## Package: SSGL (via r-universe)

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Type Package

**Title** Spike-and-Slab Group Lasso for Group-Regularized Generalized Linear Models

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**Description** Fits group-regularized generalized linear models (GLMs) using the spike-and-slab group lasso (SSGL) prior introduced by Bai et al. (2022) <doi:10.1080/01621459.2020.1765784> and extended to GLMs by Bai (2023) <arXiv:2007.07021>. This package supports fitting the SSGL model for the following GLMs with group sparsity: Gaussian linear regression, binary logistic regression, Poisson regression, negative binomial regression, and gamma regression. Stand-alone functions for group-regularized negative binomial regression and group-regularized gamma regression are also available, with the option of employing the group lasso penalty of Yuan and Lin (2006) <doi:10.1111/j.1467-9868.2005.00532.x>, the group minimax concave penalty (MCP) of Breheny and Huang <doi:10.1007/s11222-013-9424-2>, or the group smoothly clipped absolute deviation (SCAD) penalty of Breheny and Huang (2015) <doi:10.1007/s11222-013-9424-2>.

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#### **Description**

This function implements K-fold cross-validation for group-regularized gamma regression with a known shape parameter  $\nu$  and the log link. The cross-validation error (CVE) and cross-validation standard error (CVSE) are computed using the deviance for gamma regression.

For a description of group-regularized gamma regression, see the description for the gamma\_grpreg function. Our implementation is based on the least squares approximation approach of Wang and Leng (2007), and hence, the function does not allow the total number of covariates p to be greater than  $\frac{K-1}{K} \times$  sample size, where K is the number of folds.

Note that the gamma\_grpreg function also returns the generalized information criterion (GIC) of Fan and Tang (2013) for each regularization parameter in lambda, and the GIC can also be used for model selection instead of cross-validation.

#### Usage

#### **Arguments**

Υ	$n \times 1$ vector of strictly positive, continuous responses for training data.
X	$n \times p$ design matrix for training data, where the $j$ th column corresponds to the $j$ th overall feature.
groups	p-dimensional vector of group labels. The $j$ th entry in groups should contain either the group number $or$ the factor level name that the feature in the $j$ th column of X belongs to. groups must be either a vector of integers or factors.
gamma_shape	known shape parameter $\nu$ in $Gamma(\mu_i, \nu)$ distribution for the responses. Default is gamma_shape=1.
penalty	group regularization method to use on the groups of regression coefficients. The options are "gLASSO", "gSCAD", "gMCP". To implement cross-validation for gamma regression with the SSGL penalty, use the cv_SSGL function.

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n\_folds number of folds K to use in K-fold cross-validation. Default is n\_folds=10. group\_weights group-specific, nonnegative weights for the penalty. Default is to use the square

roots of the group sizes.

taper  $\gamma$  in group SCAD and group MCP controlling how rapidly the

penalty tapers off. Default is taper=4 for group SCAD and taper=3 for group

MCP. Ignored if "gLASSO" is specified as the penalty.

n\_lambda number of regularization parameters L. Default is n\_lambda=100.

lambda grid of L regularization parameters. The user may specify either a scalar or a

vector. If the user does not provide this, the program chooses the grid automati-

cally.

max\_iter maximum number of iterations in the algorithm. Default is max\_iter=10000.

tol convergence threshold for algorithm. Default is tol=1e-4.

#### Value

The function returns a list containing the following components:

lambda  $L \times 1$  vector of regularization parameters lambda used to fit the model. lambda

is displayed in descending order.

cve  $L \times 1$  vector of mean cross-validation error across all K folds. The kth entry in

cve corresponds to the kth regularization parameter in lambda.

cvse  $L \times 1$  vector of standard errors for cross-validation error across all K folds. The

kth entry in cvse corresponds to the kth regularization parameter in lambda.

lambda\_min The value in lambda that minimizes mean cross-validation error cve.

min\_index The index of lambda\_min in lambda.

#### References

Breheny, P. and Huang, J. (2015). "Group descent algorithms for nonconvex penalized linear and logistic regression models with grouped predictors." *Statistics and Computing*, **25**:173-187.

Fan, Y. and Tang, C. Y. (2013). "Tuning parameter selection in high-dimensional penalized likelihood." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **75**:531-552.

Wang, H. and Leng, C. (2007). "Unified LASSO estimation by least squares approximation." *Journal of the American Statistical Association*, **102**:1039-1048.

