

Package ‘mpath’

August 18, 2017

Title Regularized Linear Models

Version 0.3-3

Date 2017-08-15

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Description Algorithms for fitting model-based penalized coefficient paths. Currently the models include penalized Poisson, negative binomial, zero-inflated Poisson and zero-inflated negative binomial regression models. The penalties include least absolute shrinkage and selection operator (LASSO), smoothly clipped absolute deviation (SCAD) and minimax concave penalty (MCP), and each possibly combining with L₂ penalty.

Imports MASS,glmnet,pscl,numDeriv, foreach, doParallel, bst

Depends methods

Suggests zic, R.rsp

VignetteBuilder R.rsp

License GPL-2

NeedsCompilation yes

Repository CRAN

Date/Publication 2017-08-18 17:53:48 UTC

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be.zeroinfl	<i>conduct backward stepwise variable elimination for zero inflated count regression</i>
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Description

conduct backward stepwise variable elimination for zero inflated count regression from zeroinfl function

Usage

```
be.zeroinfl(object, data, dist=c("poisson", "negbin", "geometric"), alpha=0.05,
trace=FALSE)
```

Arguments

object	an object from function zeroinfl
data	argument controlling formula processing via model.frame .
dist	one of the distributions in zeroinfl function
alpha	significance level of variable elimination
trace	logical value, if TRUE, print detailed calculation results

Details

conduct backward stepwise variable elimination for zero inflated count regression from zeroinfl function

Value

an object of zeroinfl with all variables having p-values less than the significance level alpha

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

breadReg

Bread for Sandwiches in Regularized Estimators

Description

Generic function for extracting an estimator for the bread of sandwiches.

Usage

```
breadReg(x, which, ...)
```

Arguments

x	a fitted model object.
which	which penalty parameter(s)?
...	arguments passed to methods.

Value

A matrix containing an estimator for the penalized second derivative of log-likelihood function. Typically, this should be an $k \times k$ matrix corresponding to k parameters. The rows and columns should be named as in `coef` or `terms`, respectively.

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[meatReg](#), [sandwichReg](#)

Examples

```
data("bioChemists", package = "pscl")
fm_zinb <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambda=10)
breadReg(fm_zinb, which=which.min(fm_zinb$bic))
```

conv2glmreg

convert glm object to class glmreg

Description

convert glm object to class glmreg, which then can be used for other purposes

Usage

```
conv2glmreg(object, family=c("poisson", "negbin"))
```

Arguments

object	an object of class glm
family	one of families in glm class

Value

an object of class glmreg

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

conv2zipath	<i>convert zeroinfl object to class zipath</i>
-------------	--

Description

convert zeroinfl object to class zipath, which then can be used to predict new data

Usage

```
conv2zipath(object, family=c("poisson", "negbin", "geometric"))
```

Arguments

object	an object of class zeroinfl
family	one of families in zeroinfl class

Value

an object of class zipath

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

cv.glmreg	<i>Cross-validation for glmreg</i>
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Description

Does k-fold cross-validation for glmreg, produces a plot, and returns cross-validated log-likelihood values for lambda

Usage

```
## S3 method for class 'formula'  
cv.glmreg(formula, data, weights, offset=NULL, ...)  
## S3 method for class 'matrix'  
cv.glmreg(x, y, weights, offset=NULL, ...)  
## Default S3 method:  
cv.glmreg(x, ...)  
## S3 method for class 'cv.glmreg'  
plot(x,se=TRUE,ylab=NULL, main=NULL, width=0.02, col="darkgrey", ...)  
## S3 method for class 'cv.glmreg'  
coef(object,which=object$lambda.which, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via <code>model.frame</code> .
x	x matrix as in <code>glmreg</code> . It could be object of <code>cv.glmreg</code> .
y	response y as in <code>glmreg</code> .
weights	Observation weights; defaults to 1 per observation
offset	Not implemented yet
object	object of <code>cv.glmreg</code>
which	Indices of the penalty parameter lambda at which estimates are extracted. By default, the one which generates the optimal cross-validation value.
se	logical value, if TRUE, standard error curve is also plotted
ylab	ylab on y-axis
main	title of plot
width	width of lines
col	color of standard error curve
...	Other arguments that can be passed to <code>glmreg</code> .

Details

The function runs `glmreg` `nfolds+1` times; the first to compute the lambda sequence, and then to compute the fit with each of the folds omitted. The error or the log-likelihood value is accumulated, and the average value and standard deviation over the folds is computed. Note that `cv.glmreg` can be used to search for values for alpha: it is required to call `cv.glmreg` with a fixed vector `foldid` for different values of alpha.

Value

an object of class "`cv.glmreg`" is returned, which is a list with the ingredients of the cross-validation fit.

fit	a fitted <code>glmreg</code> object for the full data.
residmat	matrix of log-likelihood values with row values for lambda and column values for kth cross-validation
bic	matrix of BIC values with row values for lambda and column values for kth cross-validation
cv	The mean cross-validated log-likelihood values - a vector of length <code>length(lambda)</code> .
cv.error	estimate of standard error of cv.
foldid	an optional vector of values between 1 and <code>nfold</code> identifying what fold each observation is in.
lambda	a vector of lambda values
lambda.which	index of lambda that gives maximum cv value.
lambda.optim	value of lambda that gives maximum cv value.

Author(s)

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References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[glmreg](#) and [plot](#), [predict](#), and [coef](#) methods for "cv.glmreg" object.

Examples

```
## Not run:
data("bioChemists", package = "pscl")
fm_pois <- cv.glmreg(art ~ ., data = bioChemists, family = "poisson")
plot(fm_pois)
title("Poisson Family",line=2.5)

## End(Not run)
```

cv.glmregNB

Cross-validation for glmregNB

Description

Does k-fold cross-validation for glmregNB, produces a plot, and returns cross-validated log-likelihood values for lambda

Usage

```
cv.glmregNB(formula, data, weights, lambda=NULL,
n folds=10, foldid, plot.it=TRUE, se=TRUE, n.cores=2, ...)
```

Arguments

formula	symbolic description of the model
data	arguments controlling formula processing via model.frame .
weights	Observation weights; defaults to 1 per observation
lambda	Optional user-supplied lambda sequence; default is NULL, and glmregNB chooses its own sequence
n folds	number of folds - default is 10. Although n folds can be as large as the sample size (leave-one-out CV), it is not recommended for large datasets. Smallest value allowable is n folds=3

foldid	an optional vector of values between 1 and nfold identifying what fold each observation is in. If supplied, nfold can be missing.
plot.it	a logical value, to plot the estimated log-likelihood values if TRUE.
se	a logical value, to plot with standard errors.
n.cores	The number of CPU cores to use. The cross-validation loop will attempt to send different CV folds off to different cores.
...	Other arguments that can be passed to glmregNB.

Details

The function runs `glmregNB` `nfolds+1` times; the first to get the lambda sequence, and then the remainder to compute the fit with each of the folds omitted. The error is accumulated, and the average error and standard deviation over the folds is computed. Note that `cv.glmregNB` does NOT search for values for alpha. A specific value should be supplied, else `alpha=1` is assumed by default. If users would like to cross-validate alpha as well, they should call `cv.glmregNB` with a pre-computed vector `foldid`, and then use this same fold vector in separate calls to `cv.glmregNB` with different values of alpha.

Value

an object of class "`cv.glmregNB`" is returned, which is a list with the ingredients of the cross-validation fit.

fit	a fitted <code>glmregNB</code> object for the full data.
residmat	matrix of log-likelihood values with row values for lambda and column values for kth cross-validation
cv	The mean cross-validated log-likelihood values - a vector of length <code>length(lambda)</code> .
cv.error	The standard error of cross-validated log-likelihood values - a vector of length <code>length(lambda)</code> .
lambda	a vector of lambda values
foldid	indicators of data used in each cross-validation, for reproductive purposes
lambda.which	index of lambda that gives maximum cv value.
lambda.optim	value of lambda that gives maximum cv value.

Author(s)

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References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[glmregNB](#) and [plot](#), [predict](#), and [coef](#) methods for "`cv.glmregNB`" object.

Examples

```
## Not run:
data("bioChemists", package = "pscl")
fm_nb <- cv.glmregNB(art ~ ., data = bioChemists)
plot(fm_nb)

## End(Not run)
```

cv.glmreg_fit

Internal function of cross-validation for glmreg

Description

Internal function to conduct k-fold cross-validation for glmreg, produces a plot, and returns cross-validated log-likelihood values for lambda

Usage

