

Package: h2o4gpu (via r-universe)

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Type Package

Title Interface to 'H2O4GPU'

Version 0.3.3

Description Interface to 'H2O4GPU' <<https://github.com/h2oai/h2o4gpu>>, a collection of 'GPU' solvers for machine learning algorithms.

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URL <https://github.com/h2oai/h2o4gpu>

BugReports <https://github.com/h2oai/h2o4gpu/issues>

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Encoding UTF-8

Depends R (>= 3.1)

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Author Yuan Tang [aut] (<<https://orcid.org/0000-0001-5243-233X>>), Navdeep Gill [aut, cre], Erin LeDell [aut], Vladimir Ovsyannikov [aut], H2O.ai [cph, fnd]

Maintainer Navdeep Gill <navdeep@h2o.ai>

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 fit

Generic Method to Train an H2O4GPU Estimator

Description

Generic Method to Train an H2O4GPU Estimator

Generic Method to Transform a Dataset using Trained H2O4GPU Estimator

Usage

```
fit(object, ...)
```

```
transform(object, ...)
```

Arguments

object	The h2o4gpu model object
...	Additional arguments (unused for now).

fit.h2o4gpu_model	<i>Train an H2O4GPU Estimator</i>
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Description

This function builds the model using the training data specified.

Usage

```
## S3 method for class 'h2o4gpu_model'  
fit(object, x, y = NULL, ...)
```

Arguments

object	The h2o4gpu model object
x	The training data where each column represents a different predictor variable to be used in building the model.
y	A vector of numeric values to be used as response variable in building the model. Note that if the vector is character or factor, it will be converted to numeric column (e.g. 0, 1, 2, ...) implicitly. For unsupervised models, this argument can be ignored or specified as NULL.
...	Additional arguments (unused for now).

Examples

```
## Not run:  
  
library(h2o4gpu)  
  
# Setup dataset  
x <- iris[1:4]  
y <- as.integer(iris$Species) - 1  
  
# Train the classifier  
h2o4gpu.random_forest_classifier() %>% fit(x, y)  
  
## End(Not run)
```

h2o4gpu	<i>h2o4gpu in R</i>
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Description

h2o4gpu in R

Examples

```
## Not run:

library(h2o4gpu)

# Setup dataset
x <- iris[1:4]
y <- as.integer(iris$Species) - 1

# Initialize and train the classifier
model <- h2o4gpu.random_forest_classifier() %>% fit(x, y)

# Make predictions
predictions <- model %>% predict(x)

## End(Not run)
```

```
h2o4gpu.elastic_net_classifier
      Elastic Net Classifier
```

Description

Elastic Net Classifier

Usage

```
h2o4gpu.elastic_net_classifier(alpha = 1, l1_ratio = 0.5,
  fit_intercept = TRUE, normalize = FALSE, precompute = FALSE,
  max_iter = 5000L, copy_X = TRUE, tol = 0.01, warm_start = FALSE,
  positive = FALSE, random_state = NULL, selection = "cyclic",
  n_gpus = -1L, lambda_stop_early = TRUE, glm_stop_early = TRUE,
  glm_stop_early_error_fraction = 1, verbose = FALSE, n_threads = NULL,
  gpu_id = 0L, lambda_min_ratio = 1e-07, n_lambdas = 100L, n_folds = 5L,
  tol_seek_factor = 0.1, store_full_path = 0L, lambda_max = NULL,
  lambdas = NULL, double_precision = NULL, order = NULL,
  backend = "h2o4gpu")
```

Arguments

alpha Constant that multiplies the penalty terms. Defaults to 1.0. See the notes for the exact mathematical meaning of this parameter. $\alpha = 0$ is equivalent to an ordinary least square, solved by the `:class:LinearRegressionSklearn` object. For numerical reasons, using $\alpha = 0$ with the `LassoSklearn` object is not advised. Given this, you should use the `:class:LinearRegressionSklearn` object.

l1_ratio	The ElasticNetSklearn mixing parameter, with $0 \leq \text{l1_ratio} \leq 1$. For $\text{l1_ratio} = 0$ the penalty is an L2 penalty. For $\text{l1_ratio} = 1$ it is an L1 penalty. For $0 < \text{l1_ratio} < 1$, the penalty is a combination of L1 and L2.
fit_intercept	Whether the intercept should be estimated or not. If FALSE, the data is assumed to be already centered.
normalize	This parameter is ignored when <code>fit_intercept</code> is set to FALSE. If TRUE, the regressors X will be normalized before regression by subtracting the mean and dividing by the l2-norm. If you wish to standardize, please use <code>:class:h2o4gpu.preprocessing.Standardize</code> before calling <code>fit</code> on an estimator with <code>normalize=FALSE</code> .
precompute	Whether to use a precomputed Gram matrix to speed up calculations. The Gram matrix can also be passed as argument. For sparse input this option is always TRUE to preserve sparsity.
max_iter	The maximum number of iterations
copy_X	If TRUE, X will be copied; else, it may be overwritten.
tol	The tolerance for the optimization: if the updates are smaller than <code>tol</code> , the optimization code checks the dual gap for optimality and continues until it is smaller than <code>tol</code> .
warm_start	When set to TRUE, reuse the solution of the previous call to <code>fit</code> as initialization, otherwise, just erase the previous solution.
positive	When set to TRUE, forces the coefficients to be positive.
random_state	The seed of the pseudo random number generator that selects a random feature to update. If int, <code>random_state</code> is the seed used by the random number generator; If RandomState instance, <code>random_state</code> is the random number generator; If NULL, the random number generator is the RandomState instance used by <code>np.random</code> . Used when <code>selection == 'random'</code> .
selection	If set to 'random', a random coefficient is updated every iteration rather than looping over features sequentially by default. This (setting to 'random') often leads to significantly faster convergence especially when <code>tol</code> is higher than $1e-4$.
n_gpus	Number of gpu's to use in GLM solver.
lambda_stop_early	Stop early when there is no more relative improvement on train or validation.
glm_stop_early	Stop early when there is no more relative improvement in the primary and dual residuals for ADMM.
glm_stop_early_error_fraction	Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much).
verbose	Print verbose information to the console if set to > 0 .
n_threads	Number of threads to use in the gpu. Each thread is an independent model builder.
gpu_id	ID of the GPU on which the algorithm should run.
lambda_min_ratio	Minimum lambda ratio to maximum lambda, used in lambda search.
n_lambdas	Number of lambdas to be used in a search.

