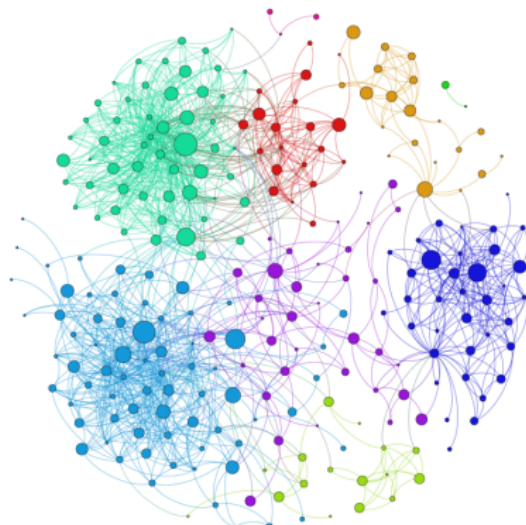


Decision Trees and Ensemble Methods

Jiaming Mao

Xiamen University



Copyright © 2017–2019, by Jiaming Mao

This version: Spring 2019

Contact: jmao@xmu.edu.cn

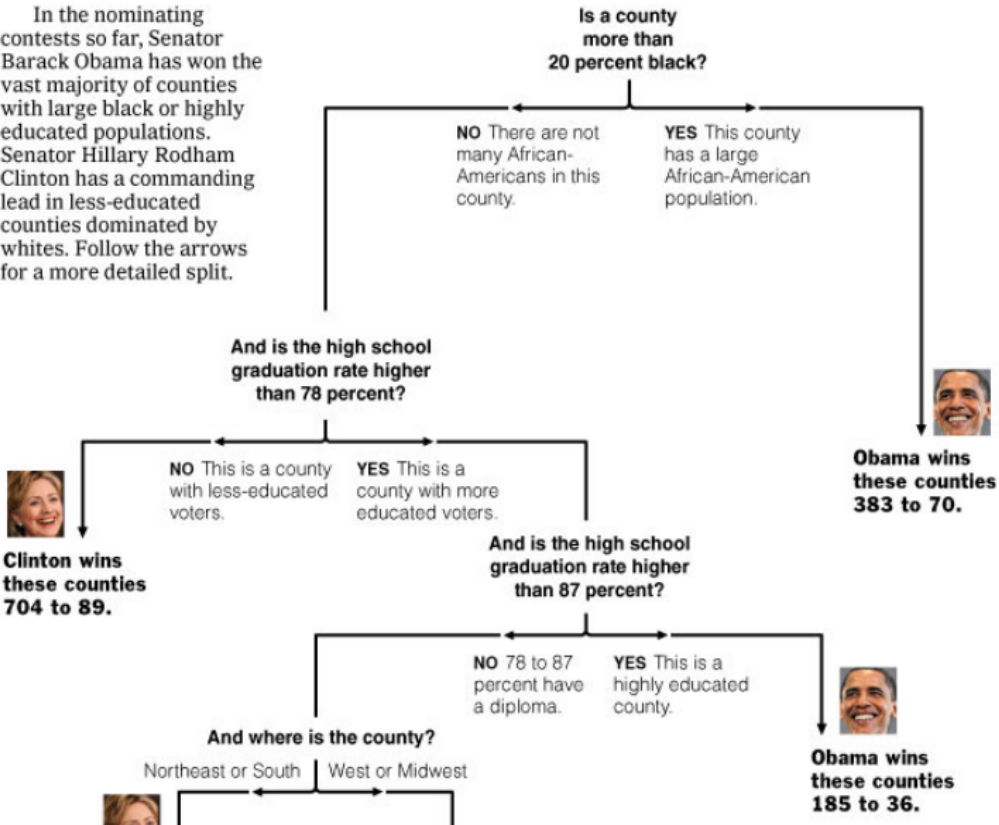
Course homepage: jiamingmao.github.io/data-analysis



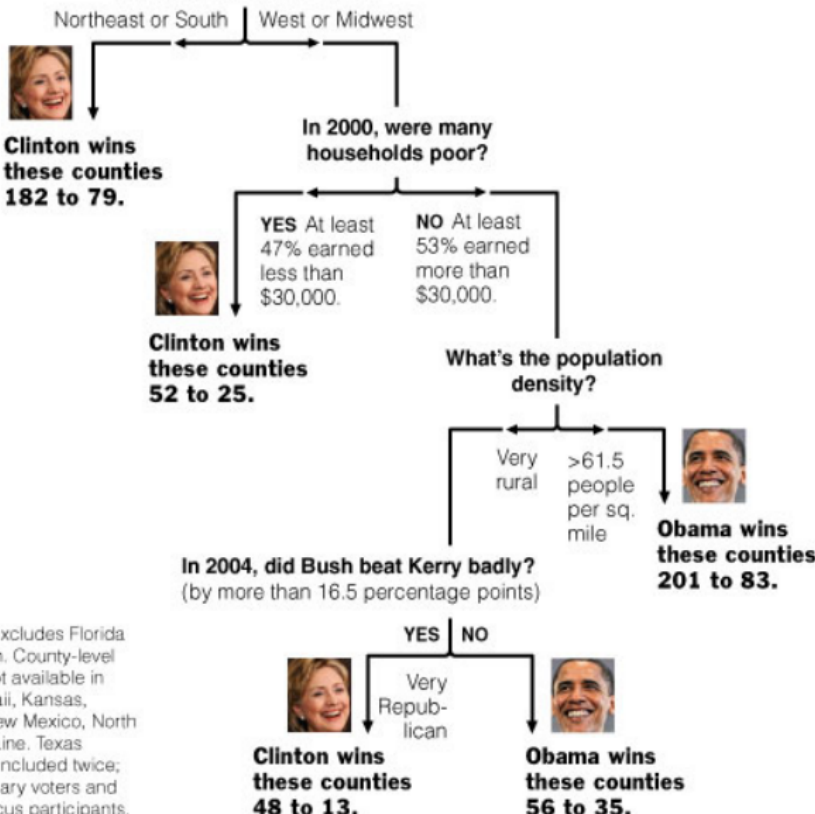
All materials are licensed under the [Creative Commons Attribution-NonCommercial 4.0 International License](#).

Decision Tree: The Obama-Clinton Divide

In the nominating contests so far, Senator Barack Obama has won the vast majority of counties with large black or highly educated populations. Senator Hillary Rodham Clinton has a commanding lead in less-educated counties dominated by whites. Follow the arrows for a more detailed split.



And where is the county?



Note. Chart excludes Florida and Michigan. County-level results are not available in Alaska, Hawaii, Kansas, Nebraska, New Mexico, North Dakota or Maine. Texas counties are included twice; once for primary voters and once for caucus participants.

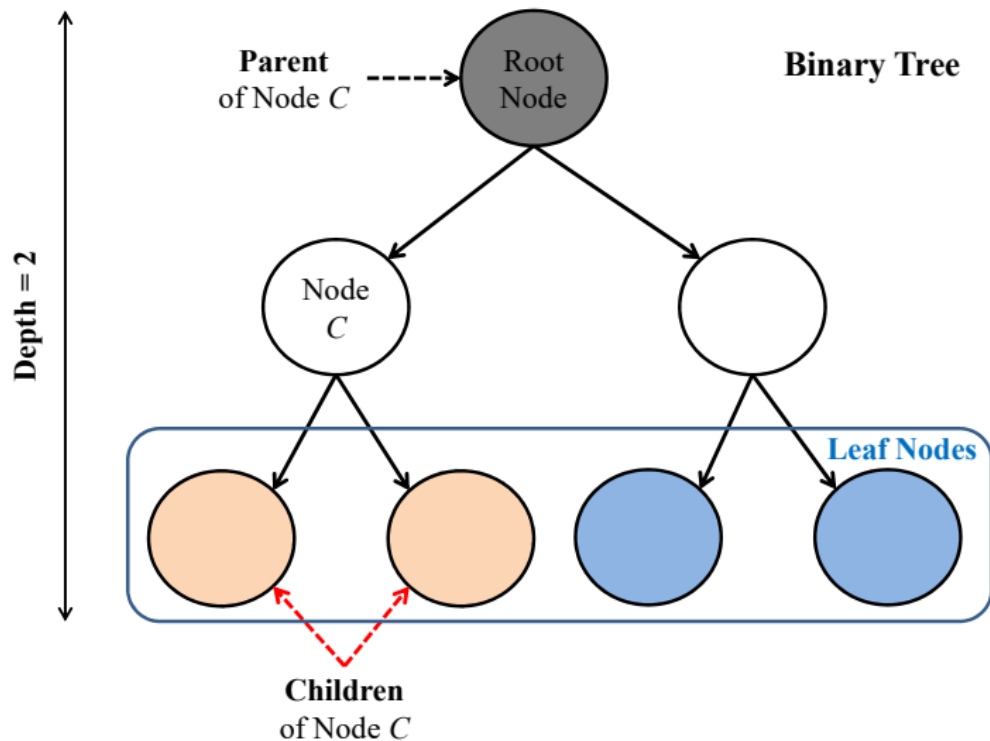
Sources: Election results via The Associated Press; Census Bureau; Dave Leip's Atlas of U.S. Presidential Elections

AMANDA COX/
THE NEW YORK TIMES

Decision Trees

- Tree-based methods divide the predictor space into different regions, and then fit a model in each region.
- The predictor space can be divided by making successive binary splits on the predictor variables x_1, \dots, x_p : we choose a variable x_j , $j = 1, \dots, p$, divide up the predictor space according to $x_j \leq c$ and $x_j > c$, and then proceed on each half.
- A decision tree thus constructed consists of **internal nodes** and **terminal nodes (leaves)**. Each internal node is a decision to split the predictor space, and the leaves are final predictions. The **size** of a tree is the number of its leaves.
- Decision trees can be applied to both regression and classification problems.

Decision Trees



Regression Tree

Let $\mathcal{D} = \{(x_i, y_i)\}_{i=1}^N$ be the training data. A regression tree of M leaves can be thought of as dividing the predictor space into M regions R_1, \dots, R_M , each corresponding to a leaf of the tree.

For each R_m , $m = 1, \dots, M$, let

$$\bar{y}_m = \frac{1}{n_m} \sum_{x_i \in R_m} y_i \quad (1)$$

, where n_m is the number of observations in R_m .

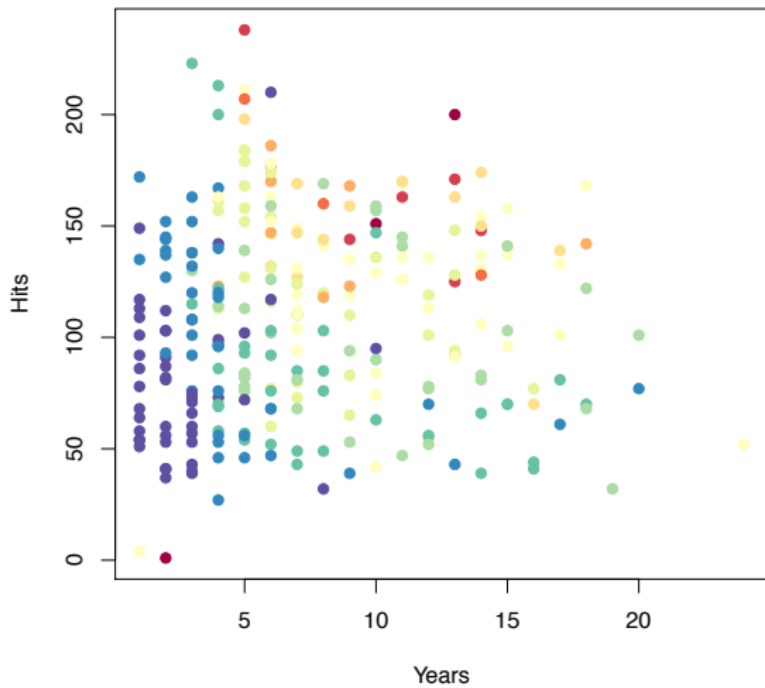
Regression Tree

The estimated regression function is:

$$\hat{f}(x) = \sum_{m=1}^M \bar{y}_m \cdot \mathcal{I}\{x \in R_m\} \quad (2)$$

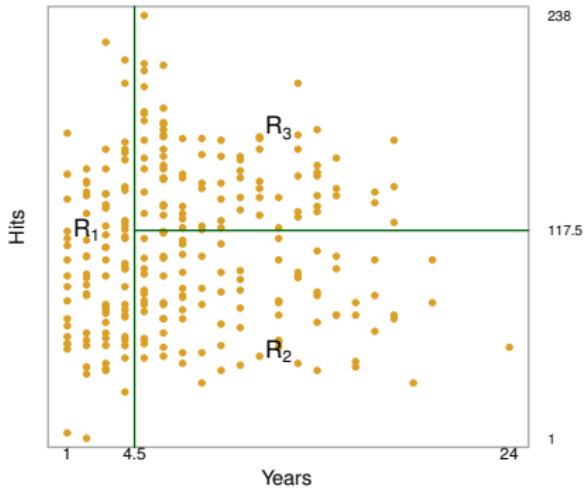
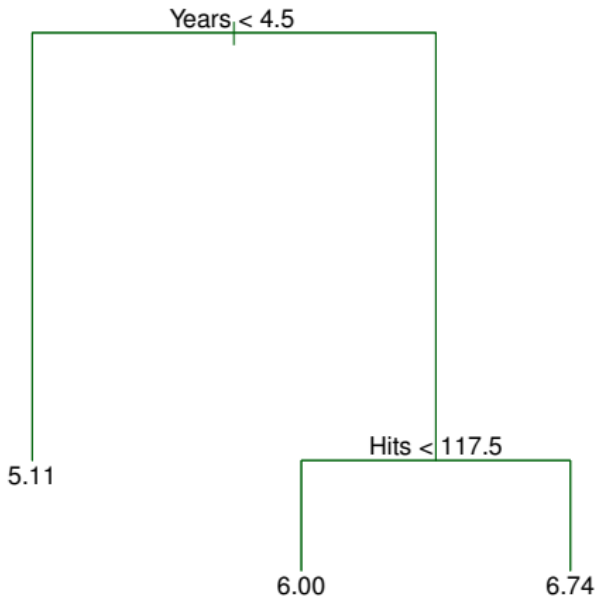
- For every observation that falls into the region R_m , we make the same prediction, which is simply the mean of the response values for the training observations in R_m .
- The result is a **piecewise constant model**.

Baseball Salary



Color-coded from low (blue, green) to high (yellow, red)

Baseball Salary



The results suggest that years is the most important factor in determining salary. Among players who have played for more than 4.5 years, the number of hits made (in the previous year) affects salary.

Building a Regression Tree

- The goal is to find R_1, \dots, R_M that minimize E_{in} ¹. In the regression setting, using the L2 loss, this means minimizing

$$\text{RSS} = \sum_{m=1}^M \sum_{x_i \in R_m} (y_i - \bar{y}_m)^2$$

- Unfortunately, it is computationally infeasible to consider every possible partition of the predictor space.
- Therefore, we adopt a **forward stepwise** procedure ...

¹Here we minimize E_{in} first and regularize it later.

The CART Algorithm

The CART algorithm is a **recursive partitioning** algorithm that divides the predictor space by making successive **binary** splits. At each step, it chooses the split to achieve the biggest drop in E_{in} .

- The algorithm begins by dividing the predictor space into two regions. Then it divides one of the two regions² further into two regions ... The process continues until a stopping criterion is reached³.
- This is a **greedy** approach because at each step of the tree-building process, the optimal split is made at that particular step, rather than looking ahead and picking a split that will lead to a better tree in some future step.

²Depending on splitting which would result in the biggest drop in E_{in} .

³For instance, we may continue until no region contains more than 10 observations.

The CART Algorithm

Given a region $R \subseteq \mathbb{R}^p$, the optimal split is choosing an x_j and a split point s , so that the two sub-regions

$$R_{\text{left}} = \{x \in R : x_j < s\} \text{ and } R_{\text{right}} = \{x \in R : x_j \geq s\}$$

are as **homogenous** in response y as possible.

For regression trees, this means choosing (j, s) to minimize:

$$\sum_{x_i \in R_{\text{left}}} (y_i - \bar{y}_{\text{left}})^2 + \sum_{x_i \in R_{\text{right}}} (y_i - \bar{y}_{\text{right}})^2$$

The CART Algorithm

The process generates a large tree T_0 . We then **prune** the tree by choosing, for each value of $\alpha \geq 0$, a subtree $T \subset T_0$ that minimizes

$$\sum_{m=1}^{|T|} \sum_{x_i \in R_m} (y_i - \bar{y}_m)^2 + \alpha |T|$$

, where $|T|$ is the size of T and α is a **tuning** parameter that controls the tradeoff between the subtree's complexity and its fit to the training data.

- $\alpha = 0 \Rightarrow T = T_0$
- $\alpha \uparrow \Rightarrow |T| \downarrow$
- The optimal α can be selected using cross-validation.

The CART Algorithm

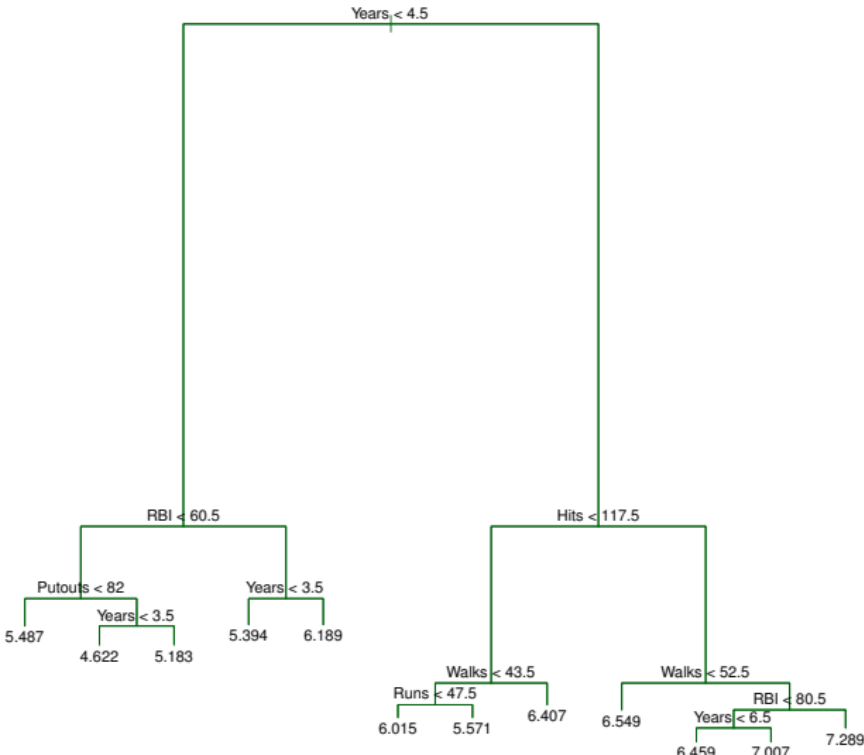
- Intuitively, a small tree might be too simple, while a large tree might overfit the data.
- The strategy of growing a large tree and then pruning it is used because any stopping rule may be short-sighted, in that a split may look bad but it may lead to a good split below it.

The CART Algorithm

Algorithm 8.1 Building a Regression Tree

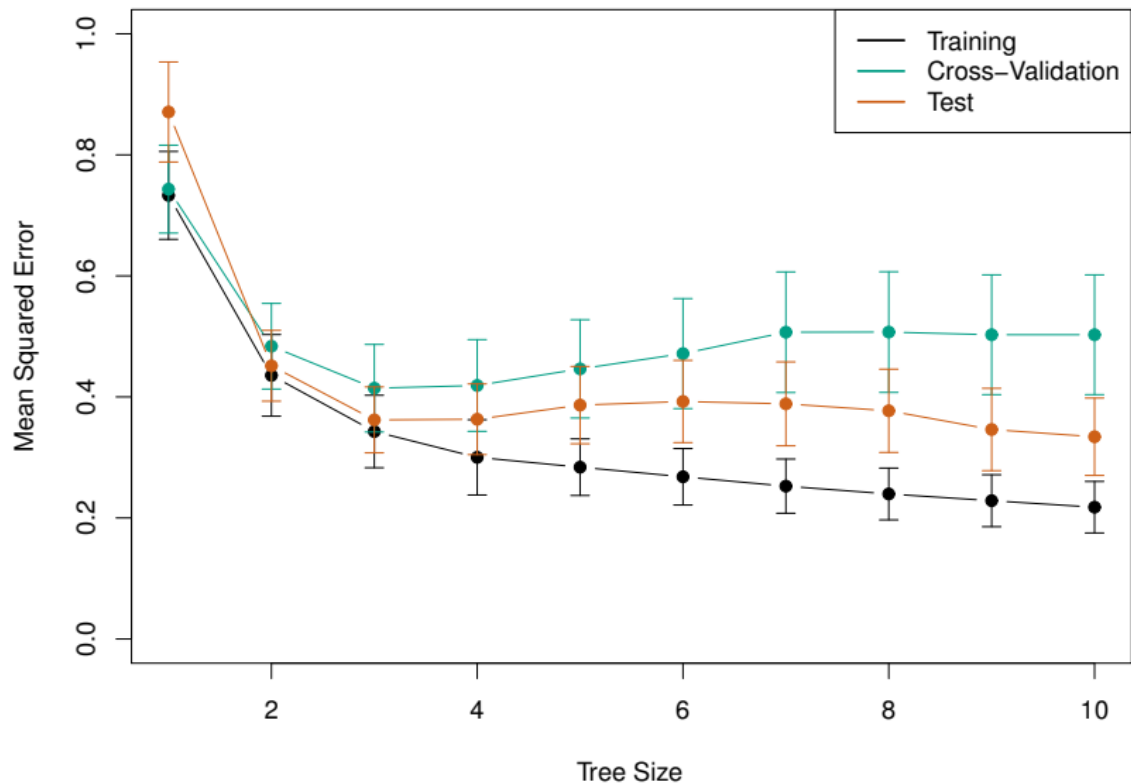
1. Use recursive binary splitting to grow a large tree on the training data, stopping only when each terminal node has fewer than some minimum number of observations.
 2. Apply cost complexity pruning to the large tree in order to obtain a sequence of best subtrees, as a function of α .
 3. Use K-fold cross-validation to choose α . That is, divide the training observations into K folds. For each $k = 1, \dots, K$:
 - (a) Repeat Steps 1 and 2 on all but the k th fold of the training data.
 - (b) Evaluate the mean squared prediction error on the data in the left-out k th fold, as a function of α .Average the results for each value of α , and pick α to minimize the average error.
 4. Return the subtree from Step 2 that corresponds to the chosen value of α .
-

Baseball Salary



The unpruned tree

Baseball Salary



Classification Tree

For classification trees, the leaves are class predictions.

Let $\mathcal{D} = \{(x_i, y_i)\}_{i=1}^N$ be the training data, where $y \in \{1, \dots, J\}$. For each $R_m, m = 1, \dots, M$, we assign a class label c_m .