```
cv.glmreg_fit(x, y, weights, lambda=NULL, balance=TRUE,
family=c("gaussian", "binomial", "poisson", "negbin"),
n folds=10, foldid, plot.it=TRUE, se=TRUE, n.cores=2, ...)
```

Arguments

x	x matrix as in glmreg.
y	response y as in glmreg.
weights	Observation weights; defaults to 1 per observation
lambda	Optional user-supplied lambda sequence; default is NULL, and glmreg chooses its own sequence
balance	for family="binomial" only
family	response variable distribution
n folds	number of folds >=3, default is 10
foldid	an optional vector of values between 1 and nfold identifying what fold each observation is in. If supplied, nfold can be missing and will be ignored.
plot.it	a logical value, to plot the estimated log-likelihood values if TRUE.
se	a logical value, to plot with standard errors.
n.cores	The number of CPU cores to use. The cross-validation loop will attempt to send different CV folds off to different cores.
...	Other arguments that can be passed to glmreg.

Details

The function runs `glmreg` `nfolds+1` times; the first to compute the `lambda` sequence, and then to compute the fit with each of the folds omitted. The error or the log-likelihood value is accumulated, and the average value and standard deviation over the folds is computed. Note that `cv.glmreg` can be used to search for values for `alpha`: it is required to call `cv.glmreg` with a fixed vector `foldid` for different values of `alpha`.

Value

an object of class `"cv.glmreg"` is returned, which is a list with the ingredients of the cross-validation fit.

<code>fit</code>	a fitted <code>glmreg</code> object for the full data.
<code>residmat</code>	matrix of log-likelihood values with row values for <code>lambda</code> and column values for <code>kth</code> cross-validation
<code>cv</code>	The mean cross-validated log-likelihood values - a vector of length <code>length(lambda)</code> .
<code>cv.error</code>	estimate of standard error of <code>cv</code> .
<code>foldid</code>	an optional vector of values between 1 and <code>nfold</code> identifying what fold each observation is in.
<code>lambda</code>	a vector of <code>lambda</code> values
<code>lambda.which</code>	index of <code>lambda</code> that gives maximum <code>cv</code> value.
<code>lambda.optim</code>	value of <code>lambda</code> that gives maximum <code>cv</code> value.

Author(s)

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References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[glmreg](#) and [plot](#), [predict](#), and [coef](#) methods for `"cv.glmreg"` object.

cv.nclreg

*Cross-validation for nclreg***Description**

Does k-fold cross-validation for nclreg, produces a plot, and returns cross-validated log-likelihood values for lambda

Usage

```
## S3 method for class 'formula'
cv.nclreg(formula, data, weights, offset=NULL, ...)
## S3 method for class 'matrix'
cv.nclreg(x, y, weights, offset=NULL, ...)
## Default S3 method:
cv.nclreg(x, ...)
## S3 method for class 'cv.nclreg'
plot(x,se=TRUE,ylab=NULL, main=NULL, width=0.02, col="darkgrey", ...)
## S3 method for class 'cv.nclreg'
coef(object,which=object$lambda.which, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via <code>model.frame</code> .
x	x matrix as in nclreg. It could be object of cv.nclreg.
y	response y as in nclreg.
weights	Observation weights; defaults to 1 per observation
offset	Not implemented yet
object	object of cv.nclreg
which	Indices of the penalty parameter lambda at which estimates are extracted. By default, the one which generates the optimal cross-validation value.
se	logical value, if TRUE, standard error curve is also plotted
ylab	ylab on y-axis
main	title of plot
width	width of lines
col	color of standard error curve
...	Other arguments that can be passed to nclreg.

Details

The function runs nclreg `nfolds+1` times; the first to compute the lambda sequence, and then to compute the fit with each of the folds omitted. The error or the loss value is accumulated, and the average value and standard deviation over the folds is computed. Note that cv.nclreg can be used to search for values for alpha: it is required to call cv.nclreg with a fixed vector `foldid` for different values of alpha.

Value

an object of class "cv.nclreg" is returned, which is a list with the ingredients of the cross-validation fit.

fit	a fitted nclreg object for the full data.
residmat	matrix of log-likelihood values with row values for lambda and column values for kth cross-validation
bic	matrix of BIC values with row values for lambda and column values for kth cross-validation
cv	The mean cross-validated log-likelihood values - a vector of length length(lambda).
cv.error	estimate of standard error of cv.
foldid	an optional vector of values between 1 and nfold identifying what fold each observation is in.
lambda	a vector of lambda values
lambda.which	index of lambda that gives minimum cv value.
lambda.optim	value of lambda that gives minimum cv value.

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

See Also

[nclreg](#) and [plot](#), [predict](#), and [coef](#) methods for "cv.nclreg" object.

cv.nclreg_fit

Internal function of cross-validation for nclreg

Description

Internal function to conduct k-fold cross-validation for nclreg, produces a plot, and returns cross-validated log-likelihood values for lambda

Usage

```
cv.nclreg_fit(x, y, weights, lambda=NULL, balance=TRUE,
rfamily=c("clossR", "closs", "gloss", "qloss"), s=1.5, nolds=10, foldid,
type = c("loss", "error"), plot.it=TRUE, se=TRUE, n.cores=2, ...)
```

Arguments

x	x matrix as in nclreg.
y	response y as in nclreg.
weights	Observation weights; defaults to 1 per observation
lambda	Optional user-supplied lambda sequence; default is NULL, and nclreg chooses its own sequence
balance	for rfamily="closs", "gloss", "qloss" only
rfamily	response variable distribution and nonconvex loss function
s	nonconvex loss tuning parameter for robust regression and classification.
nfolds	number of folds ≥ 3 , default is 10
foldid	an optional vector of values between 1 and nfold identifying what fold each observation is in. If supplied, nfold can be missing and will be ignored.
type	cross-validation criteria. For type="loss", loss function values and type="error" is misclassification error.
plot.it	a logical value, to plot the estimated log-likelihood values if TRUE.
se	a logical value, to plot with standard errors.
n.cores	The number of CPU cores to use. The cross-validation loop will attempt to send different CV folds off to different cores.
...	Other arguments that can be passed to nclreg.

Details

The function runs nclreg nfolds+1 times; the first to compute the lambda sequence, and then to compute the fit with each of the folds omitted. The error or the log-likelihood value is accumulated, and the average value and standard deviation over the folds is computed. Note that cv.nclreg can be used to search for values for alpha: it is required to call cv.nclreg with a fixed vector foldid for different values of alpha.

Value

an object of class "cv.nclreg" is returned, which is a list with the ingredients of the cross-validation fit.

fit	a fitted nclreg object for the full data.
residmat	matrix of log-likelihood values with row values for lambda and column values for kth cross-validation
cv	The mean cross-validated log-likelihood values - a vector of length length(lambda).
cv.error	estimate of standard error of cv.
foldid	an optional vector of values between 1 and nfold identifying what fold each observation is in.
lambda	a vector of lambda values
lambda.which	index of lambda that gives minimum cv value.
lambda.optim	value of lambda that gives minimum cv value.

Author(s)

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See Also

[nclreg](#) and [plot](#), [predict](#), and [coef](#) methods for "cv.nclreg" object.

cv.zipath

Cross-validation for zipath

Description

Does k-fold cross-validation for zipath, produces a plot, and returns cross-validated log-likelihood values for lambda

Usage

```
cv.zipath(formula, data, weights, nlambda=100, lambda.count=NULL, lambda.zero=NULL,
n folds=10, foldid, plot.it=TRUE, se=TRUE, n.cores=2, ...)
## S3 method for class 'cv.zipath'
coef(object, which=object$lambda.which, model = c("full", "count", "zero"), ...)
```

Arguments

formula	symbolic description of the model
data	arguments controlling formula processing via model.frame .
weights	Observation weights; defaults to 1 per observation
nlambda	number of lambda value, default value is 10.
lambda.count	Optional user-supplied lambda.count sequence; default is NULL
lambda.zero	Optional user-supplied lambda.zero sequence; default is NULL
n folds	number of folds ≥ 3 , default is 10
foldid	an optional vector of values between 1 and nfold identifying what fold each observation is in. If supplied, nfold can be missing and will be ignored.
plot.it	a logical value, to plot the estimated log-likelihood values if TRUE.
se	a logical value, to plot with standard errors.
n.cores	The number of CPU cores to use. The cross-validation loop will attempt to send different CV folds off to different cores.
...	Other arguments that can be passed to zipath.
object	object of class cv.zipath.
which	Indices of the pair of penalty parameters lambda.count and lambda.zero at which estimates are extracted. By default, the one which generates the optimal cross-validation value.
model	character specifying for which component of the model the estimated coefficients should be extracted.

Details

The function runs `zipath` `nfolds+1` times; the first to compute the `(lambda.count, lambda.zero)` sequence, and then to compute the fit with each of the folds omitted. The log-likelihood value is accumulated, and the average value and standard deviation over the folds is computed. Note that `cv.zipath` can be used to search for values for `count.alpha` or `zero.alpha`: it is required to call `cv.zipath` with a fixed vector `foldid` for different values of `count.alpha` or `zero.alpha`.