n_folds	Number of cross validation folds.
tol_seek_factor	Factor of tolerance to seek once below null model accuracy. Default is 1E-1, so seeks tolerance of 1E-3 once below null model accuracy for tol=1E-2.
store_full_path	Whether to store full solution for all alphas and lambdas. If 1, then during predict will compute best and full predictions.
lambda_max	Maximum Lambda value to use. Default is NULL, and then internally compute standard maximum
lambdas	overrides n_lambdas, lambda_max, and lambda_min_ratio.
double_precision	Internally set unless using _ptr methods. Value can either be 0 (float32) or 1(float64)
order	Order of data. Default is NULL, and internally determined (unless using _ptr methods) whether row 'r' or column 'c' major order.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.

h2o4gpu.elastic_net_regressor

Elastic Net Regressor

Description

Elastic Net Regressor

Usage

```
h2o4gpu.elastic_net_regressor(alpha = 1, l1_ratio = 0.5,
    fit_intercept = TRUE, normalize = FALSE, precompute = FALSE,
    max_iter = 5000L, copy_X = TRUE, tol = 0.01, warm_start = FALSE,
    positive = FALSE, random_state = NULL, selection = "cyclic",
    n_gpus = -1L, lambda_stop_early = TRUE, glm_stop_early = TRUE,
    glm_stop_early_error_fraction = 1, verbose = FALSE, n_threads = NULL,
    gpu_id = 0L, lambda_min_ratio = 1e-07, n_lambdas = 100L, n_folds = 5L,
    tol_seek_factor = 0.1, store_full_path = 0L, lambda_max = NULL,
    lambdas = NULL, double_precision = NULL, order = NULL,
    backend = "h2o4gpu")
```

Arguments

alpha	Constant that multiplies the penalty terms. Defaults to 1.0. See the notes for the exact mathematical meaning of this parameter.alpha = 0 is equivalent to an ordinary least square, solved by the :class:LinearRegressionSklearn object. For numerical reasons, using alpha = 0 with the LassoSklearn object is not advised. Given this, you should use the :class:LinearRegressionSklearn object.
-------	---

l1_ratio	The ElasticNetSklearn mixing parameter, with $0 \leq \text{l1_ratio} \leq 1$. For $\text{l1_ratio} = 0$ the penalty is an L2 penalty. For $\text{l1_ratio} = 1$ it is an L1 penalty. For $0 < \text{l1_ratio} < 1$, the penalty is a combination of L1 and L2.
fit_intercept	Whether the intercept should be estimated or not. If FALSE, the data is assumed to be already centered.
normalize	This parameter is ignored when <code>fit_intercept</code> is set to FALSE. If TRUE, the regressors X will be normalized before regression by subtracting the mean and dividing by the l2-norm. If you wish to standardize, please use <code>:class:h2o4gpu.preprocessing.Standardizer</code> before calling <code>fit</code> on an estimator with <code>normalize=FALSE</code> .
precompute	Whether to use a precomputed Gram matrix to speed up calculations. The Gram matrix can also be passed as argument. For sparse input this option is always TRUE to preserve sparsity.
max_iter	The maximum number of iterations
copy_X	If TRUE, X will be copied; else, it may be overwritten.
tol	The tolerance for the optimization: if the updates are smaller than <code>tol</code> , the optimization code checks the dual gap for optimality and continues until it is smaller than <code>tol</code> .
warm_start	When set to TRUE, reuse the solution of the previous call to <code>fit</code> as initialization, otherwise, just erase the previous solution.
positive	When set to TRUE, forces the coefficients to be positive.
random_state	The seed of the pseudo random number generator that selects a random feature to update. If int, <code>random_state</code> is the seed used by the random number generator; If RandomState instance, <code>random_state</code> is the random number generator; If NULL, the random number generator is the RandomState instance used by <code>np.random</code> . Used when <code>selection == 'random'</code> .
selection	If set to 'random', a random coefficient is updated every iteration rather than looping over features sequentially by default. This (setting to 'random') often leads to significantly faster convergence especially when <code>tol</code> is higher than $1e-4$.
n_gpus	Number of gpu's to use in GLM solver.
lambda_stop_early	Stop early when there is no more relative improvement on train or validation.
glm_stop_early	Stop early when there is no more relative improvement in the primary and dual residuals for ADMM.
glm_stop_early_error_fraction	Relative tolerance for metric-based stopping criterion (stop if relative improvement is not at least this much).
verbose	Print verbose information to the console if set to > 0 .
n_threads	Number of threads to use in the gpu. Each thread is an independent model builder.
gpu_id	ID of the GPU on which the algorithm should run.
lambda_min_ratio	Minimum lambda ratio to maximum lambda, used in lambda search.
n_lambdas	Number of lambdas to be used in a search.

