

Gradient Boosting

David S. Rosenberg

New York University

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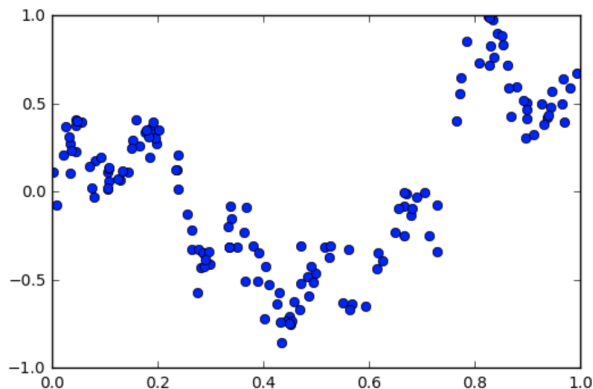
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Introduction

Nonlinear Regression

- Suppose we have the following regression problem:



- What are some options?
- basis functions, kernel methods, trees, neural nets, ...

Linear Model with Basis Functions

- Choose some basis functions on input space \mathcal{X} :

$$g_1, \dots, g_M : \mathcal{X} \rightarrow \mathbb{R}$$

- Predict with linear combination of basis functions:

$$f(x) = \sum_{m=1}^M v_m g_m(x)$$

- Can fit this using standard methods for linear models (e.g. least squares, lasso, ridge, etc.)
- In ML parlance, basis functions are called **features** or **feature functions**.

Not Limited to Regression

- Linear combination of basis functions:

$$f(x) = \sum_{m=1}^M v_m g_m(x)$$

- $f(x)$ is a number — for regression, it's exactly what we're looking for.
- Otherwise, $f(x)$ is often called a **score** function.
- It can be
 - thresholded to get a classification
 - transformed to get a probability
 - transformed to get a parameter of a probability distribution (e.g. Poisson regression)
 - used for ranking search results

Adaptive Basis Function Model

- Let's “learn” the basis functions.
- **Base hypothesis space** \mathcal{H} consisting of functions $h: \mathcal{X} \rightarrow \mathbf{R}$.
 - We will choose our “basis functions” or “features” from this set of functions.
- An **adaptive basis function expansion** over \mathcal{H} is

$$f(x) = \sum_{m=1}^M v_m h_m(x),$$

where $v_m \in \mathbf{R}$ and $h_m \in \mathcal{H}$ are chosen based on training data.

Adaptive Basis Function Model

- Base hypothesis space: \mathcal{H} of real-valued functions
- Combined hypothesis space: \mathcal{F}_M :

$$\mathcal{F}_M = \left\{ \sum_{m=1}^M v_m h_m(x) \mid v_m \in \mathbf{R}, h_m \in \mathcal{H}, m = 1, \dots, M \right\}$$

- Suppose we're given some data $\mathcal{D} = ((x_1, y_1), \dots, (x_n, y_n))$.
- Learning is choosing $v_1, \dots, v_M \in \mathbf{R}$ and $h_1, \dots, h_M \in \mathcal{H}$ to fit \mathcal{D} .

Empirical Risk Minimization

- We'll consider learning by **empirical risk minimization**:

$$\hat{f} = \arg \min_{f \in \mathcal{F}_M} \frac{1}{n} \sum_{i=1}^n \ell(y_i, f(x_i)),$$

for some **loss function** $\ell(y, \hat{y})$.

- Write ERM objective function as

$$J(v_1, \dots, v_M, h_1, \dots, h_M) = \frac{1}{n} \sum_{i=1}^n \ell \left(y_i, \sum_{m=1}^M v_m h_m(x) \right).$$

- How to optimize J ? i.e. how to learn?

- **Suppose** our base hypothesis space is parameterized by $\Theta = \mathbf{R}^b$:

$$J(v_1, \dots, v_M, \theta_1, \dots, \theta_M) = \frac{1}{n} \sum_{i=1}^n \ell \left(y_i, \sum_{m=1}^M v_m h(x; \theta_m) \right).$$

- Can we differentiate J w.r.t. v_m 's and θ_m 's? Optimize with SGD?
- For **some** hypothesis spaces and typical loss functions, yes!
- Neural networks fall into this category! (h_1, \dots, h_M are neurons of last hidden layer.)

