

# Package ‘iprior’

December 19, 2017

**Title** Regression Modelling using I-Priors

**Version** 0.7.1

**Encoding** UTF-8

**Description** Provides methods to perform and analyse I-prior regression models.  
Estimation is done either via direct optimisation of the log-likelihood or  
an EM algorithm.

**URL** <https://github.com/haziqj/iprior>

**BugReports** <https://github.com/haziqj/iprior/issues>

**License** GPL (>= 3.0)

**Depends** R (>= 3.2.5)

**Imports** doSNOW, foreach, ggplot2, mvtnorm, Rcpp (>= 0.12.5), reshape2,  
scales

**LinkingTo** Rcpp, RcppEigen

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**LazyData** yes

**VignetteBuilder** knitr, R.rsp

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Accessors *Accessor functions for ipriorMod objects.*

---

### Description

Accessor functions for ipriorMod objects.

### Usage

`get_intercept(object)`

`get_y(object)`

`get_size(object, units = "kB", standard = "SI")`

`get_hyp(object)`

`get_lambda(object)`

`get_psi(object)`

`get_lengthscales(object)`

```
get_hurst(object)
get_offset(object)
get_degree(object)
get_se(object)
get_kernels(object)
get_kern_matrix(object, theta = NULL, xstar = list(NULL))
get_prederror(object, error.type = c("RMSE", "MSE"))
get_estl(object)
get_method(object)
get_convergence(object)
get_niter(object)
get_time(object)
get_theta(object)
```

### Arguments

object	An ipriorMod object.
units	Units for object size.
standard	Standard for object size.
theta	(Optional) Value of hyperparameters to evaluate the kernel matrix.
xstar	(Optional) If not supplied, then a square, symmetric kernel matrix is returned using the data as input points. Otherwise, the kernel matrix is evaluated with respect to this set of data as well. It must be a list of vectors/matrices with similar dimensions to the original data.
error.type	(Optional) Report the mean squared error of prediction ("MSE"), or the root mean squared error of prediction ("RMSE")

### Functions

- `get_intercept`: Obtain the intercept.
- `get_y`: Obtain the response variables.
- `get_size`: Obtain the object size of the I-prior model.
- `get_hyp`: Obtain the hyperparameters of the model (both estimated and fixed ones).

- `get_lambda`: Obtain the scale parameters used.
- `get_psi`: Obtain the error precision.
- `get_lengthscales`: Obtain the lengthscales for the SE kernels used.
- `get_hurst`: Obtain the Hurst coefficient of the fBm kernels used.
- `get_offset`: Obtain the offset parameters for the polynomial kernels used.
- `get_degree`: Obtain the degree of the polynomial kernels used.
- `get_se`: Obtain the standard errors of the estimated hyperparameters.
- `get_kernels`: Obtain the kernels used.
- `get_kern_matrix`: Obtain the kernel matrix of the I-prior model.
- `get_prederror`: Obtain the training mean squared error.
- `get_est1`: Obtain information on which hyperparameters were estimated and which were fixed.
- `get_method`: Obtain the estimation method used.
- `get_convergence`: Obtain the convergence information.
- `get_niter`: Obtain the number of iterations performed.
- `get_time`: Obtain the time taken to complete the estimation procedure.
- `get_theta`: Extract the theta value at convergence. Note that this is on an unrestricted scale (see the vignette for details).

---

as.time

*Convert difftime class into time class*


---

### Description

Convert difftime class into time class

### Usage

```
as.time(x)
```

### Arguments

x                    A difftime object.

### Value

A time object which contains the time difference and units.

---

check_theta	<i>Check the structure of the hyperparameters of an I-prior model</i>
-------------	---

---

**Description**

Check the structure of the hyperparameters of an I-prior model

**Usage**

```
check_theta(object)
```

**Arguments**

object            An ipriorMod object or an ipriorKernel object.

**Value**

A printout of the structure of the hyperparameters.

---

decimal_place	<i>Cut a numeric vector to a certain number of decimal places</i>
---------------	---

---

**Description**

Cut a numeric vector to a certain number of decimal places

**Usage**

```
decimal_place(x, k = 2)
```

```
dec_plac(x, k = 2)
```

**Arguments**

x                A numeric vector.  
k                The number of decimal places.

**Value**

A character vector with the correct number of decimal places.

**Examples**

```
decimal_place(pi, 3)  
decimal_place(c(exp(1), pi, sqrt(2)), 4)
```

---

`eigenCpp`*Eigen decomposition of a matrix in C++.*

---

**Description**

Returns the eigenvalues and eigenvectors of a matrix X.

**Usage**`eigenCpp(X)`**Arguments**

X                    A symmetric, positive-definite matrix

**Details**

A fast implementation of eigen for symmetric, positive-definite matrices. This helps speed up the I-prior EM algorithm.

---

`fastSquare`*Multiplying a symmetric matrix by itself in C++.*

---

**Description**

Returns the square of a symmetric matrix X.

**Usage**`fastSquare(X)`**Arguments**

X                    A symmetric matrix

**Details**

A fast implementation of  $X^2$  for symmetric matrices. This helps speed up the I-prior EM algorithm.

---

fastVDiag	<i>Computing a quadratic matrix form in C++.</i>
-----------	--

---

**Description**

Returns XdiagXT.

**Usage**

```
fastVDiag(X, y)
```

**Arguments**

X	A symmetric, square matrix of dimension n by n
y	A vector of length n

**Details**

A fast implementation of XdiagXT. This helps speed up the I-prior EM algorithm.

