425-1439.

#### See Also

```
rlrt.pfr, predict.pfr.
```

```
## Not run:
#########
            DTI Data Example
                              #########
# For more about this example, see Swihart et al. 2013
## load and reassign the data;
data(DTI2)
Y <- DTI2$pasat ## PASAT outcome
id <- DTI2$id
       ## subject id
W1 <- DTI2$cca ## Corpus Callosum
W2 <- DTI2$rcst ## Right corticospinal
V <- DTI2$visit ## visit
```

```
## prep scalar covariate
visit.1.rest <- matrix(as.numeric(V > 1), ncol=1)
covar.in <- visit.1.rest</pre>
## note there is missingness in the functional predictors
apply(is.na(W1), 2, mean)
apply(is.na(W2), 2, mean)
## fit two univariate models
pfr.obj.t1 <- pfr(Y = Y, covariates=covar.in, funcs = list(W1), subj = id, kz = 10, kb = 50)</pre>
pfr.obj.t2 <- pfr(Y = Y, covariates=covar.in, funcs = list(W2), subj = id, kz = 10, kb = 50)
### one model with two functional predictors using "smooth.face"
### for smoothing predictors
pfr.obj.t3 <- pfr(Y = Y, covariates=covar.in, funcs = list(W1, W2),</pre>
                 subj = id, kz = 10, kb = 50, nbasis=35,smooth.option="fpca.face")
## plot the coefficient function and bounds
dev.new()
par(mfrow=c(2,2))
ran <- c(-2,.5)
matplot(cbind(pfr.obj.t1$BetaHat[[1]], pfr.obj.t1$Bounds[[1]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "CCA", xlab="Location", ylim=ran)
abline(h=0, col="blue")
matplot(cbind(pfr.obj.t2$BetaHat[[1]], pfr.obj.t2$Bounds[[1]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "RCST", xlab="Location", ylim=ran)
abline(h=0, col="blue")
matplot(cbind(pfr.obj.t3$BetaHat[[1]], pfr.obj.t3$Bounds[[1]]),
       type = '1', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "CCA - mult.", xlab="Location", ylim=ran)
abline(h=0, col="blue")
matplot(cbind(pfr.obj.t3$BetaHat[[2]], pfr.obj.t3$Bounds[[2]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "RCST - mult.", xlab="Location", ylim=ran)
abline(h=0, col="blue")
# use baseline data to regress continuous outcomes on functional
# predictors (continuous outcomes only recorded for case == 1)
data(DTI)
# subset data as needed for this example
cca = DTI$cca[which(DTI$visit ==1 & DTI$case == 1),]
rcst = DTI$rcst[which(DTI$visit ==1 & DTI$case == 1),]
DTI = DTI[which(DTI$visit ==1 & DTI$case == 1),]
# note there is missingness in the functional predictors
```

```
apply(is.na(cca), 2, mean)
apply(is.na(rcst), 2, mean)
# fit two models with single functional predictors and plot the results
fit.cca = pfr(Y=DTI$pasat, funcs = cca, kz=10, kb=50, nbasis=20)
fit.rcst = pfr(Y=DTI$pasat, funcs = rcst, kz=10, kb=50, nbasis=20)
par(mfrow = c(1,2))
matplot(cbind(fit.cca$BetaHat[[1]], fit.cca$Bounds[[1]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "CCA")
matplot(cbind(fit.rcst$BetaHat[[1]], fit.rcst$Bounds[[1]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "RCST")
# fit a model with two functional predictors and plot the results
fit.cca.rcst = pfr(Y=DTI$pasat, funcs = list(cca, rcst), kz=10, kb=30, nbasis=20)
par(mfrow = c(1,2))
matplot(cbind(fit.cca.rcst$BetaHat[[1]], fit.cca.rcst$Bounds[[1]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "CCA")
matplot(cbind(fit.cca.rcst$BetaHat[[2]], fit.cca.rcst$Bounds[[2]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "RCST")
# use baseline data to regress binary case-status outcomes on
# functional predictors
data(DTI)
# subset data as needed for this example
cca = DTI$cca[which(DTI$visit == 1),]
rcst = DTI$rcst[which(DTI$visit == 1),]
DTI = DTI[which(DTI$visit == 1),]
# fit two models with single functional predictors and plot the results
fit.cca = pfr(Y=DTI$case, funcs = cca, family = "binomial")
fit.rcst = pfr(Y=DTI$case, funcs = rcst, family = "binomial")
par(mfrow = c(1,2))
matplot(cbind(fit.cca$BetaHat[[1]], fit.cca$Bounds[[1]]),
       type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "CCA")
matplot(cbind(fit.rcst$BetaHat[[1]], fit.rcst$Bounds[[1]]),
       type = '1', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
       main = "RCST")
Octane Data Example
```

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plot.fosr

Default plotting of function-on-scalar regression objects

## **Description**

Plots the coefficient function estimates produced by fosr().

## Usage

```
## S3 method for class 'fosr'
plot(
    x,
    split = NULL,
    titles = NULL,
    xlabel = "",
    ylabel = "Coefficient function",
    set.mfrow = TRUE,
    ...
)
```

# Arguments

X	an object of class "fosr".
split	value, or vector of values, at which to divide the set of coefficient functions into groups, each plotted on a different scale. E.g., if set to 1, the first function is plotted on one scale, and all others on a different (common) scale. If NULL, all functions are plotted on the same scale.
titles	character vector of titles for the plots produced, e.g., names of the corresponding scalar predictors.
xlabel	label for the x-axes of the plots.
ylabel	label for the y-axes of the plots.

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set.mfrow logical value: if TRUE, the function will try to set an appropriate value of the

mfrow parameter for the plots. Otherwise you may wish to set mfrow outside the

function call.

... graphical parameters (see par) for the plot.

### Author(s)

```
Philip Reiss <phil.reiss@nyumc.org>
```

## See Also

fosr, which includes examples.

plot.fosr.vs

Plot for Function-on Scalar Regression with variable selection

## **Description**

Given a "fosr.vs" object, produces a figure of estimated coefficient functions.

# Usage

```
## S3 method for class 'fosr.vs' plot(x, ...)
```

## **Arguments**

```
x an object of class "fosr.vs".
```

... additional arguments.

#### Value

a figure of estimated coefficient functions.

## Author(s)

Yakuan Chen <yc2641@cumc.columbia.edu>

## See Also

```
fosr.vs
```

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### **Examples**

```
## Not run:
I = 100
p = 20
D = 50
grid = seq(0, 1, length = D)
beta.true = matrix(0, p, D)
beta.true[1,] = sin(2*grid*pi)
beta.true[2,] = cos(2*grid*pi)
beta.true[3,] = 2
psi.true = matrix(NA, 2, D)
psi.true[1,] = sin(4*grid*pi)
psi.true[2,] = cos(4*grid*pi)
lambda = c(3,1)
set.seed(100)
X = matrix(rnorm(I*p), I, p)
C = cbind(rnorm(I, mean = 0, sd = lambda[1]), rnorm(I, mean = 0, sd = lambda[2]))
fixef = X%*%beta.true
pcaef = C %*% psi.true
error = matrix(rnorm(I*D), I, D)
Yi.true = fixef
Yi.pca = fixef + pcaef
Yi.obs = fixef + pcaef + error
data = as.data.frame(X)
data$Y = Yi.obs
fit.mcp = fosr.vs(Y~., data = data[1:80,], method="grMCP")
plot(fit.mcp)
## End(Not run)
```

plot.fpcr

Default plotting for functional principal component regression output

# **Description**

Inputs an object created by fpcr, and plots the estimated coefficient function.

## Usage

```
## S3 method for class 'fpcr'
```

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```
plot(
    x,
    se = TRUE,
    col = 1,
    lty = c(1, 2, 2),
    xlab = "",
    ylab = "Coefficient function",
    ...
)
```

# Arguments

X	an object of class "fpcr".
se	if TRUE (the default), upper and lower lines are added at 2 standard errors (in the Bayesian sense; see Wood, 2006) above and below the coefficient function estimate. If a positive number is supplied, the standard error is instead multiplied by this number.
col	color for the line(s). This should be either a number, or a vector of length 3 for the coefficient function estimate, lower bound, and upper bound, respectively.
lty	line type(s) for the coefficient function estimate, lower bound, and upper bound.
xlab, ylab	x- and y-axis labels.
	other arguments passed to the underlying plotting function.

## Value

None; only a plot is produced.

# Author(s)

```
Philip Reiss <phil.reiss@nyumc.org>
```

### References

Wood, S. N. (2006). *Generalized Additive Models: An Introduction with R*. Boca Raton, FL: Chapman & Hall.

### See Also

fpcr, which includes an example.

plot.lpeer

plot.lpeer

Plotting of estimated regression functions obtained through lpeer()

### Description

Plots the estimate of components of estimated regression function obtained from an lpeer object along with pointwise confidence bands.

## Usage

```
## S3 method for class 'lpeer'
plot(x, conf = 0.95, ...)
```

### **Arguments**

```
x object of class "lpeer".

conf pointwise confidence level.