Yuan, M. and Lin, Y. (2006). "Model selection and estimation in regression with grouped variables." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **68**:49-67.

```
## Generate data
set.seed(12345)
X = matrix(runif(100*11), nrow=100)
n = dim(X)[1]
groups = c(1,1,1,2,2,2,3,3,4,5,5)
beta_true = c(-1,1,1,0,0,0,0,0,0,0,1.5,-1.5)
```

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```
## Generate responses from gamma regression with known shape parameter 1
eta = crossprod(t(X), beta_true)
shape = 1
Y = rgamma(n, rate=shape/exp(eta), shape=shape)
## 10-fold cross-validation for group-regularized gamma regression
## with the group LASSO penalty
gamma_cv = cv_gamma_grpreg(Y, X, groups, penalty="gLASSO")
## Plot cross-validation curve
plot(gamma_cv$lambda, gamma_cv$cve, type="1", xlab="lambda", ylab="CVE")
## lambda which minimizes mean CVE
gamma_cv$lambda_min
## index of lambda_min in lambda
gamma_cv$min_index
```

cv\_nb\_grpreg

Cross-validation for Group-Regularized Negative Binomial Regression

#### **Description**

This function implements K-fold cross-validation for group-regularized negative binomial regression with a known size parameter  $\alpha$  and the log link. The cross-validation error (CVE) and crossvalidation standard error (CVSE) are computed using the deviance for negative binomial regression.

For a description of group-regularized negative binomial regression, see the description for the nb\_grpreg function. Our implementation is based on the least squares approximation approach of Wang and Leng (2007), and hence, the function does not allow the total number of covariates p to be greater than  $\frac{K-1}{K} \times$  sample size, where K is the number of folds.

Note that the nb\_grpreg function also returns the generalized information criterion (GIC) of Fan and Tang (2013) for each regularization parameter in lambda, and the GIC can also be used for model selection instead of cross-validation.

#### Usage

```
cv_nb_grpreg(Y, X, groups, nb_size=1, penalty=c("gLASSO","gSCAD","gMCP"),
            n_folds=10, group_weights, taper, n_lambda=100, lambda,
            max_iter=10000, tol=1e-4)
```

#### **Arguments**

Υ  $n \times 1$  vector of strictly nonnegative integer responses for training data.

Χ  $n \times p$  design matrix for training data, where the jth column corresponds to the

jth overall feature.

p-dimensional vector of group labels. The jth entry in groups should contain eigroups

ther the group number or the factor level name that the feature in the jth column

of X belongs to. groups must be either a vector of integers or factors.

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nb_size	known size parameter $\alpha$ in $NB(\alpha,\mu_i)$ distribution for the responses. Default is nb_size=1.
penalty	group regularization method to use on the groups of regression coefficients. The options are "gLASSO", "gSCAD", "gMCP". To implement cross-validation for gamma regression with the SSGL penalty, use the cv_SSGL function.
n_folds	number of folds $K$ to use in $K$ -fold cross-validation. Default is n_folds=10.
group_weights	group-specific, nonnegative weights for the penalty. Default is to use the square roots of the group sizes.
taper	tapering term $\gamma$ in group SCAD and group MCP controlling how rapidly the penalty tapers off. Default is taper=4 for group SCAD and taper=3 for group MCP. Ignored if "gLASSO" is specified as the penalty.
n_lambda	number of regularization parameters $L$ . Default is n_lambda=100.
lambda	grid of $L$ regularization parameters. The user may specify either a scalar or a vector. If the user does not provide this, the program chooses the grid automatically.
max_iter	maximum number of iterations in the algorithm. Default is max_iter=10000.
tol	convergence threshold for algorithm. Default is tol=1e-4.

#### Value

The function returns a list containing the following components:

lambda	$L\times 1$ vector of regularization parameters 1ambda used to fit the model. 1ambda is displayed in descending order.
cve	$L \times 1$ vector of mean cross-validation error across all $K$ folds. The $k$ th entry in cve corresponds to the $k$ th regularization parameter in lambda.
cvse	$L \times 1$ vector of standard errors for cross-validation error across all $K$ folds. The $k$ th entry in cvse corresponds to the $k$ th regularization parameter in lambda.
lambda_min	The value in lambda that minimizes mean cross-validation error cve.
min_index	The index of lambda_min in lambda.

#### References

Breheny, P. and Huang, J. (2015). "Group descent algorithms for nonconvex penalized linear and logistic regression models with grouped predictors." Statistics and Computing, 25:173-187.

Fan, Y. and Tang, C. Y. (2013). "Tuning parameter selection in high dimensional penalized likelihood." Journal of the Royal Statistical Society: Series B (Statistical Methodology), 75:531-552.

Wang, H. and Leng, C. (2007). "Unified LASSO estimation by least squares approximation." Journal of the American Statistical Association, 102:1039-1048.

Yuan, M. and Lin, Y. (2006). "Model selection and estimation in regression with grouped variables." Journal of the Royal Statistical Society: Series B (Statistical Methodology), 68:49-67.

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#### **Examples**

