

Package ‘SDModels’

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Title Spectrally Deconfounded Models

Version 1.0.10

Description Screen for and analyze non-linear sparse direct effects in the presence of unobserved confounding using the spectral deconfounding techniques (Čeví, Bühlmann, and Meinshausen (2020) <jmlr.org/papers/v21/19-545.html>, Guo, Čeví, and Bühlmann (2022) <[doi:10.1214/21-AOS2152](https://doi.org/10.1214/21-AOS2152)>). These methods have been shown to be a good estimate for the true direct effect if we observe many covariates, e.g., high-dimensional settings, and we have fairly dense confounding. Even if the assumptions are violated, it seems like there is not much to lose, and the deconfounded models will, in general, estimate a function closer to the true one than classical least squares optimization. 'SDModels' provides functions SDAM() for Spectrally Deconfounded Additive Models (Scheidegger, Guo, and Bühlmann (2025) <[doi:10.1145/3711116](https://doi.org/10.1145/3711116)>) and SDForest() for Spectrally Deconfounded Random Forests (Ulmer, Scheidegger, and Bühlmann (2025) <[doi:10.48550/arXiv.2502.03969](https://doi.org/10.48550/arXiv.2502.03969)>).

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Imports data.tree, DiagrammeR, doParallel, ggplot2, GPUmatrix, gridExtra, locatexec, parallel, pbapply, Rdpack, tidyr, fda, grplasso, rlang

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URL <https://www.markus-ulmer.ch/SDModels/>

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copy.SDForest	<i>Copy a forest</i>
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Description

Returns a copy of the forest object. Might be useful if you want to keep the original forest in comparison to the pruned forest.

Usage

```
## S3 method for class 'SDForest'  
copy(object, ...)
```

Arguments

object	an SDForest object
...	Further arguments passed to or from other methods.

Value

A copy of the SDForest object

Author(s)

Markus Ulmer

See Also

[prune](#)

Examples

```
set.seed(1)  
X <- matrix(rnorm(10 * 20), nrow = 10)  
Y <- rnorm(10)  
fit <- SDForest(x = X, y = Y, nTree = 2, cp = 0.5)  
fit2 <- copy(fit)
```

`copy.SDTree`*Copy a tree*

Description

Returns a copy of the tree object. Might be useful if you want to keep the original tree in comparison to the pruned tree.

Usage

```
## S3 method for class 'SDTree'  
copy(object, ...)
```

Arguments

<code>object</code>	an SDTree object
<code>...</code>	Further arguments passed to or from other methods.

Value

A copy of the SDTree object

Author(s)

Markus Ulmer

See Also

[prune](#)

Examples

```
set.seed(1)  
X <- matrix(rnorm(10 * 20), nrow = 10)  
Y <- rnorm(10)  
fit <- SDTree(x = X, y = Y, cp = 0.5)  
fit2 <- copy(fit)
```

cvSDTree

*Cross-validation for the SDTree***Description**

Estimates the optimal complexity parameter for the SDTree using cross-validation. The transformations are estimated for each training set and validation set separately to ensure independence of the validation set.

Usage

```
cvSDTree(
  formula = NULL,
  data = NULL,
  x = NULL,
  y = NULL,
  max_leaves = NULL,
  cp = 0,
  min_sample = 5,
  mtry = NULL,
  fast = TRUE,
  Q_type = "trim",
  trim_quantile = 0.5,
  q_hat = 0,
  Qf = NULL,
  A = NULL,
  gamma = 0.5,
  gpu = FALSE,
  mem_size = 1e+07,
  max_candidates = 100,
  nfold = 3,
  cp_seq = NULL,
  mc.cores = 1,
  Q_scale = TRUE
)
```

Arguments

formula	Object of class formula or describing the model to fit of the form $y \sim x_1 + x_2 + \dots$ where y is a numeric response and x_1, x_2, \dots are vectors of covariates. Interactions are not supported.
data	Training data of class data.frame containing the variables in the model.
x	Predictor data, alternative to formula and data.
y	Response vector, alternative to formula and data.
max_leaves	Maximum number of leaves for the grown tree.

cp	Complexity parameter, minimum loss decrease to split a node. A split is only performed if the loss decrease is larger than $cp * initial_loss$, where <code>initial_loss</code> is the loss of the initial estimate using only a stump.
min_sample	Minimum number of observations per leaf. A split is only performed if both resulting leaves have at least <code>min_sample</code> observations.
mtry	Number of randomly selected covariates to consider for a split, if NULL all covariates are available for each split.
fast	If TRUE, only the optimal splits in the new leaves are evaluated and the previously optimal splits and their potential loss-decrease are reused. If FALSE all possible splits in all the leaves are reevaluated after every split.
Q_type	Type of deconfounding, one of 'trim', 'pca', 'no_deconfounding'. 'trim' corresponds to the Trim transform (Ćevič et al. 2020) as implemented in the Doubly debiased lasso (Guo et al. 2022), 'pca' to the PCA transformation (Paul et al. 2008). See get_Q .
trim_quantile	Quantile for Trim transform, only needed for trim and DDL_trim, see get_Q .
q_hat	Assumed confounding dimension, only needed for pca, see get_Q .
Qf	Spectral transformation, if NULL it is internally estimated using get_Q .
A	Numerical Anchor of class matrix. See get_W .
gamma	Strength of distributional robustness, $\gamma \in [0, \infty]$. See get_W .
gpu	If TRUE, the calculations are performed on the GPU. If it is properly set up.
mem_size	Amount of split candidates that can be evaluated at once. This is a trade-off between memory and speed can be decreased if either the memory is not sufficient or the gpu is too small.
max_candidates	Maximum number of split points that are proposed at each node for each covariate.
nfolds	Number of folds for cross-validation. It is recommended to not use more than 5 folds if the number of covariates is larger than the number of observations. In this case the spectral transformation could differ too much if the validation data is substantially smaller than the training data.
cp_seq	Sequence of complexity parameters cp to compare using cross-validation, if NULL a sequence from 0 to 0.6 with stepsize 0.002 is used.
mc.cores	Number of cores to use for parallel computation.
Q_scale	Should data be scaled to estimate the spectral transformation? Default is TRUE to not reduce the signal of high variance covariates, and we do not know of a scenario where this hurts.

Value

A list containing

cp_min	The optimal complexity parameter.
cp_table	A table containing the complexity parameter, the mean and the standard deviation of the loss on the validation sets for the complexity parameters. If multiple complexity parameters result in the same loss, only the one with the largest complexity parameter is shown.

Author(s)

Markus Ulmer

References

Guo Z, Cévid D, Bühlmann P (2022). “Doubly debiased lasso: High-dimensional inference under hidden confounding.” *The Annals of Statistics*, **50**(3). ISSN 0090-5364, doi:10.1214/21AOS2152.

Paul D, Bair E, Hastie T, Tibshirani R (2008). ““Preconditioning” for feature selection and regression in high-dimensional problems.” *The Annals of Statistics*, **36**(4). ISSN 0090-5364, doi:10.1214/009053607000000578.

Cévid D, Bühlmann P, Meinshausen N (2020). “Spectral Deconfounding via Perturbed Sparse Linear Models.” *J. Mach. Learn. Res.*, **21**(1). ISSN 1532-4435, <http://jmlr.org/papers/v21/19-545.html>.

See Also[SDTree prune](#). [SDTree regPath](#). [SDTree](#)**Examples**

```
set.seed(1)
n <- 50
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n, 0, 5)
cp <- cvSDTree(x = X, y = y, Q_type = 'no_deconfounding')
cp
```

fromList.SDForest	<i>SDForest fromList method</i>
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Description

Converts the trees in an SDForest object from class `list` to class `Node` (Glur 2023).

Usage

```
## S3 method for class 'SDForest'
fromList(object, ...)
```

Arguments

<code>object</code>	an SDForest object with the trees in list format
<code>...</code>	Further arguments passed to or from other methods.