What if Gradient Based Methods Don't Apply?

- What if base hypothesis space \mathcal{H} consists of decision trees?
- Can we even parameterize trees with $\Theta = \mathbf{R}^b$?
- Even if we could for some set of trees,
 - predictions would not change continuously w.r.t. $\theta \in \Theta$,
 - and so certainly not differentiable.
- Today we'll discuss **gradient boosting**. It applies whenever
 - our loss function is [sub]differentiable w.r.t. training predictions $f(x_i)$, and
 - we can do regression with the base hypothesis space \mathcal{H} (e.g. regression trees).

- Forward stagewise additive modeling (FSAM)
 - example: L^2 Boosting
 - example: exponential loss gives AdaBoost
 - Not clear how to do it with many other losses, including logistic loss
- Gradient Boosting
 - example: logistic loss gives BinomialBoost
- Variations on Gradient Boosting
 - step size selection
 - stochastic row/column selection
 - Newton step direction
 - XGBoost

Forward Stagewise Additive Modeling

Forward Stagewise Additive Modeling (FSAM)

- FSAM is an iterative optimization algorithm for fitting adaptive basis function models.
- Start with $f_0 \equiv 0$.
- After $m-1$ stages, we have

$$f_{m-1} = \sum_{i=1}^{m-1} \nu_i h_i.$$

- In m 'th round, we want to find
 - **step direction** $h_m \in \mathcal{H}$ (i.e. a basis function) and
 - **step size** $\nu_i > 0$
- such that

$$f_m = f_{m-1} + \nu_i h_m$$

improves objective function value by as much as possible.

Forward Stagewise Additive Modeling for ERM

① Initialize $f_0(x) = 0$.

② For $m = 1$ to M :

① Compute:

$$(\nu_m, h_m) = \arg \min_{\nu \in \mathbf{R}, h \in \mathcal{H}} \frac{1}{n} \sum \ell \left(y_i, f_{m-1}(x_i) + \underbrace{\nu h(x_i)}_{\text{new piece}} \right).$$

② Set $f_m = f_{m-1} + \nu_m h$.

③ Return: f_M .

Example: L^2 Boosting

Example: L^2 Boosting

- Suppose we use the **square loss**. Then in each step we minimize

$$J(v, h) = \frac{1}{n} \sum_{i=1}^n \left(y_i - \left[f_{m-1}(x_i) + \underbrace{vh(x_i)}_{\text{new piece}} \right] \right)^2$$

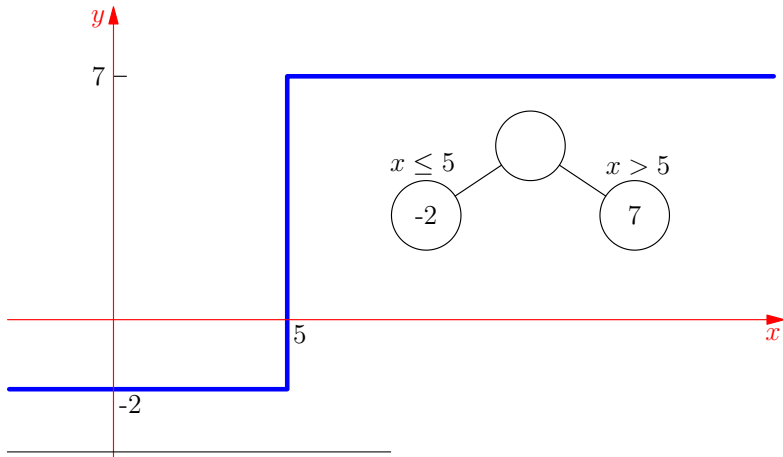
- If \mathcal{H} is closed under rescaling (i.e. if $h \in \mathcal{H}$, then $vh \in \mathcal{H}$ for all $h \in \mathbf{R}$), then don't need v .
- Take $v = 1$ and minimize

$$J(h) = \frac{1}{n} \sum_{i=1}^n ([y_i - f_{m-1}(x_i)] - h(x_i))^2$$

- This is just fitting the residuals with least-squares regression!
- If we can do regression with our base hypothesis space \mathcal{H} , then we're set!

