# Boosting

# AdaBoost, Gradient Boosting, XGboost

# SYS 6018 | Spring 2021

# boosting.pdf

# **Contents**

1	Boos	sting	2
2	AdaBoost		
	2.1	Adaboost Algorithm	4
	2.2	AdaBoost Details	7
	2.3	R package ada	8
3	Gra	dient Boosting	12
	3.1	Gradient Descent	12
	3.2	$L_2$ Boosting	12
	3.3	GBM (Gradient Boosting Machine)	15
	3.4	xgboost (Extreme Gradient Boosting)	16
	3.5	CatBoost	17
	3.6	LightGBM	18

# 1 Boosting

Boosting is a sequential ensemble method.

## **Boosting Sketch**

- A boosting model can be written as a generic ensemble
  - *M* is the number of base learners
  - $\hat{a}_k$  is the weight for the kth base learner
  - $\hat{g}_k(x)$  is the prediction from the kth base learner

$$\hat{f}_M(x) = \sum_{k=1}^M \hat{a}_k \, \hat{g}_k(x)$$

• The big difference with boosting models is that the base learners are **fitted sequentially**, and the best model at stage m is dependent on all models fit prior to stage m.

$$\hat{f}_{m+1}(x) = \underset{a, g(x)}{\arg\min} \sum_{i=1}^{n} L(y_i, \hat{f}_m(x_i) + a g(x_i))$$

- Boosting is primarily a bias reducer
  - The base models are often simple/weak (low variance, but high bias) models (like shallow trees)
- The complexity of the final model is based on i) the complexity of the base learners and ii) the number of iterations
  - Boosting models will overfit as the number of iterations increases
    - \* Early stopping is necessary
    - \* Less of a problem for hard classification problems with balanced data
  - Can apply *shrinkage*, or reducing  $|a_k|$ , to reduce complexity

- There are two main versions of boosting:
  - Gradient Boosting: fits the next model in the sequence  $\hat{g}_k(x)$  to the (pseudo) residuals calculated from the predictions on the previous models
  - AdaBoost: fits the next model to sequentially weighted observations. The weights are proportional to the how poorly the current models predict the observation.

### 2 AdaBoost

AdaBoost was motivated by the idea that many *weak* leaners can be combined to produce a *strong* aggregate model.

- AdaBoost is for binary classification problems
- Trees are a popular base learner
  - Weak learners are usually used. For trees, this means shallow depth.
- At each iteration, the current model is evaluated.
  - The *ensemble weight* of model m is based on its performance (on all the training data)
  - The observation weight of observation i is increased if it is mis-classified and decreased if it is correctly classified.
  - Thus, at each iteration, those observations that are mis-classified are weighted higher and get extra attention in the next iteration.
- Because Adaboost uses hard-classifiers, it is sensitive to unbalanced data and unequal misclassification costs.
  - Because the thresholds are set to p > .50
  - There are, of course, ways to account for unbalance and unequal costs in the algorithm
  - An improvement to AdaBoost, LogitBoost explicitly attempts to estimate the class probability during each iteration which will allow easier post-fitting adjustments for unequal costs

## 2.1 Adaboost Algorithm

## Algorithm: AdaBoost (Discrete)

#### **Inputs**:

- $D = \{(x_i, y_i)_{i=1}^n$ , where  $y_i \in \{-1, 1\}$
- Tuning parameters for base model  $\hat{q}$
- Maximum number of iterations, M

#### Algorithm:

- 1. Initialize observation weights  $w_i = 1/n$  for all i
- 2. For k = 1 to M:
  - a. Fit a *classifier*  $\hat{g}_k(x)$  that maps  $(x_i, w_i)$  to  $\{-1, 1\}$ . In other words, the classifier must make a hard classification using weighted observations.
  - b. Compute the weighted mis-classification rate

$$e_k = \frac{\sum_{i=1}^n w_i \, \mathbb{1}(y_i \neq \hat{g}_k(x_i))}{\sum_{i=1}^n w_i}$$

c. Calculate the *coefficient* for model k (ensemble weight)

$$\hat{a}_k = \log\left(\frac{1 - e_k}{e_k}\right)$$

d. Update the *observations weights*. Increase weights for observations that are mis-classified by model  $\hat{g}_k$  and decrease weights for the correctly classified observations.