... additional arguments passed to plot.
```

#### **Details**

Pointwise confidence interval is displayed only if the user set se=T in the call to lpeer, and does not reflect any multiplicity correction.

### Author(s)

Madan Gopal Kundu <mgkundu@iupui.edu>

#### References

Kundu, M. G., Harezlak, J., and Randolph, T. W. (2012). Longitudinal functional models with structured penalties. (Please contact J. Harezlak at <a href="mailto:harezlak@iupui.edu">harezlak@iupui.edu</a>).

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323–353.

## See Also

```
peer, lpeer, plot.peer
```

```
## Not run:
data(DTI)
cca = DTI$cca[which(DTI$case == 1),]
DTI = DTI[which(DTI$case == 1),]
fit.cca.lpeer1 = lpeer(Y=DTI$pasat, t=DTI$visit, subj=DTI$ID, funcs = cca)
plot(fit.cca.lpeer1)
## End(Not run)
```

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plot.peer

Plotting of estimated regression functions obtained through peer()

## **Description**

Plots the estimate of components of estimated regression function obtained from a peer object along with pointwise confidence bands.

# Usage

```
## $3 method for class 'peer'
plot(
    x,
    conf = 0.95,
    ylab = "Estimated regression function",
    main = expression(gamma),
    ...
)
```

### **Arguments**

```
x object of class "peer".

conf pointwise confidence level.

ylab y-axis label.

main title for the plot.

... additional arguments passed to plot.
```

#### **Details**

Pointwise confidence interval is displayed only if the user set se=T in the call to peer, and does not reflect any multiplicity correction.

# Author(s)

Madan Gopal Kundu <mgkundu@iupui.edu>

#### References

Kundu, M. G., Harezlak, J., and Randolph, T. W. (2012). Longitudinal functional models with structured penalties. (Please contact J. Harezlak at <a href="mailto:harezlak@iupui.edu">harezlak@iupui.edu</a>.)

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323–353.

### See Also

```
peer, lpeer, plot.lpeer
```

plot.pfr

### **Examples**

```
# See example in peer()
```

plot.pffr

Plot a pffr fit

# Description

Plot a fitted pffr-object. Simply dispatches to plot.gam.

# Usage

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

## **Arguments**

```
x a fitted pffr-object
```

... arguments handed over to plot.gam

## Value

This function only generates plots.

# Author(s)

Fabian Scheipl

plot.pfr

Plot a pfr object

# Description

This function plots the smooth coefficients of a pfr object. These include functional coefficients as well as any smooths of scalar covariates. The function dispatches to pfr\_plot.gam, which is our local copy of plot.gam with some minor changes.

# Usage

```
## S3 method for class 'pfr'
plot(x, Qtransform = FALSE, ...)
```

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## **Arguments**

x a fitted pfr-object

Qtransform For additive functional terms, TRUE indicates the coefficient should be plotted on

the quantile-transformed scale, whereas FALSE indicates the scale of the original data. Note this is different from the Qtransform arguemnt of af, which specifies

the scale on which the term is fit.

... arguments handed over to plot.gam

#### Value

This function's main purpose is its side effect of generating plots. It also silently returns a list of the data used to produce the plots, which can be used to generate customized plots.

## Author(s)

Jonathan Gellar

### See Also

```
af, pfr
```

predict.fbps

Prediction for fast bivariate P-spline (fbps)

### Description

Produces predictions given a fbps object and new data

### Usage

```
## S3 method for class 'fbps'
predict(object, newdata, ...)
```

# **Arguments**

object an object returned by fbps

newdata a data frame or list consisting of x and z values for which predicted values are

desired. vectors of x and z need to be of the same length.

... additional arguments.

## Value

A list with components

x a vector of x given in newdata z a vector of z given in newdata

fitted.values a vector of fitted values corresponding to x and z given in newdata

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### Author(s)

Luo Xiao <lxiao@jhsph.edu>

#### References

Xiao, L., Li, Y., and Ruppert, D. (2013). Fast bivariate *P*-splines: the sandwich smoother. *Journal of the Royal Statistical Society: Series B*, 75(3), 577–599.

```
#### True function #####
###############################
n1 <- 60
n2 <- 80
x <- (1: n1)/n1-1/2/n1
z <- (1: n2)/n2-1/2/n2
MY <- array(0,c(length(x),length(z)))
sigx < - .3
sigz < - .4
for(i in 1: length(x))
 for(j in 1: length(z))
   \#MY[i,j] < .75/(pi*sigx*sigz) *exp(-(x[i]-.2)^2/sigx^2-(z[j]-.3)^2/sigz^2)
  \#MY[i,j] <- MY[i,j] + .45/(pi*sigx*sigz) *exp(-(x[i]-.7)^2/sigx^2-(z[j]-.8)^2/sigz^2)
   MY[i,j] = sin(2*pi*(x[i]-.5)^3)*cos(4*pi*z[j])
 }
##############################
#### Observed data #####
sigma <- 1
Y \leftarrow MY + sigma*rnorm(n1*n2,0,1)
#############################
#### Estimation
                 #####
est <- fbps(Y,list(x=x,z=z))
mse <- mean((est$Yhat-MY)^2)</pre>
cat("mse of fbps is",mse,"\n")
cat("The smoothing parameters are:",est$lambda,"\n")
######## Compare the estimated surface with the true surface ########
par(mfrow=c(1,2))
persp(x,z,MY,zlab="f(x,z)",zlim=c(-1,2.5), phi=30,theta=45,expand=0.8,r=4,
     col="blue",main="True surface")
persp(x,z,est\$Yhat,zlab="f(x,z)",zlim=c(-1,2.5),phi=30,theta=45,
     expand=0.8,r=4,col="red",main="Estimated surface")
```

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```
#############################
#### prediction
# 1. make prediction with predict.fbps() for all pairs of x and z given in the original data
     ( it's expected to have same results as Yhat obtianed using fbps() above )
newdata <- list(x= rep(x, length(z)), z = rep(z, each=length(x)))
pred1 <- predict(est, newdata=newdata)$fitted.values</pre>
pred1.mat <- matrix(pred1, nrow=length(x))</pre>
par(mfrow=c(1,2))
image(pred1.mat); image(est$Yhat)
all.equal(as.numeric(pred1.mat), as.numeric(est$Yhat))
# 2. predict for pairs of first 10 x values and first 5 z values
newdata <- list(x= rep(x[1:10], 5), z = rep(z[1:5], each=10))
pred2 <- predict(est, newdata=newdata)$fitted.values</pre>
pred2.mat <- matrix(pred2, nrow=10)</pre>
par(mfrow=c(1,2))
image(pred2.mat); image(est$Yhat[1:10,1:5])
all.equal(as.numeric(pred2.mat), as.numeric(est$Yhat[1:10,1:5]))
# 3. predict for one pair
newdata <- list(x=x[5], z=z[3])
pred3 <- predict(est, newdata=newdata)$fitted.values</pre>
all.equal(as.numeric(pred3), as.numeric(est$Yhat[5,3]))
```

predict.fgam

Prediction from a fitted FGAM model

### **Description**

Takes a fitted fgam-object produced by fgam and produces predictions given a new set of values for the model covariates or the original values used for the model fit. Predictions can be accompanied by standard errors, based on the posterior distribution of the model coefficients. This is a wrapper function for predict.gam()

# Usage

```
## $3 method for class 'fgam'
predict(
  object,
  newdata,
  type = "response",
  se.fit = FALSE,
  terms = NULL,
  PredOutOfRange = FALSE,
  ...
)
```

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#### **Arguments**

object a fitted fgam object as produced by fgam

newdata a named list containing the values of the model covariates at which predictions

are required. If this is not provided then predictions corresponding to the original data are returned. All variables provided to newdata should be in the format supplied to fgam, i.e., functional predictors must be supplied as matrices with each row corresponding to one observed function. Index variables for the functional covariates are reused from the fitted model object or alternatively can be supplied as attributes of the matrix of functional predictor values. Any variables in the model not specified in newdata are set to their average values from the

data supplied during fitting the model

type character; see predict.gam for details
se.fit logical; see predict.gam for details
terms character see predict.gam for details

PredOutOfRange logical; if this argument is true then any functional predictor values in newdata

corresponding to fgam terms that are greater[less] than the maximum[minimum] of the domain of the marginal basis for the rows of the tensor product smooth are set to the maximum[minimum] of the domain. If this argument is false, attempting to predict a value of the functional predictor outside the range of this

basis produces an error

... additional arguments passed on to predict.gam

#### Value

If type == "lpmatrix", the design matrix for the supplied covariate values in long format. If se == TRUE, a list with entries fit and se.fit containing fits and standard errors, respectively. If type == "terms" or "iterms" each of these lists is a list of matrices of the same dimension as the response for newdata containing the linear predictor and its se for each term

## Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com> and Fabian Scheipl

#### See Also

```
fgam, predict.gam
```

```
######### Octane data example ########
data(gasoline)
N <- length(gasoline$octane)
wavelengths = 2*450:850
nir = matrix(NA, 60,401)
test <- sample(60,20)
for (i in 1:60) nir[i,] = gasoline$NIR[i, ] # changes class from AsIs to matrix
y <- gasoline$octane
#fit <- fgam(y~af(nir,xind=wavelengths,splinepars=list(k=c(6,6),m=list(c(2,2),c(2,2)))),</pre>
```

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```
# subset=(1:N)[-test])
#preds <- predict(fit,newdata=list(nir=nir[test,]),type='response')
#plot(preds,y[test])
#abline(a=0,b=1)</pre>
```

predict.fosr

Prediction from a fitted bayes\_fosr model

## **Description**

Takes a fitted fosr-object produced by bayes\_fosr and produces predictions given a new set of values for the model covariates or the original values used for the model fit.

## Usage

```
## S3 method for class 'fosr'
predict(object, newdata, ...)
```

### **Arguments**

object a fitted fosr object as produced by bayes\_fosr

newdata a named list containing the values of the model covariates at which predictions

are required. If this is not provided then predictions corresponding to the original data are returned. All variables provided to newdata should be in the format

supplied to the model fitting function.

... additional (unused) arguments

### Value

...

#### Author(s)

Jeff Goldsmith < jeff.goldsmith@columbia.edu>

#### See Also

```
bayes_fosr
```

```
## Not run:
library(reshape2)
library(dplyr)
library(ggplot2)
##### Cross-sectional real-data example #####
```

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```
## organize data
data(DTI)
DTI = subset(DTI, select = c(cca, case, pasat))
DTI = DTI[complete.cases(DTI),]
DTI$gender = factor(sample(c("male","female"), dim(DTI)[1], replace = TRUE))
DTI$status = factor(sample(c("RRMS", "SPMS", "PPMS"), dim(DTI)[1], replace = TRUE))
## fit models
VB = bayes_fosr(cca ~ pasat, data = DTI, Kp = 4, Kt = 10)
## obtain predictions
pred = predict(VB, sample_n(DTI, 10))
## End(Not run)
```

predict.fosr.vs

Prediction for Function-on Scalar Regression with variable selection

### **Description**

Given a "fosr.vs" object and new data, produces fitted values.