Value

an SDForest object with the trees in Node format

Author(s)

Markus Ulmer

References

Glur C (2023). “data.tree: General Purpose Hierarchical Data Structure.” <https://CRAN.R-project.org/package=data.tree>.

See Also

[fromList](#) [fromList.SDTree](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5, nTree = 2)
fromList(toList(model))
```

fromList.SDTree

SDTree fromList method

Description

Converts the tree in an SDTree object from class list to class Node (Glur 2023).

Usage

```
## S3 method for class 'SDTree'
fromList(object, ...)
```

Arguments

object an SDTree object with the tree in list format
 ... Further arguments passed to or from other methods.

Value

an SDTree object with the tree in Node format

Author(s)

Markus Ulmer

References

Glur C (2023). “data.tree: General Purpose Hierarchical Data Structure.” <https://CRAN.R-project.org/package=data.tree>.

See Also

[toList](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)
fromList(toList(model))
```

f_four

Function of x on a fourier basis

Description

Function of x on a fourier basis with a subset of covariates having a causal effect on Y using the parameters beta. The function is given by:

$$f(x_i) = \sum_{j=1}^p 1_{j \in js} \sum_{k=1}^K (\beta_{j,k}^{(1)} \cos(0.2kx_j) + \beta_{j,k}^{(2)} \sin(0.2kx_j))$$

Usage

```
f_four(x, beta, js)
```

Arguments

x	a vector of covariates
beta	the parameter vector for the function f(X)
js	the indices of the causal covariates in X

Value

the value of the function f(x)

Author(s)

Markus Ulmer

See Also

[simulate_data_nonlinear](#)

Examples

```
set.seed(42)
# simulation of confounded data
sim_data <- simulate_data_nonlinear(q = 2, p = 150, n = 100, m = 2)
X <- sim_data$X
j <- sim_data$j[1]
apply(X, 1, function(x) f_four(x, sim_data$beta, j))
```

get_cp_seq.SDForest *Get the sequence of complexity parameters of an SDForest*

Description

This function extracts the sequence of complexity parameters of an SDForest that result in changes of the SDForest if pruned. Only cp values that differ in the first three digits after the decimal point are returned.

Usage

```
## S3 method for class 'SDForest'
get_cp_seq(object, ...)
```

Arguments

object an SDForest object
... Further arguments passed to or from other methods.

Value

A sequence of complexity parameters

Author(s)

Markus Ulmer

See Also

[regPath](#) [stabilitySelection](#) [get_cp_seq.SDTree](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', cp = 0, nTree = 2)
get_cp_seq(model)
```

get_cp_seq.SDTree *Get the sequence of complexity parameters of an SDTree*

Description

This function extracts the sequence of complexity parameters of an SDTree that result in changes of the tree structure if pruned. Only cp values that differ in the first three digits after the decimal point are returned.

Usage

```
## S3 method for class 'SDTree'
get_cp_seq(object, ...)
```

Arguments

object	an SDTree object
...	Further arguments passed to or from other methods.

Value

A sequence of complexity parameters

Author(s)

Markus Ulmer

See Also

[regPath](#) [stabilitySelection](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0)
get_cp_seq(model)
```

 get_Q

Estimation of spectral transformation

Description

Estimates the spectral transformation Q for spectral deconfounding by shrinking the leading singular values of the covariates.

Usage

```
get_Q(X, type, trim_quantile = 0.5, q_hat = 0, gpu = FALSE, scaling = TRUE)
```

Arguments

<code>X</code>	Numerical covariates of class <code>matrix</code> .
<code>type</code>	Type of deconfounding, one of 'trim', 'pca', 'no_deconfounding'. 'trim' corresponds to the Trim transform (Ćevic et al. 2020) as implemented in the Doubly debiased lasso (Guo et al. 2022), 'pca' to the PCA transformation (Paul et al. 2008) and 'no_deconfounding' to the Identity.
<code>trim_quantile</code>	Quantile for Trim transform, only needed for trim.
<code>q_hat</code>	Assumed confounding dimension, only needed for pca.
<code>gpu</code>	If TRUE, the calculations are performed on the GPU. If it is properly set up.
<code>scaling</code>	Whether <code>X</code> should be scaled before calculating the spectral transformation.

Value

Q of class `matrix`, the spectral transformation matrix.

Author(s)

Markus Ulmer

References

Guo Z, Ćevic D, Bühlmann P (2022). "Doubly debiased lasso: High-dimensional inference under hidden confounding." *The Annals of Statistics*, **50**(3). ISSN 0090-5364, doi:10.1214/21AOS2152.

Paul D, Bair E, Hastie T, Tibshirani R (2008). "Preconditioning" for feature selection and regression in high-dimensional problems." *The Annals of Statistics*, **36**(4). ISSN 0090-5364, doi:10.1214/009053607000000578.

Ćevic D, Bühlmann P, Meinshausen N (2020). "Spectral Deconfounding via Perturbed Sparse Linear Models." *J. Mach. Learn. Res.*, **21**(1). ISSN 1532-4435, <http://jmlr.org/papers/v21/19-545.html>.

Examples

```
set.seed(1)
X <- matrix(rnorm(50 * 20), nrow = 50)
Q_trim <- get_Q(X, 'trim')
Q_pca <- get_Q(X, 'pca', q_hat = 5)
Q_plain <- get_Q(X, 'no_deconfounding')
```

get_W

*Estimation of anchor transformation***Description**

Estimates the anchor transformation for the Anchor-Objective. The anchor transformation is $W = I - (1 - \sqrt{\gamma})\Pi_A$, where $\Pi_A = A(A^T A)^{-1}A^T$. For $\gamma = 1$ this is just the identity. For $\gamma = 0$ this corresponds to residuals after orthogonal projecting onto A. For large γ this is close to the orthogonal projection onto A, scaled by γ . The estimator $\operatorname{argmin}_f \|W(Y - f(X))\|^2$ corresponds to the Anchor-Regression Estimator (Rothenhäusler et al. 2021), (Bühlmann 2020).

Usage

```
get_W(A, gamma, intercept = FALSE, gpu = FALSE)
```

Arguments

A	Numerical Anchor of class matrix.
gamma	Strength of distributional robustness, $\gamma \in [0, \infty]$.
intercept	Logical, whether to include an intercept in the anchor.
gpu	If TRUE, the calculations are performed on the GPU. If it is properly set up.

Value

W of class matrix, the anchor transformation matrix.

Author(s)

Markus Ulmer

References

Bühlmann P (2020). “Invariance, Causality and Robustness.” *Statistical Science*, **35**(3). ISSN 0883-4237, doi:10.1214/19STS721.

Rothenhäusler D, Meinshausen N, Bühlmann P, Peters J (2021). “Anchor Regression: Heterogeneous Data Meet Causality.” *Journal of the Royal Statistical Society Series B: Statistical Methodology*, **83**(2), 215–246. ISSN 1369-7412, doi:10.1111/rssb.12398.

Examples

```
set.seed(1)
n <- 50
X <- matrix(rnorm(n * 1), nrow = n)
Y <- 3 * X + rnorm(n)
W <- get_W(X, gamma = 0)
resid <- W %*% Y
```

mergeForest

Merge two forests

Description

This function merges two forests. The trees are combined and the variable importance is calculated as a weighted average of the two forests. If the forests are trained on the same data, the predictions and oob_predictions are combined as well.