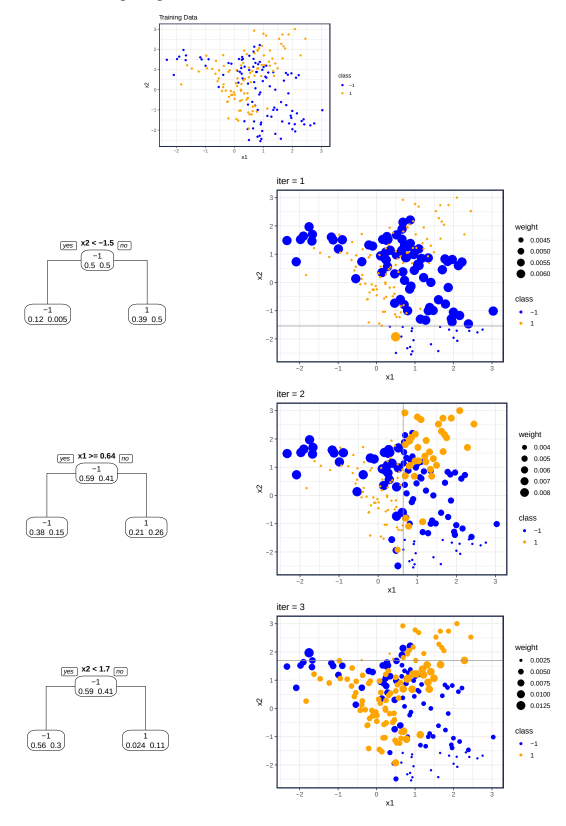
$$\begin{split} \tilde{w}_i &= w_i \cdot \exp\left(a_k \cdot \mathbb{1}(y_i \neq \hat{g}_k(x_i))\right) \\ w_i &= \frac{\tilde{w}_i}{\sum_{j=1}^n \tilde{w}_j} \end{split} \qquad \textit{(re-normalize weights)}$$

3. Output final ensemble  $\hat{f}_M(x) \in [-1, 1]$ 

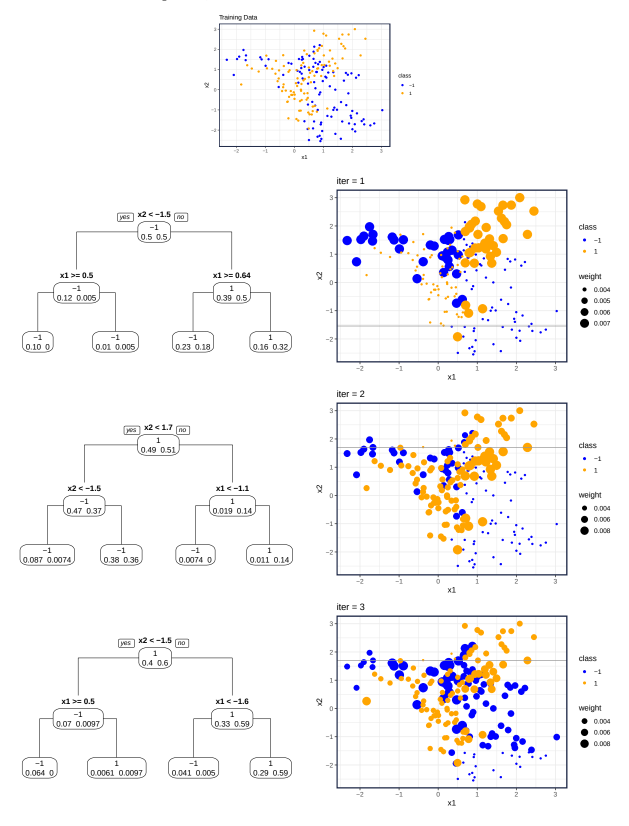
$$\hat{f}_M(x) = \sum_{k=1}^M \hat{a}_k \, \hat{g}_k(x)$$