```
## Generate data
set.seed(1234)
X = matrix(runif(100*14), nrow=100)
n = dim(X)[1]
groups = c(1,1,1,2,2,2,2,3,3,4,5,5,6,6)
beta_true = c(-1,1,1,0,0,0,0,-1,1,0,0,0,-1.5,1.5)
## Generate count responses from negative binomial regression
eta = crossprod(t(X), beta_true)
Y = rnbinom(n, size=1, mu=exp(eta))
## 10-fold cross-validation for group-regularized negative binomial
## regression with the group MCP penalty
nb_cv = cv_nb_grpreg(Y, X, groups, penalty="gMCP")
## Plot cross-validation curve
plot(nb_cv$lambda, nb_cv$cve, type="l", xlab="lambda", ylab="CVE")
## lambda which minimizes mean CVE
nb_cv$lambda_min
## index of lambda_min in lambda
nb cv$min index
```

cv\_SSGL

Cross-Validation for Spike-and-Slab Group Lasso in Group-Regularized Generalized Linear Models (GLMs)

#### Description

This function implements K-fold cross-validation for group-regularized GLMs with the spike-and-slab group lasso (SSGL) penalty of Bai et al. (2022) and Bai (2023). The identity link function is used for Gaussian regression, the logit link is used for binomial regression, and the log link is used for Poisson, negative binomial, and gamma regression.

Although one can choose lambda0 from cross-validation with this function, it can be very time-consuming to do so if the number of groups G and/or the number of total covariantes p is moderate to large. It is strongly recommended that the user simply run the SSGL function on the training dataset and select the final model according to the lambda0 that minimizes the generalized information criterion (GIC). See description of the SSGL function for more details.

#### **Usage**

```
cv_SSGL(Y, X, groups,
    family=c("gaussian","binomial","poisson","negativebinomial","gamma"),
    nb_size=1, gamma_shape=1, group_weights, n_folds=5, n_lambda0=25,
    lambda0, lambda1=1, a=1, b=dim(X)[2],
    max_iter=100, tol=1e-6, print_fold=TRUE)
```

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## Arguments

Υ	$n \times 1$ vector of responses for training data.
X	$n\times p$ design matrix for training data, where the $j{\rm th}$ column corresponds to the $j{\rm th}$ overall feature.
groups	p-dimensional vector of group labels. The $j$ th entry in groups should contain either the group number $or$ the factor level name that the feature in the $j$ th column of X belongs to. groups must be either a vector of integers or factors.
family	exponential dispersion family of the response variables. Allows for "gaussian", "binomial", "poisson", "negativebinomial", and "gamma". Note that for "negativebinomial", the size parameter must be specified in advance, while for "gamma", the shape parameter must be specified in advance.
nb_size	known size parameter $\alpha$ in $NB(\alpha,\mu_i)$ distribution for the responses if the user specifies family="negativebinomial". Default is nb_size=1. Ignored if family is not "negativebinomial".
gamma_shape	known shape parameter $\nu$ in $G(\mu_i,\nu)$ distribution for the responses if the user specifies family="gamma". Default is gamma_shape=1. Ignored if family is not "gamma".
group_weights	group-specific, nonnegative weights for the penalty. Default is to use the square roots of the group sizes.
n_folds	number of folds $K$ to use in $K$ -fold cross-validation. Default is n_folds=5.
n_lambda0	number of spike hyperparameters $L$ . Default is n_lambda0=25.
lambda0	grid of $L$ spike hyperparameters $\lambda_0$ . The user may specify either a scalar or a vector. If the user does not provide this, the program chooses the grid automatically.
lambda1	slab hyperparameter $\lambda_1$ in the SSGL prior. Default is lambda1=1.
a	shape hyperparameter for the $Beta(a,b)$ prior on the mixing proportion in the SSGL prior. Default is a=1.
b	shape hyperparameter for the $Beta(a,b)$ prior on the mixing proportion in the SSGL prior. Default is b=dim(X)[2].
max_iter	maximum number of iterations in the algorithm. Default is max_iter=100.
tol	convergence threshold for algorithm. Default is tol=1e-6.
print_fold	Boolean variable for whether or not to print the current fold in the algorithm. Default is print_fold=TRUE.

## Value

The function returns a list containing the following components:

lambda0	$L\times 1$ vector of spike hyperparameters lambda0 used to fit the model. lambda0 is displayed in descending order.
cve	$L \times 1$ vector of mean cross-validation error across all $K$ folds. The $k$ th entry in cve corresponds to the $k$ th spike hyperparameter parameter in lambda0.

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cvse  $L \times 1$  vector of standard errors for cross-validation error across all K folds.

The kth entry in cvse corresponds to the kth spike hyperparameter parameter in

lambda0.

lambda@\_min The value in lambda@ that minimizes mean cross-validation error cve.

min\_index The index of lambda0\_min in lambda0.

#### References

Bai, R. (2023). "Bayesian group regularization in generalized linear models with a continuous spike-and-slab prior." *arXiv pre-print arXiv:2007.07021*.

Bai, R., Moran, G. E., Antonelli, J. L., Chen, Y., and Boland, M.R. (2022). "Spike-and-slab group lassos for grouped regression and sparse generalized additive models." *Journal of the American Statistical Association*, **117**:184-197.

