

Package ‘ncvreg’

November 6, 2025

Title Regularization Paths for SCAD and MCP Penalized Regression Models

Version 3.16.0

Description Fits regularization paths for linear regression, GLM, and Cox regression models using lasso or nonconvex penalties, in particular the minimax concave penalty (MCP) and smoothly clipped absolute deviation (SCAD) penalty, with options for additional L2 penalties (the “elastic net” idea). Utilities for carrying out cross-validation as well as post-fitting visualization, summarization, inference, and prediction are also provided. For more information, see Breheny and Huang (2011) <doi:10.1214/10-AOAS388> or visit the ncvreg homepage <<https://pbreheny.github.io/ncvreg/>>.

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URL <https://pbreheny.github.io/ncvreg/>,
<https://github.com/pbreheny/ncvreg>

BugReports <https://github.com/pbreheny/ncvreg/issues>

Depends R (>= 4.1)

Suggests ashr, knitr, parallel, rmarkdown, survival, tinytest

LazyData true

VignetteBuilder knitr

Encoding UTF-8

RoxygenNote 7.3.3

NeedsCompilation yes

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Repository CRAN

Date/Publication 2025-11-06 06:10:33 UTC

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assign_fold	<i>Assign folds for cross-validation</i>
-------------	--

Description

If `y` has only two unique values, fold assignments are chosen so that the balance between outcomes is the same in each fold. This is useful for logistic regression and time-to-event data (to balance the fraction of observations that are censored).

Usage

```
assign_fold(y, folds, seed)
```

Arguments

<code>y</code>	Either (i) the vector of outcomes or (ii) a vector such as <code>1:n</code>
<code>folds</code>	Number of folds
<code>seed</code>	(optional) set a seed for reproducibility

Value

A vector of integers indicating fold assignments

See Also

`[cv.ncvreg()]`

Examples

```
assign_fold(rnorm(11), 2)
assign_fold(1:41, 7)
assign_fold(1:41, 7) |> table()
data(Heart)
assign_fold(Heart$y, 7) |> head()
assign_fold(Heart$y, 7) |> table(Heart$y)
```

AUC.cv.ncvsurv

AUC for cv.ncvsurv objects

Description

Calculates the cross-validated AUC (concordance) from a `cv.ncvsurv` object.

Usage

```
## S3 method for class 'cv.ncvsurv'
AUC(obj, ...)
```

Arguments

<code>obj</code>	A <code>cv.ncvsurv</code> object. You must run <code>cv.ncvsurv()</code> with the option <code>returnY=TRUE</code> in order for <code>AUC()</code> to work.
<code>...</code>	For S3 method compatibility; not used

Details

The area under the curve (AUC), or equivalently, the concordance statistic (C), is calculated according to the procedure described in van Houwelingen and Putter (2011). The function calls `survival::concordancefit()`, except cross-validated linear predictors are used to guard against overfitting. Thus, the values returned by `AUC.cv.ncvsurv()` will be lower than those you would obtain with `concordancefit()` if you fit the full (unpenalized) model.

Author(s)

Patrick Breheny, Brandon Butcher, and Lawrence Hunsicker

References

van Houwelingen H, Putter H (2011). Dynamic Prediction in Clinical Survival Analysis. CRC Press.

See Also

[cv.ncvsurv\(\)](#), [survival::concordancefit\(\)](#)

Examples

```
data(Lung)
X <- Lung$X
y <- Lung$y

cvfit <- cv.ncvsurv(X, y, returnY=TRUE)
head(AUC(cvfit))
lam <- cvfit$lambda
plot(lam, AUC(cvfit), xlim=rev(range(lam)), lwd=3, type='l',
     las=1, xlab=expression(lambda), ylab='AUC')
```

cv.ncvreg

Cross-validation for ncvreg/ncvsurv

Description

Performs k-fold cross validation for MCP- or SCAD-penalized regression models over a grid of values for the regularization parameter lambda.

Usage

```
cv.ncvreg(
  X,
  y,
  ...,
  cluster,
  nfolds = 10,
  fold,
  returnY = FALSE,
  trace = FALSE
)

cv.ncvsurv(
  X,
  y,
  ...,
  cluster,
  nfolds = 10,
  fold,
```

```

    se = c("quick", "bootstrap"),
    returnY = FALSE,
    trace = FALSE
  )

```

Arguments

<code>X</code>	The design matrix, without an intercept, as in <code>ncvreg()</code> or <code>ncvsurv()</code> .
<code>y</code>	The response, as in <code>ncvreg()</code> or <code>ncvsurv()</code> .
<code>...</code>	Additional arguments to <code>ncvreg()</code> or <code>ncvsurv()</code> .
<code>cluster</code>	<code>cv.ncvreg()</code> and <code>cv.ncvsurv()</code> can be run in parallel across a cluster using the parallel package. The cluster must be set up in advance using the <code>parallel::makeCluster()</code> function from that package. The cluster must then be passed to <code>cv.ncvreg()</code> or <code>cv.ncvsurv()</code> (see example).
<code>nfolds</code>	The number of cross-validation folds. Default is 10.
<code>fold</code>	Which fold each observation belongs to. By default the observations are randomly assigned.
<code>returnY</code>	Should <code>cv.ncvreg()/cv.ncvsurv()</code> return the linear predictors from the cross-validation folds? Default is FALSE; if TRUE, this will return a matrix in which the element for row <i>i</i> , column <i>j</i> is the fitted value for observation <i>i</i> from the fold in which observation <i>i</i> was excluded from the fit, at the <i>j</i> th value of <i>lambda</i> . NOTE: For <code>cv.ncvsurv()</code> , the rows of <i>Y</i> are ordered by time on study, and therefore will not correspond to the original order of observations passed to <code>cv.ncvsurv()</code> .
<code>trace</code>	If set to TRUE, inform the user of progress by announcing the beginning of each CV fold. Default is FALSE.
<code>se</code>	For <code>cv.ncvsurv()</code> , the method by which the cross-validation standard error (CVSE) is calculated. The 'quick' approach is based on a rough approximation, but can be calculated more or less instantly. The 'bootstrap' approach is more accurate, but requires additional computing time.

Details

The function calls `ncvreg/ncvsurv` `nfolds` times, each time leaving out $1/nfolds$ of the data. The cross-validation error is based on the deviance; [see here for more details](#).

For `family="binomial"` models, the cross-validation fold assignments are balanced across the 0/1 outcomes, so that each fold has the same proportion of 0/1 outcomes (or as close to the same proportion as it is possible to achieve if cases do not divide evenly).

For Cox models, `cv.ncvsurv()` uses the approach of calculating the full Cox partial likelihood using the cross-validated set of linear predictors. Other approaches to cross-validation for the Cox regression model have been proposed in the literature; the strengths and weaknesses of the various methods for penalized regression in the Cox model are the subject of current research. A simple approximation to the standard error is provided, although an option to bootstrap the standard error (`se='bootstrap'`) is also available.

Value

An object with S3 class `cv.ncvreg` or `cv.ncvsurv` containing:

cve The error for each value of `lambda`, averaged across the cross-validation folds.

cvse The estimated standard error associated with each value of `cve`.

fold The fold assignments for cross-validation for each observation; note that for `cv.ncvsurv()`, these are in terms of the ordered observations, not the original observations.

lambda The sequence of regularization parameter values along which the cross-validation error was calculated.

fit The fitted `ncvreg()` or `ncvsurv()` object for the whole data.

min The index of `lambda` corresponding to `lambda.min`.

lambda.min The value of `lambda` with the minimum cross-validation error.

null.dev The deviance for the intercept-only model. If you have supplied your own `lambda` sequence, this quantity may not be meaningful.

Bias The estimated bias of the minimum cross-validation error, as in Tibshirani and Tibshirani (2009) [doi:10.1214/08AOAS224](https://doi.org/10.1214/08AOAS224)

pe If `family="binomial"`, the cross-validation prediction error for each value of `lambda`.

Y If `returnY=TRUE`, the matrix of cross-validated fitted values (see above).