n_folds	Number of cross validation folds.
tol_seek_factor	Factor of tolerance to seek once below null model accuracy. Default is 1E-1, so seeks tolerance of 1E-3 once below null model accuracy for tol=1E-2.
store_full_path	Whether to store full solution for all alphas and lambdas. If 1, then during predict will compute best and full predictions.
lambda_max	Maximum Lambda value to use. Default is NULL, and then internally compute standard maximum
lambdas	overrides n_lambdas, lambda_max, and lambda_min_ratio.
double_precision	Internally set unless using _ptr methods. Value can either be 0 (float32) or 1(float64)
order	Order of data. Default is NULL, and internally determined (unless using _ptr methods) whether row 'r' or column 'c' major order.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.

h2o4gpu.gradient_boosting_classifier

Gradient Boosting Classifier

Description

Gradient Boosting Classifier

Usage

```
h2o4gpu.gradient_boosting_classifier(loss = "deviance", learning_rate = 0.1,
  n_estimators = 100L, subsample = 1, criterion = "friedman_mse",
  min_samples_split = 2L, min_samples_leaf = 1L,
  min_weight_fraction_leaf = 0, max_depth = 3L, min_impurity_decrease = 0,
  min_impurity_split = NULL, init = NULL, random_state = NULL,
  max_features = "auto", verbose = 0L, max_leaf_nodes = NULL,
  warm_start = FALSE, presort = "auto", colsample_bytree = 1,
  num_parallel_tree = 1L, tree_method = "gpu_hist", n_gpus = -1L,
  predictor = "gpu_predictor", objective = "binary:logistic",
  booster = "gbtree", n_jobs = 1L, gamma = 0L, min_child_weight = 1L,
  max_delta_step = 0L, colsample_bylevel = 1L, reg_alpha = 0L,
  reg_lambda = 1L, scale_pos_weight = 1L, base_score = 0.5,
  missing = NULL, backend = "h2o4gpu", ...)
```


Arguments

loss	loss function to be optimized. 'deviance' refers to deviance (= logistic regression) for classification with probabilistic outputs. For loss 'exponential' gradient boosting recovers the AdaBoost algorithm.
learning_rate	learning rate shrinks the contribution of each tree by learning_rate. There is a trade-off between learning_rate and n_estimators.
n_estimators	The number of boosting stages to perform. Gradient boosting is fairly robust to over-fitting so a large number usually results in better performance.
subsample	The fraction of samples to be used for fitting the individual base learners. If smaller than 1.0 this results in Stochastic Gradient Boosting. subsample interacts with the parameter n_estimators. Choosing subsample < 1.0 leads to a reduction of variance and an increase in bias.
criterion	The function to measure the quality of a split. Supported criteria are "friedman_mse" for the mean squared error with improvement score by Friedman, "mse" for mean squared error, and "mae" for the mean absolute error. The default value of "friedman_mse" is generally the best as it can provide a better approximation in some cases.
min_samples_split	The minimum number of samples required to split an internal node:
min_samples_leaf	The minimum number of samples required to be at a leaf node:
min_weight_fraction_leaf	The minimum weighted fraction of the sum total of weights (of all the input samples) required to be at a leaf node. Samples have equal weight when sample_weight is not provided.
max_depth	maximum depth of the individual regression estimators. The maximum depth limits the number of nodes in the tree. Tune this parameter for best performance; the best value depends on the interaction of the input variables.
min_impurity_decrease	A node will be split if this split induces a decrease of the impurity greater than or equal to this value.
min_impurity_split	Threshold for early stopping in tree growth. A node will split if its impurity is above the threshold, otherwise it is a leaf.
init	An estimator object that is used to compute the initial predictions. init has to provide fit and predict. If NULL it uses loss.init_estimator.
random_state	If int, random_state is the seed used by the random number generator; If RandomState instance, random_state is the random number generator; If NULL, the random number generator is the RandomState instance used by np.random.
max_features	The number of features to consider when looking for the best split:
verbose	Enable verbose output. If 1 then it prints progress and performance once in a while (the more trees the lower the frequency). If greater than 1 then it prints progress and performance for every tree.