---

gen_multilevel	<i>Generate simulated data for multilevel models</i>
----------------	--

---

**Description**

Generate simulated data for multilevel models

**Usage**

```
gen_multilevel(n = 25, m = 6, sigma_e = 2, sigma_u0 = 2, sigma_u1 = 2,
  sigma_u01 = -2, beta0 = 0, beta1 = 2, x.jitter = 0.5, seed = NULL)
```

**Arguments**

n	Sample size. Input either a single number for a balanced data set, or a vector of length m indicating the sample size in each group.
m	Number of groups/levels.
sigma_e	The standard deviation of the errors.
sigma_u0	The standard deviation of the random intercept.
sigma_u1	The standard deviation of the random slopes.
sigma_u01	The covariance of between the random intercept and the random slope.
beta0	The mean of the random intercept.
beta1	The mean of the random slope.
x.jitter	A small amount of jitter is added to the X variables generated from a normal distribution with mean zero and standard deviation equal to x.jitter.
seed	(Optional) Random seed.

**Value**

A dataframe containing the response variable  $y$ , the unidimensional explanatory variables  $X$ , and the levels/groups (factors).

**Examples**

```
gen_multilevel()
```

---

gen_smooth	<i>Generate simulated data for smoothing models</i>
------------	---

---

**Description**

Generate simulated data for smoothing models

**Usage**

```
gen_smooth(n = 150, xlim = c(0.2, 4.6), x.jitter = 0.65, seed = NULL)
```

**Arguments**

n	Sample size.
xlim	Limits of the $X$ variables to generate from.
x.jitter	A small amount of jitter is added to the $X$ variables generated from a normal distribution with mean zero and standard deviation equal to <code>x.jitter</code> .
seed	(Optional) Random seed.

**Value**

A dataframe containing the response variable  $y$  and unidimensional explanatory variable  $X$ .

**Examples**

```
gen_smooth(10)
```



---

gg_colour_hue	<i>Emulate ggplot2 default colour palette</i>
---------------	---

---

**Description**

Emulate ggplot2 default colour palette. `ipriorColPal` and `ggColPal` are DEPRECATED.

**Usage**

```
gg_colour_hue(x, h = c(0, 360) + 15, c = 100, l = 65)
```

```
gg_color_hue(x, h = c(0, 360) + 15, c = 100, l = 65)
```

```
gg_col_hue(x, h = c(0, 360) + 15, c = 100, l = 65)
```

```
ipriorColPal(x)
```

```
ggColPal(x)
```

**Arguments**

<code>x</code>	The number of colours required.
<code>h</code>	Range of hues to use, in [0, 360].
<code>c</code>	Chroma (intensity of colour), maximum value varies depending on combination of hue and luminance.
<code>l</code>	Luminance (lightness), in [0, 100].

**Details**

This is the default colour scale for categorical variables in ggplot2. It maps each level to an evenly spaced hue on the colour wheel. It does not generate colour-blind safe palettes.

`ipriorColPal()` used to provide the colour palette for the `iprior` package, but this has been changed ggplot2's colour palette instead.

---

hsb	<i>High school and beyond dataset</i>
-----	---------------------------------------

---

**Description**

A national longitudinal survey of of students from public and private high schools in the United States, with information such as students' cognitive and non-cognitive skills, high school experiences, work experiences and future plans collected.

**Usage**

hsb

**Format**

A data frame of 7185 observations on 3 variables.

mathach Math achievement.

ses Socio-Economic status.

schoolid Categorical variable indicating the school the student went to. Treated as [factor](#).

**Source**

[High School and Beyond, 1980: A Longitudinal Survey of Students in the United States \(ICPSR 7896\)](#)

**References**

Rabe-Hesketh, S., & Skrondal, A. (2008). *Multilevel and longitudinal modeling using Stata*. STATA press.

Raudenbush, S. W. (2004). *HLM 6: Hierarchical linear and nonlinear modeling*. Scientific Software International.

Raudenbush, S. W., & Bryk, A. S. (2002). *Hierarchical linear models: Applications and data analysis methods* (Vol. 1). Sage.

**Examples**

```
data(hsb)
str(hsb)
```

---

hsbsmall

*High school and beyond dataset*

---

**Description**

Smaller subset of hsb.

**Usage**

hsbsmall

**Format**

A data frame of 661 observations on 3 variables.

mathach Math achievement.

ses Socio-Economic status.

schoolid Categorical variable indicating the school the student went to. Treated as [factor](#).

**Details**

A random subset of size 16 out of the original 160 groups.

**Examples**

```
data(hsbsmall)
str(hsbsmall)
```

---

iprior *Fit an I-prior regression model*

---

**Description**

A function to perform regression using I-priors. The I-prior model parameters may be estimated in a number of ways: direct minimisation of the marginal deviance, EM algorithm, fixed hyperparameters, or using a Nystrom kernel approximation.

**Usage**

```
## Default S3 method:
iprior(y, ..., kernel = "linear", method = "direct",
       control = list(), interactions = NULL, est.lambda = TRUE,
       est.hurst = FALSE, est.lengthscale = FALSE, est.offset = FALSE,
       est.psi = TRUE, fixed.hyp = NULL, lambda = 1, psi = 1,
       nystrom = FALSE, nys.seed = NULL, model = list(), train.samp, test.samp)

## S3 method for class 'formula'
iprior(formula, data, kernel = "linear", one.lam = FALSE,
       method = "direct", control = list(), est.lambda = TRUE,
       est.hurst = FALSE, est.lengthscale = FALSE, est.offset = FALSE,
       est.psi = TRUE, fixed.hyp = NULL, lambda = 1, psi = 1,
       nystrom = FALSE, nys.seed = NULL, model = list(), train.samp, test.samp,
       ...)

## S3 method for class 'ipriorKernel'
iprior(object, method = "direct", control = list(),
       ...)

## S3 method for class 'ipriorMod'
iprior(object, method = NULL, control = list(),
       iter.update = 100, ...)
```