Author(s)

Patrick Breheny; Grant Brown helped with the parallelization support

References

Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, **5**: 232-253. [doi:10.1214/10AOAS388](https://doi.org/10.1214/10AOAS388)

See Also

`ncvreg()`, `plot.cv.ncvreg()`, `summary.cv.ncvreg()`

Examples

```
data(Prostate)

cvfit <- cv.ncvreg(Prostate$X, Prostate$y)
plot(cvfit)
summary(cvfit)

fit <- cvfit$fit
plot(fit)
beta <- fit$beta[,cvfit$min]

## requires loading the parallel package
## Not run:
```

```

library(parallel)
X <- Prostate$X
y <- Prostate$y
cl <- makeCluster(4)
cvfit <- cv.ncvreg(X, y, cluster=cl, nfolds=length(y))
## End(Not run)

# Survival
data(Lung)
X <- Lung$X
y <- Lung$y

cvfit <- cv.ncvsurv(X, y)
summary(cvfit)
plot(cvfit)
plot(cvfit, type="rsq")

```

Heart

Risk factors associated with heart disease

Description

Data from a subset of the Coronary Risk-Factor Study baseline survey, carried out in rural South Africa.

Usage

Heart

Format

A list of two objects: `y` and `X`

y Coronary heart disease at baseline; 1=Yes 0=No

X A matrix with 462 observations (rows) and 9 predictor variables (columns). The remainder of this list describes the columns of `X`

sbp Systolic blood pressure

tobacco Cumulative tobacco consumption, in kg

ldl Low-density lipoprotein cholesterol

adiposity Adipose tissue concentration

famhist Family history of heart disease (1=Present, 0=Absent)

typea Score on test designed to measure type-A behavior

obesity Obesity

alcohol Current consumption of alcohol

age Age of subject

Source

<https://web.stanford.edu/~hastie/ElemStatLearn/>

References

- Hastie T, Tibshirani R, and Friedman J. (2001). *The Elements of Statistical Learning*. Springer.
- Rousseauw J, et al. (1983). Coronary risk factor screening in three rural communities. *South African Medical Journal*, **64**: 430-436.

intervals

Projection base test statistics and intervals

Description

Constructs projection based test statistics that can be used to control FDR along with intervals for a penalized regression model.

Usage

```
intervals(
  fit,
  lambda,
  sigma,
  level = 0.95,
  posterior = TRUE,
  relaxed = FALSE,
  adjust_projection = FALSE,
  X = NULL
)
```

Arguments

<code>fit</code>	An optional fit of class <code>ncvreg</code> or <code>cv.ncvreg</code> . If supplied, <code>X</code> should only be supplied if <code>fit</code> does not contain it.
<code>lambda</code>	The penalty at which the tests and intervals are to be constructed. If left unspecified, will be selected using cross validation.
<code>sigma</code>	Standard deviation estimate used to compute test statistic and intervals. If left unspecified (default) it will be estimated using the recommendation from Reid et al. (2016)
<code>level</code>	the confidence level required.
<code>posterior</code>	whether the intervals returned should be posterior intervals (default) or debiased intervals (if <code>FALSE</code>). Posterior intervals are constructed from distributions where the coefficient estimates are the is the posterior mode. Debiased intervals are constructed around the estimates.

<code>relaxed</code>	whether the relaxed lasso based statistic / intervals should be used. Default is FALSE in which case PIPE based intervals are constructed (recommended). This affects the estimate.
<code>adjust_projection</code>	whether a Local Quadratic Approximation should be used in the projection for determining variance of the PIPE based test statistic. Default is FALSE as more research has been done without this adjustment, however without this adjustment, statistics and intervals may be over conservative in the presence of correlation. This affects the SE.
<code>X</code>	The original design matrix supplied to <code>fit</code> . Required if <code>fit</code> does not contain <code>X</code> .

Details

The function constructs test statistics and intervals based off an approximate projection onto the column space of the active features. The test statistic can be used to control FDR and the intervals generally have good coverage. However, both tend to be conservative with the introduction of correlation.

The intervals produced can either be biased (like the point estimates) or debiased by setting the parameter `posterior` accordingly. The resulting behavior is quite different. See references for more details.

Value

An `data.frame` containing the following columns:

- variable** `colnames(X)`
- coef** The original estimates at the specified parameters (`lambda`, `gamma`, `alpha`)
- estimate** The debiased estimates.
- SE** The standard errors.
- t** The PIPE / Relaxed Lasso / LQA test statistics
- lower** Interval lower bounds
- upper** Intervals upper bounds
- p.value** The unadjusted p-value
- p.adjust** The Benjamini and Hochberg corrected p-value
- penalty** The penalty used.
- lambda** The lambda value the test statistics and intervals were constructed at.
- gamma** The gamma value the test statistics and intervals were constructed at (for MCP/SCAD).
- alpha** The alpha value the test statistics and intervals were constructed at.
- level** The confidence level set for interval construction.
- sigma** The standard deviation used for constructing the test statistics and intervals.

Author(s)

Logan Harris, Patrick Breheny, and Biyue Dai

References

- Harris L and Breheny P. (2025) A new perspective on high dimensional confidence intervals. *arXiv preprint*, arXiv:2508.03504. <https://arxiv.org/abs/2508.03504>
- Harris L and Breheny P. (2025) Alternative Likelihood Approximations for High-Dimensional Intervals for Lasso. *arXiv preprint*, arXiv:2509.14971. <https://arxiv.org/abs/2509.14971>
- Dai B. (2019) Projection-based inference and model selection for penalized regression. PhD dissertation, University of Iowa, Iowa City, IA. [doi:10.17077/etd.005250](https://doi.org/10.17077/etd.005250)

Examples

```
# Linear regression (SCAD-Net penalty, PIPE intervals, pass ncvreg object)
fit <- ncvreg(Prostate$X, Prostate$y, penalty = "SCAD", alpha = 0.9)
intervals(fit) |> head()

# Logistic regression (lasso penalty, LQA intervals, pass cv.ncvreg object)
data(Heart)
cv_fit <- cv.ncvreg(Heart$X, Heart$y, family="binomial", penalty = "lasso")
intervals(cv_fit, adjust_projection = TRUE) |> head()
```

local_mfdr	<i>Estimate local mFDR for all features</i>
------------	---

Description

local_mfdr() is called by `summary.ncvreg()`, which typically offers a more convenient interface to users. If, however, you are working with local mfdRs programmatically rather than interactively, you probably want to use local_mfdr(), which skips the sorting, filtering, and print formatting of `summary.ncvreg()`.

Usage

```
local_mfdr(
  fit,
  lambda,
  X = NULL,
  y = NULL,
  method = c("ashr", "kernel"),
  sigma,
  ...
)
```

Arguments

fit	A fitted ncvreg or ncvsurv object.
lambda	The value of lambda at which inference should be carried out.

<code>X, y</code>	The design matrix and response used to fit the model; in most cases, it is not necessary to provide <code>X</code> and <code>y</code> as they are returned by <code>ncvreg</code> , but see the <code>returnX</code> argument in <code>ncvreg()</code> .
<code>method</code>	What method should be used to calculate the local <code>fdr</code> ? Options are <code>ashr</code> (which tends to be more accurate) and <code>kernel</code> (which requires no additional packages). The default is to use <code>ashr</code> if the package is installed.
<code>sigma</code>	For linear regression models, users can supply an estimate of the residual standard deviation. The default is to use <code>RSS / DF</code> , where degrees of freedom are approximated using the number of nonzero coefficients.
<code>...</code>	Additional arguments to <code>ashr::ash()</code> if using <code>method='ashr'</code> .

Value

If all features are penalized, then the object returns a data frame with one row per feature and four columns:

- `Estimate`: The coefficient estimate from the penalized regression fit
- `z`: A test statistic that approximately follows a standard normal distribution under the null hypothesis that the feature is marginally independent of the outcome
- `mfdr`: The estimated marginal local false discovery rate
- `Selected`: Features with nonzero coefficient estimates are given an asterisk

If some features are penalized and others are not, then a list is returned with two elements: `pen.vars`, which consists of the data frame described above, and `unpen.vars`, a data frame with four columns: `Estimate`, `SE`, `Statistic`, and `p.value`. The standard errors and p-values are based on a classical `lm/glm/coxph` model using the effect of the penalized features as an offset.

See Also

[summary.ncvreg\(\)](#)

Examples

```
# Linear regression
data(Prostate)
fit <- ncvreg(Prostate$X, Prostate$y)
local_mfdr(fit, 0.1)

fit <- ncvreg(Prostate$X, Prostate$y, penalty.factor=rep(0:1, each=4))
local_mfdr(fit, 0.1)

# Logistic regression
data(Heart)
X <- Heart$X
y <- Heart$y
fit <- ncvreg(X, y, family='binomial')
local_mfdr(fit, 0.1)

# Cox regression
```

```
data(Lung)
X <- Lung$X
y <- Lung$y
fit <- ncvsurv(X, y)
local_mfdr(fit, 0.1)
```

logLik.ncvreg

Extract Log-Likelihood

Description

Extract the log-likelihood of an ncvreg or ncvsurv object.