The method for `coef` by default return a single vector of coefficients, i.e., all coefficients are concatenated. By setting the `model` argument, the estimates for the corresponding model components can be extracted.

Value

an object of class "cv.zipath" is returned, which is a list with the components of the cross-validation fit.

<code>fit</code>	a fitted zipath object for the full data.
<code>residmat</code>	matrix for cross-validated log-likelihood at each <code>(count.lambda, zero.lambda)</code> sequence
<code>bic</code>	matrix of BIC values with row values for <code>lambda</code> and column values for <code>kth</code> cross-validation
<code>cv</code>	The mean cross-validated log-likelihood - a vector of length <code>length(count.lambda)</code> .
<code>cv.error</code>	estimate of standard error of <code>cv</code> .
<code>foldid</code>	an optional vector of values between 1 and <code>nfold</code> identifying what fold each observation is in.
<code>lambda.which</code>	index of <code>(count.lambda, zero.lambda)</code> that gives maximum <code>cv</code> .
<code>lambda.optim</code>	value of <code>(count.lambda, zero.lambda)</code> that gives maximum <code>cv</code> .

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

- Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]
- Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.
- Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

`zipath` and `plot`, `predict`, and `coef` methods for "cv.zipath" object.

Examples

```
## Not run:
data("bioChemists", package = "pscl")
fm_zip <- cv.zipath(art ~ . | ., data = bioChemists, family = "poisson", nlambda=10)
coef(fm_zip)
### prediction from the best model
fm_zip_predict <- predict(object=fm_zip$fit, which=fm_zip$lambda.which, type="response",
model=c("full"))
fm_znb <- cv.zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambda=10)
coef(fm_znb)

## End(Not run)
```

estfunReg

Extract Empirical First Derivative of Log-likelihood Function

Description

Generic function for extracting the empirical first derivative of log-likelihood function of a fitted regularized model.

Usage

```
estfunReg(x, ...)
```

Arguments

x	a fitted model object.
...	arguments passed to methods.

Value

A matrix containing the empirical first derivative of log-likelihood functions. Typically, this should be an $n \times k$ matrix corresponding to n observations and k parameters. The columns should be named as in [coef](#) or [terms](#), respectively.

Author(s)

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References

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[zipath](#)

Examples

```
data("bioChemists", package = "pscl")
fm_zinb <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambdas=10)
res <- estfunReg(fm_zinb, which=which.min(fm_zinb$bic))
```

glmreg

*fit a GLM with lasso (or elastic net), snet or mnet regularization***Description**

Fit a generalized linear model via penalized maximum likelihood. The regularization path is computed for the lasso (or elastic net penalty), scad (or snet) and mcp (or mnet penalty), at a grid of values for the regularization parameter lambda. Fits linear, logistic, Poisson and negative binomial (fixed scale parameter) regression models.

Usage

```
## S3 method for class 'formula'
glmreg(formula, data, weights, offset=NULL, contrasts=NULL,
x.keep=FALSE, y.keep=TRUE, ...)
## S3 method for class 'matrix'
glmreg(x, y, weights, offset=NULL, ...)
## Default S3 method:
glmreg(x, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via model.frame .
weights	optional numeric vector of weights. If standardize=TRUE, weights are renormalized to weights/sum(weights). If standardize=FALSE, weights are kept as original input
x	input matrix, of dimension nobs x nvars; each row is an observation vector
y	response variable. Quantitative for family="gaussian". Non-negative counts for family="poisson" or family="negbin". For family="binomial" should be either a factor with two levels or a vector of proportions.
x.keep, y.keep	logical values: keep response variables or keep response variable?
offset	Not implemented yet
contrasts	the contrasts corresponding to levels from the respective models
...	Other arguments passing to glmreg_fit

Details

The sequence of models implied by lambda is fit by coordinate descent. For family="gaussian" this is the lasso, mcp or scad sequence if alpha=1, else it is the enet, mnet or snet sequence. For the other families, this is a lasso (mcp, scad) or elastic net (mnet, snet) regularization path for fitting the generalized linear regression paths, by maximizing the appropriate penalized log-likelihood. Note that the objective function for "gaussian" is

$$1/2 * weights * RSS + \lambda * penalty,$$

if standardize=FALSE and

$$1/2 * \frac{weights}{\sum(weights)} * RSS + \lambda * penalty,$$

if standardize=TRUE. For the other models it is

$$- \sum(weights * loglik) + \lambda * penalty$$

if standardize=FALSE and

$$- \frac{weights}{\sum(weights)} * loglik + \lambda * penalty$$

if standardize=TRUE.

Value

An object with S3 class "glmreg" for the various types of models.

call	the call that produced this object
b0	Intercept sequence of length length(lambda)
beta	A nvars x length(lambda) matrix of coefficients.
lambda	The actual sequence of lambda values used
dev	The computed deviance (for "gaussian", this is the R-square). The deviance calculations incorporate weights if present in the model. The deviance is defined to be 2*(loglike_sat - loglike), where loglike_sat is the log-likelihood for the saturated model (a model with a free parameter per observation).
nulldev	Null deviance (per observation). This is defined to be 2*(loglike_sat - loglike(NULL)); The NULL model refers to the intercept model.
nobs	number of observations
p11	penalized log-likelihood values for standardized coefficients in the IRLS iterations. For family="gaussian", not implemented yet.
p11res	penalized log-likelihood value for the estimated model on the original scale of coefficients
fitted.values	predicted values depending on standardize, internal use only

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Breheny, P. and Huang, J. (2011) *Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection*. *Ann. Appl. Statist.*, **5**: 232-253.

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[print](#), [predict](#), [coef](#) and [plot](#) methods, and the [cv.glmreg](#) function.

Examples

```
#binomial
x=matrix(rnorm(100*20),100,20)
g2=sample(0:1,100,replace=TRUE)
fit2=glmreg(x,g2,family="binomial")
#poisson and negative binomial
data("bioChemists", package = "pscl")
fm_pois <- glmreg(art ~ ., data = bioChemists, family = "poisson")
coef(fm_pois)
fm_nb1 <- glmreg(art ~ ., data = bioChemists, family = "negbin", theta=1)
coef(fm_nb1)
## Not run:
fm_nb2 <- glmregNB(art ~ ., data = bioChemists)
coef(fm_nb2)

## End(Not run)
```

glmregNB

fit a negative binomial model with lasso (or elastic net), snet and mnet regularization

Description

Fit a negative binomial linear model via penalized maximum likelihood. The regularization path is computed for the lasso (or elastic net penalty), snet and mnet penalty, at a grid of values for the regularization parameter lambda.

Usage