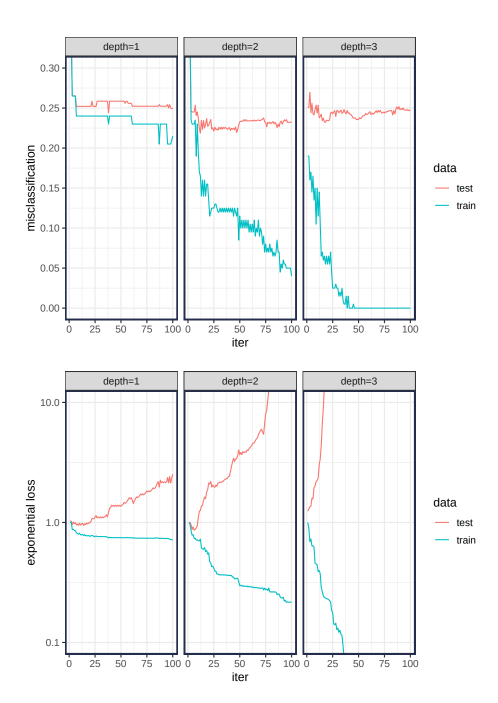
- Hard classification:  $\hat{f}_M(x) > 0$
- Or remap to a probability  $\hat{p}(x) = \frac{e^{2f}}{1 + e^{2f}}$  for thresholding

# 2.1.1 Illustration with Stumps (depth = 1, n.nodes=2)



## 2.1.2 Illustration with depth = 2, n.nodes=4





## 2.2 AdaBoost Details

- Adaboost uses a response variable of  $y \in \{-1, 1\}$
- AdaBoost uses the loss function:

$$L(y,f) = e^{-yf}$$

$$= \begin{cases} e^{-f} & y = +1 \\ e^{f} & y = -1 \end{cases}$$

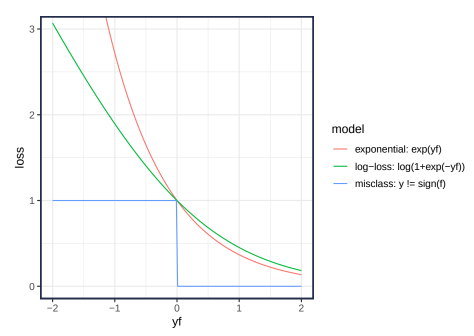
• Adaboost estimates the probability that Y = +1 as

$$\hat{p}(x) = \frac{e^{\hat{f}_M(x)}}{e^{-\hat{f}_M(x)} + e^{\hat{f}_M(x)}}$$
$$= \frac{e^{2\hat{f}_M(x)}}{1 + e^{2\hat{f}_M(x)}}$$

where 
$$p(x) = \Pr(Y = +1 \mid X = x)$$

• And  $\hat{f}(x)$  is an estimate of

$$\begin{split} \hat{f}_M(x) &= \frac{1}{2} \log \frac{\hat{p}(x)}{1 - \hat{p}(x)} \\ &= \frac{1}{2} \text{logit } \hat{p}(x) \end{split}$$



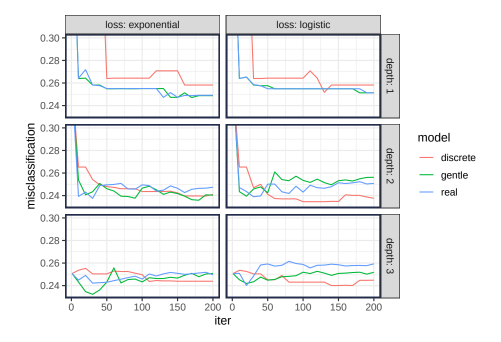
- Comparison with logistic regression (using log-loss / negative binomial log-likelihood)

  - $-\hat{f}(x) = \operatorname{logit} \hat{p}(x)$  $-\hat{p}(x) = \frac{e^{\hat{f}_{M}(x)}}{1 + e^{\hat{f}_{M}(x)}}$
  - Log-loss:  $\log(1 + e^{-yf})$  (using  $y \in \{-1, +1\}$ )

### 2.3 R package ada

The R package ada provides an implementation of AdaBoost (and related methods).