Usage

```
## S3 method for class 'ncvreg'
logLik(object, REML = FALSE, ...)

## S3 method for class 'ncvsurv'
logLik(object, ...)
```

Arguments

object	An ncvreg or ncvsurv object, as obtained from ncvreg() or ncvsurv()
REML	As in logLik.lm()
...	For S3 compatibility

See Also

[logLik\(\)](#)

Lung

VA lung cancer data set

Description

Data from a randomised trial of two treatment regimens for lung cancer. This is a standard survival analysis data set from the classic textbook by Kalbfleisch and Prentice.

Usage

Lung

Format

A list of two objects: `y` and `X`

y A two column matrix (Surv object) containing the follow-up time (in days) and an indicator variable for whether the patient died while on the study or not.

X A matrix with 137 observations (rows) and 9 predictor variables (columns). The remainder of this list describes the columns of `X`

trt Treatment indicator (1=control group, 2=treatment group)

karno Karnofsky performance score (0=bad, 100=good)

diagtime Time from diagnosis to randomization (months)

age Age (years, at baseline)

prior Prior therapy (0=no, 1=yes)

squamous Indicator for whether the cancer type is squamous cell carcinoma (0=no, 1=yes)

small Indicator for whether the cancer type is small cell lung cancer (0=no, 1=yes)

adeno Indicator for whether the cancer type is adenocarcinoma (0=no, 1=yes)

large Indicator for whether the cancer type is large cell carcinoma (0=no, 1=yes)

Source

<https://cran.r-project.org/package=survival>

References

- Kalbfleisch D and Prentice RL (1980), *The Statistical Analysis of Failure Time Data*. Wiley, New York.

See Also

[ncvsurv\(\)](#)

mfdr

Marginal false discovery rates

Description

Estimates the marginal false discovery rate (mFDR) of a penalized regression model.

Usage

```
mfdr(fit, X)
```

Arguments

<code>fit</code>	An <code>ncvreg</code> or <code>ncvsurv</code> object.
<code>X</code>	The model matrix corresponding to <code>fit</code> . This is not necessary for linear regression, but in logistic and Cox regression, the mFDR depends on <code>X</code> . It is not necessary to supply <code>X</code> if it is already contained in <code>fit</code> ; i.e., if <code>ncvreg/ncvsurv</code> was run with <code>returnX=TRUE</code> .

Details

The function estimates the marginal false discovery rate (mFDR) for a penalized regression model. The estimate tends to be accurate in most settings, but will be slightly conservative if predictors are highly correlated. For an alternative way of estimating the mFDR, typically more accurate in highly correlated cases, see [perm.ncvreg\(\)](#).

Value

An object with S3 class `mfdr` inheriting from `data.frame`, containing:

EF The number of variables selected at each value of `lambda`, averaged over the permutation fits.

S The actual number of selected variables for the non-permuted data.

mFDR The estimated marginal false discovery rate (EF/S).

Author(s)

Patrick Breheny and Ryan Miller

See Also

[ncvreg\(\)](#), [ncvsurv\(\)](#), [plot.mfdr\(\)](#), [perm.ncvreg\(\)](#)

Examples

```
# Linear regression -----
data(Prostate)
fit <- ncvreg(Prostate$X, Prostate$y)

obj <- mfdr(fit)
obj[1:10,]

# Comparison with perm.ncvreg
op <- par(mfrow=c(2,2))
plot(obj)
plot(obj, type="EF")
pmfit <- perm.ncvreg(Prostate$X, Prostate$y)
plot(pmfit)
plot(pmfit, type="EF")
par(op)

# Logistic regression -----
data(Heart)
```

```

fit <- ncvreg(Heart$X, Heart$y, family="binomial")
obj <- mfdr(fit)
head(obj)
op <- par(mfrow=c(1,2))
plot(obj)
plot(obj, type="EF")
par(op)

# Cox regression -----
data(Lung)
fit <- ncvsurv(Lung$X, Lung$y)
obj <- mfdr(fit)
head(obj)
op <- par(mfrow=c(1,2))
plot(obj)
plot(obj, type="EF")
par(op)

```

ncvfit

Direct interface for nonconvex penalized regression (non-pathwise)

Description

This function is intended for users who know exactly what they're doing and want complete control over the fitting process: no standardization is applied, no intercept is included, no path is fit. All of these things are best practices for data analysis, so if you are choosing not to do them, you are on your own – there is no guarantee that your results will be meaningful. Some things in particular that you should pay attention to:

- If your model has an intercept, it is up to you to (un)penalize it properly, typically by settings its corresponding element of `penalty.factor` to zero.
- You should provide initial values for the coefficients; in nonconvex optimization, initial values are very important in determining which local solution an algorithm converges to.

Usage

```

ncvfit(
  X,
  y,
  init = rep(0, ncol(X)),
  r,
  xtx,
  penalty = c("MCP", "SCAD", "lasso"),
  gamma = switch(penalty, SCAD = 3.7, 3),
  alpha = 1,
  lambda,
  eps = 1e-05,
  max.iter = 1000,

```

```

    penalty.factor = rep(1, ncol(X)),
    warn = TRUE
  )

```

Arguments

<code>X</code>	Design matrix; no intercept will be added, no standardization will occur (n x p matrix)
<code>y</code>	Response vector (length n vector)
<code>init</code>	Initial values for beta. Default: zero (length p vector)
<code>r</code>	Residuals corresponding to <code>init</code> ; these will be calculated if not supplied, but if they have already been calculated elsewhere, it is more efficient to pass them as an argument. WARNING: If you supply an incorrect value of <code>r</code> , the solution will be incorrect. (length n vector)
<code>xtx</code>	X scales: the <code>j</code> th element should equal <code>crossprod(X[, j])/n</code> . These will be calculated if not supplied, but if they have already been calculated elsewhere, it is more efficient to pass them as an argument. In particular, if <code>X</code> is standardized, one should pass <code>xtx = rep(1, p)</code> . WARNING: If you supply an incorrect value of <code>xtx</code> , the solution will be incorrect. (length p vector)
<code>penalty</code>	Penalty function to be applied, either "MCP" (default), "SCAD", or "lasso")
<code>gamma</code>	Tuning parameter of the MCP/SCAD penalty, as in <code>ncvreg()</code> ; default is 3 for MCP and 3.7 for SCAD.
<code>alpha</code>	Tuning parameter controlling the ridge component of penalty, as in <code>ncvreg()</code> ; default is 1 (meaning no ridge penalty)
<code>lambda</code>	Regularization parameter value at which to estimate beta; must be scalar – for pathwise optimization, see <code>ncvreg()</code>
<code>eps</code>	Convergence threshold. The algorithm iterates until the RMSD for the change in linear predictors for each coefficient is less than <code>eps</code> . Default is 1e-4.
<code>max.iter</code>	Maximum number of allowed iterations; if this number is reached, algorithm will terminate prior to convergence. Default: 1000.
<code>penalty.factor</code>	Multiplicative factor for the penalty applied to each coefficient, as in <code>ncvreg()</code> . In particular, note that if you include an intercept, you probably want to set its entry to zero here.
<code>warn</code>	Return warning messages for failures to converge and model saturation? Default is TRUE.

Details

At the moment, this function only works for least-squares loss functions. Additional functionality for other loss functions (logistic, Cox) is in development.

Value

A list containing:

- `beta`: The estimated regression coefficients

- iter: The number of iterations required to solve for ‘beta
- loss: The loss (residual sum of squares) at convergence
- resid: The residuals at convergence
- lambda: See above
- penalty: See above
- gamma: See above
- alpha: See above
- penalty.factor: See above
- n: Sample size

Examples

```
data(Prostate)
X <- cbind(1, Prostate$X)
y <- Prostate$y
fit <- ncvfit(X, y, lambda=0.1, penalty.factor=c(0, rep(1, ncol(X)-1)))
fit$beta
# Compare with:
coef(ncvreg(X, y), 0.1)
# The unstandardized version makes little sense here, as it fails to account
# for differences in the scales of the predictors.
```

ncvreg

Fit an MCP- or SCAD-penalized regression path

Description

Fit coefficients paths for MCP- or SCAD-penalized regression models over a grid of values for the regularization parameter lambda. Fits linear and logistic regression models, with option for an additional L2 penalty.