```
glmregNB(formula, data, weights, nlambda = 100, lambda=NULL, lambda.min.ratio =
ifelse(nobs<nvars,0.05,0.001), alpha=1, gamma=3, rescale=TRUE, standardize = TRUE,
penalty.factor = rep(1, nvars), thresh = 0.001, maxit.theta = 25, maxit=1000,
eps=.Machine$double.eps, trace=FALSE, start = NULL, etastart = NULL, mustart = NULL,
theta.est=TRUE, theta0=NULL, init.theta=ifelse(theta.est, theta0[1],NULL),link=log,
penalty=c("enet","mnet","snet"), method="glmreg_fit", model=TRUE,
x.keep=FALSE, y.keep=TRUE, contrasts=NULL, convex=FALSE, ...)
```

Arguments

formula	formula used to describe a model.
data	argument controlling formula processing via <code>model.frame</code> .
weights	observation weights. Default is 1 for each observation
nlambda	The number of lambda values - default is 100.
lambda	A user supplied lambda sequence
lambda.min.ratio	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero). The default depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is 0.001, close to zero. If nobs < nvars, the default is 0.05.
alpha	The L2 penalty mixing parameter, with $0 \leq \alpha \leq 1$. alpha=1 is lasso (mcp, scad) penalty; and alpha=0 the ridge penalty.
gamma	The tuning parameter of the snet or mnet penalty.
rescale	logical value, if TRUE, adaptive rescaling of the penalty parameter for penalty="mnet" or penalty="snet" with family other than "gaussian". See reference
standardize	Logical flag for x variable standardization, prior to fitting the model sequence. The coefficients are always returned on the original scale. Default is standardize=TRUE. If variables are in the same units already, you might not wish to standardize.
penalty.factor	This is a number that multiplies lambda to allow differential shrinkage of coefficients. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is same shrinkage for all variables.
thresh	Convergence threshold for coordinate descent. Defaults value is 1e-6.
maxit.theta	Maximum number of iterations for estimating theta scaling parameter
maxit	Maximum number of coordinate descent iterations for each lambda value; default is 1000.
eps	If a number is less than eps in magnitude, then this number is considered as 0
trace	If TRUE, fitting progress is reported
start, etastart, mustart, ...	arguments for the <code>link{glmreg}</code> function
init.theta	initial scaling parameter theta
theta.est	Estimate scale parameter theta? Default is TRUE. Note, the algorithm may become slow. In this case, one may use glmreg function with family="negbin", and a fixed theta.
theta0	initial scale parameter vector theta, with length nlambda if theta.est=FALSE. Default is NULL
convex	Calculate index for which objective function ceases to be locally convex? Default is FALSE and only useful if penalty="mnet" or "snet".
link	link function, default is log

penalty	Type of regularization
method	estimation method
model, x.keep, y.keep	logicals. If TRUE the corresponding components of the fit (model frame, response, model matrix) are returned.
contrasts	the contrasts corresponding to levels from the respective models

Details

The sequence of models implied by lambda is fit by coordinate descent. This is a lasso (mcp, scad) or elastic net (mnet, snet) regularization path for fitting the negative binomial linear regression paths, by maximizing the penalized log-likelihood. Note that the objective function is

$$- \sum(weights * loglik) + \lambda * penalty$$

if standardize=FALSE and

$$- \frac{weights}{\sum(weights)} * loglik + \lambda * penalty$$

if standardize=TRUE.

Value

An object with S3 class "glmreg", "glmregNB" for the various types of models.

call	the call that produced the model fit
b0	Intercept sequence of length length(lambda)
beta	A nvars x length(lambda) matrix of coefficients.
lambda	The actual sequence of lambda values used
dev	The computed deviance. The deviance calculations incorporate weights if present in the model. The deviance is defined to be 2*(loglike_sat - loglike), where loglike_sat is the log-likelihood for the saturated model (a model with a free parameter per observation).
nulldev	Null deviance (per observation). This is defined to be 2*(loglike_sat - loglike(NULL)); The NULL model refers to the intercept model.
nobs	number of observations

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

- Breheny, P. and Huang, J. (2011) *Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection*. *Ann. Appl. Statist.*, **5**: 232-253.
- Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[print](#), [predict](#), [coef](#) and [plot](#) methods, and the [cv.glmregNB](#) function.

Examples

```
## Not run:
data("bioChemists", package = "pscl")
fm_nb <- glmregNB(art ~ ., data = bioChemists)
coef(fm_nb)
### ridge regression
fm <- glmregNB(art ~ ., alpha=0, data = bioChemists, lambda=seq(0.001, 1, by=0.01))
fm <- cv.glmregNB(art ~ ., alpha=0, data = bioChemists, lambda=seq(0.001, 1, by=0.01))

## End(Not run)
```

glmreg_fit

Internal function to fit a GLM with lasso (or elastic net), snet and mnet regularization

Description

Fit a generalized linear model via penalized maximum likelihood. The regularization path is computed for the lasso (or elastic net penalty), snet and mnet penalty, at a grid of values for the regularization parameter lambda. Fits linear, logistic, Poisson and negative binomial (fixed scale parameter) regression models.

Usage

```
glmreg_fit(x, y, weights, start=NULL, etastart=NULL, mustart=NULL,
nlambda=100, lambda=NULL, lambda.min.ratio=ifelse(nobs<nvars,.05,.001),alpha=1,
gamma=3, rescale=TRUE, standardize=TRUE, penalty.factor = rep(1, nvars),thresh=1e-6,
eps.bino=1e-5, maxit=1000, eps=.Machine$double.eps, theta,
family=c("gaussian", "binomial", "poisson", "negbin"), penalty=c("enet", "mnet", "snet"),
convex=FALSE, x.keep=FALSE, y.keep=TRUE, trace=FALSE)
```

Arguments

x	input matrix, of dimension nobs x nvars; each row is an observation vector.
y	response variable. Quantitative for family="gaussian". Non-negative counts for family="poisson" or family="negbin". For family="binomial" should be either a factor with two levels or a vector of proportions.
weights	observation weights. Can be total counts if responses are proportion matrices. Default is 1 for each observation
start	starting values for the parameters in the linear predictor.
etastart	starting values for the linear predictor.
mustart	starting values for the vector of means.

nlambda	The number of lambda values - default is 100. The sequence may be truncated before nlambda is reached if a close to saturated model is fitted. See also satu.
lambda	by default, the algorithm provides a sequence of regularization values, or a user supplied lambda sequence
lambda.min.ratio	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero except the intercept). Note, there is no closed formula for lambda.max in general. If rescale=TRUE, lambda.max is the same for penalty="mnet" or "snet". Otherwise, some modifications are required. For instance, for small gamma value, half of the square root (if lambda.max is too small) of the computed lambda.max can be used when penalty="mnet" or "snet". The default of lambda.min.ratio depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is 0.001, close to zero. If nobs < nvars, the default is 0.05.
alpha	The L_2 penalty mixing parameter, with $0 \leq \alpha \leq 1$. alpha=1 is lasso (mcp, scad) penalty; and alpha=0 the ridge penalty. However, if alpha=0, one must provide lambda values.
gamma	The tuning parameter of the snet or mnet penalty.
rescale	logical value, if TRUE, adaptive rescaling of the penalty parameter for penalty="mnet" or penalty="snet" with family other than "gaussian". See reference
standardize	logical value for x variable standardization, prior to fitting the model sequence. The coefficients are always returned on the original scale. Default is standardize=TRUE.
penalty.factor	This is a number that multiplies lambda to allow differential shrinkage of coefficients. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is same shrinkage for all variables.
thresh	Convergence threshold for coordinate descent. Defaults value is 1e-6.
eps.bino	a lower bound of probabilities to be claimed as zero, for computing weights and related values when family="binomial".
maxit	Maximum number of coordinate descent iterations for each lambda value; default is 1000.
eps	If a coefficient is less than eps in magnitude, then it is reported to be 0
convex	Calculate index for which objective function ceases to be locally convex? Default is FALSE and only useful if penalty="mnet" or "snet".
theta	an overdispersion scaling parameter for family="negbin"
family	Response type (see above)
penalty	Type of regularization
x.keep, y.keep	For glmreg: logical values indicating whether the response vector and model matrix used in the fitting process should be returned as components of the returned value. For glmreg_fit: x is a design matrix of dimension n * p, and x is a vector of observations of length n.
trace	If TRUE, fitting progress is reported

Details

The sequence of models implied by lambda is fit by coordinate descent. For family="gaussian" this is the lasso, mcp or scad sequence if alpha=1, else it is the enet, mnet or snet sequence. For the other families, this is a lasso (mcp, scad) or elastic net (mnet, snet) regularization path for fitting the generalized linear regression paths, by maximizing the appropriate penalized log-likelihood. Note that the objective function for "gaussian" is

$$1/2 * weights * RSS + \lambda * penalty,$$

if standardize=FALSE and

$$1/2 * \frac{weights}{\sum(weights)} * RSS + \lambda * penalty,$$

if standardize=TRUE. For the other models it is

$$- \sum(weights * loglik) + \lambda * penalty$$

if standardize=FALSE and

$$- \frac{weights}{\sum(weights)} * loglik + \lambda * penalty$$

if standardize=TRUE.

Value

An object with S3 class "glmreg" for the various types of models.

call	the call that produced the model fit
b0	Intercept sequence of length length(lambda)
beta	A nvars x length(lambda) matrix of coefficients.
lambda	The actual sequence of lambda values used
satu	satu=1 if a saturated model (deviance/null deviance < 0.05) is fit. Otherwise satu=0. The number of nlambda sequence may be truncated before nlambda is reached if satu=1.
dev	The computed deviance (for "gaussian", this is the R-square). The deviance calculations incorporate weights if present in the model. The deviance is defined to be 2*(loglike_sat - loglike), where loglike_sat is the log-likelihood for the saturated model (a model with a free parameter per observation).
nulldev	Null deviance (per observation). This is defined to be 2*(loglike_sat - loglike(NULL)); The NULL model refers to the intercept model.
nobs	number of observations

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Breheeny, P. and Huang, J. (2011) *Coordinate descent algorithms for nonconvex penalized regression, with applications to biological feature selection*. *Ann. Appl. Statist.*, **5**: 232-253.

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[glmreg](#)

hessianReg

Hessian Matrix of Regularized Estimators

Description

Constructing Hessian matrix for regularized regression parameters.

Usage

```
hessianReg(x, which, ...)
```

Arguments

x	a fitted model object.
which	which penalty parameter(s)?
...	arguments passed to the meatReg function.

Details

hessianReg is a function to compute the Hessian matrix estimate of non-zero regularized estimators. Implemented only for zipath object with family="negbin" in the current version.

Value

A matrix containing the Hessian matrix estimate for the non-zero parameters.

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[breadReg](#), [meatReg](#)

Examples