```
## Generate data
set.seed(12345)
X = matrix(runif(50*6), nrow=50)
n = dim(X)[1]
groups = c(1,1,1,2,2,2)
beta_true = c(-2,1,1.5,0,0,0)
## Generate responses from Gaussian distribution
Y = crossprod(t(X), beta_true) + rnorm(n)
## K-fold cross-validation
## NOTE: If you do not specify lambda0, the function will automatically choose a suitable grid.
ssgl_mods = cv_SSGL(Y, X, groups, family="gaussian", lambda0=seq(from=16,to=4,by=-4))
## Plot cross-validation curve
plot(ssgl_mods$lambda0, ssgl_mods$cve, type="1", xlab="lambda0", ylab="CVE")
## lambda which minimizes mean CVE
ssgl_mods$lambda0_min
ssgl_mods$min_index
## Example with Poisson regression
## Generate count responses
eta = crossprod(t(X), beta_true)
Y = rpois(n,exp(eta))
## K-fold cross-validation
## NOTE: If you do not specify lambda0, the program will automatically choose a suitable grid.
ssgl_poisson_mods = cv_SSGL(Y, X, groups, family="poisson", lambda0=seq(from=20,to=2,by=-4))
## Plot cross-validation curve
plot(ssgl_poisson_mods$lambda0, ssgl_poisson_mods$cve, type="l", xlab="lambda0", ylab="CVE")
## lambda which minimizes mean CVE
ssgl_poisson_mods$lambda0_min
```

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ssgl\_poisson\_mods\$min\_index

gamma\_grpreg

Group-regularized Gamma Regression

## Description

This function implements group-regularized gamma regression with a known shape parameter  $\nu$ and the log link. In gamma regression, we assume that  $y_i \sim Gamma(\mu_i, \nu)$ , where

$$f(y_i|\mu_i,\nu) = \frac{1}{\Gamma(\nu)} (\frac{\nu}{\mu_i})^{\nu} \exp(-\frac{\nu}{\mu_i} y_i) y_i^{\nu-1}, y > 0.$$

Then  $E(y_i) = \mu_i$ , and we relate  $\mu_i$  to a set of p covariates  $x_i$  through the log link,

$$\log(\mu_i) = \beta_0 + x_i^T \beta, i = 1, ..., n$$

If the covariates in each  $x_i$  are grouped according to known groups g = 1, ..., G, then this function can estimate some of the G groups of coefficients as all zero, depending on the amount of regularization. Our implementation for regularized gamma regression is based on the least squares approximation approach of Wang and Leng (2007), and hence, the function does not allow the total number of covariates p to be greater than sample size.

In addition, this function has the option of returning the generalized information criterion (GIC) of Fan and Tang (2013) for each regularization parameter in the grid lambda. The GIC can be used for model selection and serves as a useful alternative to cross-validation.

#### Usage

```
gamma_grpreg(Y, X, groups, X_test, gamma_shape=1,
             penalty=c("gLASSO", "gSCAD", "gMCP"),
             group_weights, taper, n_lambda=100, lambda,
             max_iter=10000, tol=1e-4, return_GIC=TRUE)
```

#### **Arguments**

Y $n \times 1$	l vector of strictly positive	, continuous responses	for training data.
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Χ  $n \times p$  design matrix for training data, where the jth column corresponds to the

*j*th overall feature.

groups p-dimensional vector of group labels. The *j*th entry in groups should contain ei-

ther the group number or the factor level name that the feature in the jth column

of X belongs to. groups must be either a vector of integers or factors.

X\_test  $n_{test} \times p$  design matrix for test data to calculate predictions. X\_test must have

the same number of columns as X, but not necessarily the same number of rows. If no test data is provided or if in-sample predictions are desired, then the function automatically sets X\_test=X in order to calculate *in-sample* predictions.

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gamma\_shape known shape parameter  $\nu$  in  $Gamma(\mu_i, \nu)$  distribution for the responses. De-

fault is gamma\_shape=1.

penalty group regularization method to use on the groups of regression coefficients. The

options are "gLASSO", "gSCAD", "gMCP". To implement gamma regression with

the SSGL penalty, use the SSGL function.

group\_weights group-specific, nonnegative weights for the penalty. Default is to use the square

roots of the group sizes.

taper tapering term  $\gamma$  in group SCAD and group MCP controlling how rapidly the

penalty tapers off. Default is taper=4 for group SCAD and taper=3 for group

MCP. Ignored if "gLASSO" is specified as the penalty.

n\_lambda number of regularization parameters L. Default is n\_lambda=100.

lambda grid of L regularization parameters. The user may specify either a scalar or a

vector. If the user does not provide this, the program chooses the grid automati-

cally.

max\_iter maximum number of iterations in the algorithm. Default is max\_iter=10000.

tol convergence threshold for algorithm. Default is tol=1e-4.

return\_GIC Boolean variable for whether or not to return the GIC. Default is return\_GIC=TRUE.

#### Value

The function returns a list containing the following components:

lambda  $L \times 1$  vector of regularization parameters lambda used to fit the model. lambda

is displayed in descending order.

beta  $p \times L$  matrix of estimated regression coefficients. The kth column in beta

corresponds to the kth regularization parameter in lambda.

beta0  $L \times 1$  vector of estimated intercepts. The kth entry in beta0 corresponds to the

kth regularization parameter in lambda.

classifications

 $G \times L$  matrix of classifications, where G is the number of groups. An entry of "1" indicates that the group was classified as nonzero, and an entry of "0" indicates that the group was classified as zero. The kth column of classifications

corresponds to the kth regularization parameter in lambda.

Y\_pred  $n_{test} \times L$  matrix of predicted mean response values  $\mu_{test} = E(Y_{test})$  based

on the *test* data in  $X_{test}$  (or training data X if no argument was specified for  $X_{test}$ ). The kth column in  $Y_{test}$  corresponds to the predictions for the kth

regularization parameter in lambda.

GIC  $L \times 1$  vector of GIC values. The kth entry of GIC corresponds to the kth entry

in our lambda grid. This is not returned if return\_GIC=FALSE.

lambda\_min The value in lambda that minimizes GIC. This is not returned if return\_GIC=FALSE.

min\_index The index of lambda\_min in lambda. This is not returned if return\_GIC=FALSE.

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#### References

Breheny, P. and Huang, J. (2015). "Group descent algorithms for nonconvex penalized linear and logistic regression models with grouped predictors." *Statistics and Computing*, **25**:173-187.

Fan, Y. and Tang, C. Y. (2013). "Tuning parameter selection in high dimensional penalized likelihood." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **75**:531-552.

Wang, H. and Leng, C. (2007). "Unified LASSO estimation by least squares approximation." *Journal of the American Statistical Association*, **102**:1039-1048.

Yuan, M. and Lin, Y. (2006). "Model selection and estimation in regression with grouped variables." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **68**:49-67.