Usage

```
ncvreg(
  X,
  y,
  family = c("gaussian", "binomial", "poisson"),
  penalty = c("MCP", "SCAD", "lasso"),
  gamma = switch(penalty, SCAD = 3.7, 3),
  alpha = 1,
  lambda.min = ifelse(n > p, 0.001, 0.05),
  nlambdas = 100,
  lambda,
  eps = 1e-04,
  max.iter = 10000,
```

```

    convex = TRUE,
    dfmax = p + 1,
    penalty.factor = rep(1, ncol(X)),
    warn = TRUE,
    returnX,
    ...
)

```

Arguments

<code>X</code>	The design matrix, without an intercept. <code>ncvreg</code> standardizes the data and includes an intercept by default.
<code>y</code>	The response vector.
<code>family</code>	Either "gaussian", "binomial", or "poisson", depending on the response.
<code>penalty</code>	The penalty to be applied to the model. Either "MCP" (the default), "SCAD", or "lasso".
<code>gamma</code>	The tuning parameter of the MCP/SCAD penalty (see details). Default is 3 for MCP and 3.7 for SCAD.
<code>alpha</code>	Tuning parameter for the Mnet estimator which controls the relative contributions from the MCP/SCAD penalty and the ridge, or L2 penalty. <code>alpha=1</code> is equivalent to MCP/SCAD penalty, while <code>alpha=0</code> would be equivalent to ridge regression. However, <code>alpha=0</code> is not supported; <code>alpha</code> may be arbitrarily small, but not exactly 0.
<code>lambda.min</code>	The smallest value for <code>lambda</code> , as a fraction of <code>lambda.max</code> . Default is 0.001 if the number of observations is larger than the number of covariates and .05 otherwise.
<code>nlambda</code>	The number of <code>lambda</code> values. Default is 100.
<code>lambda</code>	A user-specified sequence of <code>lambda</code> values. By default, a sequence of values of length <code>nlambda</code> is computed, equally spaced on the log scale.
<code>eps</code>	Convergence threshold. The algorithm iterates until the RMSD for the change in linear predictors for each coefficient is less than <code>eps</code> . Default is $1e-4$.
<code>max.iter</code>	Maximum number of iterations (total across entire path). Default is 10000.
<code>convex</code>	Calculate index for which objective function ceases to be locally convex? Default is TRUE.
<code>dfmax</code>	Upper bound for the number of nonzero coefficients. Default is no upper bound. However, for large data sets, computational burden may be heavy for models with a large number of nonzero coefficients.
<code>penalty.factor</code>	A multiplicative factor for the penalty applied to each coefficient. If supplied, <code>penalty.factor</code> must be a numeric vector of length equal to the number of columns of <code>X</code> . The purpose of <code>penalty.factor</code> is to apply differential penalization if some coefficients are thought to be more likely than others to be in the model. In particular, <code>penalty.factor</code> can be 0, in which case the coefficient is always in the model without shrinkage.
<code>warn</code>	Return warning messages for failures to converge and model saturation? Default is TRUE.

<code>returnX</code>	Return the standardized design matrix along with the fit? By default, this option is turned on if <code>X</code> is under 100 MB, but turned off for larger matrices to preserve memory. Note that certain methods, such as <code>summary.ncvreg()</code> require access to the design matrix and may not be able to run if <code>returnX=FALSE</code> .
<code>...</code>	Not used.

Details

The sequence of models indexed by the regularization parameter `lambda` is fit using a coordinate descent algorithm. For logistic regression models, some care is taken to avoid model saturation; the algorithm may exit early in this setting. The objective function is defined to be

$$Q(\beta|X, y) = \frac{1}{n}L(\beta|X, y) + P_{\lambda}(\beta),$$

where the loss function `L` is the deviance (-2 times the log likelihood) for the specified outcome distribution (gaussian/binomial/poisson). See [here](#) for more details.

This algorithm is stable, very efficient, and generally converges quite rapidly to the solution. For GLMs, [adaptive rescaling](#) is used.

Value

An object with S3 class "ncvreg" containing:

beta The fitted matrix of coefficients. The number of rows is equal to the number of coefficients, and the number of columns is equal to `nlambda`.

iter A vector of length `nlambda` containing the number of iterations until convergence at each value of `lambda`.

lambda The sequence of regularization parameter values in the path.

penalty, family, gamma, alpha, penalty.factor Same as above.

convex.min The last index for which the objective function is locally convex. The smallest value of `lambda` for which the objective function is convex is therefore `lambda[convex.min]`, with corresponding coefficients `beta[,convex.min]`.

loss A vector containing the deviance (i.e., the loss) at each value of `lambda`. Note that for gaussian models, the loss is simply the residual sum of squares.

n Sample size.

Additionally, if `returnX=TRUE`, the object will also contain

X The standardized design matrix.

y The response, centered if `family='gaussian'`.

References

Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, 5: 232-253. [doi:10.1214/10AOAS388](https://doi.org/10.1214/10AOAS388)

See Also

[plot.ncvreg\(\)](#), [cv.ncvreg\(\)](#)

Examples

```
# Linear regression -----
data(Prostate)
X <- Prostate$X
y <- Prostate$y

op <- par(mfrow=c(2,2))
fit <- ncvreg(X, y)
plot(fit, main=expression(paste(gamma,"=",3)))
fit <- ncvreg(X, y, gamma=10)
plot(fit, main=expression(paste(gamma,"=",10)))
fit <- ncvreg(X, y, gamma=1.5)
plot(fit, main=expression(paste(gamma,"=",1.5)))
fit <- ncvreg(X, y, penalty="SCAD")
plot(fit, main=expression(paste("SCAD, ",gamma,"=",3)))
par(op)

op <- par(mfrow=c(2,2))
fit <- ncvreg(X, y)
plot(fit, main=expression(paste(alpha,"=",1)))
fit <- ncvreg(X, y, alpha=0.9)
plot(fit, main=expression(paste(alpha,"=",0.9)))
fit <- ncvreg(X, y, alpha=0.5)
plot(fit, main=expression(paste(alpha,"=",0.5)))
fit <- ncvreg(X, y, alpha=0.1)
plot(fit, main=expression(paste(alpha,"=",0.1)))
par(op)

op <- par(mfrow=c(2,2))
fit <- ncvreg(X, y)
plot(mfdr(fit)) # Independence approximation
plot(mfdr(fit), type="EF") # Independence approximation
perm.fit <- perm.ncvreg(X, y)
plot(perm.fit)
plot(perm.fit, type="EF")
par(op)

# Logistic regression -----
data(Heart)
X <- Heart$X
y <- Heart$y

op <- par(mfrow=c(2,2))
fit <- ncvreg(X, y, family="binomial")
plot(fit, main=expression(paste(gamma,"=",3)))
fit <- ncvreg(X, y, family="binomial", gamma=10)
plot(fit, main=expression(paste(gamma,"=",10)))
fit <- ncvreg(X, y, family="binomial", gamma=1.5)
```

```

plot(fit, main=expression(paste(gamma,"=",1.5)))
fit <- ncvreg(X, y, family="binomial", penalty="SCAD")
plot(fit, main=expression(paste("SCAD, ",gamma,"=",3)))
par(op)

op <- par(mfrow=c(2,2))
fit <- ncvreg(X, y, family="binomial")
plot(fit, main=expression(paste(alpha,"=",1)))
fit <- ncvreg(X, y, family="binomial", alpha=0.9)
plot(fit, main=expression(paste(alpha,"=",0.9)))
fit <- ncvreg(X, y, family="binomial", alpha=0.5)
plot(fit, main=expression(paste(alpha,"=",0.5)))
fit <- ncvreg(X, y, family="binomial", alpha=0.1)
plot(fit, main=expression(paste(alpha,"=",0.1)))
par(op)

```

ncvsurv

Fit an MCP- or SCAD-penalized survival model

Description

Fit coefficients paths for MCP- or SCAD-penalized Cox regression models over a grid of values for the regularization parameter λ , with option for an additional L2 penalty.