```
data("bioChemists", package = "pscl")
fm_zinb <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambd=10)
hessianReg(fm_zinb, which=which.min(fm_zinb$bic))
```

meatReg

Meat Matrix Estimator

Description

Estimating the variance of the first derivative of log-likelihood function

Usage

```
meatReg(x, which, ...)
```

Arguments

`x` a fitted model object. Currently only implemented for zipath object with family="negbin"
`which` which penalty parameter(s)?
`...` arguments passed to the [estfunReg](#) function.

Details

See reference below

Value

A $k \times k$
 covariance matrix of first derivative of log-likelihood function

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[sandwichReg](#), [breadReg](#), [estfunReg](#)

Examples

```
data("bioChemists", package = "pscl")
fm_zinb <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambda=10)
meatReg(fm_zinb, which=which.min(fm_zinb$bic))
```

methods

Methods for mpath Objects

Description

Methods for models fitted by coordinate descent algorithms.

Usage

```
## S3 method for class 'glmreg'
AIC(object, ..., k)
## S3 method for class 'zipath'
AIC(object, ..., k)
## S3 method for class 'glmreg'
BIC(object, ...)
## S3 method for class 'zipath'
BIC(object, ...)
```

Arguments

object	objects of class glmreg or zipath.
...	additional arguments passed to callies.
k	numeric, the <i>penalty</i> per parameter to be used; the default k = 2 is the classical AIC. k has been hard coded in the function and there is no impact to the value of AIC if k is changed

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

ncl *fit a nonconvex loss based robust linear model*

Description

Fit a linear model via penalized nonconvex loss function.

Usage

```
## S3 method for class 'formula'
ncl(formula, data, weights, offset=NULL, contrasts=NULL,
x.keep=FALSE, y.keep=TRUE, ...)
## S3 method for class 'matrix'
ncl(x, y, weights, offset=NULL, ...)
## Default S3 method:
ncl(x, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via <code>model.frame</code> .
weights	optional numeric vector of weights. If <code>standardize=TRUE</code> , weights are renormalized to <code>weights/sum(weights)</code> . If <code>standardize=FALSE</code> , weights are kept as original input
x	input matrix, of dimension <code>nobs x nvars</code> ; each row is an observation vector
y	response variable. Quantitative for <code>rfamily="clossR"</code> and <code>-1/1</code> for classification.
offset	Not implemented yet
contrasts	the contrasts corresponding to levels from the respective models
x.keep, y.keep	For <code>glmreg</code> : logical values indicating whether the response vector and model matrix used in the fitting process should be returned as components of the returned value. For <code>ncl_fit</code> : x is a design matrix of dimension <code>n * p</code> , and x is a vector of observations of length n.
...	Other arguments passing to <code>ncl_fit</code>

Details

The robust linear model is fit by majorization-minimization along with linear regression. Note that the objective function is

$$1/2 * weights * loss$$

Value

An object with S3 class "ncl" for the various types of models.

call	the call that produced this object
fitted.values	predicted values
h	pseudo response values in the MM algorithm

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

See Also

[print](#), [predict](#), [coef](#).

Examples

```
#binomial
x=matrix(rnorm(100*20),100,20)
g2=sample(c(-1,1),100,replace=TRUE)
fit=ncl(x,g2,s=1,rfamily="closs")
```

nclreg	<i>fit a nonconvex loss based robust linear model with lasso (or elastic net), snet or mnet regularization</i>
--------	--

Description

Fit a linear model via penalized nonconvex loss function. The regularization path is computed for the lasso (or elastic net penalty), scad (or snet) and mcp (or mnet penalty), at a grid of values for the regularization parameter lambda.

Usage

```
## S3 method for class 'formula'
nclreg(formula, data, weights, offset=NULL, contrasts=NULL, ...)
## S3 method for class 'matrix'
nclreg(x, y, weights, offset=NULL, ...)
## Default S3 method:
nclreg(x, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via <code>model.frame</code> .
weights	optional numeric vector of weights. If <code>standardize=TRUE</code> , weights are renormalized to <code>weights/sum(weights)</code> . If <code>standardize=FALSE</code> , weights are kept as original input
x	input matrix, of dimension <code>nobs x nvars</code> ; each row is an observation vector
y	response variable. Quantitative for <code>rfamily="clossR"</code> and <code>-1/1</code> for classification.
offset	Not implemented yet
contrasts	the contrasts corresponding to levels from the respective models
...	Other arguments passing to <code>nclreg_fit</code>

Details

The sequence of robust models implied by `lambda` is fit by majorization-minimization along with coordinate descent. Note that the objective function is

$$1/2 * weights * loss + \lambda * penalty,$$

if `standardize=FALSE` and

$$1/2 * \frac{weights}{\sum(weights)} * loss + \lambda * penalty,$$

if `standardize=TRUE`.

Value

An object with S3 class "nclreg" for the various types of models.

call	the call that produced this object
b0	Intercept sequence of length <code>length(lambda)</code>
beta	A <code>nvars x length(lambda)</code> matrix of coefficients.
lambda	The actual sequence of <code>lambda</code> values used
nobs	number of observations
risk	if <code>type.path="naive"</code> , a matrix with number of rows <code>iter</code> and number of columns <code>nlambda</code> , loss values along the regularization path. If <code>type.path="fast"</code> , a vector of length <code>nlambda</code> , loss values along the regularization path
p11	if <code>type.path="naive"</code> , a matrix with number of rows <code>iter</code> and number of columns <code>nlambda</code> , penalized loss values along the regularization path. If <code>type.path="fast"</code> , a vector of length <code>nlambda</code> , penalized loss values along the regularization path
fitted.values	predicted values depending on <code>standardize</code> , internal use only

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

See Also

[print](#), [predict](#), [coef](#) and [plot](#) methods, and the [cv.nclreg](#) function.

Examples

```
#binomial
x=matrix(rnorm(100*20),100,20)
g2=sample(c(-1,1),100,replace=TRUE)
fit=nclreg(x,g2,s=1,rfamily="closs")
```

nclreg_fit	<i>Internal function to fit a nonconvex loss based robust linear model with lasso (or elastic net), snet and mnet regularization</i>
------------	--

Description

Fit a linear model via penalized nonconvex loss function. The regularization path is computed for the lasso (or elastic net penalty), scad (or snet) and mcp (or mnet penalty), at a grid of values for the regularization parameter lambda.

Usage

```
nclreg_fit(x,y, weights, cost=0.5, rfamily=c("clossR", "closs", "gloss", "qloss"),
s=NULL, fk=NULL, iter=10, del=1e-10, nlambdas=100, lambda=NULL, lambda.min.ratio=
ifelse(nobs<nvars,.05, .001),alpha=1, gamma=3, standardize=TRUE, penalty.factor = NULL,
maxit=1000, type.init="bst", mstop.init=10, nu.init=0.1, direction="bwd",
eps=.Machine$double.eps, trace=FALSE, penalty=c("enet","mnet","snet"), type.path="naive")
```

Arguments

x	input matrix, of dimension nobs x nvars; each row is an observation vector.
y	response variable. Quantitative for rfamily="clossR" and -1/1 for classifications.
weights	observation weights. Can be total counts if responses are proportion matrices. Default is 1 for each observation
cost	price to pay for false positive, $0 < \text{cost} < 1$; price of false negative is $1 - \text{cost}$.
rfamily	Response type and relevant loss functions (see above)
s	nonconvex loss tuning parameter for robust regression and classification.
fk	predicted values at an iteration in the MM algorithm
nlambdas	The number of lambda values - default is 100. The sequence may be truncated before nlambdas is reached if a close to saturated model is fitted. See also satu.
lambda	by default, the algorithm provides a sequence of regularization values, or a user supplied lambda sequence

lambda.min.ratio	Smallest value for lambda, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero except the intercept). Note, there is no closed formula for lambda.max. The default of lambda.min.ratio depends on the sample size nobs relative to the number of variables nvars. If nobs > nvars, the default is 0.001, close to zero. If nobs < nvars, the default is 0.05.
alpha	The L_2 penalty mixing parameter, with $0 \leq \alpha \leq 1$. alpha=1 is lasso (mcp, scad) penalty; and alpha=0 the ridge penalty. However, if alpha=0, one must provide lambda values.
gamma	The tuning parameter of the snet or mnet penalty.
standardize	logical value for x variable standardization, prior to fitting the model sequence. The coefficients are always returned on the original scale. Default is standardize=TRUE.
penalty.factor	This is a number that multiplies lambda to allow differential shrinkage of coefficients. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is same shrinkage for all variables.
type.init	a method to determine the initial values. If type.init="ncl", an intercept-only model as initial parameter and run nclreg regularization path forward from lambda_max to lambda_min. If type.init="heu", heuristic initial parameters and run nclreg path backward or forward depending on direction, between lambda_min and lambda_max. If type.init="bst", run a boosting model with bst in package bst, depending on mstop.init, nu.init and run nclreg backward or forward depending on direction.
mstop.init	an integer giving the number of boosting iterations when type.init="bst"
nu.init	a small number (between 0 and 1) defining the step size or shrinkage parameter when type.init="bst".
direction	only used if lambda=NULL. direction="bwd" for backward or "fwd" for forward, used to determine regularization path direction either from lambda_max to a potentially modified lambda_min or vice versa if type.init="bst", "heu".
iter	number of iteration in the MM algorithm
maxit	Within each MM algorithm iteration, maximum number of coordinate descent iterations for each lambda value; default is 1000.
del	convergency criteria
eps	If a coefficient is less than eps in magnitude, then it is reported to be 0
penalty	Type of regularization
type.path	solution path. If type.path="naive", for each element of the lambda sequence, iterate until convergency. If type.path="onestep", update for one element of lambda depending on direction="fwd" (last element of lambda) or "bwd" (then first element of lambda) in each MM iteration, and iterate until convergency of prediction. Then fit a solution path based on the sequence of lambda.
trace	If TRUE, fitting progress is reported

Details

The sequence of robust models implied by `lambda` is fit by majorization-minimization along with coordinate descent. Note that the objective function is

$$1/2 * weights * loss + \lambda * penalty,$$

if `standardize=FALSE` and

$$1/2 * \frac{weights}{\sum(weights)} * loss + \lambda * penalty,$$

if `standardize=TRUE`.

Value

An object with S3 class "nclreg" for the various types of models.

<code>call</code>	the call that produced the model fit
<code>b0</code>	Intercept sequence of length <code>length(lambda)</code>
<code>beta</code>	A <code>nvars x length(lambda)</code> matrix of coefficients.
<code>lambda</code>	The actual sequence of <code>lambda</code> values used

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

See Also

[nclreg](#)

ncl_fit

Internal function to fit a nonconvex loss based robust linear model

Description

Fit a linear model via penalized nonconvex loss function.

Usage

