Package 'RMSS'

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Type Package

Title Robust Multi-Model Subset Selection

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Description Efficient algorithms for generating ensembles of robust, sparse and diverse models via robust multimodel subset selection (RMSS). The robust ensembles are generated by minimizing the sum of the least trimmed square loss of the models in the ensembles under constraints for the size of the models and the sharing of the predictors. Tuning parameters for the robustness, sparsity and diversity of the robust ensemble are selected by cross-validation.
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coef.cv.RMSS

Coefficients for cv.RMSS Object

Description

coef.cv.RMSS returns the coefficients for a cv.RMSS object.

Usage

```
## S3 method for class 'cv.RMSS'
coef(
  object,
  h_ind = NULL,
  t_ind = NULL,
  u_ind = NULL,
  individual_models = FALSE,
  group_index = NULL,
  ...
)
```

Arguments

object	An object of class cv.RMSS.
h_ind	Index for robustness parameter.
t_ind	Index for sparsity parameter.
u_ind	Index for diversity parameter.
individual_mode	els
	Argument to determine whether the coefficients of each model are returned. Default is FALSE.
group_index	Groups included in the ensemble. Default setting includes all the groups.
	Additional arguments for compatibility.

Value

The coefficients for the cv.RMSS object.

Author(s)

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

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See Also

cv.RMSS

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5</pre>
p.active <- 15
snr <- 2
contamination.prop <- 0.3
# Setting the seed
set.seed(0)
# Block Correlation
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.inactive</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</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 test data
m <- 2e3
x_{\text{test}} \leftarrow \text{mvnfast}::\text{rmvn}(m, mu = \text{rep}(0, p), \text{sigma} = \text{sigma.mat})
y_test <- x_test %*% true.beta + rnorm(m, 0, sigma)</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){
```

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```
a \leftarrow runif(p, min = -1, max = 1)
a \leftarrow 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_{\text{lev}}*a / p.
                           as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
 y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# CV RMSS
rmss_fit <- cv.RMSS(x = x_train, y = y_train,</pre>
                      n_{models} = 3,
                      h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                       initial_estimator = "robStepSplitReg",
                       tolerance = 1e-1,
                      max_iter = 1e3,
                      neighborhood_search = FALSE,
                      neighborhood_search_tolerance = 1e-1,
                      n_folds = 5,
                       alpha = 1/4,
                       gamma = 1,
                      n_{threads} = 1
rmss_coefs <- coef(rmss_fit,</pre>
                     h_ind = rmss_fit$h_opt,
                     t_ind = rmss_fit$t_opt,
                     u_ind = rmss_fit$u_opt,
                     group_index = 1:rmss_fit$n_models)
sens_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/p.active</pre>
spec_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/sum(rmss_coefs[-1]!=0)</pre>
rmss_preds <- predict(rmss_fit, newx = x_test,</pre>
                        h_ind = rmss_fit$h_opt,
                        t_ind = rmss_fit$t_opt,
                         u_ind = rmss_fit$u_opt,
                         group_index = 1:rmss_fit$n_models,
                         dynamic = FALSE)
rmss_mspe <- mean((y_test - rmss_preds)^2)/sigma^2</pre>
```

coef.RMSS

Coefficients for RMSS Object

Description

coef.RMSS returns the coefficients for a RMSS object.

Usage

```
## S3 method for class 'RMSS'
coef(
  object,
  h_ind,
```

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```
t_ind,
  u_ind,
  individual_models = FALSE,
  group_index = NULL,
  ...
)
```

Arguments

object An object of class RMSS.

h_ind Index for robustness parameter.

t_ind Index for sparsity parameter.

u_ind Index for diversity parameter.

individual_models

Argument to determine whether the coefficients of each model are returned.

Default is FALSE.

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

Additional arguments for compatibility.

Value

The coefficients for the RMSS object.

Author(s)

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

See Also

RMSS

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5
p.active <- 15
snr <- 2
contamination.prop <- 0.3
# Setting the seed
set.seed(0)
# Block Correlation
sigma.mat <- matrix(0, p, p)
sigma.mat[1:p.active, 1:p.active] <- rho.inactive</pre>
```