**Arguments**

**y** Vector of response variables

**...** Only used when fitting using non-formula, enter the variables (vectors or matrices) separated by commas.

kernel	<p>Character vector indicating the type of kernel for the variables. Available choices are:</p> <ul style="list-style-type: none"> <li>• "linear" - (default) for the linear kernel</li> <li>• "canonical" - alternative name for "linear"</li> <li>• "fbm", "fbm, 0.5" - for the fBm kernel with Hurst coefficient 0.5 (default)</li> <li>• "se", "se, 1" - for the SE kernel with lengthscale 1 (default)</li> <li>• "poly", "poly2", "poly2, 0" - for the polynomial kernel of degree 2 with offset 0 (default)</li> <li>• "pearson" - for the Pearson kernel</li> </ul> <p>The kernel argument can also be a vector of length equal to the number of variables, therefore it is possible to specify different kernels for each variables. Note that factor type variables are assigned the Pearson kernel by default, and that non-factor types can be forced to use the Pearson kernel (not recommended).</p>
method	<p>The estimation method. One of:</p> <ul style="list-style-type: none"> <li>• "direct" - for the direct minimisation of the marginal deviance using <code>optim()</code>'s L-BFGS method</li> <li>• "em" - for the EM algorithm</li> <li>• "mixed" - combination of the direct and EM methods</li> <li>• "fixed" - for just obtaining the posterior regression function with fixed hyperparameters (default method when setting <code>fixed.hyp = TRUE</code>)</li> <li>• "canonical" - an efficient estimation method which takes advantage of the structure of the linear kernel</li> </ul>
control	<p>(Optional) A list of control options for the estimation procedure:</p> <p><code>maxit</code> The maximum number of iterations for the quasi-Newton optimisation or the EM algorithm. Defaults to 100.</p> <p><code>em.maxit</code> For method = "mixed", the number of EM steps before switching to direct optimisation. Defaults to 5.</p> <p><code>stop.crit</code> The stopping criterion for the EM and L-BFGS algorithm, which is the difference in successive log-likelihood values. Defaults to 1e-8.</p> <p><code>theta0</code> The initial values for the hyperparameters. Defaults to random starting values.</p> <p><code>report</code> The interval of reporting for the <code>optim()</code> function.</p> <p><code>restarts</code> The number of random restarts to perform. Defaults to 0. It's also possible to set it to TRUE, in which case the number of random restarts is set to the total number of available cores.</p> <p><code>no.cores</code> The number of cores in which to do random restarts. Defaults to the total number of available cores.</p> <p><code>omega</code> The overrelaxation parameter for the EM algorithm - a value between 0 and 1.</p>
interactions	<p>Character vector to specify the interaction terms. When using formulas, this is specified automatically, so is not required. Syntax is "a:b" to indicate variable a interacts with variable b.</p>
est.lambda	<p>Logical. Estimate the scale parameters? Defaults to TRUE.</p>

<code>est.hurst</code>	Logical. Estimate the Hurst coefficients for fBm kernels? Defaults to FALSE.
<code>est.lengthscale</code>	Logical. Estimate the lengthscales for SE kernels? Defaults to FALSE.
<code>est.offset</code>	Logical. Estimate the offsets for polynomial kernels? Defaults to FALSE.
<code>est.psi</code>	Logical. Estimate the error precision? Defaults to TRUE.
<code>fixed.hyp</code>	Logical. If TRUE, then no hyperparameters are estimated, i.e. all of the above <code>est.x</code> are set to FALSE, and vice versa. If NULL (default) then all of the <code>est.x</code> defaults are respected.
<code>lambda</code>	Initial/Default scale parameters. Relevant especially if <code>est.lambda = FALSE</code> .
<code>psi</code>	Initial/Default value for error precision. Relevant especially if <code>est.psi = FALSE</code> .
<code>nystrom</code>	Either logical or an integer indicating the number of Nystrom samples to take. Defaults to FALSE. If TRUE, then approximately 10% of the sample size is used for the Nystrom approximation.
<code>nys.seed</code>	The random seed for the Nystrom sampling. Defaults to NULL, which means the random seed is not fixed.
<code>model</code>	DEPRECATED.
<code>train.samp</code>	(Optional) A vector indicating which of the data points should be used for training, and the remaining used for testing.
<code>test.samp</code>	(Optional) Similar to <code>train.samp</code> , but on test samples instead.
<code>formula</code>	The formula to fit when using formula interface.
<code>data</code>	Data frame containing variables when using formula interface.
<code>one.lam</code>	Logical. When using formula input, this is a convenient way of letting the function know to treat all variables as a single variable (i.e. shared scale parameter). Defaults to FALSE.
<code>object</code>	An <code>ipriorKernel</code> or <code>ipriorMod</code> object.
<code>iter.update</code>	The number of iterations to perform when calling the function on an <code>ipriorMod</code> object. Defaults to 100.

## Details

The `iprior()` function is able to take formula based input and non-formula. When not using formula, the syntax is as per the default S3 method. That is, the response variable is the vector `y`, and any explanatory variables should follow this, and separated by commas.

As described [here](#), the model can be loaded first into an `ipriorKernel` object, and then passed to the `iprior()` function to perform the estimation.

## Value

An `ipriorMod` object. Several accessor functions have been written to obtain pertinent things from the `ipriorMod` object. The `print()` and `summary()` methods display the relevant model information.

**Methods (by class)**

- `ipriorKernel`: Takes in object of type `ipriorKernel`, a loaded and prepared I-prior model, and proceeds to estimate it.
- `ipriorMod`: Re-run or continue running the EM algorithm from last attained parameter values in object `ipriorMod`.

