

# Package ‘stochtree’

January 28, 2026

**Title** Stochastic Tree Ensembles (XBART and BART) for Supervised Learning and Causal Inference

**Version** 0.3.0

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**Description** Flexible stochastic tree ensemble software.  
Robust implementations of Bayesian Additive Regression Trees (BART) Chipman, George, McCulloch (2010) <[doi:10.1214/09-AOAS285](https://doi.org/10.1214/09-AOAS285)> for supervised learning and Bayesian Causal Forests (BCF) Hahn, Murray, Carvalho (2020) <[doi:10.1214/19-BA1195](https://doi.org/10.1214/19-BA1195)> for causal inference. Enables model serialization and parallel sampling and provides a low-level interface for custom stochastic forest samplers.

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**LinkingTo** cpp11, BH

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**SystemRequirements** C++17

**Imports** R6, stats

**URL** <https://stochtree.ai/>, <https://github.com/StochasticTree/stochtree>

**BugReports** <https://github.com/StochasticTree/stochtree/issues>

**Config/testthat/edition** 3

**NeedsCompilation** yes

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|                   |                                                                                                           |
|-------------------|-----------------------------------------------------------------------------------------------------------|
| stochtree-package | <i>stochtree: Stochastic Tree Ensembles (XBART and BART) for Supervised Learning and Causal Inference</i> |
|-------------------|-----------------------------------------------------------------------------------------------------------|

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## Description

Flexible stochastic tree ensemble software. Robust implementations of Bayesian Additive Regression Trees (BART) Chipman, George, McCulloch (2010) [doi:10.1214/09AOAS285](https://doi.org/10.1214/09AOAS285) for supervised learning and Bayesian Causal Forests (BCF) Hahn, Murray, Carvalho (2020) [doi:10.1214/19BA1195](https://doi.org/10.1214/19BA1195) for causal inference. Enables model serialization and parallel sampling and provides a low-level interface for custom stochastic forest samplers.

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## See Also

Useful links:

- <https://stochtree.ai/>
- <https://github.com/StochasticTree/stochtree>
- Report bugs at <https://github.com/StochasticTree/stochtree/issues>

---

bart

*Run BART for Supervised Learning*

---

## Description

Run the BART algorithm for supervised learning.

## Usage

```
bart(  
  X_train,  
  y_train,  
  leaf_basis_train = NULL,  
  rfx_group_ids_train = NULL,  
  rfx_basis_train = NULL,  
  X_test = NULL,  
  leaf_basis_test = NULL,
```

```

    rfx_group_ids_test = NULL,
    rfx_basis_test = NULL,
    num_gfr = 5,
    num_burnin = 0,
    num_mcmc = 100,
    previous_model_json = NULL,
    previous_model_warmstart_sample_num = NULL,
    general_params = list(),
    mean_forest_params = list(),
    variance_forest_params = list(),
    random_effects_params = list()
)

```

## Arguments

|                                  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
|----------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>X_train</code>             | Covariates used to split trees in the ensemble. May be provided either as a dataframe or a matrix. Matrix covariates will be assumed to be all numeric. Covariates passed as a dataframe will be preprocessed based on the variable types (e.g. categorical columns stored as unordered factors will be one-hot encoded, categorical columns stored as ordered factors will be passed as integers to the core algorithm, along with the metadata that the column is ordered categorical). |
| <code>y_train</code>             | Outcome to be modeled by the ensemble.                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| <code>leaf_basis_train</code>    | (Optional) Bases used to define a regression model $y \sim W$ in each leaf of each regression tree. By default, BART assumes constant leaf node parameters, implicitly regressing on a constant basis of ones (i.e. $y \sim 1$ ).                                                                                                                                                                                                                                                         |
| <code>rfx_group_ids_train</code> | (Optional) Group labels used for an additive random effects model.                                                                                                                                                                                                                                                                                                                                                                                                                        |
| <code>rfx_basis_train</code>     | (Optional) Basis for "random-slope" regression in an additive random effects model. If <code>rfx_group_ids_train</code> is provided with a regression basis, an intercept-only random effects model will be estimated.                                                                                                                                                                                                                                                                    |
| <code>X_test</code>              | (Optional) Test set of covariates used to define "out of sample" evaluation data. May be provided either as a dataframe or a matrix, but the format of <code>X_test</code> must be consistent with that of <code>X_train</code> .                                                                                                                                                                                                                                                         |
| <code>leaf_basis_test</code>     | (Optional) Test set of bases used to define "out of sample" evaluation data. While a test set is optional, the structure of any provided test set must match that of the training set (i.e. if both <code>X_train</code> and <code>leaf_basis_train</code> are provided, then a test set must consist of <code>X_test</code> and <code>leaf_basis_test</code> with the same number of columns).                                                                                           |
| <code>rfx_group_ids_test</code>  | (Optional) Test set group labels used for an additive random effects model. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set.                                                                                                                                                                                                                                                                         |
| <code>rfx_basis_test</code>      | (Optional) Test set basis for "random-slope" regression in additive random effects model.                                                                                                                                                                                                                                                                                                                                                                                                 |

|                                     |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
|-------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| num_gfr                             | Number of "warm-start" iterations run using the grow-from-root algorithm (He and Hahn, 2021). Default: 5.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
| num_burnin                          | Number of "burn-in" iterations of the MCMC sampler. Default: 0.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               |
| num_mcmc                            | Number of "retained" iterations of the MCMC sampler. Default: 100.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| previous_model_json                 | (Optional) JSON string containing a previous BART model. This can be used to "continue" a sampler interactively after inspecting the samples or to run parallel chains "warm-started" from existing forest samples. Default: NULL.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
| previous_model_warmstart_sample_num | (Optional) Sample number from previous_model_json that will be used to warmstart this BART sampler. One-indexed (so that the first sample is used for warm-start by setting previous_model_warmstart_sample_num = 1). Default: NULL. If num_chains in the general_params list is > 1, then each successive chain will be initialized from a different sample, counting backwards from previous_model_warmstart_sample_num. That is, if previous_model_warmstart_sample_num = 10 and num_chains = 4, then chain 1 will be initialized from sample 10, chain 2 from sample 9, chain 3 from sample 8, and chain 4 from sample 7. If previous_model_json is provided but previous_model_warmstart_sample_num is NULL, the last sample in the previous model will be used to initialize the first chain, counting backwards as noted before. If more chains are requested than there are samples in previous_model_json, a warning will be raised and only the last sample will be used.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| general_params                      | <p>(Optional) A list of general (non-forest-specific) model parameters, each of which has a default value processed internally, so this argument list is optional.</p> <ul style="list-style-type: none"> <li>• <code>cutpoint_grid_size</code> Maximum size of the "grid" of potential cutpoints to consider in the GFR algorithm. Default: 100.</li> <li>• <code>standardize</code> Whether or not to standardize the outcome (and store the offset / scale in the model object). Default: TRUE.</li> <li>• <code>sample_sigma2_global</code> Whether or not to update the <math>\sigma^2</math> global error variance parameter based on <math>\text{IG}(\sigma^2_{\text{global\_shape}}, \sigma^2_{\text{global\_scale}})</math>. Default: TRUE.</li> <li>• <code>sigma2_global_init</code> Starting value of global error variance parameter. Calibrated internally as <math>1.0 \times \text{var}(y_{\text{train}})</math>, where <math>y_{\text{train}}</math> is the possibly standardized outcome, if not set.</li> <li>• <code>sigma2_global_shape</code> Shape parameter in the <math>\text{IG}(\sigma^2_{\text{global\_shape}}, \sigma^2_{\text{global\_scale}})</math> global error variance model. Default: 0.</li> <li>• <code>sigma2_global_scale</code> Scale parameter in the <math>\text{IG}(\sigma^2_{\text{global\_shape}}, \sigma^2_{\text{global\_scale}})</math> global error variance model. Default: 0.</li> <li>• <code>variable_weights</code> Numeric weights reflecting the relative probability of splitting on each variable. Does not need to sum to 1 but cannot be negative. Defaults to <math>\text{rep}(1/\text{ncol}(X_{\text{train}}), \text{ncol}(X_{\text{train}}))</math> if not set here. Note that if the propensity score is included as a covariate in either forest, its weight will default to <math>1/\text{ncol}(X_{\text{train}})</math>.</li> <li>• <code>random_seed</code> Integer parameterizing the C++ random number generator. If not specified, the C++ random number generator is seeded according to <code>std::random_device</code>.</li> </ul> |

- `keep_burnin` Whether or not "burnin" samples should be included in the stored samples of forests and other parameters. Default FALSE. Ignored if `num_mcmc = 0`.
- `keep_gfr` Whether or not "grow-from-root" samples should be included in the stored samples of forests and other parameters. Default FALSE. Ignored if `num_mcmc = 0`.
- `keep_every` How many iterations of the burned-in MCMC sampler should be run before forests and parameters are retained. Default 1. Setting `keep_every <- k` for some  $k > 1$  will "thin" the MCMC samples by retaining every  $k$ -th sample, rather than simply every sample. This can reduce the autocorrelation of the MCMC samples.
- `num_chains` How many independent MCMC chains should be sampled. If `num_mcmc = 0`, this is ignored. If `num_gfr = 0`, then each chain is run from root for `num_mcmc * keep_every + num_burnin` iterations, with `num_mcmc` samples retained. If `num_gfr > 0`, each MCMC chain will be initialized from a separate GFR ensemble, with the requirement that `num_gfr >= num_chains`. Default: 1. Note that if `num_chains > 1`, the returned model object will contain samples from all chains, stored consecutively. That is, if there are 4 chains with 100 samples each, the first 100 samples will be from chain 1, the next 100 samples will be from chain 2, etc... For more detail on working with multi-chain BART models, see [the multi chain vignette](#).
- `verbose` Whether or not to print progress during the sampling loops. Default: FALSE.
- `probit_outcome_model` Whether or not the outcome should be modeled as explicitly binary via a probit link. If TRUE, `y` must only contain the values 0 and 1. Default: FALSE.
- `num_threads` Number of threads to use in the GFR and MCMC algorithms, as well as prediction. If OpenMP is not available on a user's setup, this will default to 1, otherwise to the maximum number of available threads.

#### `mean_forest_params`

(Optional) A list of mean forest model parameters, each of which has a default value processed internally, so this argument list is optional.

- `num_trees` Number of trees in the ensemble for the conditional mean model. Default: 200. If `num_trees = 0`, the conditional mean will not be modeled using a forest, and the function will only proceed if `num_trees > 0` for the variance forest.
- `alpha` Prior probability of splitting for a tree of depth 0 in the mean model. Tree split prior combines `alpha` and `beta` via  $\alpha * (1 + \text{node\_depth})^{-\beta}$ . Default: 0.95.
- `beta` Exponent that decreases split probabilities for nodes of depth  $> 0$  in the mean model. Tree split prior combines `alpha` and `beta` via  $\alpha * (1 + \text{node\_depth})^{-\beta}$ . Default: 2.
- `min_samples_leaf` Minimum allowable size of a leaf, in terms of training samples, in the mean model. Default: 5.
- `max_depth` Maximum depth of any tree in the ensemble in the mean model. Default: 10. Can be overridden with `-1` which does not enforce any depth limits on trees.

- `sample_sigma2_leaf` Whether or not to update the leaf scale variance parameter based on  $IG(\text{sigma2\_leaf\_shape}, \text{sigma2\_leaf\_scale})$ . Cannot (currently) be set to true if  $\text{ncol}(\text{leaf\_basis\_train}) > 1$ . Default: FALSE.
- `sigma2_leaf_init` Starting value of leaf node scale parameter. Calibrated internally as  $1/\text{num\_trees}$  if not set here.
- `sigma2_leaf_shape` Shape parameter in the  $IG(\text{sigma2\_leaf\_shape}, \text{sigma2\_leaf\_scale})$  leaf node parameter variance model. Default: 3.
- `sigma2_leaf_scale` Scale parameter in the  $IG(\text{sigma2\_leaf\_shape}, \text{sigma2\_leaf\_scale})$  leaf node parameter variance model. Calibrated internally as  $0.5/\text{num\_trees}$  if not set here.
- `keep_vars` Vector of variable names or column indices denoting variables that should be included in the forest. Default: NULL.
- `drop_vars` Vector of variable names or column indices denoting variables that should be excluded from the forest. Default: NULL. If both `drop_vars` and `keep_vars` are set, `drop_vars` will be ignored.
- `num_features_subsample` How many features to subsample when growing each tree for the GFR algorithm. Defaults to the number of features in the training dataset.

#### `variance_forest_params`

(Optional) A list of variance forest model parameters, each of which has a default value processed internally, so this argument list is optional.

- `num_trees` Number of trees in the ensemble for the conditional variance model. Default: 0. Variance is only modeled using a tree / forest if `num_trees > 0`.
- `alpha` Prior probability of splitting for a tree of depth 0 in the variance model. Tree split prior combines alpha and beta via  $\alpha * (1 + \text{node\_depth})^{-\beta}$ . Default: 0.95.
- `beta` Exponent that decreases split probabilities for nodes of depth  $> 0$  in the variance model. Tree split prior combines alpha and beta via  $\alpha * (1 + \text{node\_depth})^{-\beta}$ . Default: 2.
- `min_samples_leaf` Minimum allowable size of a leaf, in terms of training samples, in the variance model. Default: 5.
- `max_depth` Maximum depth of any tree in the ensemble in the variance model. Default: 10. Can be overridden with -1 which does not enforce any depth limits on trees.
- `leaf_prior_calibration_param` Hyperparameter used to calibrate the  $IG(\text{var\_forest\_prior\_shape}, \text{var\_forest\_prior\_scale})$  conditional error variance model. If `var_forest_prior_shape` and `var_forest_prior_scale` are not set below, this calibration parameter is used to set these values to  $\text{num\_trees} / \text{leaf\_prior\_calibration\_param}^2 + 0.5$  and  $\text{num\_trees} / \text{leaf\_prior\_calibration\_param}^2$ , respectively. Default: 1.5.
- `var_forest_leaf_init` Starting value of root forest prediction in conditional (heteroskedastic) error variance model. Calibrated internally as  $\log(0.6 * \text{var}(y_{\text{train}})) / \text{num\_trees}$ , where `y_train` is the possibly standardized outcome, if not set.

- `var_forest_prior_shape` Shape parameter in the  $IG(\text{var\_forest\_prior\_shape}, \text{var\_forest\_prior\_scale})$  conditional error variance model (which is only sampled if `num_trees > 0`). Calibrated internally as `num_trees / leaf_prior_calibration_param + 0.5` if not set.
- `var_forest_prior_scale` Scale parameter in the  $IG(\text{var\_forest\_prior\_shape}, \text{var\_forest\_prior\_scale})$  conditional error variance model (which is only sampled if `num_trees > 0`). Calibrated internally as `num_trees / leaf_prior_calibration_param` if not set.
- `keep_vars` Vector of variable names or column indices denoting variables that should be included in the forest. Default: `NULL`.
- `drop_vars` Vector of variable names or column indices denoting variables that should be excluded from the forest. Default: `NULL`. If both `drop_vars` and `keep_vars` are set, `drop_vars` will be ignored.
- `num_features_subsample` How many features to subsample when growing each tree for the GFR algorithm. Defaults to the number of features in the training dataset.

#### `random_effects_params`

(Optional) A list of random effects model parameters, each of which has a default value processed internally, so this argument list is optional.

- `model_spec` Specification of the random effects model. Options are "custom" and "intercept\_only". If "custom" is specified, then a user-provided basis must be passed through `rfx_basis_train`. If "intercept\_only" is specified, a random effects basis of all ones will be dispatched internally at sampling and prediction time. If "intercept\_plus\_treatment" is specified, a random effects basis that combines an "intercept" basis of all ones with the treatment variable (`Z_train`) will be dispatched internally at sampling and prediction time. Default: "custom". If "intercept\_only" is specified, `rfx_basis_train` and `rfx_basis_test` (if provided) will be ignored.
- `working_parameter_prior_mean` Prior mean for the random effects "working parameter". Default: `NULL`. Must be a vector whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a vector.
- `group_parameters_prior_mean` Prior mean for the random effects "group parameters." Default: `NULL`. Must be a vector whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a vector.
- `working_parameter_prior_cov` Prior covariance matrix for the random effects "working parameter." Default: `NULL`. Must be a square matrix whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a diagonal matrix.
- `group_parameter_prior_cov` Prior covariance matrix for the random effects "group parameters." Default: `NULL`. Must be a square matrix whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a diagonal matrix.
- `variance_prior_shape` Shape parameter for the inverse gamma prior on the variance of the random effects "group parameter." Default: 1.

- `variance_prior_scale` Scale parameter for the inverse gamma prior on the variance of the random effects "group parameter." Default: 1.

### Value

List of sampling outputs and a wrapper around the sampled forests (which can be used for in-memory prediction on new data, or serialized to JSON on disk).

### Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]

bart_model <- bart(X_train = X_train, y_train = y_train, X_test = X_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
```

---

bcf

---

*Run BCF for Causal Effect Estimation*


---

### Description

Run the Bayesian Causal Forest (BCF) algorithm for regularized causal effect estimation.

### Usage

```
bcf(
  X_train,
  Z_train,
  y_train,
  propensity_train = NULL,
  rfx_group_ids_train = NULL,
```