- See Friedman, J., Hastie, T., and Tibshirani, R. (2000). Additive Logistic Regression: A statistical view of boosting. Annals of Statistics, 28(2), 337-374. for the details of model variations
  - {Discrete, Real, Gentle} AdaBoost
  - Logitboost



# **Algorithm: Real AdaBoost**

#### **Inputs**:

- $D = \{(x_i, y_i)_{i=1}^n$ , where  $y_i \in \{-1, 1\}$
- Tuning parameters for base model  $\hat{g}$
- Maximum number of iterations, M

#### Algorithm:

- 1. Initialize observation weights  $w_i = 1/n$  for all i
- 2. For k = 1 to M:
  - a. Fit a model  $\hat{g}_k(x)$  that uses weighted inputs  $(x_i, w_i)$  to estimate a probability  $\hat{p}_k(x) = \widehat{\Pr}(Y = 1 \mid X = x)$ . In other words, the classifier must make a soft classification using weighted observations.
  - b. Set  $f_m(x) = \frac{1}{2} \operatorname{logit} \hat{p}_k(x)$
  - c. Update the *observations weights*. Increase weights for observations that are mis-classified by model  $\hat{g}_k$  and decrease weights for the correctly classified observations.

$$ilde{w}_i = w_i \cdot \exp\left(-y_i \hat{f}_m(x_i)\right)$$
 $w_i = rac{ ilde{w}_i}{\sum_{j=1}^n ilde{w}_j}$  (re-normalize weights)

3. Output final ensemble  $\hat{f}_M(x) \in [-1, 1]$ 

$$\hat{f}_M(x) = \sum_{k=1}^M a_k \, \hat{g}_k(x)$$

• Hard classification:  $\hat{f}_M(x) > 0$ 

• Or remap to a probability  $\hat{p}(x) = \frac{e^{2f}}{1+e^{2f}}$  for thresholding

## **Algorithm: Gentle AdaBoost**

## **Inputs**:

- $D = \{(x_i, y_i)_{i=1}^n$ , where  $y_i \in \{-1, 1\}$
- Tuning parameters for base model  $\hat{g}$
- Maximum number of iterations, M

#### Algorithm:

- 1. Initialize observation weights  $w_i = 1/n$  for all i and  $f_0(x) = 0$
- 2. For k = 1 to M:
  - a. Fit a model  $\hat{g}_k(x)$  with weighted least squares that estimates  $y_i$  using features  $x_i$  and weights  $w_i$ .
  - b. Update the *observations weights*. Increase weights for observations that are mis-classified by model  $\hat{g}_k$  and decrease weights for the correctly classified observations.

$$ilde{w}_i = w_i \cdot \exp\left(-y_i \hat{f}_m(x_i)\right)$$
 $w_i = \frac{ ilde{w}_i}{\sum_{j=1}^n ilde{w}_j}$  (re-normalize weights)

3. Output final ensemble  $\hat{f}_M(x) \in [-1, 1]$ 

$$\hat{f}_M(x) = \sum_{k=1}^M \hat{g}_k(x)$$

• Hard classification:  $\hat{f}_M(x) > 0$ 

## **Algorithm: LogitBoost**

#### **Inputs**:

- $D = \{(x_i, y_i)_{i=1}^n, \text{ where } y_i \in \{-1, 1\}$
- Tuning parameters for base model  $\hat{q}$
- Maximum number of iterations, M
- Let  $y_i^* = (y+1)/2 \in \{0,1\}$