## Usage

```
## S3 method for class 'fosr.vs'
predict(object, newdata = NULL, ...)
```

# **Arguments**

object an object of class "fosr.vs".

newdata a data frame that contains the values of the model covariates at which predictors

are required.

... additional arguments.

## Value

fitted values.

### Author(s)

Yakuan Chen <yc2641@cumc.columbia.edu>

#### See Also

fosr.vs

Predict.matrix.dt.smooth 139

### **Examples**

```
## Not run:
I = 100
p = 20
D = 50
grid = seq(0, 1, length = D)
beta.true = matrix(0, p, D)
beta.true[1,] = sin(2*grid*pi)
beta.true[2,] = cos(2*grid*pi)
beta.true[3,] = 2
psi.true = matrix(NA, 2, D)
psi.true[1,] = sin(4*grid*pi)
psi.true[2,] = cos(4*grid*pi)
lambda = c(3,1)
set.seed(100)
X = matrix(rnorm(I*p), I, p)
C = cbind(rnorm(I, mean = 0, sd = lambda[1]), rnorm(I, mean = 0, sd = lambda[2]))
fixef = X%*%beta.true
pcaef = C %*% psi.true
error = matrix(rnorm(I*D), I, D)
Yi.true = fixef
Yi.pca = fixef + pcaef
Yi.obs = fixef + pcaef + error
data = as.data.frame(X)
data$Y = Yi.obs
fit.mcp = fosr.vs(Y~., data = data[1:80,], method="grMCP")
predicted.value = predict(fit.mcp, data[81:100,])
## End(Not run)
```

Predict.matrix.dt.smooth

Predict.matrix method for dt basis

## Description

Predict.matrix method for dt basis

#### Usage

```
## S3 method for class 'dt.smooth'
Predict.matrix(object, data)
```

## **Arguments**

object a dt. smooth object created by smooth.construct.dt.smooth.spec, see smooth.construct

data see smooth.construct

### Value

design matrix for domain-transformed terms

## Author(s)

Jonathan Gellar

Predict.matrix.fpc.smooth

mgcv-style constructor for prediction of FPC terms

# Description

mgcv-style constructor for prediction of FPC terms

# Usage

```
## S3 method for class 'fpc.smooth'
Predict.matrix(object, data)
```

# Arguments

object a fpc.smooth object created by smooth.construct.fpc.smooth.spec, see

smooth.construct

data see smooth.construct

### Value

design matrix for FPC terms

## Author(s)

Jonathan Gellar

```
Predict.matrix.pcre.random.effect
```

mgcv-style constructor for prediction of PC-basis functional random effects

# Description

mgcv-style constructor for prediction of PC-basis functional random effects

## Usage

```
## S3 method for class 'pcre.random.effect'
Predict.matrix(object, data)
```

### **Arguments**

object a smooth specification object, see smooth.construct

data see smooth.construct

### Value

design matrix for PC-based functional random effects

### Author(s)

Fabian Scheipl; adapted from 'Predict.matrix.random.effect' by S.N. Wood.

```
Predict.matrix.peer.smooth
```

mgcv-style constructor for prediction of PEER terms

# Description

mgcv-style constructor for prediction of PEER terms

### Usage

```
## S3 method for class 'peer.smooth'
Predict.matrix(object, data)
```

### Arguments

object a peer.smooth object created by smooth.construct.peer.smooth.spec, see

smooth.construct

data see smooth.construct

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## Value

design matrix for PEER terms

## Author(s)

Jonathan Gellar

Predict.matrix.pi.smooth

Predict.matrix method for pi basis

# Description

Predict.matrix method for pi basis

## Usage

```
## S3 method for class 'pi.smooth'
Predict.matrix(object, data)
```

## **Arguments**

object a pi.smooth object created by smooth.construct.pi.smooth.spec, see smooth.construct

data see smooth.construct

### Value

design matrix for PEER terms

### Author(s)

Jonathan Gellar

predict.pffr

Prediction for penalized function-on-function regression

# Description

Takes a fitted pffr-object produced by pffr() and produces predictions given a new set of values for the model covariates or the original values used for the model fit. Predictions can be accompanied by standard errors, based on the posterior distribution of the model coefficients. This is a wrapper function for predict.gam().

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### Usage

```
## S3 method for class 'pffr'
predict(object, newdata, reformat = TRUE, type = "link", se.fit = FALSE, ...)
```

### **Arguments**

object a fitted pffr-object

newdata A named list (or a data.frame) containing the values of the model covariates

at which predictions are required. If no newdata is provided then predictions corresponding to the original data are returned. If newdata is provided then it must contain all the variables needed for prediction, in the format supplied to pffr, i.e., functional predictors must be supplied as matrices with each row corresponding to one observed function. See Details for more on index variables

and prediction for models fit on irregular or sparse data.

reformat logical, defaults to TRUE. Should predictions be returned in matrix form (de-

fault) or in the long vector shape returned by predict.gam()?

type see predict.gam() for details. Note that type == "lpmatrix" will force reformat

to FALSE.

se.fit see predict.gam()

... additional arguments passed on to predict.gam()

#### **Details**

Index variables (i.e., evaluation points) for the functional covariates are reused from the fitted model object and cannot be supplied with newdata. Prediction is always for the entire index range of the responses as defined in the original fit. If the original fit was performed on sparse or irregular, non-gridded response data supplied via pffr's ydata-argument and no newdata was supplied, this function will simply return fitted values for the original evaluation points of the response (in list form). If the original fit was performed on sparse or irregular data and newdata was supplied, the function will return predictions on the grid of evaluation points given in object\$pffr\$yind.

#### Value

If type == "lpmatrix", the design matrix for the supplied covariate values in long format. If se == TRUE, a list with entries fit and se.fit containing fits and standard errors, respectively. If type == "terms" or "iterms" each of these lists is a list of matrices of the same dimension as the response for newdata containing the linear predictor and its se for each term.

#### Author(s)

Fabian Scheipl

#### See Also

```
predict.gam()
```

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predict.pfr

Prediction from a fitted pfr model

### **Description**

Takes a fitted pfr-object produced by pfr and produces predictions given a new set of values for the model covariates or the original values used for the model fit. Predictions can be accompanied by standard errors, based on the posterior distribution of the model coefficients. This is a wrapper function for predict.gam()

## Usage

```
## S3 method for class 'pfr'
predict(
  object,
  newdata,
  type = "response",
  se.fit = FALSE,
  terms = NULL,
  PredOutOfRange = FALSE,
  ...
)
```

## Arguments

object a fitted pfr object as produced by pfr

newdata a named list containing the values of the model covariates at which predictions

are required. If this is not provided then predictions corresponding to the original data are returned. All variables provided to newdata should be in the format supplied to pfr, i.e., functional predictors must be supplied as matrices with each row corresponding to one observed function. Index variables for the functional covariates are reused from the fitted model object or alternatively can be supplied as attributes of the matrix of functional predictor values. Any variables in the model not specified in newdata are set to their average values from the

data supplied during fitting the model

type character; see predict.gam for details
se.fit logical; see predict.gam for details
terms character see predict.gam for details

PredOutOfRange logical; if this argument is true then any functional predictor values in newdata

corresponding to pfr terms that are greater[less] than the maximum[minimum] of the domain of the marginal basis for the rows of the tensor product smooth are set to the maximum[minimum] of the domain. If this argument is false, attempting to predict a value of the functional predictor outside the range of this

basis produces an error

... additional arguments passed on to predict.gam

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### Value

If type == "lpmatrix", the design matrix for the supplied covariate values in long format. If se == TRUE, a list with entries fit and se.fit containing fits and standard errors, respectively. If type == "terms" or "iterms" each of these lists is a list of matrices of the same dimension as the response for newdata containing the linear predictor and its se for each term

#### Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com> and Fabian Scheipl

#### See Also

```
pfr, predict.gam
```

### **Examples**

print.summary.pffr

Print method for summary of a pffr fit

# Description

Pretty printing for a summary.pffr-object. See print.summary.gam() for details.

### Usage

```
## S3 method for class 'summary.pffr'
print(
    x,
    digits = max(3, getOption("digits") - 3),
    signif.stars = getOption("show.signif.stars"),
    ...
)
```

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# Arguments

```
    x a fitted pffr-object
    digits controls number of digits printed in output.
    signif.stars Should significance stars be printed alongside output?
    not used
```

# Value

```
A summary.pffr object
```

### Author(s)

Fabian Scheipl, adapted from print.summary.gam() by Simon Wood, Henric Nilsson

pwcv

Pointwise cross-validation for function-on-scalar regression

# **Description**

Estimates prediction error for a function-on-scalar regression model by leave-one-function-out cross-validation (CV), at each of a specified set of points.