```

    rfx_basis_train = NULL,
    X_test = NULL,
    Z_test = NULL,
    propensity_test = NULL,
    rfx_group_ids_test = NULL,
    rfx_basis_test = NULL,
    num_gfr = 5,
    num_burnin = 0,
    num_mcmc = 100,
    previous_model_json = NULL,
    previous_model_warmstart_sample_num = NULL,
    general_params = list(),
    prognostic_forest_params = list(),
    treatment_effect_forest_params = list(),
    variance_forest_params = list(),
    random_effects_params = list()
)

```

## Arguments

|                                  |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
|----------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>X_train</code>             | Covariates used to split trees in the ensemble. May be provided either as a dataframe or a matrix. Matrix covariates will be assumed to be all numeric. Covariates passed as a dataframe will be preprocessed based on the variable types (e.g. categorical columns stored as unordered factors will be one-hot encoded, categorical columns stored as ordered factors will be passed as integers to the core algorithm, along with the metadata that the column is ordered categorical). |
| <code>Z_train</code>             | Vector of (continuous or binary) treatment assignments.                                                                                                                                                                                                                                                                                                                                                                                                                                   |
| <code>y_train</code>             | Outcome to be modeled by the ensemble.                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
| <code>propensity_train</code>    | (Optional) Vector of propensity scores. If not provided, this will be estimated from the data.                                                                                                                                                                                                                                                                                                                                                                                            |
| <code>rfx_group_ids_train</code> | (Optional) Group labels used for an additive random effects model.                                                                                                                                                                                                                                                                                                                                                                                                                        |
| <code>rfx_basis_train</code>     | (Optional) Basis for "random-slope" regression in an additive random effects model. If <code>rfx_group_ids_train</code> is provided with a regression basis, an intercept-only random effects model will be estimated.                                                                                                                                                                                                                                                                    |
| <code>X_test</code>              | (Optional) Test set of covariates used to define "out of sample" evaluation data. May be provided either as a dataframe or a matrix, but the format of <code>X_test</code> must be consistent with that of <code>X_train</code> .                                                                                                                                                                                                                                                         |
| <code>Z_test</code>              | (Optional) Test set of (continuous or binary) treatment assignments.                                                                                                                                                                                                                                                                                                                                                                                                                      |
| <code>propensity_test</code>     | (Optional) Vector of propensity scores. If not provided, this will be estimated from the data.                                                                                                                                                                                                                                                                                                                                                                                            |
| <code>rfx_group_ids_test</code>  | (Optional) Test set group labels used for an additive random effects model. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set.                                                                                                                                                                                                                                                                         |

- `rfx_basis_test` (Optional) Test set basis for "random-slope" regression in additive random effects model.
- `num_gfr` Number of "warm-start" iterations run using the grow-from-root algorithm (He and Hahn, 2021). Default: 5.
- `num_burnin` Number of "burn-in" iterations of the MCMC sampler. Default: 0.
- `num_mcmc` Number of "retained" iterations of the MCMC sampler. Default: 100.
- `previous_model_json` (Optional) JSON string containing a previous BCF model. This can be used to "continue" a sampler interactively after inspecting the samples or to run parallel chains "warm-started" from existing forest samples. Default: NULL.
- `previous_model_warmstart_sample_num` (Optional) Sample number from `previous_model_json` that will be used to warmstart this BCF sampler. One-indexed (so that the first sample is used for warm-start by setting `previous_model_warmstart_sample_num = 1`). Default: NULL. If `num_chains` in the `general_params` list is  $> 1$ , then each successive chain will be initialized from a different sample, counting backwards from `previous_model_warmstart_sample_num`. That is, if `previous_model_warmstart_sample_num = 10` and `num_chains = 4`, then chain 1 will be initialized from sample 10, chain 2 from sample 9, chain 3 from sample 8, and chain 4 from sample 7. If `previous_model_json` is provided but `previous_model_warmstart_sample_num` is NULL, the last sample in the previous model will be used to initialize the first chain, counting backwards as noted before. If more chains are requested than there are samples in `previous_model_json`, a warning will be raised and only the last sample will be used.
- `general_params` (Optional) A list of general (non-forest-specific) model parameters, each of which has a default value processed internally, so this argument list is optional.
- `cutpoint_grid_size` Maximum size of the "grid" of potential cutpoints to consider in the GFR algorithm. Default: 100.
  - `standardize` Whether or not to standardize the outcome (and store the offset / scale in the model object). Default: TRUE.
  - `sample_sigma2_global` Whether or not to update the  $\sigma^2$  global error variance parameter based on  $IG(\sigma^2_{\text{global\_shape}}, \sigma^2_{\text{global\_scale}})$ . Default: TRUE.
  - `sigma2_global_init` Starting value of global error variance parameter. Calibrated internally as  $1.0 \times \text{var}((y_{\text{train}} - \text{mean}(y_{\text{train}})) / \text{sd}(y_{\text{train}}))$  if not set.
  - `sigma2_global_shape` Shape parameter in the  $IG(\sigma^2_{\text{global\_shape}}, \sigma^2_{\text{global\_scale}})$  global error variance model. Default: 0.
  - `sigma2_global_scale` Scale parameter in the  $IG(\sigma^2_{\text{global\_shape}}, \sigma^2_{\text{global\_scale}})$  global error variance model. Default: 0.
  - `variable_weights` Numeric weights reflecting the relative probability of splitting on each variable. Does not need to sum to 1 but cannot be negative. Defaults to  $\text{rep}(1/\text{ncol}(X_{\text{train}}), \text{ncol}(X_{\text{train}}))$  if not set here. Note that if the propensity score is included as a covariate in either forest, its weight will default to  $1/\text{ncol}(X_{\text{train}})$ . A workaround if you wish to provide a custom weight for the propensity score is to include it as a

column in `X_train` and then set `propensity_covariate` to 'none' adjust `keep_vars` accordingly for the prognostic or treatment-effect forests.

- `propensity_covariate` Whether to include the propensity score as a covariate in either or both of the forests. Enter "none" for neither, "prognostic" for the prognostic forest, "treatment\_effect" for the treatment forest, and "both" for both forests. If this is not "none" and a propensity score is not provided, it will be estimated from  $(X_{\text{train}}, Z_{\text{train}})$  using `stochtree::bart()`. Default: "mu".
- `adaptive_coding` Whether or not to use an "adaptive coding" scheme in which a binary treatment variable is not coded manually as (0,1) or (-1,1) but learned via parameters `b_0` and `b_1` that attach to the outcome model  $[b_0 (1-Z) + b_1 Z] \tau(X)$ . This is ignored when `Z` is not binary. Default: TRUE.
- `control_coding_init` Initial value of the "control" group coding parameter. This is ignored when `Z` is not binary. Default: -0.5.
- `treated_coding_init` Initial value of the "treatment" group coding parameter. This is ignored when `Z` is not binary. Default: 0.5.
- `rfx_prior_var` Prior on the (diagonals of the) covariance of the additive group-level random regression coefficients. Must be a vector of length `ncol(rfx_basis_train)`. Default: `rep(1, ncol(rfx_basis_train))`
- `random_seed` Integer parameterizing the C++ random number generator. If not specified, the C++ random number generator is seeded according to `std::random_device`.
- `keep_burnin` Whether or not "burnin" samples should be included in the stored samples of forests and other parameters. Default FALSE. Ignored if `num_mcmc = 0`.
- `keep_gfr` Whether or not "grow-from-root" samples should be included in the stored samples of forests and other parameters. Default FALSE. Ignored if `num_mcmc = 0`.
- `keep_every` How many iterations of the burned-in MCMC sampler should be run before forests and parameters are retained. Default 1. Setting `keep_every <= k` for some `k > 1` will "thin" the MCMC samples by retaining every `k`-th sample, rather than simply every sample. This can reduce the autocorrelation of the MCMC samples.
- `num_chains` How many independent MCMC chains should be sampled. If `num_mcmc = 0`, this is ignored. If `num_gfr = 0`, then each chain is run from root for `num_mcmc * keep_every + num_burnin` iterations, with `num_mcmc` samples retained. If `num_gfr > 0`, each MCMC chain will be initialized from a separate GFR ensemble, with the requirement that `num_gfr >= num_chains`. Default: 1. Note that if `num_chains > 1`, the returned model object will contain samples from all chains, stored consecutively. That is, if there are 4 chains with 100 samples each, the first 100 samples will be from chain 1, the next 100 samples will be from chain 2, etc... For more detail on working with multi-chain BCF models, see [the multi chain vignette](#).
- `verbose` Whether or not to print progress during the sampling loops. Default: FALSE.

- `probit_outcome_model` Whether or not the outcome should be modeled as explicitly binary via a probit link. If TRUE, y must only contain the values 0 and 1. Default: FALSE.
- `num_threads` Number of threads to use in the GFR and MCMC algorithms, as well as prediction. If OpenMP is not available on a user's setup, this will default to 1, otherwise to the maximum number of available threads.

#### `prognostic_forest_params`

(Optional) A list of prognostic forest model parameters, each of which has a default value processed internally, so this argument list is optional.

- `num_trees` Number of trees in the ensemble for the prognostic forest. Default: 250. Must be a positive integer.
- `alpha` Prior probability of splitting for a tree of depth 0 in the prognostic forest. Tree split prior combines alpha and beta via  $\alpha \cdot (1 + \text{node\_depth})^{-\beta}$ . Default: 0.95.
- `beta` Exponent that decreases split probabilities for nodes of depth > 0 in the prognostic forest. Tree split prior combines alpha and beta via  $\alpha \cdot (1 + \text{node\_depth})^{-\beta}$ . Default: 2.
- `min_samples_leaf` Minimum allowable size of a leaf, in terms of training samples, in the prognostic forest. Default: 5.
- `max_depth` Maximum depth of any tree in the ensemble in the prognostic forest. Default: 10. Can be overridden with -1 which does not enforce any depth limits on trees.
- `variable_weights` Numeric weights reflecting the relative probability of splitting on each variable in the prognostic forest. Does not need to sum to 1 but cannot be negative. Defaults to `rep(1/ncol(X_train), ncol(X_train))` if not set here.
- `sample_sigma2_leaf` Whether or not to update the leaf scale variance parameter based on `IG(sigma2_leaf_shape, sigma2_leaf_scale)`.
- `sigma2_leaf_init` Starting value of leaf node scale parameter. Calibrated internally as `1/num_trees` if not set here.
- `sigma2_leaf_shape` Shape parameter in the `IG(sigma2_leaf_shape, sigma2_leaf_scale)` leaf node parameter variance model. Default: 3.
- `sigma2_leaf_scale` Scale parameter in the `IG(sigma2_leaf_shape, sigma2_leaf_scale)` leaf node parameter variance model. Calibrated internally as `0.5/num_trees` if not set here.
- `keep_vars` Vector of variable names or column indices denoting variables that should be included in the forest. Default: NULL.
- `drop_vars` Vector of variable names or column indices denoting variables that should be excluded from the forest. Default: NULL. If both `drop_vars` and `keep_vars` are set, `drop_vars` will be ignored.
- `num_features_subsample` How many features to subsample when growing each tree for the GFR algorithm. Defaults to the number of features in the training dataset.

#### `treatment_effect_forest_params`

(Optional) A list of treatment effect forest model parameters, each of which has a default value processed internally, so this argument list is optional.

- `num_trees` Number of trees in the ensemble for the treatment effect forest. Default: 50. Must be a positive integer.
- `alpha` Prior probability of splitting for a tree of depth 0 in the treatment effect forest. Tree split prior combines alpha and beta via  $\alpha \cdot (1 + \text{node\_depth})^{-\beta}$ . Default: 0.25.
- `beta` Exponent that decreases split probabilities for nodes of depth  $> 0$  in the treatment effect forest. Tree split prior combines alpha and beta via  $\alpha \cdot (1 + \text{node\_depth})^{-\beta}$ . Default: 3.
- `min_samples_leaf` Minimum allowable size of a leaf, in terms of training samples, in the treatment effect forest. Default: 5.
- `max_depth` Maximum depth of any tree in the ensemble in the treatment effect forest. Default: 5. Can be overridden with -1 which does not enforce any depth limits on trees.
- `variable_weights` Numeric weights reflecting the relative probability of splitting on each variable in the treatment effect forest. Does not need to sum to 1 but cannot be negative. Defaults to `rep(1/ncol(X_train), ncol(X_train))` if not set here.
- `sample_sigma2_leaf` Whether or not to update the leaf scale variance parameter based on `IG(sigma2_leaf_shape, sigma2_leaf_scale)`. Cannot (currently) be set to true if `ncol(Z_train) > 1`. Default: FALSE.
- `sigma2_leaf_init` Starting value of leaf node scale parameter. Calibrated internally as  $1/\text{num\_trees}$  if not set here.
- `sigma2_leaf_shape` Shape parameter in the `IG(sigma2_leaf_shape, sigma2_leaf_scale)` leaf node parameter variance model. Default: 3.
- `sigma2_leaf_scale` Scale parameter in the `IG(sigma2_leaf_shape, sigma2_leaf_scale)` leaf node parameter variance model. Calibrated internally as  $0.5/\text{num\_trees}$  if not set here.
- `delta_max` Maximum plausible conditional distributional treatment effect (i.e.  $P(Y(1) = 1 | X) - P(Y(0) = 1 | X)$ ) when the outcome is binary. Only used when the outcome is specified as a probit model in `general_params`. Must be  $> 0$  and  $< 1$ . Default: 0.9. Ignored if `sigma2_leaf_init` is set directly, as this parameter is used to calibrate `sigma2_leaf_init`.
- `keep_vars` Vector of variable names or column indices denoting variables that should be included in the forest. Default: NULL.
- `drop_vars` Vector of variable names or column indices denoting variables that should be excluded from the forest. Default: NULL. If both `drop_vars` and `keep_vars` are set, `drop_vars` will be ignored.
- `num_features_subsample` How many features to subsample when growing each tree for the GFR algorithm. Defaults to the number of features in the training dataset.

#### `variance_forest_params`

(Optional) A list of variance forest model parameters, each of which has a default value processed internally, so this argument list is optional.

- `num_trees` Number of trees in the ensemble for the conditional variance model. Default: 0. Variance is only modeled using a tree / forest if `num_trees > 0`.

- `alpha` Prior probability of splitting for a tree of depth 0 in the variance model. Tree split prior combines alpha and beta via  $\alpha \cdot (1 + \text{node\_depth})^{-\beta}$ . Default: 0.95.
- `beta` Exponent that decreases split probabilities for nodes of depth  $> 0$  in the variance model. Tree split prior combines alpha and beta via  $\alpha \cdot (1 + \text{node\_depth})^{-\beta}$ . Default: 2.
- `min_samples_leaf` Minimum allowable size of a leaf, in terms of training samples, in the variance model. Default: 5.
- `max_depth` Maximum depth of any tree in the ensemble in the variance model. Default: 10. Can be overridden with -1 which does not enforce any depth limits on trees.
- `leaf_prior_calibration_param` Hyperparameter used to calibrate the  $\text{IG}(\text{var\_forest\_prior\_shape}, \text{var\_forest\_prior\_scale})$  conditional error variance model. If `var_forest_prior_shape` and `var_forest_prior_scale` are not set below, this calibration parameter is used to set these values to  $\text{num\_trees} / \text{leaf\_prior\_calibration\_param}^2 + 0.5$  and  $\text{num\_trees} / \text{leaf\_prior\_calibration\_param}^2$ , respectively. Default: 1.5.
- `variance_forest_init` Starting value of root forest prediction in conditional (heteroskedastic) error variance model. Calibrated internally as  $\log(0.6 \cdot \text{var}((y_{\text{train}} - \text{mean}(y_{\text{train}})) / \text{sd}(y_{\text{train}}))) / \text{num\_trees}$  if not set.
- `var_forest_prior_shape` Shape parameter in the  $\text{IG}(\text{var\_forest\_prior\_shape}, \text{var\_forest\_prior\_scale})$  conditional error variance model (which is only sampled if `num_trees`  $> 0$ ). Calibrated internally as  $\text{num\_trees} / 1.5^2 + 0.5$  if not set.
- `var_forest_prior_scale` Scale parameter in the  $\text{IG}(\text{var\_forest\_prior\_shape}, \text{var\_forest\_prior\_scale})$  conditional error variance model (which is only sampled if `num_trees`  $> 0$ ). Calibrated internally as  $\text{num\_trees} / 1.5^2$  if not set.
- `keep_vars` Vector of variable names or column indices denoting variables that should be included in the forest. Default: NULL.
- `drop_vars` Vector of variable names or column indices denoting variables that should be excluded from the forest. Default: NULL. If both `drop_vars` and `keep_vars` are set, `drop_vars` will be ignored.
- `num_features_subsample` How many features to subsample when growing each tree for the GFR algorithm. Defaults to the number of features in the training dataset.

#### `random_effects_params`

(Optional) A list of random effects model parameters, each of which has a default value processed internally, so this argument list is optional.

- `model_spec` Specification of the random effects model. Options are "custom", "intercept\_only", and "intercept\_plus\_treatment". If "custom" is specified, then a user-provided basis must be passed through `rfx_basis_train`. If "intercept\_only" is specified, a random effects basis of all ones will be dispatched internally at sampling and prediction time. If "intercept\_plus\_treatment" is specified, a random effects basis that combines an "intercept" basis of

all ones with the treatment variable (`Z_train`) will be dispatched internally at sampling and prediction time. Default: "custom". If either "intercept\_only" or "intercept\_plus\_treatment" is specified, `rfx_basis_train` and `rfx_basis_test` (if provided) will be ignored.

- `working_parameter_prior_mean` Prior mean for the random effects "working parameter". Default: NULL. Must be a vector whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a vector.
- `group_parameters_prior_mean` Prior mean for the random effects "group parameters." Default: NULL. Must be a vector whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a vector.
- `working_parameter_prior_cov` Prior covariance matrix for the random effects "working parameter." Default: NULL. Must be a square matrix whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a diagonal matrix.
- `group_parameter_prior_cov` Prior covariance matrix for the random effects "group parameters." Default: NULL. Must be a square matrix whose dimension matches the number of random effects bases, or a scalar value that will be expanded to a diagonal matrix.
- `variance_prior_shape` Shape parameter for the inverse gamma prior on the variance of the random effects "group parameter." Default: 1.
- `variance_prior_scale` Scale parameter for the inverse gamma prior on the variance of the random effects "group parameter." Default: 1.

## Value

List of sampling outputs and a wrapper around the sampled forests (which can be used for in-memory prediction on new data, or serialized to JSON on disk).

## Examples

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
```

```

      ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
      ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
    )
    Z <- rbinom(n, 1, pi_x)
    noise_sd <- 1
    y <- mu_x + tau_x*Z + rnorm(n, 0, noise_sd)
    test_set_pct <- 0.2
    n_test <- round(test_set_pct*n)
    n_train <- n - n_test
    test_inds <- sort(sample(1:n, n_test, replace = FALSE))
    train_inds <- (1:n)[!((1:n) %in% test_inds)]
    X_test <- X[test_inds,]
    X_train <- X[train_inds,]
    pi_test <- pi_x[test_inds]
    pi_train <- pi_x[train_inds]
    Z_test <- Z[test_inds]
    Z_train <- Z[train_inds]
    y_test <- y[test_inds]
    y_train <- y[train_inds]
    mu_test <- mu_x[test_inds]
    mu_train <- mu_x[train_inds]
    tau_test <- tau_x[test_inds]
    tau_train <- tau_x[train_inds]
    bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
                     propensity_train = pi_train, X_test = X_test, Z_test = Z_test,
                     propensity_test = pi_test, num_gfr = 10,
                     num_burnin = 0, num_mcmc = 10)

```

---

calibrateInverseGammaErrorVariance

*Calibrate Inverse Gamma Prior*


---

## Description

Calibrate the scale parameter on an inverse gamma prior for the global error variance as in Chipman et al (2022)

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](https://stochtree.ai)

Chipman, H., George, E., Hahn, R., McCulloch, R., Pratola, M. and Sparapani, R. (2022). Bayesian Additive Regression Trees, Computational Approaches. In Wiley StatsRef: Statistics Reference Online (eds N. Balakrishnan, T. Colton, B. Everitt, W. Piegorsch, F. Ruggeri and J.L. Teugels). <https://doi.org/10.1002/9781118445112.stat08288>

## Usage

```

calibrateInverseGammaErrorVariance(
  y,

```

```

X,
W = NULL,
nu = 3,
quant = 0.9,
standardize = TRUE
)

```

### Arguments

|             |                                                                                                                                                                                                                                                                                                                    |
|-------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| y           | Outcome to be modeled using BART, BCF or another nonparametric ensemble method.                                                                                                                                                                                                                                    |
| X           | Covariates to be used to partition trees in an ensemble or series of ensemble.                                                                                                                                                                                                                                     |
| W           | (Optional) Basis used to define a "leaf regression" model for each decision tree. The "classic" BART model assumes a constant leaf parameter, which is equivalent to a "leaf regression" on a basis of all ones, though it is not necessary to pass a vector of ones, here or to the BART function. Default: NULL. |
| nu          | The shape parameter for the global error variance's IG prior. The scale parameter in the Sparapani et al (2021) parameterization is defined as $\text{nu} \times \text{lambda}$ where $\text{lambda}$ is the output of this function. Default: 3.                                                                  |
| quant       | (Optional) Quantile of the inverse gamma prior distribution represented by a linear-regression-based overestimate of $\sigma^2$ . Default: 0.9.                                                                                                                                                                    |
| standardize | (Optional) Whether or not outcome should be standardized $((y - \text{mean}(y)) / \text{sd}(y))$ before calibration of $\text{lambda}$ . Default: TRUE.                                                                                                                                                            |

### Value

Value of  $\text{lambda}$  which determines the scale parameter of the global error variance prior ( $\sigma^2 \sim \text{IG}(\text{nu}, \text{nu} \times \text{lambda})$ )

### Examples

```

n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
y <- 10*X[,1] - 20*X[,2] + rnorm(n)
nu <- 3
lambda <- calibrateInverseGammaErrorVariance(y, X, nu = nu)
sigma2hat <- mean(resid(lm(y~X))^2)
mean(var(y)/rgamma(100000, nu, rate = nu*lambda) < sigma2hat)

```

## Description

Compute and return a vector representation of a forest's leaf predictions for every observation in a dataset.

The vector has a "row-major" format that can be easily re-represented as a CSR sparse matrix: elements are organized so that the first  $n$  elements correspond to leaf predictions for all  $n$  observations in a dataset for the first tree in an ensemble, the next  $n$  elements correspond to predictions for the second tree and so on. The "data" for each element corresponds to a uniquely mapped column index that corresponds to a single leaf of a single tree (i.e. if tree 1 has 3 leaves, its column indices range from 0 to 2, and then tree 2's leaf indices begin at 3, etc...).

## Usage

```
computeForestLeafIndices(
  model_object,
  covariates,
  forest_type = NULL,
  propensity = NULL,
  forest_inds = NULL
)
```

## Arguments

- |              |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|--------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| model_object | Object of type bartmodel, bcfmodel, or ForestSamples corresponding to a BART / BCF model with at least one forest sample, or a low-level ForestSamples object.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| covariates   | Covariates to use for prediction. Must have the same dimensions / column types as the data used to train a forest.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       |
| forest_type  | Which forest to use from model_object. Valid inputs depend on the model type, and whether or not a given forest was sampled in that model. <div style="margin-left: 20px;"> <b>1. BART</b> <ul style="list-style-type: none"> <li>• 'mean': Extracts leaf indices for the mean forest</li> <li>• 'variance': Extracts leaf indices for the variance forest</li> </ul> <b>2. BCF</b> <ul style="list-style-type: none"> <li>• 'prognostic': Extracts leaf indices for the prognostic forest</li> <li>• 'treatment': Extracts leaf indices for the treatment effect forest</li> <li>• 'variance': Extracts leaf indices for the variance forest</li> </ul> <b>3. ForestSamples</b> <ul style="list-style-type: none"> <li>• NULL: It is not necessary to disambiguate when this function is called directly on a ForestSamples object. This is the default value of this</li> </ul> </div> |
| propensity   | (Optional) Propensities used for prediction (BCF-only).                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |
| forest_inds  | (Optional) Indices of the forest sample(s) for which to compute leaf indices. If not provided, this function will return leaf indices for every sample of a forest. This function uses 0-indexing, so the first forest sample corresponds to forest_num = 0, and so on.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |

**Value**

Vector of size `num_obs * num_trees`, where `num_obs = nrow(covariates)` and `num_trees` is the number of trees in the relevant forest of `model_object`.

**Examples**

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
bart_model <- bart(X, y, num_gfr=0, num_mcmc=10)
computeForestLeafIndices(bart_model, X, "mean")
computeForestLeafIndices(bart_model, X, "mean", 0)
computeForestLeafIndices(bart_model, X, "mean", c(1,3,9))
```

---

`computeForestLeafVariances`

*Query Forest Leaf Scale Parameters*

---

**Description**

Return each forest's leaf node scale parameters.

If leaf scale is not sampled for the forest in question, throws an error that the leaf model does not have a stochastic scale parameter.

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the `stochtree`'s advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
computeForestLeafVariances(model_object, forest_type, forest_inds = NULL)
```

**Arguments**

- |                           |                                                                                                                                                          |
|---------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>model_object</code> | Object of type <code>bartmodel</code> or <code>bcmfmodel</code> corresponding to a BART / BCF model with at least one forest sample                      |
| <code>forest_type</code>  | Which forest to use from <code>model_object</code> . Valid inputs depend on the model type, and whether or not a given forest was sampled in that model. |
- 1. BART**
    - 'mean': Extracts leaf indices for the mean forest
    - 'variance': Extracts leaf indices for the variance forest
  - 2. BCF**
    - 'prognostic': Extracts leaf indices for the prognostic forest
    - 'treatment': Extracts leaf indices for the treatment effect forest
    - 'variance': Extracts leaf indices for the variance forest

**forest\_inds** (Optional) Indices of the forest sample(s) for which to compute leaf indices. If not provided, this function will return leaf indices for every sample of a forest. This function uses 0-indexing, so the first forest sample corresponds to `forest_num = 0`, and so on.

### Value

Vector of size `length(forest_inds)` with the leaf scale parameter for each requested forest.

### Examples

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
bart_model <- bart(X, y, num_gfr=0, num_mcmc=10)
computeForestLeafVariances(bart_model, "mean")
computeForestLeafVariances(bart_model, "mean", 0)
computeForestLeafVariances(bart_model, "mean", c(1,3,5))
```

---

computeForestMaxLeafIndex

*Query Forest Max Leaf Index*

---

### Description

Compute and return the largest possible leaf index computable by `computeForestLeafIndices` for the forests in a designated forest sample container.

### Usage

```
computeForestMaxLeafIndex(model_object, forest_type = NULL, forest_inds = NULL)
```

### Arguments

**model\_object** Object of type `bartmodel`, `bcfmodel`, or `ForestSamples` corresponding to a BART/BCF model with at least one forest sample, or a low-level `ForestSamples` object.

**forest\_type** Which forest to use from `model_object`. Valid inputs depend on the model type, and whether or not a

#### 1. BART

- 'mean': Extracts leaf indices for the mean forest
- 'variance': Extracts leaf indices for the variance forest

#### 2. BCF

- 'prognostic': Extracts leaf indices for the prognostic forest
- 'treatment': Extracts leaf indices for the treatment effect forest
- 'variance': Extracts leaf indices for the variance forest

#### 3. ForestSamples

- **NULL:** It is not necessary to disambiguate when this function is called directly on a `ForestSamples` object. This is the default value of this
- forest\_inds** (Optional) Indices of the forest sample(s) for which to compute max leaf indices. If not provided, this function will return max leaf indices for every sample of a forest. This function uses 0-indexing, so the first forest sample corresponds to `forest_num = 0`, and so on.

## Value

Vector containing the largest possible leaf index computable by `computeForestLeafIndices` for the forests in a designated forest sample container.

## Examples

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
bart_model <- bart(X, y, num_gfr=0, num_mcmc=10)
computeForestMaxLeafIndex(bart_model, "mean")
computeForestMaxLeafIndex(bart_model, "mean", 0)
computeForestMaxLeafIndex(bart_model, "mean", c(1,3,9))
```

---

`compute_bart_posterior_interval`

*Compute BART Posterior Credible Intervals*

---

## Description

Compute posterior credible intervals for specified terms from a fitted BART model. Supports intervals for mean functions, variance functions, random effects, and overall outcome predictions.

## Usage

```
compute_bart_posterior_interval(
  model_object,
  terms,
  level = 0.95,
  scale = "linear",
  X = NULL,
  leaf_basis = NULL,
  rfx_group_ids = NULL,
  rfx_basis = NULL
)
```

**Arguments**

|               |                                                                                                                                                                                                                                                                                                                                           |
|---------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| model_object  | A fitted BART or BCF model object of class bartmodel.                                                                                                                                                                                                                                                                                     |
| terms         | A character string specifying the model term(s) for which to compute intervals. Options for BART models are "mean_forest", "variance_forest", "rfx", or "y_hat".                                                                                                                                                                          |
| level         | A numeric value between 0 and 1 specifying the credible interval level (default is 0.95 for a 95% credible interval).                                                                                                                                                                                                                     |
| scale         | (Optional) Scale of mean function predictions. Options are "linear", which returns predictions on the original scale of the mean forest / RFX terms, and "probability", which transforms predictions into a probability of observing $y = 1$ . "probability" is only valid for models fit with a probit outcome model. Default: "linear". |
| X             | A matrix or data frame of covariates at which to compute the intervals. Required if the requested term depends on covariates (e.g., mean forest, variance forest, or overall predictions).                                                                                                                                                |
| leaf_basis    | An optional matrix of basis function evaluations for mean forest models with regression defined in the leaves. Required for "leaf regression" models.                                                                                                                                                                                     |
| rfx_group_ids | An optional vector of group IDs for random effects. Required if the requested term includes random effects.                                                                                                                                                                                                                               |
| rfx_basis     | An optional matrix of basis function evaluations for random effects. Required if the requested term includes random effects.                                                                                                                                                                                                              |

**Value**

A list containing the lower and upper bounds of the credible interval for the specified term. If multiple terms are requested, a named list with intervals for each term is returned.

**Examples**

```
n <- 100
p <- 5
X <- matrix(rnorm(n * p), nrow = n, ncol = p)
y <- 2 * X[,1] + rnorm(n)
bart_model <- bart(y_train = y, X_train = X)
intervals <- compute_bart_posterior_interval(
  model_object = bart_model,
  terms = c("mean_forest", "y_hat"),
  X = X,
  level = 0.90
)
```

---

```
compute_bcf_posterior_interval
```

*Compute BCF Posterior Credible Intervals*

---

## Description

Compute posterior credible intervals for specified terms from a fitted BCF model. Supports intervals for prognostic forests, CATE forests, variance forests, random effects, and overall mean outcome predictions.

## Usage

```
compute_bcf_posterior_interval(
  model_object,
  terms,
  level = 0.95,
  scale = "linear",
  X = NULL,
  Z = NULL,
  propensity = NULL,
  rfx_group_ids = NULL,
  rfx_basis = NULL
)
```

## Arguments

|                           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     |
|---------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>model_object</code> | A fitted BCF model object of class <code>bcfmodel</code> .                                                                                                                                                                                                                                                                                                                                                                                                                          |
| <code>terms</code>        | A character string specifying the model term(s) for which to compute intervals. Options for BCF models are "prognostic_function", "mu", "cate", "tau", "variance_forest", "rfx", or "y_hat". Note that "mu" is only different from "prognostic_function" if random effects are included with a model spec of "intercept_only" or "intercept_plus_treatment" and "tau" is only different from "cate" if random effects are included with a model spec of "intercept_plus_treatment". |
| <code>level</code>        | A numeric value between 0 and 1 specifying the credible interval level (default is 0.95 for a 95% credible interval).                                                                                                                                                                                                                                                                                                                                                               |
| <code>scale</code>        | (Optional) Scale of mean function predictions. Options are "linear", which returns predictions on the original scale of the mean forest / RFX terms, and "probability", which transforms predictions into a probability of observing $y = 1$ . "probability" is only valid for models fit with a probit outcome model. Default: "linear".                                                                                                                                           |
| <code>X</code>            | (Optional) A matrix or data frame of covariates at which to compute the intervals. Required if the requested term depends on covariates (e.g., prognostic forest, CATE forest, variance forest, or overall predictions).                                                                                                                                                                                                                                                            |
| <code>Z</code>            | (Optional) A vector or matrix of treatment assignments. Required if the requested term is "y_hat" (overall predictions).                                                                                                                                                                                                                                                                                                                                                            |

|               |                                                                                                                              |
|---------------|------------------------------------------------------------------------------------------------------------------------------|
| propensity    | (Optional) A vector or matrix of propensity scores. Required if the underlying model depends on user-provided propensities.  |
| rfx_group_ids | An optional vector of group IDs for random effects. Required if the requested term includes random effects.                  |
| rfx_basis     | An optional matrix of basis function evaluations for random effects. Required if the requested term includes random effects. |

**Value**

A list containing the lower and upper bounds of the credible interval for the specified term. If multiple terms are requested, a named list with intervals for each term is returned.

**Examples**

```
n <- 100
p <- 5
X <- matrix(rnorm(n * p), nrow = n, ncol = p)
pi_X <- pnorm(0.5 * X[,1])
Z <- rbinom(n, 1, pi_X)
mu_X <- X[,1]
tau_X <- 0.25 * X[,2]
y <- mu_X + tau_X * Z + rnorm(n)
bcf_model <- bcf(X_train = X, Z_train = Z, y_train = y,
                 propensity_train = pi_X)
intervals <- compute_bcf_posterior_interval(
  model_object = bcf_model,
  terms = c("prognostic_function", "cate"),
  X = X,
  Z = Z,
  propensity = pi_X,
  level = 0.90
)
```

---

compute\_contrast\_bart\_model

*Compute Contrast for BART Model*

---

**Description**

Compute a contrast using a BART model by making two sets of outcome predictions and taking their difference. This function provides the flexibility to compute any contrast of interest by specifying covariates, leaf basis, and random effects bases / IDs for both sides of a two term contrast. For simplicity, we refer to the subtrahend of the contrast as the "control" or  $Y_0$  term and the minuend of the contrast as the  $Y_1$  term, though the requested contrast need not match the "control vs treatment" terminology of a classic two-treatment causal inference problem. We mirror the function calls and terminology of the `predict.bartmodel` function, labeling each prediction data term with a 1 to denote its contribution to the treatment prediction of a contrast and 0 to denote inclusion in the control prediction.

Only valid when there is either a mean forest or a random effects term in the BART model.

**Usage**

```
compute_contrast_bart_model(
  object,
  X_0,
  X_1,
  leaf_basis_0 = NULL,
  leaf_basis_1 = NULL,
  rfx_group_ids_0 = NULL,
  rfx_group_ids_1 = NULL,
  rfx_basis_0 = NULL,
  rfx_basis_1 = NULL,
  type = "posterior",
  scale = "linear"
)
```

**Arguments**

|                 |                                                                                                                                                                                                                                                                             |
|-----------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| object          | Object of type bart containing draws of a regression forest and associated sampling outputs.                                                                                                                                                                                |
| X_0             | Covariates used for prediction in the "control" case. Must be a matrix or dataframe.                                                                                                                                                                                        |
| X_1             | Covariates used for prediction in the "treatment" case. Must be a matrix or dataframe.                                                                                                                                                                                      |
| leaf_basis_0    | (Optional) Bases used for prediction in the "control" case (by e.g. dot product with leaf values). Default: NULL.                                                                                                                                                           |
| leaf_basis_1    | (Optional) Bases used for prediction in the "treatment" case (by e.g. dot product with leaf values). Default: NULL.                                                                                                                                                         |
| rfx_group_ids_0 | (Optional) Test set group labels used for prediction from an additive random effects model in the "control" case. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set. Must be a vector.   |
| rfx_group_ids_1 | (Optional) Test set group labels used for prediction from an additive random effects model in the "treatment" case. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set. Must be a vector. |
| rfx_basis_0     | (Optional) Test set basis for used for prediction from an additive random effects model in the "control" case. Must be a matrix or vector.                                                                                                                                  |
| rfx_basis_1     | (Optional) Test set basis for used for prediction from an additive random effects model in the "treatment" case. Must be a matrix or vector.                                                                                                                                |
| type            | (Optional) Aggregation level of the contrast. Options are "mean", which averages the contrast evaluations over every draw of a BART model, and "posterior", which returns the entire matrix of posterior contrast estimates. Default: "posterior".                          |
| scale           | (Optional) Scale of the contrast. Options are "linear", which returns a contrast on the original scale of the mean forest / RFX terms, and "probability", which                                                                                                             |

transforms each contrast term into a probability of observing  $y == 1$  before taking their difference. "probability" is only valid for models fit with a probit outcome model. Default: "linear".