#### Algorithm:

- 1. Initialize observation weights  $w_i = 1/n$  for all i and  $f_0(x) = 0$
- 2. For k = 1 to M:
  - a. Like in newton-raphson for logistic regression, calculate the working response and weights for all observations

$$z_i = \frac{y_i^* - p_i}{p_i(1 - p_i)}$$

$$w_i = p_i(1 - p_i)$$

- b. Fit a model  $\hat{g}_k(x)$  with weighted least squares that estimates  $z_i$  using features  $x_i$  and weights  $w_i$ .
- c. Update  $\hat{f}_k(x)=\hat{f}_{k-1}(x)+\hat{g}_k(x)/2$  and  $p_i=e^{\hat{f}_k(x)}/(e^{\hat{f}_k(x)}+e^{-\hat{f}_k(x)})$  3. Output final ensemble  $\hat{f}_M(x)\in\mathbb{R}$

$$\hat{f}_M(x) = \sum_{k=1}^{M} \frac{1}{2} \, \hat{g}_k(x)$$

- Hard classification:  $\hat{f}_M(x) > 0$
- Or remap to a probability  $\hat{p}(x) = \frac{e^{2f}}{1+e^{2f}}$  for thresholding

# 3 Gradient Boosting

The boosting model:

$$\hat{f}_M(x) = \sum_{k=1}^M \hat{a}_k \, \hat{g}_k(x)$$

Sequential Fitting:

$$\hat{f}_{k+1}(x) = \underset{a, g(x)}{\operatorname{arg max}} \sum_{i=1}^{n} L(y_i, \hat{f}_k(x_i) + a g(x_i))$$

The concept of gradient boosting is sequentially re-fit to the negative (functional) gradients of the loss function (or *pseudo* residuals).

- The same structure can be used for many different loss functions
  - it works the same for regression and classification
  - survival analysis, ranking, etc.

#### 3.1 Gradient Descent

- Our objective is to find the model (or model parameters) that minimize the loss function
- From any starting point, we can move toward the optimum using gradient descent:

$$f_{k+1} = f_k - \nu_k L'(f_k)$$

- $\nu_k > 0$  is the step-size
- $L'(f_k)$  is the functional derivative of the loss with respect to the model  $f_k$
- Boosting fits models sequentially:

$$\hat{f}_{k+1}(x) = \hat{f}_k(x) + \hat{a}_k \, \hat{g}_k(x)$$

• So we see a parallel; each boosting model  $\hat{g}_k(x)$  can be viewed as estimating the *negative derivative* of the loss function.

### 3.2 $L_2$ Boosting

 $L_2$  boosting is based on the squared error loss function

$$L(y_i, \hat{f}(x_i)) = \frac{1}{2}(y_i - \hat{f}(x_i))^2$$

• The negative gradients are

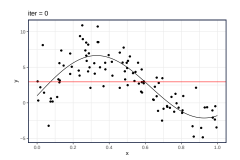
$$r_{i} = \left[ -\frac{\partial L(y_{i}, f_{i})}{\partial f_{i}} \right]_{f_{i} = \hat{f}(x_{i})}$$
$$= y_{i} - \hat{f}(x_{i})$$

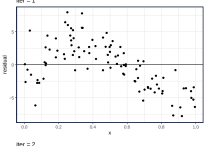
• L2 Boosting is simply re-fitting to the residuals.