**See Also**

[optim](#), [update](#), [check\\_theta](#), [print](#), [summary](#), [plot](#), [coef](#), [sigma](#), [fitted](#), [predict](#), [logLik](#), [deviance](#).

**Examples**

```
# Formula based input
(mod.stackf <- iprior(stack.loss ~ Air.Flow + Water.Temp + Acid.Conc.,
                    data = stackloss))
mod.toothf <- iprior(len ~ supp * dose, data = ToothGrowth)
summary(mod.toothf)

# Non-formula based input
mod.stacknf <- iprior(y = stackloss$stack.loss,
                    Air.Flow = stackloss$Air.Flow,
                    Water.Temp = stackloss$Water.Temp,
                    Acid.Conc. = stackloss$Acid.Conc.)
mod.toothnf <- iprior(y = ToothGrowth$len, ToothGrowth$supp, ToothGrowth$dose,
                    interactions = "1:2")

# Formula based model option one.lam = TRUE
# Sets a single scale parameter for all variables
modf <- iprior(stack.loss ~ ., data = stackloss, one.lam = TRUE)
modnf <- iprior(y = stackloss$stack.loss, X = stackloss[1:3])
all.equal(coef(modnf), coef(modf)) # both models are equivalent

# Fit models using different kernels
dat <- gen_smooth(n = 100)
mod <- iprior(y ~ X, dat, kernel = "fbm") # Hurst = 0.5 (default)
mod <- iprior(y ~ X, dat, kernel = "poly3") # polynomial degree 3

# Fit models using various estimation methods
mod1 <- iprior(y ~ X, dat)
mod2 <- iprior(y ~ X, dat, method = "em")
mod3 <- iprior(y ~ X, dat, method = "canonical")
mod4 <- iprior(y ~ X, dat, method = "mixed")
mod5 <- iprior(y ~ X, dat, method = "fixed", lambda = coef(mod1)[1],
              psi = coef(mod1)[2])
all.equal(logLik(mod1), logLik(mod2), logLik(mod3), logLik(mod4),
          logLik(mod5), tol = 1e-6)

## Not run:

# For large data sets, it is worth trying the Nystrom method
```

```

mod <- iprior(y ~ X, gen_smooth(5000), kernel = "se", nystrom = 50,
             est.lengthscale = TRUE) # a bit slow
plot_fitted(mod, ci = FALSE)

## End(Not run)

```

---

iprior\_cv

*Perform a cross-validation experiment with the iprior function*


---

## Description

A convenience function to perform a k-fold cross-validation experiment and obtain mean squared error of prediction. Most of the arguments are similar to `iprior()` and `kernL()`.

## Usage

```

## Default S3 method:
iprior_cv(y, ..., folds = 2, par.cv = TRUE,
          kernel = "linear", method = "direct", control = list(),
          interactions = NULL, est.lambda = TRUE, est.hurst = FALSE,
          est.lengthscale = FALSE, est.offset = FALSE, est.psi = TRUE,
          fixed.hyp = NULL, lambda = 1, psi = 1, nystrom = FALSE,
          nys.seed = NULL)

## S3 method for class 'formula'
iprior_cv(formula, data, folds = 2, one.lam = FALSE,
          par.cv = TRUE, kernel = "linear", method = "direct", control = list(),
          est.lambda = TRUE, est.hurst = FALSE, est.lengthscale = FALSE,
          est.offset = FALSE, est.psi = TRUE, fixed.hyp = NULL, lambda = 1,
          psi = 1, nystrom = FALSE, nys.seed = NULL, ...)

```

## Arguments

<code>y</code>	Vector of response variables
<code>...</code>	Only used when fitting using non-formula, enter the variables (vectors or matrices) separated by commas.
<code>folds</code>	The number of cross-validation folds. Set equal to sample size or <code>Inf</code> to perform leave-one-out cross-validation.
<code>par.cv</code>	Logical. Multithreading to fit the models? Defaults to <code>TRUE</code> .
<code>kernel</code>	Character vector indicating the type of kernel for the variables. Available choices are: <ul style="list-style-type: none"> <li>"linear" - (default) for the linear kernel</li> <li>"canonical" - alternative name for "linear"</li> <li>"fbm", "fbm,0.5" - for the fBm kernel with Hurst coefficient 0.5 (default)</li> <li>"se", "se,1" - for the SE kernel with lengthscale 1 (default)</li> </ul>

- "poly", "poly2", "poly2,0" - for the polynomial kernel of degree 2 with offset 0 (default)
- "pearson" - for the Pearson kernel

The kernel argument can also be a vector of length equal to the number of variables, therefore it is possible to specify different kernels for each variables. Note that factor type variables are assigned the Pearson kernel by default, and that non-factor types can be forced to use the Pearson kernel (not recommended).