Regression Stumps

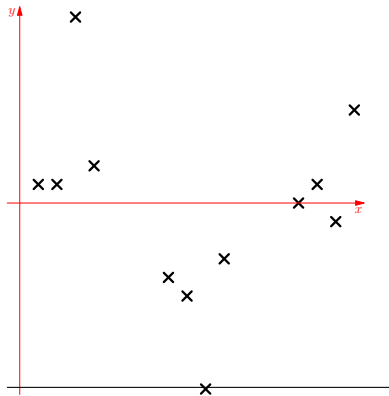
- A **regression stump** is a regression tree with a single split.
- A **regression stump** is a function of the form $h(x) = a1(x_i \leq c) + b1(x_i > c)$.



Plot courtesy of Brett Bernstein.

L^2 Boosting with Decision Stumps: Demo

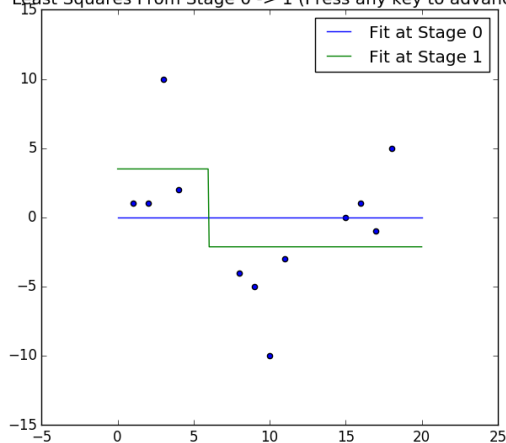
- Consider FSAM with L^2 loss (i.e. L^2 Boosting)
- For base hypothesis space of **regression stumps**
- Data we'll fit with **code**:



Plot courtesy of Brett Bernstein.

L^2 Boosting with Decision Stumps: Results

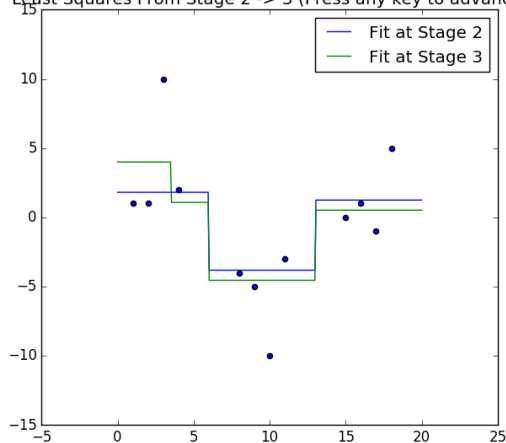
Least Squares From Stage 0 -> 1 (Press any key to advance)



Least Squares From Stage 1 -> 2 (Press any key to advance)

L^2 Boosting with Decision Stumps: Results

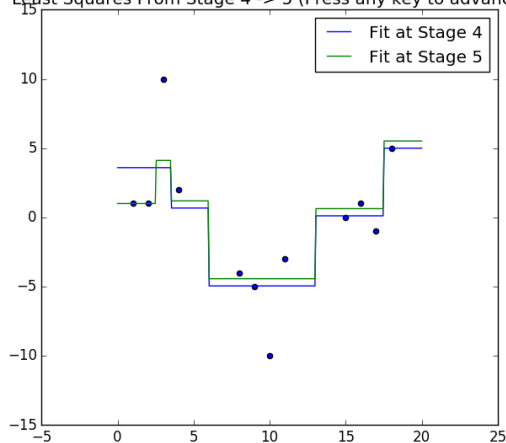
Least Squares From Stage 2 -> 3 (Press any key to advance)



Least Squares From Stage 3 -> 4 (Press any key to advance)

L^2 Boosting with Decision Stumps: Results

Least Squares From Stage 4 -> 5 (Press any key to advance)