max_leaf_nodes	Grow trees with max_leaf_nodes in best-first fashion. Best nodes are defined as relative reduction in impurity. If NULL then unlimited number of leaf nodes.
warm_start	When set to TRUE, reuse the solution of the previous call to fit and add more estimators to the ensemble, otherwise, just erase the previous solution.
presort	Whether to presort the data to speed up the finding of best splits in fitting. Auto mode by default will use presorting on dense data and default to normal sorting on sparse data. Setting presort to true on sparse data will raise an error.
colsample_bytree	Subsample ratio of columns when constructing each tree.
num_parallel_tree	Number of trees to grow per round
tree_method	The tree construction algorithm used in XGBoost Distributed and external memory version only support approximate algorithm. Choices: 'auto', 'exact', 'approx', 'hist', 'gpu_exact', 'gpu_hist' 'auto': Use heuristic to choose faster one. - For small to medium dataset, exact greedy will be used. - For very large-dataset, approximate algorithm will be chosen. - Because old behavior is always use exact greedy in single machine, - user will get a message when approximate algorithm is chosen to notify this choice. 'exact': Exact greedy algorithm. 'approx': Approximate greedy algorithm using sketching and histogram. 'hist': Fast histogram optimized approximate greedy algorithm. It uses some performance improvements such as bins caching. 'gpu_exact': GPU implementation of exact algorithm. 'gpu_hist': GPU implementation of hist algorithm.
n_gpus	Number of gpu's to use in GradientBoostingClassifier solver. Default is -1.
predictor	The type of predictor algorithm to use. Provides the same results but allows the use of GPU or CPU. - 'cpu_predictor': Multicore CPU prediction algorithm. - 'gpu_predictor': Prediction using GPU. Default for 'gpu_exact' and 'gpu_hist' tree method.
objective	Specify the learning task and the corresponding learning objective or a custom objective function to be used Note: A custom objective function can be provided for the objective parameter. In this case, it should have the signature objective(y_true, y_pred) -> grad, hess:
booster	Specify which booster to use: gbtree, gblinear or dart.
n_jobs	Number of parallel threads used to run xgboost.
gamma	Minimum loss reduction required to make a further partition on a leaf node of the tree.
min_child_weight	Minimum sum of instance weight(hessian) needed in a child.
max_delta_step	Maximum delta step we allow each tree's weight estimation to be.
colsample_bylevel	Subsample ratio of columns for each split, in each level.
reg_alpha	L1 regularization term on weights
reg_lambda	L2 regularization term on weights
scale_pos_weight	Balancing of positive and negative weights.

base_score	The initial prediction score of all instances, global bias.
missing	Value in the data which needs to be present as a missing value. If NULL, defaults to np.nan.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.
...	Other parameters for XGBoost object. Full documentation of parameters can be found here: https://github.com/dmlc/xgboost/blob/master/doc/parameter.md

h2o4gpu.gradient_boosting_regressor

Gradient Boosting Regressor

Description

Gradient Boosting Regressor

Usage

```
h2o4gpu.gradient_boosting_regressor(loss = "ls", learning_rate = 0.1,
    n_estimators = 100L, subsample = 1, criterion = "friedman_mse",
    min_samples_split = 2L, min_samples_leaf = 1L,
    min_weight_fraction_leaf = 0, max_depth = 3L, min_impurity_decrease = 0,
    min_impurity_split = NULL, init = NULL, random_state = NULL,
    max_features = "auto", alpha = 0.9, verbose = 0L,
    max_leaf_nodes = NULL, warm_start = FALSE, presort = "auto",
    colsample_bytree = 1, num_parallel_tree = 1L, tree_method = "gpu_hist",
    n_gpus = -1L, predictor = "gpu_predictor", objective = "reg:linear",
    booster = "gbtree", n_jobs = 1L, gamma = 0L, min_child_weight = 1L,
    max_delta_step = 0L, colsample_bylevel = 1L, reg_alpha = 0L,
    reg_lambda = 1L, scale_pos_weight = 1L, base_score = 0.5,
    missing = NULL, backend = "h2o4gpu", ...)
```

Arguments

loss	loss function to be optimized. 'ls' refers to least squares regression. 'lad' (least absolute deviation) is a highly robust loss function solely based on order information of the input variables. 'huber' is a combination of the two. 'quantile' allows quantile regression (use alpha to specify the quantile).
learning_rate	learning rate shrinks the contribution of each tree by learning_rate. There is a trade-off between learning_rate and n_estimators.
n_estimators	The number of boosting stages to perform. Gradient boosting is fairly robust to over-fitting so a large number usually results in better performance.
subsample	The fraction of samples to be used for fitting the individual base learners. If smaller than 1.0 this results in Stochastic Gradient Boosting. subsample interacts with the parameter n_estimators. Choosing subsample < 1.0 leads to a reduction of variance and an increase in bias.