method	<p>The estimation method. One of:</p> <ul style="list-style-type: none"> <li>• "direct" - for the direct minimisation of the marginal deviance using <code>optim()</code>'s L-BFGS method</li> <li>• "em" - for the EM algorithm</li> <li>• "mixed" - combination of the direct and EM methods</li> <li>• "fixed" - for just obtaining the posterior regression function with fixed hyperparameters (default method when setting <code>fixed.hyp = TRUE</code>)</li> <li>• "canonical" - an efficient estimation method which takes advantage of the structure of the linear kernel</li> </ul>
control	<p>(Optional) A list of control options for the estimation procedure:</p> <p><code>maxit</code> The maximum number of iterations for the quasi-Newton optimisation or the EM algorithm. Defaults to 100.</p> <p><code>em.maxit</code> For <code>method = "mixed"</code>, the number of EM steps before switching to direct optimisation. Defaults to 5.</p> <p><code>stop.crit</code> The stopping criterion for the EM and L-BFGS algorithm, which is the difference in successive log-likelihood values. Defaults to <math>1e-8</math>.</p> <p><code>theta0</code> The initial values for the hyperparameters. Defaults to random starting values.</p> <p><code>report</code> The interval of reporting for the <code>optim()</code> function.</p> <p><code>restarts</code> The number of random restarts to perform. Defaults to 0. It's also possible to set it to TRUE, in which case the number of random restarts is set to the total number of available cores.</p> <p><code>no.cores</code> The number of cores in which to do random restarts. Defaults to the total number of available cores.</p> <p><code>omega</code> The overrelaxation parameter for the EM algorithm - a value between 0 and 1.</p>
interactions	Character vector to specify the interaction terms. When using formulas, this is specified automatically, so is not required. Syntax is "a:b" to indicate variable a interacts with variable b.
<code>est.lambda</code>	Logical. Estimate the scale parameters? Defaults to TRUE.
<code>est.hurst</code>	Logical. Estimate the Hurst coefficients for fBm kernels? Defaults to FALSE.
<code>est.lengthscale</code>	Logical. Estimate the lengthscales for SE kernels? Defaults to FALSE.
<code>est.offset</code>	Logical. Estimate the offsets for polynomial kernels? Defaults to FALSE.
<code>est.psi</code>	Logical. Estimate the error precision? Defaults to TRUE.



<code>fixed.hyp</code>	Logical. If TRUE, then no hyperparameters are estimated, i.e. all of the above <code>est.x</code> are set to FALSE, and vice versa. If NULL (default) then all of the <code>est.x</code> defaults are respected.
<code>lambda</code>	Initial/Default scale parameters. Relevant especially if <code>est.lambda = FALSE</code> .
<code>psi</code>	Initial/Default value for error precision. Relevant especially if <code>est.psi = FALSE</code> .
<code>nystrom</code>	Either logical or an integer indicating the number of Nystrom samples to take. Defaults to FALSE. If TRUE, then approximately 10% of the sample size is used for the Nystrom approximation.
<code>nys.seed</code>	The random seed for the Nystrom sampling. Defaults to NULL, which means the random seed is not fixed.
<code>formula</code>	The formula to fit when using formula interface.
<code>data</code>	Data frame containing variables when using formula interface.
<code>one.lam</code>	Logical. When using formula input, this is a convenient way of letting the function know to treat all variables as a single variable (i.e. shared scale parameter). Defaults to FALSE.

### Details

Uses a multicore loop to fit the folds by default, set `par.cv = FALSE` to not use multithreading.

### Value

An `iprior_xv` object containing a data frame of the cross-validated values such as the log-likelihood, training MSE and test MSE.

### Examples

```
## Not run:

# 5-fold CV experiment
(mod.cv <- iprior_cv(y ~ X, gen_smooth(100), kernel = "se", folds = 5))

# LOOCV experiment
(mod.cv <- iprior_cv(y ~ X, gen_smooth(100), kernel = "se", folds = Inf))

# Can also get root MSE
print(mod.cv, "RMSE")

## End(Not run)
```

---

iprior_package	iprior: <i>Regression using priors with Fisher information covariance kernels.</i>
----------------	--

---

### Description

The iprior package provides methods to perform and analyse I-prior regression models. Estimation is done either via direct optimisation of the log-likelihood or an EM algorithm.

### Author(s)

**Maintainer:** Haziq Jamil

Contributors:

- Wicher Bergsma

### See Also

<http://phd.haziqj.ml/>

---

is.iprior_x	<i>Test iprior objects</i>
-------------	----------------------------

---

### Description

Test whether an object is an ipriorMod, ipriorKernel, or either object with Nystrom method enabled.

### Usage

```
is.ipriorMod(x)
```

```
is.ipriorKernel(x)
```

```
is.nystrom(x)
```

### Arguments

x                    An ipriorMod or ipriorKernel object.

### Value

Logical.

---

is.kern_x	<i>Test kernel attributes</i>
-----------	-------------------------------

---

**Description**

Test whether an object uses a specific type of kernel.

**Usage**

```
is.kern_linear(x)
```

```
is.kern_canonical(x)
```

```
is.kern_fbm(x)
```

```
is.kern_pearson(x)
```

```
is.kern_se(x)
```

```
is.kern_poly(x)
```

**Arguments**

`x` An `ipriorMod` object, `ipriorKernel` object, a kernel matrix generated from one of the `kern_x()` functions, or even simply just a character vector.

**Value**

Logical.

---

kernel	<i>Reproducing kernels for the I-prior package</i>
--------	--

---

**Description**

The kernel functions used in this package are:

- The (canonical) linear kernel
- The fractional Brownian motion (fBm) kernel with Hurst index  $\gamma$
- The Pearson kernel
- The (scaled)  $d$ -degree polynomial kernel with offset  $c$
- The squared exponential (SE) kernel with lengthscale  $l$

**Usage**

```
kern_canonical(x, y = NULL, centre = TRUE)
```

```
kern_linear(x, y = NULL, centre = TRUE)
```

```
kern_pearson(x, y = NULL)
```

```
kern_fbm(x, y = NULL, gamma = 0.5, centre = TRUE)
```

```
kern_se(x, y = NULL, l = 1, centre = TRUE)
```

```
kern_poly(x, y = NULL, c = 0, d = 2, lam.poly = 1, centre = TRUE)
```

**Arguments**

x	A vector, matrix or data frame.
y	(Optional) vector, matrix or data frame. x and y must have identical column sizes.
centre	Logical. Whether to centre the data (default) or not.
gamma	The Hurst coefficient for the fBm kernel.
l	The lengthscale for the SE kernel.
c	The offset for the polynomial kernel. This is a value greater than zero.
d	The degree for the polynomial kernel. This is an integer value greater than or equal to two.
lam.poly	The scale parameter for the polynomial kernel.