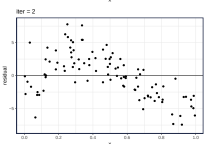
# Algorithm: $L_2$ Boosting

- 1. Initialize  $\hat{f}_0(x) = \bar{y}$
- 2. For k = 1 to M:
  - a. Calculate residuals  $r_i = y_i \hat{f}_{k-1}(x_i)$  for all i
  - b. Fit a base learner (e.g., regression trees) to the residuals  $\{(x_i,r_i)\}_{i=1}^n$  to get the model  $\hat{g}_k(x)$
  - c. Update the overall model  $\hat{f}_k(x) = f_{k-1}(x) + \nu \hat{g}_k(x)$   $0 \le \nu \le 1$  is the step-size (shrinkage)
- 3. Final model is  $\hat{f}_M(x) = \bar{y} + \sum_{k=1}^M \nu \hat{g}_k(x)$
- Like AdaBoost, emphasis is given to observations that are predicted poorly (large residuals)

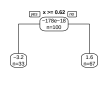
# 3.2.1 Illustration using stumps (depth=1, n.nodes=2, $\nu = .1$ )

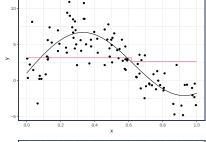


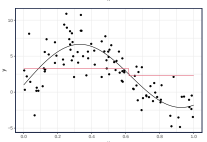


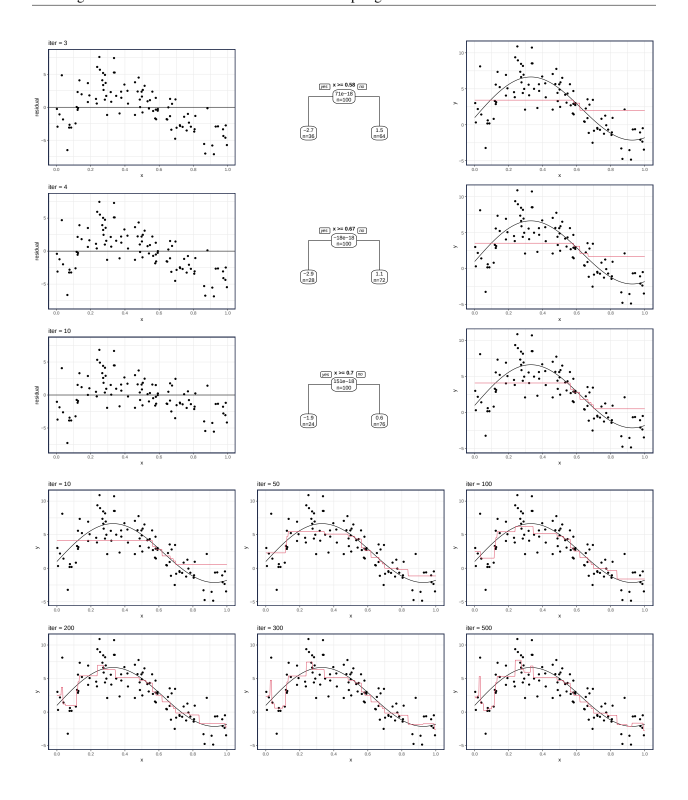












## 3.3 GBM (Gradient Boosting Machine)

- R package gbm
- GBM Documentation

#### 3.3.1 Model/Tree Tuning Parameters

- Tree depth (interaction.depth)
  - Grows trees to a depth specified by interaction.depth (unless there are not enough observations in the terminal nodes)
- Minimum number of observations allowed in the terminal nodes (n.minobsinnode)
- Sub-sampling (bag.fraction)
  - Stochastic Gradient Boosting
  - Sample (without replacement) at each iteration
- Loss Function (distribution)
  - The loss function is determined by the distribution argument
  - Use distribution="gaussian" for squared error
  - Other options are: bernoulli (for logistic regression), poisson (for Poisson regression), pairwise (for ranking/LambdaMart), adaboost (for the adaboost exponential loss), etc.