We then classify a new point $x \in \mathbb{R}^p$ by

$$\hat{f}(x) = \sum_{m=1}^M c_m \cdot \mathcal{I}\{x \in R_m\} \quad (3)$$

Classification Tree

Let $p_j^m \equiv \Pr(y = j | x \in R_m)$ be the class probabilities in R_m . p_j^m can be estimated by

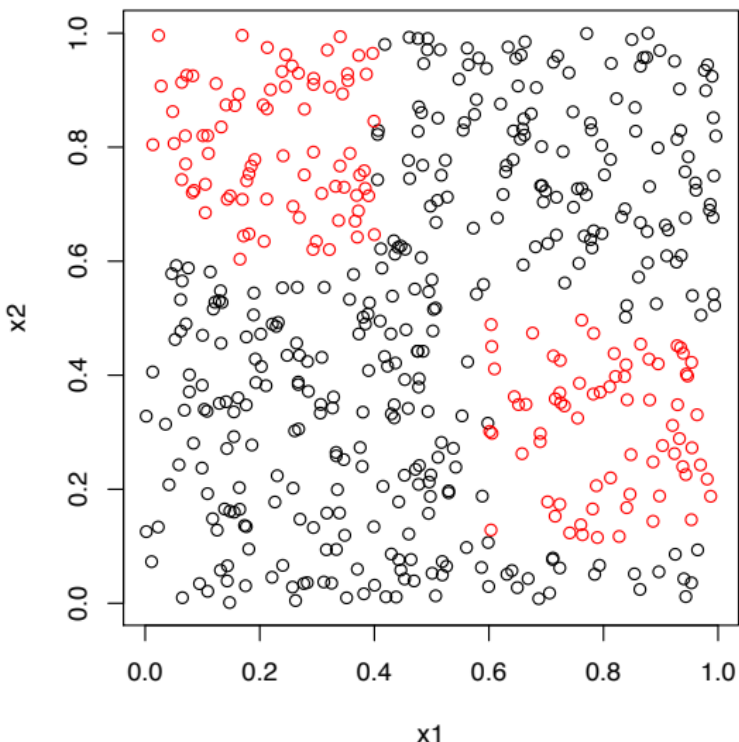
$$\hat{p}_j^m = \frac{1}{n_m} \sum_{x_i \in R_m} \mathcal{I}\{y_i = j\} \quad (4)$$

Then

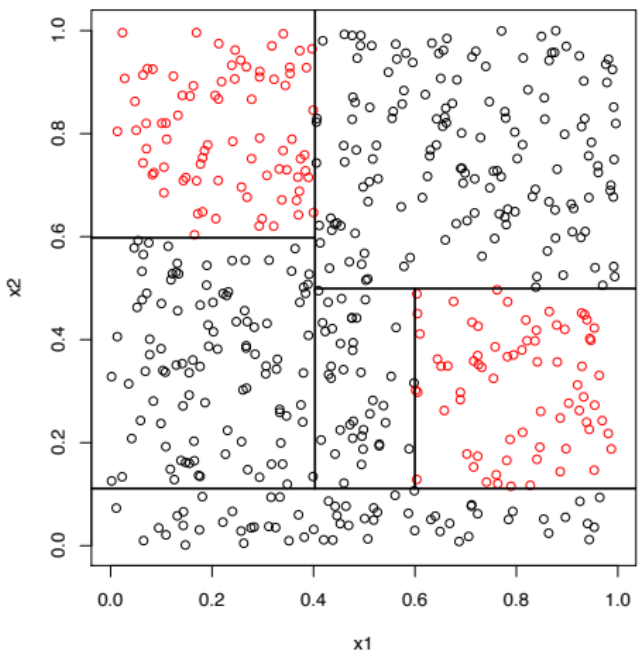
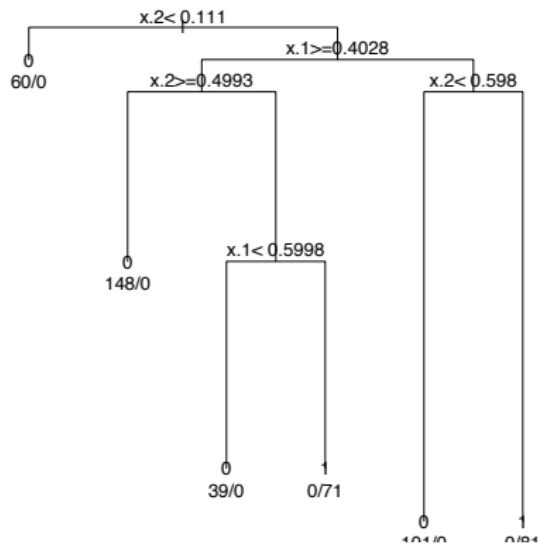
$$c_m = \arg \max_j \hat{p}_j^m \quad (5)$$

, i.e., c_m is the most common occurring class in R_m .

Classification Tree



Classification Tree



Building a Classification Tree

As in building regression trees, we first find R_1, \dots, R_M that minimize E_{in} . Let $Q_m = E_{in}(R_m)$. Then at each step, the CART algorithm chooses the optimal split to minimize:

$$n_{\text{left}} Q_{\text{left}} + n_{\text{right}} Q_{\text{right}}$$

For classification trees, Q_m is also called the **node impurity** measure.

Node Impurity Measures

Commonly used node impurity measures are:

Misclassification rate:

$$Q_m = 1 - \hat{p}_{c_m}^m$$

, where $\hat{p}_{c_m}^m$ is the proportion of observations in R_m that belong to class c_m .

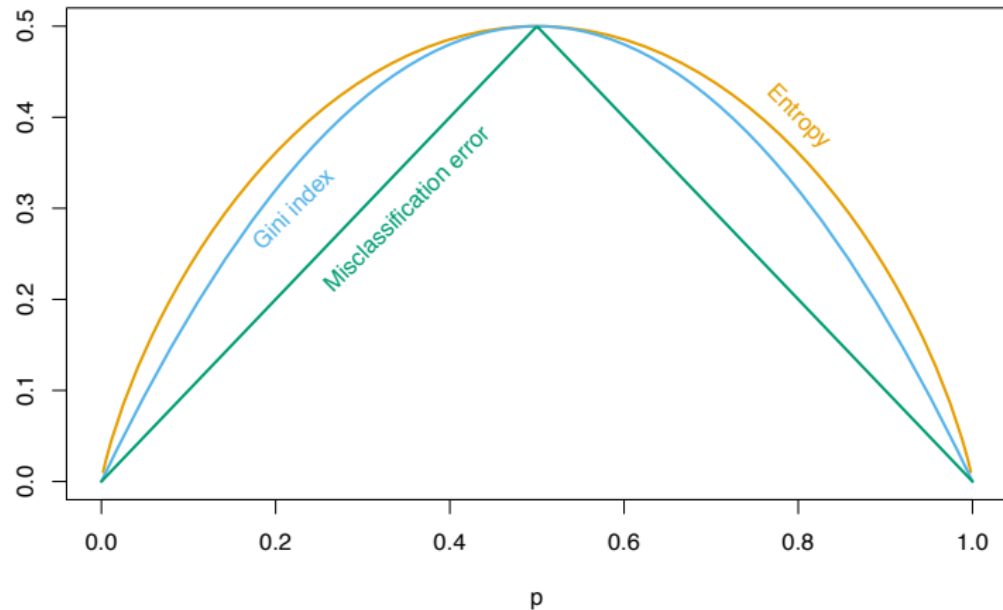
Gini index:

$$Q_m = \sum_{j=1}^J \hat{p}_j^m (1 - \hat{p}_j^m)$$

Cross-entropy:

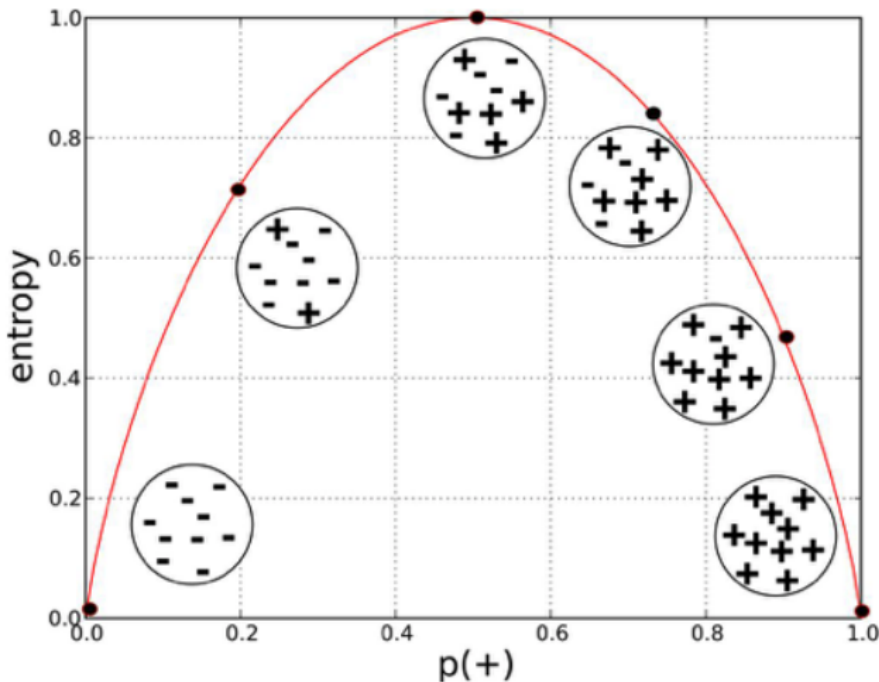
$$Q_m = - \sum_{j=1}^J \hat{p}_j^m \log \hat{p}_j^m$$

Node Impurity Measures



Node impurity measures for binary classification (cross-entropy has been scaled to pass through $(0.5, 0.5)$). Note that the Gini index and cross-entropy are differentiable and thus easier to optimize numerically.

Node Impurity Measures



Mode of Transportation

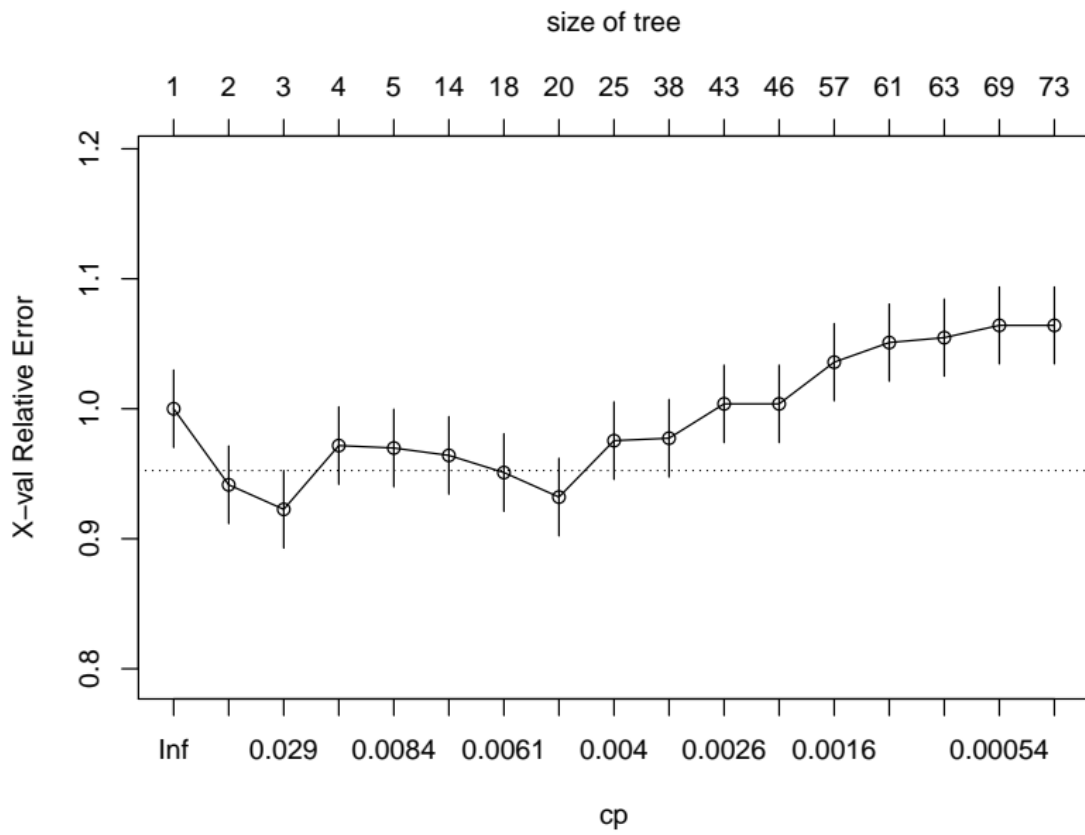
```
transport = read.csv("Transport.txt")
names(transport) = c("loginc", "distance", "mode")

#####
# Classification Tree #
#####

require(rpart)
# step 1: grow a tree until stopping criteria:
#          cp: cost complexity parameter; any split must lower in-sample
#                  error by a factor of cp in order to continue grow
fit0 = rpart(mode~., transport, control=rpart.control(cp=0))

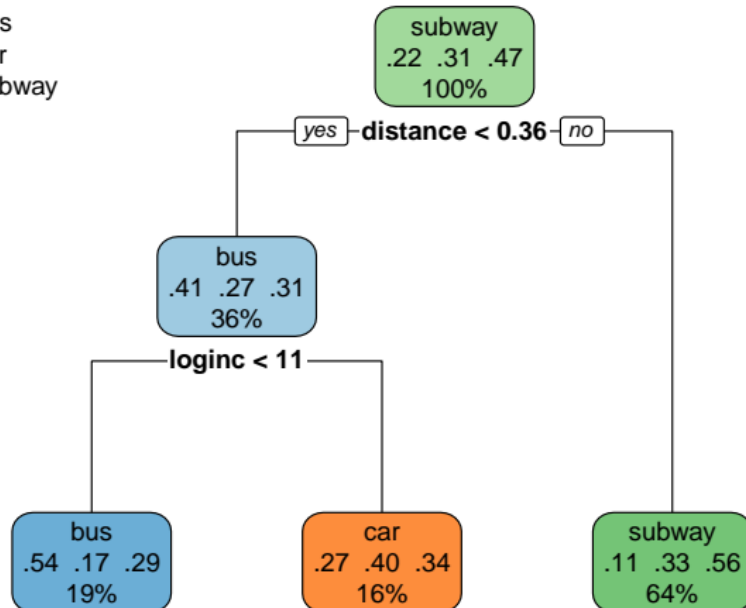
# step 2: prune the tree
# select the cost-complexity parameter ("cp") with the smallest
# (relative) cross-validation error ("xerror")
fit = prune(fit0, cp=fit0$cptable[which.min(fit0$cptable[, "xerror"]),"CP"])
```