```
ncl_fit(x,y, weights, cost=0.5, rfamily=c("clossR", "closs", "gloss", "qloss"), s=NULL,
fk=NULL, iter=10, del=1e-10, trace=FALSE)
```

Arguments

x	input matrix, of dimension nobs x nvars; each row is an observation vector.
y	response variable. Quantitative for rfamily="clossR" and -1/1 for classifications.
weights	observation weights. Can be total counts if responses are proportion matrices. Default is 1 for each observation
cost	price to pay for false positive, $0 < \text{cost} < 1$; price of false negative is $1 - \text{cost}$.
rfamily	Response type and relevant loss functions (see above)
s	nonconvex loss tuning parameter for robust regression and classification.
fk	predicted values at an iteration in the MM algorithm
iter	number of iteration in the MM algorithm
del	convergency criteria
trace	If TRUE, fitting progress is reported

Details

The robust linear model is fit by majorization-minimization along with least squares. Note that the objective function is

$$1/2 * weights * loss$$

.

Value

An object with S3 class "ncl" for the various types of models.

call	the call that produced the model fit
fitted.values	predicted values
h	pseudo response values in the MM algorithm

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

See Also

[ncl](#)

plot.glmreg *plot coefficients from a "glmreg" object*

Description

Produces a coefficient profile plot of the coefficient paths for a fitted "glmreg" object.

Usage

```
## S3 method for class 'glmreg'  
plot(x, xvar = c("norm", "lambda", "dev"), label = FALSE, shade=TRUE, ...)
```

Arguments

x	fitted "glmreg" model
xvar	What is on the X-axis. "norm" plots against the L1-norm of the coefficients, "lambda" against the log-lambda sequence, and "dev" against the percent deviance explained.
label	If TRUE, label the curves with variable sequence numbers.
shade	Should nonconvex region be shaded? Default is TRUE. Code developed for all weights=1 only
...	Other graphical parameters to plot

Details

A coefficient profile plot is produced.

Author(s)

Zhu Wang zwang@connecticutchildrens.org

See Also

glmreg, and print, predict and coef methods.

Examples

```
x=matrix(rnorm(100*20),100,20)  
y=rnorm(100)  
fit1=glmreg(x,y)  
plot(fit1)  
plot(fit1,xvar="lambda",label=TRUE)
```

predict.glmreg *Model predictions based on a fitted "glmreg" object.*

Description

This function returns predictions from a fitted "glmreg" object.

Usage

```
## S3 method for class 'glmreg'
predict(object,newx,which=1:length(object$lambda),
type=c("link","response","class","coefficients","nonzero"), na.action=na.pass, ...)
## S3 method for class 'glmreg'
coef(object,which=1:length(object$lambda),...)
```

Arguments

object	Fitted "glmreg" model object.
newx	Matrix of values at which predictions are to be made. Not used for type="coefficients"
which	Indices of the penalty parameter lambda at which predictions are required. By default, all indices are returned.
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.
na.action	action for missing data value
...	arguments for predict

Value

The returned object depends on type.

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[glmreg](#)

Examples

```
## Dobson (1990) Page 93: Randomized Controlled Trial :
counts <- c(18,17,15,20,10,20,25,13,12)
outcome <- gl(3,1,9)
treatment <- gl(3,3)
print(d.AD <- data.frame(treatment, outcome, counts))
fit <- glmreg(counts ~ outcome + treatment, data=d.AD, family="poisson")
summary(fit)
coef(fit)
```

predict.zipath *Methods for zipath Objects*

Description

Methods for extracting information from fitted penalized zero-inflated regression model objects of class "zipath".

Usage

```
## S3 method for class 'zipath'
predict(object, newdata, which = 1:object$nlambda,
        type = c("response", "prob", "count", "zero", "nonzero"), na.action = na.pass,
        at = NULL, ...)
## S3 method for class 'zipath'
residuals(object, type = c("pearson", "response"), ...)

## S3 method for class 'zipath'
coef(object, which=1:object$nlambda, model = c("full", "count", "zero"), ...)

## S3 method for class 'zipath'
terms(x, model = c("count", "zero"), ...)
## S3 method for class 'zipath'
model.matrix(object, model = c("count", "zero"), ...)
```

Arguments

object, x	an object of class "zipath" as returned by zipath .
newdata	optionally, a data frame in which to look for variables with which to predict. If omitted, the original observations are used.
which	Indices of the penalty parameters lambda at which predictions are required. By default, all indices are returned.
type	character specifying the type of predictions or residuals, respectively. For details see below.
na.action	function determining what should be done with missing values in newdata. The default is to predict NA.

at	optionally, if type = "prob", a numeric vector at which the probabilities are evaluated. By default $0:\max(y)$ is used where y is the original observed response.
model	character specifying for which component of the model the terms or model matrix should be extracted.
...	currently not used.

Details

Re-uses the design of function `zeroinfl` in package `pscl` (see reference). A set of standard extractor functions for fitted model objects is available for objects of class "zipath", including methods to the generic functions `print` and `summary` which print the estimated coefficients along with some further information. As usual, the `summary` method returns an object of class "summary.zipath" containing the relevant summary statistics which can subsequently be printed using the associated `print` method.

The methods for `coef` by default return a single vector of coefficients and their associated covariance matrix, respectively, i.e., all coefficients are concatenated. By setting the `model` argument, the estimates for the corresponding model components can be extracted.

Both the `fitted` and `predict` methods can compute fitted responses. The latter additionally provides the predicted density (i.e., probabilities for the observed counts), the predicted mean from the count component (without zero inflation) and the predicted probability for the zero component. The `residuals` method can compute raw residuals (observed - fitted) and Pearson residuals (raw residuals scaled by square root of variance function).

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[zipath](#)

Examples

```
## Not run:
data("bioChemists", package = "pscl")
fm_zip <- zipath(art ~ . | ., data = bioChemists, nlambda=10)
plot(residuals(fm_zip) ~ fitted(fm_zip))
coef(fm_zip, model = "count")
coef(fm_zip, model = "zero")
summary(fm_zip)
logLik(fm_zip)

## End(Not run)
```

pval.zipath	<i>compute p-values from penalized zero-inflated model with multi-split data</i>
-------------	--

Description

compute p-values from penalized zero-inflated Poisson, negative binomial and geometric model with multi-split data

Usage