# Usage

```
pwcv(
  fdobj,
  Z,
  L = NULL,
  lambda,
  eval.pts = seq(min(fdobj$basis$range), max(fdobj$basis$range), length.out = 201),
  scale = FALSE
)
```

# **Arguments**

fdobj	a functional data object (class fd) giving the functional responses.
Z	the model matrix, whose columns represent scalar predictors.
L	a row vector or matrix of linear contrasts of the coefficient functions, to be restricted to equal zero.
lambda	smoothing parameter: either a nonnegative scalar or a vector, of length nco1(Z), of nonnegative values.
eval.pts	argument values at which the CV score is to be evaluated.
scale	logical value or vector determining scaling of the matrix Z (see scale, to which the value of this argument is passed).

qq.pffr

### **Details**

Integrating the pointwise CV estimate over the function domain yields the *cross-validated integrated squared error*, the standard overall model fit score returned by lofocv.

It may be desirable to derive the value of lambda from an appropriate call to fosr, as in the example below.

#### Value

A vector of the same length as eval.pts giving the CV scores.

### Author(s)

```
Philip Reiss <phil.reiss@nyumc.org>
```

#### References

Reiss, P. T., Huang, L., and Mennes, M. (2010). Fast function-on-scalar regression with penalized basis expansions. *International Journal of Biostatistics*, 6(1), article 28. Available at https://pubmed.ncbi.nlm.nih.gov/21969982/

#### See Also

```
fosr, lofocv
```

qq.pffr

QQ plots for pffr model residuals

# Description

This is simply a wrapper for qq.gam().

### Usage

```
qq.pffr(
  object,
  rep = 0,
  level = 0.9,
  s.rep = 10,
  type = c("deviance", "pearson", "response"),
  pch = ".",
  rl.col = 2,
  rep.col = "gray80",
  ...
)
```

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# Arguments

object	a fitted pffr-object
rep	How many replicate datasets to generate to simulate quantiles of the residual distribution. 0 results in an efficient simulation free method for direct calculation, if this is possible for the object family.
level	If simulation is used for the quantiles, then reference intervals can be provided for the QQ-plot, this specifies the level. 0 or less for no intervals, 1 or more to simply plot the QQ plot for each replicate generated.
s.rep	how many times to randomize uniform quantiles to data under direct computation.
type	what sort of residuals should be plotted? See residuals.gam.
pch	plot character to use. 19 is good.
rl.col	color for the reference line on the plot.
rep.col	color for reference bands or replicate reference plots.
	extra graphics parameters to pass to plotting functions.

quadWeights	Compute quadrature weights	

# Description

Utility function for numerical integration.

# Usage

```
quadWeights(argvals, method = "trapezoidal")
```

# Arguments

argvals function arguments.

method quadrature method. Can be either trapedoidal or midpoint.

# Value

a vector of quadrature weights for the points supplied in argvals.

# Author(s)

Clara Happ, with modifications by Philip Reiss

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re

Random effects constructor for fgam

# Description

Sets up a random effect for the levels of x. Use the by-argument to request random slopes.

### Usage

```
re(x, ...)
```

### Arguments

```
x a grouping variable: must be a factor... further arguments handed over to s, see random.effects
```

# **Details**

See random. effects in mgcv.

### See Also

random.effects

residuals.pffr

Obtain residuals and fitted values for a pffr models

# Description

See predict.pffr for alternative options to extract estimated values from a pffr object. "Fitted values" here refers to the estimated additive predictor values, these will not be on the scale of the response for models with link functions.

# Usage

```
## S3 method for class 'pffr'
residuals(object, reformat = TRUE, ...)
## S3 method for class 'pffr'
fitted(object, reformat = TRUE, ...)
```

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# **Arguments**

object a fitted pffr-object

reformat logical, defaults to TRUE. Should residuals be returned in n x yindex matrix

form (regular grid data) or, respectively, in the shape of the originally supplied ydata argument (sparse/irregular data), or, if FALSE, simply as a long vector as

returned by resid.gam()?

... other arguments, passed to residuals.gam.

#### Value

A matrix or ydata-like data. frame or a vector of residuals / fitted values (see reformat-argument)

### Author(s)

Fabian Scheipl

rlrt.pfr Likelihood Ratio Test and Restricted Likelihood Ratio Test for infer-

ence of functional predictors

# **Description**

NOTE: this function is designed to work with pfr\_old() rather than pfr(). Given a pfr object of family="gaussian", tests whether the function is identically equal to its mean (constancy), or whether the functional predictor significantly improves the model (inclusion). Based on zero-variance-component work of Crainiceanu et al. (2004), Scheipl et al. (2008), and Swihart et al. (2012).

### Usage

```
rlrt.pfr(pfr.obj = pfr.obj, test = NULL, ...)
```

# **Arguments**

pfr.obj an object returned by pfr\_old()

test "constancy" will test functional form of the coefficient function of the last func-

tion listed in funcs in pfr.obj against the null of a constant line: the average of the functional predictor. "inclusion" will test functional form of the coefficient function of the last function listed in funcs in pfr.obj against the null of 0: that

is, whether the functional predictor should be included in the model.

... additional arguments

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#### **Details**

A Penalized Functional Regression of family="gaussian" can be represented as a linear mixed model dependent on variance components. Testing whether certain variance components and (potentially) fixed effect coefficients are 0 correspond to tests of constancy and inclusion of functional predictors.

For rlrt.pfr, Restricted Likelihood Ratio Test is preferred for the constancy test as under the special B-splines implementation of pfr for the coefficient function basis the test involves only the variance component. Therefore, the constancy test is best for pfr objects with method="REML"; if the method was something else, a warning is printed and the model refit with "REML" and a test is then conducted.

For rlrt.pfr, the Likelihood Ratio Test is preferred for the inclusion test as under the special B-splines implementation of pfr for the coefficient function basis the test involves both the variance component and a fixed effect coefficient in the linear mixed model representation. Therefore, the inclusion test is best for pfr objects with method="ML"; if the method was something else, a warning is printed and the model refit with "ML" and a test is then conducted.

#### Value

p.val the p-value for the full model (alternative) against the null specified by the test

test.stat the test statistic, see Scheipl et al. 2008 and Swihart et al 2012

ma the alternative model as fit with mgcv::gam

m0 the null model as fit with mgcv::gam

m the model containing only the parameters being tested as fit with mgcv::gam

### Author(s)

Jeff Goldsmith <jeff.goldsmith@columbia.edu> and Bruce Swihart <bswihart@jhsph.edu>

### References

Goldsmith, J., Bobb, J., Crainiceanu, C., Caffo, B., and Reich, D. (2011). Penalized functional regression. *Journal of Computational and Graphical Statistics*, 20(4), 830–851.

Goldsmith, J., Crainiceanu, C., Caffo, B., and Reich, D. (2012). Longitudinal penalized functional regression for cognitive outcomes on neuronal tract measurements. *Journal of the Royal Statistical Society: Series C*, 61(3), 453–469.

Crainiceanu, C. and Ruppert, D. (2004) Likelihood ratio tests in linear mixed models with one variance component. *Journal of the Royal Statistical Society: Series B*, 66, 165–185.

Scheipl, F. (2007) Testing for nonparametric terms and random effects in structured additive regression. Diploma thesis.\ https://www.statistik.lmu.de/~scheipl/downloads/DIPLOM.zip.

Scheipl, F., Greven, S. and Kuechenhoff, H (2008) Size and power of tests for a zero random effect variance or polynomial regression in additive and linear mixed models. *Computational Statistics & Data Analysis*, 52(7), 3283–3299.

Swihart, Bruce J., Goldsmith, Jeff; and Crainiceanu, Ciprian M. (2012). Testing for functional effects. Johns Hopkins University Dept. of Biostatistics Working Paper 247. Available at https://biostats.bepress.com/jhubiostat/paper247/

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#### See Also

```
pfr, predict.pfr, package RLRsim
```

# **Examples**

```
## Not run:
DTI Data Example
                                              #########
# For more about this example, see Swihart et al. 2012
# Testing for Functional Effects
## load and reassign the data;
data(DTI2)
O <- DTI2$pasat ## PASAT outcome
id <- DTI2$id ## subject id
W1 <- DTI2$cca ## Corpus Callosum
W2 <- DTI2$rcst ## Right corticospinal
V <- DTI2$visit ## visit
## prep scalar covariate
visit.1.rest <- matrix(as.numeric(V > 1), ncol=1)
covar.in <- visit.1.rest</pre>
## note there is missingness in the functional predictors
apply(is.na(W1), 2, mean)
apply(is.na(W2), 2, mean)
## fit two univariate models, then one model with both functional predictors
pfr.obj.t2 <- pfr_old(Y = 0, covariates=covar.in, funcs = list(W2), subj = id, kz = 10, kb = 50)
pfr.obj.t3 <- pfr_old(Y = 0, covariates=covar.in, funcs = list(W1, W2), subj = id, kz = 10, kb = 50)
## plot the coefficient function and bounds
dev.new()
par(mfrow=c(2,2))
ran <- c(-2,.5)
matplot(cbind(pfr.obj.t1$BetaHat[[1]], pfr.obj.t1$Bounds[[1]]),
 type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
 main = "CCA", xlab="Location", ylim=ran)
abline(h=0, col="blue")
matplot(cbind(pfr.obj.t2$BetaHat[[1]], pfr.obj.t2$Bounds[[1]]),
 type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
 main = "RCST", xlab="Location", ylim=ran)
abline(h=0, col="blue")
matplot(cbind(pfr.obj.t3$BetaHat[[1]], pfr.obj.t3$Bounds[[1]]),
 type = '1', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
 main = "CCA - mult.", xlab="Location", ylim=ran)
```