Mode of Transportation

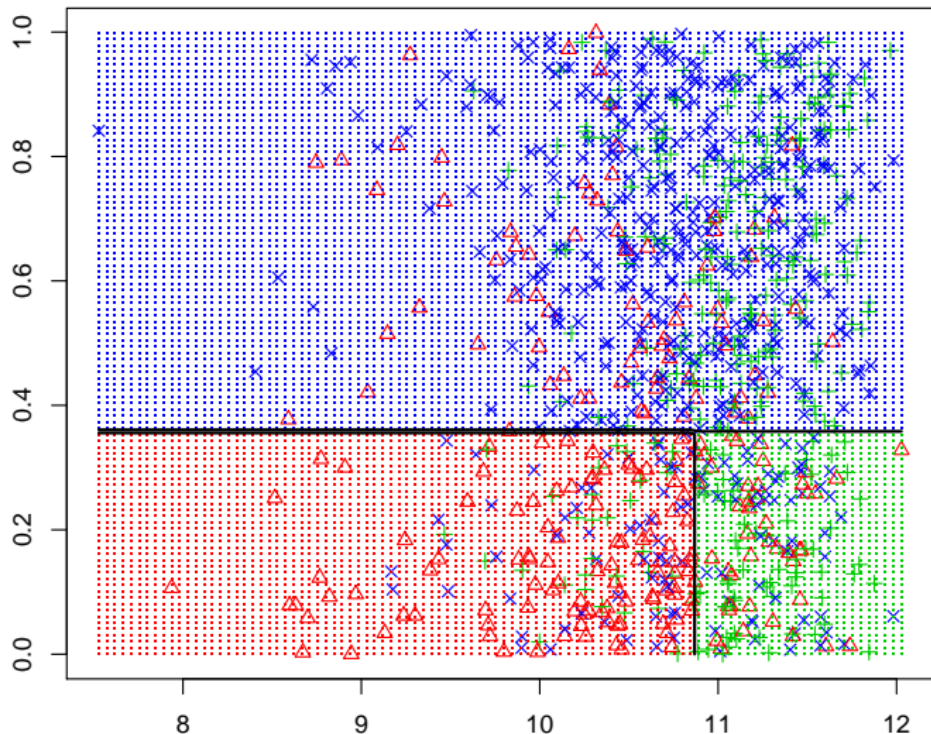


Mode of Transportation

bus
car
subway



Mode of Transportation



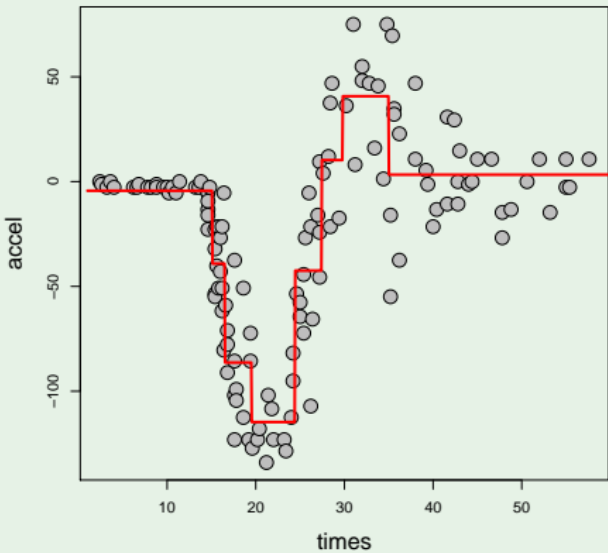
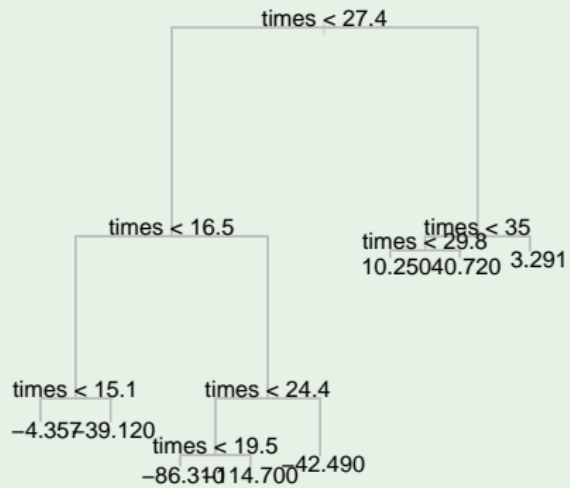
Advantages and Disadvantages of Trees

Pros:

- Highly interpretable
- Automatically detecting nonlinear relationships
- Automatically modeling interactions

Advantages and Disadvantages of Trees

Motorcycle crash test dummy data



x : time from impact; y : head acceleration

Advantages and Disadvantages of Trees

Cons:

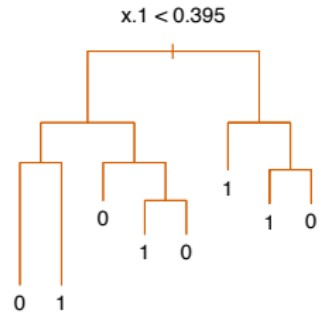
- Relatively poor predictive performance
 - ▶ Trees generally suffer from high variance and are **unstable**: a small change in the observed data often results in a completely different tree.
 - ▶ This is due to their hierarchical nature: once a split is made, all the splits under it are changed as well.
- Difficulty in capturing simple relationships
 - ▶ Trees require a large number of parameters (splits) to capture simple linear and additive relationships.
 - ▶ In other words, the weaknesses of tree-based methods are the strengths of linear models, and vice versa.

Bagging

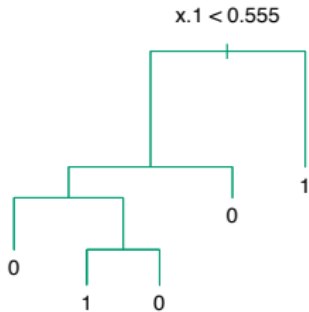
- **Bootstrap aggregation**, or **bagging**, is a general-purpose procedure for reducing the variance of a statistical learning method.
- We know that given a set of N independent random variables each with variance σ^2 , the variance of the mean is $\frac{\sigma^2}{N}$.
- Hence if we can average the predictions made on many independent training data sets drawn from the population, then we can reduce the variance and hence increase the prediction accuracy of our method.
- In practice, we do not have access to multiple independent training data sets from the population. Instead, we can draw multiple samples from the **empirical distribution** using bootstrap.

Bagging

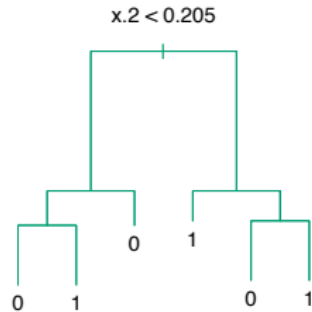
Original Tree



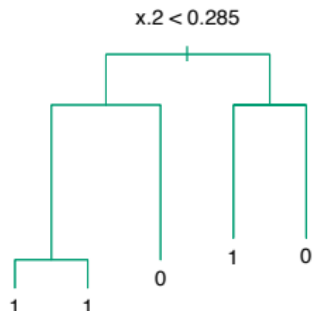
$b = 1$



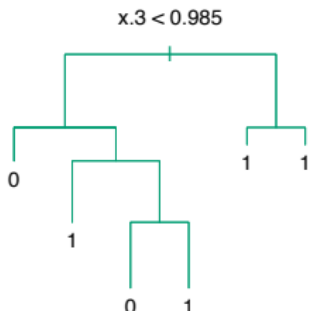
$b = 2$



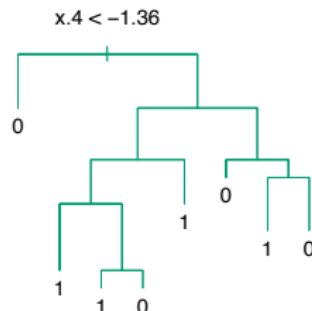
$b = 3$



$b = 4$



$b = 5$



Bagging for Regression

Let $b = 1, \dots, B$ index the bootstrap samples drawn from the training data. We fit our model on each bootstrap sample.

For regression problems, let $\hat{f}^{(b)}(x)$ be the model's prediction trained on sample b . We then average all the predictions to obtain

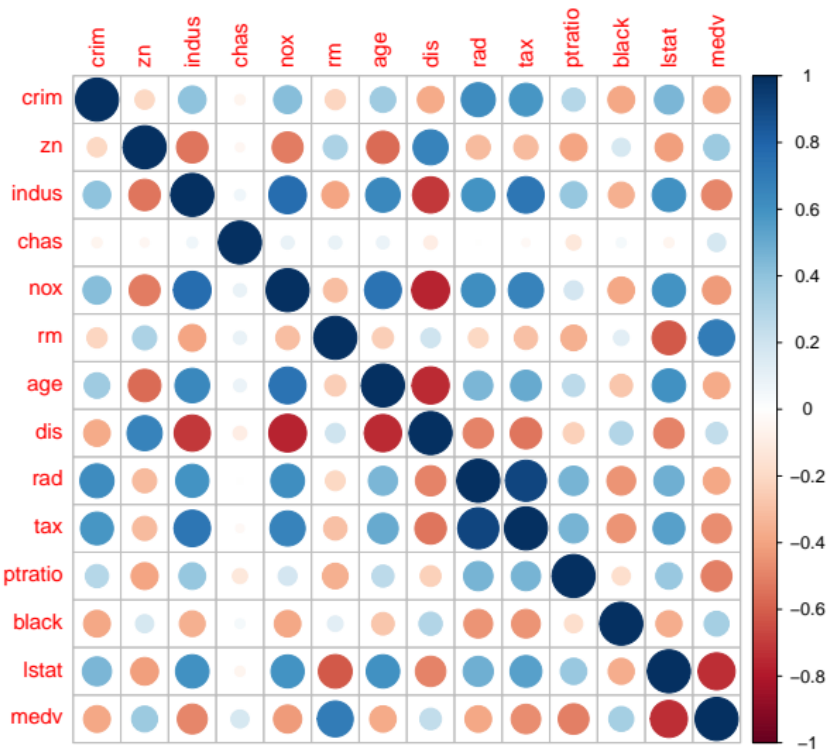
$$\hat{f}(x) = \frac{1}{B} \sum_{b=1}^B \hat{f}^{(b)}(x) \quad (6)$$

Boston Housing Price

Data: median value of owner-occupied homes in 506 census tracts in the Boston area from the 1970 census. Other variables include average rooms per house, distance to employment centers, crime rate, percentage of population with lower socioeconomic status, etc.

```
require(MASS) # contains the data set 'Boston'  
require(rpart)  
head(Boston,3)  
  
##      crim zn indus chas   nox     rm    age     dis rad tax ptratio  black  
## 1 0.00632 18  2.31     0 0.538 6.575 65.2 4.0900    1 296 15.3 396.90  
## 2 0.02731  0  7.07     0 0.469 6.421 78.9 4.9671    2 242 17.8 396.90  
## 3 0.02729  0  7.07     0 0.469 7.185 61.1 4.9671    2 242 17.8 392.83  
##      lstat medv  
## 1 4.98 24.0  
## 2 9.14 21.6  
## 3 4.03 34.7
```

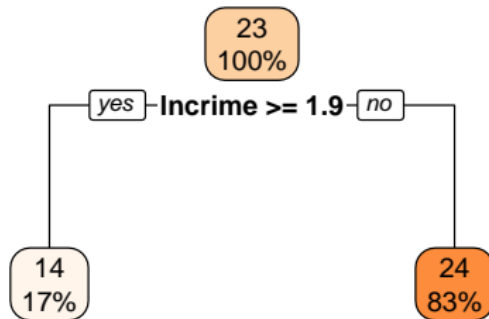
Boston Housing Price



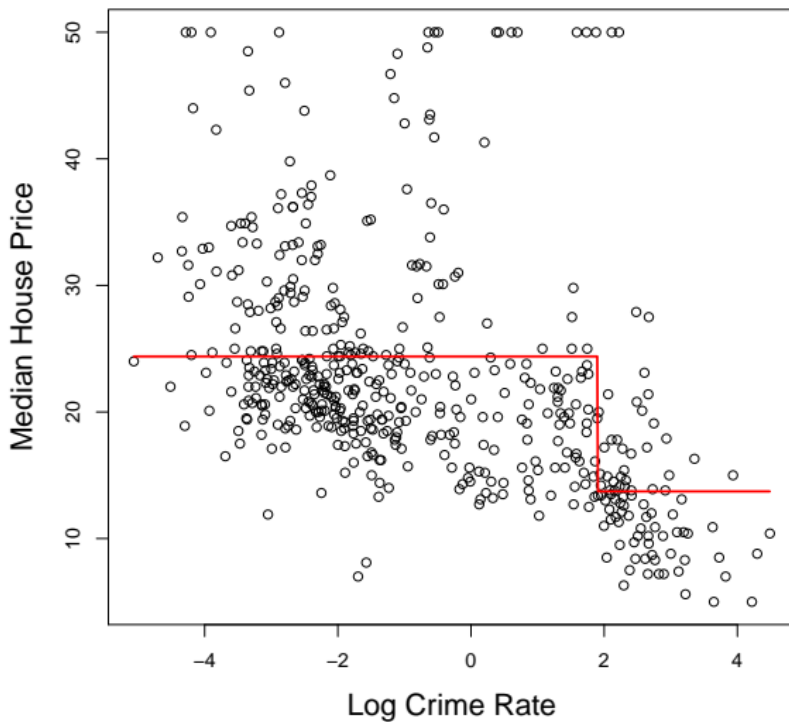
Boston Housing Price

Let's first predict median house prices based on crime rates only, using a single **decision stump** – the simplest tree with a single split at root.

```
# Fitting a single strump
n = nrow(Boston)
Boston$lncrime = log(Boston$crim)
fit = rpart(medv~lncrime,Boston,control=rpart.control(maxdepth=1))
```



Boston Housing Price

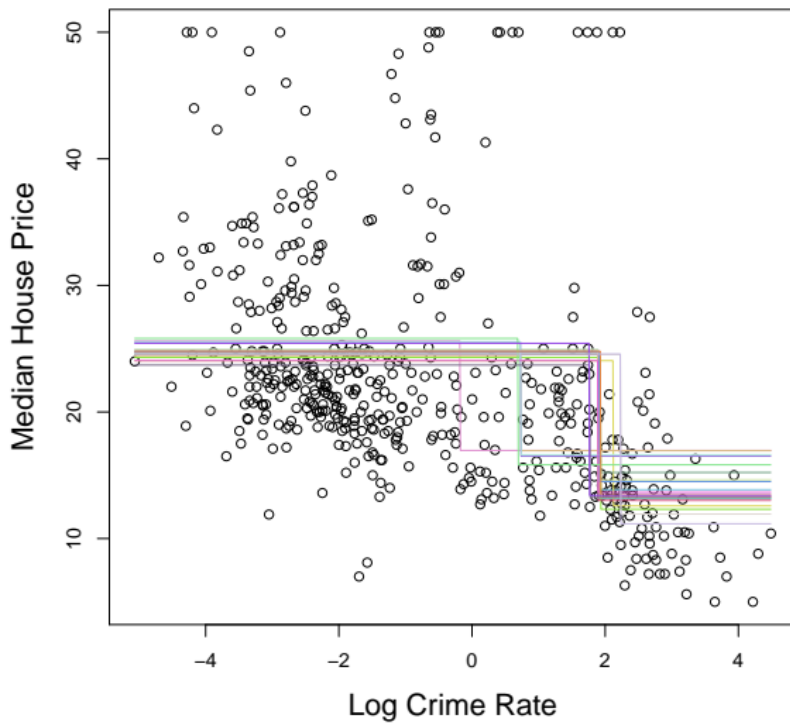


Boston Housing Price

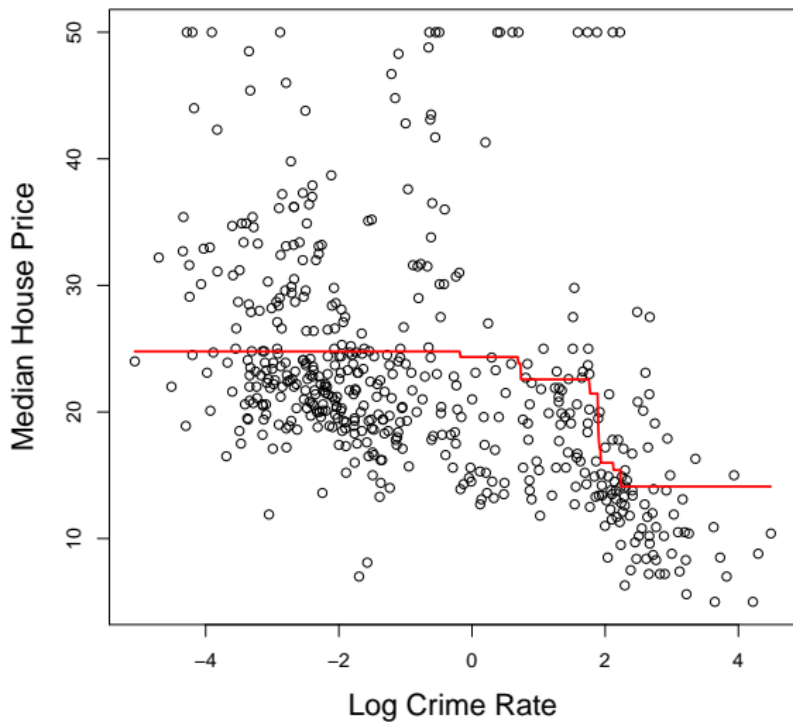
Now let's perform bagging using $B = 20$ bootstrap samples and make predictions of housing price on a range of crime rate values:

```
crimelims = range(Boston$lncrime)
crime.grid = seq(crimelims[1],crimelims[2],0.01)
#
# Function to fit a single stump
stump = function(index,b){
  fit = rpart(medv~lncrime,data=Boston,subset=indx,
              control=rpart.control(maxdepth=1))
  return(predict(fit,newdata=list(lncrime=crime.grid)))
}
# Bagging
B = 20
yhat_bag = 0
for (b in 1:B){
  bootsample = sample(n,n,replace=T)
  yhat_bag = yhat_bag + stump(bootsample,b)/B
}
```

Boston Housing Price



Boston Housing Price



Bagging for Classification

For classification problems, let $\hat{p}_j^{(b)}(x)$ be the class probabilities estimated on bootstrap sample b . We have:

$$\hat{f}(x) = \arg \max_{j \in \{1, \dots, J\}} \hat{p}_j(x) \quad (7)$$

, where

$$\hat{p}_j(x) = \frac{1}{B} \sum_{b=1}^B \hat{p}_j^{(b)}(x)$$

This is the *probability approach*.

Bagging for Classification

Alternatively, for classification problems, let

$$\hat{f}(x) = \arg \max_{j \in \{1, \dots, J\}} \sum_{b=1}^B \mathcal{I}(\hat{f}^{(b)}(x) = j) \quad (8)$$

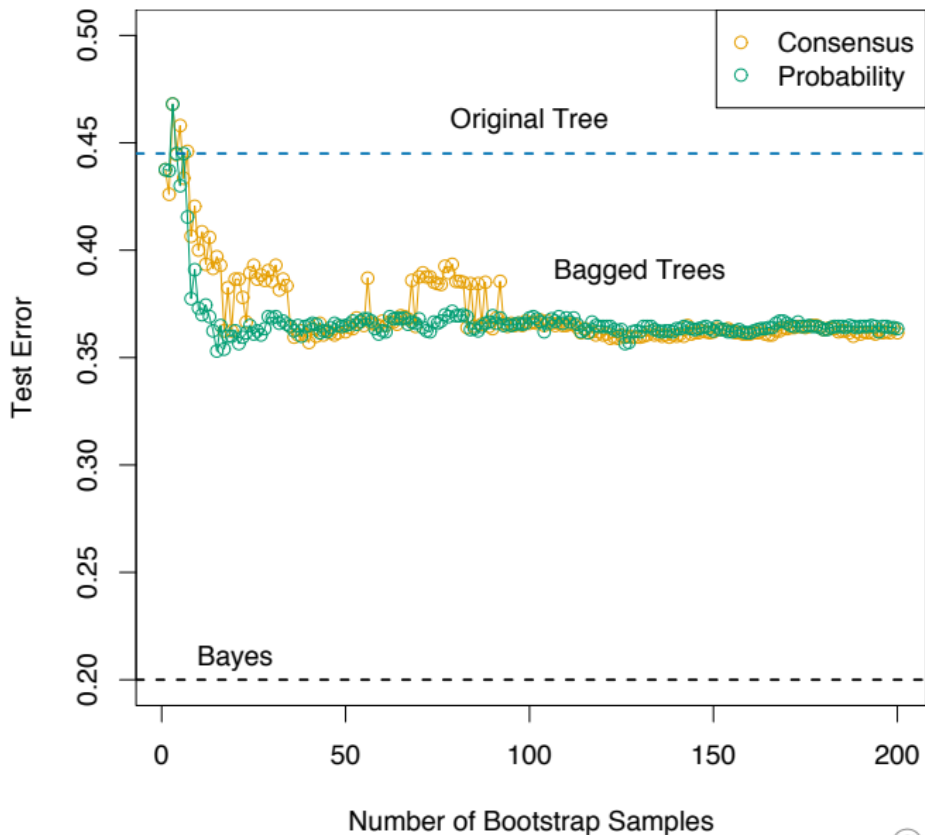
This is the *consensus approach* – decision by “**majority vote**”.

Bagging for Classification

Simulation: $N = 30, p = 5, J = 2$. Each predictor has a standard Gaussian distribution with pairwise correlation 0.95. y is generated according to $p(y = 1 | x_1 \leq 0.5) = 0.2$ and $p(y = 1 | x_1 > 0.5) = 0.8$.

Bagging is performed by fitting classification trees (with no pruning) to 200 bootstrap samples.

Bagging for Classification



The Wisdom of Crowds

Consider a binary classification problem where $y \in \{F, T\}$. Suppose that for a given input x , the true value of y is T . We have B independent classifiers, $f^{(b)}(x)$, $b = 1, \dots, B$, and each has a misclassification rate of 0.4, i.e., $\Pr(f^{(b)}(x) = F) = 0.4$.

Let $\mathbb{V} = \sum_{b=1}^B \mathcal{I}(f^{(b)}(x) = T)$ be the total number of “votes” for $y = T$ among the B classifiers. Then

$$\mathbb{V} \sim \text{Binomial}(B, 0.6)$$

The bagged classifier (consensus approach) is

$$f^{\text{bag}}(x) = \arg \max_{j \in \{F, T\}} \sum_{b=1}^B \mathcal{I}(f^{(b)}(x) = j)$$

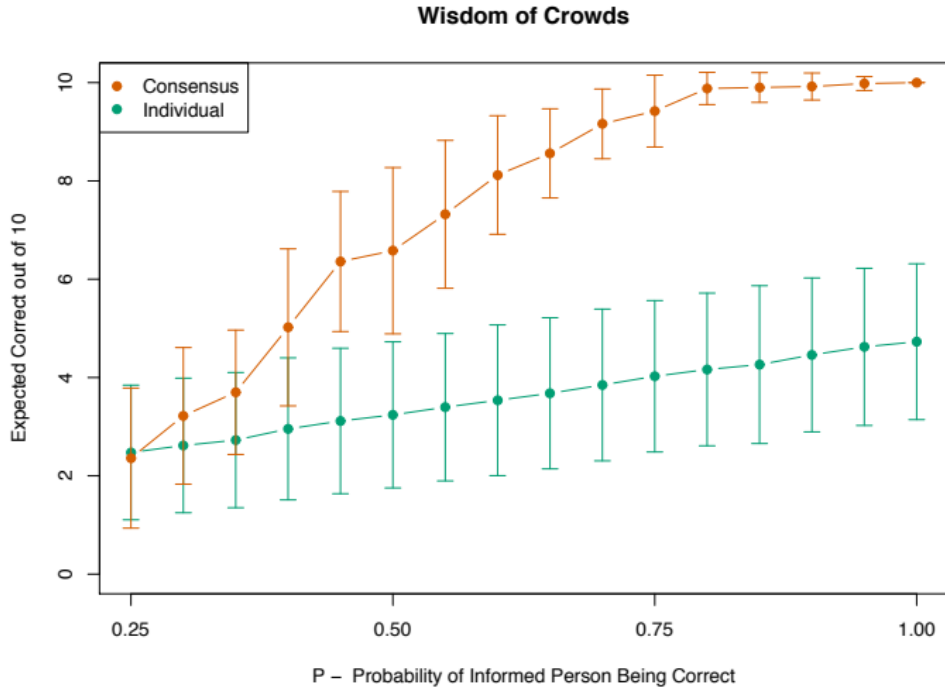
The Wisdom of Crowds

Hence the misclassification rate of the bagged classifier is

$$\Pr(f^{\text{bag}}(x) = F) = \Pr\left(\frac{\text{votes}}{B} < \frac{B}{2}\right) \rightarrow 0 \quad \text{as } B \rightarrow \infty \quad (9)$$

- (9) is true in this example as long as the misclassification rate of the individual classifier is less than 0.5. If $\Pr(f^{(b)}(x) = F) > 0.5$, then $f^{\text{bag}}(x)$ will become perfectly inaccurate as $B \rightarrow \infty$.
- Bagging a good classifier can improve predictive accuracy, but bagging a bad one hurts.

The Wisdom of Crowds



50 members vote in 10 categories, each with 4 nominations. For each category, only 15 (randomly selected) voters are informed (probability of selecting the "correct" candidate > 0.25).

Bagging

- Bagging can dramatically reduce the variance and **stabilize** the predictions of unstable procedures, and is therefore particularly useful for **high-variance, low-bias** procedures.
- Trees are ideal candidates for bagging, since they can capture complex interaction structures in the data, and if grown sufficiently deep, have relatively low bias.
- Real structure that persists across data sets shows up in the average. Noisy useless signals will average out to have no effect.
- This technique of **model averaging** is central to many advanced nonparametric learning algorithms.
- Downside? Loss of interpretability. *A bagged tree is no longer a tree.* Bagging improves prediction accuracy at the expense of interpretability.

Random Forests

- **Random forests** improve on bagging by reducing the correlation between the sampled trees.
- We know that given a set of N identically distributed random variables with variance σ^2 and positive pairwise correlation ρ , the variance of the mean is $\rho\sigma^2 + (1 - \rho)\frac{\sigma^2}{N}$.
- Hence, in the context of decision trees, we can improve the variance reduction of bagging by reducing the correlation between the trees, without increasing the variance too much.
- This is achieved in random forests through random selection of the input variables in the tree-growing process.

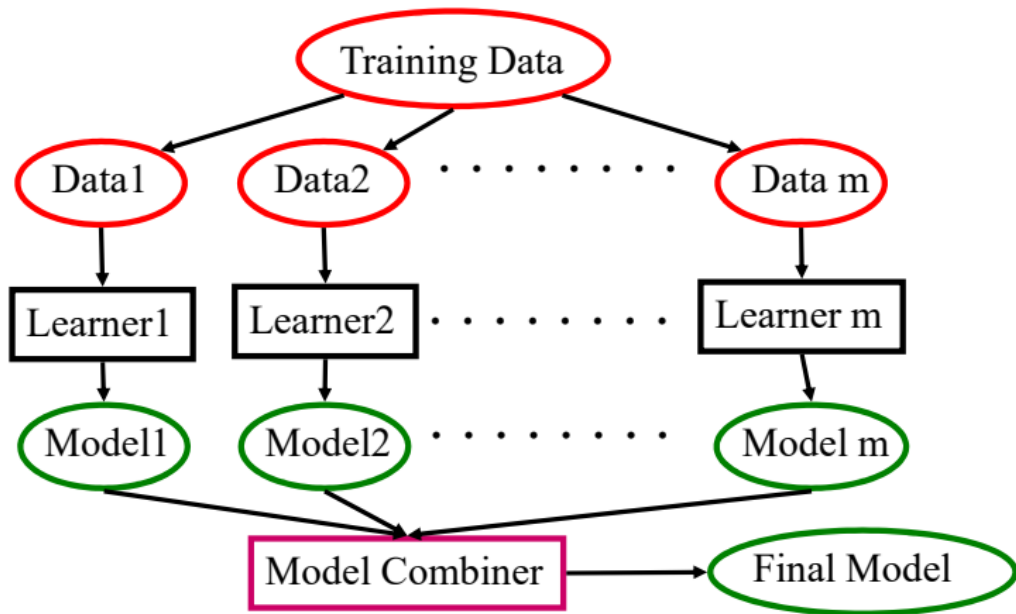
Random Forests

- As in bagging, random forests build a number of trees on bootstrapped samples.
- When building these trees, each time a split in a tree is considered, a random selection of m ($m < p$) predictors is chosen as split candidates from the full set of p predictors. The split is allowed to use only one of these m predictors.
- A fresh selection of m predictors is taken at each split.
 - ▶ Typical choice of m : $m = \sqrt{p}$.

Ensemble Learning

- Bagging and random forests are examples of **ensemble methods**.
- Instead of learning a single hypothesis from a hypothesis set, ensemble methods select a collection (ensemble) of hypotheses and combine their predictions.
- Doing so often improves prediction accuracy at the expense of interpretability.

Ensemble Learning



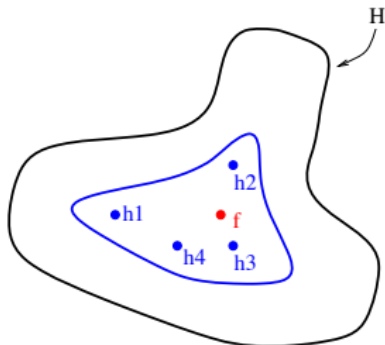
Ensemble Learning

Three reasons why ensemble methods may work well:

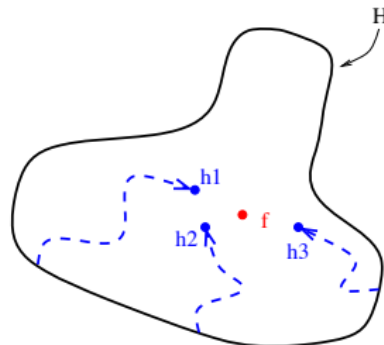
- ① **Statistical**: for a given hypothesis set \mathcal{H} , when sample size is small, many different hypotheses in \mathcal{H} can all give the same accuracy on the training data. Combining their predictions helps lower the variance and improve prediction accuracy.
- ② **Computational**: combining local optima produced by local search
- ③ **Representational**: by forming weighted sums of hypotheses drawn from \mathcal{H} , it might be possible to expand \mathcal{H} and produce a better approximation to target function f .

Ensemble Learning

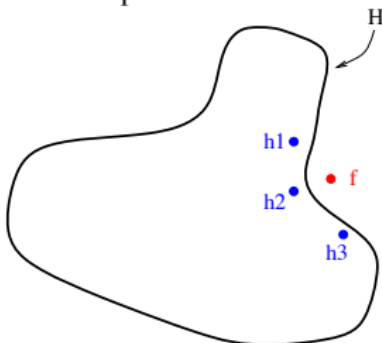
Statistical



Computational



Representational



Boosting

- Like bagging, boosting is a general and powerful tool that can be applied to many statistical learning methods for regression or classification.
- While bagging fits a model (**base learner**) independently to multiple bootstrap samples, boosting fits a simple model (**weak learner**) to the training data **sequentially** to construct a more complex model (**strong learner**).
- Basic idea: in each round, the training data are **re-weighted** according to the performance of the model in the previous round. Observations that were not accurately predicted see their weights increase. The model is then fit to the re-weighted data. In this way, we **slowly** improve the performance of the model in areas where it does not perform well.

L2 Boost (Regression)

① Set $\hat{f}(x) = 0$ and $r_i = y_i \forall i$

② For $b = 1, \dots, B$:

① Fit a regression model $\hat{f}^{(b)}$ to the training data $\{(x_1, r_1), \dots, (x_N, r_N)\}$.

② Update \hat{f} by adding in a *shrunken* version of $\hat{f}^{(b)}$:

$$\hat{f}(x) \leftarrow \hat{f}(x) + \lambda \hat{f}^{(b)}(x)$$

③ Update the residuals:

$$r_i \leftarrow r_i - \lambda \hat{f}^{(b)}(x_i)$$

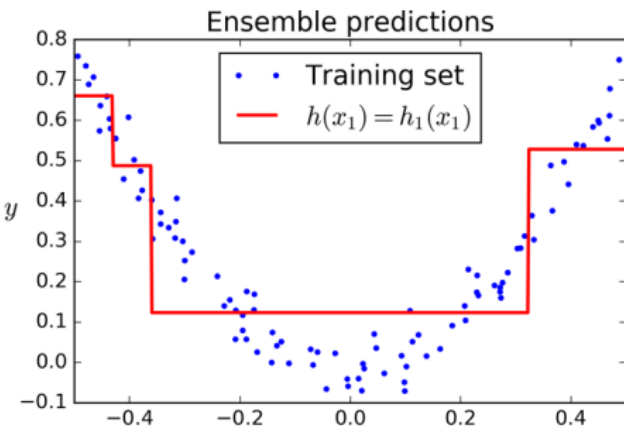
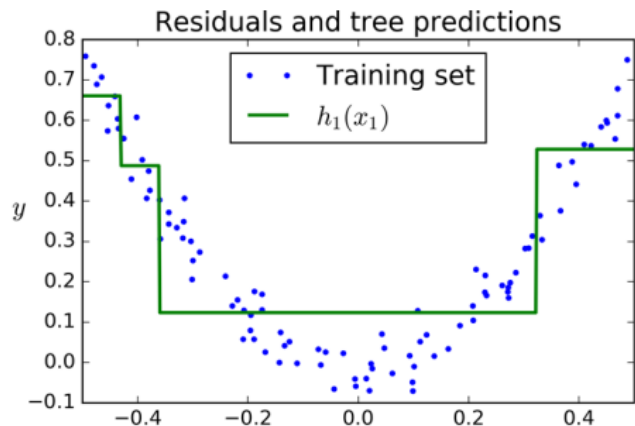
④ Output

$$\hat{f}(x) = \sum_{b=1}^B \lambda \hat{f}^{(b)}(x)$$

Boosting for Regression

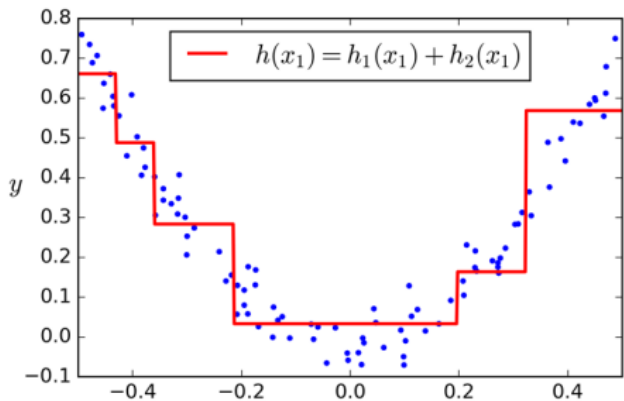
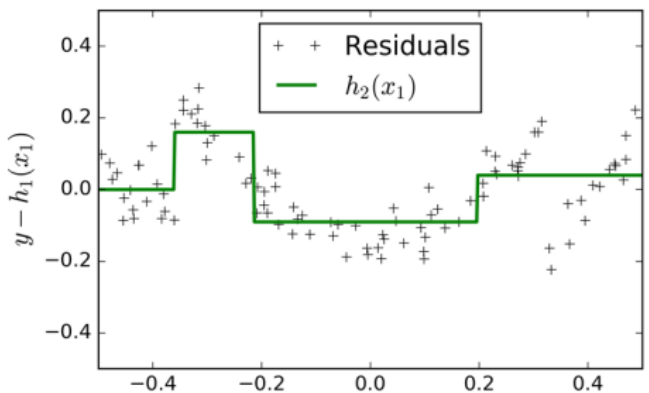
- When applying boosting to regression algorithms, in each round b , we simply fit the base learner, $\hat{f}^{(b)}$, to the residuals of $\hat{f}^{(b-1)}$. This is akin to re-weighting the training data: observations that were fit well in the previous round see their importance diminished (r_i small), while those that were not fit well become more important (r_i large) in this round.
- The base learner $\hat{f}^{(b)}$ is usually a *weak learner*, e.g., a stump.
- The shrinkage parameter λ is a small positive number (typical values: 0.01, 0.001, etc.). The boosting approach is to learn *slowly* and λ controls the rate of learning.

Boosting for Regression



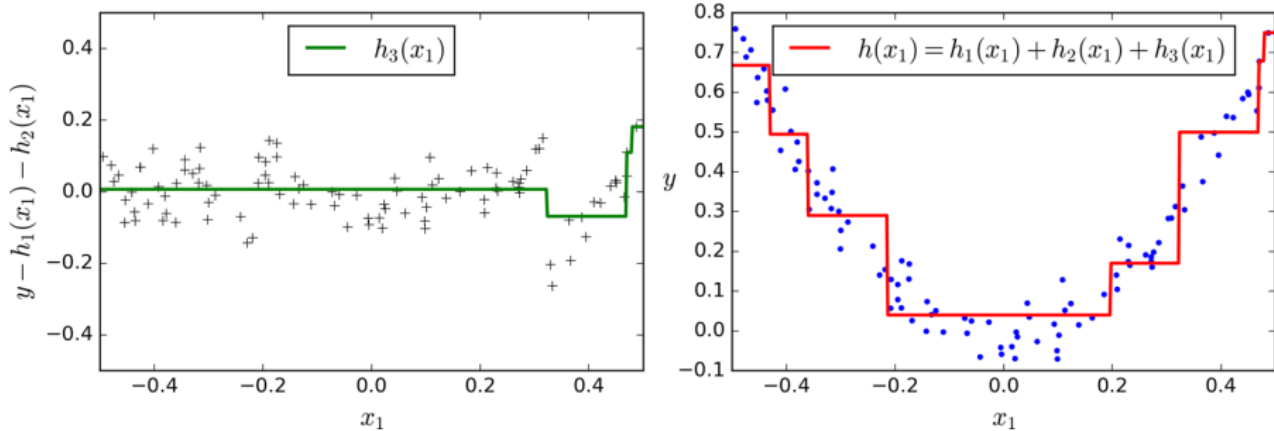
L2 Boost ($\lambda = 1$)

Boosting for Regression



L2 Boost ($\lambda = 1$)

Boosting for Regression



L2 Boost ($\lambda = 1$)

AdaBoost (Binary Classification)

- ① Let $y \in \{-1, 1\}$. Initialize $w_i = \frac{1}{N} \forall i$.
- ② For $b = 1, \dots, B$:
 - ① Fit a classifier $\hat{f}^{(b)}$ to the training data by minimizing the weighted error

$$\frac{\sum_{i=1}^N w_i \mathcal{I}(y_i \neq f^{(b)}(x_i))}{\sum_{i=1}^N w_i}$$

- ② Let $\alpha_b = \log((1 - \epsilon_b)/\epsilon_b)$, where ϵ_b is the weighted error of $\hat{f}^{(b)}$, and update w_i as follows:

$$w_i \leftarrow w_i \exp(\alpha_b \mathcal{I}(y_i \neq \hat{f}^{(b)}(x_i)))$$

- ③ Output

$$\hat{f}(x) = \text{sign} \left(\sum_{b=1}^B \alpha_b \hat{f}^{(b)}(x) \right)$$

Boosting for Classification

- For classification problems, the boosted classifier is a weighted sum of individual classifiers, with weights proportional to each classifier's accuracy on the training set (good classifiers get more weight).
- In AdaBoost, If an individual classifier has accuracy $< 50\%$ ($\epsilon_b > 0.5$), we flip the sign of its predictions and turn it into a classifier with accuracy $> 50\%$. This is achieved by making $\alpha_b < 0$ so that the classifier enters negatively into the final hypothesis.

Boosting for Classification

- In each round of the boosting process, we re-weight the observations in the training data: those that were misclassified in the previous round⁴ see their weights increase relative to those that were correctly classified. In this way, successive classifiers are forced to place greater emphasis on points that have been misclassified by previous classifiers, and data points that continue to be misclassified by successive classifiers receive ever greater weight⁵.

⁴ After classifiers with $\epsilon_b > 0.5$ are “flipped”.

⁵ This adaptive data weighting scheme gives the algorithm its name: AdaBoost stands for “Adaptive Boosting”.

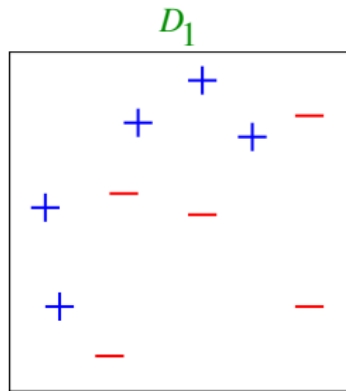
Boosting for Classification

Now suppose we use decision tree as the base classifier. To fit classification trees to weighted data, we calculate the weighted \hat{p}_j^m for each region:

$$\hat{p}_j^m = \frac{\sum_{x_i \in R_m} w_i \mathcal{I}(y_i = j)}{\sum_{x_i \in R_m} w_i} \quad (10)$$

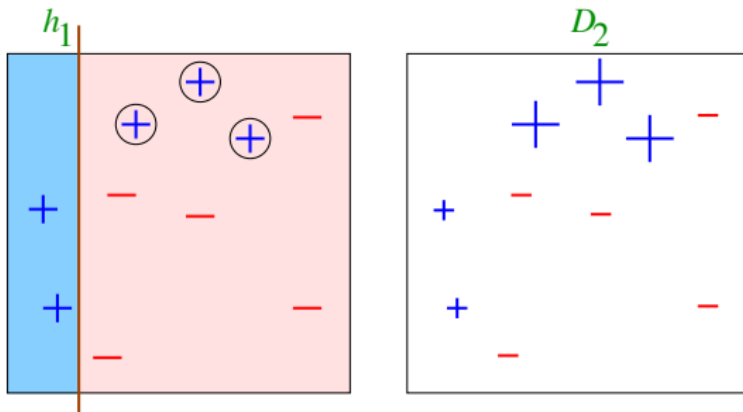
and let (10) replace (4) in the CART algorithm.

Boosting for Classification



Weak classifiers = vertical or horizontal half-planes

Boosting for Classification

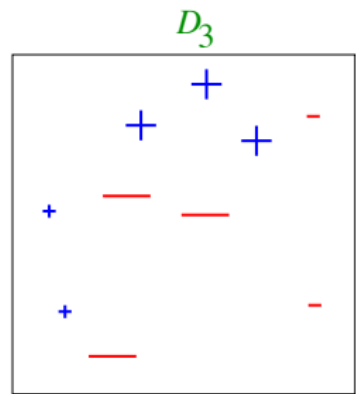
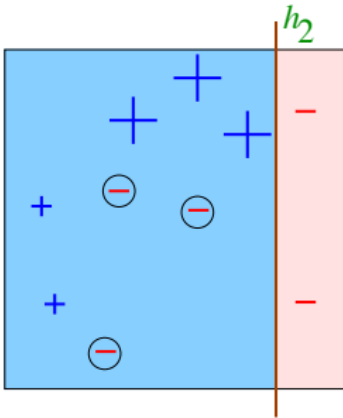
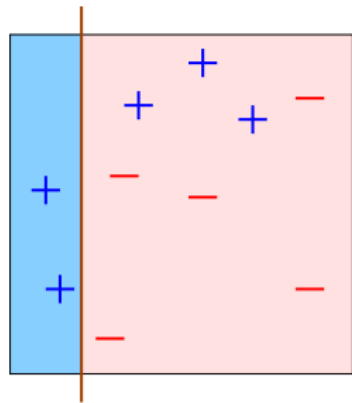


$$\varepsilon_1 = 0.30$$

$$\alpha_1 = 0.42$$

Round 1

Boosting for Classification

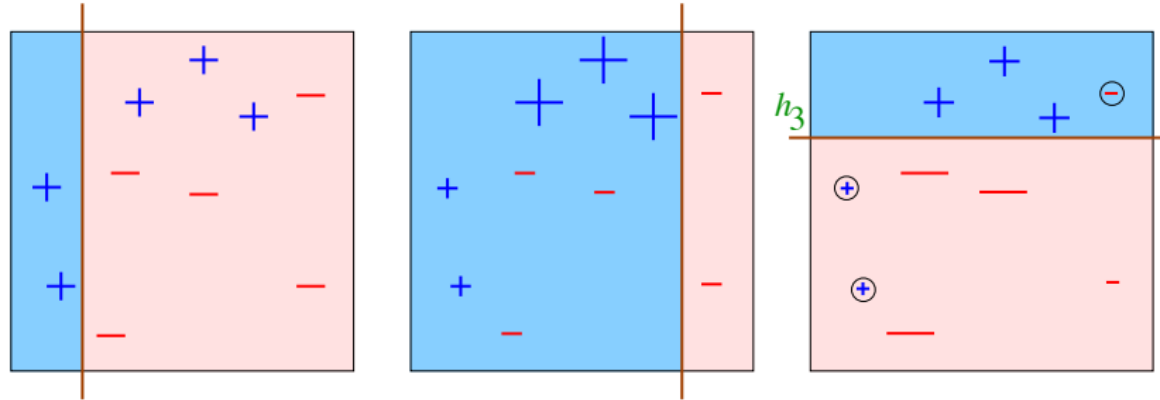


$$\varepsilon_2 = 0.21$$

$$\alpha_2 = 0.65$$

Round 2

Boosting for Classification

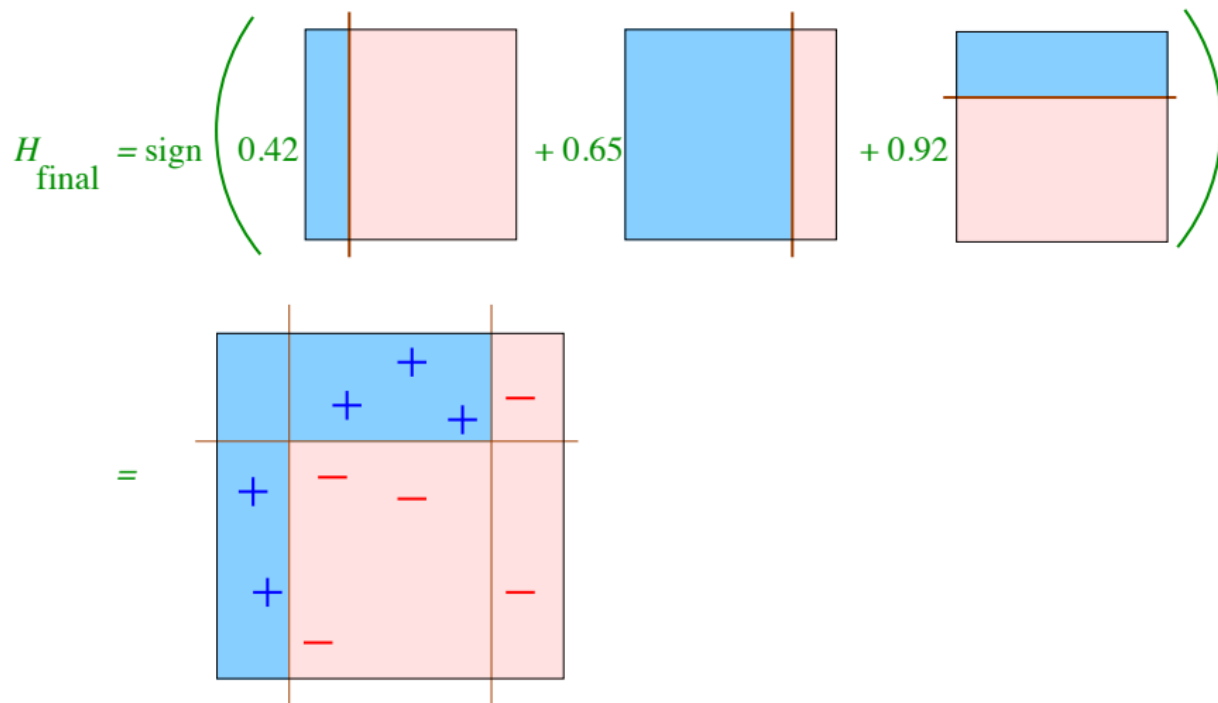


$$\varepsilon_3 = 0.14$$

$$\alpha_3 = 0.92$$

Round 3

Boosting for Classification



Final Classifier

Boosting

In essence, boosting is a **forward stagewise** algorithm to fit an adaptive additive model:

$$f(x) = \alpha_1\phi(x; \beta_1) + \cdots + \alpha_M\phi(x; \beta_M) \quad (11)$$

(11) is equivalent in functional form to a linear basis function model. The difference is that here the basis functions are *base learners* (e.g., decision trees) whose functional form are not chosen before seeing the data but are *learned* from the data – they are called **adaptive basis functions**.