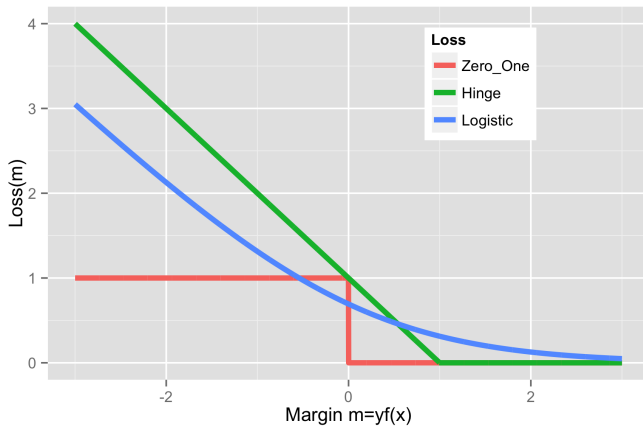
Least Squares From Stage 49 -> 50 (Press any key to advance)

Example: AdaBoost

The Classification Problem

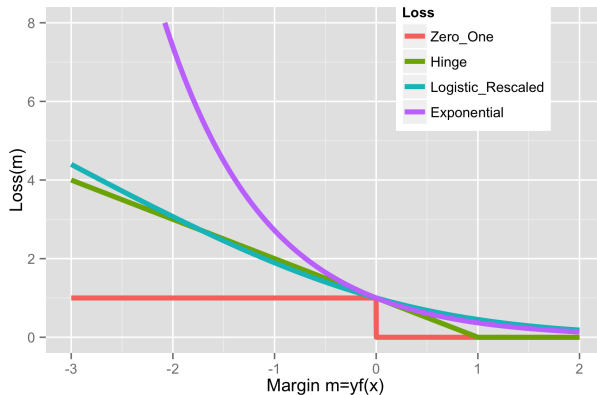
- Outcome space $\mathcal{Y} = \{-1, 1\}$
- Action space $\mathcal{A} = \mathbf{R}$
- Score function $f : \mathcal{X} \rightarrow \mathcal{A}$.
- Margin for example (x, y) is $m = yf(x)$.
 - $m > 0 \iff$ classification correct
 - Larger m is better.

Margin-Based Losses for Classification



Exponential Loss

- Introduce the **exponential loss**: $\ell(y, f(x)) = \exp(-yf(x))$.



FSAM with Exponential Loss

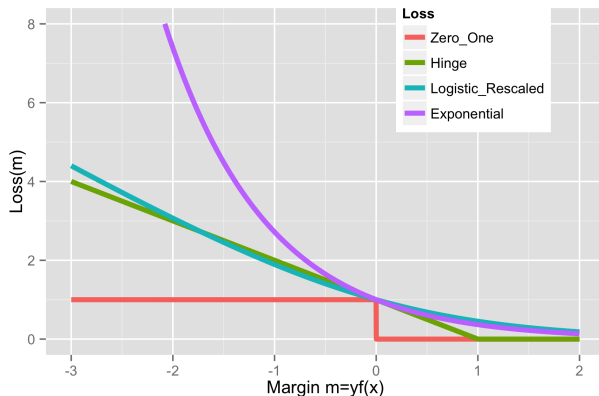
- Consider classification setting: $\mathcal{Y} = \{-1, 1\}$.
- Take loss function to be the **exponential loss**:

$$\ell(y, f(x)) = \exp(-yf(x)).$$

- Let \mathcal{H} be a base hypothesis space of classifiers $h: \mathcal{X} \rightarrow \{-1, 1\}$.
 - (Also assume \mathcal{H} closed under negation: $h \in \mathcal{H} \implies -h \in \mathcal{H}$)
- Then Forward Stagewise Additive Modeling (FSAM) reduces to a version of **AdaBoost**.
- Proof on [Spring 2017 Homework #6, Problem 4](#) (and see HTF Section 10.4).

Exponential Loss

- Note that exponential loss puts a very large weight on bad misclassifications.