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```
abline(h=0, col="blue")
matplot(cbind(pfr.obj.t3$BetaHat[[2]], pfr.obj.t3$Bounds[[2]]),
  type = 'l', lty = c(1,2,2), col = c(1,2,2), ylab = "BetaHat",
  main = "RCST - mult.", xlab="Location", ylim=ran)
abline(h=0, col="blue")
## do some testing
t1 <- rlrt.pfr(pfr.obj.t1, "constancy")</pre>
t2 <- rlrt.pfr(pfr.obj.t2, "constancy")</pre>
t3 <- rlrt.pfr(pfr.obj.t3, "inclusion")
t1$test.stat
t1$p.val
t2$test.stat
t2$p.val
t3$test.stat
t3$p.val
## do some testing with rlrt.pfr(); same as above but subj = NULL
pfr.obj.t1 <- pfr(Y = 0, covariates=covar.in, funcs = list(W1),</pre>
                                                                    subj = NULL, kz = 10, kb = 50)
pfr.obj.t2 \leftarrow pfr(Y = 0, covariates = covar.in, funcs = list(W2), subj = NULL, kz = 10, kb = 50)
pfr.obj.t3 \leftarrow pfr(Y = 0, covariates = covar.in, funcs = list(W1, W2), subj = NULL, kz = 10, kb = 50)
t1 <- rlrt.pfr(pfr.obj.t1, "constancy")</pre>
t2 <- rlrt.pfr(pfr.obj.t2, "constancy")
t3 <- rlrt.pfr(pfr.obj.t3, "inclusion")
t1$test.stat
t1$p.val
t2$test.stat
t2$p.val
t3$test.stat
t3$p.val
## End(Not run)
```

sff

Construct a smooth function-on-function regression term

# Description

Defines a term  $\int_{s_{lo,i}}^{s_{hi,i}} f(X_i(s),s,t)ds$  for inclusion in an mgcv::gam-formula (or bam or gamm or gamm4:::gamm) as constructed by pffr. Defaults to a cubic tensor product B-spline with marginal second differences penalties for  $f(X_i(s),s,t)$  and integration over the entire range  $[s_{lo,i},s_{hi,i}]$ 

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 $[\min(s_i), \max(s_i)]$ . Can't deal with any missing X(s), unequal lengths of  $X_i(s)$  not (yet?) possible. Unequal ranges for different  $X_i(s)$  should work.  $X_i(s)$  is assumed to be numeric. sff() IS AN EXPERIMENTAL FEATURE AND NOT WELL TESTED YET – USE AT YOUR OWN RISK.

# Usage

```
sff(
   X,
   yind,
   xind = seq(0, 1, 1 = ncol(X)),
   basistype = c("te", "t2", "s"),
   integration = c("simpson", "trapezoidal"),
   L = NULL,
   limits = NULL,
   splinepars = list(bs = "ps", m = c(2, 2, 2))
)
```

# **Arguments**

_	
X	an n by ncol(xind) matrix of function evaluations $X_i(s_{i1}),\dots,X_i(s_{iS}); i=1,\dots,n.$
yind	<i>DEPRECATED</i> matrix (or vector) of indices of evaluations of $Y_i(t)$ ; i.e. matrix with rows $(t_{i1}, \ldots, t_{iT})$ ; no longer used.
xind	vector of indices of evaluations of $X_i(s)$ , i.e, $(s_1, \ldots, s_S)$
basistype	defaults to "te", i.e. a tensor product spline to represent $f(X_i(s),t)$ . Alternatively, use "s" for bivariate basis functions (see s) or "t2" for an alternative parameterization of tensor product splines (see t2).
integration	method used for numerical integration. Defaults to "simpson"'s rule. Alternatively and for non-equidistant grids, "trapezoidal".
L	optional: an n by $ncol(xind)$ giving the weights for the numerical integration over $s$ .
limits	defaults to NULL for integration across the entire range of $X(s)$ , otherwise specifies the integration limits $s_{hi,i}, s_{lo,i}$ : either one of "s <t" "s<='t"' <math="" for="" or="">(s_{hi,i}, s_{lo,i}) = (0,t) or a function that takes s as the first and t as the second argument and returns TRUE for combinations of values (s,t) if s falls into the integration range for the given t. This is an experimental feature and not well tested yet; use at your own risk.</t">
splinepars	optional arguments supplied to the basistype-term. Defaults to a cubic tensor product B-spline with marginal second differences, i.e. list(bs="ps", $m=c(2,2,2)$ ). See te or s for details

# Value

# a list containing

• call a "call" to te (or s, t2) using the appropriately constructed covariate and weight matrices (see linear.functional.terms)

data a list containing the necessary covariate and weight matrices

### Author(s)

Fabian Scheipl, based on Sonja Greven's trick for fitting functional responses.

```
smooth.construct.dt.smooth.spec

Domain Transformation basis constructor
```

# Description

The dt basis allows for any of the standard mgcv (or user-defined) bases to be aplied to a transformed version of the original terms. Smooths may be of any number of terms. Transformations are specified by supplying a function of any or all of the original terms. "by" variables are not transformed.

### Usage

```
## S3 method for class 'dt.smooth.spec'
smooth.construct(object, data, knots)
```

### **Arguments**

object	a smooth specification object, generated by $s()$ , $te()$ , $ti()$ , or $t2()$ , with $bs="dt"$
data	a list containing just the data (including any by variable) required by this term, with names corresponding to object\$term (and object\$by). The by variable is the last element.
knots	a list containing any knots supplied for basis setup - in same order and with same names as data. Can be NULL.

### **Details**

object should be creaated with an xt argument. For non-tensor-product smooths, this will be a list with the following elements:

- 1. tf (required): a function or character string (or list of functions and/or character strings) defining the coordinate transformations; see further details below.
- 2. bs (optional): character string indicating the bs for the basis applied to the transformed coordinates; if empty, the appropriate defaults are used.
- 3. basistype (optional): character string indicating type of bivariate basis used. Options include "s" (the default), "te", "ti", and "t2", which correspond to s, te, ti, and t2.
- 4. ... (optional): for tensor product smooths, additional arguments to the function specified by basistype that are not available in s() can be included here, e.g. d, np, etc.

For tensor product smooths, we recommend using s() to set up the basis, and specifying the tensor product using xt\$basistype as described above. If the basis is set up using te(), then the variables in object\$term will be split up, meaning all transformation functions would have to be univariate.

#### Value

An object of class "dt.smooth". This will contain all the elements associated with the smooth.construct object from the inner smooth (defined by xt\$bs), in addition to an xt element used by the Predict.matrix method.

#### **Transformation Functions**

Let nterms = length(object\$term). The tf element can take one of the following forms:

- 1. a function of nargs arguments, where nargs <= nterms. If nterms > 1, it is assumed that this function will be applied to the first term of object\$term. If all argument names of the function are term names, then those arguments will correspond to those terms; otherwise, they will correspond to the first nargs terms in object\$term.
- 2. a character string corresponding to one of the built-in transformations (listed below).
- 3. A list of length ntfuncs, where ntfuncs<=nterms, containing either the functions or character strings described above. If this list is named with term names, then the transformation functions will be applied to those terms; otherwise, they will be applied to the first ntfuncs terms in object\$term.

The following character strings are recognized as built-in transformations:

- "log": log transformation (univariate)
- "ecdf": empirical cumulative distribution function (univariate)
- "linear01": linearly rescale from 0 to 1 (univariate)
- "s-t": first term ("s") minus the second term ("t") (bivariate)
- "s/t": first term ("s") divided by the second term ("t") (bivariate)
- "QTransform": performs a time-specific eddf transformation for a bivariate smooth, where time is indicated by the first term, and x by the second term. Primarily for use with refund::af.

Some transformations rely on a fixed "pivot point" based on the data used to fit the model, e.g. quantiles (such as the min or max) of this data. When making predictions based on these transformations, the transformation function will need to know what the pivot points are, based on the original (not prediction) data. In order to accomplish this, we allow the user to specify that they want their transformation function to refer to the original data (as opposed to whatever the "current" data is). This is done by appending a zero ("0") to the argument name.

For example, suppose you want to scale the term linearly so that the data used to define the basis ranges from 0 to 1. The wrong way to define this transformation function: function(x)  $\{(x - \min(x))/(\max(x) - \min(x))\}$ . This function will result in incorrect predictions if the range of data for which predictions are being made is not the same as the range of data that was used to define the basis. The proper way to define this function: function(x)  $\{(x - \min(x0))/(\max(x0) - \min(x0))\}$ . By referring to x0 instead of x, you are indicating that you want to use the original data instead of the current data. This may seem strange to refer to a variable that is not one of the arguments, but the "dt" constructor explicitly places these variables in the environment of the transformation function to make them available.

# Author(s)

Jonathan Gellar

### See Also

smooth.construct

```
smooth.construct.fpc.smooth.spec

**Basis constructor for FPC terms**
```

# **Description**

Basis constructor for FPC terms

### Usage

```
## S3 method for class 'fpc.smooth.spec'
smooth.construct(object, data, knots)
```

### **Arguments**

object a fpc.smooth.spec object, usually generated by a term s(x, bs="fpc"); see

Details.

data a list containing the data (including any by variable) required by this term, with

names corresponding to object\$term (and object\$by). Only the first element

of this list is used.

knots not used, but required by the generic smooth.construct.

### **Details**

object must contain an xt element. This is a list that can contain the following elements:

X (required) matrix of functional predictors

**method** (required) the method of finding principal components; options include "svd" (unconstrained), "fpca.sc", "fpca.face", or "fpca.ssvd"

**npc** (optional) the number of PC's to retain

**pve** (only needed if npc not supplied) the percent variance explained used to determine npc

penalize (required) if FALSE, the smoothing parameter is set to 0

bs the basis class used to pre-smooth X; default is "ps"

Any additional options for the pre-smoothing basis (e.g. k, m, etc.) can be supplied in the corresponding elements of object. See s for a full list of options.

### Value

An object of class "fpc.smooth". In additional to the elements listed in smooth.construct, the object will contain

sm the smooth that is fit in order to generate the basis matrix over object\$term

V.A the matrix of principal components

### Author(s)

Jonathan Gellar < JGellar@mathematica-mpr.com>

# References

Reiss, P. T., and Ogden, R. T. (2007). Functional principal component regression and functional partial least squares. *Journal of the American Statistical Association*, 102, 984-996.

#### See Also

fpcr