```
## Generate data
set.seed(1234)
X = matrix(runif(100*11), nrow=100)
n = dim(X)[1]
groups = c(1,1,1,2,2,2,3,3,4,5,5)
beta_true = c(-1,1,1,0,0,0,0,0,0,1.5,-1.5)
## Generate responses from gamma regression with known shape parameter 1
eta = crossprod(t(X), beta_true)
shape = 1
Y = rgamma(n, rate=shape/exp(eta), shape=shape)
## Generate test data
n_{\text{test}} = 50
X_test = matrix(runif(n_test*11), nrow=n_test)
## Fit gamma regression models with the group SCAD penalty
gamma_mod = gamma_grpreg(Y, X, groups, X_test, penalty="gSCAD")
## Tuning parameters used to fit models
gamma_mod$lambda
## Predicted n_test-dimensional vectors mu=E(Y_test) based on test data, X_test.
## The kth column of 'Y_pred' corresponds to the kth entry in 'lambda.'
gamma_mod$Y_pred
## Classifications of the 5 groups. The kth column of 'classifications'
# corresponds to the kth entry in 'lambda.'
gamma_mod$classifications
## Plot lambda vs. GIC
plot(gamma_mod$lambda, gamma_mod$GIC, type='l')
## Model selection with the lambda that minimizes GIC
gamma_mod$lambda_min
gamma_mod$min_index
gamma_mod$classifications[, gamma_mod$min_index]
gamma_mod$beta[, gamma_mod$min_index]
```

nb\_grpreg

nb\_grpreg

Group-regularized Negative Binomial Regression

#### **Description**

This function implements group-regularized negative binomial regression with a known size parameter  $\alpha$  and the log link. In negative binomial regression, we assume that  $y_i \sim NB(\alpha, \mu_i)$ , where

$$f(y_i|\alpha,\mu_i) = \frac{\Gamma(y_i+\alpha)}{y_i |\Gamma(\alpha)} \left(\frac{\mu_i}{\mu_i+\alpha}\right)^{y_i} \left(\frac{\alpha}{\mu_i+\alpha}\right)^{\alpha}, y_i = 0, 1, 2, \dots$$

Then  $E(y_i) = \mu_i$ , and we relate  $\mu_i$  to a set of p covariates  $x_i$  through the log link,

$$\log(\mu_i) = \beta_0 + x_i^T \beta, i = 1, ..., n$$

If the covariates in each  $x_i$  are grouped according to known groups g=1,...,G, then this function can estimate some of the G groups of coefficients as all zero, depending on the amount of regularization. Our implementation for regularized negative binomial regression is based on the least squares approximation approach of Wang and Leng (2007), and hence, the function does not allow the total number of covariates p to be greater than sample size.

In addition, this function has the option of returning the generalized information criterion (GIC) of Fan and Tang (2013) for each regularization parameter in the grid lambda. The GIC can be used for model selection and serves as a useful alternative to cross-validation.

#### Usage

#### **Arguments**

Υ	$n \times 1$ vector of strictly nonnegative integer responses for training data.
X	$n\times p$ design matrix for training data, where the $j{\rm th}$ column corresponds to the $j{\rm th}$ overall feature.
groups	p-dimensional vector of group labels. The $j$ th entry in groups should contain either the group number $or$ the factor level name that the feature in the $j$ th column of X belongs to. groups must be either a vector of integers or factors.
X_test	$n_{test}  imes p$ design matrix for test data to calculate predictions. X_test must have the $\mathit{same}$ number of columns as X, but not necessarily the same number of rows. If $\mathit{no}$ test data is provided or if in-sample predictions are desired, then the function automatically sets X_test=X in order to calculate $\mathit{in-sample}$ predictions.
nb_size	known size parameter $\alpha$ in $NB(\alpha, \mu_i)$ distribution for the responses. Default is

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penalty group regularization method to use on the groups of regression coefficients. The

options are "gLASSO", "gSCAD", "gMCP". To implement gamma regression with

the SSGL penalty, use the SSGL function.

group\_weights group-specific, nonnegative weights for the penalty. Default is to use the square

roots of the group sizes.

taper  $\gamma$  tapering term  $\gamma$  in group SCAD and group MCP controlling how rapidly the

penalty tapers off. Default is taper=4 for group SCAD and taper=3 for group

MCP. Ignored if "gLASSO" is specified as the penalty.

n\_lambda number of regularization parameters L. Default is n\_lambda=100.

lambda grid of L regularization parameters. The user may specify either a scalar or a

vector. If the user does not provide this, the program chooses the grid automati-

cally.

max\_iter maximum number of iterations in the algorithm. Default is max\_iter=10000.

tol convergence threshold for algorithm. Default is tol=1e-4.

return\_GIC Boolean variable for whether or not to return the GIC. Default is return\_GIC=TRUE.

#### Value

The function returns a list containing the following components:

lambda  $L \times 1$  vector of regularization parameters lambda used to fit the model. lambda

is displayed in descending order.

beta  $p \times L$  matrix of estimated regression coefficients. The kth column in beta

corresponds to the kth regularization parameter in lambda.

beta0  $L \times 1$  vector of estimated intercepts. The kth entry in beta0 corresponds to the

kth regularization parameter in lambda.

classifications

 $G \times L$  matrix of classifications, where G is the number of groups. An entry of "1" indicates that the group was classified as nonzero, and an entry of "0" indicates that the group was classified as zero. The kth column of classifications

corresponds to the kth regularization parameter in lambda.

Y\_pred  $n_{test} \times L$  matrix of predicted mean response values  $\mu_{test} = E(Y_{test})$  based

on the test data in X\_test (or training data X if no argument was specified for X\_test). The kth column in Y\_pred corresponds to the predictions for the kth

regularization parameter in lambda.

GIC  $L \times 1$  vector of GIC values. The kth entry of GIC corresponds to the kth entry

in our lambda grid. This is not returned if return\_GIC=FALSE.

lambda\_min The value in lambda that minimizes GIC. This is not returned if return\_GIC=FALSE.

min\_index The index of lambda\_min in lambda. This is not returned if return\_GIC=FALSE.

#### References

Breheny, P. and Huang, J. (2015). "Group descent algorithms for nonconvex penalized linear and logistic regression models with grouped predictors." *Statistics and Computing*, **25**:173-187.

nb\_grpreg

Fan, Y. and Tang, C. Y. (2013). "Tuning parameter selection in high dimensional penalized likelihood." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **75**:531-552.

Wang, H. and Leng, C. (2007). "Unified LASSO estimation by least squares approximation." *Journal of the American Statistical Association*, **102**:1039-1048.

Yuan, M. and Lin, Y. (2006). "Model selection and estimation in regression with grouped variables." *Journal of the Royal Statistical Society: Series B (Statistical Methodology)*, **68**:49-67.