Usage

```
mergeForest(fit1, fit2)
```

Arguments

fit1	first SDForest object
fit2	second SDForest object

Value

```
merged SDForest object set.seed(1) n <- 50 X <- matrix(rnorm(n * 5), nrow = n) y <- sign(X[, 1])
* 3 + rnorm(n) fit1 <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', nTree = 5, cp = 0.5)
fit2 <- SDForest(x = X, y = y, nTree = 5, cp = 0.5) mergeForest(fit1, fit2)
```

Author(s)

Markus Ulmer

partDependence	<i>Partial dependence</i>
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Description

This function calculates the partial dependence of a model on a single variable. For that predictions are made for all observations in the dataset while varying the value of the variable of interest. The overall partial effect is the average of all predictions. (Friedman 2001)

Usage

```
partDependence(object, j, X = NULL, subSample = NULL, mc.cores = 1)
```

Arguments

object	A model object that has a predict method that takes newdata as argument and returns predictions.
j	The variable for which the partial dependence should be calculated. Either the column index of the variable in the dataset or the name of the variable.
X	The dataset on which the partial dependence should be calculated. Should contain the same variables as the dataset used to train the model. If NULL, tries to extract the dataset from the model object.
subSample	Number of samples to draw from the original data for the empirical partial dependence. If NULL, all the observations are used.
mc.cores	Number of cores to use for parallel computation. Parallel computing is only supported for unix.

Value

An object of class `partDependence` containing

preds_mean	The average prediction for each value of the variable of interest.
x_seq	The sequence of values for the variable of interest.
preds	The predictions for each value of the variable of interest for each observation.
j	The name of the variable of interest.
xj	The values of the variable of interest in the dataset.

Author(s)

Markus Ulmer

References

Friedman JH (2001). "Greedy Function Approximation: A Gradient Boosting Machine." *The Annals of Statistics*, **29**(5), 1189–1232. ISSN 00905364, <http://www.jstor.org/stable/2699986>.

See Also

[SDForest](#), [SDTree](#)

Examples

```
set.seed(1)
x <- rnorm(100)
y <- sign(x) * 3 + rnorm(100)
model <- SDTree(x = x, y = y, Q_type = 'no_deconfounding')
pd <- partDependence(model, 1, X = x, subSample = 10)
plot(pd)
```

plot.partDependence *Plot partial dependence*

Description

This function plots the partial dependence of a model on a single variable.

Usage

```
## S3 method for class 'partDependence'
plot(x, n_examples = 19, ...)
```

Arguments

x	An object of class partDependence returned by partDependence .
n_examples	Number of examples to plot in addition to the average prediction.
...	Further arguments passed to or from other methods.

Value

A ggplot object.

Author(s)

Markus Ulmer

See Also

[partDependence](#) set.seed(1) x <- rnorm(10) y <- sign(x) * 3 + rnorm(10) model <- SDTree(x = x, y = y, Q_type = 'no_deconfounding', cp = 0.5) pd <- partDependence(model, 1, X = x) plot(pd)

plot.paths	<i>Visualize the paths of an SDTree or SDForest</i>
------------	---

Description

This function visualizes the variable importance of an SDTree or SDForest for different complexity parameters. Both the regularization path and the stability selection path can be visualized.

Usage

```
## S3 method for class 'paths'  
plot(x, plotly = FALSE, selection = NULL, sqrt_scale = FALSE, ...)
```

Arguments

x	A paths object
plotly	If TRUE the plot is returned interactive using plotly. Might be slow for large data.
selection	A vector of indices of the covariates to be plotted. Can be used to plot only a subset of the covariates in case of many covariates.
sqrt_scale	If TRUE the y-axis is on a square root scale.
...	Further arguments passed to or from other methods.

Value

A ggplot object with the variable importance for different regularization. If the path object includes a cp_min value, a black dashed line is added to indicate the out-of-bag optimal variable selection.

Author(s)

Markus Ulmer

See Also

[regPath stabilitySelection](#)

Examples

```
set.seed(1)  
n <- 10  
X <- matrix(rnorm(n * 5), nrow = n)  
y <- sign(X[, 1]) * 3 + sign(X[, 2]) + rnorm(n)  
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)  
paths <- regPath(model)  
plot(paths)
```

```
plot(paths, plotly = TRUE)
```

```
plot.SDTree
```

```
Plot SDTree
```

Description

Plot the SDTree.

Usage

```
## S3 method for class 'SDTree'
plot(x, ...)
```

Arguments

`x` Fitted object of class SDTree.
`...` Further arguments for DiagrammeR::render_graph()

Value

graph object from DiagrammeR::render_graph()

Author(s)

Markus Ulmer

See Also

[SDTree](#) `set.seed(1) n <- 10 X <- matrix(rnorm(n * 5), nrow = n) y <- sign(X[, 1]) * 3 + rnorm(n)`
`model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5) plot(model)`

```
plotOOB
```

```
Visualize the out-of-bag performance of an SDForest
```

Description

This function visualizes the out-of-bag performance of an SDForest for different complexity parameters. Can be used to choose the optimal complexity parameter.

Usage

```
plotOOB(object, sqrt_scale = FALSE)
```

Arguments

`object` A paths object with `loss_path` matrix with the out-of-bag performance for each complexity parameter.

`sqrt_scale` If TRUE the x-axis is on a square root scale.

Value

A ggplot object

Author(s)

Markus Ulmer

See Also

[regPath.SDForest](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + sign(X[, 2]) + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)
paths <- regPath(model)
plotOOB(paths)
```

predict.SDAM

Predictions for SDAM

Description

Predicts the response for new data using a fitted SDAM.

Usage

```
## S3 method for class 'SDAM'
predict(object, newdata, ...)
```

Arguments

`object` Fitted object of class SDAM.

`newdata` New test data of class `data.frame` containing the covariates for which to predict the response.

`...` Further arguments passed to or from other methods.

Value

A vector of predictions for the new data.

Author(s)

Cyrill Scheidegger

See Also

[SDAM](#)

Examples

```
set.seed(1)
X <- matrix(rnorm(10 * 5), ncol = 5)
Y <- sin(X[, 1]) - X[, 2] + rnorm(10)
model <- SDAM(x = X, y = Y, Q_type = "trim", trim_quantile = 0.5, nfold = 2, n_K = 1)
predict(model, newdata = data.frame(X))
```

predict.SDForest

Predictions for the SDForest

Description

Predicts the response for new data using a fitted SDForest.

Usage

```
## S3 method for class 'SDForest'
predict(object, newdata, mc.cores = 1, ...)
```

Arguments

object	Fitted object of class SDForest.
newdata	New test data of class data.frame containing the covariates for which to predict the response.
mc.cores	Number of cores to use for parallel processing, if mc.cores > 1 the trees predict in parallel.
...	Further arguments passed to or from other methods.

Value

A vector of predictions for the new data.

Author(s)

Markus Ulmer

See Also[SDForest](#)**Examples**

```
set.seed(1)
n <- 50
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', nTree = 5, cp = 0.5)
predict(model, newdata = data.frame(X))
```

predict.SDTree

Predictions for the SDTree

Description

Predicts the response for new data using a fitted SDTree.

Usage

```
## S3 method for class 'SDTree'
predict(object, newdata, ...)
```

Arguments

object	Fitted object of class SDTree.
newdata	New test data of class data.frame containing the covariates for which to predict the response.
...	Further arguments passed to or from other methods.

Value

A vector of predictions for the new data.

Author(s)

Markus Ulmer

See Also[SDTree](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)
predict(model, newdata = data.frame(X))
```

predictOOB

Out-of-bag predictions for the SDForest

Description

Predicts the response for the training data using only the trees in the SDForest that were not trained on the observation.

Usage

```
predictOOB(object, X = NULL)
```

Arguments

object	Fitted object of class SDForest.
X	Covariates of the training data. If NULL, the data saved in the object is used.

Value

A vector of out-of-bag predictions for the training data. #' set.seed(1) n <- 50 X <- matrix(rnorm(n * 5), nrow = n) y <- sign(X[, 1]) * 3 + rnorm(n) model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', nTree = 5, cp = 0.5) predictOOB(model)

Author(s)

Markus Ulmer

See Also

[SDForest](#) [prune.SDForest](#) [plotOOB](#)

predict_individual_fj *Predictions of individual component functions for SDAM*

Description

Predicts the contribution of an individual component j using a fitted SDAM.