**Details**

The Pearson kernel is used for nominal-type variables, and thus `factor`-type variables are treated with the Pearson kernel automatically when fitting I-prior models. The other kernels are for continuous variables, and each emits different properties of functions.

The linear kernel is used for "straight-line" functions. In addition, if squared, cubic, or higher order terms are to be modelled, then the polynomial kernel is suitable for this purpose. For smoothing models, the fBm kernel is preferred, although the SE kernel may be used as well.

**Value**

A matrix whose  $[i, j]$  entries are given by  $h(x[i], y[j])$ , with  $h$  being the appropriate kernel function. The matrix has dimensions  $m$  by  $n$  according to the lengths of  $y$  and  $x$  respectively. When a single argument  $x$  is supplied, then  $y$  is taken to be equal to  $x$ , and a symmetric  $n$  by  $n$  matrix is returned.

The matrix has a "kernel" attribute indicating which type of kernel function was called.

**References**

<http://phd.haziqj.ml/intro/>

**Examples**

```
kern_linear(1:3)
kern_fbm(1:5, 1:3, gamma = 0.7)
```

---

kernL

*Load the kernel matrices for I-prior models*


---

**Description**

Load the kernel matrices for I-prior models

**Usage**

```
kernL(y, ..., kernel = "linear", interactions = NULL, est.lambda = TRUE,
      est.hurst = FALSE, est.lengthscale = FALSE, est.offset = FALSE,
      est.psi = TRUE, fixed.hyp = NULL, lambda = 1, psi = 1,
      nystrom = FALSE, nys.seed = NULL, model = list(), train.samp, test.samp)
```

```
## S3 method for class 'formula'
```

```
kernL(formula, data, kernel = "linear", one.lam = FALSE,
      est.lambda = TRUE, est.hurst = FALSE, est.lengthscale = FALSE,
      est.offset = FALSE, est.psi = TRUE, fixed.hyp = NULL, lambda = 1,
      psi = 1, nystrom = FALSE, nys.seed = NULL, model = list(), train.samp,
      test.samp, ...)
```

**Arguments**

y	Vector of response variables
...	Only used when fitting using non-formula, enter the variables (vectors or matrices) separated by commas.
kernel	Character vector indicating the type of kernel for the variables. Available choices are: <ul style="list-style-type: none"> <li>• "linear" - (default) for the linear kernel</li> <li>• "canonical" - alternative name for "linear"</li> <li>• "fbm", "fbm,0.5" - for the fBm kernel with Hurst coefficient 0.5 (default)</li> <li>• "se", "se,1" - for the SE kernel with lengthscale 1 (default)</li> <li>• "poly", "poly2", "poly2,0" - for the polynomial kernel of degree 2 with offset 0 (default)</li> <li>• "pearson" - for the Pearson kernel</li> </ul>

The kernel argument can also be a vector of length equal to the number of variables, therefore it is possible to specify different kernels for each variables. Note that factor type variables are assigned the Pearson kernel by default, and that non-factor types can be forced to use the Pearson kernel (not recommended).

<code>interactions</code>	Character vector to specify the interaction terms. When using formulas, this is specified automatically, so is not required. Syntax is "a:b" to indicate variable a interacts with variable b.
<code>est.lambda</code>	Logical. Estimate the scale parameters? Defaults to TRUE.
<code>est.hurst</code>	Logical. Estimate the Hurst coefficients for fBm kernels? Defaults to FALSE.
<code>est.lengthscale</code>	Logical. Estimate the lengthscales for SE kernels? Defaults to FALSE.
<code>est.offset</code>	Logical. Estimate the offsets for polynomial kernels? Defaults to FALSE.
<code>est.psi</code>	Logical. Estimate the error precision? Defaults to TRUE.
<code>fixed.hyp</code>	Logical. If TRUE, then no hyperparameters are estimated, i.e. all of the above <code>est.x</code> are set to FALSE, and vice versa. If NULL (default) then all of the <code>est.x</code> defaults are respected.
<code>lambda</code>	Initial/Default scale parameters. Relevant especially if <code>est.lambda = FALSE</code> .
<code>psi</code>	Initial/Default value for error precision. Relevant especially if <code>est.psi = FALSE</code> .
<code>nystrom</code>	Either logical or an integer indicating the number of Nystrom samples to take. Defaults to FALSE. If TRUE, then approximately 10% of the sample size is used for the Nystrom approximation.
<code>nys.seed</code>	The random seed for the Nystrom sampling. Defaults to NULL, which means the random seed is not fixed.
<code>model</code>	DEPRECATED.
<code>train.samp</code>	(Optional) A vector indicating which of the data points should be used for training, and the remaining used for testing.
<code>test.samp</code>	(Optional) Similar to <code>train.samp</code> , but on test samples instead.
<code>formula</code>	The formula to fit when using formula interface.
<code>data</code>	Data frame containing variables when using formula interface.
<code>one.lam</code>	Logical. When using formula input, this is a convenient way of letting the function know to treat all variables as a single variable (i.e. shared scale parameter). Defaults to FALSE.

### Value

An `ipriorKernel` object which contains the relevant material to be passed to the `iprior` function for model fitting.