```
## Generate data
set.seed(1234)
X = matrix(runif(100*15), nrow=100)
n = dim(X)[1]
groups = as.factor(groups)
beta_true = c(-1.5, 1.5, -1.5, 1.5, 0, 0, 0, 0, 0, 0, 2, -2, 0, 0, 0)
## Generate count responses from negative binomial regression
eta = crossprod(t(X), beta_true)
Y = rnbinom(n, size=1, mu=exp(eta))
## Generate test data
n_{test} = 50
X_test = matrix(runif(n_test*15), nrow=n_test)
## Fit negative binomial regression models with the group MCP penalty
nb_mod = nb_grpreg(Y, X, groups, X_test, penalty="gMCP")
## Tuning parameters used to fit models
nb_mod$lambda
# Predicted n_test-dimensional vectors mu=E(Y_test) based on test data, X_test.
# The kth column of 'Y_pred' corresponds to the kth entry in 'lambda.'
# Classifications of the 8 groups. The kth column of 'classifications'
# corresponds to the kth entry in lambda.
nb mod$classifications
## Plot lambda vs. GIC
plot(nb_mod$lambda, nb_mod$GIC, type='l')
## Model selection with the lambda that minimizes GIC
nb_mod$lambda_min
nb_mod$min_index
nb_mod$classifications[, nb_mod$min_index]
nb_mod$beta[, nb_mod$min_index]
```

SSGL Spike-and-Slab Group Lasso for Group-Regularized Generalized Linear Models (GLMs)

#### **Description**

This is a function to implement group-regularized GLMs with the spike-and-slab group lasso (SSGL) penalty of Bai et al. (2022) and Bai (2023). The identity link function is used for Gaussian regression, the logit link is used for binomial regression, and the log link is used for Poisson, negative binomial, and gamma regression. If the covariates in each  $x_i$  are grouped according to known groups g=1,...,G, then this function can estimate some of the G groups of coefficients as all zero, depending on the amount of regularization.

In addition, this function has the option of returning the generalized information criterion (GIC) of Fan and Tang (2013) for each regularization parameter in the grid lambda0. The GIC can be used for model selection and serves as a useful alternative to cross-validation. The formula for the GIC and a given  $\lambda_0$  is

$$DIC(\lambda_0) = \frac{1}{n} Deviance_{\lambda_0} + a_n \times \nu),$$

where  $Deviance_{\lambda_0}$  is the deviance computed with the estimate of beta based on spike hyperparameter  $\lambda_0$ ,  $\nu_0$  is the number of nonzero elements in the estimated beta, and  $a_n$  is a sequence that diverges at a suitable rate relative to n. As recommended by Fan and Tang (2013), we set  $a_n = \{\log(\log(n))\} \log(p)$ .

#### Usage