```
smooth.construct.pco.smooth.spec

Principal coordinate ridge regression
```

### Description

Smooth constructor function for principal coordinate ridge regression fitted by gam. When the principal coordinates are defined by a relevant distance among functional predictors, this is a form of nonparametric scalar-on-function regression. Reiss et al. (2016) describe the approach and apply it to dynamic time warping distances among functional predictors.

# Usage

```
## S3 method for class 'pco.smooth.spec'
smooth.construct(object, data, knots)
```

# **Arguments**

object a smooth specification object, usually generated by a term of the form s(dummy,

bs="pco", k, xt); see Details.

data a list containing just the data.

knots IGNORED!

#### Value

An object of class pco. smooth. The resulting object has an xt element which contains details of the multidimensional scaling, which may be interesting.

#### **Details**

The constructor is not normally called directly, but is rather used internally by gam.

In a gam term of the above form s(dummy, bs="pco", k, xt),

- dummy is an arbitrary vector (or name of a column in data) whose length is the number of observations. This is not actually used, but is required as part of the input to s. Note that if multiple pco terms are used in the model, there must be multiple unique term names (e.g., "dummy1", "dummy2", etc).
- k is the number of principal coordinates (e.g., k=9 will give a 9-dimensional projection of the data).
- xt is a list supplying the distance information, in one of two ways. (i) A matrix Dmat of distances can be supplied directly via xt=list(D=Dmat,...). (ii) Alternatively, one can use xt=list(realdata=..., dist\_fn=..., ...) to specify a data matrix realdata and distance function dist\_fn, whereupon a distance matrix dist\_fn(realdata) is created.

The list xt also has the following optional elements:

- add: Passed to cmdscale when performing multidimensional scaling; for details, see the help for that function. (Default FALSE.)
- fastcmd: if TRUE, multidimensional scaling is performed by cmdscale\_lanczos, which uses
  Lanczos iteration to eigendecompose the distance matrix; if FALSE, MDS is carried out by
  cmdscale. Default is FALSE, to use cmdscale.

#### Author(s)

David L Miller, based on code from Lan Huo and Phil Reiss

### References

Reiss, P. T., Miller, D. L., Wu, P.-S., and Wen-Yu Hua, W.-Y. Penalized nonparametric scalar-on-function regression via principal coordinates. Under revision. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC5714326/.

### **Examples**

```
## Not run:
# a simulated example
library(refund)
library(mgcv)
require(dtw)

## First generate the data
Xnl <- matrix(0, 30, 101)
set.seed(813)
tt <- sort(sample(1:90, 30))
for(i in 1:30){
    Xnl[i, tt[i]:(tt[i]+4)] <- -1
    Xnl[i, (tt[i]+5):(tt[i]+9)] <- 1
}</pre>
```

```
X.toy <- Xnl + matrix(rnorm(30*101, ,0.05), 30)
y.toy <- tt + rnorm(30, 0.05)
y.rainbow <- rainbow(30, end=0.9)[(y.toy-min(y.toy))/</pre>
                                    diff(range(y.toy))*29+1]
## Display the toy data
par(mfrow=c(2, 2))
matplot((0:100)/100, t(Xnl[c(4, 25), ]), type="1", xlab="t", ylab="",
        ylim=range(X.toy), main="Noiseless functions")
matplot((0:100)/100, t(X.toy[c(4, 25), ]), type="l", xlab="t", ylab="",
        ylim=range(X.toy), main="Observed functions")
matplot((0:100)/100, t(X.toy), type="l", lty=1, col=y.rainbow, xlab="t",
        ylab="", main="Rainbow plot")
## Obtain DTW distances
D.dtw <- dist(X.toy, method="dtw", window.type="sakoechiba", window.size=5)</pre>
## Compare PC vs. PCo ridge regression
# matrix to store results
GCVmat <- matrix(NA, 15, 2)
# dummy response variable
dummy \leftarrow rep(1,30)
# loop over possible projection dimensions
for (k. in 1:15){
  # fit PC (m1) and PCo (m2) ridge regression
  m1 <- gam(y.toy ~ s(dummy, bs="pco", k=k.,
            xt=list(realdata=X.toy, dist_fn=dist)), method="REML")
  m2 \leftarrow gam(y.toy \sim s(dummy, bs="pco", k=k., xt=list(D=D.dtw)), method="REML")
  # calculate and store GCV scores
  GCVmat[k., ] \leftarrow length(y.toy) * c(sum(m1$residuals^2)/m1$df.residual^2,
                   sum(m2$residuals^2)/m2$df.residual^2)
}
## plot the GCV scores per dimension for each model
matplot(GCVmat, lty=1:2, col=1, pch=16:17, type="o", ylab="GCV",
        xlab="Number of principal components / coordinates",
        main="GCV score")
legend("right", c("PC ridge regression", "DTW-based PCoRR"), lty=1:2, pch=16:17)
## example of making a prediction
# fit a model to the toy data
m <- gam(y.toy ~ s(dummy, bs="pco", k=2, xt=list(D=D.dtw)), method="REML")</pre>
# first build the distance matrix
# in this case we just subsample the original matrix
# see ?pco_predict_preprocess for more information on formatting this data
dist_list \leftarrow list(dummy = as.matrix(D.dtw)[, c(1:5,10:15)])
# preprocess the prediction data
pred_data <- pco_predict_preprocess(m, newdata=NULL, dist_list)</pre>
```

```
# make the prediction
p <- predict(m, pred_data)

# check that these are the same as the corresponding fitted values
print(cbind(fitted(m)[ c(1:5,10:15)],p))

## End(Not run)</pre>
```

smooth.construct.pcre.smooth.spec

mgcv-style constructor for PC-basis functional random effects

# Description

Sets up design matrix for functional random effects based on the PC scores of the covariance operator of the random effect process. See smooth.construct.re.smooth.spec for more details on mgcv-style smoother specification and pcre for the corresponding pffr()-formula wrapper.

# Usage

```
## S3 method for class 'pcre.smooth.spec'
smooth.construct(object, data, knots)
```

# Arguments

object a smooth specification object, see smooth.construct

data see smooth.construct
knots see smooth.construct

### Value

An object of class "random.effect". See smooth.construct for the elements that this object will contain.

# Author(s)

Fabian Scheipl; adapted from 're' constructor by S.N. Wood.

```
smooth.construct.peer.smooth.spec

**Basis constructor for PEER terms**
```

### Description

Smooth basis constructor to define structured penalties (Randolph et al., 2012) for smooth terms.

### Usage

```
## S3 method for class 'peer.smooth.spec'
smooth.construct(object, data, knots)
```

### **Arguments**

object a peer.smooth.spec object, usually generated by a term s(x, bs="peer"); see Details.

data a list containing the data (including any by variable) required by this term, with

names corresponding to object\$term (and object\$by). Only the first element

of this list is used.

knots not used, but required by the generic smooth.construct.

#### **Details**

The smooth specification object, defined using s(), should contain an xt element. xt will be a list that contains additional information needed to specify the penalty. The type of penalty is indicated by xt\$pentype. There are four types of penalties available:

- 1. xt\$pentype=="RIDGE" for a ridge penalty, the default
- 2. xt\$pentype=="D" for a difference penalty. The order of the difference penalty is specified by the m argument of s().
- 3. xt\$pentype=="DECOMP" for a decomposition-based penalty,  $bP_Q+a(I-P_Q)$ , where  $P_Q=Q^t(QQ^t)^{-1}Q$ . The Q matrix must be specified by xt\$Q, and the scalar a by xt\$phia. The number of columns of Q must be equal to the length of the data. Each row represents a basis function where the functional predictor is expected to lie, according to prior belief.
- 4. xtpentype=="USER" for a user-specified penalty matrix L, supplied by xtL.

### Value

An object of class "peer.smooth". See smooth.construct for the elements that this object will contain.

### Author(s)

Madan Gopal Kundu <mgkundu@iupui.edu> and Jonathan Gellar

#### References

Randolph, T. W., Harezlak, J, and Feng, Z. (2012). Structured penalties for functional linear models - partially empirical eigenvectors for regression. *Electronic Journal of Statistics*, 6, 323-353.

#### See Also

peer

```
smooth.construct.pi.smooth.spec
```

Parametric Interaction basis constructor

# Description

The pi basis is appropriate for smooths of multiple variables. Its purpose is to parameterize the way in which the basis changes with one of those variables. For example, suppose the smooth is over three variables, x, y, and t, and we want to parameterize the effect of t. Then the pi basis will assume  $f(x,y,t) = \sum_k g_k(t) * f_k(x,y)$ , where the  $g_k(t)$  functions are pre-specified and the  $f_k(x,y)$  functions are estimated using a bivariate basis. An example of a parametric interaction is a linear interaction, which would take the form  $f(x,y,t) = f_1(x,y) + t * f_2(x,y)$ .

#### Usage