### See Also

[iprior](#)

### Examples

```
str(ToothGrowth)
(mod <- kernL(y = ToothGrowth$len,
             supp = ToothGrowth$supp,
             dose = ToothGrowth$dose,
```

```

        interactions = "1:2"))
kernL(len ~ supp * dose, data = ToothGrowth) # equivalent formula call

# Choosing different kernels
str(stackloss)
kernL(stack.loss ~ ., stackloss, kernel = "fbm") # all fBm kernels
kernL(stack.loss ~ ., stackloss, kernel = "FBm") # cApS dOn't MatTeR
kernL(stack.loss ~ ., stackloss,
       kernel = c("linear", "se", "poly3")) # different kernels

# Sometimes the print output is too long, can use str() options here
print(mod, strict.width = "cut", width = 30)

```

---

logLik.ipriorMod	<i>Obtain the log-likelihood and deviance of an I-prior model</i>
------------------	---

---

## Description

This function calculates the log-likelihood value or deviance (twice the negative log-likelihood) for I-prior models. It works for both `ipriorMod` and `ipriorKernel` class objects.

## Usage

```

## S3 method for class 'ipriorMod'
logLik(object, theta = NULL, ...)

## S3 method for class 'ipriorMod'
deviance(object, theta = NULL, ...)

## S3 method for class 'ipriorKernel'
logLik(object, theta = NULL, ...)

## S3 method for class 'ipriorKernel'
deviance(object, theta = NULL, ...)

```

## Arguments

object	An object of class <code>ipriorMod</code> or <code>ipriorKernel</code> .
theta	(Optional) Evaluates the log-likelihood at theta.
...	Not used.

## Details

For `ipriorKernel` objects, the log-likelihood or deviance is calculated at the default parameter values: scale parameters and error precision are equal to one, while hyperparameters of the kernels (e.g. Hurst index, lengthscale, etc.) are the default values (see [here](#) for details) or ones that has been

specified. For `ipriorMod` objects, the log-likelihood or deviance is calculated at the last obtained value from the estimation method.

For both types of objects, it is possible to supply parameter values at which to calculate the log-likelihood/deviance. This makes estimating an I-prior model more flexible, by first loading the variables into an `ipriorKernel` object, and then using an optimiser such as `optim`. Parameters have been transformed so that they can be optimised unconstrained.

### See Also

[check\\_theta](#).

---

plot.ipriorMod	<i>Plots for I-prior models</i>
----------------	---------------------------------

---

### Description

There are three types of plots that are currently written in the package:

`plot_fitted` Plot the fitted regression line with credibility bands.

`plot_predict` Plot residuals against fitted values.

`plot_iter` Plot the progression of the log-likelihood value over time.

The S3 method `plot` for class `ipriorMod` currently returns `plot_fitted`.

### Usage

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

plot_resid(x)

plot_fitted_multilevel(x, X.var = 1, grp.var = 1, facet = c(2, 3),
  cred.bands = TRUE, show.legend = TRUE, show.points = TRUE,
  x.lab = NULL, y.lab = NULL, grp.lab = NULL, extrapolate = FALSE)

plot_fitted(x, X.var = 1, cred.bands = TRUE, size = 1,
  linetype = "solid")

plot_iter(x, niter.plot = NULL, lab.pos = c("up", "down"))

plot_ppc(x, draws = 100)
```



**Arguments**

x	An ipriorMod object.
...	Not used
X.var	The index of the X variable to plot.
grp.var	Index of the grouping variable for multilevel plots.
facet	The index of the X variable in which to facet. This is a vector of maximum length 2.
cred.bands	Logical. Plot the confidence intervals? Defaults to TRUE.
show.legend	Logical. Show legend?
show.points	Logical. Show data points?
x.lab	(Optional) X axis label.
y.lab	(Optional) Y axis label.
grp.lab	(Optional) The name for the groups, which is also the legend title.
extrapolate	Logical. Extend the fitted regression line to fill the plot?
size	Size of the fitted line
linetype	Type of the fitted line
niter.plot	(Optional) Vector of length at most two, indicating the start and end points of the iterations to plot.
lab.pos	Adjust the position of the log-likelihood label.
draws	Number of draws for posterior predictive check.
grp	The index of the groups.

**Details**

See `ggplot2` documentation for the plotting parameters.

---

pollution	<i>Air pollution and mortality</i>
-----------	------------------------------------

---

**Description**

Data on the relation between weather, socioeconomic, and air pollution variables and mortality rates in 60 Standard Metropolitan Statistical Areas (SMSAs) of the USA, for the years 1959-1961.

**Usage**

```
pollution
```

**Format**

A data frame of 16 observations on 16 variables.

Mortality Total age-adjusted mortality rate per 100,000.

Rain Mean annual precipitation in inches.

Humid Mean annual precipitation in inches.

JanTemp Mean annual precipitation in inches.

JulTemp Mean annual precipitation in inches.

Over65 Mean annual precipitation in inches.

Popn Mean annual precipitation in inches.

Educ Mean annual precipitation in inches.

Hous Mean annual precipitation in inches.

Dens Mean annual precipitation in inches.

NonW Mean annual precipitation in inches.

WhiteCol Mean annual precipitation in inches.

Poor Mean annual precipitation in inches.

HC Mean annual precipitation in inches.

NOx Mean annual precipitation in inches.

SO2 Mean annual precipitation in inches.

**References**

McDonald, G. C. and Schwing, R. C. (1973). Instabilities of regression estimates relating air pollution to mortality. *Technometrics*, 15(3):463-481.

**Examples**

```
data(pollution)
str(pollution)
```

---

predict

*Obtain predicted values from ipriorMod objects*

---

**Description**

Obtain predicted values from ipriorMod objects

**Usage**

```
## S3 method for class 'ipriorMod'
fitted(object, intervals = FALSE, alpha = 0.05, ...)

## S3 method for class 'ipriorMod'
predict(object, newdata = list(), y.test = NULL,
        intervals = FALSE, alpha = 0.05, ...)