Usage

```

ncvsurv(
  X,
  y,
  penalty = c("MCP", "SCAD", "lasso"),
  gamma = switch(penalty, SCAD = 3.7, 3),
  alpha = 1,
  lambda.min = ifelse(n > p, 0.001, 0.05),
  nlambdas = 100,
  lambda,
  eps = 1e-04,
  max.iter = 10000,
  convex = TRUE,
  dfmax = p,
  penalty.factor = rep(1, ncol(X)),
  warn = TRUE,
  returnX,
  ...
)

```

Arguments

<code>X</code>	The design matrix of predictor values. <code>ncvsurv</code> standardizes the data prior to fitting.
<code>y</code>	The time-to-event outcome, as a two-column matrix or <code>survival::Surv()</code> object. The first column should be time on study (follow up time); the second column should be a binary variable with 1 indicating that the event has occurred and 0 indicating (right) censoring.
<code>penalty</code>	The penalty to be applied to the model. Either "MCP" (the default), "SCAD", or "lasso".
<code>gamma</code>	The tuning parameter of the MCP/SCAD penalty (see details). Default is 3 for MCP and 3.7 for SCAD.
<code>alpha</code>	Tuning parameter for the Mnet estimator which controls the relative contributions from the MCP/SCAD penalty and the ridge, or L2 penalty. <code>alpha=1</code> is equivalent to MCP/SCAD penalty, while <code>alpha=0</code> would be equivalent to ridge regression. However, <code>alpha=0</code> is not supported; <code>alpha</code> may be arbitrarily small, but not exactly 0.
<code>lambda.min</code>	The smallest value for <code>lambda</code> , as a fraction of <code>lambda.max</code> . Default is .001 if the number of observations is larger than the number of covariates and .05 otherwise.
<code>nlambda</code>	The number of <code>lambda</code> values. Default is 100.
<code>lambda</code>	A user-specified sequence of <code>lambda</code> values. By default, a sequence of values of length <code>nlambda</code> is computed, equally spaced on the log scale.
<code>eps</code>	Convergence threshold. The algorithm iterates until the RMSD for the change in linear predictors for any coefficient is less than <code>eps</code> . Default is $1e-4$.
<code>max.iter</code>	Maximum number of iterations (total across entire path). Default is 1000.
<code>convex</code>	Calculate index for which objective function ceases to be locally convex? Default is TRUE.
<code>dfmax</code>	Upper bound for the number of nonzero coefficients. Default is no upper bound. However, for large data sets, computational burden may be heavy for models with a large number of nonzero coefficients.
<code>penalty.factor</code>	A multiplicative factor for the penalty applied to each coefficient. If supplied, <code>penalty.factor</code> must be a numeric vector of length equal to the number of columns of <code>X</code> . The purpose of <code>penalty.factor</code> is to apply differential penalization if some coefficients are thought to be more likely than others to be in the model. In particular, <code>penalty.factor</code> can be 0, in which case the coefficient is always in the model without any penalization/shrinkage.
<code>warn</code>	Return warning messages for failures to converge and model saturation? Default is TRUE.
<code>returnX</code>	Return the standardized design matrix along with the fit? By default, this option is turned on if <code>X</code> is under 100 MB, but turned off for larger matrices to preserve memory. Note that certain methods, such as <code>summary.ncvreg()</code> , require access to the design matrix and may not be able to run if <code>returnX=FALSE</code> .
<code>...</code>	Not used.

Details

The sequence of models indexed by the regularization parameter `lambda` is fit using a coordinate descent algorithm. In order to accomplish this, the second derivative (Hessian) of the Cox partial log-likelihood is diagonalized (see references for details). The objective function is defined to be

$$Q(\beta|X, y) = \frac{1}{n}L(\beta|X, y) + P_\lambda(\beta),$$

where the loss function `L` is the deviance (-2 times the partial log-likelihood) from the Cox regression mode. See [here](#) for more details.

Presently, ties are not handled by `ncvsurv` in a particularly sophisticated manner. This will be improved upon in a future release of **ncvreg**.

Value

An object with S3 class `ncvsurv` containing:

- beta** The fitted matrix of coefficients. The number of rows is equal to the number of coefficients, and the number of columns is equal to `nlambda`.
- iter** A vector of length `nlambda` containing the number of iterations until convergence at each value of `lambda`.
- lambda** The sequence of regularization parameter values in the path.
- penalty, gamma, penalty.factor, alpha, model** Same as above.
- convex.min** The last index for which the objective function is locally convex. The smallest value of `lambda` for which the objective function is convex is therefore `lambda[convex.min]`, with corresponding coefficients `beta[, convex.min]`.
- loss** The deviance of the fitted model at each value of `lambda`.
- n** The number of instances.

For Cox models, the following objects are also returned (and are necessary to estimate baseline survival conditional on the estimated regression coefficients), all of which are ordered by time on study. I.e., the `ith` row of `W` does not correspond to the `ith` row of `X`):

- W** Matrix of `exp(beta)` values for each subject over all `lambda` values.
- time** Times on study.
- fail** Failure event indicator.

Additionally, if `returnX=TRUE`, the object will also contain

- X** The standardized design matrix.

References

- Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, **5**: 232-253. doi:10.1214/10AOAS388
- Simon N, Friedman JH, Hastie T, and Tibshirani R. (2011) Regularization Paths for Cox's Proportional Hazards Model via Coordinate Descent. *Journal of Statistical Software*, **39**: 1-13. doi:10.18637/jss.v039.i05

See Also

[plot.ncvreg\(\)](#), [cv.ncvsurv\(\)](#)

Examples

```
data(Lung)
X <- Lung$X
y <- Lung$y

op <- par(mfrow=c(2,2))
fit <- ncvsurv(X, y)
plot(fit, main=expression(paste(gamma,"=",3)))
fit <- ncvsurv(X, y, gamma=10)
plot(fit, main=expression(paste(gamma,"=",10)))
fit <- ncvsurv(X, y, gamma=1.5)
plot(fit, main=expression(paste(gamma,"=",1.5)))
fit <- ncvsurv(X, y, penalty="SCAD")
plot(fit, main=expression(paste("SCAD, ",gamma,"=",3)))
par(op)

fit <- ncvsurv(X,y)
ll <- log(fit$lambda)
op <- par(mfrow=c(2,1))
plot(ll, BIC(fit), type="l", xlim=rev(range(ll)))
lam <- fit$lambda[which.min(BIC(fit))]
b <- coef(fit, lambda=lam)
b[b!=0]
plot(fit)
abline(v=lam)
par(op)

S <- predict(fit, X, type='survival', lambda=lam)
plot(S, xlim=c(0,200))
```

perm.ncvreg

Permutation fitting for ncvreg

Description

Fits multiple penalized regression models in which the outcome is randomly permuted, thereby allowing estimation of the marginal false discovery rate.

Usage

```
perm.ncvreg(
  X,
  y,
  ...,
  permute = c("outcome", "residuals"),
```



```

    N = 10,
    seed,
    trace = FALSE
  )

```

Arguments

<code>x</code>	The design matrix, without an intercept, as in <code>ncvreg</code> .
<code>y</code>	The response vector, as in <code>ncvreg</code> .
<code>...</code>	Additional arguments to <code>ncvreg</code> .
<code>permute</code>	What to permute. If 'outcome', the response vector, <code>y</code> , is permuted. If 'residuals', the residuals are permuted. This is only available for linear regression (i.e., for <code>family='gaussian'</code>). Note that permuting the residuals may take a long time, as the residuals differ for each value of <code>lambda</code> , so separate permutations are required at every value of <code>lambda</code> . See also permres() .
<code>N</code>	The number of permutation replications. Default is 10.
<code>seed</code>	You may set the seed of the random number generator in order to obtain reproducible results.
<code>trace</code>	If set to TRUE, <code>perm.ncvreg</code> will inform the user of its progress by announcing the beginning of each permutation fit. Default is FALSE.

Details

The function fits a penalized regression model to the actual data, then repeats the process `N` times with a permuted version of the response vector. This allows estimation of the expected number of variables included by chance for each value of `lambda`. The ratio of this expected quantity to the number of selected variables using the actual (non-permuted) response is called the marginal false discovery rate (mFDR).

Value

An object with S3 class "perm.ncvreg" containing:

<code>EF</code>	The number of variables selected at each value of <code>lambda</code> , averaged over the permutation fits.
<code>S</code>	The actual number of selected variables for the non-permuted data.
<code>mFDR</code>	The estimated marginal false discovery rate (<code>EF/S</code>).
<code>fit</code>	The fitted <code>ncvreg</code> object for the original (non-permuted) data.
<code>loss</code>	The loss/deviance for each value of <code>lambda</code> , averaged over the permutation fits. This is an estimate of the explanatory power of the model under null conditions, and can be used to adjust the loss of the fitted model in a manner akin to the idea of an adjusted R-squared in classical regression.

