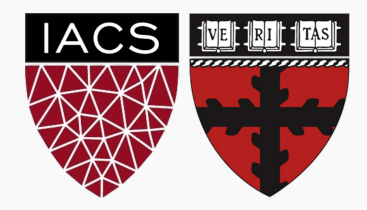
Lecture 17: Boosting

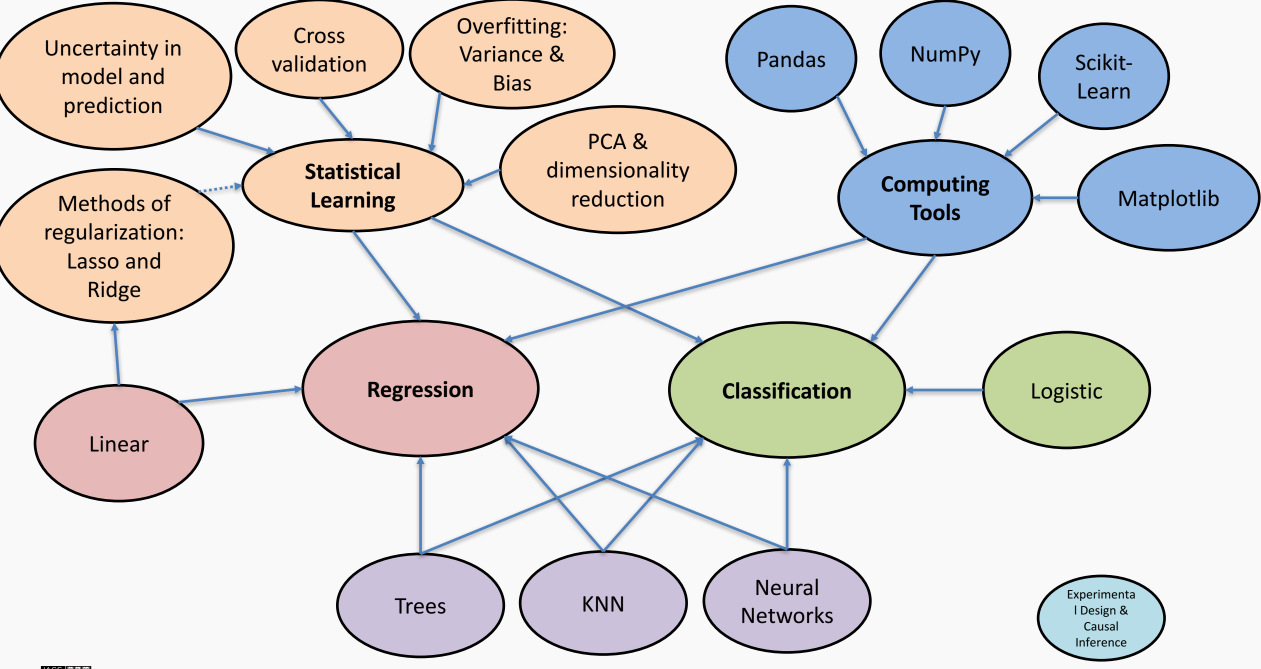
CS109A Introduction to Data Science Pavlos Protopapas, Kevin Rader and Chris Tanner



ANNOUNCEMENTS

- Homework 5 (209) due on Wednesday 11:59 pm, Nov 6
- Homework 4 grades. They should out soon 😳





Outline

- Review of Ensemble Methods
 - Finish Random Forest
- Boosting
 - Gradient Boosting
 - Set-up and intuition
 - Connection to Gradient Descent
 - The Algorithm
 - AdaBoost
- Other boosting algorithms



Bags and Forests of Trees

- Last time we examined how the short-comings of single decision tree models can be overcome by ensemble methods - making one model out of many trees.
- We focused on the problem of training large trees, these models have low bias but high variance.
- We compensated by training an ensemble of full decision trees and then averaging their predictions - thereby reducing the variance of our final model.



Bags and Forests of Trees (cont.)

<u>Bagging:</u>

- create an ensemble of trees, each trained on a bootstrap sample of the training set
- average the predictions.

Random forest:

- create an ensemble of trees, each trained on a bootstrap sample of the training set
- in each tree and each split, randomly select a subset of predictors, choose a predictor from this subset for splitting
- average the predictions

Note that the ensemble building aspects of both methods are embarrassingly parallel!



Random forest models have multiple hyper-parameters to tune:

- 1. the number of predictors to randomly select at each split
- 2. the total number of trees in the ensemble
- 3. the minimum leaf node size

In theory, each tree in the random forest is full, but in practice this can be computationally expensive (and added redundancies in the model), thus, imposing a minimum node size is not unusual.



There are standard (default) values for each of random forest hyperparameters recommended by long time practitioners, but generally these parameters should be tuned through **OOB** (making them data and problem dependent).

e.g. number of predictors to randomly select at each split:

- $\sqrt{N_j}$ for classification
- $-\frac{N}{3}$ for regression

Using out-of-bag errors, training and cross validation can be done in a single sequence - we cease training once the out-of-bag error stabilizes