## S3 method for class 'ipriorPredict'
print(x, rows = 10, dp = 3, ...)
```

**Arguments**

object, x	An ipriorMod object.
intervals	Logical. Calculate the credibility intervals for the fitted values. Defaults to FALSE.
alpha	The significance level for the credibility intervals. This is a number between 0 and 1.
...	Not used.
newdata	Either a data frame when using formula method, or a list of vectors/matrices if using default method. Either way, the new data must be structurally similar to the original data used to fit the model.
y.test	(Optional) Test data, in order to compute test error rates.
rows	(Optional) The number of values/rows to display.
dp	(Optional) Decimal places for the values.

**Value**

A list of class `ipriorPredict` containing the fitted values, residuals (observed minus fitted), the training mean squared error, and the lower and upper intervals (if called).

**Examples**

```
dat <- gen_smooth(20)
mod <- iprior(y ~ ., dat, kernel = "se")
fitted(mod)
fitted(mod, intervals = TRUE)
predict(mod, gen_smooth(5))

with(dat, mod <-< iprior(y, X, kernel = "poly"))
newdat <- gen_smooth(30)
mod.pred <- predict(mod, list(newdat$X), y.test = newdat$y, intervals = TRUE)
str(mod.pred)
print(mod.pred, row = 5)
```

---

sigma	<i>Obtain the standard deviation of the residuals 'sigma'</i>
-------	---

---

**Description**

Extract the standard deviation of the residuals. For I-prior models, this is  $\sigma = 1 / \sqrt{\psi}$ .

**Usage**

```
## S3 method for class 'ipriorMod'
sigma(object, ...)
```

**Arguments**

object	An object of class ipriorMod.
...	Not used.

---

summary.ipriorMod	<i>Print and summary method for I-prior models</i>
-------------------	--

---

**Description**

Print and summary method for I-prior models

**Usage**

```
## S3 method for class 'ipriorMod'
print(x, digits = 5, ...)
```

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

**Arguments**

digits	Number of decimal places for the printed coefficients.
...	Not used.
object, x	An ipriorMod object.

---

 tecator.cv

*Results of I-prior cross-validation experiment on Tecator data set*


---

## Description

Results of I-prior cross-validation experiment on Tecator data set

## Usage

```
tecator.cv
```

## Format

Results from `iprior_cv` cross validation experiment. This is a list of seven, with each component bearing the results for the linear, quadratic, cubic, fBm-0.5, fBm-MLE and SE I-prior models. The seventh is a summarised table of the results.

## Details

For the fBm and SE kernels, it seems numerical issues arise when using a direct optimisation approach. Terminating the algorithm early (say using a relaxed stopping criterion) seems to help.

## Examples

```
# Results from the six experiments
print(tecator.cv[[1]], "RMSE")
print(tecator.cv[[2]], "RMSE")
print(tecator.cv[[3]], "RMSE")
print(tecator.cv[[4]], "RMSE")
print(tecator.cv[[5]], "RMSE")
print(tecator.cv[[6]], "RMSE")

# Summary of results
print(tecator.cv[[7]])

## Not run:

# Prepare data set
data(tecator, package = "caret")
endpoints <- as.data.frame(endpoints)
colnames(endpoints) <- c("water", "fat", "protein")
absorp <- -t(diff(t(absorp))) # this takes first differences using diff()
fat <- endpoints$fat

# Here is the code to replicate the results
mod1.cv <- iprior_cv(fat, absorp, folds = Inf)
mod2.cv <- iprior_cv(fat, absorp, folds = Inf, kernel = "poly2",
  est.offset = TRUE)
mod3.cv <- iprior_cv(fat, absorp, folds = Inf, kernel = "poly3",
```

```

        est.offset = TRUE)
mod4.cv <- iprior_cv(fat, absorp, method = "em", folds = Inf, kernel = "fbm",
                    control = list(stop.crit = 1e-2))
mod5.cv <- iprior_cv(fat, absorp, folds = Inf, kernel = "fbm",
                    est.hurst = TRUE, control = list(stop.crit = 1e-2))
mod6.cv <- iprior_cv(fat, absorp, folds = Inf, kernel = "se",
                    est.lengthscale = TRUE, control = list(stop.crit = 1e-2))

tecator_res_cv <- function(mod) {
  res <- as.numeric(apply(mod$res[, -1], 2, mean)) # Calculate RMSE
  c("Training RMSE" = res[1], "Test RMSE" = res[2])
}

tecator_tab_cv <- function() {
  tab <- t(sapply(list(mod1.cv, mod2.cv, mod3.cv, mod4.cv, mod5.cv, mod6.cv),
                 tecator_res_cv))
  rownames(tab) <- c("Linear", "Quadratic", "Cubic", "fBm-0.5", "fBm-MLE",
                   "SE-MLE")

  tab
}

tecator.cv <- list(
  "linear" = mod1.cv,
  "quadratic" = mod2.cv,
  "cubic" = mod3.cv,
  "fbm-0.5" = mod4.cv,
  "fbm-MLE" = mod5.cv,
  "SE" = mod6.cv,
  "summary" = tecator_tab_cv()
)

## End(Not run)

```

---

update.ipriorMod

*Update an I-prior model*


---

## Description

Update an I-prior model

## Usage

```

## S3 method for class 'ipriorMod'
update(object, method = NULL, control = list(),
       iter.update = 100, ...)

```

**Arguments**

<code>object</code>	An <code>ipriorMod</code> object.
<code>method</code>	An optional method. See <a href="#">here</a> for details.
<code>control</code>	An optional list of controls for the estimation procedure. See <a href="#">here</a> for details.
<code>iter.update</code>	The number of additional iterations to update the I-prior model.
<code>...</code>	Not used.

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