```
## S3 method for class 'pi.smooth.spec'
smooth.construct(object, data, knots)
```

### **Arguments**

object	a smooth specification object, generated by, e.g., s(x, y, t, bs="pi", xt=list(g=list(g1, g2, g3))). For transformation functions g1, g2, and g3, see Details below.
data	a list containing the variables of the smooth (x, y, and t above), as well as any by variable.
knots	a list containing any knots supplied for basis setup - in same order and with same names as data. Can be NULL.

#### **Details**

All functions  $f_k()$  are defined using the same basis set. Accordingly, they are penalized using a single block-diagonal penalty matrix and one smoothing parameter. Future versions of this function may be able to relax this assumption.

object should be defined (using s()) with an xt argument. This argument is a list that could contain any of the following elements:

- 1. g: the functions  $g_k(t)$ , specified as described below.
- 2. bs: the basis code used for the functions  $f_k()$ ; defaults to thin-plate regression splines, which is mgcv's default. The same basis will be used for all k.

- 3. idx: an integer index indicating which variable from object\$term is to be parameterized, i.e., the *t* variable; defaults to length(object\$term)
- 4. mp: flag to indicate whether multiple penalties should be estimated, one for each  $f_k()$ . Defaults to TRUE. If FALSE, the penalties for each k are combined into a single block-diagonal penalty matrix (with one smoothing parameter).
- 5. ...: further xt options to be passed onto the basis for  $f_k()$ .

xt\$g can be entered in one of the following forms:

- 1. a list of functions of length k, where each function is of one argument (assumed to be t)
- 2. one of the following recognized character strings: "linear", indicating a linear interaction, i.e.  $f(x,t) = f_1(x) + t * f_2(x)$ ; "quadratic", indicating a quadratic interaction, i.e.  $f(x,t) = f_1(x) + t * f_2(x) + t^2 * f_3(x)$ ; or "none", indicating no interaction with t, i.e.  $f(x,t) = f_1(x)$ .

The only one of the above elements that is required is xt. If default values for bs, idx, and mp are desired, xt may also be entered as the g element itself; i.e. xt=g, where g is either the list of functions or an acceptable character string.

Additional arguments for the lower-dimensional basis over f\_k may be entered using the corresponding arguments of s(), e.g. k, m, sp, etc. For example, s(x, t, bs="pi", k=15, xt=list(g="linear", bs="ps")) will define a linear interaction with t of a univariate p-spline basis of dimension 15 over x.

#### Value

An object of class "pi.smooth". See smooth. construct for the elements it will contain.

#### Author(s)

Fabian Scheipl and Jonathan Gellar

```
smooth.construct.pss.smooth.spec
```

P-spline constructor with modified 'shrinkage' penalty

### Description

Construct a B-spline basis with a modified difference penalty of full rank (i.e., that also penalizes low-order polynomials).

### Usage

```
## S3 method for class 'pss.smooth.spec'
smooth.construct(object, data, knots)
```

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### **Arguments**

object see smooth.construct. The shrinkage factor can be specified via object\$xt\$shrink

data see smooth.construct.
knots see smooth.construct.

#### **Details**

This penalty-basis combination is useful to avoid non-identifiability issues for ff terms. See 'ts' or 'cs' in smooth.terms for similar "shrinkage penalties" for thin plate and cubic regression splines. The basic idea is to replace the k-th zero eigenvalue of the original penalty by  $s^k \nu_m$ , where s is the shrinkage factor (defaults to 0.1) and  $\nu_m$  is the smallest non-zero eigenvalue. See reference for the original idea, implementation follows that in the 'ts' and 'cs' constructors (see smooth.terms).

#### Author(s)

Fabian Scheipl; adapted from 'ts' and 'cs' constructors by S.N. Wood.

#### References

Marra, G., & Wood, S. N. (2011). Practical variable selection for generalized additive models. *Computational Statistics & Data Analysis*, 55(7), 2372-2387.

sofa SOFA (Sequential Organ Failure Assessment) Data

### **Description**

A dataset containing the SOFA scores (Vincent et al, 1996). for 520 patients, hospitalized in the intensive care unit (ICU) with Acute Lung Injury. Daily measurements are available for as long as each one remains in the ICU. This is an example of variable-domain functional data, as described by Gellar et al. (2014).

# Usage

sofa

# **Format**

A data frame with 520 rows (subjects) and 7 variables:

death binary indicator that the subject died in the ICU

**SOFA** 520 x 173 matrix in variable-domain format (a ragged array). Each column represents an ICU day. Each row contains the SOFA scores for a subject, one per day, for as long as the subject remained in the ICU. The remaining cells of each row are padded with NAs. SOFA scores range from 0 to 24, increasing with severity of organ failure. Missing values during one's ICU stay have been imputed using LOCF.

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**SOFA\_raw** Identical to the SOFA element, except that it contains some missing values during one's hospitalization. These missing values arise when a subject leaves the ICU temporarily, only to be re-admitted. SOFA scores are not monitored outside the ICU.

**los** ICU length of stay, i.e., the number of days the patient remained in the ICU prior to death or final discharge.

age Patient age

male Binary indicator for male gender

**Charlson** Charlson co-morbidity index, a measure of baseline health status (before hospitalization and ALI).

#### **Details**

The data was collected as part of the Improving Care of ALI Patients (ICAP) study (Needham et al., 2006). If you use this dataset as an example in written work, please cite the study protocol.

### References

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summary.pffr

Summary for a pffr fit

### **Description**

Take a fitted pffr-object and produce summaries from it. See summary.gam() for details.

# Usage

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

### **Arguments**

```
object a fitted pffr-object
... see summary.gam() for options.
```

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# Value

A list with summary information, see summary.gam()

### Author(s)

Fabian Scheipl, adapted from summary.gam() by Simon Wood, Henric Nilsson

summary.pfr

Summary for a pfr fit

# **Description**

Take a fitted pfr-object and produce summaries from it. See summary.gam() for details.

# Usage

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

# **Arguments**

```
object a fitted pfr-object
... see summary.gam() for options.
```

### **Details**

This function currently simply strips the "pfr" class label and calls summary.gam.

### Value

A list with summary information, see summary.gam()

# Author(s)

Jonathan Gellar < JGellar@mathematica-mpr.com>, Fabian Scheipl

vb\_cs\_fpca

vb\_cs\_fpca

Cross-sectional FoSR using Variational Bayes and FPCA

# Description

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using a VB and estimates the residual covariance surface using FPCA.

# Usage

```
vb_cs_fpca(
  formula,
  data = NULL,
  verbose = TRUE,
  Kt = 5,
  Kp = 2,
  alpha = 0.1,
  Aw = NULL,
  Bw = NULL,
  Apsi = NULL,
  Bpsi = NULL,
  argvals = NULL)
```

# Arguments

formula	a formula indicating the structure of the proposed model.
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
verbose	logical defaulting to TRUE – should updates on progress be printed?
Kt	number of spline basis functions used to estimate coefficient functions
Кр	number of FPCA basis functions to be estimated
alpha	tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha = 0 is all second-derivative penalty)
Aw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
Bw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
Apsi	hyperparameter for inverse gamma controlling variance of spline terms for FPC effects
Bpsi	hyperparameter for inverse gamma controlling variance of spline terms for FPC effects
argvals	not currently implemented

vb\_cs\_wish

### Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

vb\_cs\_wish

Cross-sectional FoSR using Variational Bayes and Wishart prior

# **Description**

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using VB and estimates the residual covariance surface using a Wishart prior.

# Usage

```
vb_cs_wish(
  formula,
  data = NULL,
  verbose = TRUE,
  Kt = 5,
  alpha = 0.1,
  min.iter = 10,
  max.iter = 50,
  Aw = NULL,
  Bw = NULL,
  v = NULL
)
```

# Arguments

formula	a formula indicating the structure of the proposed model.
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
verbose	logical defaulting to TRUE – should updates on progress be printed?
Kt	number of spline basis functions used to estimate coefficient functions
alpha	tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha $= 0$ is all second-derivative penalty)
min.iter	minimum number of iterations of VB algorithm
max.iter	maximum number of iterations of VB algorithm

vb\_mult\_fpca

Aw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects; if NULL, defaults to $Kt/2$ .
Bw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects; if NULL, defaults to $1/2$ tr(mu.q.beta of the model
V	hyperparameter for inverse Wishart prior on residual covariance; if NULL, Psi defaults to an FPCA decomposition of the residual covariance in which residuals are estimated based on an OLS fit of the model (note the "nugget effect" on this covariance is assumed to be constant over the time domain).

### Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

vb\_mult\_fpca

Multilevel FoSR using Variational Bayes and FPCA

### **Description**

Fitting function for function-on-scalar regression for multilevel data. This function estimates model parameters using a VB and estimates the residual covariance surface using FPCA.

# Usage

```
vb_mult_fpca(
  formula,
  data = NULL,
  verbose = TRUE,
  Kt = 5,
  Kp = 2,
  alpha = 0.1,
  argvals = NULL
)
```

# **Arguments**

formula a formula indicating the structure of the proposed model.

data an optional data frame, list or environment containing the variables in the model.

If not found in data, the variables are taken from environment(formula), typi-

cally the environment from which the function is called.

verbose logical defaulting to TRUE – should updates on progress be printed?

vb\_mult\_wish

Kt	number of spline basis functions used to estimate coefficient functions
Кр	number of FPCA basis functions to be estimated
alpha	tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha = $0$ is all second-derivative penalty)
argvals	not currently implemented

# Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

vb\_mult\_wish

Multilevel FoSR using Variational Bayes and Wishart prior

# Description

Fitting function for function-on-scalar regression for cross-sectional data. This function estimates model parameters using VB and estimates the residual covariance surface using a Wishart prior. If prior hyperparameters are NULL they are estimated using the data.

# Usage

