# Package ‘sparkxgb’

March 13, 2019

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<th>Type</th>
<th>Package</th>
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<tbody>
<tr>
<td>Title</td>
<td>Interface for 'XGBoost' on 'Apache Spark'</td>
</tr>
<tr>
<td>Version</td>
<td>0.1.0</td>
</tr>
<tr>
<td>Maintainer</td>
<td>Kevin Kuo <a href="mailto:kevin.kuo@rstudio.com">kevin.kuo@rstudio.com</a></td>
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<tr>
<td>Description</td>
<td>A 'sparklyr' <a href="https://spark.rstudio.com/">https://spark.rstudio.com/</a> extension that provides an interface for 'XGBoost' <a href="https://github.com/dmlc/xgboost">https://github.com/dmlc/xgboost</a> on 'Apache Spark'. 'XGBoost' is an optimized distributed gradient boosting library.</td>
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<td>SystemRequirements</td>
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<td>R (&gt;= 3.1.2)</td>
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<td>Imports</td>
<td>sparklyr (&gt;= 1.0), forge (&gt;= 0.2.0)</td>
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<tr>
<td>Suggests</td>
<td>testthat, dplyr, purrr, rlang</td>
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<tr>
<td>NeedsCompilation</td>
<td>no</td>
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<tr>
<td>Author</td>
<td>Kevin Kuo [aut, cre] (<a href="https://orcid.org/0000-0001-7803-7901">https://orcid.org/0000-0001-7803-7901</a>)</td>
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<td>Repository</td>
<td>CRAN</td>
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XGBoost Classifier

**Description**

XGBoost classifier for Spark.

**Usage**

```r
xgboost_classifier(x, formula = NULL, eta = 0.3, gamma = 0,
max_depth = 6, min_child_weight = 1, max_delta_step = 0,
grow_policy = "depthwise", max_bins = 16, subsample = 1,
colsample_bytree = 1, colsample_bylevel = 1, lambda = 1,
alpha = 0, tree_method = "auto", sketch_eps = 0.03,
scale_pos_weight = 1, sample_type = "uniform",
normalize_type = "tree", rate_drop = 0, skip_drop = 0,
lambda_bias = 0, tree_limit = 0, num_round = 1, num_workers = 1,
nthread = 1, use_external_memory = FALSE, silent = 0,
custom_obj = NULL, custom_eval = NULL, missing = NaN, seed = 0,
timeout_request_workers = 30 * 60 * 1000, checkpoint_path = "",
checkpoint_interval = -1, objective = "multi:softprob",
base_score = 0.5, train_test_ratio = 1,
num_early_stopping_rounds = 0, objective_type = "classification",
eval_metric = NULL, maximize_evaluation_metrics = FALSE,
num_class = NULL, base_margin_col = NULL, thresholds = NULL,
weight_col = NULL, features_col = "features", label_col = "label",
prediction_col = "prediction", probability_col = "probability",
raw_prediction_col = "rawPrediction",
uid = random_string("xgboost_classifier_"), ...)
```

**Arguments**

- **x**  
  A spark_connection, ml_pipeline, or a tbl_spark.

- **formula**  
  Used when x is a tbl_spark. R formula as a character string or a formula. 
  This is used to transform the input dataframe before fitting, see `ft_r_formula` for details.

- **eta**  
  Step size shrinkage used in update to prevents overfitting. After each boosting step, we can directly get the weights of new features and eta actually shrinks the feature weights to make the boosting process more conservative. [default=0.3] 
  range: [0,1]

- **gamma**  
  Minimum loss reduction required to make a further partition on a leaf node of the tree. the larger, the more conservative the algorithm will be. [default=0]

- **max_depth**  
  Maximum depth of a tree, increase this value will make model more complex / likely to be overfitting. [default=6]
**min_child_weight**
Minimum sum of instance weight(hessian) needed in a child. If the tree partition step results in a leaf node with the sum of instance weight less than min_child_weight, then the building process will give up further partitioning. In linear regression mode, this simply corresponds to minimum number of instances needed to be in each node. The larger, the more conservative the algorithm will be. [default=1]