# 3.3.2 Boosting Tuning Parameters

- Number of iterations/trees (n.trees)
  - Use cross-validation (or out-of-bag) to find optimal value
  - Can use the helper function gbm.perf() to get the optimal value
- Shrinkage parameter (shrinkage)
  - Set small, but the smaller the shrinkage, the more iterations/trees need to be used
  - "Ranges from 0.001 to 0.100 usually work"
- Cross-validation (cv.folds)
  - gbm has a built in cross-validation
  - no way to manually set the folds

#### 3.3.3 Computational Settings

- Number of Cores (n.cores)
  - Only used when cross-validation is implemented

### 3.4 xgboost (Extreme Gradient Boosting)

- R package xgboost
- xgboost Documentation
- xgboost Model
- xbgoost Paper

#### 3.4.1 Model/Tree Tuning Parameters

- Different base leaners (booster)
  - gbtree is a tree
  - gblinear creates a (generalized) liner model (forward stagewise linear model)
- Tree building (tree\_method)
  - To speed up the fitting, only consider making splits at certain quantiles of the input vector (rather than considering every unique value)
- Sub-sampling (subsample)
  - Stochastic Gradient Boosting
  - Sample (without replacement) at each iteration
- Feature sampling (colsample\_bytree, colsample\_bylevel, colsample\_bynode)
  - Like used in Random Forest, the features/columns are subsampled
  - Can use a subsample of features for each tree, level, or node

#### **Model Complexity Parameters**

- Tree depth (max depth)
  - Grows trees to a depth specified by max\_depth (unless there are not enough observations in the terminal nodes)
  - Trees may not reach max\_depth if the gamma or min\_child\_weight arguments are set.
- Minimum number of observations (or sum of weights) allowed in the terminal nodes (min\_child\_weight)
- Pruning (gamma or min\_split\_loss)
  - Minimum loss reduction required to make a further partition on a leaf node of the tree
  - The larger gamma is, the more conservative the algorithm will be
- ElasticNet type penalty (lambda and alpha)
  - lambda is an  $L_2$  penalty
  - alpha is an  $L_1$  penalty
  - Recall that trees model the response as a *constant* in each region

$$\hat{f}_T(x) = \sum_{m=1}^M \hat{c}_m \, \mathbb{1}(x \in \hat{R}_m)$$

· Cost-complexity pruning found the optimal tree as the one that minimized the penalized loss objective

function:

$$C_{\gamma}(T) = \sum_{m=1}^{|T|} \operatorname{Loss}(T) + \gamma |T|$$

• XGBoost selects a tree at each iteration using the following penalized loss:

$$C_{\gamma,\lambda,\alpha}(T) = \sum_{m=1}^{|T|} \operatorname{Loss}(T) + \gamma |T| + \frac{\lambda}{2} \sum_{m=1}^{|T|} \hat{c}_m^2 + \alpha \sum_{m=1}^{|T|} |\hat{c}_m|$$

- Loss Function (objective)
  - The loss function is determined by the objective argument
  - Use req: squarederror for squared error
  - Other options are: reg:logistic or binary:logistic (for logistic regression), count:poisson (for Poisson regression), rank:pairwise (for ranking/LambdaMart), etc.

#### 3.4.2 Boosting Tuning Parameters

- Shrinkage parameter (eta or learning\_rate)
  - Set small, but the smaller the eta, the more iterations/trees need to be used
- Number of iterations/trees (num\_rounds)
  - Use cross-validation (or out-of-bag) to find optimal value
- Cross-validation (xqb.cv)
  - xgboost has a built in cross-validation
  - It is possible to manually set the folds

#### 3.4.3 Computational Settings

- Number of Threads (nthread)
- GPU Support (https://xgboost.readthedocs.io/en/latest/gpu/index.html)
  - Used for finding tree split points and evaluating/calculating the loss function

### 3.5 CatBoost

- R package: (https://github.com/catboost/catboost/tree/master/catboost/R-package)
- CatBoost Documentation
- Model/Tree Tuning Parameters:

• Boosting Tuning Parameters:

# 3.6 LightGBM

- $\bullet \ \ R \ Package: \ https://github.com/microsoft/LightGBM/tree/master/R-package$
- LightGBM Documentation
- Model/Tree Tuning Parameters:

• Boosting Tuning Parameters: