# Package 'bartcs'

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Title Bayesian Additive Regression Trees for Confounder Selection
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<b>Description</b> Fit Bayesian Regression Additive Trees (BART) models to select true confounders from a large set of potential confounders and to estimate average treatment effect. For more information, see Kim et al. (2023) <doi:10.1111 biom.13833="">.</doi:10.1111>
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bartcs-package

bartcs: Bayesian Additive Regression Trees for Confounder Selection

## **Description**

Fit Bayesian Regression Additive Trees (BART) models to select true confounders from a large set of potential confounders and to estimate average treatment effect. For more information, see Kim et al. (2023) doi:10.1111/biom.13833.

# **Details**

Functions in bartcs serve one of three purposes.

- 1. Functions for fitting: separate\_bart() and single\_bart().
- 2. Functions for summary: summary() and plot().
- 3. Utility function for OpenMP: count\_omp\_thread().

The code of BART model are based on the 'BART' package by Sparapani et al. (2021) under the GPL license, with modifications. The modifications from the BART package include (but are not limited to):

- · Add CHANGE step.
- · Add Single and Separate Model.
- · Add causal effect estimation.
- Add confounder selection.

# References

Sparapani R, Spanbauer C, McCulloch R (2021). "Nonparametric Machine Learning and Efficient Computation with Bayesian Additive Regression Trees: The BART R Package." *Journal of Statistical Software*, 97(1), 1–66. doi:10.18637/jss.v097.i01

Kim, C., Tec, M., & Zigler, C. M. (2023). Bayesian Nonparametric Adjustment of Confounding, *Biometrics* doi:10.1111/biom.13833

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bart

Fit BART models to select confounders and estimate treatment effect

# Description

Fit Bayesian Regression Additive Trees (BART) models to select relevant confounders among a large set of potential confounders and to estimate average treatment effect E[Y(1) - Y(0)].

# Usage

```
separate_bart(
 Y, trt, X,
                  = 1,
  trt_treated
  trt_control
                  = 0,
                  = 50,
 num_tree
                  = 4,
 num_chain
 num_burn_in
                  = 100,
 num_thin
                  = 1,
 num_post_sample = 100,
                 = c(0.28, 0.28, 0.44),
  step_prob
                  = 0.95,
 alpha
 beta
                  = 2,
 nu
                  = 3,
                  = 0.95,
                  = 5,
 dir_alpha
 parallel
                  = FALSE,
                  = TRUE
 verbose
)
single_bart(
 Y, trt, X,
  trt_treated
                  = 1,
  trt_control
                  = 0,
                  = 50,
 num_tree
                  = 4,
 num_chain
                  = 100,
 num_burn_in
 num_thin
                  = 1,
 num_post_sample = 100,
                  = c(0.28, 0.28, 0.44),
 step_prob
                  = 0.95,
 alpha
 beta
                  = 2,
                  = 3,
 nu
                  = 0.95,
 dir_alpha
                  = 5,
 parallel
                  = FALSE,
  verbose
                  = TRUE
)
```

## **Arguments**

Υ

A vector of outcome values.

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trt A vector of treatment values. Binary treatment works for both model and continuous treatment works for single bart(). For binary treatment, use 1 to indicate the treated group and 0 for the control group. Χ A matrix of potential confounders. trt\_treated Value of trt for the treated group. The default value is set to 1. trt\_control Value of trt for the control group. The default value is set to 0. Number of trees in BART model. The default value is set to 100. num\_tree Number of MCMC chains. Need to set num\_chain > 1 for the Gelman-Rubin num\_chain diagnostic. The default value is set to 4. num\_burn\_in Number of MCMC samples to be discarded per chain as initial burn-in periods. The default value is set to 100. Number of thinning per chain. One in every num\_thin samples are selected. num\_thin The default value is set to 1. num\_post\_sample Final number of posterior samples per chain. Number of MCMC iterations per chain is burn\_in + num\_thin \* num\_post\_sample. The default value is set to 100. A vector of tree alteration probabilities (GROW, PRUNE, CHANGE). Each alstep\_prob teration is proposed to change the tree structure. The default setting is (0.28, 0.28, 0.44). Hyperparameters for tree regularization prior. A terminal node of depth d will alpha, beta split with probability of alpha  $* (1 + d)^{-beta}$ . The default setting is (alpha, beta) = (0.95, 2) from Chipman et al. (2010). Values to calibrate hyperparameter of sigma prior. nu, q The default setting is (nu, q) = (3, 0.95) from Chipman et al. (2010). Hyperparameter of Dirichlet prior for selection probabilities. The default value dir\_alpha is 5. parallel If TRUE, model fitting will be parallelized with respect to N = nrow(X). Parallelization is recommended for very high n only. The default setting is FALSE. verbose If TRUE, message will be printed during training. If FALSE, message will be suppressed.

## **Details**

separate\_bart() and single\_bart() fit an exposure model and outcome model(s) for estimating treatment effect with adjustment of confounders in the presence of a large set of potential confounders (Kim et al. 2023).

The exposure model E[A|X] and the outcome model(s) E[Y|A,X] are linked together with a common Dirichlet prior that accrues posterior selection probabilities to the corresponding confounders (X) on the basis of association with both the exposure (A) and the outcome (Y).

There is a distinction between fitting separate outcome models for the treated and control groups and fitting a single outcome model for both groups.

- separate\_bart() specifies two "separate" outcome models for two binary treatment levels. Thus, it fits three models: one exposure model and two separate outcome models for A=0,1.
- single\_bart() specifies one "single" outcome model. Thus, it fits two models: one exposure model and one outcome model for the entire sample.

All inferences are made with outcome model(s).

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#### Value

A bartcs object. A list object contains the following components.

mcmc\_outcome A mcmc.list object from **coda** package. mcmc\_outcome contains the following items.

- ATE Posterior sample of average treatment effect E[Y(1) Y(0)].
- Y1 Posterior sample of potential outcome E[Y(1)].
- Y0 Posterior sample of potential outcome E[Y(0)].

mcmc\_param A mcmc.list object from **coda** package. mcmc\_param contains the following items.

- dir\_alpha Posterior sample of dir\_alpha.
- sigma2\_out Posterior sample of sigma2 in the outcome model.

var\_prob Aggregated posterior inclusion probability of each variable.
var\_count Number of selection of each variable in each MCMC iteration. Its dimension is num\_post\_sample \* ncol(X).
chains A list of results from each MCMC chain.
model separate or single.

model separate or single Column names of X.

params Parameters used in the model.

## References

Chipman, H. A., George, E. I., & McCulloch, R. E. (2010). BART: Bayesian additive regression trees. *The Annals of Applied Statistics*, 4(1), 266-298. doi:10.1214/09AOAS285

Kim, C., Tec, M., & Zigler, C. M. (2023). Bayesian Nonparametric Adjustment of Confounding, *Biometrics* doi:10.1111/biom.13833

## **Examples**

```
data(ihdp, package = "bartcs")
single_bart(
 Υ
                 = ihdp$y_factual,
 trt
                 = ihdp$treatment,
                = ihdp[, 6:30],
 Χ
                = 10,
 num_tree
                = 2,
 num chain
 num_post_sample = 20,
 num_burn_in = 10,
 verbose
                 = FALSE
separate_bart(
 Υ
                 = ihdp$y_factual,
 trt
                 = ihdp$treatment,
 Χ
                = ihdp[, 6:30],
 num_tree
                = 10.
 num_chain
                 = 2.
 num_post_sample = 20,
 num_burn_in = 10,
 verbose
                 = FALSE
```

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count\_omp\_thread

Count the number of OpenMP threads for parallel computation

## **Description**

count\_omp\_thread() counts the number of OpenMP threads for parallel computation. If it returns 1, OpenMP is not viable.

## Usage

```
count_omp_thread()
```

# Value

Number of OpenMP thread(s).

## **Examples**

```
count_omp_thread()
```

ihdp

Infant Health and Development Program Data

# Description

Infant Health and Development Program (IHDP) is a randomized experiment from 1985 to 1988 which studied the effect of home visits on cognitive test scores for infants.

# Usage

ihdp

# **Format**

treatment Given treatment.

y\_factual Observed outcome.

**y\_cfactual** Potential outcome given the opposite treatment.

mu0 Control conditional means.

mu1 Treated conditional means.

X1 ~ X6 Confounders with continuous values.

X7 ~ X25 Confounders with binary values.

# **Details**

This dataset was first used by Hill (2011), then used by other researchers (Shalit et al. 2017, Louizos et al. 2017).

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#### Source

Our version of dataset is the dataset used by Louizos et al. (2017). This is the first realization of 10 generated datasets and you can find other realizations from <a href="https://github.com/AMLab-Amsterdam/CEVAE">https://github.com/AMLab-Amsterdam/CEVAE</a>.

#### References

Hill, J. L. (2011). Bayesian nonparametric modeling for causal inference. *Journal of Computational and Graphical Statistics*, 20(1), 217-240. doi:10.1198/jcgs.2010.08162

Louizos, C., Shalit, U., Mooij, J. M., Sontag, D., Zemel, R., & Welling, M. (2017). Causal effect inference with deep latent-variable models. *Advances in neural information processing systems*, 30. doi:10.48550/arXiv.1705.08821 https://github.com/AMLab-Amsterdam/CEVAE

Shalit, U., Johansson, F. D., & Sontag, D. (2017, July). Estimating individual treatment effect: generalization bounds and algorithms. In *International Conference on Machine Learning* (pp. 3076-3085). PMLR. doi:10.48550/arXiv.1606.03976

plot.bartcs

Draw plot for bartcs object

## **Description**

Two options are available: posterior inclusion probability (PIP) plot and trace plot.

# Usage

```
## S3 method for class 'bartcs'
plot(x, method = NULL, parameter = NULL, ...)
```

### **Arguments**

x A bartcs object.

method "pip" for posterior inclusion probability plot or "trace" for trace plot.

parameter Parameter for traceplot.

... Additional arguments for PIP plot. Check ?ggcharts::bar\_chart for possible

arguments.

## **Details**

## PIP plot:

When a posterior sample is sampled during training, separate\_bart() or single\_bart() also counts which variables are included in the model and compute PIP for each variable. For bartcs object x, this is stored in x\$var\_count and x\$var\_prob respectively. plot(method = "pip") uses this information and draws plot using ggcharts::bar\_chart().

### Traceplot:

Parameters are recorded for each MCMC iterations. Parameters include "ATE", "Y1", "Y0", "dir\_alpha", and either "sigma2\_out" from single\_bart() or "sigma2\_out1" and "sigma2\_out0" from separate\_bart(). Vertical line indicates burn-in.

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## Value

A ggplot object of either PIP plot or trace plot.

# **Examples**

```
data(ihdp, package = "bartcs")
x <- single_bart(</pre>
  Υ
                   = ihdp$y_factual,
  trt
                  = ihdp$treatment,
                  = ihdp[, 6:30],
  Χ
  num_tree = 10,
num_chain = 2,
  num_post_sample = 20,
  num_burn_in = 10,
                 = FALSE
  verbose
# PIP plot
plot(x, method = "pip")
plot(x, method = "pip", top_n = 10)
plot(x, method = "pip", threshold = 0.5)
# Check `?ggcharts::bar_chart` for other possible arguments.
# trace plot
plot(x, method = "trace")
plot(x, method = "trace", "Y1")
plot(x, method = "trace", "dir_alpha")
```

summary.bartcs

Summary for bartcs object

# Description

Provide summary for bartcs object.

# Usage

```
## S3 method for class 'bartcs'
summary(object, ...)
```

# Arguments

```
object A bartcs object.
... Additional arguments. Not yet supported.
```

# **Details**

summary() provides 95% posterior credible interval for both aggregated outcome and individual outcomes from each MCMC chain.

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## Value

Provide list with the following components

model separate\_bart or single\_bart.

trt\_value Treatment values for each treatment group: trt\_treated for the treatment

group and trt\_control for the control group.

tree\_params Parameters for the tree structure. chain\_params Parameters for MCMC chains.

outcome Summary of outcomes from the model. This includes both aggregated outcome

and individual outcomes from each MCMC chain.

# **Examples**

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