```
pval.zipath(formula, data, weights, subset, na.action, offset, standardize=TRUE,
family = c("poisson", "negbin", "geometric"),penalty = c("enet", "mnet", "snet"),
gamma.count = 3, gamma.zero = 3, prop=0.5, trace=TRUE, B=10, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via <code>model.frame</code> .
weights	optional numeric vector of weights. If <code>standardize=TRUE</code> , weights are renormalized to <code>weights/sum(weights)</code> . If <code>standardize=FALSE</code> , weights are kept as original input
subset	subset of data
na.action	how to deal with missing data
offset	Not implemented yet
standardize	logical value, should variables be standardized?
family	family to fit zipath
penalty	penalty considered as one of <code>enet</code> , <code>mnet</code> , <code>snet</code> .
gamma.count	The tuning parameter of the <code>snet</code> or <code>mnet</code> penalty for the count part of model.
gamma.zero	The tuning parameter of the <code>snet</code> or <code>mnet</code> penalty for the zero part of model.
prop	proportion of data split, default is 50/50 split
trace	logical value, if <code>TRUE</code> , print detailed calculation results
B	number of repeated multi-split replications
...	Other arguments passing to <code>glmreg_fit</code>

Details

compute p-values from penalized zero-inflated Poisson, negative binomial and geometric model with multi-split data

Value

count.pval	raw p-values in the count component
zero.pval	raw p-values in the zero component
count.pval.q	Q value for the count component
zero.pval.q	Q value for the zero component

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Nicolai Meinshausen, Lukas Meier and Peter Buehlmann (2013) *p-Values for High-Dimensional Regression*, *Journal of the American Statistical Association*, **104(488)**, 1671–1681

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

rzi

random number generation of zero-inflated count response

Description

random number generation of zero-inflated count response

Usage

```
rzi(n, x, z, a, b, theta=1, family=c("poisson", "negbin", "geometric"), infl=TRUE)
```

Arguments

n	sample size of random number generation
x	design matrix of count model
z	design matrix of zero model
a	coefficient vector for x, length must be the same as column size of x
b	coefficient vector for z, length must be the same as column size of z

theta	dispersion parameter for family="negbin"
family	distribution of count model
infl	logical value, if TRUE, zero-inflated count response

Details

random number generation of zero-inflated count response

Value

numeric vector of zero-inflated count response

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

sandwichReg *Making Sandwiches with Bread and Meat for Regularized Estimators*

Description

Constructing sandwich covariance matrix estimators by multiplying bread and meat matrices for regularized regression parameters.

Usage

```
sandwichReg(x, breadreg.=breadReg, meatreg.=meatReg, which, log=FALSE, ...)
```

Arguments

x	a fitted model object.
breadreg.	either a breadReg matrix or a function for computing this via breadreg.(x).
meatreg.	either a breadReg matrix or a function for computing this via meatreg.(x, ...).
which	which penalty parameters(s) to compute?
log	if TRUE, the corresponding element is with respect to log(theta) in negative binomial regression. Otherwise, for theta
...	arguments passed to the meatReg function.

Details

`sandwichReg` is a function to compute an estimator for the covariance of the non-zero parameters. It takes a `breadReg` matrix (i.e., estimator of the expectation of the negative derivative of the penalized estimating functions) and a `meatReg` matrix (i.e., estimator of the variance of the log-likelihood function) and multiplies them to a sandwich with meat between two slices of bread. By default `breadReg` and `meatReg` are called. Implemented only for `zipath` object with `family="negbin"` in the current version.

Value

A matrix containing the sandwich covariance matrix estimate for the non-zero parameters.

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[breadReg](#), [meatReg](#)

Examples

```
data("bioChemists", package = "pscl")
fm_zinb <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambda=10)
sandwichReg(fm_zinb, which=which.min(fm_zinb$bic))
```

se

Standard Error of Regularized Estimators

Description

Generic function for computing standard errors of non-zero regularized estimators

Usage

```
se(x, which, log=TRUE, ...)
```

Arguments

x	a fitted model object.
which	which penalty parameter(s)?
log	if TRUE, the computed standard error is for log(theta) for negative binomial regression, otherwise, for theta.
...	arguments passed to methods.

Value

A vector containing standard errors of non-zero regularized estimators.

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[zipath](#)

Examples

```
data("bioChemists", package = "pscl")
fm_zinb <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambd=10)
res <- se(fm_zinb, which=which.min(fm_zinb$bic))
```

stan

standardize variables

Description

Standardize variables. For each column, return mean 0 and mean value of sum of squares = 1.

Usage

```
stan(x, weights)
```

Arguments

x	numeric variables, can be a matrix or vector
weights	numeric positive vector of weights

Value

A list with the following items.

x	standardized variables with each column: mean value 0 and mean value of sum of squares = 1.
meanx	a vector of means for each column in the original x
normx	a vector of scales for each column in the original x

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

summary.glmregNB

Summary Method Function for Objects of Class 'glmregNB'

Description

Summary results of fitted penalized negative binomial regression model

Usage

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

Arguments

object	fitted model object of class glmregNB.
...	arguments passed to or from other methods.

Details

This function is a method for the generic function `summary()` for class "glmregNB". It can be invoked by calling `summary(x)` for an object `x` of the appropriate class, or directly by calling `summary.glmregNB(x)` regardless of the class of the object.

Value

Summary of fitted penalized negative binomial model

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

See Also

[summary](#), [glm.nb](#)

Examples

```
## Not run:
data(quine, package="MASS")
summary(glmregNB(Days ~ Eth*Age*Lrn*Sex, quine, link = log))

## End(Not run)
```

tuning.zipath

find optimal penalized zero-inflated model

Description

Fit penalized zero-inflated models, generate multiple paths with varying penalty parameters, therefore determine optimal penalty parameters

Usage

```
tuning.zipath(formula, data, weights, subset, na.action, offset, standardize=TRUE,
family = c("poisson", "negbin", "geometric"), penalty = c("enet", "mnet", "snet"),
lambdaCountRatio = .0001, lambdaZeroRatio = c(.1, .01, .001),
maxit.theta=1, gamma.count=3, gamma.zero=3, ...)
```

Arguments

formula	symbolic description of the model, see details.
data	argument controlling formula processing via model.frame .
weights	optional numeric vector of weights. If standardize=TRUE, weights are renormalized to weights/sum(weights). If standardize=FALSE, weights are kept as original input
subset	subset of data
na.action	how to deal with missing data
offset	Not implemented yet
standardize	logical value, should variables be standardized?
family	family to fit
penalty	penalty considered as one of enet, mnet, snet.
lambdaCountRatio, lambdaZeroRatio	Smallest value for lambda.count and lambda.zero, respectively, as a fraction of lambda.max, the (data derived) entry value (i.e. the smallest value for which all coefficients are zero except the intercepts). This lambda.max can be a surrogate value for penalty="mnet" or "snet"

maxit.theta	For family="negbin", the maximum iteration allowed for estimating scale parameter theta. Note, the default value 1 is for computing speed purposes, and is typically too small and less desirable in real data analysis
gamma.count	The tuning parameter of the snet or mnet penalty for the count part of model.
gamma.zero	The tuning parameter of the snet or mnet penalty for the zero part of model.
...	Other arguments passing to zipath

Details

find optimal lambdaZeroRatio for penalized zero-inflated Poisson, negative binomial and geometric model

Value

An object of class zipath with the optimal lambdaZeroRatio

Author(s)

Zhu Wang <zwang@connecticutchildrens.org>

References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[zipath](#)

Examples

```
## Not run:
## data
data("bioChemists", package = "pscl")

## inflation with regressors
## ("art ~ . | ." is "art ~ fem + mar + kid5 + phd + ment | fem + mar + kid5 + phd + ment")
fm_zip2 <- tuning.zipath(art ~ . | ., data = bioChemists, nlambd=10)
summary(fm_zip2)
fm_zinb2 <- tuning.zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambd=10)
summary(fm_zinb2)

## End(Not run)
```

zipath	<i>Fit zero-inflated count data linear model with lasso (or elastic net), snet or mnet regularization</i>
--------	---

Description

Fit zero-inflated regression models for count data via penalized maximum likelihood.

Usage

```
zipath(formula, data, weights, subset, na.action, offset,
       standardize = TRUE, family = c("poisson", "negbin", "geometric"),
       link = c("logit", "probit", "cloglog", "cauchit", "log"),
       penalty = c("enet", "mnet", "snet"), start = NULL, model = TRUE,
       y = TRUE, x = FALSE, nlambda = 100, lambda.count = NULL, lambda.zero = NULL,
       penalty.factor.count=NULL, penalty.factor.zero=NULL,
       lambda.count.min.ratio = .0001, lambda.zero.min.ratio = .1,
       alpha.count = 1, alpha.zero = alpha.count, gamma.count = 3,
       gamma.zero = gamma.count, rescale=FALSE, init.theta, theta.fixed=FALSE,
       EM = TRUE, maxit.em=200, convtype=c("count", "both"), maxit = 1000,
       maxit.theta = 1, reltol = 1e-5, eps.bino=1e-5, shortlist=FALSE, trace = FALSE, ...)
```

Arguments

formula	symbolic description of the model, see details.
weights	optional numeric vector of weights.
data, subset, na.action	arguments controlling formula processing via model.frame .
offset	optional numeric vector with an a priori known component to be included in the linear predictor of the count model. See below for more information on offsets.
standardize	Logical flag for x variable standardization, prior to fitting the model sequence. The coefficients are always returned on the original scale. Default is standardize=TRUE.
family	character specification of count model family (a log link is always used).
link	character specification of link function in the binary zero-inflation model (a binomial family is always used).
model, y, x	logicals. If TRUE the corresponding components of the fit (model frame, response, model matrix) are returned.
penalty	penalty considered as one of enet, mnet, snet.
start	starting values for the parameters in the linear predictor.
nlambda	number of lambda value, default value is 100. The sequence may be truncated before nlambda is reached if a close to saturated model for the zero component is fitted.

<code>lambda.count</code>	A user supplied <code>lambda.count</code> sequence. Typical usage is to have the program compute its own <code>lambda.count</code> and <code>lambda.zero</code> sequence based on <code>nlambda</code> and <code>lambda.min.ratio</code> .
<code>lambda.zero</code>	A user supplied <code>lambda.zero</code> sequence.
<code>penalty.factor.count</code> , <code>penalty.factor.zero</code>	These are numeric vectors with the same length as predictor variables. that multiply <code>lambda.count</code> , <code>lambda.zero</code> , respectively, to allow differential shrinkage of coefficients. Can be 0 for some variables, which implies no shrinkage, and that variable is always included in the model. Default is same shrinkage for all variables.
<code>lambda.count.min.ratio</code> , <code>lambda.zero.min.ratio</code>	Smallest value for <code>lambda.count</code> and <code>lambda.zero</code> , respectively, as a fraction of <code>lambda.max</code> , the (data derived) entry value (i.e. the smallest value for which all coefficients are zero except the intercepts). Note, there is a closed formula for <code>lambda.max</code> for <code>penalty="enet"</code> . If <code>rescale=TRUE</code> , <code>lambda.max</code> is the same for <code>penalty="mnet"</code> or <code>"snet"</code> . Otherwise, some modifications are required. In the current implementation, for small gamma value, the square root of the computed <code>lambda.zero[1]</code> is used when <code>penalty="mnet"</code> or <code>"snet"</code> .
<code>alpha.count</code>	The elastic net mixing parameter for the count part of model.
<code>alpha.zero</code>	The elastic net mixing parameter for the zero part of model.
<code>gamma.count</code>	The tuning parameter of the <code>snet</code> or <code>mnet</code> penalty for the count part of model.
<code>gamma.zero</code>	The tuning parameter of the <code>snet</code> or <code>mnet</code> penalty for the zero part of model.
<code>rescale</code>	logical value, if TRUE, adaptive rescaling
<code>init.theta</code>	The initial value of theta for <code>family="negbin"</code> .
<code>theta.fixed</code>	Logical value only used for <code>family="negbin"</code> . If TRUE, theta is not updated.
EM	Using EM algorithm. Not implemented otherwise
<code>convtype</code>	convergency type, default is for count component only for speedy computation
<code>maxit.em</code>	Maximum number of EM algorithm
<code>maxit</code>	Maximum number of coordinate descent algorithm
<code>maxit.theta</code>	Maximum number of iterations for estimating theta scaling parameter if <code>family="negbin"</code> . Default value <code>maxit.theta</code> may be increased, yet may slow the algorithm
<code>eps.bino</code>	a lower bound of probabilities to be claimed as zero, for computing weights and related values when <code>family="binomial"</code> .
<code>reltol</code>	Convergence criteria, default value $1e-5$ may be reduced to make more accurate yet slow
<code>shortlist</code>	logical value, if TRUE, limited results return
<code>trace</code>	If TRUE, progress of algorithm is reported
...	Other arguments which can be passed to from <code>glmreg</code>

Details

The algorithm fits penalized zero-inflated count data regression models using the coordinate descent algorithm within the EM algorithm. The returned fitted model object is of class "zipath" and is similar to fitted "glm" and "zeroinfl" objects. For elements such as "coefficients" a list is returned with elements for the zero and count component, respectively. For details see below.

A set of standard extractor functions for fitted model objects is available for objects of class "zipath", including methods to the generic functions `print`, `coef`, `logLik`, `residuals`, `predict`. See `predict.zipath` for more details on all methods.

The program may terminate with the following message:

```
Error in: while (j <= maxit.em && !converged) { :
Missing value, where TRUE/FALSE is necessary
Calls: zipath
Additionally: Warning:
In glmreg_fit(Znew, probi, weights = weights, standardize = standardize, :
saturated model, exiting ...
Execution halted
```

One possible reason is that the fitted model is too complex for the data. There are two suggestions to overcome the error. One is to reduce the number of variables. Second, find out what lambda values caused the problem and omit them. Try with other lambda values instead.

Value

An object of class "zipath", i.e., a list with components including

<code>coefficients</code>	a list with elements "count" and "zero" containing the coefficients from the respective models,
<code>residuals</code>	a vector of raw residuals (observed - fitted),
<code>fitted.values</code>	a vector of fitted means,
<code>weights</code>	the case weights used,
<code>terms</code>	a list with elements "count", "zero" and "full" containing the terms objects for the respective models,
<code>theta</code>	estimate of the additional θ parameter of the negative binomial model (if a negative binomial regression is used),
<code>loglik</code>	log-likelihood of the fitted model,
<code>family</code>	character string describing the count distribution used,
<code>link</code>	character string describing the link of the zero-inflation model,
<code>linkinv</code>	the inverse link function corresponding to link,
<code>converged</code>	logical value, TRUE indicating successful convergence of zipath, FALSE indicating otherwise
<code>call</code>	the original function call
<code>formula</code>	the original formula
<code>levels</code>	levels of the categorical regressors

contrasts	a list with elements "count" and "zero" containing the contrasts corresponding to levels from the respective models,
model	the full model frame (if model = TRUE),
y	the response count vector (if y = TRUE),
x	a list with elements "count" and "zero" containing the model matrices from the respective models (if x = TRUE),

Author(s)

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References

Zhu Wang, Shuangge Ma, Michael Zappitelli, Chirag Parikh, Ching-Yun Wang and Prasad Devarajan (2014) *Penalized Count Data Regression with Application to Hospital Stay after Pediatric Cardiac Surgery*, *Statistical Methods in Medical Research*. 2014 Apr 17. [Epub ahead of print]

Zhu Wang, Shuangge Ma, Ching-Yun Wang, Michael Zappitelli, Prasad Devarajan and Chirag R. Parikh (2014) *EM for Regularized Zero Inflated Regression Models with Applications to Postoperative Morbidity after Cardiac Surgery in Children*, *Statistics in Medicine*. 33(29):5192-208.

Zhu Wang, Shuangge Ma and Ching-Yun Wang (2015) *Variable selection for zero-inflated and overdispersed data with application to health care demand in Germany*, *Biometrical Journal*. 57(5):867-84.

See Also

[glm](#), [glmreg](#), [glmregNB](#)

Examples

```
## Not run:
## data
data("bioChemists", package = "pscl")

## without inflation
## ("art ~ ." is "art ~ fem + mar + kid5 + phd + ment")
fm_pois <- glmreg(art ~ ., data = bioChemists, family = "poisson")
coef(fm_pois)
fm_nb <- glmregNB(art ~ ., data = bioChemists)
coef(fm_nb)
## with simple inflation (no regressors for zero component)
fm_zip <- zipath(art ~ . | 1, data = bioChemists, nlambda=10)
summary(fm_zip)
fm_zinb <- zipath(art ~ . | 1, data = bioChemists, family = "negbin", nlambda=10)
summary(fm_zinb)
## inflation with regressors
## ("art ~ . | ." is "art ~ fem + mar + kid5 + phd + ment | fem + mar + kid5 + phd + ment")
fm_zip2 <- zipath(art ~ . | ., data = bioChemists, nlambda=10)
summary(fm_zip2)
fm_zinb2 <- zipath(art ~ . | ., data = bioChemists, family = "negbin", nlambda=10)
summary(fm_zinb2)
```

```
### non-penalized regression, compare with zeroinfl
fm_zinb3 <- zipath(art ~ . | ., data = bioChemists, family = "negbin",
lambda.count=0, lambda.zero=0, reltol=1e-12)
summary(fm_zinb3)
fm_zinb4 <- zeroinfl(art ~ . | ., data = bioChemists, dist = "negbin")
summary(fm_zinb4)

## End(Not run)
```

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