Author(s)

Patrick Breheny patrick-breheny@uiowa.edu

See Also

`ncvreg()`, `plot.mfdr()`, `mfdr()`

Examples

```
# Linear regression -----
data(Prostate)
pmfit <- perm.ncvreg(Prostate$X, Prostate$y)

op <- par(mfcol=c(2,2))
plot(pmfit)
plot(pmfit, type="EF")
plot(pmfit$fit)
lam <- pmfit$fit$lambda

pmfit.r <- perm.ncvreg(Prostate$X, Prostate$y, permute='residuals')
plot(pmfit.r, col="red")           # Permuting residuals is
lines(lam, pmfit$mFDR, col="gray60") # less conservative
par(op)

# Logistic regression -----
data(Heart)
pmfit <- perm.ncvreg(Heart$X, Heart$y, family="binomial")

op <- par(mfcol=c(2,2))
plot(pmfit)
plot(pmfit, type="EF")
plot(pmfit$fit)
par(op)
```

permres

Permute residuals for a fitted ncvreg model

Description

Fits multiple penalized regression models in which the residuals are randomly permuted, thereby allowing estimation of the marginal false discovery rate.

Usage

```
permres(fit, ...)

## S3 method for class 'ncvreg'
permres(fit, lambda, N = 10, seed, trace = FALSE, ...)
```

Arguments

<code>fit</code>	A fitted <code>ncvreg</code> model, as produced by <code>ncvreg()</code> . To use with <code>permres</code> , the model must be fit using the <code>returnX=TRUE</code> option.
<code>...</code>	Not used.
<code>lambda</code>	The regularization parameter to use for estimating residuals. Unlike <code>perm.ncvreg()</code> , <code>permres()</code> calculates EF and mFDR for a specific <code>lambda</code> value, not an entire path. As a result, it runs much faster.
<code>N</code>	The number of permutation replications. Default is 10.
<code>seed</code>	You may set the seed of the random number generator in order to obtain reproducible results.
<code>trace</code>	If set to <code>TRUE</code> , <code>perm.ncvreg</code> will inform the user of its progress by announcing the beginning of each permutation fit. Default is <code>FALSE</code> .

Details

The function fits a penalized regression model to the actual data, then repeats the process `N` times with a permuted version of the response vector. This allows estimation of the expected number of variables included by chance for each value of `lambda`. The ratio of this expected quantity to the number of selected variables using the actual (non-permuted) response is called the marginal false discovery rate (mFDR).

Value

A list with the following components:

<code>EF</code>	The number of variables selected at each value of <code>lambda</code> , averaged over the permutation fits.
<code>S</code>	The actual number of selected variables for the non-permuted data.
<code>mFDR</code>	The estimated marginal false discovery rate (EF/S).
<code>loss</code>	The loss/deviance, averaged over the permutation fits. This is an estimate of the explanatory power of the model under null conditions, and can be used to adjust the loss of the fitted model in a manner akin to the idea of an adjusted R-squared in classical regression.

Author(s)

Patrick Breheny patrick-breheny@uiowa.edu

See Also

`ncvreg()`, `'mfdr()`, `perm.ncvreg()`

Examples

```
data(Prostate)
fit <- ncvreg(Prostate$X, Prostate$y, N=50)
permres(fit, lambda=0.15)
```

plot.cv.ncvreg

Plots the cross-validation curve from a cv.ncvreg object

Description

Plots the cross-validation curve from a `cv.ncvreg` or `cv.ncvsurv` object, along with standard error bars.

Usage

```
## S3 method for class 'cv.ncvreg'
plot(
  x,
  log.l = TRUE,
  type = c("cve", "rsq", "scale", "snr", "pred", "all"),
  selected = TRUE,
  vertical.line = TRUE,
  col = "red",
  ...
)
```

Arguments

<code>x</code>	A <code>cv.ncvreg</code> or <code>cv.ncvsurv</code> object.
<code>log.l</code>	Should horizontal axis be on the log scale? Default is TRUE.
<code>type</code>	What to plot on the vertical axis: • <code>cve</code> plots the cross-validation error (deviance) • <code>rsq</code> plots an estimate of the fraction of the deviance explained by the model (R-squared) • <code>snr</code> plots an estimate of the signal-to-noise ratio • <code>scale</code> plots, for <code>family="gaussian"</code>, an estimate of the scale parameter (standard deviation) • <code>pred</code> plots, for <code>family="binomial"</code>, the estimated prediction error • <code>all</code> produces all of the above
<code>selected</code>	If TRUE (the default), places an axis on top of the plot denoting the number of variables in the model (i.e., that have a nonzero regression coefficient) at that value of <code>lambda</code> .
<code>vertical.line</code>	If TRUE (the default), draws a vertical line at the value where cross-validation error is minimized.
<code>col</code>	Controls the color of the dots (CV estimates).
<code>...</code>	Other graphical parameters to <code>plot()</code>

Details

Error bars representing approximate 68% confidence intervals are plotted along with the estimates across values of λ . For `rsq` and `snr` applied to models other than linear regression, the Cox-Snell R-squared is used.

Author(s)

Patrick Breheny

References

Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, **5**: 232-253. [doi:10.1214/10AOAS388](https://doi.org/10.1214/10AOAS388)

See Also

[ncvreg\(\)](#), [cv.ncvreg\(\)](#)

Examples

```
# Linear regression -----
data(Prostate)
cvfit <- cv.ncvreg(Prostate$X, Prostate$y)
plot(cvfit)
op <- par(mfrow=c(2,2))
plot(cvfit, type="all")
par(op)

# Logistic regression -----
data(Heart)
cvfit <- cv.ncvreg(Heart$X, Heart$y, family="binomial")
plot(cvfit)
op <- par(mfrow=c(2,2))
plot(cvfit, type="all")
par(op)

# Cox regression -----
data(Lung)
cvfit <- cv.ncvsurv(Lung$X, Lung$y)
op <- par(mfrow=c(1,2))
plot(cvfit)
plot(cvfit, type="rsq")
par(op)
```

plot.mfdr

*Plot marginal false discovery rate curves***Description**

Plot marginal false discovery rate curves from an mfdr or perm.ncvreg object.

Usage

```
## S3 method for class 'mfdr'
plot(
  x,
  type = c("mFDR", "EF"),
  log.l = FALSE,
  selected = TRUE,
  legend = TRUE,
  ...
)
```

Arguments

x	A perm.ncvreg or mfdr object.
type	What to plot on the vertical axis. mFDR plots the marginal false discovery rate; EF plots the expected number of false discoveries along with the actual number of variables included in the model.
log.l	Should horizontal axis be on the log scale? Default is FALSE.
selected	If TRUE (the default), places an axis on top of the plot denoting the number of variables in the model (i.e., that have a nonzero regression coefficient) at that value of lambda.
legend	For type="EF" plots, draw a legend to indicate which line is for the actual selections and which line is for the expected number of false discoveries? Default is TRUE.
...	Other graphical parameters to pass to plot()

Author(s)

Patrick Breheny

References

Breheny P (2019). Marginal false discovery rates for penalized regression models. *Biostatistics*, 20: 299-314.

See Also

[mfdr\(\)](#), [perm.ncvreg\(\)](#)

Examples

```

data(Prostate)
fit <- ncvreg(Prostate$X, Prostate$y)

obj <- mfdr(fit)
obj[1:10,]

# Some plotting options
plot(obj)
plot(obj, type="EF")
plot(obj, log=TRUE)

# Comparison with perm.ncvreg
op <- par(mfrow=c(2,2))
plot(obj)
plot(obj, type="EF")
pmfit <- perm.ncvreg(Prostate$X, Prostate$y)
plot(pmfit)
plot(pmfit, type="EF")
par(op)

```

plot.ncvreg

*Plot coefficients from a ncvreg object***Description**

Produces a plot of the coefficient paths for a fitted ncvreg object.

Usage

```

## S3 method for class 'ncvreg'
plot(x, alpha = 1, log.l = FALSE, shade = TRUE, col, ...)

```

Arguments

x	Fitted "ncvreg" model.
alpha	Controls alpha-blending, helpful when the number of features is large. Default is alpha=1.
log.l	Should horizontal axis be on the log scale? Default is FALSE.
shade	Should nonconvex region be shaded? Default is TRUE.
col	Vector of colors for coefficient lines. By default, evenly spaced colors are selected automatically.
...	Other graphical parameters to plot()

Author(s)

Patrick Breheny

References

Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, **5**: 232-253. doi:10.1214/10AOAS388

See Also

[ncvreg\(\)](#)

Examples

```
data(Prostate)
fit <- ncvreg(Prostate$X, Prostate$y)
plot(fit)
plot(fit, col="black")
plot(fit, log=TRUE)
fit <- ncvreg(Prostate$X, Prostate$y, penalty.factor=rep(c(1, 1, 1, Inf), 2))
plot(fit, col=c('red', 'black', 'green')) # Recycled among nonzero paths
```

plot.ncvsurv.func

Plot survival curve for ncvsurv model

Description

Plot survival curve for a model that has been fit using [ncvsurv\(\)](#) followed by a prediction of the survival function using [predict.ncvsurv\(\)](#).