- When Bayes error rate is high (e.g. $\mathbb{P}(f^*(X) \neq Y) = 0.25$)
 - e.g. there's some intrinsic randomness in the label
 - e.g. training examples with same input, but different classifications.
- Best we can do is predict the most likely class for each X .
- Some training predictions **should be wrong**,
 - because example doesn't have majority class
 - AdaBoost / exponential loss puts a lot of focus on getting those right
- Empirically, AdaBoost has degraded performance in situations with
 - high Bayes error rate, or when there's
 - high “**label noise**”
- Logistic loss performs better in settings with high Bayes error

FSAM for Other Loss Functions

- We know how to do FSAM for certain loss functions
 - e.g square loss, absolute loss, exponential loss,...
- In each case, happens to reduce to another problem we know how to solve, at least approximately.
- However, not clear how to do FSAM in general.
- For example, logistic loss / cross-entropy loss?

Gradient Boosting / “Anyboost”

FSAM Is Iterative Optimization

- The FSAM step

$$(\nu_m, h_m) = \arg \min_{\nu \in \mathbf{R}, h \in \mathcal{H}} \sum_{i=1}^n \ell \left(y_i, f_{m-1}(x_i) + \underbrace{\nu h(x_i)}_{\text{new piece}} \right).$$

- Hard part: finding the **best step direction** h .
- What if we looked for the **locally best** step direction?
 - like in gradient descent

“Functional” Gradient Descent

- We want to minimize

$$J(f) = \sum_{i=1}^n \ell(y_i, f(x_i)).$$

- In some sense, we want to take the gradient w.r.t. “ f ”, whatever that means.
- $J(f)$ only depends on f at the n training points.
- Define

$$\mathbf{f} = (f(x_1), \dots, f(x_n))^T$$

and write the objective function as

$$J(\mathbf{f}) = \sum_{i=1}^n \ell(y_i, \mathbf{f}_i).$$

Functional Gradient Descent: Unconstrained Step Direction

- Consider gradient descent on

$$J(\mathbf{f}) = \sum_{i=1}^n \ell(y_i, \mathbf{f}_i).$$

- The **negative gradient step direction** at \mathbf{f} is

$$\begin{aligned} -\mathbf{g} &= -\nabla_{\mathbf{f}} J(\mathbf{f}) \\ &= -(\partial_{\mathbf{f}_1} \ell(y_1, \mathbf{f}_1), \dots, \partial_{\mathbf{f}_n} \ell(y_n, \mathbf{f}_n)) \end{aligned}$$

which we can easily calculate.

- $-\mathbf{g} \in \mathbf{R}^n$ is the direction we want to change each of our n predictions on training data.
- Eventually we need more than just \mathbf{f} , which is just predictions on training.

Functional Gradient Descent: Projection Step

- Unconstrained step direction is

$$-\mathbf{g} = -\nabla_{\mathbf{f}} J(\mathbf{f}) = -(\partial_{\mathbf{f}_1} \ell(y_1, \mathbf{f}_1), \dots, \partial_{\mathbf{f}_n} \ell(y_n, \mathbf{f}_n)).$$

- Also called the “**pseudo-residuals**”
 - (for square loss, they’re exactly the residuals)
- Find the closest base hypothesis $h \in \mathcal{H}$ (in the ℓ^2 sense):

$$\min_{h \in \mathcal{H}} \sum_{i=1}^n (-\mathbf{g}_i - h(x_i))^2.$$

- This is a least squares regression problem over hypothesis space \mathcal{H} .
- Take the $h \in \mathcal{H}$ that best approximates $-\mathbf{g}$ as our step direction.

Functional Gradient Descent: Step Size

- Finally, we choose a stepsize.
- Option 1 (Line search):

$$\nu_m = \arg \min_{\nu > 0} \sum_{i=1}^n \ell\{y_i, f_{m-1}(x_i) + \nu h_m(x_i)\}.$$

- Option 2: (Shrinkage parameter – **more common**)
 - We consider $\nu = 1$ to be the full gradient step.
 - Choose a fixed $\nu \in (0, 1)$ – called a **shrinkage parameter**.
 - A value of $\nu = 0.1$ is typical – optimize as a hyperparameter .

The Gradient Boosting Machine Ingredients (Recap)

- Take any loss function [sub]differentiable w.r.t. the prediction
- Choose a base hypothesis space for regression.
- Choose number of steps (or a stopping criterion).
- Choose step size methodology.
- Then you're good to go!