Usage

```
predict_individual_fj(object, j, x = NULL)
```

Arguments

object	Fitted object of class SDAM.
j	Which component to evaluate.
x	New numeric data to predict for.

Value

A vector of predictions for f_j evaluated at $X_{j\text{new}}$.

Author(s)

Cyrill Scheidegger

See Also

[SDAM](#)

Examples

```
set.seed(1)
X <- matrix(rnorm(10 * 5), ncol = 5)
Y <- sin(X[, 1]) - X[, 2] + rnorm(10)
model <- SDAM(x = X, y = Y, Q_type = "trim", trim_quantile = 0.5, nfold = 2, n_K = 1)
predict_individual_fj(model, j = 1, seq(-2, 2, length.out = 100))
```

`print.partDependence` *Print partDependence*

Description

Print contents of the `partDependence`.

Usage

```
## S3 method for class 'partDependence'  
print(x, ...)
```

Arguments

`x` Fitted object of class `partDependence`.
`...` Further arguments passed to or from other methods.

Value

No return value, called for side effects

Author(s)

Markus Ulmer

See Also

[partDependence](#), [plot.partDependence](#)

Examples

```
set.seed(1)  
x <- rnorm(10)  
y <- sign(x) * 3 + rnorm(10)  
model <- SDTree(x = x, y = y, Q_type = 'no_deconfounding', cp = 0.5)  
pd <- partDependence(model, 1, X = x)  
print(pd)
```

print.SDAM	<i>Print SDAM</i>
------------	-------------------

Description

Print number of covariates and number of active covariates for SDAM object.

Usage

```
## S3 method for class 'SDAM'  
print(x, ...)
```

Arguments

x	Fitted object of class SDAM.
...	Further arguments passed to or from other methods.

Value

No return value, called for side effects

Author(s)

Cyrill Scheidegger

See Also

[SDAM](#)

Examples

```
set.seed(1)  
X <- matrix(rnorm(10 * 5), ncol = 5)  
Y <- sin(X[, 1]) - X[, 2] + rnorm(10)  
model <- SDAM(x = X, y = Y, Q_type = "trim", trim_quantile = 0.5, nfold = 2, n_K = 1)  
print(model)
```

print.SDForest	<i>Print SDForest</i>
----------------	-----------------------

Description

Print contents of the SDForest.

Usage

```
## S3 method for class 'SDForest'  
print(x, ...)
```

Arguments

x	Fitted object of class SDForest.
...	Further arguments passed to or from other methods.

Value

No return value, called for side effects

Author(s)

Markus Ulmer

See Also

[SDForest](#)

Examples

```
set.seed(1)  
n <- 50  
X <- matrix(rnorm(n * 5), nrow = n)  
y <- sign(X[, 1]) * 3 + rnorm(n)  
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', nTree = 5, cp = 0.5)  
print(model)
```

print.SDTree	<i>Print a SDTree</i>
--------------	-----------------------

Description

Print contents of the SDTree.

Usage

```
## S3 method for class 'SDTree'  
print(x, ...)
```

Arguments

x	Fitted object of class SDTree.
...	Further arguments passed to or from other methods.

Value

No return value, called for side effects

Author(s)

Markus Ulmer

See Also

[SDTree](#)

Examples

```
set.seed(1)  
n <- 10  
X <- matrix(rnorm(n * 5), nrow = n)  
y <- sign(X[, 1]) * 3 + rnorm(n)  
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)  
print(model)
```

```
prune.SDForest      Prune an SDForest
```

Description

Prunes all trees in the forest and re-calculates the out-of-bag predictions and performance measures. The training data is needed to calculate the out-of-bag statistics. Note that the forest is pruned in place. If you intend to keep the original forest, make a copy of it before pruning.

Usage

```
## S3 method for class 'SDForest'
prune(object, cp, X = NULL, Y = NULL, Q = NULL, pred = TRUE, ...)
```

Arguments

object	an SDForest object
cp	Complexity parameter, the higher the value the more nodes are pruned.
X	The training data, if NULL the data from the forest object is used.
Y	The training response variable, if NULL the data from the forest object is used.
Q	The transformation function, if NULL the data from the forest object is used.
pred	If TRUE the predictions are calculated, if FALSE only the out-of-bag statistics are calculated. This can set to FALSE to save computation time if only the out-of-bag statistics are needed.
...	Further arguments passed to or from other methods.

Value

A pruned SDForest object

Author(s)

Markus Ulmer

See Also

[copy](#) [prune.SDTree](#) [regPath](#)

Examples

```
set.seed(1)
X <- matrix(rnorm(10 * 20), nrow = 10)
Y <- rnorm(10)
fit <- SDForest(x = X, y = Y, nTree = 2)
pruned_fit <- prune(copy(fit), 0.2)
```

`prune.SDTree`*Prune an SDTree*

Description

Removes all nodes that did not improve the loss by more than `cp` times the initial loss. Either by themselves or by one of their successors. Note that the tree is pruned in place. If you intend to keep the original tree, make a copy of it before pruning.

Usage

```
## S3 method for class 'SDTree'  
prune(object, cp, ...)
```

Arguments

<code>object</code>	an SDTree object
<code>cp</code>	Complexity parameter, the higher the value the more nodes are pruned.
<code>...</code>	Further arguments passed to or from other methods.

Value

A pruned SDTree object

Author(s)

Markus Ulmer

See Also

[copy](#)

Examples

```
set.seed(1)  
X <- matrix(rnorm(10 * 20), nrow = 10)  
Y <- rnorm(10)  
tree <- SDTree(x = X, y = Y)  
pruned_tree <- prune(copy(tree), 0.2)  
tree  
pruned_tree
```

regPath.SDForest *Calculate the regularization path of an SDForest*

Description

This function calculates the variable importance of an SDForest and the out-of-bag performance for different complexity parameters.

Usage

```
## S3 method for class 'SDForest'
regPath(object, cp_seq = NULL, X = NULL, Y = NULL, Q = NULL, copy = TRUE, ...)
```

Arguments

object	an SDForest object
cp_seq	A sequence of complexity parameters. If NULL, the sequence is calculated automatically using only relevant values.
X	The training data, if NULL the data from the forest object is used.
Y	The training response variable, if NULL the data from the forest object is used.
Q	The transformation matrix, if NULL the data from the forest object is used.
copy	Whether the tree should be copied for the regularization path. If FALSE, the pruning is done in place and will change the SDForest. This might be reasonable, if the SDForest is to large to copy.
...	Further arguments passed to or from other methods.

Value

An object of class paths containing

cp	The sequence of complexity parameters.
varImp_path	A matrix with the variable importance for each complexity parameter.
loss_path	A matrix with the out-of-bag performance for each complexity parameter.
cp_min	The complexity parameter with the lowest out-of-bag performance.
type	Path type

Author(s)

Markus Ulmer

See Also

[plot.paths](#) [plotOOB](#) [regPath.SDTree](#) [prune](#) [get_cp_seq](#) [SDForest](#)

Examples

```

set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + sign(X[, 2]) + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)
paths <- regPath(model)
plotOOB(paths)
plot(paths)

plot(paths, plotly = TRUE)

```

regPath.SDTree

Calculate the regularization path of an SDTree

Description

This function calculates the variable importance of an SDTree for different complexity parameters.

Usage

```

## S3 method for class 'SDTree'
regPath(object, cp_seq = NULL, ...)

```

Arguments

object	an SDTree object
cp_seq	A sequence of complexity parameters. If NULL, the sequence is calculated automatically using only relevant values.
...	Further arguments passed to or from other methods.

Value

An object of class paths containing

cp	The sequence of complexity parameters.
varImp_path	A matrix with the variable importance for each complexity parameter.
type	Path type

Author(s)

Markus Ulmer

See Also

[plot.paths](#) [prune](#) [get_cp_seq](#) [SDTree](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + sign(X[, 2]) + rnorm(n)
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)
paths <- regPath(model)
plot(paths)

plot(paths, plotly = TRUE)
```

SDAM

Spectrally Deconfounded Additive Models

Description

Estimate high-dimensional additive models using spectral deconfounding (Scheidegger et al. 2025). The covariates are expanded into B-spline basis functions. A spectral transformation is used to remove bias arising from hidden confounding and a group lasso objective is minimized to enforce component-wise sparsity. Optimal number of basis functions per component and sparsity penalty are chosen by cross validation.

Usage

```
SDAM(
  formula = NULL,
  data = NULL,
  x = NULL,
  y = NULL,
  Q_type = "trim",
  trim_quantile = 0.5,
  q_hat = 0,
  nfolds = 5,
  cv_method = "1se",
  n_K = 4,
  n_lambda1 = 10,
  n_lambda2 = 20,
  Q_scale = TRUE,
  ind_lin = NULL,
  mc.cores = 1,
  verbose = TRUE,
  notRegularized = NULL
)
```


Arguments

formula	Object of class <code>formula</code> or describing the model to fit of the form $y \sim x_1 + x_2 + \dots$ where y is a numeric response and x_1, x_2, \dots are vectors of covariates. Interactions are not supported.
data	Training data of class <code>data.frame</code> containing the variables in the model.
x	Matrix of covariates, alternative to <code>formula</code> and <code>data</code> .
y	Vector of responses, alternative to <code>formula</code> and <code>data</code> .
Q_type	Type of deconfounding, one of <code>'trim'</code> , <code>'pca'</code> , <code>'no_deconfounding'</code> . <code>'trim'</code> corresponds to the Trim transform (Ćevič et al. 2020) as implemented in the Doubly debiased lasso (Guo et al. 2022), <code>'pca'</code> to the PCA transformation (Paul et al. 2008). See get_Q .
trim_quantile	Quantile for Trim transform, only needed for <code>trim</code> , see get_Q .
q_hat	Assumed confounding dimension, only needed for <code>pca</code> , see get_Q .
nfolds	The number of folds for cross-validation. Default is 5.
cv_method	The method for selecting the regularization parameter during cross-validation. One of <code>"min"</code> (minimum cv-loss) and <code>"1se"</code> (one-standard-error rule) Default is <code>"1se"</code> .
n_K	The number of candidate values for the number of basis functions for B-splines. Default is 4.
n_lambda1	The number of candidate values for the regularization parameter in the initial cross-validation step. Default is 10.
n_lambda2	The number of candidate values for the regularization parameter in the second stage of cross-validation (once the optimal number of basis function K is decided, a second stage of cross-validation for the regularization parameter is performed on a finer grid). Default is 20.
Q_scale	Should data be scaled to estimate the spectral transformation? Default is <code>TRUE</code> to not reduce the signal of high variance covariates.
ind_lin	A vector of indices specifying which covariates to model linearly (i.e. not expanded into basis function). Default is <code>'NULL'</code> .
mc.cores	Number of cores to use for parallel processing, if <code>mc.cores > 1</code> the cross validation is parallelized. Default is <code>'1'</code> . (only supported for unix)
verbose	If <code>TRUE</code> fitting information is shown.
notRegularized	A vector of indices specifying which covariates not to regularize. Default is <code>'NULL'</code> .

Value

An object of class `'SDAM'` containing the following elements:

X	The original design matrix.
p	The number of covariates in <code>'X'</code> .
var_names	Names of the covariates in the training data.
intercept	The intercept term of the fitted model.

K	A vector of the number of basis functions for each covariate, where 1 corresponds to a linear term. The entries of the vector will mostly be the same, but some entries might be lower if the corresponding component of X contains only few unique values.
breaks	A list of breakpoints used for the B-splines. Used to reconstruct the B-spline basis functions.
coefs	A list of coefficients for the B-spline basis functions for each component.
active	A vector of active covariates that contribute to the model.

Author(s)

Cyrill Scheidegger

References

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See Also

[get_Q](#), [predict.SDAM](#), [varImp.SDAM](#), [predict_individual_fj](#), [partDependence](#)

Examples

```
set.seed(1)
X <- matrix(rnorm(10 * 5), ncol = 5)
Y <- sin(X[, 1]) - X[, 2] + rnorm(10)
model <- SDAM(x = X, y = Y, Q_type = "trim", trim_quantile = 0.5, nfold = 2, n_K = 1)

# if we know that the first covariate one is relevant, we can also choose to not regularize it
model <- SDAM(x = X, y = Y, Q_type = "trim", trim_quantile = 0.5, nfold = 2,
              n_K = 1, notRegularized = c(1))

library(HDclassif)
data(wine)
names(wine) <- c("class", "alcohol", "malicAcid", "ash", "alcalinityAsh", "magnesium",
                "totPhenols", "flavanoids", "nonFlavPhenols", "proanthocyanins",
                "colIntens", "hue", "OD", "proline")
```

```

wine <- log(wine)

# estimate model
# do not use class in the model and restrict proline to be linear
model <- SDAM(alcohol ~ ., wine, ind_lin = "proline", nfold = 3)

# extract variable importance
varImp(model)

# most important variable
mostImp <- names(which.max(varImp(model)))
mostImp

# predict for individual Xj
x <- seq(min(wine[, mostImp]), max(wine[, mostImp]), length.out = 100)
predJ <- predict_individual_fj(object = model, j = mostImp, x = x)

plot(x, predJ,
      xlab = paste0("log ", mostImp), ylab = "log alcohol")

# partial dependence
plot(partDependence(model, mostImp))

# predict
predict(model, newdata = wine[42, ])

## alternative function call
mod_none <- SDAM(x = as.matrix(wine[1:10, -c(1, 2)]), y = wine$alcohol[1:10],
                 Q_type = "no_deconfounding", nolds = 2, n_K = 4,
                 n_lambda1 = 4, n_lambda2 = 8)

```

SDForest

*Spectrally Deconfounded Random Forests***Description**

Estimate regression Random Forest using spectral deconfounding. The spectrally deconfounded Random Forest (SDForest) combines SDTrees in the same way, as in the original Random Forest (Breiman 2001). The idea is to combine multiple regression trees into an ensemble in order to decrease variance and get a smooth function. Ensembles work best if the different models are independent of each other. To decorrelate the regression trees as much as possible from each other, we have two mechanisms. The first one is bagging (Breiman 1996), where we train each regression tree on an independent bootstrap sample of the observations, e.g., we draw a random sample of size n with replacement from the observations. The second mechanic to decrease the correlation is that only a random subset of the covariates is available for each split. Before each split, we sample m covariates from all the covariates and choose the one that reduces the loss the most only from those.

$$\widehat{f(X)} = \frac{1}{N_{tree}} \sum_{t=1}^{N_{tree}} SDTree_t(X)$$

Usage

```

SDForest(
  formula = NULL,
  data = NULL,
  x = NULL,
  y = NULL,
  nTree = 100,
  cp = 0,
  min_sample = 5,
  mtry = NULL,
  mc.cores = 1,
  Q_type = "trim",
  trim_quantile = 0.5,
  q_hat = 0,
  Qf = NULL,
  A = NULL,
  gamma = 7,
  max_size = NULL,
  gpu = FALSE,
  return_data = TRUE,
  mem_size = 1e+07,
  leave_out_ind = NULL,
  envs = NULL,
  nTree_leave_out = NULL,
  nTree_env = NULL,
  max_candidates = 100,
  Q_scale = TRUE,
  verbose = TRUE,
  predictors = NULL
)

```

Arguments

formula	Object of class formula or describing the model to fit of the form $y \sim x_1 + x_2 + \dots$ where y is a numeric response and x_1, x_2, \dots are vectors of covariates. Interactions are not supported.
data	Training data of class data.frame containing the variables in the model.
x	Matrix of covariates, alternative to formula and data.
y	Vector of responses, alternative to formula and data.
nTree	Number of trees to grow.
cp	Complexity parameter, minimum loss decrease to split a node. A split is only performed if the loss decrease is larger than $cp * \text{initial_loss}$, where initial_loss is the loss of the initial estimate using only a stump.
min_sample	Minimum number of observations per leaf. A split is only performed if both resulting leaves have at least min_sample observations.
mtry	Number of randomly selected covariates to consider for a split, if NULL half of the covariates are available for each split. $mtry = \lfloor \frac{p}{2} \rfloor$

<code>mc.cores</code>	Number of cores to use for parallel processing, if <code>mc.cores > 1</code> the trees are estimated in parallel.
<code>Q_type</code>	Type of deconfounding, one of <code>'trim'</code> , <code>'pca'</code> , <code>'no_deconfounding'</code> . <code>'trim'</code> corresponds to the Trim transform (Ćevič et al. 2020) as implemented in the Doubly debiased lasso (Guo et al. 2022), <code>'pca'</code> to the PCA transformation (Paul et al. 2008). See get_Q .
<code>trim_quantile</code>	Quantile for Trim transform, only needed for trim, see get_Q .
<code>q_hat</code>	Assumed confounding dimension, only needed for pca, see get_Q .
<code>Qf</code>	Spectral transformation, if NULL it is internally estimated using get_Q .
<code>A</code>	Numerical Anchor of class matrix. See get_W .
<code>gamma</code>	Strength of distributional robustness, $\gamma \in [0, \infty]$. See get_W .
<code>max_size</code>	Maximum number of observations used for a bootstrap sample. If NULL <code>n</code> samples with replacement are drawn.
<code>gpu</code>	If TRUE, the calculations are performed on the GPU. If it is properly set up.
<code>return_data</code>	If TRUE, the training data is returned in the output. This is needed for prune.SDForest , regPath.SDForest , and for mergeForest .
<code>mem_size</code>	Amount of split candidates that can be evaluated at once. This is a trade-off between memory and speed can be decreased if either the memory is not sufficient or the <code>gpu</code> is too small.
<code>leave_out_ind</code>	Indices of observations that should not be used for training.
<code>envs</code>	Vector of environments of class <code>factor</code> which can be used for stratified tree fitting.
<code>nTree_leave_out</code>	Number of trees that should be estimated while leaving one of the environments out. Results in number of environments times number of trees.
<code>nTree_env</code>	Number of trees that should be estimated for each environment. Results in number of environments times number of trees.
<code>max_candidates</code>	Maximum number of split points that are proposed at each node for each covariate.
<code>Q_scale</code>	Should data be scaled to estimate the spectral transformation? Default is TRUE to not reduce the signal of high variance covariates, and we do not know of a scenario where this hurts.
<code>verbose</code>	If TRUE fitting information is shown.
<code>predictors</code>	Subset of <code>colnames(X)</code> or numerical indices of the covariates for which an effect on <code>y</code> should be estimated. All the other covariates are only used for deconfounding.

Value

Object of class `SDForest` containing:

<code>predictions</code>	Vector of predictions for each observation.
<code>forest</code>	List of <code>SDTree</code> objects.

var_names	Names of the covariates.
oob_loss	Out-of-bag loss. MSE
oob_SDloss	Out-of-bag loss using the spectral transformation.
var_importance	Variable importance. The variable importance is calculated as the sum of the decrease in the loss function resulting from all splits that use a covariate for each tree. The mean of the variable importance of all trees results in the variable importance for the forest.
oob_ind	List of indices of trees that did not contain the observation in the training set.
oob_predictions	Out-of-bag predictions.

If return_data is TRUE the following are also returned:

X	Matrix of covariates.
Y	Vector of responses.
Q	Spectral transformation.

If envs is provided the following are also returned:

envs	Vector of environments.
nTree_env	Number of trees for each environment.
ooEnv_ind	List of indices of trees that did not contain the observation or the same environment in the training set for each observation.
ooEnv_loss	Out-of-bag loss using only trees that did not contain the observation or the same environment.
ooEnv_SDloss	Out-of-bag loss using the spectral transformation and only trees that did not contain the observation or the same environment.
ooEnv_predictions	Out-of-bag predictions using only trees that did not contain the observation or the same environment.
nTree_leave_out	If environments are left out, the environment for each tree, that was left out.
nTree_env	If environments are provided, the environment each tree is trained with.

Author(s)

Markus Ulmer

References

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See Also

[get_Q](#), [get_W](#), [SDTree](#), [simulate_data_nonlinear](#), [regPath](#), [stabilitySelection](#), [prune](#), [partDependence](#)

Examples

```
set.seed(1)
n <- 50
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', nTree = 5, cp = 0.5)
predict(model, newdata = data.frame(X))

##### subset of predictors
# if we know, that only the first covariate has an effect on y,
# we can estimate only its effect and use the others just for deconfounding
model <- SDForest(x = X, y = y, cp = 0.5, nTree = 5, predictors = c(1))

set.seed(42)
# simulation of confounded data
sim_data <- simulate_data_nonlinear(q = 2, p = 150, n = 100, m = 2)
X <- sim_data$X
Y <- sim_data$Y
train_data <- data.frame(X, Y)
# causal parents of y
sim_data$j

# comparison to classical random forest
fit_ranger <- ranger::ranger(Y ~ ., train_data, importance = 'impurity')

fit <- SDForest(x = X, y = Y, nTree = 10, Q_type = 'pca', q_hat = 2)
fit <- SDForest(Y ~ ., nTree = 10, train_data)
fit

# comparison of variable importance
imp_ranger <- fit_ranger$variable.importance
imp_sdf <- fit$var_importance
imp_col <- rep('black', length(imp_ranger))
imp_col[sim_data$j] <- 'red'

plot(imp_ranger, imp_sdf, col = imp_col, pch = 20,
```

```

xlab = 'ranger', ylab = 'SDForest',
main = 'Variable Importance')

# check regularization path of variable importance
path <- regPath(fit)
# out of bag error for different regularization
plotOOB(path)
plot(path)

# detection of causal parent using stability selection
stablePath <- stabilitySelection(fit)
plot(stablePath)

# pruning of forest according to optimal out-of-bag performance
fit <- prune(fit, cp = path$cp_min)

# partial functional dependence of y on the most important covariate
most_imp <- which.max(fit$var_importance)
dep <- partDependence(fit, most_imp)
plot(dep, n_examples = 100)

```

SDTree

Spectrally Deconfounded Tree

Description

Estimates a regression tree using spectral deconfounding. A regression tree is part of the function class of step functions $f(X) = \sum_{m=1}^M 1_{\{X \in R_m\}} c_m$, where (R_m) with $m = 1, \dots, M$ are regions dividing the space of \mathbb{R}^p into M rectangular parts. Each region has response level $c_m \in \mathbb{R}$. For the training data, we can write the step function as $f(\mathbf{X}) = \mathcal{P}c$ where $\mathcal{P} \in \{0, 1\}^{n \times M}$ is an indicator matrix encoding to which region an observation belongs and $c \in \mathbb{R}^M$ is a vector containing the levels corresponding to the different regions. This function then minimizes

$$(\hat{\mathcal{P}}, \hat{c}) = \operatorname{argmin}_{\mathcal{P}' \in \{0, 1\}^{n \times M}, c' \in \mathbb{R}^M} \frac{\|Q(\mathbf{Y} - \mathcal{P}'c')\|_2^2}{n}$$

We find $\hat{\mathcal{P}}$ by using the tree structure and repeated splitting of the leaves, similar to the original cart algorithm (Breiman et al. 2017). Since comparing all possibilities for \mathcal{P} is impossible, we let a tree grow greedily. Given the current tree, we iterate over all leaves and all possible splits. We choose the one that reduces the spectral loss the most and estimate after each split all the leaf estimates $\hat{c} = \operatorname{argmin}_{c' \in \mathbb{R}^M} \frac{\|Q\mathbf{Y} - Q\mathcal{P}'c'\|_2^2}{n}$ which is just a linear regression problem. This is repeated until the loss decreases less than a minimum loss decrease after a split. The minimum loss decrease equals a cost-complexity parameter cp times the initial loss when only an overall mean is estimated. The cost-complexity parameter cp controls the complexity of a regression tree and acts as a regularization parameter.

Usage