Usage

```
## S3 method for class 'ncvsurv.func'
plot(x, alpha = 1, ...)
```

Arguments

x	A ncvsurv.func object, which is returned by predict.ncvsurv() if type='survival' is specified. See examples.
alpha	Controls alpha-blending (i.e., transparency). Useful if many overlapping lines are present.
...	Other graphical parameters to pass to plot

Author(s)

Patrick Breheny

See Also

[ncvsurv\(\)](#), [predict.ncvsurv\(\)](#)

Examples

```

data(Lung)
X <- Lung$X
y <- Lung$y

fit <- ncvsurv(X, y)

# A single survival curve
S <- predict(fit, X[1,], type='survival', lambda=.15)
plot(S, xlim=c(0,200))

# Lots of survival curves
S <- predict(fit, X, type='survival', lambda=.08)
plot(S, xlim=c(0,200), alpha=0.3)

```

predict.cv.ncvreg	<i>Model predictions based on a fitted ncvreg object.</i>
-------------------	---

Description

Similar to other predict methods, this function returns predictions from a fitted ncvreg object.

Usage

```

## S3 method for class 'cv.ncvreg'
predict(
  object,
  X,
  type = c("link", "response", "class", "coefficients", "vars", "nvars"),
  which = object$min,
  ...
)

## S3 method for class 'cv.ncvreg'
coef(object, which = object$min, ...)

## S3 method for class 'cv.ncvsurv'
predict(
  object,
  X,
  type = c("link", "response", "survival", "median", "coefficients", "vars", "nvars"),
  which = object$min,
  ...
)

## S3 method for class 'ncvreg'
predict(

```

```

    object,
    X,
    type = c("link", "response", "class", "coefficients", "vars", "nvars"),
    lambda,
    which = 1:length(object$lambda),
    ...
)

## S3 method for class 'ncvreg'
coef(object, lambda, which = 1:length(object$lambda), drop = TRUE, ...)

```

Arguments

object	Fitted ncvreg model object.
X	Matrix of values at which predictions are to be made. Not used for type="coefficients" or for some of the type settings in predict.
type	Type of prediction: • link returns the linear predictors • response gives the fitted values • class returns the binomial outcome with the highest probability • coefficients returns the coefficients • vars returns a list containing the indices and names of the nonzero variables at each value of lambda • nvars returns the number of nonzero coefficients at each value of lambda.
which	Indices of the penalty parameter lambda at which predictions are required. By default, all indices are returned. If lambda is specified, this will override which.
...	Not used.
lambda	Values of the regularization parameter lambda at which predictions are requested. For values of lambda not in the sequence of fitted models, linear interpolation is used.
drop	If coefficients for a single value of lambda are to be returned, reduce dimensions to a vector? Setting drop=FALSE returns a 1-column matrix.

Value

The object returned depends on type.

Author(s)

Patrick Breheny

References

Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, 5: 232-253. [doi:10.1214/10AOAS388](https://doi.org/10.1214/10AOAS388)

See Also[ncvreg\(\)](#)**Examples**

```
data(Heart)

fit <- ncvreg(Heart$X, Heart$y, family="binomial")
coef(fit, lambda=0.05)
head(predict(fit, Heart$X, type="link", lambda=0.05))
head(predict(fit, Heart$X, type="response", lambda=0.05))
head(predict(fit, Heart$X, type="class", lambda=0.05))
predict(fit, type="vars", lambda=c(0.05, 0.01))
predict(fit, type="nvars", lambda=c(0.05, 0.01))
```

predict.ncvsurv

*Model predictions based on a fitted ncvsurv object.***Description**

Similar to other predict methods, this function returns predictions from a fitted ncvsurv object.

Usage

```
## S3 method for class 'ncvsurv'
predict(
  object,
  X,
  type = c("link", "response", "survival", "hazard", "median", "coefficients", "vars",
    "nvars"),
  lambda,
  which = 1:length(object$lambda),
  ...
)
```

Arguments

object	Fitted "ncvsurv" model object.
X	Matrix of values at which predictions are to be made. Not used for type="coefficients" or for some of the type settings in predict.
type	Type of prediction: • link returns the linear predictors • response gives the risk (i.e., exp(link)) • survival returns the estimated survival function • median estimates median survival times The other options are all identical to their ncvreg() counterparts:

	• <code>coefficients</code> returns the coefficients • <code>vars</code> returns a list containing the indices and names of the nonzero variables at each value of <code>lambda</code> • <code>nvars</code> returns the number of nonzero coefficients at each value of <code>lambda</code>.
<code>lambda</code>	Values of the regularization parameter <code>lambda</code> at which predictions are requested. For values of <code>lambda</code> not in the sequence of fitted models, linear interpolation is used.
<code>which</code>	Indices of the penalty parameter <code>lambda</code> at which predictions are required. By default, all indices are returned. If <code>lambda</code> is specified, this will override <code>which</code> .
<code>...</code>	Not used.

Details

Estimation of baseline survival function conditional on the estimated values of β is carried out according to the method described in Chapter 4.3 of Kalbfleish and Prentice. In particular, it agrees exactly the results returned by `survfit.coxph(..., type='kalbfleisch-prentice')` in the `survival` package.

Value

The object returned depends on `type`.

Author(s)

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References

- Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, **5**: 232-253. doi:[10.1214/10AOAS388](https://doi.org/10.1214/10AOAS388)
- Kalbfleish JD and Prentice RL (2002). *The Statistical Analysis of Failure Time Data*, 2nd edition. Wiley.

See Also

[ncvsurv\(\)](#)

Examples

```
data(Lung)
X <- Lung$X
y <- Lung$y

fit <- ncvsurv(X,y)
coef(fit, lambda=0.05)
head(predict(fit, X, type="link", lambda=0.05))
head(predict(fit, X, type="response", lambda=0.05))
```

```
# Survival function
S <- predict(fit, X[1,], type="survival", lambda=0.05)
S(100)
S <- predict(fit, X, type="survival", lambda=0.05)
plot(S, xlim=c(0,200))

# Medians
predict(fit, X[1,], type="median", lambda=0.05)
M <- predict(fit, X, type="median")
M[1:10, 1:10]

# Nonzero coefficients
predict(fit, type="vars", lambda=c(0.1, 0.01))
predict(fit, type="nvars", lambda=c(0.1, 0.01))
```

Prostate

*Factors associated with prostate specific antigen***Description**

Data from a study by Stamey et al. (1989) to examine the association between prostate specific antigen (PSA) and several clinical measures that are potentially associated with PSA in men who were about to receive a radical prostatectomy.

Usage

Prostate

Format

A list of two objects: **y** and **X**

y Log PSA

X A matrix with 97 instances (rows) and 8 predictor variables (columns). The remainder of this list describes the columns of **X**

lcavol Log cancer volume

lweight Log prostate weight

age The man's age (years)

lbph Log of the amount of benign hyperplasia

svi Seminal vesicle invasion (1=Yes, 0=No)

lcp Log of capsular penetration

gleason Gleason score

pgg45 Percent of Gleason scores 4 or 5

Source

<https://web.stanford.edu/~hastie/ElemStatLearn/>

References

- Hastie T, Tibshirani R, and Friedman J. (2001). *The Elements of Statistical Learning*. Springer.
- Stamey T, et al. (1989). Prostate specific antigen in the diagnosis and treatment of adenocarcinoma of the prostate. II. Radical prostatectomy treated patients. *Journal of Urology*, **16**: 1076-1083.

residuals.ncvreg	<i>Extract residuals from a ncvreg or ncvsurv fit</i>
------------------	---

Description

Currently, only deviance residuals are supported.

Usage

```
## S3 method for class 'ncvreg'
residuals(object, lambda, which = 1:length(object$lambda), drop = TRUE, ...)
```

Arguments

object	Object of class ncvreg or ncvsurv.
lambda	Values of the regularization parameter at which residuals are requested (numeric vector). For values of lambda not in the sequence of fitted models, linear interpolation is used.
which	Index of the penalty parameter at which residuals are requested (default = all indices). If lambda is specified, this take precedence over which.
drop	By default, if a single value of lambda is supplied, a vector of residuals is returned (logical; default=TRUE). Set drop=FALSE if you wish to have the function always return a matrix (see drop()).
...	Not used.