### Value

Contrast matrix or vector, depending on whether type = "mean" or "posterior".

### Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
W <- matrix(runif(n*1), ncol = 1)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5*W[,1]) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5*W[,1]) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5*W[,1]) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5*W[,1])
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
W_test <- W[test_inds,]
W_train <- W[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, leaf_basis_train = W_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
contrast_test <- compute_contrast_bart_model(
  bart_model,
  X_0 = X_test,
  X_1 = X_test,
  leaf_basis_0 = matrix(0, nrow = n_test, ncol = 1),
  leaf_basis_1 = matrix(1, nrow = n_test, ncol = 1),
  type = "posterior",
  scale = "linear"
)
```

## Description

Compute a contrast using a BCF model by making two sets of outcome predictions and taking their difference. For simple BCF models with binary treatment, this will yield the same prediction as requesting `terms = "cate"` in the `predict.bcfmodel` function. For more general models, such as models with continuous / multivariate treatments or an additive random effects term with a coefficient on the treatment, this function provides the flexibility to compute a any contrast of interest by specifying covariates, treatment, and random effects bases and IDs for both sides of a two term contrast. For simplicity, we refer to the subtrahend of the contrast as the "control" or  $Y_0$  term and the minuend of the contrast as the  $Y_1$  term, though the requested contrast need not match the "control vs treatment" terminology of a classic two-arm experiment. We mirror the function calls and terminology of the `predict.bcfmodel` function, labeling each prediction data term with a 1 to denote its contribution to the treatment prediction of a contrast and 0 to denote inclusion in the control prediction.

## Usage

```
compute_contrast_bcf_model(
  object,
  X_0,
  X_1,
  Z_0,
  Z_1,
  propensity_0 = NULL,
  propensity_1 = NULL,
  rfx_group_ids_0 = NULL,
  rfx_group_ids_1 = NULL,
  rfx_basis_0 = NULL,
  rfx_basis_1 = NULL,
  type = "posterior",
  scale = "linear"
)
```

## Arguments

|                           |                                                                                                                          |
|---------------------------|--------------------------------------------------------------------------------------------------------------------------|
| <code>object</code>       | Object of type <code>bcfmodel</code> containing draws of a Bayesian causal forest model and associated sampling outputs. |
| <code>X_0</code>          | Covariates used for prediction in the "control" case. Must be a matrix or dataframe.                                     |
| <code>X_1</code>          | Covariates used for prediction in the "treatment" case. Must be a matrix or dataframe.                                   |
| <code>Z_0</code>          | Treatments used for prediction in the "control" case. Must be a matrix or vector.                                        |
| <code>Z_1</code>          | Treatments used for prediction in the "treatment" case. Must be a matrix or vector.                                      |
| <code>propensity_0</code> | (Optional) Propensities used for prediction in the "control" case. Must be a matrix or vector.                           |
| <code>propensity_1</code> | (Optional) Propensities used for prediction in the "treatment" case. Must be a matrix or vector.                         |

|                 |                                                                                                                                                                                                                                                                                                                                                                   |
|-----------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| rfx_group_ids_0 | (Optional) Test set group labels used for prediction from an additive random effects model in the "control" case. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set. Must be a vector.                                                                                         |
| rfx_group_ids_1 | (Optional) Test set group labels used for prediction from an additive random effects model in the "treatment" case. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set. Must be a vector.                                                                                       |
| rfx_basis_0     | (Optional) Test set basis for used for prediction from an additive random effects model in the "control" case. Must be a matrix or vector.                                                                                                                                                                                                                        |
| rfx_basis_1     | (Optional) Test set basis for used for prediction from an additive random effects model in the "treatment" case. Must be a matrix or vector.                                                                                                                                                                                                                      |
| type            | (Optional) Aggregation level of the contrast. Options are "mean", which averages the contrast evaluations over every draw of a BCF model, and "posterior", which returns the entire matrix of posterior contrast estimates. Default: "posterior".                                                                                                                 |
| scale           | (Optional) Scale of the contrast. Options are "linear", which returns a contrast on the original scale of the mean forest / RFX terms, and "probability", which transforms each contrast term into a probability of observing $y == 1$ before taking their difference. "probability" is only valid for models fit with a probit outcome model. Default: "linear". |

## Value

List of prediction matrices or single prediction matrix / vector, depending on the terms requested.

## Examples

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
```

```

)
Z <- rbinom(n, 1, pi_x)
noise_sd <- 1
y <- mu_x + tau_x*Z + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
                 propensity_train = pi_train, num_gfr = 10,
                 num_burnin = 0, num_mcmc = 10)
tau_hat_test <- compute_contrast_bcf_model(
  bcf_model, X_0=X_test, X_1=X_test, Z_0=rep(0, n_test), Z_1=rep(1, n_test),
  propensity_0 = pi_test, propensity_1 = pi_test
)

```

---

convertPreprocessorToJson

*Convert Covariate Preprocessor to CppJson*

---

## Description

Convert the persistent aspects of a covariate preprocessor to (in-memory) C++ JSON object

## Usage

```
convertPreprocessorToJson(object)
```

## Arguments

|        |                                                                                                    |
|--------|----------------------------------------------------------------------------------------------------|
| object | List containing information on variables, including train set categories for categorical variables |
|--------|----------------------------------------------------------------------------------------------------|

## Value

wrapper around in-memory C++ JSON object

**Examples**

```
cov_mat <- matrix(1:12, ncol = 3)
preprocess_list <- preprocessTrainData(cov_mat)
preprocessor_json <- convertPreprocessorToJson(preprocess_list$metadata)
```

---

 CppJson

---

*JSON C++ Object Wrapper*


---

**Description**

Wrapper around a C++ `nlohmann::json` object

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Public fields**

`json_ptr` External pointer to a C++ `nlohmann::json` object  
`num_forests` Number of forests in the `nlohmann::json` object  
`forest_labels` Names of forest objects in the overall `nlohmann::json` object  
`num_rfx` Number of random effects terms in the `nlohman::json` object  
`rfx_container_labels` Names of rfx container objects in the overall `nlohmann::json` object  
`rfx_mapper_labels` Names of rfx label mapper objects in the overall `nlohmann::json` object  
`rfx_groupid_labels` Names of rfx group id objects in the overall `nlohmann::json` object

**Methods****Public methods:**

- `CppJson$new()`
- `CppJson$add_forest()`
- `CppJson$add_random_effects()`
- `CppJson$add_scalar()`
- `CppJson$add_integer()`
- `CppJson$add_boolean()`
- `CppJson$add_string()`
- `CppJson$add_vector()`
- `CppJson$add_integer_vector()`
- `CppJson$add_string_vector()`
- `CppJson$add_list()`
- `CppJson$add_string_list()`
- `CppJson$get_scalar()`

- `CppJson$get_integer()`
- `CppJson$get_boolean()`
- `CppJson$get_string()`
- `CppJson$get_vector()`
- `CppJson$get_integer_vector()`
- `CppJson$get_string_vector()`
- `CppJson$get_numeric_list()`
- `CppJson$get_string_list()`
- `CppJson$return_json_string()`
- `CppJson$save_file()`
- `CppJson$load_from_file()`
- `CppJson$load_from_string()`

**Method** `new()`: Create a new CppJson object.

*Usage:*

`CppJson$new()`

*Returns:* A new CppJson object.

**Method** `add_forest()`: Convert a forest container to json and add to the current CppJson object

*Usage:*

`CppJson$add_forest(forest_samples)`

*Arguments:*

`forest_samples` ForestSamples R class

*Returns:* None

**Method** `add_random_effects()`: Convert a random effects container to json and add to the current CppJson object

*Usage:*

`CppJson$add_random_effects(rfx_samples)`

*Arguments:*

`rfx_samples` RandomEffectSamples R class

*Returns:* None

**Method** `add_scalar()`: Add a scalar to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

`CppJson$add_scalar(field_name, field_value, subfolder_name = NULL)`

*Arguments:*

`field_name` The name of the field to be added to json

`field_value` Numeric value of the field to be added to json

`subfolder_name` (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_integer()`: Add a scalar to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$add_integer(field_name, field_value, subfolder_name = NULL)
```

*Arguments:*

`field_name` The name of the field to be added to json

`field_value` Integer value of the field to be added to json

`subfolder_name` (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_boolean()`: Add a boolean value to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$add_boolean(field_name, field_value, subfolder_name = NULL)
```

*Arguments:*

`field_name` The name of the field to be added to json

`field_value` Numeric value of the field to be added to json

`subfolder_name` (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_string()`: Add a string value to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$add_string(field_name, field_value, subfolder_name = NULL)
```

*Arguments:*

`field_name` The name of the field to be added to json

`field_value` Numeric value of the field to be added to json

`subfolder_name` (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_vector()`: Add a vector to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$add_vector(field_name, field_vector, subfolder_name = NULL)
```

*Arguments:*

`field_name` The name of the field to be added to json

`field_vector` Vector to be stored in json

`subfolder_name` (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_integer_vector()`: Add an integer vector to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$add_integer_vector(field_name, field_vector, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be added to json

field\_vector Vector to be stored in json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_string_vector()`: Add an array to the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$add_string_vector(field_name, field_vector, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be added to json

field\_vector Character vector to be stored in json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which to place the value

*Returns:* None

**Method** `add_list()`: Add a list of vectors (as an object map of arrays) to the json object under the name "field\_name"

*Usage:*

```
CppJson$add_list(field_name, field_list)
```

*Arguments:*

field\_name The name of the field to be added to json

field\_list List to be stored in json

*Returns:* None

**Method** `add_string_list()`: Add a list of vectors (as an object map of arrays) to the json object under the name "field\_name"

*Usage:*

```
CppJson$add_string_list(field_name, field_list)
```

*Arguments:*

field\_name The name of the field to be added to json

field\_list List to be stored in json

*Returns:* None

**Method** `get_scalar()`: Retrieve a scalar value from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_scalar(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_integer()`: Retrieve a integer value from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_integer(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_boolean()`: Retrieve a boolean value from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_boolean(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_string()`: Retrieve a string value from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_string(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_vector()`: Retrieve a vector from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_vector(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_integer_vector()`: Retrieve an integer vector from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_integer_vector(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_string_vector()`: Retrieve a character vector from the json object under the name "field\_name" (with optional subfolder "subfolder\_name")

*Usage:*

```
CppJson$get_string_vector(field_name, subfolder_name = NULL)
```

*Arguments:*

field\_name The name of the field to be accessed from json

subfolder\_name (Optional) Name of the subfolder / hierarchy under which the field is stored

*Returns:* None

**Method** `get_numeric_list()`: Reconstruct a list of numeric vectors from the json object stored under "field\_name"

*Usage:*

```
CppJson$get_numeric_list(field_name, key_names)
```

*Arguments:*

field\_name The name of the field to be added to json

key\_names Vector of names of list elements (each of which is a vector)

*Returns:* None

**Method** `get_string_list()`: Reconstruct a list of string vectors from the json object stored under "field\_name"

*Usage:*

```
CppJson$get_string_list(field_name, key_names)
```

*Arguments:*

field\_name The name of the field to be added to json

key\_names Vector of names of list elements (each of which is a vector)

*Returns:* None

**Method** `return_json_string()`: Convert a JSON object to in-memory string

*Usage:*

```
CppJson$return_json_string()
```

*Returns:* JSON string

**Method** `save_file()`: Save a json object to file

*Usage:*

```
CppJson$save_file(filename)
```

*Arguments:*

filename String of filepath, must end in ".json"

Returns: None

**Method** load\_from\_file(): Load a json object from file

Usage:

CppJson\$load\_from\_file(filename)

Arguments:

filename String of filepath, must end in ".json"

Returns: None

**Method** load\_from\_string(): Load a json object from string

Usage:

CppJson\$load\_from\_string(json\_string)

Arguments:

json\_string JSON string dump

Returns: None

---

CppRNG

Random Number Generator C++ Wrapper

---

## Description

Wrapper around a C++ random number generator object (for reproducibility). The class persists a C++ random number generator throughout an R session to ensure a given seed generates the same outputs (on the same OS). If no seed is provided, the C++ random number generator is initialized using `std::random_device`.

## Public fields

rng\_ptr External pointer to a C++ `std::mt19937` class

## Methods

### Public methods:

- [CppRNG\\$new\(\)](#)

**Method** new(): Create a new CppRNG object.

Usage:

CppRNG\$new(random\_seed = -1)

Arguments:

random\_seed (Optional) random seed for sampling

Returns: A new CppRNG object.

---

```
createBARTModelFromCombinedJson
```

*Convert JSON List to Single BART Model*

---

## Description

Convert a list of (in-memory) JSON representations of a BART model to a single combined BART model object which can be used for prediction, etc...

## Usage

```
createBARTModelFromCombinedJson(json_object_list)
```

## Arguments

```
json_object_list
```

List of objects of type CppJson containing Json representation of a BART model

## Value

Object of type bartmodel

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bart_json <- list(saveBARTModelToJson(bart_model))
bart_model_roundtrip <- createBARTModelFromCombinedJson(bart_json)
```

---

```
createBARTModelFromCombinedJsonString
```

*Convert JSON String List to Single BART Model*

---

**Description**

Convert a list of (in-memory) JSON strings that represent BART models to a single combined BART model object which can be used for prediction, etc...

**Usage**

```
createBARTModelFromCombinedJsonString(json_string_list)
```

**Arguments**

`json_string_list`  
List of JSON strings which can be parsed to objects of type CppJson containing  
Json representation of a BART model

**Value**

Object of type `bartmodel`

**Examples**

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bart_json_string_list <- list(saveBARTModelToJsonString(bart_model))
bart_model_roundtrip <- createBARTModelFromCombinedJsonString(bart_json_string_list)
```

---

```
createBARTModelFromJson
```

*Convert JSON to BART Model*

---

## Description

Convert an (in-memory) JSON representation of a BART model to a BART model object which can be used for prediction, etc...

## Usage

```
createBARTModelFromJson(json_object)
```

## Arguments

`json_object`      Object of type CppJson containing Json representation of a BART model

## Value

Object of type `bartmodel`

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bart_json <- saveBARTModelToJson(bart_model)
bart_model_roundtrip <- createBARTModelFromJson(bart_json)
```

---

```
createBARTModelFromJsonFile
```

*Convert JSON File to BART Model*

---

## Description

Convert a JSON file containing sample information on a trained BART model to a BART model object which can be used for prediction, etc...

## Usage

```
createBARTModelFromJsonFile(json_filename)
```

## Arguments

`json_filename` String of filepath, must end in ".json"

## Value

Object of type `bartmodel`

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
tmpjson <- tempfile(fileext = ".json")
saveBARTModelToJsonFile(bart_model, file.path(tmpjson))
bart_model_roundtrip <- createBARTModelFromJsonFile(file.path(tmpjson))
unlink(tmpjson)
```

---

```
createBARTModelFromJsonString
```

*Convert JSON String to BART Model*

---

## Description

Convert a JSON string containing sample information on a trained BART model to a BART model object which can be used for prediction, etc...

## Usage

```
createBARTModelFromJsonString(json_string)
```

## Arguments

json\_string      JSON string dump

## Value

Object of type bartmodel

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bart_json <- saveBARTModelToJsonString(bart_model)
bart_model_roundtrip <- createBARTModelFromJsonString(bart_json)
y_hat_mean_roundtrip <- rowMeans(predict(bart_model_roundtrip, X=X_train)$y_hat)
```

---

```
createBCFModelFromCombinedJson
```

*Convert JSON List to BCF Model*

---

### Description

Convert a list of (in-memory) JSON strings that represent BCF models to a single combined BCF model object which can be used for prediction, etc...

### Usage

```
createBCFModelFromCombinedJson(json_object_list)
```

### Arguments

`json_object_list`  
List of objects of type CppJson containing Json representation of a BCF model

### Value

Object of type `bcfmodel`

### Examples

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
```

```

rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bcf_json_list <- list(saveBCFModelToJson(bcf_model))
bcf_model_roundtrip <- createBCFModelFromCombinedJson(bcf_json_list)

```

---

```
createBCFModelFromCombinedJsonString
```

*Convert JSON String List to BCF Model*

---

## Description

Convert a list of (in-memory) JSON strings that represent BCF models to a single combined BCF model object which can be used for prediction, etc...

## Usage

```
createBCFModelFromCombinedJsonString(json_string_list)
```

**Arguments**

json\_string\_list

List of JSON strings which can be parsed to objects of type CppJson containing  
Json representation of a BCF model

**Value**

Object of type bcfmodel

**Examples**

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
```

```

y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bcf_json_string_list <- list(saveBCFModelToJsonString(bcf_model))
bcf_model_roundtrip <- createBCFModelFromCombinedJsonString(bcf_json_string_list)

```

---

createBCFModelFromJson

*Convert JSON to BCF Model*

---

## Description

Convert an (in-memory) JSON representation of a BCF model to a BCF model object which can be used for prediction, etc...

## Usage

```
createBCFModelFromJson(json_object)
```

## Arguments

`json_object`      Object of type CppJson containing Json representation of a BCF model

## Value

Object of type `bcfmodel`

## Examples

```

n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +

```

```

      ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
      ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
      ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
    )
    pi_x <- (
      ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
      ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
      ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
      ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
    )
    tau_x <- (
      ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
      ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
      ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
      ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
    )
    Z <- rbinom(n, 1, pi_x)
    E_XZ <- mu_x + Z*tau_x
    snr <- 3
    rfx_group_ids <- rep(c(1,2), n %% 2)
    rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
    rfx_basis <- cbind(1, runif(n, -1, 1))
    rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
    y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
    test_set_pct <- 0.2
    n_test <- round(test_set_pct*n)
    n_train <- n - n_test
    test_inds <- sort(sample(1:n, n_test, replace = FALSE))
    train_inds <- (1:n)[!((1:n) %in% test_inds)]
    X_test <- X[test_inds,]
    X_train <- X[train_inds,]
    pi_test <- pi_x[test_inds]
    pi_train <- pi_x[train_inds]
    Z_test <- Z[test_inds]
    Z_train <- Z[train_inds]
    y_test <- y[test_inds]
    y_train <- y[train_inds]
    mu_test <- mu_x[test_inds]
    mu_train <- mu_x[train_inds]
    tau_test <- tau_x[test_inds]
    tau_train <- tau_x[train_inds]
    rfx_group_ids_test <- rfx_group_ids[test_inds]
    rfx_group_ids_train <- rfx_group_ids[train_inds]
    rfx_basis_test <- rfx_basis[test_inds,]
    rfx_basis_train <- rfx_basis[train_inds,]
    rfx_term_test <- rfx_term[test_inds]
    rfx_term_train <- rfx_term[train_inds]
    mu_params <- list(sample_sigma2_leaf = TRUE)
    tau_params <- list(sample_sigma2_leaf = FALSE)
    bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
      propensity_train = pi_train,
      rfx_group_ids_train = rfx_group_ids_train,
      rfx_basis_train = rfx_basis_train, X_test = X_test,

```

```

      Z_test = Z_test, propensity_test = pi_test,
      rfx_group_ids_test = rfx_group_ids_test,
      rfx_basis_test = rfx_basis_test,
      num_gfr = 10, num_burnin = 0, num_mcmc = 10,
      prognostic_forest_params = mu_params,
      treatment_effect_forest_params = tau_params)
bcf_json <- saveBCFModelToJson(bcf_model)
bcf_model_roundtrip <- createBCFModelFromJson(bcf_json)

```

---

```
createBCFModelFromJsonFile
```

*Convert JSON File to BCF Model*

---

## Description

Convert a JSON file containing sample information on a trained BCF model to a BCF model object which can be used for prediction, etc...

## Usage

```
createBCFModelFromJsonFile(json_filename)
```

## Arguments

json\_filename    String of filepath, must end in ".json"

## Value

Object of type bcfmodel

## Examples

```

n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +

```

```

      ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
      ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
    )
    Z <- rbinom(n, 1, pi_x)
    E_XZ <- mu_x + Z*tau_x
    snr <- 3
    rfx_group_ids <- rep(c(1,2), n %/% 2)
    rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
    rfx_basis <- cbind(1, runif(n, -1, 1))
    rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
    y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
    test_set_pct <- 0.2
    n_test <- round(test_set_pct*n)
    n_train <- n - n_test
    test_inds <- sort(sample(1:n, n_test, replace = FALSE))
    train_inds <- (1:n)[!((1:n) %in% test_inds)]
    X_test <- X[test_inds,]
    X_train <- X[train_inds,]
    pi_test <- pi_x[test_inds]
    pi_train <- pi_x[train_inds]
    Z_test <- Z[test_inds]
    Z_train <- Z[train_inds]
    y_test <- y[test_inds]
    y_train <- y[train_inds]
    mu_test <- mu_x[test_inds]
    mu_train <- mu_x[train_inds]
    tau_test <- tau_x[test_inds]
    tau_train <- tau_x[train_inds]
    rfx_group_ids_test <- rfx_group_ids[test_inds]
    rfx_group_ids_train <- rfx_group_ids[train_inds]
    rfx_basis_test <- rfx_basis[test_inds,]
    rfx_basis_train <- rfx_basis[train_inds,]
    rfx_term_test <- rfx_term[test_inds]
    rfx_term_train <- rfx_term[train_inds]
    mu_params <- list(sample_sigma2_leaf = TRUE)
    tau_params <- list(sample_sigma2_leaf = FALSE)
    bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
      propensity_train = pi_train,
      rfx_group_ids_train = rfx_group_ids_train,
      rfx_basis_train = rfx_basis_train, X_test = X_test,
      Z_test = Z_test, propensity_test = pi_test,
      rfx_group_ids_test = rfx_group_ids_test,
      rfx_basis_test = rfx_basis_test,
      num_gfr = 10, num_burnin = 0, num_mcmc = 10,
      prognostic_forest_params = mu_params,
      treatment_effect_forest_params = tau_params)
    tmpjson <- tempfile(fileext = ".json")
    saveBCFModelToJsonFile(bcf_model, file.path(tmpjson))
    bcf_model_roundtrip <- createBCFModelFromJsonFile(file.path(tmpjson))
    unlink(tmpjson)

```

---

```
createBCFModelFromJsonString
```

*Convert JSON String to BCF Model*

---

**Description**

Convert a JSON string containing sample information on a trained BCF model to a BCF model object which can be used for prediction, etc...

**Usage**

```
createBCFModelFromJsonString(json_string)
```

**Arguments**

json\_string      JSON string dump

**Value**

Object of type bcfmodel

**Examples**

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
```

```

rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bcf_json <- saveBCFModelToJsonString(bcf_model)
bcf_model_roundtrip <- createBCFModelFromJsonString(bcf_json)

```

---

createCppJson

*Create CppJson Object*


---

## Description

Create a new (empty) C++ Json object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Usage

```
createCppJson()
```

**Value**

CppJson object

**Examples**

```
example_vec <- runif(10)
example_json <- createCppJson()
example_json$add_vector("myvec", example_vec)
```

---

|                   |                                        |
|-------------------|----------------------------------------|
| createCppJsonFile | <i>Create CppJson Object from File</i> |
|-------------------|----------------------------------------|

---

**Description**

Create a C++ Json object from a Json file

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createCppJsonFile(json_filename)
```

**Arguments**

json\_filename    Name of file to read. Must end in .json.

**Value**

CppJson object

**Examples**

```
example_vec <- runif(10)
example_json <- createCppJson()
example_json$add_vector("myvec", example_vec)
tmpjson <- tempfile(fileext = ".json")
example_json$save_file(file.path(tmpjson))
example_json_roundtrip <- createCppJsonFile(file.path(tmpjson))
unlink(tmpjson)
```

---

|                     |                                          |
|---------------------|------------------------------------------|
| createCppJsonString | <i>Create CppJson Object from String</i> |
|---------------------|------------------------------------------|

---

**Description**

Create a C++ Json object from a Json string

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createCppJsonString(json_string)
```

**Arguments**

|             |                  |
|-------------|------------------|
| json_string | JSON string dump |
|-------------|------------------|

**Value**

CppJson object

**Examples**

```
example_vec <- runif(10)
example_json <- createCppJson()
example_json$add_vector("myvec", example_vec)
example_json_string <- example_json$return_json_string()
example_json_roundtrip <- createCppJsonString(example_json_string)
```

---

|              |                             |
|--------------|-----------------------------|
| createCppRNG | <i>Create CppRNG Object</i> |
|--------------|-----------------------------|

---

**Description**

Create an R class that wraps a C++ random number generator

**Usage**

```
createCppRNG(random_seed = -1)
```

**Arguments**

|             |                                     |
|-------------|-------------------------------------|
| random_seed | (Optional) random seed for sampling |
|-------------|-------------------------------------|

**Value**

CppRng object

**Examples**

```
rng <- createCppRNG(1234)
rng <- createCppRNG()
```

---

|              |                             |
|--------------|-----------------------------|
| createForest | <i>Create Forest Object</i> |
|--------------|-----------------------------|

---

**Description**

Create a forest

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createForest(
  num_trees,
  leaf_dimension = 1,
  is_leaf_constant = FALSE,
  is_exponentiated = FALSE
)
```

**Arguments**

|                  |                                                                          |
|------------------|--------------------------------------------------------------------------|
| num_trees        | Number of trees in the forest                                            |
| leaf_dimension   | Dimensionality of the outcome model                                      |
| is_leaf_constant | Whether leaf is constant                                                 |
| is_exponentiated | Whether forest predictions should be exponentiated before being returned |

**Value**

Forest object

**Examples**

```
num_trees <- 100
leaf_dimension <- 2
is_leaf_constant <- FALSE
is_exponentiated <- FALSE
forest <- createForest(num_trees, leaf_dimension, is_leaf_constant, is_exponentiated)
```

---

|                     |                             |
|---------------------|-----------------------------|
| createForestDataset | Create ForestDataset Object |
|---------------------|-----------------------------|

---

### Description

Create a forest dataset object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

### Usage