**max_delta_step**
Maximum delta step we allow each tree’s weight estimation to be. If the value is set to 0, it means there is no constraint. If it is set to a positive value, it can help making the update step more conservative. Usually this parameter is not needed, but it might help in logistic regression when class is extremely imbalanced. Set it to value of 1-10 might help control the update. [default=0]

**grow_policy**
Growth policy for fast histogram algorithm.

**max_bins**
Maximum number of bins in histogram.

**subsample**
Subsample ratio of the training instance. Setting it to 0.5 means that XGBoost randomly collected half of the data instances to grow trees and this will prevent overfitting. [default=1] range:(0,1]

**colsample_bytree**
Subsample ratio of columns when constructing each tree. [default=1] range: (0,1]

**colsample_bylevel**
Subsample ratio of columns for each split, in each level. [default=1] range: (0,1]

**lambda**
L2 regularization term on weights, increase this value will make model more conservative. [default=1]

**alpha**
L1 regularization term on weights, increase this value will make model more conservative, defaults to 0.

**tree_method**
The tree construction algorithm used in XGBoost. options: 'auto', 'exact', 'approx' [default='auto']

**sketch_eps**
This is only used for approximate greedy algorithm. This roughly translated into O(1 / sketch_eps) number of bins. Compared to directly select number of bins, this comes with theoretical guarantee with sketch accuracy. [default=0.03] range: (0, 1)

**scale_pos_weight**
Control the balance of positive and negative weights, useful for unbalanced classes. A typical value to consider: sum(negative cases) / sum(positive cases). [default=1]

**sample_type**
Parameter for Dart booster. Type of sampling algorithm. "uniform": dropped trees are selected uniformly. "weighted": dropped trees are selected in proportion to weight. [default=“uniform”]

**normalize_type**
Parameter of Dart booster. type of normalization algorithm, options: 'tree', 'forest'. [default=“tree”]

**rate_drop**
Parameter of Dart booster. dropout rate. [default=0.0] range: [0.0, 1.0]

**skip_drop**
Parameter of Dart booster. probability of skip dropout. If a dropout is skipped, new trees are added in the same manner as gbtree. [default=0.0] range: [0.0, 1.0]
lambda_bias  Parameter of linear booster L2 regularization term on bias, default 0 (no L1 reg on bias because it is not important.)
tree_limit  Limit number of trees in the prediction; defaults to 0 (use all trees.)
num_round  The number of rounds for boosting.
num_workers  number of workers used to train xgboost model. Defaults to 1.
nthread  Number of threads used by per worker. Defaults to 1.
use_external_memory  The tree construction algorithm used in XGBoost. options: 'auto', 'exact', 'approx' [default='auto']
silent  0 means printing running messages, 1 means silent mode. default: 0
custom_obj  Customized objective function provided by user. Currently unsupported.
custom_eval  Customized evaluation function provided by user. Currently unsupported.
missing  The value treated as missing. default: Float.NaN
seed  Random seed for the C++ part of XGBoost and train/test splitting.
timeout_request_workers  the maximum time to wait for the job requesting new workers. default: 30 minutes
checkpoint_path  The hdfs folder to load and save checkpoint boosters.
checkpoint_interval  Param for set checkpoint interval (>= 1) or disable checkpoint (-1). E.g. 10 means that the trained model will get checkpointed every 10 iterations. Note: checkpoint_path must also be set if the checkpoint interval is greater than 0.
base_score  Param for initial prediction (aka base margin) column name. Defaults to 0.5.
train_test_ratio  Fraction of training points to use for testing.
num_early_stopping_rounds  If non-zero, the training will be stopped after a specified number of consecutive increases in any evaluation metric.
optimize_type  The learning objective type of the specified custom objective and eval. Corresponding type will be assigned if custom objective is defined options: regression, classification.
eval_metric  Evaluation metrics for validation data, a default metric will be assigned according to objective(rmse for regression, and error for classification, mean average precision for ranking). options: rmse, mae, logloss, error, merror, mlogloss, auc, aucpr, ndcg, map, gamma-deviance
maximize_evaluation_metrics  Whether to maximize evaluation metrics. Defaults to FALSE (for minization.)
num_class  Number of classes.
base_margin_col
Param for initial prediction (aka base margin) column name.