```
SDTree(
  formula = NULL,
  data = NULL,
  x = NULL,
  y = NULL,
  max_leaves = NULL,
  cp = 0.01,
  min_sample = 5,
  mtry = NULL,
  fast = TRUE,
  Q_type = "trim",
  trim_quantile = 0.5,
  q_hat = 0,
  Qf = NULL,
  A = NULL,
  gamma = 0.5,
  gpu = FALSE,
  mem_size = 1e+07,
  max_candidates = 100,
  Q_scale = TRUE,
  predictors = NULL
)
```

Arguments

formula	Object of class formula or describing the model to fit of the form $y \sim x_1 + x_2 + \dots$ where y is a numeric response and x_1, x_2, \dots are vectors of covariates. Interactions are not supported.
data	Training data of class data.frame containing the variables in the model.
x	Matrix of covariates, alternative to formula and data.
y	Vector of responses, alternative to formula and data.
max_leaves	Maximum number of leaves for the grown tree.
cp	Complexity parameter, minimum loss decrease to split a node. A split is only performed if the loss decrease is larger than $cp * initial_loss$, where $initial_loss$ is the loss of the initial estimate using only a stump.
min_sample	Minimum number of observations per leaf. A split is only performed if both resulting leaves have at least <code>min_sample</code> observations.
mtry	Number of randomly selected covariates to consider for a split, if NULL all covariates are available for each split.
fast	If TRUE, only the optimal splits in the new leaves are evaluated and the previously optimal splits and their potential loss-decrease are reused. If FALSE all possible splits in all the leaves are reevaluated after every split.
Q_type	Type of deconfounding, one of 'trim', 'pca', 'no_deconfounding'. 'trim' corresponds to the Trim transform (Čevič et al. 2020) as implemented in the Doubly

	debiased lasso (Guo et al. 2022), 'pca' to the PCA transformation (Paul et al. 2008). See <code>get_Q</code> .
<code>trim_quantile</code>	Quantile for Trim transform, only needed for trim, see <code>get_Q</code> .
<code>q_hat</code>	Assumed confounding dimension, only needed for pca, see <code>get_Q</code> .
<code>Qf</code>	Spectral transformation, if NULL it is internally estimated using <code>get_Q</code> .
<code>A</code>	Numerical Anchor of class matrix. See <code>get_W</code> .
<code>gamma</code>	Strength of distributional robustness, $\gamma \in [0, \infty]$. See <code>get_W</code> .
<code>gpu</code>	If TRUE, the calculations are performed on the GPU. If it is properly set up.
<code>mem_size</code>	Amount of split candidates that can be evaluated at once. This is a trade-off between memory and speed can be decreased if either the memory is not sufficient or the gpu is too small.
<code>max_candidates</code>	Maximum number of split points that are proposed at each node for each covariate.
<code>Q_scale</code>	Should data be scaled to estimate the spectral transformation? Default is TRUE to not reduce the signal of high variance covariates, and we do not know of a scenario where this hurts.
<code>predictors</code>	Subset of <code>colnames(X)</code> or numerical indices of the covariates for which an effect on <code>y</code> should be estimated. All the other covariates are only used for deconfounding.

Value

Object of class `SDTree` containing

<code>predictions</code>	Predictions for the training set.
<code>tree</code>	The estimated tree of class <code>Node</code> from (Glur 2023). The tree contains the information about all the splits and the resulting estimates.
<code>var_names</code>	Names of the covariates in the training data.
<code>var_importance</code>	Variable importance of the covariates. The variable importance is calculated as the sum of the decrease in the loss function resulting from all splits that use this covariate.

Author(s)

Markus Ulmer

References

- Breiman L, Friedman JH, Olshen RA, Stone CJ (2017). *Classification And Regression Trees*. Routledge. ISBN 9781315139470, doi:10.1201/9781315139470.
- Glur C (2023). "data.tree: General Purpose Hierarchical Data Structure." <https://CRAN.R-project.org/package=data.tree>.
- Guo Z, Cévid D, Bühlmann P (2022). "Doubly debiased lasso: High-dimensional inference under

hidden confounding.” *The Annals of Statistics*, **50**(3). ISSN 0090-5364, doi:10.1214/21AOS2152.

Paul D, Bair E, Hastie T, Tibshirani R (2008). ““Preconditioning” for feature selection and regression in high-dimensional problems.” *The Annals of Statistics*, **36**(4). ISSN 0090-5364, doi:10.1214/009053607000000578.

Ćevid D, Bühlmann P, Meinshausen N (2020). “Spectral Deconfounding via Perturbed Sparse Linear Models.” *J. Mach. Learn. Res.*, **21**(1). ISSN 1532-4435, <http://jmlr.org/papers/v21/19-545.html>.

See Also

[simulate_data_nonlinear](#), [regPath.SDTree](#), [prune.SDTree](#), [partDependence](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDTree(x = X, y = y, cp = 0.5)

##### subset of predictors
# if we know, that only the first covariate has an effect on y,
# we can estimate only its effect and use the others just for deconfounding
model <- SDTree(x = X, y = y, cp = 0.5, predictors = c(1))

set.seed(42)
# simulation of confounded data
sim_data <- simulate_data_step(q = 2, p = 15, n = 100, m = 2)
X <- sim_data$X
Y <- sim_data$Y
train_data <- data.frame(X, Y)
# causal parents of y
sim_data$j

tree_plain_cv <- cvSDTree(Y ~ ., train_data, Q_type = "no_deconfounding")
tree_plain <- SDTree(Y ~ ., train_data, Q_type = "no_deconfounding", cp = 0)

tree_causal_cv <- cvSDTree(Y ~ ., train_data)
tree_causal <- SDTree(y = Y, x = X, cp = 0)

# check regularization path of variable importance
path <- regPath(tree_causal)
plot(path)

tree_plain <- prune(tree_plain, cp = tree_plain_cv$cp_min)
tree_causal <- prune(tree_causal, cp = tree_causal_cv$cp_min)
plot(tree_causal)
plot(tree_plain)
```

 simulate_data_nonlinear

Simulate data with linear confounding and non-linear causal effect

Description

Simulation of data from a confounded non-linear model. The data generating process is given by:

$$Y = f(X) + \delta^T H + \nu$$

$$X = \Gamma^T H + E$$

where $f(X)$ is a random function on the fourier basis with a subset of size m covariates X_j having a causal effect on Y .

$$f(x_i) = \sum_{j=1}^p 1_{j \in j_s} \sum_{k=1}^K (\beta_{j,k}^{(1)} \cos(0.2kx_j) + \beta_{j,k}^{(2)} \sin(0.2kx_j))$$

E, ν are random error terms and $H \in \mathbb{R}^{n \times q}$ is a matrix of random confounding covariates. $\Gamma \in \mathbb{R}^{q \times p}$ and $\delta \in \mathbb{R}^q$ are random coefficient vectors. For the simulation, all the above parameters are drawn from a standard normal distribution, except for ν which is drawn from a normal distribution with standard deviation 0.1. The parameters β are drawn from a uniform distribution between -1 and 1.

Usage

```
simulate_data_nonlinear(q, p, n, m, K = 2, eff = NULL, fixEff = FALSE)
```

Arguments

q	number of confounding covariates in H
p	number of covariates in X
n	number of observations
m	number of covariates with a causal effect on Y
K	number of fourier basis functions $K \in \mathbb{N}$, e.g. complexity of causal function
eff	the number of affected covariates in X by the confounding, if NULL all covariates are affected
fixEff	if eff is smaller than p: If fixEff = TRUE, the causal parents are always affected by confounding if fixEff = FALSE, affected covariates are chosen completely at random.