Example: BinomialBoost

BinomialBoost: Gradient Boosting with Logistic Loss

- Recall the logistic loss for classification, with $\mathcal{Y} = \{-1, 1\}$:

$$\ell(y, f(x)) = \log(1 + e^{-yf(x)})$$

- Pseudoresidual for i 'th example is negative derivative of loss w.r.t. prediction:

$$\begin{aligned} r_i &= -\partial_{f(x_i)} \left[\log(1 + e^{-y_i f(x_i)}) \right] \\ &= \frac{y_i e^{-y_i f(x_i)}}{1 + e^{-y_i f(x_i)}} \\ &= \frac{y_i}{1 + e^{y_i f(x_i)}} \end{aligned}$$

BinomialBoost: Gradient Boosting with Logistic Loss

- Pseudoresidual for i th example:

$$r_i = -\partial_{f(x_i)} \left[\log \left(1 + e^{-y_i f(x_i)} \right) \right] = \frac{y_i}{1 + e^{y_i f(x_i)}}$$

- So if $f_{m-1}(x)$ is prediction after $m-1$ rounds, step direction for m 'th round is

$$h_m = \arg \min_{h \in \mathcal{H}} \sum_{i=1}^n \left[\left(\frac{y_i}{1 + e^{y_i f_{m-1}(x_i)}} \right) - h(x_i) \right]^2.$$

- And $f_m(x) = f_{m-1}(x) + \eta h_m(x)$.

Gradient Tree Boosting

Gradient Tree Boosting

- One common form of gradient boosting machine takes

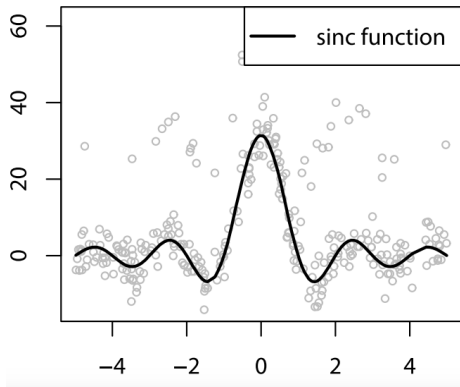
$$\mathcal{H} = \{\text{regression trees of size } J\},$$

where J is the number of terminal nodes.

- $J = 2$ gives decision stumps
- HTF recommends $4 \leq J \leq 8$ (but more recent results use much larger trees)
- Software packages:
 - Gradient tree boosting is implemented by the **gbm package** for R
 - as `GradientBoostingClassifier` and `GradientBoostingRegressor` in **sklearn**
 - **xgboost** and **lightGBM** are state of the art for speed and performance

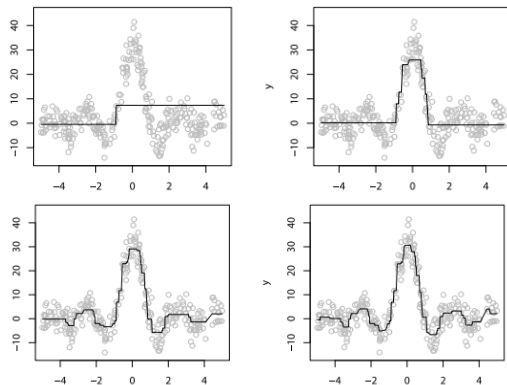
GBM Regression with Stumps

Sinc Function: Our Dataset



From Natekin and Knoll's "Gradient boosting machines, a tutorial"