<code>criterion</code>	The function to measure the quality of a split. Supported criteria are "friedman_mse" for the mean squared error with improvement score by Friedman, "mse" for mean squared error, and "mae" for the mean absolute error. The default value of "friedman_mse" is generally the best as it can provide a better approximation in some cases.
<code>min_samples_split</code>	The minimum number of samples required to split an internal node:
<code>min_samples_leaf</code>	The minimum number of samples required to be at a leaf node:
<code>min_weight_fraction_leaf</code>	The minimum weighted fraction of the sum total of weights (of all the input samples) required to be at a leaf node. Samples have equal weight when <code>sample_weight</code> is not provided.
<code>max_depth</code>	maximum depth of the individual regression estimators. The maximum depth limits the number of nodes in the tree. Tune this parameter for best performance; the best value depends on the interaction of the input variables.
<code>min_impurity_decrease</code>	A node will be split if this split induces a decrease of the impurity greater than or equal to this value.
<code>min_impurity_split</code>	Threshold for early stopping in tree growth. A node will split if its impurity is above the threshold, otherwise it is a leaf.
<code>init</code>	An estimator object that is used to compute the initial predictions. <code>init</code> has to provide <code>fit</code> and <code>predict</code> . If NULL it uses <code>loss.init_estimator</code> .
<code>random_state</code>	If int, <code>random_state</code> is the seed used by the random number generator; If <code>RandomState</code> instance, <code>random_state</code> is the random number generator; If NULL, the random number generator is the <code>RandomState</code> instance used by <code>np.random</code> .
<code>max_features</code>	The number of features to consider when looking for the best split:
<code>alpha</code>	The alpha-quantile of the huber loss function and the quantile loss function. Only if <code>loss='huber'</code> or <code>loss='quantile'</code> .
<code>verbose</code>	Enable verbose output. If 1 then it prints progress and performance once in a while (the more trees the lower the frequency). If greater than 1 then it prints progress and performance for every tree.
<code>max_leaf_nodes</code>	Grow trees with <code>max_leaf_nodes</code> in best-first fashion. Best nodes are defined as relative reduction in impurity. If NULL then unlimited number of leaf nodes.
<code>warm_start</code>	When set to TRUE, reuse the solution of the previous call to fit and add more estimators to the ensemble, otherwise, just erase the previous solution.
<code>presort</code>	Whether to presort the data to speed up the finding of best splits in fitting. Auto mode by default will use presorting on dense data and default to normal sorting on sparse data. Setting <code>presort</code> to true on sparse data will raise an error.
<code>colsample_bytree</code>	Subsample ratio of columns when constructing each tree.
<code>num_parallel_tree</code>	Number of trees to grow per round

tree_method	The tree construction algorithm used in XGBoost Distributed and external memory version only support approximate algorithm. Choices: 'auto', 'exact', 'approx', 'hist', 'gpu_exact', 'gpu_hist' 'auto': Use heuristic to choose faster one. - For small to medium dataset, exact greedy will be used. - For very large-dataset, approximate algorithm will be chosen. - Because old behavior is always use exact greedy in single machine, - user will get a message when approximate algorithm is chosen to notify this choice. 'exact': Exact greedy algorithm. 'approx': Approximate greedy algorithm using sketching and histogram. 'hist': Fast histogram optimized approximate greedy algorithm. It uses some performance improvements such as bins caching. 'gpu_exact': GPU implementation of exact algorithm. 'gpu_hist': GPU implementation of hist algorithm.
n_gpus	Number of gpu's to use in GradientBoostingRegressor solver. Default is -1.
predictor	The type of predictor algorithm to use. Provides the same results but allows the use of GPU or CPU. - 'cpu_predictor': Multicore CPU prediction algorithm. - 'gpu_predictor': Prediction using GPU. Default for 'gpu_exact' and 'gpu_hist' tree method.
objective	Specify the learning task and the corresponding learning objective or a custom objective function to be used Note: A custom objective function can be provided for the objective parameter. In this case, it should have the signature objective(y_true, y_pred) -> grad, hess:
booster	Specify which booster to use: gbtree, gblinear or dart.
n_jobs	Number of parallel threads used to run xgboost.
gamma	Minimum loss reduction required to make a further partition on a leaf node of the tree.
min_child_weight	Minimum sum of instance weight(hessian) needed in a child.
max_delta_step	Maximum delta step we allow each tree's weight estimation to be.
colsample_bylevel	Subsample ratio of columns for each split, in each level.
reg_alpha	L1 regularization term on weights
reg_lambda	L2 regularization term on weights
scale_pos_weight	Balancing of positive and negative weights.
base_score	The initial prediction score of all instances, global bias.
missing	Value in the data which needs to be present as a missing value. If NULL, defaults to np.nan.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.
...	Other parameters for XGBoost object. Full documentation of parameters can be found here: https://github.com/dmlc/xgboost/blob/master/doc/parameter.md

h2o4gpu.kmeans

K-means Clustering

Description

K-means Clustering

Usage

```
h2o4gpu.kmeans(n_clusters = 8L, init = "k-means++", n_init = 1L,
               max_iter = 300L, tol = 1e-04, precompute_distances = "auto",
               verbose = 0L, random_state = NULL, copy_x = TRUE, n_jobs = 1L,
               algorithm = "auto", gpu_id = 0L, n_gpus = -1L, do_checks = 1L,
               backend = "h2o4gpu")
```

Arguments

n_clusters	The number of clusters to form as well as the number of centroids to generate.
init	Method for initialization, defaults to 'random': 'k-means++': selects initial cluster centers for k-mean clustering in a smart way to speed up convergence. <i>Not supported yet</i> - if chosen we will use SKLearn's methods. 'random': choose k observations (rows) at random from data for the initial centroids. If an ndarray is passed, it should be of shape (n_clusters, n_features) and gives the initial centers. <i>Not supported yet</i> - if chosen we will use SKLearn's methods.
n_init	Number of time the k-means algorithm will be run with different centroid seeds. The final results will be the best output of n_init consecutive runs in terms of inertia. <i>Not supported yet</i> - always runs 1.
max_iter	Maximum number of iterations of the algorithm.
tol	Relative tolerance to declare convergence.
precompute_distances	Precompute distances (faster but takes more memory). 'auto' : do not precompute distances if n_samples * n_clusters > 12 million. This corresponds to about 100MB overhead per job using double precision. TRUE : always precompute distances FALSE : never precompute distances <i>Not supported yet</i> - always uses auto if running h2o4gpu version.
verbose	Logger verbosity level.
random_state	random_state for RandomState. Must be convertible to 32 bit unsigned integers.
copy_x	When pre-computing distances it is more numerically accurate to center the data first. If copy_x is TRUE, then the original data is not modified. If FALSE, the original data is modified, and put back before the function returns, but small numerical differences may be introduced by subtracting and then adding the data mean. <i>Not supported yet</i> - always uses TRUE if running h2o4gpu version.

n_jobs	The number of jobs to use for the computation. This works by computing each of the n_init runs in parallel. If -1 all CPUs are used. If 1 is given, no parallel computing code is used at all, which is useful for debugging. For n_jobs below -1, (n_cpus + 1 + n_jobs) are used. Thus for n_jobs = -2, all CPUs but one are used. <i>Not supported yet</i> - CPU backend not yet implemented.
algorithm	K-means algorithm to use. The classical EM-style algorithm is "full". The "elkan" variation is more efficient by using the triangle inequality, but currently doesn't support sparse data. "auto" chooses "elkan" for dense data and "full" for sparse data. <i>Not supported yet</i> - always uses full if running h2o4gpu version.
gpu_id	ID of the GPU on which the algorithm should run.
n_gpus	Number of GPUs on which the algorithm should run. < 0 means all possible GPUs on the machine. 0 means no GPUs, run on CPU.
do_checks	If set to 0 GPU error check will not be performed.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.

h2o4gpu.pca

*Principal Component Analysis (PCA)***Description**

Principal Component Analysis (PCA)

Usage

```
h2o4gpu.pca(n_components = 2L, copy = TRUE, whiten = FALSE,
            svd_solver = "arpack", tol = 0, iterated_power = "auto",
            random_state = NULL, verbose = FALSE, backend = "h2o4gpu",
            gpu_id = 0L)
```

Arguments

n_components	Desired dimensionality of output data
copy	If FALSE, data passed to fit are overwritten and running fit(X).transform(X) will not yield the expected results, use fit_transform(X) instead.
whiten	When TRUE (FALSE by default) the components_ vectors are multiplied by the square root of (n_samples) and divided by the singular values to ensure uncorrelated outputs with unit component-wise variances.
svd_solver	'auto' is selected by a default policy based on X.shape and n_components: if the input data is larger than 500x500 and the number of components to extract is lower than 80 percent of the smallest dimension of the data, then the more efficient 'randomized' method is enabled. Otherwise the exact full SVD is computed and optionally truncated afterwards. 'full' runs exact full SVD calling the standard LAPACK solver via scipy.linalg.svd and select the components by postprocessing 'arpack' runs SVD truncated to n_components calling ARPACK solver via scipy.sparse.linalg.svds. It requires strictly 0 < n_components < columns. 'randomized' runs randomized SVD by the method of Halko et al.

tol	Tolerance for singular values computed by svd_solver == 'arpack'.
iterated_power	Number of iterations for the power method computed by svd_solver == 'randomized'.
random_state	If int, random_state is the seed used by the random number generator; If RandomState instance, random_state is the random number generator; If NULL, the random number generator is the RandomState instance used by np.random. Used when svd_solver == 'arpack' or 'randomized'.
verbose	Verbose or not
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.
gpu_id	ID of the GPU on which the algorithm should run. Only used by h2o4gpu backend.

h2o4gpu.random_forest_classifier

Random Forest Classifier

Description

Random Forest Classifier

Usage

```
h2o4gpu.random_forest_classifier(n_estimators = 100L, criterion = "gini",
    max_depth = 3L, min_samples_split = 2L, min_samples_leaf = 1L,
    min_weight_fraction_leaf = 0, max_features = "auto",
    max_leaf_nodes = NULL, min_impurity_decrease = 0,
    min_impurity_split = NULL, bootstrap = TRUE, oob_score = FALSE,
    n_jobs = 1L, random_state = NULL, verbose = 0L, warm_start = FALSE,
    class_weight = NULL, subsample = 1, colsample_bytree = 1,
    num_parallel_tree = 1L, tree_method = "gpu_hist", n_gpus = -1L,
    predictor = "gpu_predictor", backend = "h2o4gpu")
```

Arguments

n_estimators	The number of trees in the forest.
criterion	The function to measure the quality of a split. Supported criteria are "gini" for the Gini impurity and "entropy" for the information gain. Note: this parameter is tree-specific.
max_depth	The maximum depth of the tree. If NULL, then nodes are expanded until all leaves are pure or until all leaves contain less than min_samples_split samples.
min_samples_split	The minimum number of samples required to split an internal node:
min_samples_leaf	The minimum number of samples required to be at a leaf node:

<code>min_weight_fraction_leaf</code>	The minimum weighted fraction of the sum total of weights (of all the input samples) required to be at a leaf node. Samples have equal weight when <code>sample_weight</code> is not provided.
<code>max_features</code>	The number of features to consider when looking for the best split:
<code>max_leaf_nodes</code>	Grow trees with <code>max_leaf_nodes</code> in best-first fashion. Best nodes are defined as relative reduction in impurity. If NULL then unlimited number of leaf nodes.
<code>min_impurity_decrease</code>	A node will be split if this split induces a decrease of the impurity greater than or equal to this value.
<code>min_impurity_split</code>	Threshold for early stopping in tree growth. A node will split if its impurity is above the threshold, otherwise it is a leaf.
<code>bootstrap</code>	Whether bootstrap samples are used when building trees.
<code>oob_score</code>	whether to use out-of-bag samples to estimate the R^2 on unseen data.
<code>n_jobs</code>	The number of jobs to run in parallel for both <code>fit</code> and <code>predict</code> . If -1, then the number of jobs is set to the number of cores.
<code>random_state</code>	If int, <code>random_state</code> is the seed used by the random number generator; If <code>RandomState</code> instance, <code>random_state</code> is the random number generator; If NULL, the random number generator is the <code>RandomState</code> instance used by <code>np.random</code> .
<code>verbose</code>	Controls the verbosity of the tree building process.
<code>warm_start</code>	When set to <code>TRUE</code> , reuse the solution of the previous call to <code>fit</code> and add more estimators to the ensemble, otherwise, just fit a whole new forest.
<code>class_weight</code>	"balanced_subsample" or NULL, optional (default=NULL) Weights associated with classes in the form <code>{class_label: weight}</code> . If not given, all classes are supposed to have weight one. For multi-output problems, a list of dicts can be provided in the same order as the columns of <code>y</code> .
<code>subsample</code>	Subsample ratio of the training instance.
<code>colsample_bytree</code>	Subsample ratio of columns when constructing each tree.
<code>num_parallel_tree</code>	Number of trees to grow per round
<code>tree_method</code>	The tree construction algorithm used in XGBoost Distributed and external memory version only support approximate algorithm. Choices: 'auto', 'exact', 'approx', 'hist', 'gpu_exact', 'gpu_hist' 'auto': Use heuristic to choose faster one. - For small to medium dataset, exact greedy will be used. - For very large-dataset, approximate algorithm will be chosen. - Because old behavior is always use exact greedy in single machine, - user will get a message when approximate algorithm is chosen to notify this choice. 'exact': Exact greedy algorithm. 'approx': Approximate greedy algorithm using sketching and histogram. 'hist': Fast histogram optimized approximate greedy algorithm. It uses some performance improvements such as bins caching. 'gpu_exact': GPU implementation of exact algorithm. 'gpu_hist': GPU implementation of hist algorithm.
<code>n_gpus</code>	Number of gpu's to use in RandomForestClassifier solver. Default is -1.

predictor	The type of predictor algorithm to use. Provides the same results but allows the use of GPU or CPU. - 'cpu_predictor': Multicore CPU prediction algorithm. - 'gpu_predictor': Prediction using GPU. Default for 'gpu_exact' and 'gpu_hist' tree method.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.

h2o4gpu.random_forest_regressor

Random Forest Regressor

Description

Random Forest Regressor

Usage

```
h2o4gpu.random_forest_regressor(n_estimators = 100L, criterion = "mse",
  max_depth = 3L, min_samples_split = 2L, min_samples_leaf = 1L,
  min_weight_fraction_leaf = 0, max_features = "auto",
  max_leaf_nodes = NULL, min_impurity_decrease = 0,
  min_impurity_split = NULL, bootstrap = TRUE, oob_score = FALSE,
  n_jobs = 1L, random_state = NULL, verbose = 0L, warm_start = FALSE,
  subsample = 1, colsample_bytree = 1, num_parallel_tree = 1L,
  tree_method = "gpu_hist", n_gpus = -1L, predictor = "gpu_predictor",
  backend = "h2o4gpu")
```

Arguments

n_estimators	The number of trees in the forest.
criterion	The function to measure the quality of a split. Supported criteria are "mse" for the mean squared error, which is equal to variance reduction as feature selection criterion, and "mae" for the mean absolute error.
max_depth	The maximum depth of the tree. If NULL, then nodes are expanded until all leaves are pure or until all leaves contain less than min_samples_split samples.
min_samples_split	The minimum number of samples required to split an internal node:
min_samples_leaf	The minimum number of samples required to be at a leaf node:
min_weight_fraction_leaf	The minimum weighted fraction of the sum total of weights (of all the input samples) required to be at a leaf node. Samples have equal weight when sample_weight is not provided.
max_features	The number of features to consider when looking for the best split:

max_leaf_nodes	Grow trees with max_leaf_nodes in best-first fashion. Best nodes are defined as relative reduction in impurity. If NULL then unlimited number of leaf nodes.
min_impurity_decrease	A node will be split if this split induces a decrease of the impurity greater than or equal to this value.
min_impurity_split	Threshold for early stopping in tree growth. A node will split if its impurity is above the threshold, otherwise it is a leaf.
bootstrap	Whether bootstrap samples are used when building trees.
oob_score	whether to use out-of-bag samples to estimate the R ² on unseen data.
n_jobs	The number of jobs to run in parallel for both fit and predict. If -1, then the number of jobs is set to the number of cores.
random_state	If int, random_state is the seed used by the random number generator; If RandomState instance, random_state is the random number generator; If NULL, the random number generator is the RandomState instance used by np.random.
verbose	Controls the verbosity of the tree building process.
warm_start	When set to TRUE, reuse the solution of the previous call to fit and add more estimators to the ensemble, otherwise, just fit a whole new forest.
subsample	Subsample ratio of the training instance.
colsample_bytree	Subsample ratio of columns when constructing each tree.
num_parallel_tree	Number of trees to grow per round
tree_method	The tree construction algorithm used in XGBoost Distributed and external memory version only support approximate algorithm. Choices: 'auto', 'exact', 'approx', 'hist', 'gpu_exact', 'gpu_hist' 'auto': Use heuristic to choose faster one. - For small to medium dataset, exact greedy will be used. - For very large-dataset, approximate algorithm will be chosen. - Because old behavior is always use exact greedy in single machine, - user will get a message when approximate algorithm is chosen to notify this choice. 'exact': Exact greedy algorithm. 'approx': Approximate greedy algorithm using sketching and histogram. 'hist': Fast histogram optimized approximate greedy algorithm. It uses some performance improvements such as bins caching. 'gpu_exact': GPU implementation of exact algorithm. 'gpu_hist': GPU implementation of hist algorithm.
n_gpus	Number of gpu's to use in RandomForestRegressor solver. Default is -1.
predictor	The type of predictor algorithm to use. Provides the same results but allows the use of GPU or CPU. - 'cpu_predictor': Multicore CPU prediction algorithm. - 'gpu_predictor': Prediction using GPU. Default for 'gpu_exact' and 'gpu_hist' tree method.
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.

h2o4gpu.truncated_svd *Truncated Singular Value Decomposition (TruncatedSVD)*

Description

Truncated Singular Value Decomposition (TruncatedSVD)

Usage

```
h2o4gpu.truncated_svd(n_components = 2L, algorithm = "power",
                      n_iter = 100L, random_state = NULL, tol = 1e-05, verbose = FALSE,
                      backend = "h2o4gpu", n_gpus = 1L, gpu_id = 0L)
```

Arguments

n_components	Desired dimensionality of output data
algorithm	SVD solver to use. H2O4GPU options: Either "cusolver" (similar to ARPACK) or "power" for the power method. SKlearn options: Either "arpack" for the ARPACK wrapper in SciPy (scipy.sparse.linalg.svds), or "randomized" for the randomized algorithm due to Halko (2009).
n_iter	number of iterations (only relevant for power method) Should be at most 2147483647 due to INT_MAX in C++ backend.
random_state	seed (NULL for auto-generated)
tol	Tolerance for "power" method. Ignored by "cusolver". Should be > 0.0 to ensure convergence. Should be 0.0 to effectively ignore and only base convergence upon n_iter
verbose	Verbose or not
backend	Which backend to use. Options are 'auto', 'sklearn', 'h2o4gpu'. Saves as attribute for actual backend used.
n_gpus	How many gpus to use. If 0, use CPU backup method. Currently SVD only uses 1 GPU, so >1 has no effect compared to 1.
gpu_id	ID of the GPU on which the algorithm should run.

predict.h2o4gpu_model *Make Predictions using Trained H2O4GPU Estimator*

Description

This function makes predictions from new data using a trained H2O4GPU model and returns class predictions for classification and predicted values for regression.

Usage

```
## S3 method for class 'h2o4gpu_model'
predict(object, x, type = "raw", ...)
```

Arguments

object	The h2o4gpu model object
x	The new data where each column represents a different predictor variable to be used in generating predictions.
type	One of "raw" or "prob", indicating the type of output: predicted values or probabilities
...	Additional arguments (unused for now).

Examples

```
## Not run:

library(h2o4gpu)

# Setup dataset
x <- iris[1:4]
y <- as.integer(iris$Species) - 1

# Initialize and train the classifier
model <- h2o4gpu.random_forest_classifier() %>% fit(x, y)

# Make predictions
predictions <- model %>% predict(x)

## End(Not run)
```

```
transform.h2o4gpu_model
```

Transform a Dataset using Trained H2O4GPU Estimator

Description

This function transforms the given new data using a trained H2O4GPU model.

Usage

```
## S3 method for class 'h2o4gpu_model'
transform(object, x, ...)
```

Arguments

object	The h2o4gpu model object
x	The new data where each column represents a different predictor variable to be used in generating predictions.
...	Additional arguments (unused for now).

Examples

```
## Not run:

library(h2o4gpu)

# Prepare data
iris$Species <- as.integer(iris$Species) # convert to numeric data

# Randomly sample 80% of the rows for the training set
set.seed(1)
train_idx <- sample(1:nrow(iris), 0.8*nrow(iris))
train <- iris[train_idx, ]
test <- iris[-train_idx, ]

# Train a K-Means model
model_km <- h2o4gpu.kmeans(n_clusters = 3L) %>% fit(train)

# Transform test data
test_dist <- model_km %>% transform(test)

## End(Not run)
```

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