```
SSGL(Y, X, groups,
    family=c("gaussian","binomial","poisson","negativebinomial","gamma"),
    X_test, nb_size=1, gamma_shape=1, group_weights, n_lambda0=25,
    lambda0, lambda1=1, a=1, b=dim(X)[2],
    max_iter=100, tol = 1e-6, return_GIC=TRUE, print_lambda0=TRUE)
```

#### **Arguments**

Υ	$n \times 1$ vector of responses for training data.
X	$n\times p$ design matrix for training data, where the $j{\rm th}$ column of X corresponds to the $j{\rm th}$ overall covariate.
groups	p-dimensional vector of group labels. The $j$ th entry in groups should contain either the group number $or$ the factor level name that the feature in the $j$ th column of X belongs to. groups must be either a vector of integers or factors.
family	exponential dispersion family of the response variables. Allows for "gaussian", "binomial", "poisson", "negativebinomial", and "gamma". Note that for "negativebinomial", the size parameter must be specified in advance, while for "gamma", the shape parameter must be specified in advance.

X_test	$n_{test}  imes p$ design matrix for test data to calculate predictions. X_test must have the <i>same</i> number of columns as X, but not necessarily the same number of rows. If <i>no</i> test data is provided or if in-sample predictions are desired, then the function automatically sets X_test=X in order to calculate <i>in-sample</i> predictions.
nb_size	known size parameter $\alpha$ in $NB(\alpha,\mu_i)$ distribution for the responses if the user specifies family="negativebinomial". Default is nb_size=1. Ignored if family is not "gamma".
gamma_shape	known shape parameter $\nu$ in $Gamma(\mu_i, \nu)$ distribution for the responses if the user specifies family="gamma". Default is gamma_shape=1.
group_weights	group-specific, nonnegative weights for the penalty. Default is to use the square roots of the group sizes.
n_lambda0	number of spike hyperparameters $L$ . Default is n_lambda0=25.
lambda0	grid of $L$ spike hyperparameters $\lambda_0$ . The user may specify either a scalar or a vector. If the user does not provide this, the program chooses the grid automatically.
lambda1	slab hyperparameter $\lambda_1$ in the SSGL prior. Default is lambda1=1.
а	shape hyperparameter for the $Beta(a,b)$ prior on the mixing proportion in the SSGL prior. Default is a=1.
b	shape hyperparameter for the $Beta(a,b)$ prior on the mixing proportion in the SSGL prior. Default is b=dim(X)[2].
max_iter	maximum number of iterations in the algorithm. Default is max_iter=100.
tol	convergence threshold for algorithm. Default is tol=1e-6.
return_GIC	$Boolean\ variable\ for\ whether\ or\ not\ to\ return\ the\ GIC.\ Default\ is\ {\tt return\_GIC=TRUE}.$
print_lambda0	Boolean variable for whether or not to print the current value in lambda0. Default is print_lambda0=TRUE.

#### Value

The function returns a list containing the following components:

lambda0  $L\times 1$  vector of spike hyperpameters lambda0 used to fit the model. lambda0 is

displayed in descending order.

beta  $p \times L$  matrix of estimated regression coefficients. The kth column in beta

corresponds to the kth spike hyperparameter in lambda0.

beta0  $L \times 1$  vector of estimated intercepts. The kth entry in beta0 corresponds to the

kth spike hyperparameter in lambda0.

classifications

 $G \times L$  matrix of classifications, where G is the number of groups. An entry of "1" indicates that the group was classified as nonzero, and an entry of "0" indicates that the group was classified as zero. The kth column of classifications

corresponds to the  $k{\rm th}$  spike hyperparameter in lambda0.

Y\_pred  $n_{test} \times L$  matrix of predicted mean response values  $\mu_{test} = E(Y_{test})$  based

on the test data in X\_test (or training data X if no argument was specified for X\_test). The kth column in Y\_pred corresponds to the predictions for the kth

spike hyperparameter in lambda0.

 $L \times 1$  vector of GIC values. The kth entry of GIC corresponds to the kth entry GIC

in our lambda0 grid. This is not returned if return\_GIC=FALSE.

The value in lambda0 that minimizes GIC. This is not returned if return\_GIC=FALSE. lambda0\_min The index of lambda0\_min in lambda0. This is not returned if return\_GIC=FALSE. min\_index

#### References

Bai, R. (2023). "Bayesian group regularization in generalized linear models with a continuous spike-and-slab prior." arXiv pre-print arXiv:2007.07021.

Bai, R., Moran, G. E., Antonelli, J. L., Chen, Y., and Boland, M.R. (2022). "Spike-and-slab group lassos for grouped regression and sparse generalized additive models." Journal of the American Statistical Association, 117:184-197.

Fan, Y. and Tang, C. Y. (2013). "Tuning parameter selection in high dimensional penalized likelihood." Journal of the Royal Statistical Society: Series B (Statistical Methodology), 75:531-552.

```
## Generate data
set.seed(12345)
X = matrix(runif(100*10), nrow=100)
n = dim(X)[1]
groups = c("A", "A", "A", "B", "B", "B", "C", "C", "D", "D")
groups = as.factor(groups)
beta_true = c(-2.5, 1.5, 1.5, 0, 0, 0, 2, -2, 0, 0)
## Generate responses from Gaussian distribution
Y = crossprod(t(X), beta_true) + rnorm(n)
## Generate test data
n_{test} = 50
X_test = matrix(runif(n_test*10), nrow=n_test)
## Fit SSGL model with 10 spike hyperparameters
## NOTE: If you do not specify lambda0, the program will automatically choose a suitable grid.
SSGL_mod = SSGL(Y, X, groups, family="gaussian", X_test, lambda0=seq(from=50,to=5,by=-5))
## Regression coefficient estimates
SSGL_mod$beta
## Predicted n_test-dimensional vectors mu=E(Y.test) based on test data, X_test.
## The kth column of 'Y_pred' corresponds to the kth entry in 'lambda.'
SSGL_mod$Y_pred
## Classifications of the 8 groups. The kth column of 'classifications'
## corresponds to the kth entry in 'lambda.'
SSGL_mod$classifications
## Plot lambda vs. GIC
plot(SSGL_mod$lambda0, SSGL_mod$GIC, type='1')
## Model selection with the lambda that minimizes GIC
```

```
SSGL_mod$lambda0_min
SSGL_mod$min_index
SSGL_mod$classifications[, SSGL_mod$min_index]
SSGL_mod$beta[, SSGL_mod$min_index]
## Example with binary logistic regression
set.seed(12345)
X = matrix(runif(100*8), nrow=100)
n = \dim(X)[1]
groups = c("A", "A", "A", "B", "B", "B", "C", "C")
groups = as.factor(groups)
beta_true = c(-2.5, 1.5, 1.5, 0, 0, 0, 2, -2)
## Generate binary responses
eta = crossprod(t(X), beta_true)
Y = rbinom(n, size=1, prob=1/(1+exp(-eta)))
## Generate test data
n_{\text{test}} = 50
X_test = matrix(runif(n_test*8), nrow=n_test)
## Fit SSGL logistic regression model with 10 spike hyperparameters
## NOTE: If you do not specify lambda0, the program will automatically choose a suitable grid.
SSGL_logistic_mod = SSGL(Y, X, groups, family="binomial", X_test, lambda0=seq(from=10,to=1,by=-1.5))
## Regression coefficient estimates
{\tt SSGL\_logistic\_mod\$beta}
## Predicted n_test-dimensional vectors mu=E(Y_test) based on test data, X_test.
## The kth column of 'Y_pred' corresponds to the kth entry in 'lambda.'
SSGL\_logistic\_mod\$Y\_pred
## Classifications of the 8 groups. The kth column of 'classifications'
## corresponds to the kth entry in 'lambda.'
SSGL_logistic_mod$classifications
## Plot lambda vs. GIC
plot(SSGL_logistic_mod$lambda0, SSGL_logistic_mod$GIC, type='l')
## Model selection with the lambda that minimizes GIC
{\tt SSGL\_logistic\_mod\$lambda0\_min}
SSGL_logistic_mod$min_index
SSGL_logistic_mod$classifications[, SSGL_logistic_mod$min_index]
SSGL_logistic_mod$beta[, SSGL_logistic_mod$min_index]
```

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