Examples

```
data(Prostate)
X <- Prostate$X
y <- Prostate$y
fit <- ncvreg(X, y)
residuals(fit)[1:5, 1:5]
head(residuals(fit, lambda=0.1))
```

std	<i>Standardizes a design matrix</i>
-----	-------------------------------------

Description

Accepts a design matrix and returns a standardized version of that matrix (i.e., each column will have mean 0 and mean sum of squares equal to 1).

Usage

```
std(X, Xnew)
```

Arguments

X	A matrix (or object that can be coerced to a matrix, such as a data frame or numeric vector).
Xnew	Optional. If supplied, X must be the output of std() and Xnew is to be standardized in the same way. See examples for why this might be useful.

Details

This function centers and scales each column of X so that

$$\sum_{i=1}^n x_{ij} = 0$$

and

$$n^{-1} \sum_{i=1}^n x_{ij}^2 = 1$$

for all j. This is usually not necessary to call directly, as **ncvreg** internally standardizes the design matrix, but inspection of the standardized design matrix can sometimes be useful. This differs from the base R function [scale\(\)](#) in two ways:

1. `scale()` uses the sample standard deviation $\sqrt{\text{sum}(x^2)/(n-1)}$, while `std()` uses the root-mean-square standard deviation $\sqrt{\text{mean}(\text{sum}(x^2))}$ without the $n/(n-1)$ correction
2. `std` is faster.

Value

The standardized design matrix, with the following attributes:

center, scale	mean and standard deviation used to scale the columns
nonsingular	A vector indicating which columns of the original design matrix were able to be standardized (constant columns cannot be standardized to have a standard deviation of 1)

Examples

```
data(Prostate)
S <- std(Prostate$X)
apply(S, 2, sum)
apply(S, 2, function(x) mean(x^2))

# Standardizing new observations
X1 <- Prostate$X[1:90,]
X2 <- Prostate$X[91:97,]
S <- std(X1)
head(std(S, X2))
# Useful if you fit to a standardized X, but then get new obs:
y <- Prostate$y[1:90]
fit <- ncvreg(S, y)
predict(fit, std(S, X2), lambda=0.1)
# Same as
predict(ncvreg(X1, y), X2, lambda=0.1)
```

summary.cv.ncvreg

Summarizing cross-validation-based inference

Description

Summary method for cv.ncvreg objects

Usage

```
## S3 method for class 'cv.ncvreg'
summary(object, include_fit = FALSE, ...)

## S3 method for class 'summary.cv.ncvreg'
print(x, digits, include_fit = FALSE, ...)
```

Arguments

object	A cv.ncvreg or cv.ncvsurv object.
include_fit	In addition to summarizing the cross-validation object, also summarize the model with the lowest CV error? (default: FALSE)
...	Further arguments passed to or from other methods.
x	A summary.cv.ncvreg object.
digits	Number of digits past the decimal point to print out. Can be a vector specifying different display digits for each of the five non-integer printed values.

Value

An object with S3 class `summary.cv.ncvreg`. The class has its own print method and contains the following list elements:

penalty The penalty used by `ncvreg`.

model Either "linear" or "logistic", depending on the family option in `ncvreg`.

n Number of instances

p Number of regression coefficients (not including the intercept).

min The index of `lambda` with the smallest cross-validation error.

lambda The sequence of `lambda` values used by `cv.ncvreg`.

cve Cross-validation error (deviance).

r.squared Proportion of variance explained by the model, as estimated by cross-validation. For models outside of linear regression, the Cox-Snell approach to defining R-squared is used.

snr Signal to noise ratio, as estimated by cross-validation.

sigma For linear regression models, the scale parameter estimate.

pe For logistic regression models, the prediction error (misclassification error).

Author(s)

Patrick Breheny

References

Breheny P and Huang J. (2011) Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection. *Annals of Applied Statistics*, 5: 232-253.
doi:[10.1214/10AOAS388](https://doi.org/10.1214/10AOAS388)

See Also

[ncvreg\(\)](#), [cv.ncvreg\(\)](#), [plot.cv.ncvreg\(\)](#), [summary.ncvreg\(\)](#)

Examples

```
# Linear regression -----
data(Prostate)
cvfit <- cv.ncvreg(Prostate$X, Prostate$y)
summary(cvfit)

# Logistic regression -----
data(Heart)
cvfit <- cv.ncvreg(Heart$X, Heart$y, family="binomial")
summary(cvfit)

# Cox regression -----
data(Lung)
cvfit <- cv.ncvsurv(Lung$X, Lung$y)
summary(cvfit)
```

summary.ncvreg

*Summary method for ncvreg objects***Description**

Inferential summaries for ncvreg and ncvsurv objects based on local marginal false discovery rates.

Usage

```
## S3 method for class 'ncvreg'
summary(object, lambda, which, number, cutoff, sort = TRUE, sigma, ...)

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

Arguments

object	An ncvreg or ncvsurv object.
lambda	The regularization parameter value at which inference should be reported.
which	Alternatively, lambda may be specified by index; which=10 means: report inference for the 10th value of lambda along the regularization path. If both lambda and which are specified, lambda takes precedence.
number	By default, summary will provide an inferential summary for each variable that has been selected (i.e., each variable with a nonzero coefficient). Specifying number=5, for example, means that the summary table will include the 5 features with the lowest mfdR values, regardless of whether they were selected. To see all features, number=Inf.
cutoff	Alternatively, specifying for example cutoff=0.3 will report inference for all features with mfdR under 30%. If both number and cutoff are specified, the intersection between both sets of features is reported.
sort	Should the results be sorted by mfdR? (default: TRUE)
sigma	For linear regression models, users can supply an estimate of the residual standard deviation. The default is to use RSS / DF, where degrees of freedom are approximated using the number of nonzero coefficients.
...	Further arguments; in particular, if you have set returnX=FALSE, you will need to supply X and y in order to calculate local mFDRs.
x	A summary.ncvreg object.
digits	Number of digits past the decimal point to print out. Can be a vector specifying different display digits for each of the five non-integer printed values.

Value

An object with S3 class `summary.ncvreg`. The class has its own print method and contains the following list elements:

<code>penalty</code>	The penalty used by <code>ncvreg</code> or <code>ncvsurv</code>
<code>model</code>	Either "linear", "logistic", or "Cox".
<code>n</code>	Number of instances.
<code>p</code>	Number of regression coefficients (not including the intercept).
<code>lambda</code>	The lambda value at which inference is being reported.
<code>nvars</code>	The number of nonzero coefficients (again, not including the intercept) at that value of lambda.
<code>table</code>	A table containing estimates, normalized test statistics (z), and an estimate of the local mfdR for each coefficient. The mfdR may be loosely interpreted, in an empirical Bayes sense, as the probability that the given feature is null.
<code>unpen.table</code>	If there are any unpenalized coefficients, a separate inferential summary is given for them. Currently, this is based on <code>lm/glm/coxph</code> using the penalized coefficients to provide an offset. This is useful and more or less accurate, but not ideal; we hope to improve the inferential methods for unpenalized variables in the future.

Author(s)

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See Also

[ncvreg\(\)](#), [cv.ncvreg\(\)](#), [plot.cv.ncvreg\(\)](#), [local_mfdR\(\)](#)

Examples

```
# Linear regression -----
data(Prostate)
fit <- ncvreg(Prostate$X, Prostate$y)
summary(fit, lambda=0.08)

# Logistic regression -----
data(Heart)
fit <- ncvreg(Heart$X, Heart$y, family="binomial")
summary(fit, lambda=0.05)

# Cox regression -----
data(Lung)
fit <- ncvsurv(Lung$X, Lung$y)
summary(fit, lambda=0.1)

# Options -----
fit <- ncvreg(Heart$X, Heart$y, family="binomial")
summary(fit, lambda=0.08, number=3)
```

```
summary(fit, lambda=0.08, number=Inf)
summary(fit, lambda=0.08, cutoff=0.5)
summary(fit, lambda=0.08, number=3, cutoff=0.5)
summary(fit, lambda=0.08, number=5, cutoff=0.1)
summary(fit, lambda=0.08, number=Inf, sort=FALSE)
summary(fit, lambda=0.08, number=3, cutoff=0.5, sort=FALSE)

# If X and y are not returned with the fit, they must be supplied
fit <- ncvreg(Heart$X, Heart$y, family="binomial", returnX=FALSE)
summary(fit, X=Heart$X, y=Heart$y, lambda=0.08)
```

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