

Package ‘srlars’

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Type Package

Title Fast and Scalable Cellwise-Robust Ensemble

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Description Functions to perform robust variable selection and regression using the Fast and Scalable Cellwise-Robust Ensemble (FSCORE) algorithm. The approach establishes a robust foundation using the Detect Deviating Cells (DDC) algorithm and robust correlation estimates. It then employs a competitive ensemble architecture where a robust Least Angle Regression (LARS) engine proposes candidate variables and cross-validation arbitrates their assignment. A final robust MM-estimator is applied to the selected predictors.

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Encoding UTF-8

Biarch true

Imports cellWise, robustbase, mvnfast

Suggests testthat

RoxygenNote 7.3.3

NeedsCompilation no

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coef.srlars *Coefficients for srlars Object*

Description

coef.srlars returns the averaged coefficients for a srlars object.

Usage

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

Arguments

object	An object of class srlars.
model_index	Indices of the sub-models to include in the ensemble average. Default is NULL, which includes all models.
...	Additional arguments for compatibility.

Value

A numeric vector containing the averaged intercept (first element) and slope coefficients.

Author(s)

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

See Also

[srlars](#), [predict.srlars](#)

Examples

```
# Required libraries  
library(mvnfast)  
library(cellwise)  
library(robustbase)  
  
# Simulation parameters  
n <- 50  
p <- 100  
rho.within <- 0.8  
rho.between <- 0.2  
p.active <- 20  
group.size <- 5  
snr <- 3  
contamination.prop <- 0.1
```

```

# Setting the seed
set.seed(0)

# Block correlation structure
sigma.mat <- matrix(0, p, p)
sigma.mat[1:p.active, 1:p.active] <- rho.between
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
diag(sigma.mat) <- 1

# 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))

# Simulation of uncontaminated data
x <- mvnfast::rmvn(n, mu = rep(0, p), sigma = sigma.mat)
y <- x %% true.beta + rnorm(n, 0, sigma)

# Cellwise contamination
contamination_indices <- sample(1:(n * p), round(n * p * contamination.prop))
x_train <- x
x_train[contamination_indices] <- runif(length(contamination_indices), -10, 10)

# FSCORE Ensemble model
ensemble_fit <- srlars(x_train, y,
                      n_models = 5,
                      tolerance = 0.01,
                      robust = TRUE,
                      compute_coef = TRUE)

# Ensemble coefficients
# Default: Average over all models
ensemble_coefs <- coef(ensemble_fit)

# Sensitivity (Recall)
active_selected <- which(ensemble_coefs[-1] != 0)
true_active <- which(true.beta != 0)
recall <- length(intersect(active_selected, true_active)) / length(true_active)
print(paste("Recall:", recall))

# Precision
if(length(active_selected) > 0){
  precision <- length(intersect(active_selected, true_active)) / length(active_selected)
} else {
  precision <- 0
}
print(paste("Precision:", precision))

```

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, model_index = NULL, dynamic = TRUE, ...)
```

Arguments

object	An object of class srlars.
newx	New data matrix for predictions.
model_index	Indices of the sub-models to include in the ensemble. Default is NULL (all models).
dynamic	Logical. If TRUE, and the model was trained robustly, the new data newx is cleaned using DDCpredict before prediction. This ensures consistency with the robust training phase. Default is TRUE.
...	Additional arguments for compatibility.

Value

A numeric vector of predictions.

Author(s)

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

See Also

[srlars](#)

Examples

```
# Required libraries  
library(mvnfast)  
library(cellwise)  
library(robustbase)  
  
# Simulation parameters  
n <- 50  
p <- 100  
rho.within <- 0.8  
rho.between <- 0.2
```

```

p.active <- 20
group.size <- 5
snr <- 3
contamination.prop <- 0.1

# Setting the seed
set.seed(0)

# Block correlation structure
sigma.mat <- matrix(0, p, p)
sigma.mat[1:p.active, 1:p.active] <- rho.between
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
diag(sigma.mat) <- 1

# 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))

# Simulation of uncontaminated data
x <- mvnfast::rmvn(n, mu = rep(0, p), sigma = sigma.mat)
y <- x %% true.beta + rnorm(n, 0, sigma)

# Cellwise contamination
contamination_indices <- sample(1:(n * p), round(n * p * contamination.prop))
x_train <- x
x_train[contamination_indices] <- runif(length(contamination_indices), -10, 10)

# FSCORE Ensemble model
ensemble_fit <- srlars(x_train, y,
                      n_models = 5,
                      tolerance = 0.01,
                      robust = TRUE,
                      compute_coef = TRUE)

# Simulation of test data
m <- 50
x_test <- mvnfast::rmvn(m, mu = rep(0, p), sigma = sigma.mat)
y_test <- x_test %% true.beta + rnorm(m, 0, sigma)

# Prediction of test samples
# Default: Averaged prediction from all models
# Dynamic imputation is used by default if the model is robust
ensemble_preds <- predict(ensemble_fit, newx = x_test)

# Evaluate MSPE
mspe <- mean((y_test - ensemble_preds)^2) / sigma^2
print(paste("MSPE:", mspe))

```

`srlars`*Fast and Scalable Cellwise-Robust Ensemble (FSCRE)*

Description

`srlars` performs the FSCRE algorithm for robust variable selection and regression.

Usage

```
srlars(  
  x,  
  y,  
  n_models = 5,  
  tolerance = 1e-08,  
  max_predictors = NULL,  
  robust = TRUE,  
  compute_coef = TRUE  
)
```

Arguments

<code>x</code>	Design matrix (n x p).
<code>y</code>	Response vector (n x 1).
<code>n_models</code>	Number of models in the ensemble (K). Default is 5.
<code>tolerance</code>	Relative improvement tolerance for stopping (tau). Default is 1e-8.
<code>max_predictors</code>	Maximum total number of variables to select across all models. Default is n * n_models.
<code>robust</code>	Logical. If TRUE (default), performs DDC imputation and robust initialization.
<code>compute_coef</code>	Logical. If TRUE, fits the final robust MM-models. Default is TRUE.

Value

An object of class `srlars` containing the selected variables and coefficients.

Author(s)

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

See Also

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

Examples

```
# Required libraries
library(mvnfast)
library(cellwise)
library(robustbase)

# Simulation parameters
n <- 50
p <- 100
rho.within <- 0.8
rho.between <- 0.2
p.active <- 20
group.size <- 5
snr <- 3
contamination.prop <- 0.1

# Setting the seed
set.seed(0)

# Block correlation structure
sigma.mat <- matrix(0, p, p)
sigma.mat[1:p.active, 1:p.active] <- rho.between
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
diag(sigma.mat) <- 1

# 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))

# Simulation of uncontaminated data
x <- mvnfast::rmvn(n, mu = rep(0, p), sigma = sigma.mat)
y <- x %*% true.beta + rnorm(n, 0, sigma)

# Cellwise contamination
contamination_indices <- sample(1:(n * p), round(n * p * contamination.prop))
x_train <- x
x_train[contamination_indices] <- runif(length(contamination_indices), -10, 10)

# FSCORE Ensemble model
ensemble_fit <- srlars(x_train, y,
                      n_models = 5,
                      tolerance = 1e-8,
                      robust = TRUE,
                      compute_coef = TRUE)

# Check selected variables
print(ensemble_fit$active.sets)
```

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