```
vb_mult_wish(
  formula,
  data = NULL,
  verbose = TRUE,
  Kt = 5,
  alpha = 0.1,
  min.iter = 10,
  max.iter = 50,
  Az = NULL,
  Bz = NULL,
  Bw = NULL,
  v = NULL
)
```

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### **Arguments**

formula	a formula indicating the structure of the proposed model.
data	an optional data frame, list or environment containing the variables in the model. If not found in data, the variables are taken from environment(formula), typically the environment from which the function is called.
verbose	logical defaulting to TRUE – should updates on progress be printed?
Kt	number of spline basis functions used to estimate coefficient functions
alpha	tuning parameter balancing second-derivative penalty and zeroth-derivative penalty (alpha = 0 is all second-derivative penalty)
min.iter	minimum number of iterations of VB algorithm
max.iter	maximum number of iterations of VB algorithm
Az	hyperparameter for inverse gamma controlling variance of spline terms for subject-level effects
Bz	hyperparameter for inverse gamma controlling variance of spline terms for subject-level effects
Aw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
Bw	hyperparameter for inverse gamma controlling variance of spline terms for population-level effects
V	hyperparameter for inverse Wishart prior on residual covariance

# Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

### References

Goldsmith, J., Kitago, T. (2016). Assessing Systematic Effects of Stroke on Motor Control using Hierarchical Function-on-Scalar Regression. *Journal of the Royal Statistical Society: Series C*, 65 215-236.

vis.fgam	Visualization of FGAM objects	

# Description

Produces perspective or contour plot views of an estimated surface corresponding to af terms fit using fgam or plots "slices" of the estimated surface or estimated second derivative surface with one of its arguments fixed and corresponding twice-standard error "Bayesian" confidence bands constructed using the method in Marra and Wood (2012). See the details.

vis.fgam

# Usage

```
vis.fgam(
  object,
  af.term,
  xval = NULL,
  tval = NULL,
  deriv2 = FALSE,
  theta = 50,
  plot.type = "persp",
  ticktype = "detailed",
  ...
)
```

# Arguments

object	an fgam object, produced by fgam
af.term	character; the name of the functional predictor to be plotted. Only important if multiple af terms are fit. Defaults to the first af term in object\$call
xval	a number in the range of functional predictor to be plotted. The surface will be plotted with the first argument of the estimated surface fixed at this value
tval	a number in the domain of the functional predictor to be plotted. The surface will be plotted with the second argument of the estimated surface fixed at this value. Ignored if xval is specified
deriv2	logical; if TRUE, plot the estimated second derivative surface along with Bayesian confidence bands. Only implemented for the "slices" plot from either xval or tval being specified
theta	numeric; viewing angle; see persp
plot.type	one of "contour" (to use levelplot) or "persp" (to use persp). Ignored if either xval or tval is specified
ticktype	how to draw the tick marks if plot.type="persp". Defaults to "detailed"
• • •	other options to be passed to persp, levelplot, or plot

### **Details**

The confidence bands used when plotting slices of the estimated surface or second derivative surface are the ones proposed in Marra and Wood (2012). These are a generalization of the "Bayesian" intervals of Wahba (1983) with an adjustment for the uncertainty about the model intercept. The estimated covariance matrix of the model parameters is obtained from assuming a particular Bayesian model on the parameters.

# Value

Simply produces a plot

# Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com>

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#### References

McLean, M. W., Hooker, G., Staicu, A.-M., Scheipl, F., and Ruppert, D. (2014). Functional generalized additive models. *Journal of Computational and Graphical Statistics*, **23(1)**, pp. 249-269. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3982924/.

Marra, G., and Wood, S. N. (2012) Coverage properties of confidence intervals for generalized additive model components. *Scandinavian Journal of Statistics*, **39(1)**, pp. 53–74.

Wabha, G. (1983) "Confidence intervals" for the cross-validated smoothing spline. *Journal of the Royal Statistical Society, Series B*, **45**(1), pp. 133–150.

#### See Also

```
vis.gam, plot.gam, fgam, persp, levelplot
```

### **Examples**

```
data(DTI)
## only consider first visit and cases (since no PASAT scores for controls)
y <- DTI$pasat[DTI$visit==1 & DTI$case==1]</pre>
X <- DTI$cca[DTI$visit==1 & DTI$case==1,]</pre>
## remove samples containing missing data
ind <- rowSums(is.na(X))>0
y \leftarrow y[!ind]
X \leftarrow X[!ind,]
## fit the fgam using FA measurements along corpus
## callosum as functional predictor with PASAT as response
## using 8 cubic B-splines for each marginal bases with
## third order marginal difference penalties
## specifying gamma>1 enforces more smoothing when using GCV
## to choose smoothing parameters
#fit <- fgam(y\sim f(X,splinepars=list(k=c(8,8),m=list(c(2,3),c(2,3)))),gamma=1.2)
## contour plot of the fitted surface
#vis.fgam(fit,plot.type='contour')
## similar to Figure 5 from McLean et al.
## Bands seem too conservative in some cases
#xval <- runif(1, min(fit$fgam$ft[[1]]$Xrange), max(fit$fgam$ft[[1]]$Xrange))</pre>
#tval <- runif(1, min(fit$fgam$ft[[1]]$xind), max(fit$fgam$ft[[1]]$xind))</pre>
\#par(mfrow=c(4, 1))
#vis.fgam(fit, af.term='X', deriv2=FALSE, xval=xval)
#vis.fgam(fit, af.term='X', deriv2=FALSE, tval=tval)
#vis.fgam(fit, af.term='X', deriv2=TRUE, xval=xval)
#vis.fgam(fit, af.term='X', deriv2=TRUE, tval=tval)
```

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vis.pfr

Visualization of PFR objects

# Description

Produces perspective or contour plot views of an estimated surface corresponding smooths over two or more dimensions. Alternatively plots "slices" of the estimated surface or estimated second derivative surface with one of its arguments fixed. Corresponding twice-standard error "Bayesian" confidence bands are constructed using the method in Marra and Wood (2012). See the details.

# Usage

```
vis.pfr(
  object,
  select = 1,
  xval = NULL,
  tval = NULL,
  deriv2 = FALSE,
  theta = 50,
  plot.type = "persp",
  ticktype = "detailed",
  ...
)
```

# Arguments

object	an pfr object, produced by pfr
select	index for the smooth term to be plotted, according to its position in the model formula (and in object\$smooth). Not needed if only one multivariate term is present.
xval	a number in the range of functional predictor to be plotted. The surface will be plotted with the first argument of the estimated surface fixed at this value
tval	a number in the domain of the functional predictor to be plotted. The surface will be plotted with the second argument of the estimated surface fixed at this value. Ignored if xval is specified.
deriv2	logical; if TRUE, plot the estimated second derivative surface along with Bayesian confidence bands. Only implemented for the "slices" plot from either xval or tval being specified
theta	numeric; viewing angle; see persp
plot.type	one of "contour" (to use levelplot) or "persp" (to use persp). Ignored if either xval or tval is specified
ticktype	how to draw the tick marks if plot.type="persp". Defaults to "detailed"
	other options to be passed to persp, levelplot, or plot

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#### **Details**

The confidence bands used when plotting slices of the estimated surface or second derivative surface are the ones proposed in Marra and Wood (2012). These are a generalization of the "Bayesian" intervals of Wahba (1983) with an adjustment for the uncertainty about the model intercept. The estimated covariance matrix of the model parameters is obtained from assuming a particular Bayesian model on the parameters.

#### Value

Simply produces a plot

#### Author(s)

Mathew W. McLean <mathew.w.mclean@gmail.com>

#### References

McLean, M. W., Hooker, G., Staicu, A.-M., Scheipl, F., and Ruppert, D. (2014). Functional generalized additive models. *Journal of Computational and Graphical Statistics*, **23**(1), pp. 249-269. Available at https://www.ncbi.nlm.nih.gov/pmc/articles/PMC3982924/.

Marra, G., and Wood, S. N. (2012) Coverage properties of confidence intervals for generalized additive model components. *Scandinavian Journal of Statistics*, **39(1)**, pp. 53–74.

Wabha, G. (1983) "Confidence intervals" for the cross-validated smoothing spline. *Journal of the Royal Statistical Society, Series B*, **45**(1), pp. 133–150.

### See Also

```
vis.gam, plot.gam, pfr, persp, levelplot
```

### **Examples**

Xt\_siginv\_X

```
## similar to Figure 5 from McLean et al.
## Bands seem too conservative in some cases
xval <- runif(1, min(fit$pfr$ft[[1]]$Xrange), max(fit$pfr$ft[[1]]$Xrange))
tval <- runif(1, min(fit$pfr$ft[[1]]$xind), max(fit$pfr$ft[[1]]$xind))
par(mfrow=c(2, 2))
vis.pfr(fit, deriv2=FALSE, xval=xval)
vis.pfr(fit, deriv2=FALSE, tval=tval)
vis.pfr(fit, deriv2=TRUE, xval=xval)
vis.pfr(fit, deriv2=TRUE, tval=tval)</pre>
```

Xt\_siginv\_X

Internal computation function

# Description

Internal function used compute the products in cross-sectional VB algorithm and Gibbs sampler

# Usage

```
Xt_siginv_X(tx, siginv, y = NULL)
```

### **Arguments**

tx transpose of the X design matrix
siginv inverse variance matrix
y outcome matrix. if NULL, function computes first product; if not, function computes second product.

# Author(s)

Jeff Goldsmith <ajg2202@cumc.columbia.edu>

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