Package 'srlars'

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Type Package

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Maintainer Anthony Christidis <anthony.christidis@stat.ubc.ca></anthony.christidis@stat.ubc.ca>	
Description Functions to perform split robust least angle regression. The approach first uses the least angle regression algorithm to split the variables into the models of an ensemble and robust estimates of the correlation between predictors. An elastic net estimator is then applied to the selected predictors in each model using the imputed data from the detect deviating cell (DDC) method.	
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Author Anthony Christidis [aut, cre], Gabriela Cohen-Freue [aut]	
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coef.srlars

Coefficients for srlars Object

Description

coef.srlars returns the coefficients for a srlars object.

Usage

```
## S3 method for class 'srlars'
coef(object, group_index = NULL, ...)
```

Arguments

object An object of class srlars

group_index Groups included in the ensemble. Default setting includes all the groups.

... Additional arguments for compatibility.

Value

The coefficients for the srlars object.

Author(s)

Anthony-Alexander Christidis, <anthony.christidis@stat.ubc.ca>

See Also

srlars

Examples

```
# Required library
library(mvnfast)

# Simulation parameters
n <- 50
p <- 500
rho.within <- 0.8
rho.between <- 0.2
p.active <- 100
group.size <- 25
snr <- 3
contamination.prop <- 0.2

# Setting the seed
set.seed(0)

# Block correlation structure</pre>
```

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```
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.between</pre>
for(group in 0:(p.active/group.size - 1))
  sigma.mat[(group*group.size+1):(group*group.size+group.size),
  (group*group.size+1):(group*group.size+group.size)] <- rho.within</pre>
diag(sigma.mat) <- 1</pre>
# Simulation of beta vector
true.beta <- c(runif(p.active, 0, 5)*(-1)^rbinom(p.active, 1, 0.7), rep(0, p - p.active))
# Setting the SD of the variance
sigma <- as.numeric(sqrt(t(true.beta) %*% sigma.mat %*% true.beta)/sqrt(snr))</pre>
# Simulation of uncontaminated data
x \leftarrow mvnfast::rmvn(n, mu = rep(0, p), sigma = sigma.mat)
y <- x %*% true.beta + rnorm(n, 0, sigma)
# Contamination of data
contamination_indices <- 1:floor(n*contamination.prop)</pre>
k_lev <- 2
k_slo <- 100
x_train <- x
y_train <- y
beta_cont <- true.beta</pre>
beta_cont[true.beta!=0] <- beta_cont[true.beta!=0]*(1 + k_slo)</pre>
beta_cont[true.beta==0] <- k_slo*max(abs(true.beta))</pre>
for(cont_id in contamination_indices){
  a \leftarrow runif(p, min = -1, max = 1)
  a \leftarrow a - as.numeric((1/p)*t(a) %*% rep(1, p))
  x_{train[cont_id,]} \leftarrow mvnfast::rmvn(1, rep(0, p), 0.1^2*diag(p)) +
    k_lev * a / as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# Ensemble models
ensemble_fit <- srlars(x_train, y_train,</pre>
                         n_{models} = 5,
                         model_saturation = c("fixed", "p-value")[1],
                         alpha = 0.05, model_size = n - 1,
                         robust = TRUE,
                         compute_coef = TRUE,
                         en_alpha = 1/4)
# Ensemble coefficients
ensemble_coefs <- coef(ensemble_fit, group_index = 1:ensemble_fit$n_models)</pre>
sens_ensemble <- sum(which((ensemble_coefs[-1]!=0)) <= p.active)/p.active</pre>
spec_ensemble <- sum(which((ensemble_coefs[-1]!=0)) <= p.active)/sum(ensemble_coefs[-1]!=0)</pre>
# Simulation of test data
m <- 2e3
x_{test} \leftarrow mvnfast::rmvn(m, mu = rep(0, p), sigma = sigma.mat)
y_test <- x_test %*% true.beta + rnorm(m, 0, sigma)</pre>
```

4 predict.srlars

predict.srlars

Predictions for srlars Object

Description

predict.srlars returns the predictions for a srlars object.

Usage

```
## S3 method for class 'srlars'
predict(object, newx, group_index = NULL, dynamic = FALSE, ...)
```

Arguments

object An object of class srlars
newx New data for predictions.

group_index Groups included in the ensemble. Default setting includes all the groups.

dynamic Argument to determine whether dynamic predictions are used based on deviat-

ing cells. Default is FALSE.

... Additional arguments for compatibility.

Value

The predictions for the srlars object.

Author(s)

Anthony-Alexander Christidis, <anthony.christidis@stat.ubc.ca>

See Also

srlars

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Examples

```
# Required library
library(mvnfast)
# Simulation parameters
n <- 50
p <- 500
rho.within <- 0.8
rho.between <- 0.2
p.active <- 100
group.size <- 25
snr <- 3
contamination.prop <- 0.2</pre>
# Setting the seed
set.seed(0)
# Block correlation structure
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.between</pre>
for(group in 0:(p.active/group.size - 1))
  sigma.mat[(group*group.size+1):(group*group.size+group.size),
  (group*group.size+1):(group*group.size+group.size)] <- rho.within</pre>
diag(sigma.mat) <- 1</pre>
# Simulation of beta vector
true.beta <- c(runif(p.active, 0, 5)*(-1)^rbinom(p.active, 1, 0.7), rep(0, p - p.active))
# Setting the SD of the variance
sigma <- as.numeric(sqrt(t(true.beta) %*% sigma.mat %*% true.beta)/sqrt(snr))</pre>
# Simulation of uncontaminated data
x \leftarrow mvnfast::rmvn(n, mu = rep(0, p), sigma = sigma.mat)
y <- x %*% true.beta + rnorm(n, 0, sigma)
# Contamination of data
contamination_indices <- 1:floor(n*contamination.prop)</pre>
k_lev <- 2
k_slo <- 100
x_train <- x
y_train <- y
beta_cont <- true.beta</pre>
beta_cont[true.beta!=0] <- beta_cont[true.beta!=0]*(1 + k_slo)</pre>
beta_cont[true.beta==0] <- k_slo*max(abs(true.beta))</pre>
for(cont_id in contamination_indices){
  a \leftarrow runif(p, min = -1, max = 1)
  a <- a - as.numeric((1/p)*t(a) %*% rep(1, p))
  x_{\text{train}}[\text{cont}_{\text{id}}] \leftarrow \text{mvnfast}::\text{rmvn}(1, \text{rep}(0, p), 0.1^2*\text{diag}(p)) +
    k_lev * a / as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
```

```
# Ensemble models
ensemble_fit <- srlars(x_train, y_train,</pre>
                        n_{models} = 5,
                        model_saturation = c("fixed", "p-value")[1],
                        alpha = 0.05, model_size = n - 1,
                        robust = TRUE,
                        compute_coef = TRUE,
                        en_alpha = 1/4)
# Ensemble coefficients
ensemble_coefs <- coef(ensemble_fit, group_index = 1:ensemble_fit$n_models)</pre>
sens_ensemble <- sum(which((ensemble_coefs[-1]!=0)) <= p.active)/p.active</pre>
spec\_ensemble <- sum(which((ensemble\_coefs[-1]!=0)) <= p.active)/sum(ensemble\_coefs[-1]!=0)
# Simulation of test data
m <- 2e3
x_{test} \leftarrow mvnfast::rmvn(m, mu = rep(0, p), sigma = sigma.mat)
y_test <- x_test %*% true.beta + rnorm(m, 0, sigma)</pre>
# Prediction of test samples
ensemble_preds <- predict(ensemble_fit, newx = x_test,</pre>
                           group_index = 1:ensemble_fit$n_models,
                            dynamic = FALSE)
mspe_ensemble <- mean((y_test - ensemble_preds)^2)/sigma^2</pre>
```

srlars

Robust Split Least Angle Regression

Description

srlars performs split robust least angle regression.

Usage

```
srlars(
    x,
    y,
    n_models = 1,
    model_saturation = c("fixed", "p-value")[1],
    alpha = 0.05,
    model_size = NULL,
    robust = TRUE,
    compute_coef = FALSE,
    en_alpha = 1/4
)
```

Arguments

x Design matrix.y Response vector.

n_models Number of models into which the variables are split.

model_saturation

Criterion to determine if a model is saturated. Must be one of "fixed" (default)

or "p-value".

alpha P-value used to determine when the model is saturated

model_size Size of the models in the ensemble.

robust Argument to determine if robust measures of location, scale and correlation are

used. Default is TRUE.

compute_coef Argument to determine if coefficients are computed (via the elastic net) for each

model. Default is FALSE.

en_alpha Elastic net mixing parmeter for parameters shrinkage. Default is 1/4.

Value

An object of class srlars

Author(s)

Anthony-Alexander Christidis, <anthony.christidis@stat.ubc.ca>

See Also

```
coef.srlars, predict.srlars
```

Examples

```
# Required library
library(mvnfast)
# Simulation parameters
n <- 50
p <- 500
rho.within <- 0.8
rho.between <- 0.2
p.active <- 100
group.size <- 25
snr <- 3
contamination.prop <- 0.2
# Setting the seed
set.seed(0)
# Block correlation structure
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.between</pre>
```

```
for(group in 0:(p.active/group.size - 1))
  sigma.mat[(group*group.size+1):(group*group.size+group.size),
  (group*group.size+1):(group*group.size+group.size)] <- rho.within</pre>
diag(sigma.mat) <- 1</pre>
# Simulation of beta vector
true.beta <- c(runif(p.active, 0, 5)*(-1)^rbinom(p.active, 1, 0.7), rep(0, p - p.active))
# Setting the SD of the variance
sigma <- as.numeric(sqrt(t(true.beta) %*% sigma.mat %*% true.beta)/sqrt(snr))</pre>
# Simulation of uncontaminated data
x \leftarrow mvnfast::rmvn(n, mu = rep(0, p), sigma = sigma.mat)
y <- x %*% true.beta + rnorm(n, 0, sigma)
# Contamination of data
contamination_indices <- 1:floor(n*contamination.prop)</pre>
k_lev <- 2
k_slo <- 100
x_train <- x
y_train <- y
beta_cont <- true.beta</pre>
beta\_cont[true.beta!=0] <- beta\_cont[true.beta!=0]*(1 + k\_slo)
beta_cont[true.beta==0] <- k_slo*max(abs(true.beta))</pre>
for(cont_id in contamination_indices){
  a \leftarrow runif(p, min = -1, max = 1)
  a <- a - as.numeric((1/p)*t(a) %*% rep(1, p))</pre>
  x_{\text{train}}[\text{cont}_{\text{id}}] \leftarrow \text{mvnfast}::\text{rmvn}(1, \text{rep}(0, p), 0.1^2*\text{diag}(p)) +
    k_lev * a / as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# Ensemble models
ensemble_fit <- srlars(x_train, y_train,</pre>
                         n_{models} = 5,
                         model_saturation = c("fixed", "p-value")[1],
                         alpha = 0.05, model_size = n - 1,
                         robust = TRUE,
                         compute_coef = TRUE,
                         en_alpha = 1/4)
# Ensemble coefficients
ensemble_coefs <- coef(ensemble_fit, group_index = 1:ensemble_fit$n_models)</pre>
sens_ensemble \leftarrow sum(which((ensemble_coefs[-1]!=0)) \leftarrow p.active)/p.active
spec_ensemble <- sum(which((ensemble_coefs[-1]!=0)) <= p.active)/sum(ensemble_coefs[-1]!=0)</pre>
# Simulation of test data
m < - 2e3
x_{test} \leftarrow mvnfast::rmvn(m, mu = rep(0, p), sigma = sigma.mat)
y_test <- x_test %*% true.beta + rnorm(m, 0, sigma)</pre>
# Prediction of test samples
```

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