Value

a list containing the simulated data:

X	a matrix of covariates
Y	a vector of responses
f_X	a vector of the true function $f(X)$
j	the indices of the causal covariates in X
beta	the parameter vector for the function $f(X)$, see f_four
H	the matrix of confounding covariates

Author(s)

Markus Ulmer

See Also

[f_four](#)

Examples

```
set.seed(42)
# simulation of confounded data
sim_data <- simulate_data_nonlinear(q = 2, p = 150, n = 100, m = 2)
X <- sim_data$X
Y <- sim_data$Y
```

simulate_data_step	<i>Simulate data with linear confounding and causal effect following a step-function</i>
--------------------	--

Description

Simulation of data from a confounded non-linear model. Where the non-linear function is a random regression tree. The data generating process is given by:

$$Y = f(X) + \delta^T H + \nu$$

$$X = \Gamma^T H + E$$

where $f(X)$ is a random regression tree with m random splits of the data. Resulting in a random step-function with $m + 1$ levels, i.e. leaf-levels.

$$f(x_i) = \sum_{k=1}^K 1_{\{x_i \in R_k\}} c_k$$

E, ν are random error terms and $H \in \mathbb{R}^{n \times q}$ is a matrix of random confounding covariates. $\Gamma \in \mathbb{R}^{q \times p}$ and $\delta \in \mathbb{R}^q$ are random coefficient vectors. For the simulation, all the above parameters are drawn from a standard normal distribution, except for δ which is drawn from a normal distribution with standard deviation 10. The leaf levels c_k are drawn from a uniform distribution between -50 and 50.

Usage

```
simulate_data_step(q, p, n, m, make_tree = FALSE)
```

Arguments

q	number of confounding covariates in H
p	number of covariates in X
n	number of observations
m	number of covariates with a causal effect on Y
make_tree	Whether the random regression tree should be returned.

Value

a list containing the simulated data:

X	a matrix of covariates
Y	a vector of responses
f_X	a vector of the true function $f(X)$
j	the indices of the causal covariates in X
tree	If <code>make_tree</code> , the random regression tree of class <code>Node</code> from (Glur 2023)

Author(s)

Markus Ulmer

References

Glur C (2023). “data.tree: General Purpose Hierarchical Data Structure.” <https://CRAN.R-project.org/package=data.tree>.

See Also

[simulate_data_nonlinear](#)

Examples

```
set.seed(42)
# simulation of confounded data
sim_data <- simulate_data_step(q = 2, p = 15, n = 100, m = 2)
X <- sim_data$X
Y <- sim_data$Y
```

`stabilitySelection.SDForest`*Calculate the stability selection of an SDForest*

Description

This function calculates the stability selection of an SDForest (Meinshausen and Bühlmann 2010). Stability selection is calculated as the fraction of trees in the forest that select a variable for a split at each complexity parameter.

Usage

```
## S3 method for class 'SDForest'  
stabilitySelection(object, cp_seq = NULL, ...)
```

Arguments

<code>object</code>	an SDForest object
<code>cp_seq</code>	A sequence of complexity parameters. If NULL, the sequence is calculated automatically using only relevant values.
<code>...</code>	Further arguments passed to or from other methods.

Value

An object of class `paths` containing

<code>cp</code>	The sequence of complexity parameters.
<code>varImp_path</code>	A matrix with the stability selection for each complexity parameter.
<code>type</code>	Path type

Author(s)

Markus Ulmer

References

Meinshausen N, Bühlmann P (2010). “Stability Selection.” *Journal of the Royal Statistical Society Series B: Statistical Methodology*, **72**(4), 417–473. ISSN 1369-7412, doi:[10.1111/j.1467-9868.2010.00740.x](https://doi.org/10.1111/j.1467-9868.2010.00740.x).

See Also

[plot.paths](#) [regPath](#) [prune](#) [get_cp_seq](#) [SDForest](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + sign(X[, 2]) + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', nTree = 2, cp = 0.5)
paths <- stabilitySelection(model)
plot(paths)

plot(paths, plotly = TRUE)
```

toList.SDForest

SDForest toList method

Description

Converts the trees in an SDForest object from class Node (Glur 2023) to class list. This makes it substantially easier to save the forest to disk.

Usage

```
## S3 method for class 'SDForest'
toList(object, ...)
```

Arguments

object an SDForest object with the trees in Node format
... Further arguments passed to or from other methods.

Value

an SDForest object with the trees in list format

Author(s)

Markus Ulmer

References

Glur C (2023). “data.tree: General Purpose Hierarchical Data Structure.” <https://CRAN.R-project.org/package=data.tree>.

See Also

[fromList toList.SDTree](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDForest(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5, nTree = 2)
toList(model)
```

toList.SDTree	<i>SDTree toList method</i>
---------------	-----------------------------

Description

Converts the tree in an SDTree object from class Node (Glur 2023) to class list. This makes it substantially easier to save the tree to disk.

Usage

```
## S3 method for class 'SDTree'
toList(object, ...)
```

Arguments

object	an SDTree object with the tree in Node format
...	Further arguments passed to or from other methods.

Value

an SDTree object with the tree in list format

Author(s)

Markus Ulmer

References

Glur C (2023). “data.tree: General Purpose Hierarchical Data Structure.” <https://CRAN.R-project.org/package=data.tree>.

See Also

[fromList](#)

Examples

```
set.seed(1)
n <- 10
X <- matrix(rnorm(n * 5), nrow = n)
y <- sign(X[, 1]) * 3 + rnorm(n)
model <- SDTree(x = X, y = y, Q_type = 'no_deconfounding', cp = 0.5)
toList(model)
```

varImp.SDAM

Extract Variable importance for SDAM

Description

This function extracts the variable importance of an SDAM. The variable importance is calculated as the empirical squared L2 norm of fj. The measure is not standardized.

Usage

```
## S3 method for class 'SDAM'
varImp(object)
```

Arguments

object an SDAM object

Value

A vector of variable importance

Author(s)

Cyrill Scheidegger

See Also

[SDAM](#)

Examples

```
set.seed(1)
X <- matrix(rnorm(10 * 5), ncol = 5)
Y <- sin(X[, 1]) - X[, 2] + rnorm(10)
model <- SDAM(x = X, y = Y, Q_type = "trim", trim_quantile = 0.5, nfold = 2)
varImp(model)
```

varImp.SDForest	<i>Extract variable importance of an SDForest</i>
-----------------	---

Description

This function extracts the variable importance of an SDForest. The variable importance is calculated as the sum of the decrease in the loss function resulting from all splits that use a covariate for each tree. The mean of the variable importance of all trees results in the variable importance for the forest.

Usage

```
## S3 method for class 'SDForest'  
varImp(object)
```

Arguments

object an SDForest object

Value

A named vector of variable importance

Author(s)

Markus Ulmer

See Also

[varImp.SDTree SDForest](#)

Examples

```
data(iris)  
fit <- SDForest(Sepal.Length ~ Sepal.Width + Petal.Length + Petal.Width,  
               iris, nTree = 10, cp = 0.5)  
varImp(fit)
```

varImp.SDTree	<i>Extract variable importance of an SDTree</i>
---------------	---

Description

This function extracts the variable importance of an SDTree. The variable importance is calculated as the sum of the decrease in the loss function resulting from all splits that use this covariate.

Usage

```
## S3 method for class 'SDTree'  
varImp(object)
```

Arguments

object an SDTree object

Value

A named vector of variable importance

Author(s)

Markus Ulmer

See Also

[varImp.SDForest](#) [SDTree](#)

Examples

```
data(iris)  
tree <- SDTree(Sepal.Length ~ Sepal.Width + Petal.Length + Petal.Width, iris, cp = 0.5)  
varImp(tree)
```

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