```
createForestDataset(covariates, basis = NULL, variance_weights = NULL)
```

### Arguments

|                  |                                                             |
|------------------|-------------------------------------------------------------|
| covariates       | Matrix of covariates                                        |
| basis            | (Optional) Matrix of bases used to define a leaf regression |
| variance_weights | (Optional) Vector of observation-specific variance weights  |

### Value

ForestDataset object

### Examples

```
covariate_matrix <- matrix(runif(10*100), ncol = 10)
basis_matrix <- matrix(rnorm(3*100), ncol = 3)
weight_vector <- rnorm(100)
forest_dataset <- createForestDataset(covariate_matrix)
forest_dataset <- createForestDataset(covariate_matrix, basis_matrix)
forest_dataset <- createForestDataset(covariate_matrix, basis_matrix, weight_vector)
```

---

|                   |                           |
|-------------------|---------------------------|
| createForestModel | Create ForestModel Object |
|-------------------|---------------------------|

---

### Description

Create a forest model object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createForestModel(forest_dataset, forest_model_config, global_model_config)
```

**Arguments**

`forest_dataset` ForestDataset object, used to initialize forest sampling data structures

`forest_model_config` ForestModelConfig object containing forest model parameters and settings

`global_model_config` GlobalModelConfig object containing global model parameters and settings

**Value**

ForestModel object

**Examples**

```
num_trees <- 100
n <- 100
p <- 10
alpha <- 0.95
beta <- 2.0
min_samples_leaf <- 2
max_depth <- 10
feature_types <- as.integer(rep(0, p))
X <- matrix(runif(n*p), ncol = p)
forest_dataset <- createForestDataset(X)
forest_model_config <- createForestModelConfig(feature_types=feature_types,
                                              num_trees=num_trees, num_features=p,
                                              num_observations=n, alpha=alpha, beta=beta,
                                              min_samples_leaf=min_samples_leaf,
                                              max_depth=max_depth, leaf_model_type=1)
global_model_config <- createGlobalModelConfig(global_error_variance=1.0)
forest_model <- createForestModel(forest_dataset, forest_model_config, global_model_config)
```

---

```
createForestModelConfig
```

*Create ForestModelConfig Object*

---

**Description**

Create a forest model config object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](https://stochtree.ai)

**Usage**

```
createForestModelConfig(
  feature_types = NULL,
  sweep_update_indices = NULL,
  num_trees = NULL,
  num_features = NULL,
  num_observations = NULL,
  variable_weights = NULL,
  leaf_dimension = 1,
  alpha = 0.95,
  beta = 2,
  min_samples_leaf = 5,
  max_depth = -1,
  leaf_model_type = 1,
  leaf_model_scale = NULL,
  variance_forest_shape = 1,
  variance_forest_scale = 1,
  cutpoint_grid_size = 100,
  num_features_subsample = NULL
)
```

**Arguments**

|                                   |                                                                                                                                                        |
|-----------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>feature_types</code>        | Vector of integer-coded feature types (integers where 0 = numeric, 1 = ordered categorical, 2 = unordered categorical)                                 |
| <code>sweep_update_indices</code> | Vector of (0-indexed) indices of trees to update in a sweep                                                                                            |
| <code>num_trees</code>            | Number of trees in the forest being sampled                                                                                                            |
| <code>num_features</code>         | Number of features in training dataset                                                                                                                 |
| <code>num_observations</code>     | Number of observations in training dataset                                                                                                             |
| <code>variable_weights</code>     | Vector specifying sampling probability for all p covariates in ForestDataset                                                                           |
| <code>leaf_dimension</code>       | Dimension of the leaf model (default: 1)                                                                                                               |
| <code>alpha</code>                | Root node split probability in tree prior (default: 0.95)                                                                                              |
| <code>beta</code>                 | Depth prior penalty in tree prior (default: 2.0)                                                                                                       |
| <code>min_samples_leaf</code>     | Minimum number of samples in a tree leaf (default: 5)                                                                                                  |
| <code>max_depth</code>            | Maximum depth of any tree in the ensemble in the model. Setting to -1 does not enforce any depth limits on trees. Default: -1.                         |
| <code>leaf_model_type</code>      | Integer specifying the leaf model type (0 = constant leaf, 1 = univariate leaf regression, 2 = multivariate leaf regression). Default: 0.              |
| <code>leaf_model_scale</code>     | Scale parameter used in Gaussian leaf models (can either be a scalar or a q x q matrix, where q is the dimensionality of the basis and is only >1 when |

leaf\_model\_int = 2). Calibrated internally as  $1/\text{num\_trees}$ , propagated along diagonal if needed for multivariate leaf models.

variance\_forest\_shape  
Shape parameter for IG leaf models (applicable when leaf\_model\_type = 3). Default: 1.

variance\_forest\_scale  
Scale parameter for IG leaf models (applicable when leaf\_model\_type = 3). Default: 1.

cutpoint\_grid\_size  
Number of unique cutpoints to consider (default: 100)

num\_features\_subsample  
Number of features to subsample for the GFR algorithm

**Value**

ForestModelConfig object

**Examples**

```
config <- createForestModelConfig(num_trees = 10, num_features = 5, num_observations = 100)
```

---

createForestSamples     *Create ForestSamples Object*

---

**Description**

Create a container of forest samples

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](https://stochtree.ai)

**Usage**

```
createForestSamples(
  num_trees,
  leaf_dimension = 1,
  is_leaf_constant = FALSE,
  is_exponentiated = FALSE
)
```

**Arguments**

num\_trees            Number of trees

leaf\_dimension      Dimensionality of the outcome model

is\_leaf\_constant    Whether leaf is constant

is\_exponentiated    Whether forest predictions should be exponentiated before being returned

**Value**

ForestSamples object

**Examples**

```
num_trees <- 100
leaf_dimension <- 2
is_leaf_constant <- FALSE
is_exponentiated <- FALSE
forest_samples <- createForestSamples(num_trees, leaf_dimension, is_leaf_constant, is_exponentiated)
```

---

`createGlobalModelConfig`*Create GlobalModelConfig Object*

---

**Description**

Create a global model config object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createGlobalModelConfig(global_error_variance = 1)
```

**Arguments**

`global_error_variance`  
Global error variance parameter (default: 1.0)

**Value**

GlobalModelConfig object

**Examples**

```
config <- createGlobalModelConfig(global_error_variance = 100)
```

---

|               |                              |
|---------------|------------------------------|
| createOutcome | <i>Create Outcome Object</i> |
|---------------|------------------------------|

---

### Description

Create an outcome object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

### Usage

```
createOutcome(outcome)
```

### Arguments

|         |                          |
|---------|--------------------------|
| outcome | Vector of outcome values |
|---------|--------------------------|

### Value

Outcome object

### Examples

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
outcome <- createOutcome(y)
```

---

|                            |                                                       |
|----------------------------|-------------------------------------------------------|
| createPreprocessorFromJson | <i>Reload Covariate Preprocessor from JSON String</i> |
|----------------------------|-------------------------------------------------------|

---

### Description

Reload a covariate preprocessor object from a JSON string containing a serialized preprocessor

### Usage

```
createPreprocessorFromJson(json_object)
```

### Arguments

|             |                                                                                     |
|-------------|-------------------------------------------------------------------------------------|
| json_object | in-memory wrapper around JSON C++ object containing covariate preprocessor metadata |
|-------------|-------------------------------------------------------------------------------------|

**Value**

Preprocessor object that can be used with the preprocessPredictionData function

**Examples**

```
cov_mat <- matrix(1:12, ncol = 3)
preprocess_list <- preprocessTrainData(cov_mat)
preprocessor_json <- convertPreprocessorToJson(preprocess_list$metadata)
preprocessor_roundtrip <- createPreprocessorFromJson(preprocessor_json)
```

---

`createPreprocessorFromJsonString`

*Reload Covariate Preprocessor from JSON String*

---

**Description**

Reload a covariate preprocessor object from a JSON string containing a serialized preprocessor

**Usage**

```
createPreprocessorFromJsonString(json_string)
```

**Arguments**

`json_string`      in-memory JSON string containing covariate preprocessor metadata

**Value**

Preprocessor object that can be used with the preprocessPredictionData function

**Examples**

```
cov_mat <- matrix(1:12, ncol = 3)
preprocess_list <- preprocessTrainData(cov_mat)
preprocessor_json_string <- savePreprocessorToJsonString(preprocess_list$metadata)
preprocessor_roundtrip <- createPreprocessorFromJsonString(preprocessor_json_string)
```

`createRandomEffectSamples`*Create RandomEffectSamples Object*

---

**Description**

Create a RandomEffectSamples object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createRandomEffectSamples(num_components, num_groups, random_effects_tracker)
```

**Arguments**

`num_components` Number of "components" or bases defining the random effects regression

`num_groups` Number of random effects groups

`random_effects_tracker`  
Object of type RandomEffectsTracker

**Value**

RandomEffectSamples object

**Examples**

```
n <- 100
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_tracker <- createRandomEffectsTracker(rfx_group_ids)
rfx_samples <- createRandomEffectSamples(num_components, num_groups, rfx_tracker)
```

---

`createRandomEffectsDataset`*Create RandomEffectsDataset Object*

---

**Description**

Create a random effects dataset object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createRandomEffectsDataset(group_labels, basis, variance_weights = NULL)
```

**Arguments**

|                               |                                                                                                                   |
|-------------------------------|-------------------------------------------------------------------------------------------------------------------|
| <code>group_labels</code>     | Vector of group labels                                                                                            |
| <code>basis</code>            | Matrix of bases used to define the random effects regression (for an intercept-only model, pass an array of ones) |
| <code>variance_weights</code> | (Optional) Vector of observation-specific variance weights                                                        |

**Value**

RandomEffectsDataset object

**Examples**

```
rfx_group_ids <- sample(1:2, size = 100, replace = TRUE)
rfx_basis <- matrix(rnorm(3*100), ncol = 3)
weight_vector <- rnorm(100)
rfx_dataset <- createRandomEffectsDataset(rfx_group_ids, rfx_basis)
rfx_dataset <- createRandomEffectsDataset(rfx_group_ids, rfx_basis, weight_vector)
```

---

`createRandomEffectsModel`*Create RandomEffectsModel Object*

---

**Description**

Create a RandomEffectsModel object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createRandomEffectsModel(num_components, num_groups)
```

**Arguments**

num\_components    Number of "components" or bases defining the random effects regression  
 num\_groups        Number of random effects groups

**Value**

RandomEffectsModel object

**Examples**

```
n <- 100
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_model <- createRandomEffectsModel(num_components, num_groups)
```

---

```
createRandomEffectsTracker
```

*Create RandomEffectsTracker Object*

---

**Description**

Create a RandomEffectsTracker object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
createRandomEffectsTracker(rfx_group_indices)
```

**Arguments**

rfx\_group\_indices  
                     Integer indices indicating groups used to define random effects

**Value**

RandomEffectsTracker object

**Examples**

```
n <- 100
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_tracker <- createRandomEffectsTracker(rfx_group_ids)
```

---

|                   |                                                   |
|-------------------|---------------------------------------------------|
| extract_parameter | <i>Extract Parameter Samples Generic Function</i> |
|-------------------|---------------------------------------------------|

---

**Description**

Generic function for extracting parameter samples from a model object (BCF, BART, etc...)

**Usage**

```
extract_parameter(object, term)
```

**Arguments**

|        |                                                                        |
|--------|------------------------------------------------------------------------|
| object | Fitted model object from which to extract parameter samples            |
| term   | Name of the parameter to extract (e.g., "sigma2", "y_hat_train", etc.) |

**Value**

Parameter sample array

**Examples**

```
n <- 100
p <- 10
X <- matrix(runif(n*p), ncol = p)
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- rep(1.0, n)
y <- (-5 + 10*(X[,1] > 0.5)) + (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
bart_model <- bart(X_train=X, y_train=y, rfx_group_ids_train=rfx_group_ids,
  rfx_basis_train = rfx_basis, num_gfr=0, num_mcmc=10)
sigma2_samples <- extract_parameter(bart_model, "sigma2")
```

---

```
extract_parameter.bartmodel
```

*Extract BART Parameter Samples.*

---

## Description

Extract a vector, matrix or array of parameter samples from a BART model by name. Random effects are handled by a separate `getRandomEffectSamples` function due to the complexity of the random effects parameters. If the requested model term is not found, an error is thrown. The following conventions are used for parameter names:

- Global error variance: "sigma2", "global\_error\_scale", "sigma2\_global"
- Leaf scale: "sigma2\_leaf", "leaf\_scale"
- In-sample mean function predictions: "y\_hat\_train"
- Test set mean function predictions: "y\_hat\_test"
- In-sample variance forest predictions: "sigma2\_x\_train", "var\_x\_train"
- Test set variance forest predictions: "sigma2\_x\_test", "var\_x\_test"

## Usage

```
## S3 method for class 'bartmodel'
extract_parameter(object, term)
```

## Arguments

|                     |                                                                                                         |
|---------------------|---------------------------------------------------------------------------------------------------------|
| <code>object</code> | Object of type <code>bartmodel</code> containing draws of a BART model and associated sampling outputs. |
| <code>term</code>   | Name of the parameter to extract (e.g., "sigma2", "y_hat_train", etc.)                                  |

## Value

Array of parameter samples. If the underlying parameter is a scalar, this will be a vector of length `num_samples`. If the underlying parameter is vector-valued, this will be (`parameter_dimension` x `num_samples`) matrix, and if the underlying parameter is multidimensional, this will be an array of dimension (`parameter_dimension_1` x `parameter_dimension_2` x ... x `num_samples`).

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
```

```

)
snr <- 3
group_ids <- rep(c(1,2), n %% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[group_ids,] * rfx_basis)
E_y <- f_XW + rfx_term
y <- E_y + rnorm(n, 0, 1)*(sd(E_y)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
rfx_group_ids_test <- group_ids[test_inds]
rfx_group_ids_train <- group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train, X_test = X_test,
                  rfx_group_ids_train = rfx_group_ids_train,
                  rfx_group_ids_test = rfx_group_ids_test,
                  rfx_basis_train = rfx_basis_train,
                  rfx_basis_test = rfx_basis_test,
                  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
sigma2_samples <- extract_parameter(bart_model, "sigma2")

```

---

```
extract_parameter.bcfmodel
```

*Extract BCF Parameter Samples*

---

## Description

Extract a vector, matrix or array of parameter samples from a BCF model by name. Random effects are handled by a separate `getRandomEffectSamples` function due to the complexity of the random effects parameters. If the requested model term is not found, an error is thrown. The following conventions are used for parameter names:

- Global error variance: "sigma2", "global\_error\_scale", "sigma2\_global"
- Prognostic forest leaf scale: "sigma2\_leaf\_mu", "leaf\_scale\_mu", "mu\_leaf\_scale"
- Treatment effect forest leaf scale: "sigma2\_leaf\_tau", "leaf\_scale\_tau", "tau\_leaf\_scale"
- Adaptive coding parameters: "adaptive\_coding" (returns both the control and treated parameters jointly, with control in the first row and treated in the second row)
- In-sample mean function predictions: "y\_hat\_train"

- Test set mean function predictions: "y\_hat\_test"
- In-sample treatment effect forest predictions: "tau\_hat\_train"
- Test set treatment effect forest predictions: "tau\_hat\_test"
- In-sample variance forest predictions: "sigma2\_x\_train", "var\_x\_train"
- Test set variance forest predictions: "sigma2\_x\_test", "var\_x\_test"

## Usage

```
## S3 method for class 'bcfmodel'
extract_parameter(object, term)
```

## Arguments

|        |                                                                                          |
|--------|------------------------------------------------------------------------------------------|
| object | Object of type bcfmodel containing draws of a BCF model and associated sampling outputs. |
| term   | Name of the parameter to extract (e.g., "sigma2", "y_hat_train", etc.)                   |

## Value

Array of parameter samples. If the underlying parameter is a scalar, this will be a vector of length num\_samples. If the underlying parameter is vector-valued, this will be (parameter\_dimension x num\_samples) matrix, and if the underlying parameter is multidimensional, this will be an array of dimension (parameter\_dimension\_1 x parameter\_dimension\_2 x ... x num\_samples).

## Examples

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
```

```

rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
bcf_model <- bcf(X_train = X, y_train = y, Z_train = Z,
                 rfx_group_ids_train = rfx_group_ids,
                 rfx_basis_train = rfx_basis,
                 num_gfr = 10, num_burnin = 0, num_mcmc = 10)
sigma2_samples <- extract_parameter(bcf_model, "sigma2")

```

---

Forest

*Forest C++ Wrapper*


---

## Description

Wrapper around a C++ class that stores a single ensemble of decision trees (often treated as the "active forest" / current state of a forest term in a sampling loop in R)

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Public fields

`forest_ptr` External pointer to a C++ `TreeEnsemble` class

`internal_forest_is_empty` Whether the forest has not yet been "initialized" such that its `predict` function can be called.

## Methods

### Public methods:

- `Forest$new()`
- `Forest$merge_forest()`
- `Forest$add_constant()`
- `Forest$multiply_constant()`
- `Forest$predict()`
- `Forest$predict_raw()`
- `Forest$set_root_leaves()`
- `Forest$prepare_for_sampler()`
- `Forest$adjust_residual()`
- `Forest$num_trees()`
- `Forest$leaf_dimension()`
- `Forest$is_constant_leaf()`
- `Forest$is_exponentiated()`

- `Forest$add_numeric_split_tree()`
- `Forest$get_tree_leaves()`
- `Forest$get_tree_split_counts()`
- `Forest$get_forest_split_counts()`
- `Forest$tree_max_depth()`
- `Forest$average_max_depth()`
- `Forest$is_empty()`

**Method** `new()`: Create a new Forest object.

*Usage:*

```
Forest$new(
  num_trees,
  leaf_dimension = 1,
  is_leaf_constant = FALSE,
  is_exponentiated = FALSE
)
```

*Arguments:*

`num_trees` Number of trees in the forest

`leaf_dimension` Dimensionality of the outcome model

`is_leaf_constant` Whether leaf is constant

`is_exponentiated` Whether forest predictions should be exponentiated before being returned

*Returns:* A new Forest object.

**Method** `merge_forest()`: Create a larger forest by merging the trees of this forest with those of another forest

*Usage:*

```
Forest$merge_forest(forest)
```

*Arguments:*

`forest` Forest to be merged into this forest

**Method** `add_constant()`: Add a constant value to every leaf of every tree in an ensemble. If leaves are multi-dimensional, `constant_value` will be added to every dimension of the leaves.

*Usage:*

```
Forest$add_constant(constant_value)
```

*Arguments:*

`constant_value` Value that will be added to every leaf of every tree

**Method** `multiply_constant()`: Multiply every leaf of every tree by a constant value. If leaves are multi-dimensional, `constant_multiple` will be multiplied through every dimension of the leaves.

*Usage:*

```
Forest$multiply_constant(constant_multiple)
```

*Arguments:*

`constant_multiple` Value that will be multiplied by every leaf of every tree

**Method** `predict()`: Predict forest on every sample in `forest_dataset`

*Usage:*

```
Forest$predict(forest_dataset)
```

*Arguments:*

`forest_dataset` `ForestDataset` R class

*Returns:* vector of predictions with as many rows as in `forest_dataset`

**Method** `predict_raw()`: Predict "raw" leaf values (without being multiplied by basis) for every sample in `forest_dataset`

*Usage:*

```
Forest$predict_raw(forest_dataset)
```

*Arguments:*

`forest_dataset` `ForestDataset` R class

*Returns:* Array of predictions for each observation in `forest_dataset` and each sample in the `ForestSamples` class with each prediction having the dimensionality of the forests' leaf model. In the case of a constant leaf model or univariate leaf regression, this array is a vector (length is the number of observations). In the case of a multivariate leaf regression, this array is a matrix (number of observations by leaf model dimension, number of samples).

**Method** `set_root_leaves()`: Set a constant predicted value for every tree in the ensemble. Stops program if any tree is more than a root node.

*Usage:*

```
Forest$set_root_leaves(leaf_value)
```

*Arguments:*

`leaf_value` Constant leaf value(s) to be fixed for each tree in the ensemble indexed by `forest_num`. Can be either a single number or a vector, depending on the forest's leaf dimension.

**Method** `prepare_for_sampler()`: Set a constant predicted value for every tree in the ensemble. Stops program if any tree is more than a root node.

*Usage:*

```
Forest$prepare_for_sampler(
  dataset,
  outcome,
  forest_model,
  leaf_model_int,
  leaf_value
)
```

*Arguments:*

`dataset` `ForestDataset` `Dataset` class (covariates, basis, etc...)

`outcome` `Outcome` `Outcome` class (residual / partial residual)

`forest_model` `ForestModel` object storing tracking structures used in training / sampling

**leaf\_model\_int** Integer value encoding the leaf model type (0 = constant gaussian, 1 = univariate gaussian, 2 = multivariate gaussian, 3 = log linear variance).

**leaf\_value** Constant leaf value(s) to be fixed for each tree in the ensemble indexed by **forest\_num**. Can be either a single number or a vector, depending on the forest's leaf dimension.

**Method** `adjust_residual()`: Adjusts residual based on the predictions of a forest

This is typically run just once at the beginning of a forest sampling algorithm. After trees are initialized with constant root node predictions, their root predictions are subtracted out of the residual.

*Usage:*

```
Forest$adjust_residual(dataset, outcome, forest_model, requires_basis, add)
```

*Arguments:*

**dataset** ForestDataset object storing the covariates and bases for a given forest

**outcome** Outcome object storing the residuals to be updated based on forest predictions

**forest\_model** ForestModel object storing tracking structures used in training / sampling

**requires\_basis** Whether or not a forest requires a basis for prediction

**add** Whether forest predictions should be added to or subtracted from residuals

**Method** `num_trees()`: Return number of trees in each ensemble of a Forest object

*Usage:*

```
Forest$num_trees()
```

*Returns:* Tree count

**Method** `leaf_dimension()`: Return output dimension of trees in a Forest object

*Usage:*

```
Forest$leaf_dimension()
```

*Returns:* Leaf node parameter size

**Method** `is_constant_leaf()`: Return constant leaf status of trees in a Forest object

*Usage:*

```
Forest$is_constant_leaf()
```

*Returns:* TRUE if leaves are constant, FALSE otherwise

**Method** `is_exponentiated()`: Return exponentiation status of trees in a Forest object

*Usage:*

```
Forest$is_exponentiated()
```

*Returns:* TRUE if leaf predictions must be exponentiated, FALSE otherwise

**Method** `add_numeric_split_tree()`: Add a numeric (i.e.  $X[,i] \leq c$ ) split to a given tree in the ensemble

*Usage:*

```
Forest$add_numeric_split_tree(  
  tree_num,  
  leaf_num,  
  feature_num,  
  split_threshold,  
  left_leaf_value,  
  right_leaf_value  
)
```

*Arguments:*

tree\_num Index of the tree to be split

leaf\_num Leaf to be split

feature\_num Feature that defines the new split

split\_threshold Value that defines the cutoff of the new split

left\_leaf\_value Value (or vector of values) to assign to the newly created left node

right\_leaf\_value Value (or vector of values) to assign to the newly created right node

**Method** `get_tree_leaves()`: Retrieve a vector of indices of leaf nodes for a given tree in a given forest

*Usage:*

```
Forest$get_tree_leaves(tree_num)
```

*Arguments:*

tree\_num Index of the tree for which leaf indices will be retrieved

**Method** `get_tree_split_counts()`: Retrieve a vector of split counts for every training set variable in a given tree in the forest

*Usage:*

```
Forest$get_tree_split_counts(tree_num, num_features)
```

*Arguments:*

tree\_num Index of the tree for which split counts will be retrieved

num\_features Total number of features in the training set

**Method** `get_forest_split_counts()`: Retrieve a vector of split counts for every training set variable in the forest

*Usage:*

```
Forest$get_forest_split_counts(num_features)
```

*Arguments:*

num\_features Total number of features in the training set

**Method** `tree_max_depth()`: Maximum depth of a specific tree in the forest

*Usage:*

```
Forest$tree_max_depth(tree_num)
```

*Arguments:*

tree\_num Tree index within forest

*Returns:* Maximum leaf depth

**Method** `average_max_depth()`: Average the maximum depth of each tree in the forest

*Usage:*

`Forest$average_max_depth()`

*Returns:* Average maximum depth

**Method** `is_empty()`: When a forest object is created, it is "empty" in the sense that none of its component trees have leaves with values. There are two ways to "initialize" a Forest object. First, the `set_root_leaves()` method simply initializes every tree in the forest to a single node carrying the same (user-specified) leaf value. Second, the `prepare_for_sampler()` method initializes every tree in the forest to a single node with the same value and also propagates this information through to a ForestModel object, which must be synchronized with a Forest during a forest sampler loop.

*Usage:*

`Forest$is_empty()`

*Returns:* TRUE if a Forest has not yet been initialized with a constant root value, FALSE otherwise if the forest has already been initialized / grown.

---

ForestDataset

*Forest Dataset C++ Wrapper*

---

## Description

Wrapper around a C++ dataset class used to sample a forest. A dataset consists of three matrices / vectors: covariates, bases, and variance weights. Both the basis vector and variance weights are optional.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Public fields

`data_ptr` External pointer to a C++ ForestDataset class

## Methods

### Public methods:

- `ForestDataset$new()`
- `ForestDataset$update_basis()`
- `ForestDataset$update_variance_weights()`
- `ForestDataset$num_observations()`
- `ForestDataset$num_covariates()`

- `ForestDataset$num_basis()`
- `ForestDataset$get_covariates()`
- `ForestDataset$get_basis()`
- `ForestDataset$get_variance_weights()`
- `ForestDataset$has_basis()`
- `ForestDataset$has_variance_weights()`

**Method** `new()`: Create a new ForestDataset object.

*Usage:*

```
ForestDataset$new(covariates, basis = NULL, variance_weights = NULL)
```

*Arguments:*

`covariates` Matrix of covariates

`basis` (Optional) Matrix of bases used to define a leaf regression

`variance_weights` (Optional) Vector of observation-specific variance weights

*Returns:* A new ForestDataset object.

**Method** `update_basis()`: Update basis matrix in a dataset

*Usage:*

```
ForestDataset$update_basis(basis)
```

*Arguments:*

`basis` Updated matrix of bases used to define a leaf regression

**Method** `update_variance_weights()`: Update variance\_weights in a dataset

*Usage:*

```
ForestDataset$update_variance_weights(variance_weights, exponentiate = F)
```

*Arguments:*

`variance_weights` Updated vector of variance weights used to define individual variance / case weights

`exponentiate` Whether or not input vector should be exponentiated before being written to the Dataset's variance weights. Default: F.