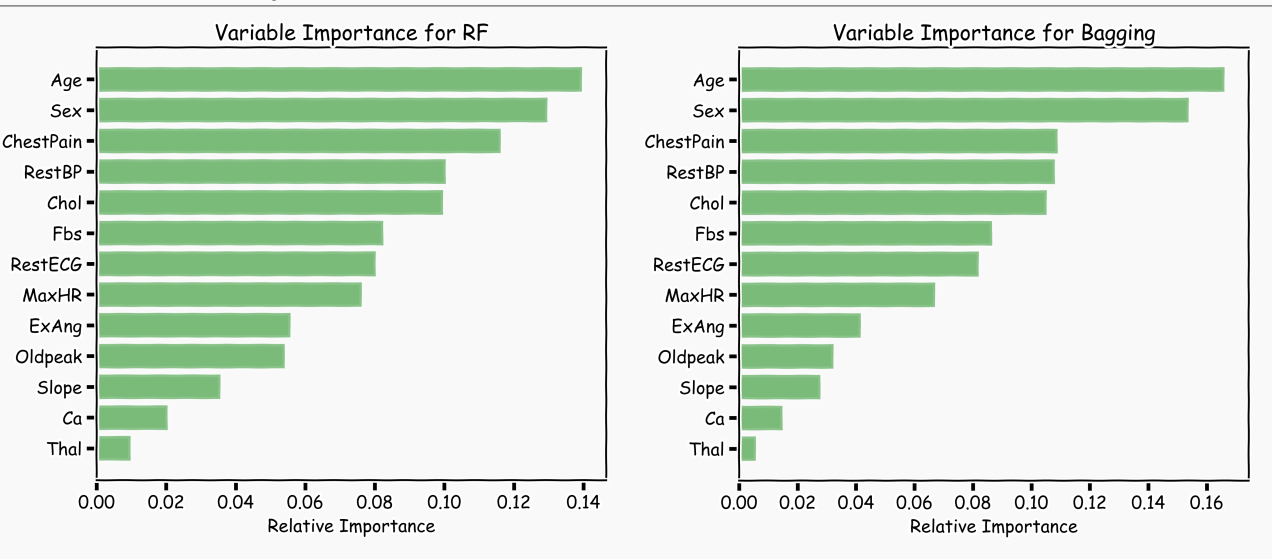


Same as with Bagging:

Calculate the total amount that the RSS (for regression) or Gini index (for classification) is decreased due to splits over a given predictor, averaged over all *B* trees.



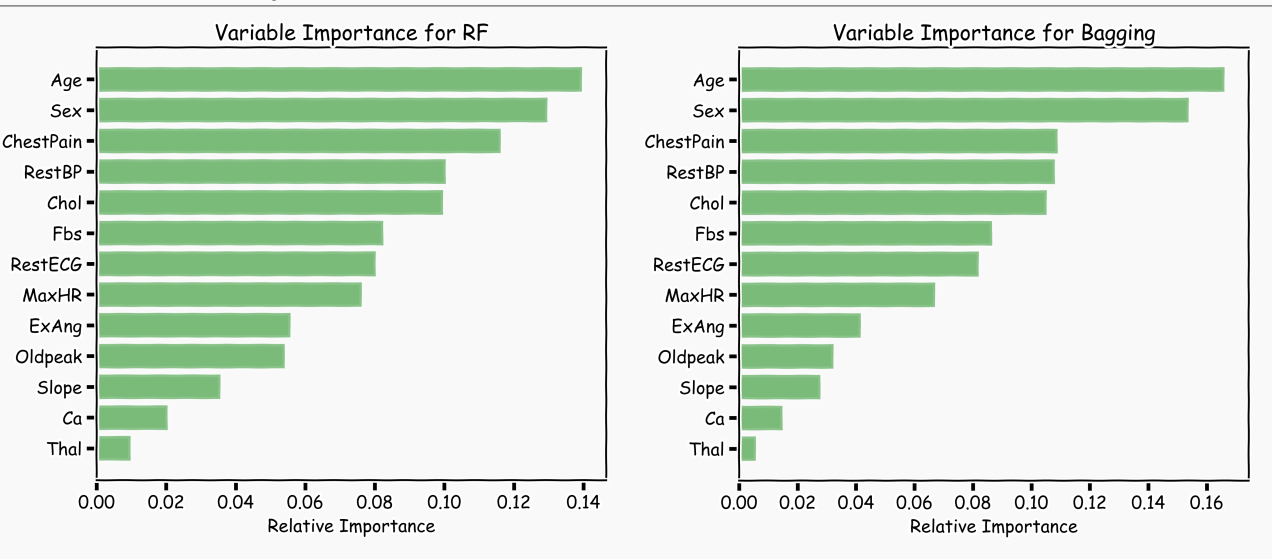
Variable Importance for RF





100 trees, max_depth=10

Variable Importance for RF





100 trees, max_depth=10

When the number of predictors is large, but the number of relevant predictors is small, random forests can perform poorly.

Question: Why?

In each split, the chances of selected a relevant predictor will be low and hence most trees in the ensemble will be weak models.



Final Thoughts on Random Forests (cont.)

Increasing the number of trees in the ensemble generally does not increase the risk of overfitting.

Again, by decomposing the generalization error in terms of bias and variance, we see that increasing the number of trees produces a model that is at least as robust as a single tree.

However, if the number of trees is too large, then the trees in the ensemble may become more correlated, and therefore increase the variance.



Final Thoughts on Random Forests (cont.)

Probabilities:

- Random Forrest Classifier (and bagging) can return probabilities.
- Question: How?



- Unbalance dataset
- Weighted samples
- Categorical data
- Missing data
- Why did we reject misclassification error?



Boosting



Question: Could we address the shortcomings of single decision trees models in some other way?

For example, rather than performing variance reduction on complex trees, can we decrease the bias of simple trees - make them more expressive?

Can we learn from our mistakes?

A solution to this problem, making an expressive model from simple trees, is another class of ensemble methods called **boosting**.



Boosting Algorithms



The key intuition behind boosting is that one can take an ensemble of simple models $\{T_h\}_{h \in H}$ and additively combine them into a single, more complex model.

Each model *T_h* might be a poor fit for the data, but a linear combination of the ensemble

$$T = \sum_{h} \lambda_h T_H$$

can be expressive/flexible.

Question: But which models should we include in our ensemble? What should the coefficients or weights in the linear combination be?



Gradient Boosting: the algorithm

Gradient boosting is a method for iteratively building a complex regression model *T* by adding simple models. Each new simple model added to the ensemble compensates for the weaknesses of the current ensemble.

1. Fit a simple model $T^{(0)}$ on the training data

 $\{(x_1, y_1), \dots, (x_N, y_N)\}\$

Set $T \leftarrow T^{(0)}$. Compute the residuals $\{r_1, \ldots, r_N\}$ for T.

2. Fit a simple model, $T^{(1)}$, to the current **residuals**, i.e. train using

 $\{(x_1, r_1), \dots, (x_N, r_N)\}$

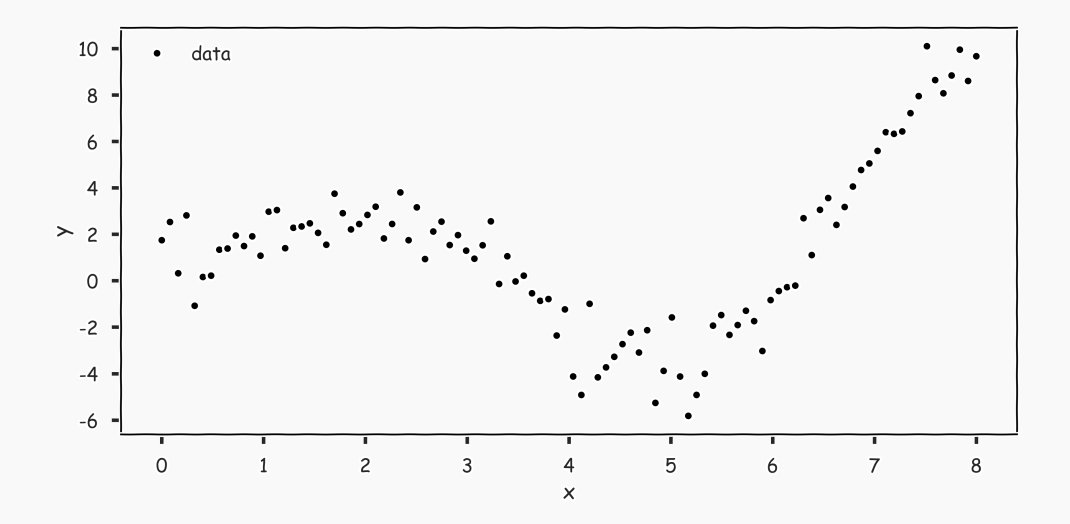
3. Set $T \leftarrow T + \lambda T^{(1)}$

4. Compute residuals, set $r_n \leftarrow r_n - \lambda T^i(x_n)$, n = 1, ..., N

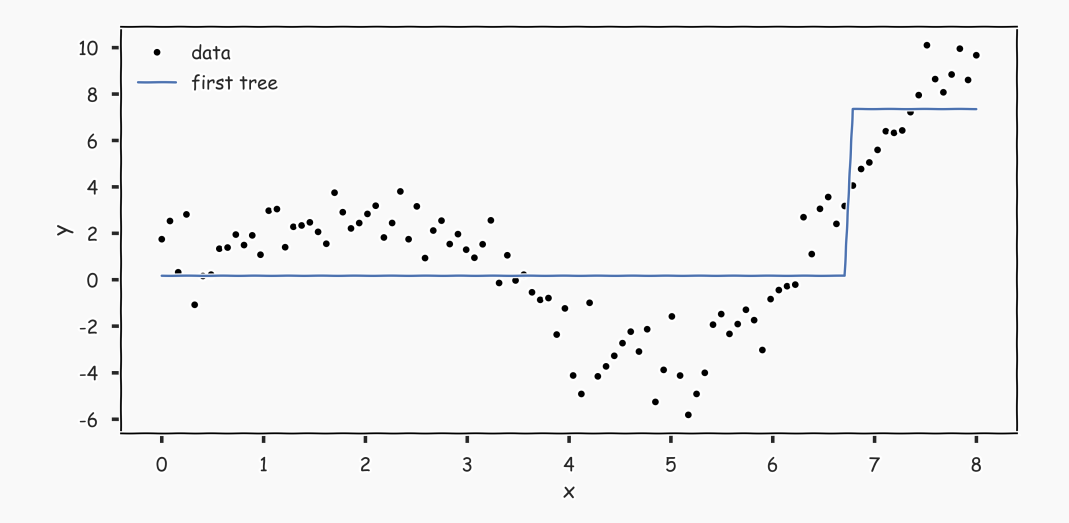
5. Repeat steps 2-4 until **stopping** condition met.

where λ is a constant called the learning rate.

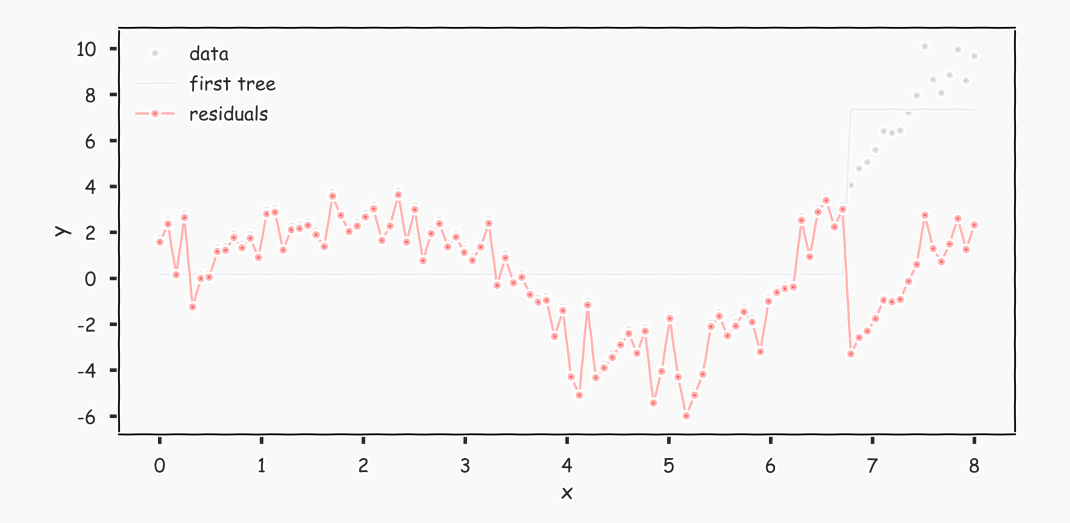




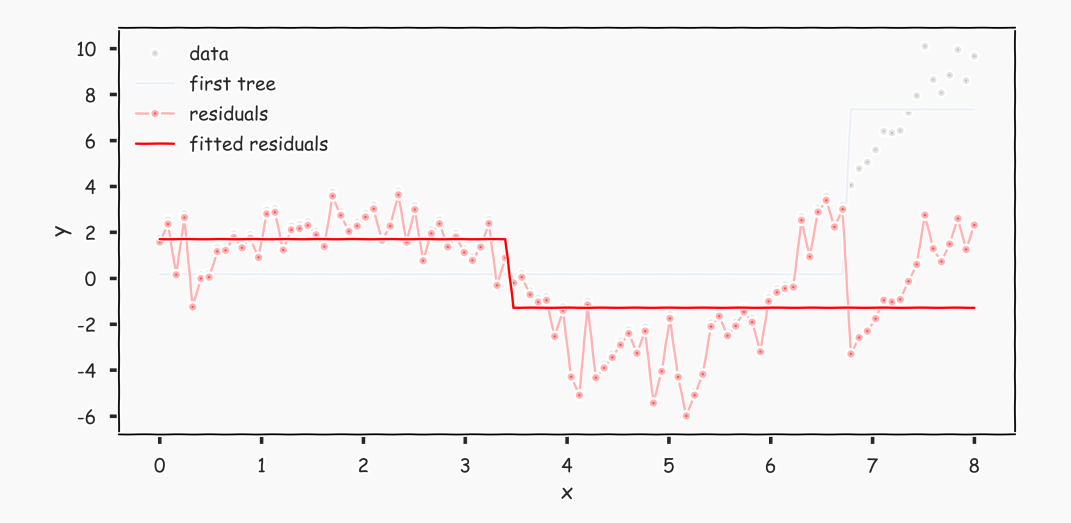




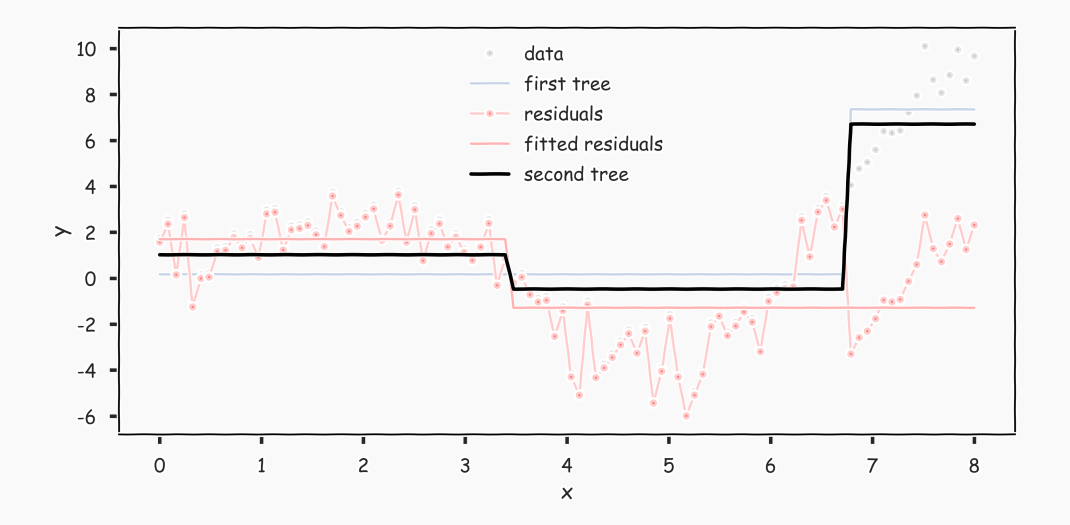




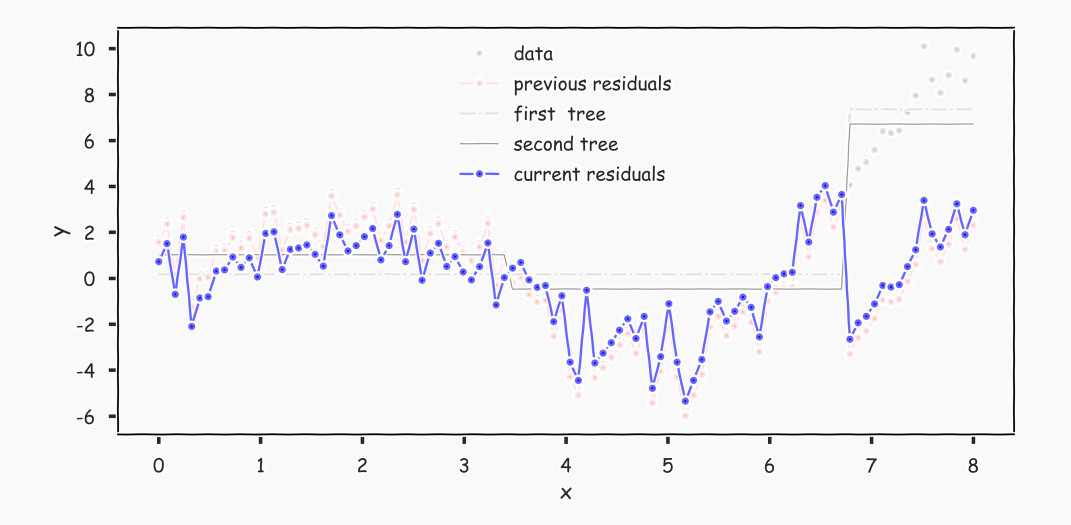






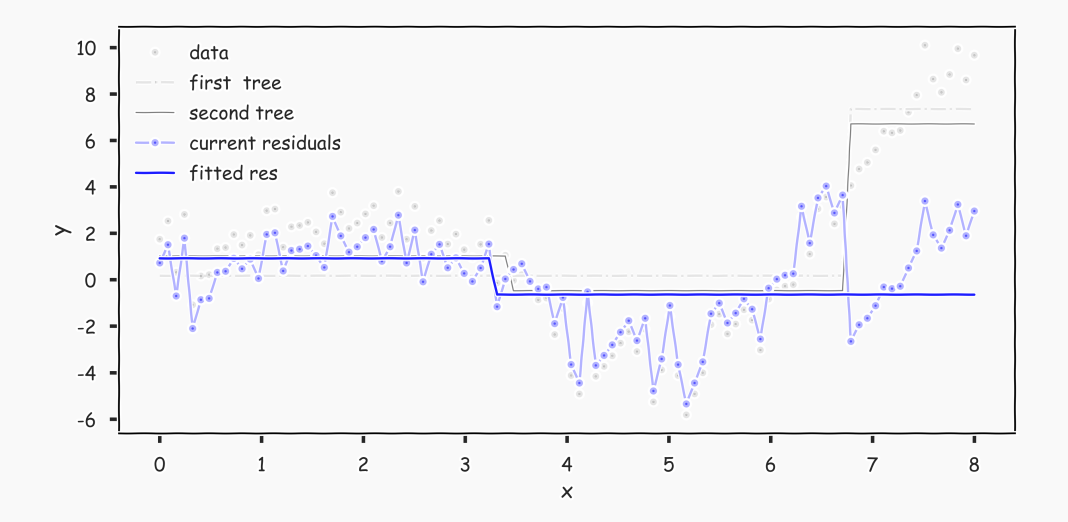




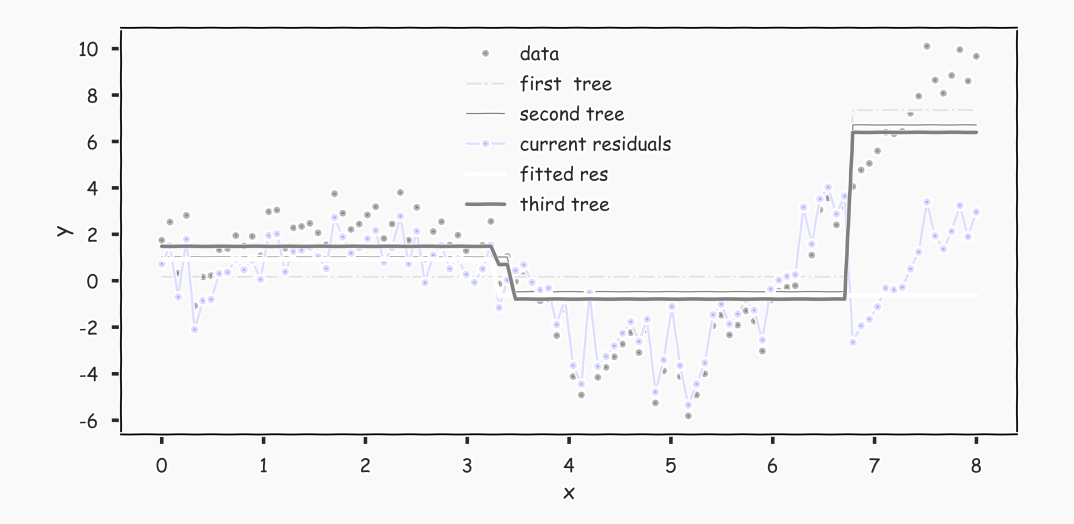




Gradient Boosting: illustration









Intuitively, each simple model *T*^(*i*) we add to our ensemble model *T*, models the errors of *T*.

Thus, with each addition of $T^{(i)}$, the residual is reduced

 $r_n - \lambda T^{(i)}(x_n)$

Note that gradient boosting has a tuning parameter, λ .

If we want to easily reason about how to choose λ and investigate the effect of λ on the model T, we need a bit more mathematical formalism. In particular, how can we effectively descend through this optimization via an iterative algorithm?

We need to formulate gradient boosting as a type of *gradient descent*.



In optimization, when we wish to minimize a function, called the **objective function**, over a set of variables, we compute the partial derivatives of this function with respect to the variables.

If the partial derivatives are sufficiently simple, one can analytically find a common root - i.e. a point at which all the partial derivatives vanish; this is called a **stationary point.**

If the objective function has the property of being **convex**, then the stationary point is precisely the min.



In practice, our objective functions are complicated and analytically find the stationary point is intractable.

Instead, we use an iterative method called *gradient descent* (as discussed in lecture 5):

1. Initialize the variables at any value:

$$x = [x_1, \dots, x_J]$$

2. Take the gradient of the objective function at the current variable values:

$$\nabla f(x) = \left[\frac{\partial f}{\partial x_1}(x), \dots, \frac{\partial f}{\partial x_J}(x)\right]$$

3. Adjust the variables values by some negative multiple of the gradient:

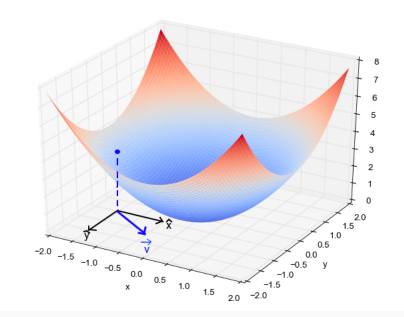
$$x \leftarrow x - \lambda \nabla f(x)$$

The factor λ is often called the learning rate.



Claim: If the function is convex, this iterative methods will eventually move x close enough to the minimum, for an appropriate choice of λ .

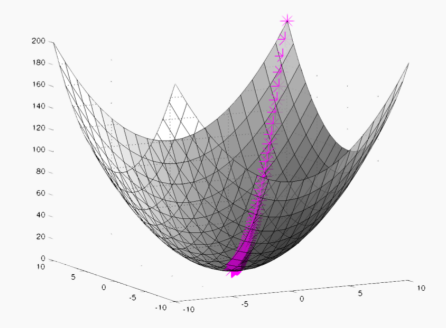
Why does this work? Recall, that as a vector, the gradient at at point gives the direction for the greatest possible rate of increase.





Subtracting a λ multiple of the gradient from x, moves x in the **opposite** direction of the gradient (hence towards the steepest decline) by a step of size λ .

If *f* is convex, and we keep taking steps descending on the graph of *f*, we will eventually reach the minimum.





Often in regression, our objective is to minimize the MSE

$$MSE(\hat{y}_1, \dots, \hat{y}_N) = \frac{1}{N} \sum_{i=1}^{N} (y_i - \hat{y}_i)^2$$

Treating this as an optimization problem, we can try to directly minimize the MSE with respect to the predictions

$$\nabla \mathsf{MSE} = \begin{bmatrix} \frac{\partial \mathsf{MSE}}{\partial \hat{y}_1}, \dots, \frac{\partial \mathsf{MSE}}{\partial \hat{y}_N} \end{bmatrix}$$
$$= -2 \left[y_1 - \hat{y}_1, \dots, y_N - \hat{y}_N \right]$$
$$= -2 \left[r_1, \dots, r_N \right]$$

The update step for gradient descent would look like

$$\hat{y}_n \leftarrow \hat{y}_n + \lambda r_n, \quad n = 1, \dots, N$$



There are two reasons why minimizing the MSE with respect to \hat{y}_n 's is not interesting:

- We know where the minimum MSE occurs: $\hat{y}_n = y_n$, for every *n*.
- Learning sequences of predictions, ŷ_n¹, ..., ŷ_nⁱ, ..., does not produce a model. The predictions in the sequences do not depend on the predictors!



The solution is to change the update step in gradient descent. Instead of using the gradient - the residuals - we use an **approximation** of the gradient that depends on the predictors:

$$\hat{y} \leftarrow \hat{y}_n + \lambda \, \hat{r}_n(x_n), \qquad n = 1, \dots, N$$

In gradient boosting, we use a simple model to approximate the residuals, $\hat{r}_n(x_n)$, in each iteration.

Motto: gradient boosting is a form of gradient descent with the MSE as the objective function.

Technical note: note that gradient boosting is descending in a space of models or functions relating x_n to y_n !



Gradient Boosting as Gradient Descent (cont.)

But why do we care that gradient boosting is gradient descent?

By making this connection, we can import the massive amount of techniques for studying gradient descent to analyze gradient boosting.

For example, we can easily reason about how to choose the learning rate λ in gradient boosting.



Under ideal conditions, gradient descent iteratively approximates and converges to the optimum.

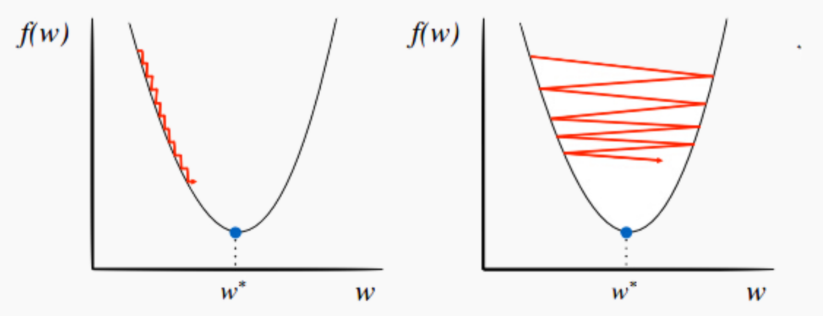
When do we terminate gradient descent?

- We can limit the number of iterations in the descent. But for an arbitrary choice of maximum iterations, we cannot guarantee that we are sufficiently close to the optimum in the end.
- If the descent is stopped when the updates are sufficiently small (e.g. the residuals of T are small), we encounter a new problem: the algorithm may never terminate!

Both problems have to do with the magnitude of the learning rate, λ .



For a constant learning rate, λ , if λ is too small, it takes too many iterations to reach the optimum.



If λ is too large, the algorithm may 'bounce' around the optimum and never get sufficiently close.



Choosing λ :

- If λ is a constant, then it should be tuned through cross validation.
- For better results, use a variable λ. That is, let the value of λ depend on the gradient

$$\lambda = h(\|\nabla f(x)\|),$$

where $\|\nabla f(x)\|$ is the magnitude of the gradient, $\nabla f(x)$. So

- around the optimum, when the gradient is small, $\boldsymbol{\lambda}$ should be small
- far from the optimum, when the gradient is large, λ should be larger



PPppBoost

AdaBoost



Using the language of gradient descent also allow us to connect gradient boosting for regression to a boosting algorithm often used for classification, AdaBoost.

In classification, we typically want to minimize the classification error:

$$\mathsf{Error} = \frac{1}{N} \sum_{n=1}^{N} \mathbb{1}(y_n \neq \hat{y}_n), \quad \mathbb{1}(y_n \neq \hat{y}_n) = \begin{cases} 0, & y_n = \hat{y}_n \\ 1, & y_n \neq \hat{y}_n \end{cases}$$

Naively, we can try to minimize Error via gradient descent, just like we did for MSE in gradient boosting.

Unfortunately, Error is not differentiable 🛞



Our solution: we replace the Error function with a differentiable function that is a good indicator of classification error.

The function we choose is called **exponential loss**

ExpLoss =
$$\frac{1}{N} \sum_{n=1}^{N} \exp(-y_n \hat{y}_n), y_n \in \{-1, 1\}$$

Exponential loss is differentiable with respect to \hat{y}_n and it is an upper bound of Error.



We first compute the gradient for ExpLoss:

$$\nabla_{\hat{y}} \operatorname{Exp} = \left[-y_1 \exp\left(-y_1 \hat{y}_1\right), \dots, -y_N \exp\left(-y_N \hat{y}_N\right)\right]$$

It's easier to decompose each $y_n \exp(-y_n \hat{y}_n)$ as $w_n y_n$, where $w_n = \exp(-y_n \hat{y}_n)$.

This way, we see that the gradient is just a re-weighting applied the target values

$$abla \mathsf{Exp} = [-w_1 y_1, \dots, -w_N y_N]$$

Notice that when $y_n = \hat{y}_n$, the weight w_n is small; when $y_n \neq \hat{y}_n$, the weight is larger.



The update step in the gradient descent is

$$\hat{y}_n \leftarrow \hat{y}_n + \lambda w_n y_n, \qquad n = 1, \dots, N$$

Just like in gradient boosting, we approximate the gradient, $\lambda w_n y_n$ with a simple model, $T^{(i)}$, that depends on x_n .

This means training *T*^(*i*) on a re-weighted set of target values,

 $\{(x_1, w_1y_1), \dots, (x_N, w_Ny_N)\}$

That is, gradient descent with exponential loss means iteratively training simple models that **focuses on the points misclassified by the previous model.**



With a minor adjustment to the exponential loss function, we have the algorithm for gradient descent:

- 1. Choose an initial distribution over the training data, $w_n = 1/N$.
- 2. At the *i*th step, fit a simple classifier $T^{(i)}$ on weighted training data $\{(x_1, w_1y_1), \dots, (x_N, w_Ny_N)\}$
- 3. Update the weights:

$$w_n \leftarrow \frac{w_n \exp(-\lambda^{(i)} y_n T^{(i)}(x_n))}{Z}$$

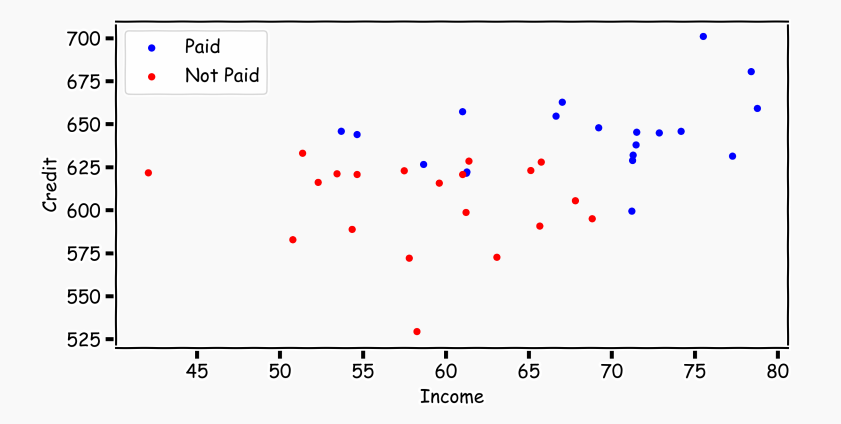
where Z is the normalizing constant for the collection of updated weights

4. Update $T: T \leftarrow T + \lambda^{(i)}T^{(i)}$

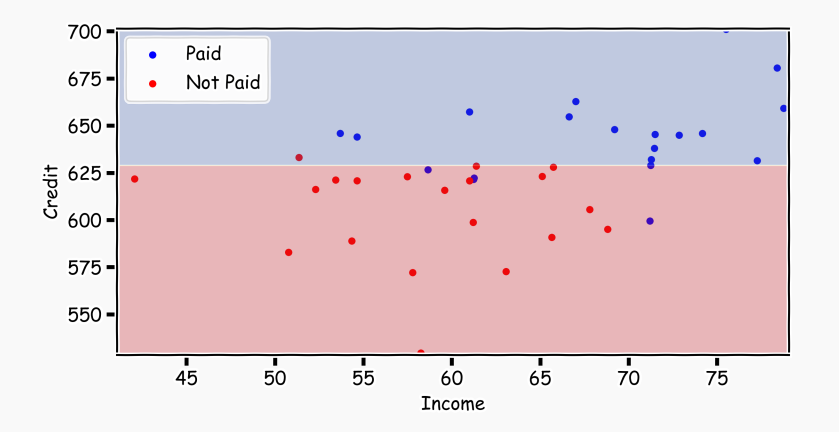
where λ is the learning rate.



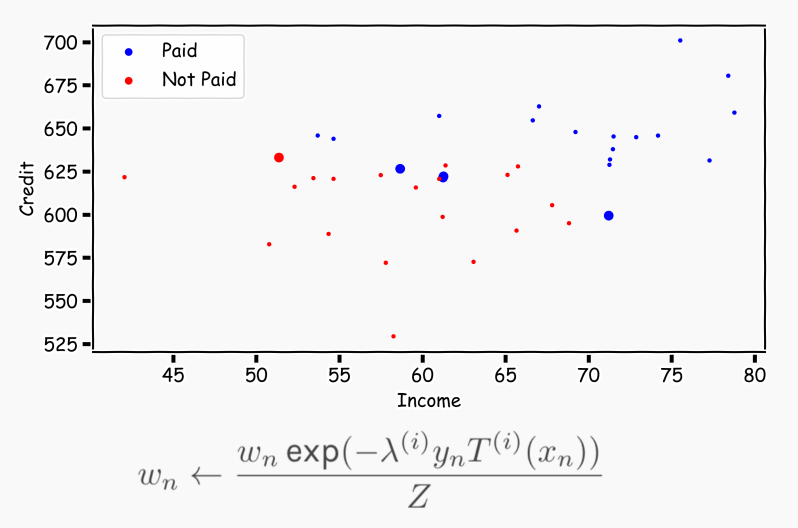
AdaBoost: start with equal weights







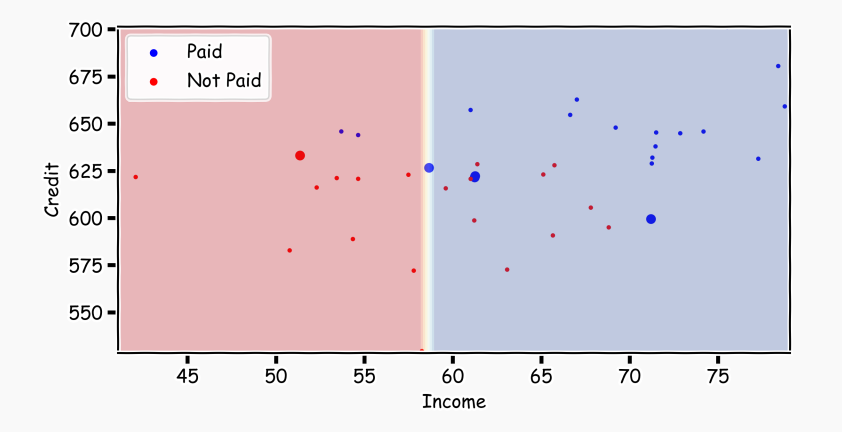






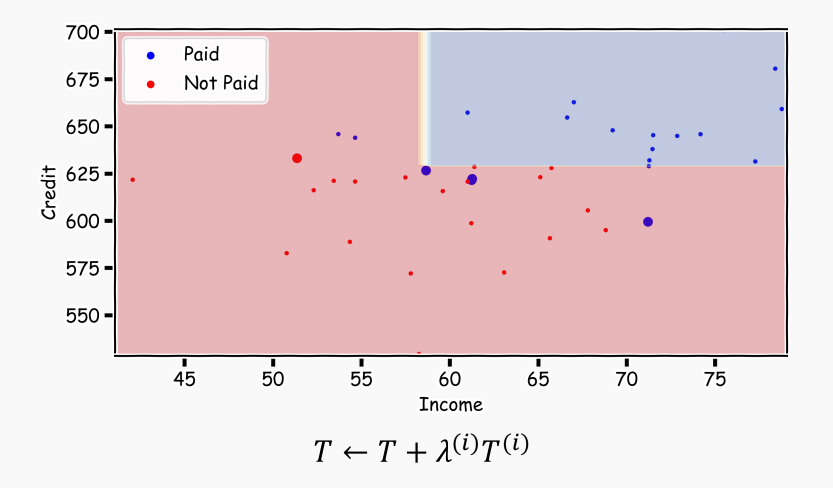
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AdaBoost: fit another simple decision tree on re-weighted data

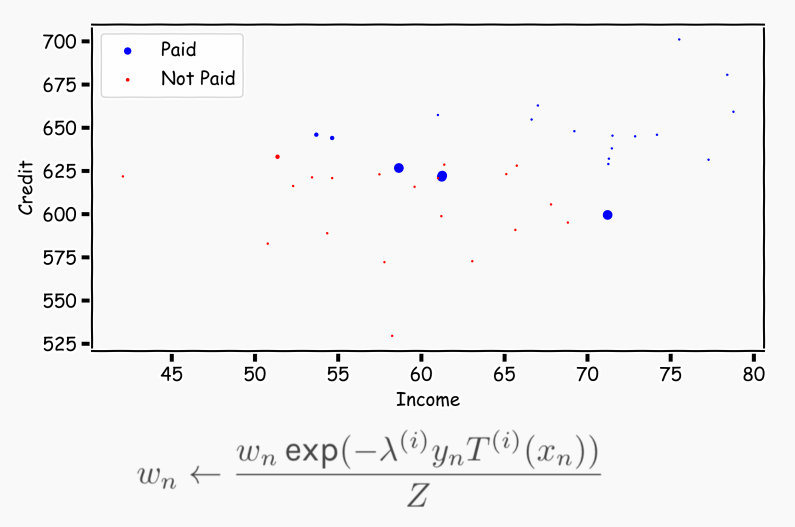




AdaBoost: add the new model to the ensemble



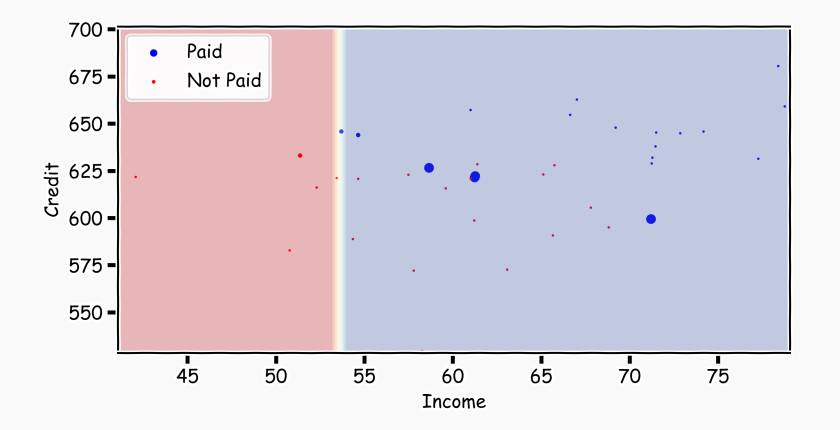






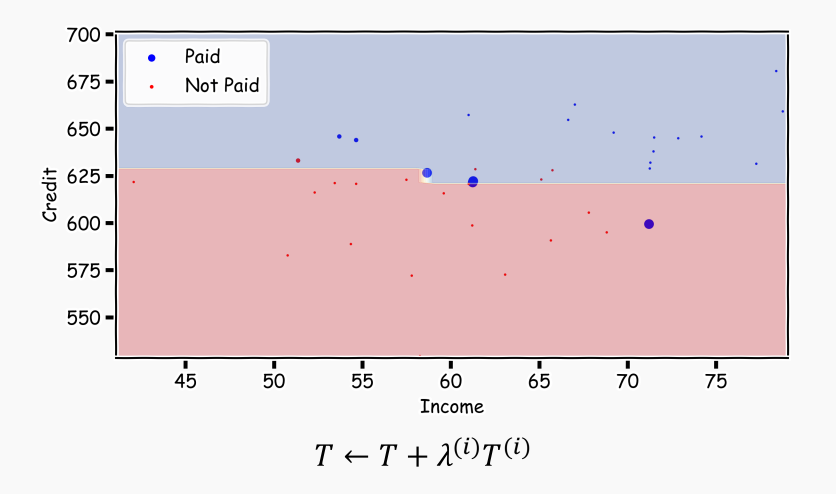
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AdaBoost: fit a third, simple decision tree on re-weighted data





AdaBoost: add the new model to the ensemble, repeat...





Unlike in the case of gradient boosting for regression, we can analytically solve for the optimal learning rate for AdaBoost, by optimizing:

$$\underset{\lambda}{\operatorname{argmin}} \frac{1}{N} \sum_{n=1}^{N} \exp\left[-y_n(T+\lambda^{(i)}T^{(i)}(x_n))\right]$$

Doing so, we get that

$$\lambda^{(i)} = \frac{1}{2} \ln \frac{1-\epsilon}{\epsilon}, \quad \epsilon = \sum_{n=1}^{N} w_n \mathbb{1}(y_n \neq T^{(i)}(x_n))$$



There are few implementations on boosting:

- XGBoost: An efficient Gradient Boosting Decision
- LGBM: Light Gradient Boosted Machines. It is a library for training GBMs developed by Microsoft, and it competes with XGBoost
- CatBoost: A new library for Gradient Boosting Decision Trees, offering appropriate handling of categorical features

ADVANCED TOPICS



Final thoughts on Boosting

Increasing the number of trees can lead to overfitting.

Question: Why?