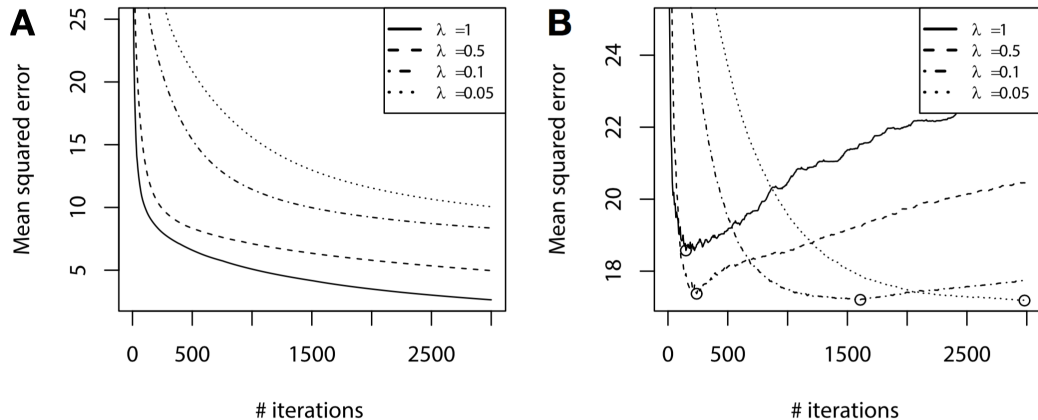
Minimizing Square Loss with Ensemble of Decision Stumps



Decision stumps with 1, 10, 50, and 100 steps, step size $\lambda = 1$.

Figure 3 from Natekin and Knoll's "Gradient boosting machines, a tutorial"

Step Size as Regularization



Performance vs rounds of boosting and step size. (Left is training set, right is validation set)

Figure 5 from Natekin and Knoll's "Gradient boosting machines, a tutorial"

Rule of Thumb

- The smaller the step size, the more steps you'll need.
- But never seems to make results worse, and often better.
- So set your step size as small as you have patience for.

Variations on Gradient Boosting

Stochastic Gradient Boosting

- For each stage,
 - choose random subset of data for computing projected gradient step.
 - “Typically, about 50% of the dataset size, can be much smaller for large training set.”
 - Fraction is called the **bag fraction**.
- Why do this?
 - Subsample percentage is additional regularization parameter – may help overfitting.
 - Faster.
- We can view this is a **minibatch method**.
 - we’re estimating the “true” step direction (the projected gradient) using a subset of data

Introduced by Friedman (1999) in [Stochastic Gradient Boosting](#).

Bag as Minibatch

- Just as we argued for minibatch SGD,
 - sample size needed for a good estimate of step direction is independent of training set size
- Minibatch size should depend on
 - the complexity of base hypothesis space
 - the complexity of the target function (Bayes decision function)
- Seems like an interesting area for both practical and theoretical pursuit.

Column / Feature Subsampling for Regularization

- Similar to random forest, randomly choose a subset of features for each round.
- XGBoost paper says: “According to user feedback, using column sub-sampling prevents overfitting even more so than the traditional row sub-sampling.”
- Zhao Xing (top Kaggle competitor) finds optimal percentage to be 20%-100%

Newton Step Direction

- For GBM, we find the closest $h \in \mathcal{F}$ to the negative gradient

$$-\mathbf{g} = -\nabla_{\mathbf{f}} J(\mathbf{f}).$$

- This is a “first order” method.
- Newton’s method is a “second order method”:
 - Find 2nd order (quadratic) approximation to J at \mathbf{f} .
 - Requires computing gradient and Hessian of J .
 - Newton step direction points towards minimizer of the quadratic.
 - Minimizer of quadratic is easy to find in closed form
- Boosting methods with projected Newton step direction:
 - LogitBoost (logistic loss function)
 - XGBoost (any loss – uses regression trees for base classifier)

Newton Step Direction for GBM

- Generically, second order Taylor expansion of J at \mathbf{f} in direction \mathbf{r}

$$J(\mathbf{f} + \mathbf{r}) = J(\mathbf{f}) + [\nabla_{\mathbf{f}} J(\mathbf{f})]^T \mathbf{r} + \frac{1}{2} \mathbf{r}^T [\nabla_{\mathbf{f}}^2 J(\mathbf{f})] \mathbf{r}$$

- For $J(\mathbf{f}) = \sum_{i=1}^n \ell(y_i, \mathbf{f}_i)$,

$$J(\mathbf{f} + \mathbf{r}) = \sum_{i=1}^n \left[\ell(y_i, \mathbf{f}_i) + g_i \mathbf{r}_i + \frac{1}{2} h_i \mathbf{r}_i^2 \right],$$

where $g_i = \partial_{\mathbf{f}_i} \ell(y_i, \mathbf{f}_i)$ and $h_i = \partial_{\mathbf{f}_i}^2 \ell(y_i, \mathbf{f}_i)$.