**Method** `num_observations()`: Return number of observations in a ForestDataset object

*Usage:*

```
ForestDataset$num_observations()
```

*Returns:* Observation count

**Method** `num_covariates()`: Return number of covariates in a ForestDataset object

*Usage:*

```
ForestDataset$num_covariates()
```

*Returns:* Covariate count

**Method** `num_basis()`: Return number of bases in a ForestDataset object

*Usage:*

ForestDataset\$num\_basis()

*Returns:* Basis count

**Method** get\_covariates(): Return covariates as an R matrix

*Usage:*

ForestDataset\$get\_covariates()

*Returns:* Covariate data

**Method** get\_basis(): Return bases as an R matrix

*Usage:*

ForestDataset\$get\_basis()

*Returns:* Basis data

**Method** get\_variance\_weights(): Return variance weights as an R vector

*Usage:*

ForestDataset\$get\_variance\_weights()

*Returns:* Variance weight data

**Method** has\_basis(): Whether or not a dataset has a basis matrix

*Usage:*

ForestDataset\$has\_basis()

*Returns:* True if basis matrix is loaded, false otherwise

**Method** has\_variance\_weights(): Whether or not a dataset has variance weights

*Usage:*

ForestDataset\$has\_variance\_weights()

*Returns:* True if variance weights are loaded, false otherwise

---

ForestModel

*Forest Model C++ Wrapper*

---

## Description

Wraps the C++ data structures needed to sample an ensemble of decision trees and exposes functionality to run a forest sampler (using either MCMC or the grow-from-root algorithm).

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Public fields

tracker\_ptr External pointer to a C++ ForestTracker class

tree\_prior\_ptr External pointer to a C++ TreePrior class

## Methods

### Public methods:

- `ForestModel$new()`
- `ForestModel$sample_one_iteration()`
- `ForestModel$get_cached_forest_predictions()`
- `ForestModel$propagate_basis_update()`
- `ForestModel$propagate_residual_update()`
- `ForestModel$update_alpha()`
- `ForestModel$update_beta()`
- `ForestModel$update_min_samples_leaf()`
- `ForestModel$update_max_depth()`
- `ForestModel$get_alpha()`
- `ForestModel$get_beta()`
- `ForestModel$get_min_samples_leaf()`
- `ForestModel$get_max_depth()`

**Method** `new()`: Create a new ForestModel object.

*Usage:*

```
ForestModel$new(
  forest_dataset,
  feature_types,
  num_trees,
  n,
  alpha,
  beta,
  min_samples_leaf,
  max_depth = -1
)
```

*Arguments:*

`forest_dataset` ForestDataset object, used to initialize forest sampling data structures

`feature_types` Feature types (integers where 0 = numeric, 1 = ordered categorical, 2 = unordered categorical)

`num_trees` Number of trees in the forest being sampled

`n` Number of observations in `forest_dataset`

`alpha` Root node split probability in tree prior

`beta` Depth prior penalty in tree prior

`min_samples_leaf` Minimum number of samples in a tree leaf

`max_depth` Maximum depth that any tree can reach

*Returns:* A new ForestModel object.

**Method** `sample_one_iteration()`: Run a single iteration of the forest sampling algorithm (MCMC or GFR)

*Usage:*

```
ForestModel$sample_one_iteration(
  forest_dataset,
  residual,
  forest_samples,
  active_forest,
  rng,
  forest_model_config,
  global_model_config,
  num_threads = -1,
  keep_forest = TRUE,
  gfr = TRUE
)
```

*Arguments:*

`forest_dataset` Dataset used to sample the forest  
`residual` Outcome used to sample the forest  
`forest_samples` Container of forest samples  
`active_forest` "Active" forest updated by the sampler in each iteration  
`rng` Wrapper around C++ random number generator  
`forest_model_config` ForestModelConfig object containing forest model parameters and settings  
`global_model_config` GlobalModelConfig object containing global model parameters and settings  
`num_threads` Number of threads to use in the GFR and MCMC algorithms, as well as prediction. If OpenMP is not available on a user's system, this will default to 1, otherwise to the maximum number of available threads.  
`keep_forest` (Optional) Whether the updated forest sample should be saved to `forest_samples`. Default: TRUE.  
`gfr` (Optional) Whether or not the forest should be sampled using the "grow-from-root" (GFR) algorithm. Default: TRUE.

**Method** `get_cached_forest_predictions()`: Extract an internally-cached prediction of a forest on the training dataset in a sampler.

*Usage:*

```
ForestModel$get_cached_forest_predictions()
```

*Returns:* Vector with as many elements as observations in the training dataset

**Method** `propagate_basis_update()`: Propagates basis update through to the (full/partial) residual by iteratively (a) adding back in the previous prediction of each tree, (b) recomputing predictions for each tree (caching on the C++ side), (c) subtracting the new predictions from the residual.

This is useful in cases where a basis (for e.g. leaf regression) is updated outside of a tree sampler (as with e.g. adaptive coding for binary treatment BCF). Once a basis has been updated, the overall "function" represented by a tree model has changed and this should be reflected through to the residual before the next sampling loop is run.

*Usage:*

```
ForestModel$propagate_basis_update(dataset, outcome, active_forest)
```

*Arguments:*

dataset ForestDataset object storing the covariates and bases for a given forest  
outcome Outcome object storing the residuals to be updated based on forest predictions  
active\_forest "Active" forest updated by the sampler in each iteration

**Method** propagate\_residual\_update(): Update the current state of the outcome (i.e. partial residual) data by subtracting the current predictions of each tree. This function is run after the Outcome class's update\_data method, which overwrites the partial residual with an entirely new stream of outcome data.

*Usage:*

```
ForestModel$propagate_residual_update(residual)
```

*Arguments:*

residual Outcome used to sample the forest

*Returns:* None

**Method** update\_alpha(): Update alpha in the tree prior

*Usage:*

```
ForestModel$update_alpha(alpha)
```

*Arguments:*

alpha New value of alpha to be used

*Returns:* None

**Method** update\_beta(): Update beta in the tree prior

*Usage:*

```
ForestModel$update_beta(beta)
```

*Arguments:*

beta New value of beta to be used

*Returns:* None

**Method** update\_min\_samples\_leaf(): Update min\_samples\_leaf in the tree prior

*Usage:*

```
ForestModel$update_min_samples_leaf(min_samples_leaf)
```

*Arguments:*

min\_samples\_leaf New value of min\_samples\_leaf to be used

*Returns:* None

**Method** update\_max\_depth(): Update max\_depth in the tree prior

*Usage:*

```
ForestModel$update_max_depth(max_depth)
```

*Arguments:*

max\_depth New value of max\_depth to be used

*Returns:* None

**Method** `get_alpha()`: Update alpha in the tree prior

*Usage:*

`ForestModel$get_alpha()`

*Returns:* Value of alpha in the tree prior

**Method** `get_beta()`: Update beta in the tree prior

*Usage:*

`ForestModel$get_beta()`

*Returns:* Value of beta in the tree prior

**Method** `get_min_samples_leaf()`: Query min\_samples\_leaf in the tree prior

*Usage:*

`ForestModel$get_min_samples_leaf()`

*Returns:* Value of min\_samples\_leaf in the tree prior

**Method** `get_max_depth()`: Query max\_depth in the tree prior

*Usage:*

`ForestModel$get_max_depth()`

*Returns:* Value of max\_depth in the tree prior

---

ForestModelConfig

*Forest Model Configuration Object*

---

## Description

Object used to get / set parameters and other model configuration options for a forest model in the "low-level" stochtree interface. The "low-level" stochtree interface enables a high degree of sampler customization, in which users employ R wrappers around C++ objects like ForestDataset, Outcome, CppRng, and ForestModel to run the Gibbs sampler of a BART model with custom modifications. ForestModelConfig allows users to specify / query the parameters of a forest model they wish to run.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Value

Vector of integer-coded feature types (integers where 0 = numeric, 1 = ordered categorical, 2 = unordered categorical)

Vector of (0-indexed) indices of trees to update in a sweep

Vector specifying sampling probability for all p covariates in ForestDataset

Number of trees in a forest

Number of features in a forest model training set  
 Number of observations in a forest model training set  
 Root node split probability in tree prior  
 Depth prior penalty in tree prior  
 Minimum number of samples in a tree leaf  
 Maximum depth of any tree in the ensemble in the model  
 Integer coded leaf model type  
 Scale parameter used in Gaussian leaf models  
 Shape parameter for IG leaf models  
 Scale parameter for IG leaf models  
 Number of unique cutpoints to consider  
 Number of features to subsample for the GFR algorithm

### Public fields

feature\_types Vector of integer-coded feature types (integers where 0 = numeric, 1 = ordered categorical, 2 = unordered categorical)  
 sweep\_update\_indices Vector of trees to update in a sweep  
 num\_trees Number of trees in the forest being sampled  
 num\_features Number of features in training dataset  
 num\_observations Number of observations in training dataset  
 leaf\_dimension Dimension of the leaf model  
 alpha Root node split probability in tree prior  
 beta Depth prior penalty in tree prior  
 min\_samples\_leaf Minimum number of samples in a tree leaf  
 max\_depth Maximum depth of any tree in the ensemble in the model. Setting to -1 does not enforce any depth limits on trees.  
 leaf\_model\_type Integer specifying the leaf model type (0 = constant leaf, 1 = univariate leaf regression, 2 = multivariate leaf regression)  
 leaf\_model\_scale Scale parameter used in Gaussian leaf models  
 variable\_weights Vector specifying sampling probability for all p covariates in ForestDataset  
 variance\_forest\_shape Shape parameter for IG leaf models (applicable when leaf\_model\_type = 3)  
 variance\_forest\_scale Scale parameter for IG leaf models (applicable when leaf\_model\_type = 3)  
 cutpoint\_grid\_size Number of unique cutpoints to consider  
 num\_features\_subsample Number of features to subsample for the GFR algorithm Create a new ForestModelConfig object.

## Methods

### Public methods:

- `ForestModelConfig$new()`
- `ForestModelConfig$update_feature_types()`
- `ForestModelConfig$update_sweep_indices()`
- `ForestModelConfig$update_variable_weights()`
- `ForestModelConfig$update_alpha()`
- `ForestModelConfig$update_beta()`
- `ForestModelConfig$update_min_samples_leaf()`
- `ForestModelConfig$update_max_depth()`
- `ForestModelConfig$update_leaf_model_scale()`
- `ForestModelConfig$update_variance_forest_shape()`
- `ForestModelConfig$update_variance_forest_scale()`
- `ForestModelConfig$update_cutpoint_grid_size()`
- `ForestModelConfig$update_num_features_subsample()`
- `ForestModelConfig$get_feature_types()`
- `ForestModelConfig$get_sweep_indices()`
- `ForestModelConfig$get_variable_weights()`
- `ForestModelConfig$get_num_trees()`
- `ForestModelConfig$get_num_features()`
- `ForestModelConfig$get_num_observations()`
- `ForestModelConfig$get_alpha()`
- `ForestModelConfig$get_beta()`
- `ForestModelConfig$get_min_samples_leaf()`
- `ForestModelConfig$get_max_depth()`
- `ForestModelConfig$get_leaf_model_type()`
- `ForestModelConfig$get_leaf_model_scale()`
- `ForestModelConfig$get_variance_forest_shape()`
- `ForestModelConfig$get_variance_forest_scale()`
- `ForestModelConfig$get_cutpoint_grid_size()`
- `ForestModelConfig$get_num_features_subsample()`

### Method `new()`:

*Usage:*

```
ForestModelConfig$new(  
  feature_types = NULL,  
  sweep_update_indices = NULL,  
  num_trees = NULL,  
  num_features = NULL,  
  num_observations = NULL,  
  variable_weights = NULL,  
  leaf_dimension = 1,  
  alpha = 0.95,
```

```

    beta = 2,
    min_samples_leaf = 5,
    max_depth = -1,
    leaf_model_type = 1,
    leaf_model_scale = NULL,
    variance_forest_shape = 1,
    variance_forest_scale = 1,
    cutpoint_grid_size = 100,
    num_features_subsample = NULL
)

```

*Arguments:*

**feature\_types** Vector of integer-coded feature types (where 0 = numeric, 1 = ordered categorical, 2 = unordered categorical)

**sweep\_update\_indices** Vector of (0-indexed) indices of trees to update in a sweep

**num\_trees** Number of trees in the forest being sampled

**num\_features** Number of features in training dataset

**num\_observations** Number of observations in training dataset

**variable\_weights** Vector specifying sampling probability for all p covariates in ForestDataset

**leaf\_dimension** Dimension of the leaf model (default: 1)

**alpha** Root node split probability in tree prior (default: 0.95)

**beta** Depth prior penalty in tree prior (default: 2.0)

**min\_samples\_leaf** Minimum number of samples in a tree leaf (default: 5)

**max\_depth** Maximum depth of any tree in the ensemble in the model. Setting to -1 does not enforce any depth limits on trees. Default: -1.

**leaf\_model\_type** Integer specifying the leaf model type (0 = constant leaf, 1 = univariate leaf regression, 2 = multivariate leaf regression). Default: 0.

**leaf\_model\_scale** Scale parameter used in Gaussian leaf models (can either be a scalar or a  $q \times q$  matrix, where  $q$  is the dimensionality of the basis and is only  $>1$  when `leaf_model_type` = 2). Calibrated internally as  $1/\text{num\_trees}$ , propagated along diagonal if needed for multivariate leaf models.

**variance\_forest\_shape** Shape parameter for IG leaf models (applicable when `leaf_model_type` = 3). Default: 1.

**variance\_forest\_scale** Scale parameter for IG leaf models (applicable when `leaf_model_type` = 3). Default: 1.

**cutpoint\_grid\_size** Number of unique cutpoints to consider (default: 100)

**num\_features\_subsample** Number of features to subsample for the GFR algorithm

*Returns:* A new ForestModelConfig object.

**Method** `update_feature_types()`: Update feature types

*Usage:*

```
ForestModelConfig$update_feature_types(feature_types)
```

*Arguments:*

**feature\_types** Vector of integer-coded feature types (integers where 0 = numeric, 1 = ordered categorical, 2 = unordered categorical)

**Method** `update_sweep_indices()`: Update sweep update indices

*Usage:*

```
ForestModelConfig$update_sweep_indices(sweep_update_indices)
```

*Arguments:*

`sweep_update_indices` Vector of (0-indexed) indices of trees to update in a sweep

**Method** `update_variable_weights()`: Update variable weights

*Usage:*

```
ForestModelConfig$update_variable_weights(variable_weights)
```

*Arguments:*

`variable_weights` Vector specifying sampling probability for all  $p$  covariates in ForestDataset

**Method** `update_alpha()`: Update root node split probability in tree prior

*Usage:*

```
ForestModelConfig$update_alpha(alpha)
```

*Arguments:*

`alpha` Root node split probability in tree prior

**Method** `update_beta()`: Update depth prior penalty in tree prior

*Usage:*

```
ForestModelConfig$update_beta(beta)
```

*Arguments:*

`beta` Depth prior penalty in tree prior

**Method** `update_min_samples_leaf()`: Update minimum number of samples per leaf node in the tree prior

*Usage:*

```
ForestModelConfig$update_min_samples_leaf(min_samples_leaf)
```

*Arguments:*

`min_samples_leaf` Minimum number of samples in a tree leaf

**Method** `update_max_depth()`: Update max depth in the tree prior

*Usage:*

```
ForestModelConfig$update_max_depth(max_depth)
```

*Arguments:*

`max_depth` Maximum depth of any tree in the ensemble in the model

**Method** `update_leaf_model_scale()`: Update scale parameter used in Gaussian leaf models

*Usage:*

```
ForestModelConfig$update_leaf_model_scale(leaf_model_scale)
```

*Arguments:*

`leaf_model_scale` Scale parameter used in Gaussian leaf models

**Method** `update_variance_forest_shape()`: Update shape parameter for IG leaf models

*Usage:*

```
ForestModelConfig$update_variance_forest_shape(variance_forest_shape)
```

*Arguments:*

`variance_forest_shape` Shape parameter for IG leaf models

**Method** `update_variance_forest_scale()`: Update scale parameter for IG leaf models

*Usage:*

```
ForestModelConfig$update_variance_forest_scale(variance_forest_scale)
```

*Arguments:*

`variance_forest_scale` Scale parameter for IG leaf models

**Method** `update_cutpoint_grid_size()`: Update number of unique cutpoints to consider

*Usage:*

```
ForestModelConfig$update_cutpoint_grid_size(cutpoint_grid_size)
```

*Arguments:*

`cutpoint_grid_size` Number of unique cutpoints to consider

**Method** `update_num_features_subsample()`: Update number of features to subsample for the GFR algorithm

*Usage:*

```
ForestModelConfig$update_num_features_subsample(num_features_subsample)
```

*Arguments:*

`num_features_subsample` Number of features to subsample for the GFR algorithm

**Method** `get_feature_types()`: Query feature types for this ForestModelConfig object

*Usage:*

```
ForestModelConfig$get_feature_types()
```

**Method** `get_sweep_indices()`: Query sweep update indices for this ForestModelConfig object

*Usage:*

```
ForestModelConfig$get_sweep_indices()
```

**Method** `get_variable_weights()`: Query variable weights for this ForestModelConfig object

*Usage:*

```
ForestModelConfig$get_variable_weights()
```

**Method** `get_num_trees()`: Query number of trees

*Usage:*

```
ForestModelConfig$get_num_trees()
```

**Method** `get_num_features()`: Query number of features

*Usage:*

```
ForestModelConfig$get_num_features()
```

**Method** `get_num_observations()`: Query number of observations

*Usage:*

`ForestModelConfig$get_num_observations()`

**Method** `get_alpha()`: Query root node split probability in tree prior for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_alpha()`

**Method** `get_beta()`: Query depth prior penalty in tree prior for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_beta()`

**Method** `get_min_samples_leaf()`: Query root node split probability in tree prior for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_min_samples_leaf()`

**Method** `get_max_depth()`: Query root node split probability in tree prior for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_max_depth()`

**Method** `get_leaf_model_type()`: Query (integer-coded) type of leaf model

*Usage:*

`ForestModelConfig$get_leaf_model_type()`

**Method** `get_leaf_model_scale()`: Query scale parameter used in Gaussian leaf models for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_leaf_model_scale()`

**Method** `get_variance_forest_shape()`: Query shape parameter for IG leaf models for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_variance_forest_shape()`

**Method** `get_variance_forest_scale()`: Query scale parameter for IG leaf models for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_variance_forest_scale()`

**Method** `get_cutpoint_grid_size()`: Query number of unique cutpoints to consider for this ForestModelConfig object

*Usage:*

`ForestModelConfig$get_cutpoint_grid_size()`

**Method** `get_num_features_subsample()`: Query number of features to subsample for the GFR algorithm

*Usage:*

```
ForestModelConfig$get_num_features_subsample()
```

---

ForestSamples

*Forest Container C++ Wrapper*


---

## Description

Wrapper around a C++ class that stores draws from an random ensemble of decision trees.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Public fields

`forest_container_ptr` External pointer to a C++ ForestContainer class

## Methods

### Public methods:

- `ForestSamples$new()`
- `ForestSamples$collapse()`
- `ForestSamples$combine_forests()`
- `ForestSamples$add_to_forest()`
- `ForestSamples$multiply_forest()`
- `ForestSamples$load_from_json()`
- `ForestSamples$append_from_json()`
- `ForestSamples$load_from_json_string()`
- `ForestSamples$append_from_json_string()`
- `ForestSamples$predict()`
- `ForestSamples$predict_raw()`
- `ForestSamples$predict_raw_single_forest()`
- `ForestSamples$predict_raw_single_tree()`
- `ForestSamples$set_root_leaves()`
- `ForestSamples$prepare_for_sampler()`
- `ForestSamples$adjust_residual()`
- `ForestSamples$save_json()`
- `ForestSamples$load_json()`
- `ForestSamples$num_samples()`
- `ForestSamples$num_trees()`

- `ForestSamples$leaf_dimension()`
- `ForestSamples$is_constant_leaf()`
- `ForestSamples$is_exponentiated()`
- `ForestSamples$add_forest_with_constant_leaves()`
- `ForestSamples$add_numeric_split_tree()`
- `ForestSamples$get_tree_leaves()`
- `ForestSamples$get_tree_split_counts()`
- `ForestSamples$get_forest_split_counts()`
- `ForestSamples$get_aggregate_split_counts()`
- `ForestSamples$get_granular_split_counts()`
- `ForestSamples$ensemble_tree_max_depth()`
- `ForestSamples$average_ensemble_max_depth()`
- `ForestSamples$average_max_depth()`
- `ForestSamples$num_forest_leaves()`
- `ForestSamples$sum_leaves_squared()`
- `ForestSamples$is_leaf_node()`
- `ForestSamples$is_numeric_split_node()`
- `ForestSamples$is_categorical_split_node()`
- `ForestSamples$parent_node()`
- `ForestSamples$left_child_node()`
- `ForestSamples$right_child_node()`
- `ForestSamples$node_depth()`
- `ForestSamples$node_split_index()`
- `ForestSamples$node_split_threshold()`
- `ForestSamples$node_split_categories()`
- `ForestSamples$node_leaf_values()`
- `ForestSamples$num_nodes()`
- `ForestSamples$num_leaves()`
- `ForestSamples$num_leaf_parents()`
- `ForestSamples$num_split_nodes()`
- `ForestSamples$nodes()`
- `ForestSamples$leaves()`
- `ForestSamples$delete_sample()`

**Method** `new()`: Create a new `ForestContainer` object.

*Usage:*

```
ForestSamples$new(
  num_trees,
  leaf_dimension = 1,
  is_leaf_constant = FALSE,
  is_exponentiated = FALSE
)
```

*Arguments:*

num\_trees Number of trees  
 leaf\_dimension Dimensionality of the outcome model  
 is\_leaf\_constant Whether leaf is constant  
 is\_exponentiated Whether forest predictions should be exponentiated before being returned

*Returns:* A new ForestContainer object.

**Method collapse():** Collapse forests in this container by a pre-specified batch size. For example, if we have a container of twenty 10-tree forests, and we specify a batch\_size of 5, then this method will yield four 50-tree forests. "Excess" forests remaining after the size of a forest container is divided by batch\_size will be pruned from the beginning of the container (i.e. earlier sampled forests will be deleted). This method has no effect if batch\_size is larger than the number of forests in a container.

*Usage:*

```
ForestSamples$collapse(batch_size)
```

*Arguments:*

batch\_size Number of forests to be collapsed into a single forest

**Method combine\_forests():** Merge specified forests into a single forest

*Usage:*

```
ForestSamples$combine_forests(forest_inds)
```

*Arguments:*

forest\_inds Indices of forests to be combined (0-indexed)

**Method add\_to\_forest():** Add a constant value to every leaf of every tree of a given forest

*Usage:*

```
ForestSamples$add_to_forest(forest_index, constant_value)
```

*Arguments:*

forest\_index Index of forest whose leaves will be modified (0-indexed)

constant\_value Value to add to every leaf of every tree of the forest at forest\_index

**Method multiply\_forest():** Multiply every leaf of every tree of a given forest by constant value

*Usage:*

```
ForestSamples$multiply_forest(forest_index, constant_multiple)
```

*Arguments:*

forest\_index Index of forest whose leaves will be modified (0-indexed)

constant\_multiple Value to multiply through by every leaf of every tree of the forest at forest\_index

**Method load\_from\_json():** Create a new ForestContainer object from a json object

*Usage:*

```
ForestSamples$load_from_json(json_object, json_forest_label)
```

*Arguments:*

`json_object` Object of class CppJson

`json_forest_label` Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy

*Returns:* A new ForestContainer object.

**Method** `append_from_json()`: Append to a ForestContainer object from a json object

*Usage:*

```
ForestSamples$append_from_json(json_object, json_forest_label)
```

*Arguments:*

`json_object` Object of class CppJson

`json_forest_label` Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy

*Returns:* None

**Method** `load_from_json_string()`: Create a new ForestContainer object from a json object

*Usage:*

```
ForestSamples$load_from_json_string(json_string, json_forest_label)
```

*Arguments:*

`json_string` JSON string which parses into object of class CppJson

`json_forest_label` Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy

*Returns:* A new ForestContainer object.

**Method** `append_from_json_string()`: Append to a ForestContainer object from a json object

*Usage:*

```
ForestSamples$append_from_json_string(json_string, json_forest_label)
```

*Arguments:*

`json_string` JSON string which parses into object of class CppJson

`json_forest_label` Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy

*Returns:* None

**Method** `predict()`: Predict every tree ensemble on every sample in `forest_dataset`

*Usage:*

```
ForestSamples$predict(forest_dataset)
```

*Arguments:*

`forest_dataset` ForestDataset R class

*Returns:* matrix of predictions with as many rows as in `forest_dataset` and as many columns as samples in the ForestContainer

**Method** `predict_raw()`: Predict "raw" leaf values (without being multiplied by basis) for every tree ensemble on every sample in `forest_dataset`

*Usage:*

```
ForestSamples$predict_raw(forest_dataset)
```

*Arguments:*

forest\_dataset ForestDataset R class

*Returns:* Array of predictions for each observation in forest\_dataset and each sample in the ForestSamples class with each prediction having the dimensionality of the forests' leaf model. In the case of a constant leaf model or univariate leaf regression, this array is two-dimensional (number of observations, number of forest samples). In the case of a multivariate leaf regression, this array is three-dimension (number of observations, leaf model dimension, number of samples).

**Method** predict\_raw\_single\_forest(): Predict "raw" leaf values (without being multiplied by basis) for a specific forest on every sample in forest\_dataset

*Usage:*

```
ForestSamples$predict_raw_single_forest(forest_dataset, forest_num)
```

*Arguments:*

forest\_dataset ForestDataset R class

forest\_num Index of the forest sample within the container

*Returns:* matrix of predictions with as many rows as in forest\_dataset and as many columns as dimensions in the leaves of trees in ForestContainer

**Method** predict\_raw\_single\_tree(): Predict "raw" leaf values (without being multiplied by basis) for a specific tree in a specific forest on every observation in forest\_dataset

*Usage:*

```
ForestSamples$predict_raw_single_tree(forest_dataset, forest_num, tree_num)
```

*Arguments:*

forest\_dataset ForestDataset R class

forest\_num Index of the forest sample within the container

tree\_num Index of the tree to be queried

*Returns:* matrix of predictions with as many rows as in forest\_dataset and as many columns as dimensions in the leaves of trees in ForestContainer

**Method** set\_root\_leaves(): Set a constant predicted value for every tree in the ensemble. Stops program if any tree is more than a root node.

*Usage:*

```
ForestSamples$set_root_leaves(forest_num, leaf_value)
```

*Arguments:*

forest\_num Index of the forest sample within the container.

leaf\_value Constant leaf value(s) to be fixed for each tree in the ensemble indexed by forest\_num.  
Can be either a single number or a vector, depending on the forest's leaf dimension.

**Method** prepare\_for\_sampler(): Set a constant predicted value for every tree in the ensemble. Stops program if any tree is more than a root node.

*Usage:*

```
ForestSamples$prepare_for_sampler(
  dataset,
  outcome,
  forest_model,
  leaf_model_int,
  leaf_value
)
```

*Arguments:*

**dataset** ForestDataset Dataset class (covariates, basis, etc...)  
**outcome** Outcome Outcome class (residual / partial residual)  
**forest\_model** ForestModel object storing tracking structures used in training / sampling  
**leaf\_model\_int** Integer value encoding the leaf model type (0 = constant gaussian, 1 = univariate gaussian, 2 = multivariate gaussian, 3 = log linear variance).  
**leaf\_value** Constant leaf value(s) to be fixed for each tree in the ensemble indexed by forest\_num.  
 Can be either a single number or a vector, depending on the forest's leaf dimension.

**Method** `adjust_residual()`: Adjusts residual based on the predictions of a forest

This is typically run just once at the beginning of a forest sampling algorithm. After trees are initialized with constant root node predictions, their root predictions are subtracted out of the residual.

*Usage:*

```
ForestSamples$adjust_residual(
  dataset,
  outcome,
  forest_model,
  requires_basis,
  forest_num,
  add
)
```

*Arguments:*

**dataset** ForestDataset object storing the covariates and bases for a given forest  
**outcome** Outcome object storing the residuals to be updated based on forest predictions  
**forest\_model** ForestModel object storing tracking structures used in training / sampling  
**requires\_basis** Whether or not a forest requires a basis for prediction  
**forest\_num** Index of forest used to update residuals  
**add** Whether forest predictions should be added to or subtracted from residuals

**Method** `save_json()`: Store the trees and metadata of ForestDataset class in a json file

*Usage:*

```
ForestSamples$save_json(json_filename)
```

*Arguments:*

**json\_filename** Name of output json file (must end in ".json")

**Method** `load_json()`: Load trees and metadata for an ensemble from a json file. Note that any trees and metadata already present in ForestDataset class will be overwritten.

*Usage:*

ForestSamples\$load\_json(json\_filename)

*Arguments:*

json\_filename Name of model input json file (must end in ".json")

**Method** num\_samples(): Return number of samples in a ForestContainer object

*Usage:*

ForestSamples\$num\_samples()

*Returns:* Sample count

**Method** num\_trees(): Return number of trees in each ensemble of a ForestContainer object

*Usage:*

ForestSamples\$num\_trees()

*Returns:* Tree count

**Method** leaf\_dimension(): Return output dimension of trees in a ForestContainer object

*Usage:*

ForestSamples\$leaf\_dimension()

*Returns:* Leaf node parameter size

**Method** is\_constant\_leaf(): Return constant leaf status of trees in a ForestContainer object

*Usage:*

ForestSamples\$is\_constant\_leaf()

*Returns:* TRUE if leaves are constant, FALSE otherwise

**Method** is\_exponentiated(): Return exponentiation status of trees in a ForestContainer object

*Usage:*

ForestSamples\$is\_exponentiated()

*Returns:* TRUE if leaf predictions must be exponentiated, FALSE otherwise

**Method** add\_forest\_with\_constant\_leaves(): Add a new all-root ensemble to the container, with all of the leaves set to the value / vector provided

*Usage:*

ForestSamples\$add\_forest\_with\_constant\_leaves(leaf\_value)

*Arguments:*

leaf\_value Value (or vector of values) to initialize root nodes in tree

**Method** add\_numeric\_split\_tree(): Add a numeric (i.e.  $X[,i] \leq c$ ) split to a given tree in the ensemble

*Usage:*

```
ForestSamples$add_numeric_split_tree(
  forest_num,
  tree_num,
  leaf_num,
  feature_num,
  split_threshold,
  left_leaf_value,
  right_leaf_value
)
```

*Arguments:*

forest\_num Index of the forest which contains the tree to be split

tree\_num Index of the tree to be split

leaf\_num Leaf to be split

feature\_num Feature that defines the new split

split\_threshold Value that defines the cutoff of the new split

left\_leaf\_value Value (or vector of values) to assign to the newly created left node

right\_leaf\_value Value (or vector of values) to assign to the newly created right node

**Method** `get_tree_leaves()`: Retrieve a vector of indices of leaf nodes for a given tree in a given forest

*Usage:*

```
ForestSamples$get_tree_leaves(forest_num, tree_num)
```

*Arguments:*

forest\_num Index of the forest which contains tree tree\_num

tree\_num Index of the tree for which leaf indices will be retrieved

**Method** `get_tree_split_counts()`: Retrieve a vector of split counts for every training set variable in a given tree in a given forest

*Usage:*

```
ForestSamples$get_tree_split_counts(forest_num, tree_num, num_features)
```

*Arguments:*

forest\_num Index of the forest which contains tree tree\_num

tree\_num Index of the tree for which split counts will be retrieved

num\_features Total number of features in the training set

**Method** `get_forest_split_counts()`: Retrieve a vector of split counts for every training set variable in a given forest

*Usage:*

```
ForestSamples$get_forest_split_counts(forest_num, num_features)
```

*Arguments:*

forest\_num Index of the forest for which split counts will be retrieved

num\_features Total number of features in the training set

**Method** `get_aggregate_split_counts()`: Retrieve a vector of split counts for every training set variable in a given forest, aggregated across ensembles and trees

*Usage:*

```
ForestSamples$get_aggregate_split_counts(num_features)
```

*Arguments:*

num\_features Total number of features in the training set

**Method** `get_granular_split_counts()`: Retrieve a vector of split counts for every training set variable in a given forest, reported separately for each ensemble and tree

*Usage:*

```
ForestSamples$get_granular_split_counts(num_features)
```

*Arguments:*

num\_features Total number of features in the training set

**Method** `ensemble_tree_max_depth()`: Maximum depth of a specific tree in a specific ensemble in a ForestSamples object

*Usage:*

```
ForestSamples$ensemble_tree_max_depth(ensemble_num, tree_num)
```

*Arguments:*

ensemble\_num Ensemble number

tree\_num Tree index within ensemble ensemble\_num

*Returns:* Maximum leaf depth

**Method** `average_ensemble_max_depth()`: Average the maximum depth of each tree in a given ensemble in a ForestSamples object

*Usage:*

```
ForestSamples$average_ensemble_max_depth(ensemble_num)
```

*Arguments:*

ensemble\_num Ensemble number

*Returns:* Average maximum depth

**Method** `average_max_depth()`: Average the maximum depth of each tree in each ensemble in a ForestContainer object

*Usage:*

```
ForestSamples$average_max_depth()
```

*Returns:* Average maximum depth

**Method** `num_forest_leaves()`: Number of leaves in a given ensemble in a ForestSamples object

*Usage:*

```
ForestSamples$num_forest_leaves(forest_num)
```

*Arguments:*

forest\_num Index of the ensemble to be queried

*Returns:* Count of leaves in the ensemble stored at forest\_num

**Method** `sum_leaves_squared()`: Sum of squared (raw) leaf values in a given ensemble in a ForestSamples object

*Usage:*

```
ForestSamples$sum_leaves_squared(forest_num)
```

*Arguments:*

`forest_num` Index of the ensemble to be queried

*Returns:* Average maximum depth

**Method** `is_leaf_node()`: Whether or not a given node of a given tree in a given forest in the ForestSamples is a leaf

*Usage:*

```
ForestSamples$is_leaf_node(forest_num, tree_num, node_id)
```

*Arguments:*

`forest_num` Index of the forest to be queried

`tree_num` Index of the tree to be queried

`node_id` Index of the node to be queried

*Returns:* TRUE if node is a leaf, FALSE otherwise

**Method** `is_numeric_split_node()`: Whether or not a given node of a given tree in a given forest in the ForestSamples is a numeric split node

*Usage:*

```
ForestSamples$is_numeric_split_node(forest_num, tree_num, node_id)
```

*Arguments:*

`forest_num` Index of the forest to be queried

`tree_num` Index of the tree to be queried

`node_id` Index of the node to be queried

*Returns:* TRUE if node is a numeric split node, FALSE otherwise

**Method** `is_categorical_split_node()`: Whether or not a given node of a given tree in a given forest in the ForestSamples is a categorical split node

*Usage:*

```
ForestSamples$is_categorical_split_node(forest_num, tree_num, node_id)
```

*Arguments:*

`forest_num` Index of the forest to be queried

`tree_num` Index of the tree to be queried

`node_id` Index of the node to be queried

*Returns:* TRUE if node is a categorical split node, FALSE otherwise

**Method** `parent_node()`: Parent node of given node of a given tree in a given forest in a ForestSamples object

*Usage:*

```
ForestSamples$parent_node(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried  
tree\_num Index of the tree to be queried  
node\_id Index of the node to be queried

*Returns:* Integer ID of the parent node

**Method** left\_child\_node(): Left child node of given node of a given tree in a given forest in a ForestSamples object

*Usage:*

```
ForestSamples$left_child_node(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried  
tree\_num Index of the tree to be queried  
node\_id Index of the node to be queried

*Returns:* Integer ID of the left child node

**Method** right\_child\_node(): Right child node of given node of a given tree in a given forest in a ForestSamples object

*Usage:*

```
ForestSamples$right_child_node(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried  
tree\_num Index of the tree to be queried  
node\_id Index of the node to be queried

*Returns:* Integer ID of the right child node

**Method** node\_depth(): Depth of given node of a given tree in a given forest in a ForestSamples object, with 0 depth for the root node.

*Usage:*

```
ForestSamples$node_depth(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried  
tree\_num Index of the tree to be queried  
node\_id Index of the node to be queried

*Returns:* Integer valued depth of the node

**Method** node\_split\_index(): Split index of given node of a given tree in a given forest in a ForestSamples object. Returns -1 if node is a leaf.

*Usage:*

```
ForestSamples$node_split_index(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

node\_id Index of the node to be queried

*Returns:* Integer valued depth of the node

**Method** node\_split\_threshold(): Threshold that defines a numeric split for a given node of a given tree in a given forest in a ForestSamples object. Returns Inf if the node is a leaf or a categorical split node.

*Usage:*

```
ForestSamples$node_split_threshold(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

node\_id Index of the node to be queried

*Returns:* Threshold defining a split for the node

**Method** node\_split\_categories(): Array of category indices that define a categorical split for a given node of a given tree in a given forest in a ForestSamples object. Returns c(Inf) if the node is a leaf or a numeric split node.

*Usage:*

```
ForestSamples$node_split_categories(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

node\_id Index of the node to be queried

*Returns:* Categories defining a split for the node

**Method** node\_leaf\_values(): Leaf node value(s) for a given node of a given tree in a given forest in a ForestSamples object. Values are stale if the node is a split node.

*Usage:*

```
ForestSamples$node_leaf_values(forest_num, tree_num, node_id)
```

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

node\_id Index of the node to be queried

*Returns:* Vector (often univariate) of leaf values

**Method** num\_nodes(): Number of nodes in a given tree in a given forest in a ForestSamples object.

*Usage:*

```
ForestSamples$num_nodes(forest_num, tree_num)
```

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

*Returns:* Count of total tree nodes

**Method** num\_leaves(): Number of leaves in a given tree in a given forest in a ForestSamples object.

*Usage:*

ForestSamples\$num\_leaves(forest\_num, tree\_num)

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

*Returns:* Count of total tree leaves

**Method** num\_leaf\_parents(): Number of leaf parents (split nodes with two leaves as children) in a given tree in a given forest in a ForestSamples object.

*Usage:*

ForestSamples\$num\_leaf\_parents(forest\_num, tree\_num)

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

*Returns:* Count of total tree leaf parents

**Method** num\_split\_nodes(): Number of split nodes in a given tree in a given forest in a ForestSamples object.

*Usage:*

ForestSamples\$num\_split\_nodes(forest\_num, tree\_num)

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

*Returns:* Count of total tree split nodes

**Method** nodes(): Array of node indices in a given tree in a given forest in a ForestSamples object.

*Usage:*

ForestSamples\$nodes(forest\_num, tree\_num)

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

*Returns:* Indices of tree nodes

**Method** leaves(): Array of leaf indices in a given tree in a given forest in a ForestSamples object.

*Usage:*

```
ForestSamples$leaves(forest_num, tree_num)
```

*Arguments:*

forest\_num Index of the forest to be queried

tree\_num Index of the tree to be queried

*Returns:* Indices of leaf nodes

**Method** delete\_sample(): Modify the ForestSamples object by removing the forest sample indexed by 'forest\_num'

*Usage:*

```
ForestSamples$delete_sample(forest_num)
```

*Arguments:*

forest\_num Index of the forest to be removed

---

getRandomEffectSamples

*Extract Random Effect Samples Generic Function*

---

## Description

Generic function for extracting random effect samples from a model object (BCF, BART, etc...)

## Usage

```
getRandomEffectSamples(object, ...)
```

## Arguments

|        |                                                          |
|--------|----------------------------------------------------------|
| object | Fitted model object from which to extract random effects |
| ...    | Other parameters to be used in random effects extraction |

## Value

List of random effect samples

## Examples

```
n <- 100
p <- 10
X <- matrix(runif(n*p), ncol = p)
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- rep(1.0, n)
y <- (-5 + 10*(X[,1] > 0.5)) + (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
bart_model <- bart(X_train=X, y_train=y, rfx_group_ids_train=rfx_group_ids,
                  rfx_basis_train = rfx_basis, num_gfr=0, num_mcmc=10)
rfx_samples <- getRandomEffectSamples(bart_model)
```

---

getRandomEffectSamples.bartmodel

*Extract Random Effects Samples*


---

## Description

Extract raw sample values for each of the random effect parameter terms.

## Usage

```
## S3 method for class 'bartmodel'
getRandomEffectSamples(object, ...)
```

## Arguments

|        |                                                                                            |
|--------|--------------------------------------------------------------------------------------------|
| object | Object of type bartmodel containing draws of a BART model and associated sampling outputs. |
| ...    | Other parameters to be used in random effects extraction                                   |

## Value

List of arrays. The alpha array has dimension (num\_components, num\_samples) and is simply a vector if num\_components = 1. The xi and beta arrays have dimension (num\_components, num\_groups, num\_samples) and is simply a matrix if num\_components = 1. The sigma array has dimension (num\_components, num\_samples) and is simply a vector if num\_components = 1.

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
snr <- 3
group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[group_ids,] * rfx_basis)
E_y <- f_XW + rfx_term
y <- E_y + rnorm(n, 0, 1)*(sd(E_y)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
```

```

X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
rfx_group_ids_test <- group_ids[test_inds]
rfx_group_ids_train <- group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train, X_test = X_test,
                  rfx_group_ids_train = rfx_group_ids_train,
                  rfx_group_ids_test = rfx_group_ids_test,
                  rfx_basis_train = rfx_basis_train,
                  rfx_basis_test = rfx_basis_test,
                  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
rfx_samples <- getRandomEffectSamples(bart_model)

```

---

```
getRandomEffectSamples.bcfmodel
```

*Extract Random Effect Samples from BCF Model*

---

## Description

Extract raw sample values for each of the random effect parameter terms.

## Usage

```
## S3 method for class 'bcfmodel'
getRandomEffectSamples(object, ...)
```

## Arguments

|                     |                                                                                                                          |
|---------------------|--------------------------------------------------------------------------------------------------------------------------|
| <code>object</code> | Object of type <code>bcfmodel</code> containing draws of a Bayesian causal forest model and associated sampling outputs. |
| <code>...</code>    | Other parameters to be used in random effects extraction                                                                 |

## Value

List of arrays. The alpha array has dimension (num\_components, num\_samples) and is simply a vector if num\_components = 1. The xi and beta arrays have dimension (num\_components, num\_groups, num\_samples) and is simply a matrix if num\_components = 1. The sigma array has dimension (num\_components, num\_samples) and is simply a vector if num\_components = 1.

**Examples**

```

n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]

```

```

rfx_term_train <- rfx_term[train_inds]
mu_params <- list(sample_sigma2_leaf = TRUE)
tau_params <- list(sample_sigma2_leaf = FALSE)
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10,
  prognostic_forest_params = mu_params,
  treatment_effect_forest_params = tau_params)
rfx_samples <- getRandomEffectSamples(bcf_model)

```

GlobalModelConfig

*Global Model Configuration Object***Description**

Object used to get / set global parameters and other global model configuration options in the "low-level" stochtree interface. The "low-level" stochtree interface enables a high degree of sampler customization, in which users employ R wrappers around C++ objects like ForestDataset, Outcome, CppRng, and ForestModel to run the Gibbs sampler of a BART model with custom modifications. GlobalModelConfig allows users to specify / query the global parameters of a model they wish to run.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Value**

Global error variance parameter

**Public fields**

`global_error_variance` Global error variance parameter Create a new GlobalModelConfig object.

**Methods****Public methods:**

- `GlobalModelConfig$new()`
- `GlobalModelConfig$update_global_error_variance()`
- `GlobalModelConfig$get_global_error_variance()`

**Method new():**

*Usage:*

```
GlobalModelConfig$new(global_error_variance = 1)
```

*Arguments:*

global\_error\_variance Global error variance parameter (default: 1.0)

*Returns:* A new GlobalModelConfig object.

**Method** update\_global\_error\_variance(): Update global error variance parameter

*Usage:*

```
GlobalModelConfig$update_global_error_variance(global_error_variance)
```

*Arguments:*

global\_error\_variance Global error variance parameter

**Method** get\_global\_error\_variance(): Query global error variance parameter for this GlobalModelConfig object

*Usage:*

```
GlobalModelConfig$get_global_error_variance()
```

---

loadForestContainerCombinedJson

*Combine JSON Model Objects into ForestSamples*

---

**Description**

Combine multiple JSON model objects containing forests (with the same hierarchy / schema) into a single forest\_container

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
loadForestContainerCombinedJson(json_object_list, json_forest_label)
```

**Arguments**

json\_object\_list

List of objects of class CppJson

json\_forest\_label

Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy (must exist in every json object in the list)

**Value**

ForestSamples object

**Examples**

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
bart_model <- bart(X, y, num_gfr=0, num_mcmc=10)
bart_json <- list(saveBARTModelToJson(bart_model))
mean_forest <- loadForestContainerCombinedJson(bart_json, "forest_0")
```

---

loadForestContainerCombinedJsonString

*Combine JSON Strings into ForestSamples*


---

**Description**

Combine multiple JSON strings representing model objects containing forests (with the same hierarchy / schema) into a single forest\_container

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
loadForestContainerCombinedJsonString(json_string_list, json_forest_label)
```

**Arguments**

```
json_string_list
```

List of strings that parse into objects of type CppJson

```
json_forest_label
```

Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy (must exist in every json object in the list)

**Value**

ForestSamples object

**Examples**

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
bart_model <- bart(X, y, num_gfr=0, num_mcmc=10)
bart_json_string <- list(saveBARTModelToJsonString(bart_model))
mean_forest <- loadForestContainerCombinedJsonString(bart_json_string, "forest_0")
```

---

loadForestContainerJson

*Load Forest Samples from JSON*


---

### Description

Load a container of forest samples from json

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

### Usage

```
loadForestContainerJson(json_object, json_forest_label)
```

### Arguments

json\_object      Object of class CppJson

json\_forest\_label

Label referring to a particular forest (i.e. "forest\_0") in the overall json hierarchy

### Value

ForestSamples object

### Examples

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
bart_model <- bart(X, y, num_gfr=0, num_mcmc=10)
bart_json <- saveBARTModelToJson(bart_model)
mean_forest <- loadForestContainerJson(bart_json, "forest_0")
```

---

loadRandomEffectSamplesCombinedJson

*Combine JSON Model Objects into RandomEffectSamples*


---

### Description

Combine multiple JSON model objects containing random effects (with the same hierarchy / schema) into a single container

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
loadRandomEffectSamplesCombinedJson(json_object_list, json_rfx_num)
```

**Arguments**

```
json_object_list      List of objects of class CppJson
json_rfx_num          Integer index indicating the position of the random effects term to be unpacked
```

**Value**

RandomEffectSamples object

**Examples**

```
n <- 100
p <- 10
X <- matrix(runif(n*p), ncol = p)
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- rep(1.0, n)
y <- (-5 + 10*(X[,1] > 0.5)) + (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
bart_model <- bart(X_train=X, y_train=y, rfx_group_ids_train=rfx_group_ids,
  rfx_basis_train = rfx_basis, num_gfr=0, num_mcmc=10)
bart_json <- list(saveBARTModelToJson(bart_model))
rfx_samples <- loadRandomEffectSamplesCombinedJson(bart_json, 0)
```

---

```
loadRandomEffectSamplesCombinedJsonString
```

*Combine JSON Strings into RandomEffectSamples*

---

**Description**

Combine multiple JSON strings representing model objects containing random effects (with the same hierarchy / schema) into a single container

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
loadRandomEffectSamplesCombinedJsonString(json_string_list, json_rfx_num)
```

**Arguments**

```
json_string_list      List of objects of class CppJson
json_rfx_num          Integer index indicating the position of the random effects term to be unpacked
```

**Value**

RandomEffectSamples object

**Examples**

```
n <- 100
p <- 10
X <- matrix(runif(n*p), ncol = p)
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- rep(1.0, n)
y <- (-5 + 10*(X[,1] > 0.5)) + (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
bart_model <- bart(X_train=X, y_train=y, rfx_group_ids_train=rfx_group_ids,
                  rfx_basis_train = rfx_basis, num_gfr=0, num_mcmc=10)
bart_json_string <- list(saveBARTModelToJsonString(bart_model))
rfx_samples <- loadRandomEffectSamplesCombinedJsonString(bart_json_string, 0)
```

---

loadRandomEffectSamplesJson

*Load Random Effect Samples from JSON*

---

**Description**

Load a container of random effect samples from json

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
loadRandomEffectSamplesJson(json_object, json_rfx_num)
```

**Arguments**

json\_object      Object of class CppJson

json\_rfx\_num      Integer index indicating the position of the random effects term to be unpacked

**Value**

RandomEffectSamples object

**Examples**

```

n <- 100
p <- 10
X <- matrix(runif(n*p), ncol = p)
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- rep(1.0, n)
y <- (-5 + 10*(X[,1] > 0.5)) + (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
bart_model <- bart(X_train=X, y_train=y, rfx_group_ids_train=rfx_group_ids,
                  rfx_basis_train = rfx_basis, num_gfr=0, num_mcmc=10)
bart_json <- saveBARTModelToJson(bart_model)
rfx_samples <- loadRandomEffectSamplesJson(bart_json, 0)

```

loadScalarJson

*Load Scalar from JSON***Description**

Load a scalar from json

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
loadScalarJson(json_object, json_scalar_label, subfolder_name = NULL)
```

**Arguments**

`json_object`      Object of class CppJson

`json_scalar_label`      Label referring to a particular scalar / string value (i.e. "num\_samples") in the overall json hierarchy

`subfolder_name`      (Optional) Name of the subfolder / hierarchy under which vector sits

**Value**

R vector

**Examples**

```

example_scalar <- 5.4
example_json <- createCppJson()
example_json$add_scalar("myscalar", example_scalar)
roundtrip_scalar <- loadScalarJson(example_json, "myscalar")

```

---

|                |                              |
|----------------|------------------------------|
| loadVectorJson | <i>Load Vector from JSON</i> |
|----------------|------------------------------|

---

### Description

Load a vector from json

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

### Usage

```
loadVectorJson(json_object, json_vector_label, subfolder_name = NULL)
```

### Arguments

`json_object`      Object of class CppJson

`json_vector_label`      Label referring to a particular vector (i.e. "sigma2\_global\_samples") in the overall json hierarchy

`subfolder_name`      (Optional) Name of the subfolder / hierarchy under which vector sits

### Value

R vector

### Examples

```
example_vec <- runif(10)
example_json <- createCppJson()
example_json$add_vector("myvec", example_vec)
roundtrip_vec <- loadVectorJson(example_json, "myvec")
```

---

|         |                                 |
|---------|---------------------------------|
| Outcome | <i>Outcome Data C++ Wrapper</i> |
|---------|---------------------------------|

---

### Description

Outcome / partial residual used to sample an additive model. The outcome class is wrapper around a vector of (mutable) outcomes for ML tasks (supervised learning, causal inference). When an additive tree ensemble is sampled, the outcome used to sample a specific model term is the "partial residual" consisting of the outcome minus the predictions of every other model term (trees, group random effects, etc...).

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

### Public fields

`data_ptr` External pointer to a C++ Outcome class

### Methods

#### Public methods:

- `Outcome$new()`
- `Outcome$get_data()`
- `Outcome$add_vector()`
- `Outcome$subtract_vector()`
- `Outcome$update_data()`

**Method** `new()`: Create a new Outcome object.

*Usage:*

`Outcome$new(outcome)`

*Arguments:*

`outcome` Vector of outcome values

*Returns:* A new Outcome object.

**Method** `get_data()`: Extract raw data in R from the underlying C++ object

*Usage:*

`Outcome$get_data()`

*Returns:* R vector containing (copy of) the values in Outcome object

**Method** `add_vector()`: Update the current state of the outcome (i.e. partial residual) data by adding the values of `update_vector`

*Usage:*

`Outcome$add_vector(update_vector)`

*Arguments:*

`update_vector` Vector to be added to outcome

*Returns:* None

**Method** `subtract_vector()`: Update the current state of the outcome (i.e. partial residual) data by subtracting the values of `update_vector`

*Usage:*

`Outcome$subtract_vector(update_vector)`

*Arguments:*

`update_vector` Vector to be subtracted from outcome

*Returns:* None

**Method** `update_data()`: Update the current state of the outcome (i.e. partial residual) data by replacing each element with the elements of `new_vector`

*Usage:*

```
Outcome$update_data(new_vector)
```

*Arguments:*

`new_vector` Vector from which to overwrite the current data

*Returns:* None

---

plot.bartmodel

*Plot BART Model Fit.*


---

## Description

Plot the BART model fit and any relevant sampled quantities. This will default to a traceplot of the global error scale and the in-sample mean forest predictions for the first train set observation. Since `stochtree::bart()` is flexible and it's possible to sample a model with a fixed global error scale and no mean forest, this procedure is adaptive and will attempt to plot a trace of whichever model terms are included if these two default terms are omitted.

## Usage

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

## Arguments

|                  |                       |
|------------------|-----------------------|
| <code>x</code>   | The BART model object |
| <code>...</code> | Additional arguments  |

## Value

BART model object unchanged after summarizing

---

plot.bcfmodel

*Plot BCF Model*


---

### Description

Plot the BCF model fit and any relevant sampled quantities. This will default to a traceplot of the global error scale and the in-sample mean forest predictions for the first train set observation. Since `stochtree::bcf()` is flexible and it's possible to sample a model with a fixed global error scale and no mean forest, this procedure will throw an error if these two default terms are omitted.

### Usage

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

### Arguments

|     |                      |
|-----|----------------------|
| x   | The BCF model object |
| ... | Additional arguments |

### Value

BCF model object unchanged after summarizing

---

predict.bartmodel

*Predict From a BART Model*


---

### Description

Predict from a sampled BART model on new data

### Usage