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```
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</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 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>
# 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_{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>
}
# RMSS
rmss_fit <- RMSS(x = x_train, y = y_train,</pre>
                  n_{models} = 3,
                  h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                  initial_estimator = "robStepSplitReg",
                  tolerance = 1e-1,
                  max_iter = 1e3,
                  neighborhood_search = FALSE,
                  neighborhood_search_tolerance = 1e-1)
rmss_coefs <- coef(rmss_fit,</pre>
                    h_{ind} = 1, t_{ind} = 2, u_{ind} = 1,
                    group_index = 1:rmss_fit$n_models)
sens_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/p.active</pre>
spec_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/sum(rmss_coefs[-1]!=0)</pre>
```

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cv.RMSS

Cross-Validatoin for Robust Multi-Model Subset Selection

Description

cv. RMSS performs the cross-validation procedure for robust multi-model subset selection.

Usage

```
cv.RMSS(
  Х,
  у,
  n_models,
  h_grid,
  t_grid,
  u_grid,
  initial_estimator = c("robStepSplitReg", "srlars")[1],
  tolerance = 0.1,
  max_iter = 1000,
  neighborhood_search = FALSE,
  neighborhood_search_tolerance = 0.1,
  cv_criterion = c("tau", "trimmed")[1],
  n_folds = 5,
  alpha = 1/4,
  gamma = 1,
  n_{threads} = 1
)
```

Arguments

```
x Design matrix.

y Response vector.

n_models Number of models into which the variables are split.

h_grid Grid for robustness parameter.

t_grid Grid for sparsity parameter.

u_grid Grid for diversity parameter.

initial_estimator

Method used for initial estimator. Must be one of "robStepSplitReg" (default) or "srlars".
```

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Tolerance level for convergence of PSBGD algorithm. tolerance max_iter Maximum number of iterations in PSBGD algorithm. neighborhood_search Neighborhood search to improve solution. Default is FALSE. neighborhood_search_tolerance Tolerance parameter for neighborhood search. Default is 1e-1. Criterion to use for cross-validation procedure. Must be one of "tau" (default) cv_criterion or "trimmed". n_folds Number of folds for cross-validation procedure. Default is 5. alpha Proportion of trimmed samples for cross-validation procedure. Default is 1/4. Weight parameter for ensemble MSPE (gamma) and average MSPE of individgamma ual models (1 - gamma). Default is 1.

n_threads Number of threads used by OpenMP for multithreading over the folds. Default

is 1.

Value

An object of class cv.RMSS

Author(s)

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

See Also

```
coef.cv.RMSS, predict.cv.RMSS
```

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5</pre>
p.active <- 15
snr <- 2
contamination.prop <- 0.3</pre>
# Setting the seed
set.seed(0)
# Block Correlation
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.inactive</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</pre>
diag(sigma.mat) <- 1</pre>
```

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```
# 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 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>
# 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))</pre>
 x_{\text{train}}[\text{cont\_id,}] \leftarrow \text{mvnfast::rmvn}(1, \text{rep}(0, p), 0.1^2*\text{diag}(p)) + k_{\text{lev}}*a / p
                          as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# CV RMSS
rmss_fit <- cv.RMSS(x = x_train, y = y_train,</pre>
                      n_{models} = 3,
                      h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                      initial_estimator = "robStepSplitReg",
                      tolerance = 1e-1,
                      max_iter = 1e3,
                      neighborhood_search = FALSE,
                      neighborhood_search_tolerance = 1e-1,
                      n_folds = 5,
                      alpha = 1/4,
                      gamma = 1,
                      n_{threads} = 1
rmss_coefs <- coef(rmss_fit,</pre>
                     h_ind = rmss_fit$h_opt,
                     t_ind = rmss_fit$t_opt,
                     u_ind = rmss_fit$u_opt,
                     group_index = 1:rmss_fit$n_models)
```

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predict.cv.RMSS

Predictions for cv.RMSS Object

Description

predict.cv.RMSS returns the predictions for a cv.RMSS object.

Usage

```
## $3 method for class 'cv.RMSS'
predict(
  object,
  newx,
  h_ind = NULL,
  t_ind = NULL,
  u_ind = NULL,
  group_index = NULL,
  dynamic = FALSE,
  ...
)
```

Arguments

An object of class cv.RMSS. object newx New data for predictions. h_ind Index for robustness parameter. t_ind Index for sparsity parameter. u_ind Index for diversity parameter. group_index Groups included in the ensemble. Default setting includes all the groups. dynamic Argument to determine whether dynamic predictions are used based on deviating cells. Default is FALSE. Additional arguments for compatibility.

Value

The predictions for the cv.RMSS object.

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Author(s)

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

See Also

```
cv.RMSS
```

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5</pre>
p.active <- 15
snr <- 2
contamination.prop <- 0.3
# Setting the seed
set.seed(0)
# Block Correlation
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.inactive
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</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 test data
m <- 2e3
x_{test} < -mvnfast::rmvn(m, mu = rep(0, p), sigma = sigma.mat)
y_test <- x_test %*% true.beta + rnorm(m, 0, sigma)</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_{ev} < -2
k_slo <- 100
x_train <- x
y_train <- y
```

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```
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_{\text{train}}[\text{cont}_{\text{id}}] \leftarrow \text{mvnfast}::\text{rmvn}(1, \text{rep}(0, p), 0.1^2*\text{diag}(p)) + k_{\text{lev}}*a / p.
                           as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# CV RMSS
rmss_fit <- cv.RMSS(x = x_train, y = y_train,</pre>
                      n_{models} = 3,
                      h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                      initial_estimator = "robStepSplitReg",
                      tolerance = 1e-1,
                      max_iter = 1e3,
                      neighborhood_search = FALSE,
                      neighborhood_search_tolerance = 1e-1,
                      n_folds = 5,
                      alpha = 1/4,
                      gamma = 1,
                      n_{threads} = 1
rmss_coefs <- coef(rmss_fit,</pre>
                     h_ind = rmss_fit$h_opt,
                     t_ind = rmss_fit$t_opt,
                     u_ind = rmss_fit$u_opt,
                     group_index = 1:rmss_fit$n_models)
sens_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/p.active</pre>
spec_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/sum(rmss_coefs[-1]!=0)</pre>
rmss_preds <- predict(rmss_fit, newx = x_test,</pre>
                        h_ind = rmss_fit$h_opt,
                         t_ind = rmss_fit$t_opt,
                         u_ind = rmss_fit$u_opt,
                         group_index = 1:rmss_fit$n_models,
                         dynamic = FALSE)
rmss_mspe <- mean((y_test - rmss_preds)^2)/sigma^2</pre>
```

predict.RMSS

Predictions for RMSS Object

Description

predict.RMSS returns the predictions for a RMSS object.

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Usage

```
## S3 method for class 'RMSS'
predict(
  object,
  newx,
  h_ind,
  t_ind,
  u_ind,
  group_index = NULL,
  dynamic = FALSE,
  ...
)
```

Arguments

object	An object of class RMSS.
newx	New data for predictions.
h_ind	Index for robustness parameter.
t_ind	Index for sparsity parameter.
u_ind	Index for diversity parameter.
group_index	Groups included in the ensemble. Default setting includes all the groups.
dynamic	Argument to determine whether dynamic predictions are used based on deviating cells. Default is FALSE.
	Additional arguments for compatibility.

Value

The predictions for the RMSS object.

Author(s)

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

See Also

RMSS

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5
p.active <- 15
snr <- 2
contamination.prop <- 0.3</pre>
```

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```
# Setting the seed
set.seed(0)
# Block Correlation
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.inactive</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</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 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>
# 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_{\text{train}}[\text{cont\_id},] \leftarrow \text{mvnfast}::rmvn(1, rep(0, p), 0.1^2*diag(p)) + k_lev * a / lev * 
                                                        as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
    y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# RMSS
rmss_fit <- RMSS(x = x_train, y = y_train,</pre>
                                        n_{models} = 3,
                                        h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                                        initial_estimator = "robStepSplitReg",
                                        tolerance = 1e-1,
                                        max_iter = 1e3,
```

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RMSS

Robust Multi-Model Subset Selection

Description

RMSS performs robust multi-model subset selection.

Usage

```
RMSS(
    x,
    y,
    n_models,
    h_grid,
    t_grid,
    u_grid,
    initial_estimator = c("robStepSplitReg", "srlars")[1],
    tolerance = 0.1,
    max_iter = 1000,
    neighborhood_search = FALSE,
    neighborhood_search_tolerance = 0.1
)
```

Arguments

Х	Design matrix.
у	Response vector.
n_models	Number of models into which the variables are split.
h_grid	Grid for robustness parameter.
t_grid	Grid for sparsity parameter.
u_grid	Grid for diversity parameter.

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```
initial_estimator
```

Method used for initial estimator. Must be one of "robStepSplitReg" (default)

or "srlars".

tolerance Tolerance level for convergence of PSBGD algorithm.

max_iter Maximum number of iterations in PSBGD algorithm.

neighborhood_search

Neighborhood search to improve solution. Default is FALSE.

neighborhood_search_tolerance

Tolerance parameter for neighborhood search. Default is 1e-1.

Value

An object of class RMSS

Author(s)

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

See Also

```
coef.RMSS, predict.RMSS
```

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5
p.active <- 15
snr <- 2
contamination.prop <- 0.3
# Setting the seed
set.seed(0)
# Block Correlation
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.inactive</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</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 test data
m <- 2e3
x_{\text{test}} < - mvnfast::rmvn(m, mu = rep(0, p), sigma = sigma.mat)
y_test <- x_test %*% true.beta + rnorm(m, 0, sigma)</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_{\text{lev}}*a / p.
                           as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
# RMSS
rmss_fit <- RMSS(x = x_train, y = y_train,
                   n_{models} = 3,
                   h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                   initial_estimator = "robStepSplitReg",
                   tolerance = 1e-1,
                   max_iter = 1e3,
                   neighborhood_search = FALSE,
                   neighborhood_search_tolerance = 1e-1)
rmss_coefs <- coef(rmss_fit,</pre>
                     h_{ind} = 1, t_{ind} = 2, u_{ind} = 1,
                     group_index = 1:rmss_fit$n_models)
sens_rmss <- sum(which((rmss_coefs[-1]!=0)) <= p.active)/p.active</pre>
spec\_rmss <- sum(which((rmss\_coefs[-1]!=0)) <= p.active)/sum(rmss\_coefs[-1]!=0)
rmss_preds <- predict(rmss_fit, newx = x_test,
                        h_{ind} = 1, t_{ind} = 2, u_{ind} = 1,
                        group_index = 1:rmss_fit$n_models,
                        dynamic = FALSE)
rmss_mspe <- mean((y_test - rmss_preds)^2)/sigma^2</pre>
```

trimmed_samples

Description

trimmed_samples returns the coefficients for a RMSS or cv.RMSS object.

Usage

```
trimmed_samples(
  object,
h_ind = NULL,
t_ind = NULL,
u_ind = NULL,
group_index = NULL,
...
)
```

Arguments

object	An object of class RMSS
h_ind	Index for robustness parameter.
t_ind	Index for sparsity parameter.
u_ind	Index for diversity parameter.
group_index	Groups included in the ensemble. Default setting includes all the groups.
	Additional arguments for compatibility.

Value

The trimmed samples for the RMSS or cv.RMSS object.

Author(s)

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See Also

RMSS

```
# Simulation parameters
n <- 50
p <- 100
rho <- 0.8
rho.inactive <- 0.2
group.size <- 5
p.active <- 15
snr <- 2
contamination.prop <- 0.3
# Setting the seed
set.seed(0)</pre>
```

```
# Block Correlation
sigma.mat <- matrix(0, p, p)</pre>
sigma.mat[1:p.active, 1:p.active] <- rho.inactive</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</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 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>
# 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 / (a)
                          as.numeric(sqrt(t(a) %*% solve(sigma.mat) %*% a))
  y_train[cont_id] <- t(x_train[cont_id,]) %*% beta_cont</pre>
}
# RMSS
rmss_fit <- RMSS(x = x_train, y = y_train,
                  n_{models} = 3,
                  h_{grid} = c(35), t_{grid} = c(6, 8, 10), u_{grid} = c(1:3),
                  tolerance = 1e-1,
                  max_iter = 1e3,
                  neighborhood_search = FALSE,
                  neighborhood_search_tolerance = 1e-1)
rmss_coefs <- coef(rmss_fit,</pre>
                    h_{ind} = 1, t_{ind} = 2, u_{ind} = 1,
```

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