- Can find \mathbf{r} that minimizes $J(\mathbf{f} + \mathbf{r})$ in closed form.
- Can take step direction to be “projection” of \mathbf{r} into base hypothesis space \mathcal{H} .

XGBoost: Objective Function with Tree Penalty Term

- Adds explicit penalty term on tree complexity to the empirical risk:

$$\Omega(r) = \gamma T + \frac{1}{2} \lambda \sum_{j=1}^T w_j^2,$$

where $r \in \mathcal{H}$ is a regression tree from our base hypothesis space and

- T is the number of leaf nodes and
- w_j is the prediction in the j 'th node
- Objective function at step m :

$$J(r) = \sum_{i=1}^n \left[g_i r(x_i) + \frac{1}{2} h_i r(x_i)^2 \right] + \Omega(r)$$

- In XGBoost, they also use this objective to decide on tree splits
- See [XGBoost Introduction](#) for a nice introduction.

XGBoost: Rewriting objective function

- For a given tree, let $q(x_i)$ be x_i 's node assignment and w_j the prediction for node j .
- In each step of XGBoost we're looking for a tree that minimizes

$$\begin{aligned} & \sum_{i=1}^n \left[g_i w_{q(x_i)} + \frac{1}{2} h_i w_{q(x_i)}^2 \right] + \gamma T + \frac{1}{2} \lambda \sum_{j=1}^T w_j^2 \\ &= \sum_{\text{leaf node } j=1}^T \left[\left(\underbrace{\sum_{i \in I_j} g_i}_{G_j} \right) w_j + \frac{1}{2} \left(\underbrace{\sum_{i \in I_j} h_i + \lambda}_{H_j} \right) w_j^2 \right] + \gamma T, \end{aligned}$$

where $I_j = \{i \mid q(x_i) = j\}$ is set of training example indices landing in leaf j .

XGBoost: Simple Expression for Tree Penalty/Loss

- Simplifies to

$$\sum_{j=1}^T \left[G_j w_j + \frac{1}{2} (H_j + \lambda) w_j^2 \right] + \gamma T$$

- For fixed $q(x)$ (i.e. fixed tree partitioning), objective minimized when leaf node values are

$$w_j^* = -G_j / (H_j + \lambda).$$

- Plugging w_j^* back in, this objective reduces to

$$-\frac{1}{2} \sum_{j=1}^T \frac{G_j^2}{H_j + \lambda} + \gamma T,$$

which we can think of as the loss for tree partitioning function $q(x)$.

- If time were no issue, we could search over all trees to minimize this objective.

XGBoost: Building Tree Using Objective Function

- Expression to evaluate a tree's node assignment function $q(x)$:

$$-\frac{1}{2} \sum_{j=1}^T \frac{G_j^2}{H_j + \lambda} + \gamma T,$$

where $G_j = \sum_{i \in I_j} g_i$ for examples i assigned to leaf node j . And $H_j = \sum_{i \in I_j} h_i$.

- Suppose we're considering splitting some data into two nodes: L and R .
- Loss of tree with this one split is

$$-\frac{1}{2} \left[\frac{G_L^2}{H_L + \lambda} + \frac{G_R^2}{H_R + \lambda} \right] + 2\gamma.$$

- Without the split – i.e. a tree with a single leaf node, loss is

$$-\frac{1}{2} \left[\frac{(G_L + G_R)^2}{H_L + H_R + \lambda} \right] + \gamma.$$

XGBoost: Node Splitting Criterion

- We can define the **gain** of a split to be the reduction in objective between tree with and without split:

$$\text{Gain} = \frac{1}{2} \left[\frac{G_L^2}{H_L + \lambda} + \frac{G_R^2}{H_R + \lambda} - \frac{(G_L + G_R)^2}{H_L + H_R + \lambda} \right] - \gamma.$$

- Tree building method:
 - recursively choose split that maximizes the gain.