```
## S3 method for class 'bartmodel'
predict(
  object,
  X,
  leaf_basis = NULL,
  rfx_group_ids = NULL,
  rfx_basis = NULL,
  type = "posterior",
  terms = "all",
  scale = "linear",
  ...
)
```

**Arguments**

|               |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    |
|---------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| object        | Object of type bart containing draws of a regression forest and associated sampling outputs.                                                                                                                                                                                                                                                                                                                                                                                       |
| X             | Covariates used to determine tree leaf predictions for each observation. Must be passed as a matrix or dataframe.                                                                                                                                                                                                                                                                                                                                                                  |
| leaf_basis    | (Optional) Bases used for prediction (by e.g. dot product with leaf values). Default: NULL.                                                                                                                                                                                                                                                                                                                                                                                        |
| rfx_group_ids | (Optional) Test set group labels used for an additive random effects model. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set.                                                                                                                                                                                                                                                                  |
| rfx_basis     | (Optional) Test set basis for "random-slope" regression in additive random effects model.                                                                                                                                                                                                                                                                                                                                                                                          |
| type          | (Optional) Type of prediction to return. Options are "mean", which averages the predictions from every draw of a BART model, and "posterior", which returns the entire matrix of posterior predictions. Default: "posterior".                                                                                                                                                                                                                                                      |
| terms         | (Optional) Which model terms to include in the prediction. This can be a single term or a list of model terms. Options include "y_hat", "mean_forest", "rfx", "variance_forest", or "all". If a model doesn't have mean forest, random effects, or variance forest predictions, but one of those terms is request, the request will simply be ignored. If none of the requested terms are present in a model, this function will return NULL along with a warning. Default: "all". |
| scale         | (Optional) Scale of mean function predictions. Options are "linear", which returns predictions on the original scale of the mean forest / RFX terms, and "probability", which transforms predictions into a probability of observing $y = 1$ . "probability" is only valid for models fit with a probit outcome model. Default: "linear".                                                                                                                                          |
| ...           | (Optional) Other prediction parameters.                                                                                                                                                                                                                                                                                                                                                                                                                                            |

**Value**

List of prediction matrices or single prediction matrix / vector, depending on the terms requested.

**Examples**

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
```

```

n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
                  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
y_hat_test <- predict(bart_model, X=X_test)$y_hat

```

---

|                  |                               |
|------------------|-------------------------------|
| predict.bcfmodel | <i>Predict From BCF Model</i> |
|------------------|-------------------------------|

---

## Description

Predict from a sampled BCF model on new data

## Usage

```

## S3 method for class 'bcfmodel'
predict(
  object,
  X,
  Z,
  propensity = NULL,
  rfx_group_ids = NULL,
  rfx_basis = NULL,
  type = "posterior",
  terms = "all",
  scale = "linear",
  ...
)

```

## Arguments

|               |                                                                                                                                                                                                                   |
|---------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| object        | Object of type bcfmodel containing draws of a Bayesian causal forest model and associated sampling outputs.                                                                                                       |
| X             | Covariates used to determine tree leaf predictions for each observation. Must be passed as a matrix or dataframe.                                                                                                 |
| Z             | Treatments used for prediction.                                                                                                                                                                                   |
| propensity    | (Optional) Propensities used for prediction.                                                                                                                                                                      |
| rfx_group_ids | (Optional) Test set group labels used for an additive random effects model. We do not currently support (but plan to in the near future), test set evaluation for group labels that were not in the training set. |

|           |                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          |
|-----------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| rfx_basis | (Optional) Test set basis for "random-slope" regression in additive random effects model. If the model was sampled with a random effects model_spec of "intercept_only" or "intercept_plus_treatment", this is optional, but if it is provided, it will be used.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         |
| type      | (Optional) Type of prediction to return. Options are "mean", which averages the predictions from every draw of a BCF model, and "posterior", which returns the entire matrix of posterior predictions. Default: "posterior".                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             |
| terms     | (Optional) Which model terms to include in the prediction. This can be a single term or a list of model terms. Options include "y_hat", "prognostic_function", "mu", "cate", "tau", "rfx", "variance_forest", or "all". If a model doesn't have random effects or variance forest predictions, but one of those terms is request, the request will simply be ignored. If a model has random effects fit with either "intercept_only" or "intercept_plus_treatment" model_spec, then "prognostic_function" refers to the predictions of the prognostic forest plus the random intercept and "cate" refers to the predictions of the treatment effect forest plus the random slope on the treatment variable. For these models, the forest predictions alone can be requested via "mu" (prognostic forest) and "tau" (treatment effect forest). In all other cases, "mu" will return exactly the same result as "prognostic_function" and "tau" will return exactly the same result as "cate". If none of the requested terms are present in a model, this function will return NULL along with a warning. Default: "all". |
| scale     | (Optional) Scale of mean function predictions. Options are "linear", which returns predictions on the original scale of the mean forest / RFX terms, and "probability", which transforms predictions into a probability of observing $y = 1$ . "probability" is only valid for models fit with a probit outcome model. Default: "linear".                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                |
| ...       | (Optional) Other prediction parameters.                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  |

## Value

List of prediction matrices or single prediction matrix / vector, depending on the terms requested.

## Examples

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
```

```

tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
noise_sd <- 1
y <- mu_x + tau_x*Z + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train, num_gfr = 10,
  num_burnin = 0, num_mcmc = 10)
preds <- predict(bcf_model, X_test, Z_test, pi_test)

```

---

preprocessPredictionData

*Preprocess Covariates for Model Prediction*


---

## Description

Preprocess covariates for use in a ForestDataset at prediction time. DataFrames will be preprocessed based on their column types. Matrices will be passed through assuming all columns are numeric.

## Usage

```
preprocessPredictionData(input_data, metadata)
```

## Arguments

|            |                                                                                                    |
|------------|----------------------------------------------------------------------------------------------------|
| input_data | Covariates, provided as either a dataframe or a matrix                                             |
| metadata   | List containing information on variables, including train set categories for categorical variables |

**Value**

Preprocessed data with categorical variables appropriately handled

**Examples**

```
cov_df <- data.frame(x1 = 1:5, x2 = 5:1, x3 = 6:10)
metadata <- list(num_ordered_cat_vars = 0, num_unordered_cat_vars = 0,
                 num_numeric_vars = 3, numeric_vars = c("x1", "x2", "x3"))
X_preprocessed <- preprocessPredictionData(cov_df, metadata)
```

---

|                     |                                                 |
|---------------------|-------------------------------------------------|
| preprocessTrainData | <i>Preprocess Covariates for Model Training</i> |
|---------------------|-------------------------------------------------|

---

**Description**

Preprocess covariates for use in a ForestDataset at train time. DataFrames will be preprocessed based on their column types. Matrices will be passed through assuming all columns are numeric.

**Usage**

```
preprocessTrainData(input_data)
```

**Arguments**

|            |                                                        |
|------------|--------------------------------------------------------|
| input_data | Covariates, provided as either a dataframe or a matrix |
|------------|--------------------------------------------------------|

**Value**

List with preprocessed (unmodified) data and details on the number of each type of variable, unique categories associated with categorical variables, and the vector of feature types needed for calls to BART and BCF.

**Examples**

```
cov_mat <- matrix(1:12, ncol = 3)
preprocess_list <- preprocessTrainData(cov_mat)
X <- preprocess_list$X
```

---

|                 |                               |
|-----------------|-------------------------------|
| print.bartmodel | <i>Summarize a BART Model</i> |
|-----------------|-------------------------------|

---

**Description**

Prints a summary of the BART model, including the model terms and their specifications.

**Usage**

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

**Arguments**

|     |                       |
|-----|-----------------------|
| x   | The BART model object |
| ... | Additional arguments  |

**Value**

BART model object unchanged after printing summary

---

|                |                                   |
|----------------|-----------------------------------|
| print.bcfmodel | <i>Print Summary of BCF Model</i> |
|----------------|-----------------------------------|

---

**Description**

Prints a summary of the BCF model, including the model terms and their specifications.

**Usage**

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

**Arguments**

|     |                                         |
|-----|-----------------------------------------|
| x   | The BCF model object                    |
| ... | Additional arguments (currently unused) |

**Value**

BCF model object unchanged after printing summary

---

RandomEffectSamples     *Random Effect Container C++ Wrapper*


---

## Description

Class that wraps the "persistent" aspects of a C++ random effects model, including draws of the parameters and a map from the original label indices to the 0-indexed label numbers used to place group samples in memory (i.e. the first label is stored in column 0 of the sample matrix, the second label is store in column 1 of the sample matrix, etc...)

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

Coordinates various C++ random effects classes and persists those needed for prediction / serialization

## Public fields

rfx\_container\_ptr External pointer to a C++ StochTree::RandomEffectsContainer class

label\_mapper\_ptr External pointer to a C++ StochTree::LabelMapper class

training\_group\_ids Unique vector of group IDs that were in the training dataset

## Methods

### Public methods:

- [RandomEffectSamples\\$new\(\)](#)
- [RandomEffectSamples\\$load\\_in\\_session\(\)](#)
- [RandomEffectSamples\\$load\\_from\\_json\(\)](#)
- [RandomEffectSamples\\$append\\_from\\_json\(\)](#)
- [RandomEffectSamples\\$load\\_from\\_json\\_string\(\)](#)
- [RandomEffectSamples\\$append\\_from\\_json\\_string\(\)](#)
- [RandomEffectSamples\\$predict\(\)](#)
- [RandomEffectSamples\\$extract\\_parameter\\_samples\(\)](#)
- [RandomEffectSamples\\$delete\\_sample\(\)](#)
- [RandomEffectSamples\\$extract\\_label\\_mapping\(\)](#)

**Method** `new()`: Create a new RandomEffectSamples object.

*Usage:*

```
RandomEffectSamples$new()
```

*Returns:* A new RandomEffectSamples object.

**Method** `load_in_session()`: Construct RandomEffectSamples object from other "in-session" R objects

*Usage:*

```
RandomEffectSamples$load_in_session(
  num_components,
  num_groups,
  random_effects_tracker
)
```

*Arguments:*

`num_components` Number of "components" or bases defining the random effects regression

`num_groups` Number of random effects groups

`random_effects_tracker` Object of type RandomEffectsTracker

*Returns:* None

**Method** `load_from_json()`: Construct RandomEffectSamples object from a json object

*Usage:*

```
RandomEffectSamples$load_from_json(
  json_object,
  json_rfx_container_label,
  json_rfx_mapper_label,
  json_rfx_groupids_label
)
```

*Arguments:*

`json_object` Object of class CppJson

`json_rfx_container_label` Label referring to a particular rfx sample container (i.e. "random\_effect\_container\_0") in the overall json hierarchy

`json_rfx_mapper_label` Label referring to a particular rfx label mapper (i.e. "random\_effect\_label\_mapper\_0") in the overall json hierarchy

`json_rfx_groupids_label` Label referring to a particular set of rfx group IDs (i.e. "random\_effect\_groupids\_0") in the overall json hierarchy

*Returns:* A new RandomEffectSamples object.

**Method** `append_from_json()`: Append random effect draws to RandomEffectSamples object from a json object

*Usage:*

```
RandomEffectSamples$append_from_json(
  json_object,
  json_rfx_container_label,
  json_rfx_mapper_label,
  json_rfx_groupids_label
)
```

*Arguments:*

`json_object` Object of class CppJson

`json_rfx_container_label` Label referring to a particular rfx sample container (i.e. "random\_effect\_container\_0") in the overall json hierarchy

`json_rfx_mapper_label` Label referring to a particular rfx label mapper (i.e. "random\_effect\_label\_mapper\_0") in the overall json hierarchy

json\_rfx\_groupids\_label Label referring to a particular set of rfx group IDs (i.e. "random\_effect\_groupids\_0") in the overall json hierarchy

*Returns:* None

**Method** load\_from\_json\_string(): Construct RandomEffectSamples object from a json object

*Usage:*

```
RandomEffectSamples$load_from_json_string(
  json_string,
  json_rfx_container_label,
  json_rfx_mapper_label,
  json_rfx_groupids_label
)
```

*Arguments:*

json\_string JSON string which parses into object of class CppJson

json\_rfx\_container\_label Label referring to a particular rfx sample container (i.e. "random\_effect\_container\_0") in the overall json hierarchy

json\_rfx\_mapper\_label Label referring to a particular rfx label mapper (i.e. "random\_effect\_label\_mapper\_0") in the overall json hierarchy

json\_rfx\_groupids\_label Label referring to a particular set of rfx group IDs (i.e. "random\_effect\_groupids\_0") in the overall json hierarchy

*Returns:* A new RandomEffectSamples object.

**Method** append\_from\_json\_string(): Append random effect draws to RandomEffectSamples object from a json object

*Usage:*

```
RandomEffectSamples$append_from_json_string(
  json_string,
  json_rfx_container_label,
  json_rfx_mapper_label,
  json_rfx_groupids_label
)
```

*Arguments:*

json\_string JSON string which parses into object of class CppJson

json\_rfx\_container\_label Label referring to a particular rfx sample container (i.e. "random\_effect\_container\_0") in the overall json hierarchy

json\_rfx\_mapper\_label Label referring to a particular rfx label mapper (i.e. "random\_effect\_label\_mapper\_0") in the overall json hierarchy

json\_rfx\_groupids\_label Label referring to a particular set of rfx group IDs (i.e. "random\_effect\_groupids\_0") in the overall json hierarchy

*Returns:* None

**Method** predict(): Predict random effects for each observation implied by rfx\_group\_ids and rfx\_basis. If a random effects model is "intercept-only" the rfx\_basis will be a vector of ones of size length(rfx\_group\_ids).

*Usage:*

```
RandomEffectSamples$predict(rfx_group_ids, rfx_basis = NULL)
```

*Arguments:*

`rfx_group_ids` Indices of random effects groups in a prediction set

`rfx_basis` (Optional) Basis used for random effects prediction

*Returns:* Matrix with as many rows as observations provided and as many columns as samples drawn of the model.

**Method** `extract_parameter_samples()`: Extract the random effects parameters sampled. With the "redundant parameterization" of Gelman et al (2008), this includes four parameters: alpha (the "working parameter" shared across every group), xi (the "group parameter" sampled separately for each group), beta (the product of alpha and xi, which corresponds to the overall group-level random effects), and sigma (group-independent prior variance for each component of xi).

*Usage:*

```
RandomEffectSamples$extract_parameter_samples()
```

*Returns:* List of arrays. The alpha array has dimension (num\_components, num\_samples) and is simply a vector if num\_components = 1. The xi and beta arrays have dimension (num\_components, num\_groups, num\_samples) and are simply matrices if num\_components = 1. The sigma array has dimension (num\_components, num\_samples) and is simply a vector if num\_components = 1.

**Method** `delete_sample()`: Modify the RandomEffectsSamples object by removing the parameter samples index by sample\_num.

*Usage:*

```
RandomEffectSamples$delete_sample(sample_num)
```

*Arguments:*

`sample_num` Index of the RFX sample to be removed

**Method** `extract_label_mapping()`: Convert the mapping of group IDs to random effect components indices from C++ to R native format

*Usage:*

```
RandomEffectSamples$extract_label_mapping()
```

*Returns:* List mapping group ID to random effect components.

---

RandomEffectsDataset    *Random Effects Dataset C++ Wrapper*

---

## Description

Dataset used to sample a random effects model. A random effects dataset consists of three matrices / vectors: group labels, bases, and variance weights. Variance weights are optional.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Public fields**

data\_ptr External pointer to a C++ RandomEffectsDataset class

**Methods****Public methods:**

- `RandomEffectsDataset$new()`
- `RandomEffectsDataset$update_basis()`
- `RandomEffectsDataset$update_variance_weights()`
- `RandomEffectsDataset$num_observations()`
- `RandomEffectsDataset$num_basis()`
- `RandomEffectsDataset$get_group_labels()`
- `RandomEffectsDataset$get_basis()`
- `RandomEffectsDataset$get_variance_weights()`
- `RandomEffectsDataset$has_group_labels()`
- `RandomEffectsDataset$has_basis()`
- `RandomEffectsDataset$has_variance_weights()`

**Method** `new()`: Create a new RandomEffectsDataset object.

*Usage:*

```
RandomEffectsDataset$new(group_labels, basis, variance_weights = NULL)
```

*Arguments:*

group\_labels Vector of group labels

basis Matrix of bases used to define the random effects regression (for an intercept-only model, pass an array of ones)

variance\_weights (Optional) Vector of observation-specific variance weights

*Returns:* A new RandomEffectsDataset object.

**Method** `update_basis()`: Update basis matrix in a dataset

*Usage:*

```
RandomEffectsDataset$update_basis(basis)
```

*Arguments:*

basis Updated matrix of bases used to define random slopes / intercepts

**Method** `update_variance_weights()`: Update variance\_weights in a dataset

*Usage:*

```
RandomEffectsDataset$update_variance_weights(  
  variance_weights,  
  exponentiate = F  
)
```

*Arguments:*

variance\_weights Updated vector of variance weights used to define individual variance / case weights

**exponentiate** Whether or not input vector should be exponentiated before being written to the RandomEffectsDataset's variance weights. Default: F.

**Method** num\_observations(): Return number of observations in a RandomEffectsDataset object

*Usage:*

RandomEffectsDataset\$num\_observations()

*Returns:* Observation count

**Method** num\_basis(): Return dimension of the basis matrix in a RandomEffectsDataset object

*Usage:*

RandomEffectsDataset\$num\_basis()

*Returns:* Basis vector count

**Method** get\_group\_labels(): Return group labels as an R vector

*Usage:*

RandomEffectsDataset\$get\_group\_labels()

*Returns:* Group label data

**Method** get\_basis(): Return bases as an R matrix

*Usage:*

RandomEffectsDataset\$get\_basis()

*Returns:* Basis data

**Method** get\_variance\_weights(): Return variance weights as an R vector

*Usage:*

RandomEffectsDataset\$get\_variance\_weights()

*Returns:* Variance weight data

**Method** has\_group\_labels(): Whether or not a dataset has group label indices

*Usage:*

RandomEffectsDataset\$has\_group\_labels()

*Returns:* True if group label vector is loaded, false otherwise

**Method** has\_basis(): Whether or not a dataset has a basis matrix

*Usage:*

RandomEffectsDataset\$has\_basis()

*Returns:* True if basis matrix is loaded, false otherwise

**Method** has\_variance\_weights(): Whether or not a dataset has variance weights

*Usage:*

RandomEffectsDataset\$has\_variance\_weights()

*Returns:* True if variance weights are loaded, false otherwise

---

RandomEffectsModel      *Random Effects Model C++ Wrapper*


---

## Description

The core "model" class for sampling random effects. Stores current model state, prior parameters, and procedures for sampling from the conditional posterior of each parameter.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Public fields

`rfx_model_ptr` External pointer to a C++ `StochTree::RandomEffectsModel` class

`num_groups` Number of groups in the random effects model

`num_components` Number of components (i.e. dimension of basis) in the random effects model

## Methods

### Public methods:

- `RandomEffectsModel$new()`
- `RandomEffectsModel$sample_random_effect()`
- `RandomEffectsModel$predict()`
- `RandomEffectsModel$set_working_parameter()`
- `RandomEffectsModel$set_group_parameters()`
- `RandomEffectsModel$set_working_parameter_cov()`
- `RandomEffectsModel$set_group_parameter_cov()`
- `RandomEffectsModel$set_variance_prior_shape()`
- `RandomEffectsModel$set_variance_prior_scale()`

**Method** `new()`: Create a new `RandomEffectsModel` object.

*Usage:*

```
RandomEffectsModel$new(num_components, num_groups)
```

*Arguments:*

`num_components` Number of "components" or bases defining the random effects regression

`num_groups` Number of random effects groups

*Returns:* A new `RandomEffectsModel` object.

**Method** `sample_random_effect()`: Sample from random effects model.

*Usage:*

```
RandomEffectsModel$sample_random_effect(
  rfx_dataset,
  residual,
  rfx_tracker,
  rfx_samples,
  keep_sample,
  global_variance,
  rng
)
```

*Arguments:*

`rfx_dataset` Object of type RandomEffectsDataset  
`residual` Object of type Outcome  
`rfx_tracker` Object of type RandomEffectsTracker  
`rfx_samples` Object of type RandomEffectSamples  
`keep_sample` Whether sample should be retained in `rfx_samples`. If FALSE, the state of `rfx_tracker` will be updated, but the parameter values will not be added to the sample container. Samples are commonly discarded due to burn-in or thinning.  
`global_variance` Scalar global variance parameter  
`rng` Object of type CppRNG

*Returns:* None

**Method** `predict()`: Predict from (a single sample of a) random effects model.

*Usage:*

```
RandomEffectsModel$predict(rfx_dataset, rfx_tracker)
```

*Arguments:*

`rfx_dataset` Object of type RandomEffectsDataset  
`rfx_tracker` Object of type RandomEffectsTracker

*Returns:* Vector of predictions with size matching number of observations in `rfx_dataset`

**Method** `set_working_parameter()`: Set value for the "working parameter." This is typically used for initialization, but could also be used to interrupt or override the sampler.

*Usage:*

```
RandomEffectsModel$set_working_parameter(value)
```

*Arguments:*

`value` Parameter input

*Returns:* None

**Method** `set_group_parameters()`: Set value for the "group parameters." This is typically used for initialization, but could also be used to interrupt or override the sampler.

*Usage:*

```
RandomEffectsModel$set_group_parameters(value)
```

*Arguments:*

`value` Parameter input

*Returns:* None

**Method** `set_working_parameter_cov()`: Set value for the working parameter covariance. This is typically used for initialization, but could also be used to interrupt or override the sampler.

*Usage:*

```
RandomEffectsModel$set_working_parameter_cov(value)
```

*Arguments:*

value Parameter input

*Returns:* None

**Method** `set_group_parameter_cov()`: Set value for the group parameter covariance. This is typically used for initialization, but could also be used to interrupt or override the sampler.

*Usage:*

```
RandomEffectsModel$set_group_parameter_cov(value)
```

*Arguments:*

value Parameter input

*Returns:* None

**Method** `set_variance_prior_shape()`: Set shape parameter for the group parameter variance prior.

*Usage:*

```
RandomEffectsModel$set_variance_prior_shape(value)
```

*Arguments:*

value Parameter input

*Returns:* None

**Method** `set_variance_prior_scale()`: Set shape parameter for the group parameter variance prior.

*Usage:*

```
RandomEffectsModel$set_variance_prior_scale(value)
```

*Arguments:*

value Parameter input

*Returns:* None

---

RandomEffectsTracker     *Random Effects Tracker C++ Wrapper*


---

### Description

Class that defines a "tracker" for random effects models, most notably storing the data indices available in each group for quicker posterior computation and sampling of random effects terms. The class stores a mapping from every observation to its group index, a mapping from group indices to the training sample observations available in that group, and predictions for each observation.

This class is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

### Public fields

`rfx_tracker_ptr` External pointer to a C++ `StochTree::RandomEffectsTracker` class

### Methods

#### Public methods:

- [RandomEffectsTracker\\$new\(\)](#)

**Method** `new()`: Create a new `RandomEffectsTracker` object.

*Usage:*

`RandomEffectsTracker$new(rfx_group_indices)`

*Arguments:*

`rfx_group_indices` Integer indices indicating groups used to define random effects

*Returns:* A new `RandomEffectsTracker` object.

---

resetActiveForest     *Reset Active Forest*


---

### Description

Reset an active forest, either from a specific forest in a `ForestContainer` or to an ensemble of single-node (i.e. root) trees

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
resetActiveForest(active_forest, forest_samples = NULL, forest_num = NULL)
```

**Arguments**

`active_forest` Current active forest

`forest_samples` (Optional) Container of forest samples from which to re-initialize active forest. If not provided, active forest will be reset to an ensemble of single-node (i.e. root) trees.

`forest_num` (Optional) Index of forest samples from which to initialize active forest. If not provided, active forest will be reset to an ensemble of single-node (i.e. root) trees.

**Value**

None

**Examples**

```
num_trees <- 100
leaf_dimension <- 1
is_leaf_constant <- TRUE
is_exponentiated <- FALSE
active_forest <- createForest(num_trees, leaf_dimension, is_leaf_constant, is_exponentiated)
forest_samples <- createForestSamples(num_trees, leaf_dimension, is_leaf_constant, is_exponentiated)
forest_samples$add_forest_with_constant_leaves(0.0)
forest_samples$add_numeric_split_tree(0, 0, 0, 0, 0.5, -1.0, 1.0)
forest_samples$add_numeric_split_tree(0, 1, 0, 1, 0.75, 3.4, 0.75)
active_forest$set_root_leaves(0.1)
resetActiveForest(active_forest, forest_samples, 0)
resetActiveForest(active_forest)
```

---

resetForestModel

*Reset Forest Model*


---

**Description**

Re-initialize a forest model (tracking data structures) from a specific forest in a ForestContainer

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
resetForestModel(forest_model, forest, dataset, residual, is_mean_model)
```



```

active_forest$prepare_for_sampler(forest_dataset, outcome, forest_model, 0, 0.)
forest_model$sample_one_iteration(
  forest_dataset, outcome, forest_samples, active_forest,
  rng, forest_model_config, global_model_config,
  keep_forest = TRUE, gfr = FALSE
)
resetActiveForest(active_forest, forest_samples, 0)
resetForestModel(forest_model, active_forest, forest_dataset, outcome, TRUE)

```

---

resetRandomEffectsModel

*Reset RandomEffectsModel Object*


---

## Description

Reset a RandomEffectsModel object based on the parameters indexed by sample\_num in a RandomEffectsSamples object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Usage

```
resetRandomEffectsModel(rfx_model, rfx_samples, sample_num, sigma_alpha_init)
```

## Arguments

|                  |                                                                                                                                                                                           |
|------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| rfx_model        | Object of type RandomEffectsModel.                                                                                                                                                        |
| rfx_samples      | Object of type RandomEffectSamples.                                                                                                                                                       |
| sample_num       | Index of sample stored in rfx_samples from which to reset the state of a random effects model. Zero-indexed, so resetting based on the first sample would require setting sample_num = 0. |
| sigma_alpha_init | Initial value of the "working parameter" scale parameter.                                                                                                                                 |

## Value

None

## Examples

```

n <- 100
p <- 10
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
rfx_dataset <- createRandomEffectsDataset(rfx_group_ids, rfx_basis)

```

```

y <- (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
y_std <- (y-mean(y))/sd(y)
outcome <- createOutcome(y_std)
rng <- createCppRNG(1234)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_model <- createRandomEffectsModel(num_components, num_groups)
rfx_tracker <- createRandomEffectsTracker(rfx_group_ids)
rfx_samples <- createRandomEffectSamples(num_components, num_groups, rfx_tracker)
alpha_init <- rep(1,num_components)
xi_init <- matrix(rep(alpha_init, num_groups),num_components,num_groups)
sigma_alpha_init <- diag(1,num_components,num_components)
sigma_xi_init <- diag(1,num_components,num_components)
sigma_xi_shape <- 1
sigma_xi_scale <- 1
rfx_model$set_working_parameter(alpha_init)
rfx_model$set_group_parameters(xi_init)
rfx_model$set_working_parameter_cov(sigma_alpha_init)
rfx_model$set_group_parameter_cov(sigma_xi_init)
rfx_model$set_variance_prior_shape(sigma_xi_shape)
rfx_model$set_variance_prior_scale(sigma_xi_scale)
for (i in 1:3) {
  rfx_model$sample_random_effect(rfx_dataset=rfx_dataset, residual=outcome,
                                rfx_tracker=rfx_tracker, rfx_samples=rfx_samples,
                                keep_sample=TRUE, global_variance=1.0, rng=rng)
}
resetRandomEffectsModel(rfx_model, rfx_samples, 0, 1.0)

```

---

resetRandomEffectsTracker

*Reset RandomEffectsTracker Object*

---

## Description

Reset a RandomEffectsTracker object based on the parameters indexed by sample\_num in a RandomEffectSamples object

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Usage

```

resetRandomEffectsTracker(
  rfx_tracker,
  rfx_model,
  rfx_dataset,
  residual,
  rfx_samples
)

```

**Arguments**

|             |                                      |
|-------------|--------------------------------------|
| rfx_tracker | Object of type RandomEffectsTracker. |
| rfx_model   | Object of type RandomEffectsModel.   |
| rfx_dataset | Object of type RandomEffectsDataset. |
| residual    | Object of type Outcome.              |
| rfx_samples | Object of type RandomEffectSamples.  |

**Value**

None

**Examples**

```

n <- 100
p <- 10
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
rfx_dataset <- createRandomEffectsDataset(rfx_group_ids, rfx_basis)
y <- (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
y_std <- (y-mean(y))/sd(y)
outcome <- createOutcome(y_std)
rng <- createCppRNG(1234)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_model <- createRandomEffectsModel(num_components, num_groups)
rfx_tracker <- createRandomEffectsTracker(rfx_group_ids)
rfx_samples <- createRandomEffectSamples(num_components, num_groups, rfx_tracker)
alpha_init <- rep(1,num_components)
xi_init <- matrix(rep(alpha_init, num_groups),num_components,num_groups)
sigma_alpha_init <- diag(1,num_components,num_components)
sigma_xi_init <- diag(1,num_components,num_components)
sigma_xi_shape <- 1
sigma_xi_scale <- 1
rfx_model$set_working_parameter(alpha_init)
rfx_model$set_group_parameters(xi_init)
rfx_model$set_working_parameter_cov(sigma_alpha_init)
rfx_model$set_group_parameter_cov(sigma_xi_init)
rfx_model$set_variance_prior_shape(sigma_xi_shape)
rfx_model$set_variance_prior_scale(sigma_xi_scale)
for (i in 1:3) {
  rfx_model$sample_random_effect(rfx_dataset=rfx_dataset, residual=outcome,
                                rfx_tracker=rfx_tracker, rfx_samples=rfx_samples,
                                keep_sample=TRUE, global_variance=1.0, rng=rng)
}
resetRandomEffectsModel(rfx_model, rfx_samples, 0, 1.0)
resetRandomEffectsTracker(rfx_tracker, rfx_model, rfx_dataset, outcome, rfx_samples)

```

---

rootResetRandomEffectsModel

*Reset RandomEffectsModel Object to Default State*


---

## Description

Reset a RandomEffectsModel object to its "default" state

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Usage

```
rootResetRandomEffectsModel(
  rfx_model,
  alpha_init,
  xi_init,
  sigma_alpha_init,
  sigma_xi_init,
  sigma_xi_shape,
  sigma_xi_scale
)
```

## Arguments

|                  |                                                                               |
|------------------|-------------------------------------------------------------------------------|
| rfx_model        | Object of type RandomEffectsModel.                                            |
| alpha_init       | Initial value of the "working parameter".                                     |
| xi_init          | Initial value of the "group parameters".                                      |
| sigma_alpha_init | Initial value of the "working parameter" scale parameter.                     |
| sigma_xi_init    | Initial value of the "group parameters" scale parameter.                      |
| sigma_xi_shape   | Shape parameter for the inverse gamma variance model on the group parameters. |
| sigma_xi_scale   | Scale parameter for the inverse gamma variance model on the group parameters. |

## Value

None

## Examples

```
n <- 100
p <- 10
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
rfx_dataset <- createRandomEffectsDataset(rfx_group_ids, rfx_basis)
```

```

y <- (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
y_std <- (y-mean(y))/sd(y)
outcome <- createOutcome(y_std)
rng <- createCppRNG(1234)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_model <- createRandomEffectsModel(num_components, num_groups)
rfx_tracker <- createRandomEffectsTracker(rfx_group_ids)
rfx_samples <- createRandomEffectSamples(num_components, num_groups, rfx_tracker)
alpha_init <- rep(1,num_components)
xi_init <- matrix(rep(alpha_init, num_groups),num_components,num_groups)
sigma_alpha_init <- diag(1,num_components,num_components)
sigma_xi_init <- diag(1,num_components,num_components)
sigma_xi_shape <- 1
sigma_xi_scale <- 1
rfx_model$set_working_parameter(alpha_init)
rfx_model$set_group_parameters(xi_init)
rfx_model$set_working_parameter_cov(sigma_alpha_init)
rfx_model$set_group_parameter_cov(sigma_xi_init)
rfx_model$set_variance_prior_shape(sigma_xi_shape)
rfx_model$set_variance_prior_scale(sigma_xi_scale)
for (i in 1:3) {
  rfx_model$sample_random_effect(rfx_dataset=rfx_dataset, residual=outcome,
                                rfx_tracker=rfx_tracker, rfx_samples=rfx_samples,
                                keep_sample=TRUE, global_variance=1.0, rng=rng)
}
rootResetRandomEffectsModel(rfx_model, alpha_init, xi_init, sigma_alpha_init,
                             sigma_xi_init, sigma_xi_shape, sigma_xi_scale)

```

---

rootResetRandomEffectsTracker

*Reset RandomEffectsTracker Object to Default State*


---

## Description

Reset a RandomEffectsTracker object to its "default" state

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Usage

```
rootResetRandomEffectsTracker(rfx_tracker, rfx_model, rfx_dataset, residual)
```

## Arguments

|             |                                      |
|-------------|--------------------------------------|
| rfx_tracker | Object of type RandomEffectsTracker. |
| rfx_model   | Object of type RandomEffectsModel.   |

`rfx_dataset`      Object of type `RandomEffectsDataset`.  
`residual`        Object of type `Outcome`.

### Value

None

### Examples

```
n <- 100
p <- 10
rfx_group_ids <- sample(1:2, size = n, replace = TRUE)
rfx_basis <- matrix(rep(1.0, n), ncol=1)
rfx_dataset <- createRandomEffectsDataset(rfx_group_ids, rfx_basis)
y <- (-2*(rfx_group_ids==1)+2*(rfx_group_ids==2)) + rnorm(n)
y_std <- (y-mean(y))/sd(y)
outcome <- createOutcome(y_std)
rng <- createCppRNG(1234)
num_groups <- length(unique(rfx_group_ids))
num_components <- ncol(rfx_basis)
rfx_model <- createRandomEffectsModel(num_components, num_groups)
rfx_tracker <- createRandomEffectsTracker(rfx_group_ids)
rfx_samples <- createRandomEffectSamples(num_components, num_groups, rfx_tracker)
alpha_init <- rep(1,num_components)
xi_init <- matrix(rep(alpha_init, num_groups),num_components,num_groups)
sigma_alpha_init <- diag(1,num_components,num_components)
sigma_xi_init <- diag(1,num_components,num_components)
sigma_xi_shape <- 1
sigma_xi_scale <- 1
rfx_model$set_working_parameter(alpha_init)
rfx_model$set_group_parameters(xi_init)
rfx_model$set_working_parameter_cov(sigma_alpha_init)
rfx_model$set_group_parameter_cov(sigma_xi_init)
rfx_model$set_variance_prior_shape(sigma_xi_shape)
rfx_model$set_variance_prior_scale(sigma_xi_scale)
for (i in 1:3) {
  rfx_model$sample_random_effect(rfx_dataset=rfx_dataset, residual=outcome,
                                rfx_tracker=rfx_tracker, rfx_samples=rfx_samples,
                                keep_sample=TRUE, global_variance=1.0, rng=rng)
}
rootResetRandomEffectsModel(rfx_model, alpha_init, xi_init, sigma_alpha_init,
                             sigma_xi_init, sigma_xi_shape, sigma_xi_scale)
rootResetRandomEffectsTracker(rfx_tracker, rfx_model, rfx_dataset, outcome)
```

**Description**

Sample one iteration of the (inverse gamma) global variance model

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
sampleGlobalErrorVarianceOneIteration(residual, dataset, rng, a, b)
```

**Arguments**

|          |                                 |
|----------|---------------------------------|
| residual | Outcome class                   |
| dataset  | ForestDataset class             |
| rng      | C++ random number generator     |
| a        | Global variance shape parameter |
| b        | Global variance scale parameter |

**Value**

None

**Examples**

```
X <- matrix(runif(10*100), ncol = 10)
y <- -5 + 10*(X[,1] > 0.5) + rnorm(100)
y_std <- (y-mean(y))/sd(y)
forest_dataset <- createForestDataset(X)
outcome <- createOutcome(y_std)
rng <- createCppRNG(1234)
a <- 1.0
b <- 1.0
sigma2 <- sampleGlobalErrorVarianceOneIteration(outcome, forest_dataset, rng, a, b)
```

---

```
sampleLeafVarianceOneIteration
```

*Sample Leaf Scale*

---

**Description**

Sample one iteration of the leaf parameter variance model (only for univariate basis and constant leaf!)

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

**Usage**

```
sampleLeafVarianceOneIteration(forest, rng, a, b)
```

**Arguments**

|        |                               |
|--------|-------------------------------|
| forest | C++ forest                    |
| rng    | C++ random number generator   |
| a      | Leaf variance shape parameter |
| b      | Leaf variance scale parameter |

**Value**

None

**Examples**

```
num_trees <- 100
leaf_dimension <- 1
is_leaf_constant <- TRUE
is_exponentiated <- FALSE
active_forest <- createForest(num_trees, leaf_dimension, is_leaf_constant, is_exponentiated)
rng <- createCppRNG(1234)
a <- 1.0
b <- 1.0
tau <- sampleLeafVarianceOneIteration(active_forest, rng, a, b)
```

---

```
sample_bart_posterior_predictive
```

*Sample BART Posterior Predictive*

---

**Description**

Sample from the posterior predictive distribution for outcomes modeled by BART

**Usage**

```
sample_bart_posterior_predictive(
  model_object,
  X = NULL,
  leaf_basis = NULL,
  rfx_group_ids = NULL,
  rfx_basis = NULL,
  num_draws_per_sample = NULL
)
```

**Arguments**

|                      |                                                                                                                                                                                                                                                                                                                                               |
|----------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| model_object         | A fitted BART model object of class bartmodel.                                                                                                                                                                                                                                                                                                |
| X                    | A matrix or data frame of covariates. Required if the BART model depends on covariates (e.g., contains a mean or variance forest).                                                                                                                                                                                                            |
| leaf_basis           | A matrix of bases for mean forest models with regression defined in the leaves. Required for "leaf regression" models.                                                                                                                                                                                                                        |
| rfx_group_ids        | A vector of group IDs for random effects model. Required if the BART model includes random effects.                                                                                                                                                                                                                                           |
| rfx_basis            | A matrix of bases for random effects model. Required if the BART model includes random effects.                                                                                                                                                                                                                                               |
| num_draws_per_sample | The number of posterior predictive samples to draw for each posterior sample. Defaults to a heuristic based on the number of samples in a BART model (i.e. if the BART model has >1000 draws, we use 1 draw from the likelihood per sample, otherwise we upsample to ensure intervals are based on at least 1000 posterior predictive draws). |

**Value**

Array of posterior predictive samples with dimensions (num\_observations, num\_posterior\_samples, num\_draws\_per\_sample) if num\_draws\_per\_sample > 1, otherwise (num\_observations, num\_posterior\_samples).

**Examples**

```
n <- 100
p <- 5
X <- matrix(rnorm(n * p), nrow = n, ncol = p)
y <- 2 * X[,1] + rnorm(n)
bart_model <- bart(y_train = y, X_train = X)
ppd_samples <- sample_bart_posterior_predictive(
  model_object = bart_model, X = X
)
```

---

sample\_bcf\_posterior\_predictive

*Sample BCF Posterior Predictive*

---

**Description**

Sample from the posterior predictive distribution for outcomes modeled by BCF

**Usage**

```
sample_bcf_posterior_predictive(
  model_object,
  X = NULL,
  Z = NULL,
  propensity = NULL,
  rfx_group_ids = NULL,
  rfx_basis = NULL,
  num_draws_per_sample = NULL
)
```

**Arguments**

|                                   |                                                                                                                                                                                                                                                                                                                                     |
|-----------------------------------|-------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| <code>model_object</code>         | A fitted BCF model object of class <code>bcfmodel</code> .                                                                                                                                                                                                                                                                          |
| <code>X</code>                    | A matrix or data frame of covariates.                                                                                                                                                                                                                                                                                               |
| <code>Z</code>                    | A vector or matrix of treatment assignments.                                                                                                                                                                                                                                                                                        |
| <code>propensity</code>           | (Optional) A vector or matrix of propensity scores. Required if the underlying model depends on user-provided propensities.                                                                                                                                                                                                         |
| <code>rfx_group_ids</code>        | (Optional) A vector of group IDs for random effects model. Required if the BCF model includes random effects.                                                                                                                                                                                                                       |
| <code>rfx_basis</code>            | (Optional) A matrix of bases for random effects model. Required if the BCF model includes random effects.                                                                                                                                                                                                                           |
| <code>num_draws_per_sample</code> | (Optional) The number of samples to draw from the likelihood for each draw of the posterior. Defaults to a heuristic based on the number of samples in a BCF model (i.e. if the BCF model has >1000 draws, we use 1 draw from the likelihood per sample, otherwise we upsample to ensure at least 1000 posterior predictive draws). |

**Value**

Array of posterior predictive samples with dimensions `(num_observations, num_posterior_samples, num_draws_per_sample)` if `num_draws_per_sample > 1`, otherwise `(num_observations, num_posterior_samples)`.

**Examples**

```
n <- 100
p <- 5
X <- matrix(rnorm(n * p), nrow = n, ncol = p)
pi_X <- pnorm(X[,1] / 2)
Z <- rbinom(n, 1, pi_X)
y <- 2 * X[,2] + 0.5 * X[,2] * Z + rnorm(n)
bcf_model <- bcf(X_train = X, Z_train = Z, y_train = y, propensity_train = pi_X)
ppd_samples <- sample_bcf_posterior_predictive(
  model_object = bcf_model, X = X,
  Z = Z, propensity = pi_X
)
```

---

`sample_without_replacement`*Sample Without Replacement*

---

## Description

Draw `sample_size` samples from `population_vector` without replacement, weighted by `sampling_probabilities`

This function is intended for advanced use cases in which users require detailed control of sampling algorithms and data structures. Minimal input validation and error checks are performed – users are responsible for providing the correct inputs. For tutorials on the "proper" usage of the stochtree's advanced workflow, we provide several vignettes at [stochtree.ai](http://stochtree.ai)

## Usage

```
sample_without_replacement(  
  population_vector,  
  sampling_probabilities,  
  sample_size  
)
```

## Arguments

|                                     |                                                                                                                                      |
|-------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------|
| <code>population_vector</code>      | Vector from which to draw samples.                                                                                                   |
| <code>sampling_probabilities</code> | Vector of probabilities of drawing each element of <code>population_vector</code> .                                                  |
| <code>sample_size</code>            | Number of samples to draw from <code>population_vector</code> . Must be less than or equal to <code>length(population_vector)</code> |

## Value

Vector of size `sample_size`

## Examples

```
a <- as.integer(c(4,3,2,5,1,9,7))  
p <- c(0.7,0.2,0.05,0.02,0.01,0.01,0.01)  
num_samples <- 5  
sample_without_replacement(a, p, num_samples)
```

---

|                     |                                   |
|---------------------|-----------------------------------|
| saveBARTModelToJson | <i>Convert BART Model to JSON</i> |
|---------------------|-----------------------------------|

---

### Description

Convert the persistent aspects of a BART model to (in-memory) JSON

### Usage

```
saveBARTModelToJson(object)
```

### Arguments

|        |                                                                                                         |
|--------|---------------------------------------------------------------------------------------------------------|
| object | Object of type <code>bartmodel</code> containing draws of a BART model and associated sampling outputs. |
|--------|---------------------------------------------------------------------------------------------------------|

### Value

Object of type `CppJson`

### Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bart_json <- saveBARTModelToJson(bart_model)
```

---

saveBARTModelToJsonFile

*Save BART Model to JSON File*


---

## Description

Convert the persistent aspects of a BART model to (in-memory) JSON and save to a file

## Usage

```
saveBARTModelToJsonFile(object, filename)
```

## Arguments

|          |                                                                                            |
|----------|--------------------------------------------------------------------------------------------|
| object   | Object of type bartmodel containing draws of a BART model and associated sampling outputs. |
| filename | String of filepath, must end in ".json"                                                    |

## Value

None

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
tmpjson <- tempfile(fileext = ".json")
saveBARTModelToJsonFile(bart_model, file.path(tmpjson))
unlink(tmpjson)
```

---

saveBARTModelToJsonString

*Convert BART Model to JSON String*


---

## Description

Convert the persistent aspects of a BART model to (in-memory) JSON string

## Usage

```
saveBARTModelToJsonString(object)
```

## Arguments

|        |                                                                                            |
|--------|--------------------------------------------------------------------------------------------|
| object | Object of type bartmodel containing draws of a BART model and associated sampling outputs. |
|--------|--------------------------------------------------------------------------------------------|

## Value

in-memory JSON string

## Examples

```
n <- 100
p <- 5
X <- matrix(runif(n*p), ncol = p)
f_XW <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
noise_sd <- 1
y <- f_XW + rnorm(n, 0, noise_sd)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
y_test <- y[test_inds]
y_train <- y[train_inds]
bart_model <- bart(X_train = X_train, y_train = y_train,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10)
bart_json_string <- saveBARTModelToJsonString(bart_model)
```

---

|                    |                                  |
|--------------------|----------------------------------|
| saveBCFModelToJson | <i>Convert BCF Model to JSON</i> |
|--------------------|----------------------------------|

---

**Description**

Convert the persistent aspects of a BCF model to (in-memory) JSON

**Usage**

```
saveBCFModelToJson(object)
```

**Arguments**

|        |                                                                                                                          |
|--------|--------------------------------------------------------------------------------------------------------------------------|
| object | Object of type <code>bcfmodel</code> containing draws of a Bayesian causal forest model and associated sampling outputs. |
|--------|--------------------------------------------------------------------------------------------------------------------------|

**Value**

Object of type `CppJson`

**Examples**

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
```

```

y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
mu_params <- list(sample_sigma2_leaf = TRUE)
tau_params <- list(sample_sigma2_leaf = FALSE)
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10,
  prognostic_forest_params = mu_params,
  treatment_effect_forest_params = tau_params)
bcf_json <- saveBCFModelToJson(bcf_model)

```

---

saveBCFModelToJsonFile

*Save BCF Model to JSON File*

---

## Description

Convert the persistent aspects of a BCF model to (in-memory) JSON and save to a file

## Usage

```
saveBCFModelToJsonFile(object, filename)
```

**Arguments**

|          |                                                                                                                          |
|----------|--------------------------------------------------------------------------------------------------------------------------|
| object   | Object of type <code>bcfmodel</code> containing draws of a Bayesian causal forest model and associated sampling outputs. |
| filename | String of filepath, must end in ".json"                                                                                  |

**Value**

in-memory JSON string

**Examples**

```
n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %/% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
```

```

y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]
rfx_term_train <- rfx_term[train_inds]
mu_params <- list(sample_sigma2_leaf = TRUE)
tau_params <- list(sample_sigma2_leaf = FALSE)
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10,
  prognostic_forest_params = mu_params,
  treatment_effect_forest_params = tau_params)
tmpjson <- tempfile(fileext = ".json")
saveBCFModelToJsonFile(bcf_model, file.path(tmpjson))
unlink(tmpjson)

```

---

saveBCFModelToJsonString

*Convert BCF Model to JSON String*

---

## Description

Convert the persistent aspects of a BCF model to (in-memory) JSON string

## Usage

```
saveBCFModelToJsonString(object)
```

## Arguments

|        |                                                                                                                          |
|--------|--------------------------------------------------------------------------------------------------------------------------|
| object | Object of type <code>bcfmodel</code> containing draws of a Bayesian causal forest model and associated sampling outputs. |
|--------|--------------------------------------------------------------------------------------------------------------------------|

## Value

JSON string

**Examples**

```

n <- 500
p <- 5
X <- matrix(runif(n*p), ncol = p)
mu_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (-7.5) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (-2.5) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (2.5) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (7.5)
)
pi_x <- (
  ((0 <= X[,1]) & (0.25 > X[,1])) * (0.2) +
  ((0.25 <= X[,1]) & (0.5 > X[,1])) * (0.4) +
  ((0.5 <= X[,1]) & (0.75 > X[,1])) * (0.6) +
  ((0.75 <= X[,1]) & (1 > X[,1])) * (0.8)
)
tau_x <- (
  ((0 <= X[,2]) & (0.25 > X[,2])) * (0.5) +
  ((0.25 <= X[,2]) & (0.5 > X[,2])) * (1.0) +
  ((0.5 <= X[,2]) & (0.75 > X[,2])) * (1.5) +
  ((0.75 <= X[,2]) & (1 > X[,2])) * (2.0)
)
Z <- rbinom(n, 1, pi_x)
E_XZ <- mu_x + Z*tau_x
snr <- 3
rfx_group_ids <- rep(c(1,2), n %% 2)
rfx_coefs <- matrix(c(-1, -1, 1, 1), nrow=2, byrow=TRUE)
rfx_basis <- cbind(1, runif(n, -1, 1))
rfx_term <- rowSums(rfx_coefs[rfx_group_ids,] * rfx_basis)
y <- E_XZ + rfx_term + rnorm(n, 0, 1)*(sd(E_XZ)/snr)
test_set_pct <- 0.2
n_test <- round(test_set_pct*n)
n_train <- n - n_test
test_inds <- sort(sample(1:n, n_test, replace = FALSE))
train_inds <- (1:n)[!((1:n) %in% test_inds)]
X_test <- X[test_inds,]
X_train <- X[train_inds,]
pi_test <- pi_x[test_inds]
pi_train <- pi_x[train_inds]
Z_test <- Z[test_inds]
Z_train <- Z[train_inds]
y_test <- y[test_inds]
y_train <- y[train_inds]
mu_test <- mu_x[test_inds]
mu_train <- mu_x[train_inds]
tau_test <- tau_x[test_inds]
tau_train <- tau_x[train_inds]
rfx_group_ids_test <- rfx_group_ids[test_inds]
rfx_group_ids_train <- rfx_group_ids[train_inds]
rfx_basis_test <- rfx_basis[test_inds,]
rfx_basis_train <- rfx_basis[train_inds,]
rfx_term_test <- rfx_term[test_inds]

```

```

rfx_term_train <- rfx_term[train_inds]
mu_params <- list(sample_sigma2_leaf = TRUE)
tau_params <- list(sample_sigma2_leaf = FALSE)
bcf_model <- bcf(X_train = X_train, Z_train = Z_train, y_train = y_train,
  propensity_train = pi_train,
  rfx_group_ids_train = rfx_group_ids_train,
  rfx_basis_train = rfx_basis_train, X_test = X_test,
  Z_test = Z_test, propensity_test = pi_test,
  rfx_group_ids_test = rfx_group_ids_test,
  rfx_basis_test = rfx_basis_test,
  num_gfr = 10, num_burnin = 0, num_mcmc = 10,
  prognostic_forest_params = mu_params,
  treatment_effect_forest_params = tau_params)
saveBCFModelToJsonString(bcf_model)

```

---

savePreprocessorToJsonString

*Convert Covariate Preprocessor to JSON String*

---

## Description

Convert the persistent aspects of a covariate preprocessor to (in-memory) JSON string

## Usage

```
savePreprocessorToJsonString(object)
```

## Arguments

|        |                                                                                                    |
|--------|----------------------------------------------------------------------------------------------------|
| object | List containing information on variables, including train set categories for categorical variables |
|--------|----------------------------------------------------------------------------------------------------|

## Value

in-memory JSON string

## Examples

```

cov_mat <- matrix(1:12, ncol = 3)
preprocess_list <- preprocessTrainData(cov_mat)
preprocessor_json_string <- savePreprocessorToJsonString(preprocess_list$metadata)

```

---

|                   |                                                        |
|-------------------|--------------------------------------------------------|
| summary.bartmodel | <i>Summarize the BART model fit and sampled terms.</i> |
|-------------------|--------------------------------------------------------|

---

**Description**

Summarize the BART with a description of the model that was fit and numeric summaries of any sampled quantities.

**Usage**

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

**Arguments**

|        |                       |
|--------|-----------------------|
| object | The BART model object |
| ...    | Additional arguments  |

**Value**

BART model object unchanged after summarizing

---

|                  |                            |
|------------------|----------------------------|
| summary.bcfmodel | <i>Summarize BCF Model</i> |
|------------------|----------------------------|

---

**Description**

Summarize the BCF with a description of the model that was fit and numeric summaries of any sampled quantities.

**Usage**

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

**Arguments**

|        |                      |
|--------|----------------------|
| object | The BCF model object |
| ...    | Additional arguments |

**Value**

BCF model object unchanged after summarizing

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