thresholds
Thresholds in multi-class classification to adjust the probability of predicting each class. Array must have length equal to the number of classes, with values > 0 excepting that at most one value may be 0. The class with largest value p/t is predicted, where p is the original probability of that class and t is the class’s threshold.

weight_col
Weight column.

features_col
Features column name, as a length-one character vector. The column should be single vector column of numeric values. Usually this column is output by `ft_r_formula`.

label_col
Label column name. The column should be a numeric column. Usually this column is output by `ft_r_formula`.

prediction_col
Prediction column name.

probability_col
Column name for predicted class conditional probabilities.

raw_prediction_col
Raw prediction (a.k.a. confidence) column name.

uid
A character string used to uniquely identify the ML estimator.

Examples

```r
library(sparkxgb)
library(sparklyr)

sc <- spark_connect(master = "local")
iris_tbl <- sdf_copy_to(sc, iris)
xgb_model <- xgboost_classifier(
  iris_tbl,
  Species ~ .,
  num_class = 3,
  num_round = 50,
  max_depth = 4
)
```

**xgboost_regressor**

XGBoost Regressor

**Description**

XGBoost regressor for Spark.
Usage

xgboost_regressor(x, formula = NULL, eta = 0.3, gamma = 0,
max_depth = 6, min_child_weight = 1, max_delta_step = 0,
grow_policy = "depthwise", max_bins = 16, subsample = 1,
colsample_bytree = 1, colsample_bylevel = 1, lambda = 1,
alpha = 0, tree_method = "auto", sketch_eps = 0.03,
scale_pos_weight = 1, sample_type = "uniform",
normalize_type = "tree", rate_drop = 0, skip_drop = 0,
lambda_bias = 0, tree_limit = 0, num_round = 1, num_workers = 1,
nthread = 1, use_external_memory = FALSE, silent = 0,
custom_obj = NULL, custom_eval = NULL, missing = NaN, seed = 0,
timeout_request_workers = 30 * 60 * 1000, checkpoint_path = "",
checkpoint_interval = -1, objective = "reg:linear",
base_score = 0.5, train_test_ratio = 1,
num_early_stopping_rounds = 0, objective_type = "regression",
eval_metric = NULL, maximize_evaluation_metrics = FALSE,
base_margin_col = NULL, weight_col = NULL,
features_col = "features", label_col = "label",
prediction_col = "prediction",
uid = random_string("xgboost_regressor_"), ...

Arguments

x  
A spark_connection, ml_pipeline, or a tbl_spark.

formula  
Used when x is a tbl_spark. R formula as a character string or a formula. This is used to transform the input dataframe before fitting, see ft_r_formula for details.

eta  
Step size shrinkage used in update to prevents overfitting. After each boosting step, we can directly get the weights of new features and eta actually shrinks the feature weights to make the boosting process more conservative. [default=0.3] range: [0,1]

gamma  
Minimum loss reduction required to make a further partition on a leaf node of the tree. the larger, the more conservative the algorithm will be. [default=0]

max_depth  
Maximum depth of a tree, increase this value will make model more complex / likely to be overfitting. [default=6]

min_child_weight  
Minimum sum of instance weight(hessian) needed in a child. If the tree partition step results in a leaf node with the sum of instance weight less than min_child_weight, then the building process will give up further partitioning. In linear regression mode, this simply corresponds to minimum number of instances needed to be in each node. The larger, the more conservative the algorithm will be. [default=1]

max_delta_step  
Maximum delta step we allow each tree’s weight estimation to be. If the value is set to 0, it means there is no constraint. If it is set to a positive value, it can help making the update step more conservative. Usually this parameter is not needed, but it might help in logistic regression when class is extremely imbalanced. Set it to value of 1-10 might help control the update. [default=0]
<table>
<thead>
<tr>
<th>Parameter</th>
<th>Description</th>
</tr>
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<tbody>
<tr>
<td><code>grow_policy</code></td>
<td>Growth policy for fast histogram algorithm.</td>
</tr>
<tr>
<td><code>max_bins</code></td>
<td>Maximum number of bins in histogram.</td>
</tr>
<tr>
<td><code>subsample</code></td>
<td>Subsample ratio of the training instance. Setting it to 0.5 means that XGBoost randomly collected half of the data instances to grow trees and this will prevent overfitting. [default=1] range:(0,1]</td>
</tr>
<tr>
<td><code>colsample_bytree</code></td>
<td>Subsample ratio of columns when constructing each tree. [default=1] range: (0,1]</td>
</tr>
<tr>
<td><code>colsample_bylevel</code></td>
<td>Subsample ratio of columns for each split, in each level. [default=1] range: (0,1]</td>
</tr>
<tr>
<td><code>lambda</code></td>
<td>L2 regularization term on weights, increase this value will make model more conservative. [default=1]</td>
</tr>
<tr>
<td><code>alpha</code></td>
<td>L1 regularization term on weights, increase this value will make model more conservative, defaults to 0.</td>
</tr>
<tr>
<td><code>tree_method</code></td>
<td>The tree construction algorithm used in XGBoost. options: 'auto', 'exact', 'approx' [default='auto']</td>
</tr>
<tr>
<td><code>sketch_eps</code></td>
<td>This is only used for approximate greedy algorithm. This roughly translated into O(1 / sketch_eps) number of bins. Compared to directly select number of bins, this comes with theoretical guarantee with sketch accuracy. [default=0.03] range: (0, 1]</td>
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<tr>
<td><code>scale_pos_weight</code></td>
<td>Control the balance of positive and negative weights, useful for unbalanced classes. A typical value to consider: sum(negative cases) / sum(positive cases). [default=1]</td>
</tr>
<tr>
<td><code>sample_type</code></td>
<td>Parameter for Dart booster. Type of sampling algorithm. &quot;uniform&quot;: dropped trees are selected uniformly. &quot;weighted&quot;: dropped trees are selected in proportion to weight. [default=&quot;uniform&quot;]</td>
</tr>
<tr>
<td><code>normalize_type</code></td>
<td>Parameter of Dart booster. type of normalization algorithm, options: 'tree', 'forest'. [default=&quot;tree&quot;]</td>
</tr>
<tr>
<td><code>rate_drop</code></td>
<td>Parameter of Dart booster. dropout rate. [default=0.0] range: [0.0, 1.0]</td>
</tr>
<tr>
<td><code>skip_drop</code></td>
<td>Parameter of Dart booster. probability of skip dropout. If a dropout is skipped, new trees are added in the same manner as gbtree. [default=0.0] range: [0.0, 1.0]</td>
</tr>
<tr>
<td><code>lambda_bias</code></td>
<td>Parameter of linear booster L2 regularization term on bias, default 0 (no L1 reg on bias because it is not important.)</td>
</tr>
<tr>
<td><code>tree_limit</code></td>
<td>Limit number of trees in the prediction; defaults to 0 (use all trees.)</td>
</tr>
<tr>
<td><code>num_round</code></td>
<td>The number of rounds for boosting.</td>
</tr>
<tr>
<td><code>num_workers</code></td>
<td>number of workers used to train xgboost model. Defaults to 1.</td>
</tr>
<tr>
<td><code>nthread</code></td>
<td>Number of threads used by per worker. Defaults to 1.</td>
</tr>
<tr>
<td><code>use_external_memory</code></td>
<td>The tree construction algorithm used in XGBoost. options: 'auto', 'exact', 'approx' [default='auto']</td>
</tr>
<tr>
<td><code>silent</code></td>
<td>0 means printing running messages, 1 means silent mode. default: 0</td>
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</tbody>
</table>
custom_obj | Customized objective function provided by user. Currently unsupported.
custom_eval | Customized evaluation function provided by user. Currently unsupported.
missing | The value treated as missing. default: Float.NaN
seed | Random seed for the C++ part of XGBoost and train/test splitting.
timeout_request_workers | the maximum time to wait for the job requesting new workers. default: 30 minutes
checkpoint_path | The hdfs folder to load and save checkpoint boosters.
checkpoint_interval | Param for set checkpoint interval (>= 1) or disable checkpoint (-1). E.g. 10 means that the trained model will get checkpointed every 10 iterations. Note: checkpoint_path must also be set if the checkpoint interval is greater than 0.
base_score | Param for initial prediction (aka base margin) column name. Defaults to 0.5.
train_test_ratio | Fraction of training points to use for testing.
num_early_stopping_rounds | If non-zero, the training will be stopped after a specified number of consecutive increases in any evaluation metric.
objective_type | The learning objective type of the specified custom objective and eval. Corresponding type will be assigned if custom objective is defined options: regression, classification.
eval_metric | Evaluation metrics for validation data, a default metric will be assigned according to objective(rmse for regression, and error for classification, mean average precision for ranking). options: rmse, mae, logloss, error, merror, mlogloss, auc, aucpr, ndcg, map, gamma-deviance
maximize_evaluation_metrics | Whether to maximize evaluation metrics. Defaults to FALSE (for minimization.)
base_margin_col | Param for initial prediction (aka base margin) column name.
weight_col | Weight column.
features_col | Features column name, as a length-one character vector. The column should be single vector column of numeric values. Usually this column is output by ft_r_formula.
label_col | Label column name. The column should be a numeric column. Usually this column is output by ft_r_formula.
prediction_col | Prediction column name.
uid | A character string used to uniquely identify the ML estimator.
... | Optional arguments; see Details.
**Details**

When `x` is a `tbl_spark` and `formula` (alternatively, `response` and `features`) is specified, the function returns a `ml_model` object wrapping a `ml_pipeline_model` which contains data pre-processing transformers, the ML predictor, and, for classification models, a post-processing transformer that converts predictions into class labels. For classification, an optional argument `predicted_label_col` (defaults to "predicted_label") can be used to specify the name of the predicted label column. In addition to the fitted `ml_pipeline_model`, `ml_model` objects also contain a `ml_pipeline` object where the ML predictor stage is an estimator ready to be fit against data. This is utilized by `ml_save` with `type = "pipeline"` to facilitate model refresh workflows.

**Value**

The object returned depends on the class of `x`.

- `spark_connection`: When `x` is a `spark_connection`, the function returns an instance of a `ml_estimator` object. The object contains a pointer to a Spark Predictor object and can be used to compose Pipeline objects.
- `ml_pipeline`: When `x` is a `ml_pipeline`, the function returns a `ml_pipeline` with the predictor appended to the pipeline.
- `tbl_spark`: When `x` is a `tbl_spark`, a predictor is constructed then immediately fit with the input `tbl_spark`, returning a prediction model.
- `tbl_spark`, with `formula` specified: When `formula` is specified, the input `tbl_spark` is first transformed using a `RFormula` transformer before being fit by the predictor. The object returned in this case is a `ml_model` which is a wrapper of a `ml_pipeline_model`.

**See Also**

See [http://spark.apache.org/docs/latest/ml-classification-regression.html](http://spark.apache.org/docs/latest/ml-classification-regression.html) for more information on the set of supervised learning algorithms.

**Examples**

```r
library(sparkxgb)
library(sparklyr)
sc <- spark_connect(master = "local")
mtcars_tbl <- copy_to(sc, mtcars)

xgb_model <- xgboost_regressor(
  mtcars_tbl,
  mpg ~ .
)
```
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