Boosting

Fitting (11) requires minimizing the in-sample error:

$$E_{in}(\theta) = \frac{1}{N} \sum_{i=1}^N \ell(y_i, f(x_i))$$

, where $\theta = \{\alpha_m, \beta_m\}_{m=1}^M$.

- L2 loss: $\ell(y, f(x)) = (y - f(x))^2 \Rightarrow$ L2 Boost
- Exponential loss: $\ell(y, f(x)) = \exp(-y \cdot f(x)) \Rightarrow$ AdaBoost⁶

⁶See [Appendix](#) for why AdaBoost corresponds to forward stagewise additive modeling with exponential loss.

Boosting

Instead of doing a global minimization, the boosting strategy is to follow a forward stagewise procedure by adding basis functions *one by one*⁷:

- Choose the $\phi(x; \beta_1)$ and α_1 that gives the smallest in-sample error.
Let $\hat{f}(x) = \hat{\alpha}_1 \phi(x; \hat{\beta}_1)$.
- Choose the $\phi(x; \beta_2)$ and α_2 that gives the largest *additional* reduction in in-sample error:

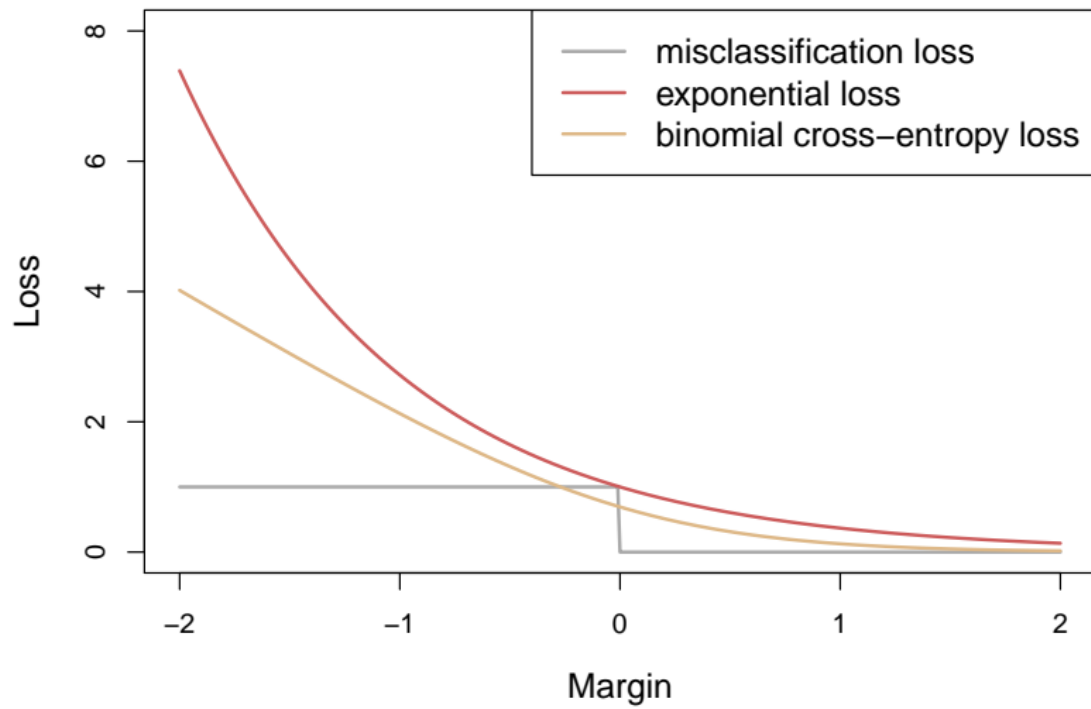
$$(\hat{\alpha}_2, \hat{\beta}_2) = \arg \min_{\alpha_2, \beta_2} \sum_{i=1}^N \ell(y_i, \hat{f}(x_i) + \alpha_2 \phi(x_i; \beta_2))$$

$$\text{Let } \hat{f}(x) = \hat{\alpha}_1 \phi(x; \hat{\beta}_1) + \hat{\alpha}_2 \phi(x; \hat{\beta}_2).$$

- And continue ...

⁷Similar to forward stepwise regression. The difference between the forward stagewise and the forward stepwise is that all coefficients can change at each step in a stepwise algorithm, while a stagewise procedure updates only one coefficient at each step.

Boosting



Simulation 1

Let's first generate some data for a two-dimensional classification problem:

$$y \sim \text{Bernoulli}(p) \quad (12)$$

, where

$$p = \frac{1}{1 + \exp\{- (2x_1 + 2x_2)\}}$$

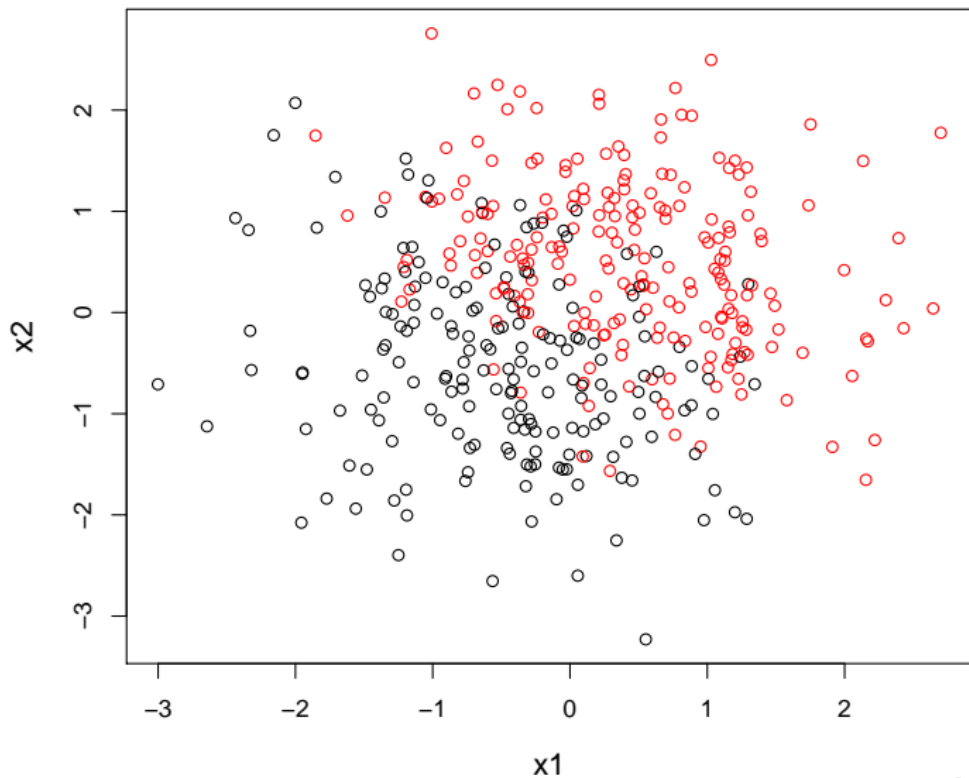
Then (12) implies the true Bayes decision boundary is: $x_1 + x_2 = 0$.

Simulation 1

```
# Simulation
n = 1000
x = matrix(rnorm(n*2), ncol=2)
z = 2*x[,1] + 2*x[,2]
p = exp(z)/(1+exp(z)) #p(y=1)
y = (p>runif(n))

# Create training and test sets
mydata = data.frame(x1=x[,1], x2=x[,2], y=as.factor(y))
train = sample(n, n*0.4)
data_train = mydata[train,]
data_test = mydata[-train,]
```

Simulation 1



Simulation 1

```
#####
# Logistic Regression #
#####

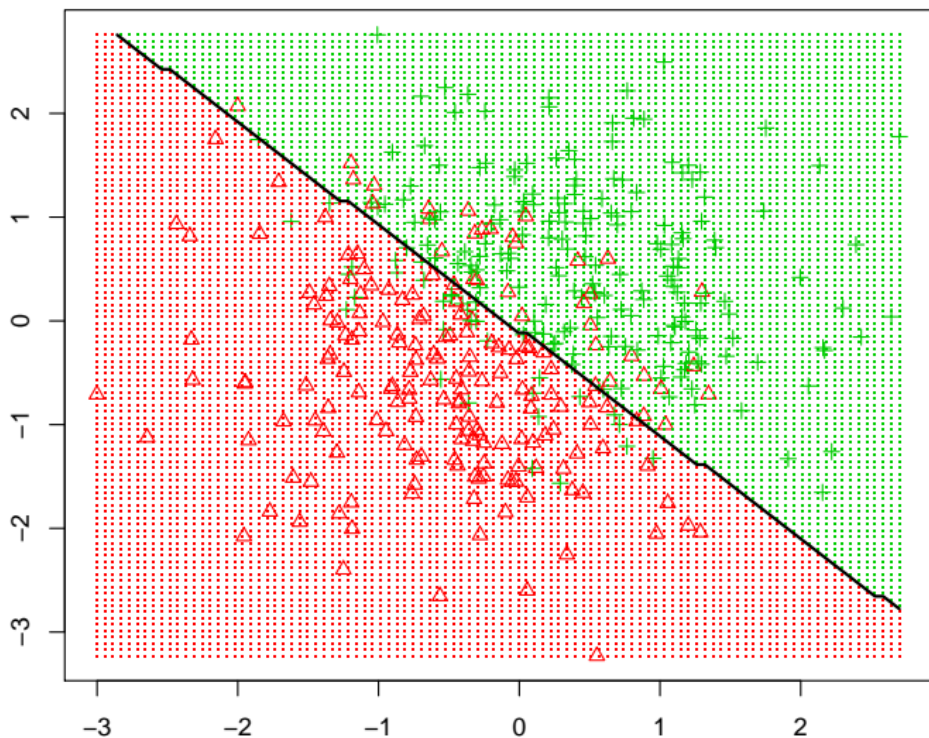
require(nnet)
require(AER)
fit = multinom(y~.,data_train)

coeftest(fit)

##
## z test of coefficients:
##
##           Estimate Std. Error z value Pr(>|z|)
## (Intercept)  0.18475   0.14825  1.2462  0.2127
## x1          2.00503   0.22415  8.9450 <2e-16 ***
## x2          1.99984   0.22210  9.0041 <2e-16 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Simulation 1

Logistic Regression



Simulation 1

```
# test err
ytrue = data_test[, "y"]
yhat = predict(fit, data_test)
table(ytrue, yhat)

##          yhat
## ytrue   FALSE  TRUE
##   FALSE    223    61
##   TRUE      43   273

err = 1 - mean(yhat == ytrue)
err

## [1] 0.1733333
```

Simulation 1

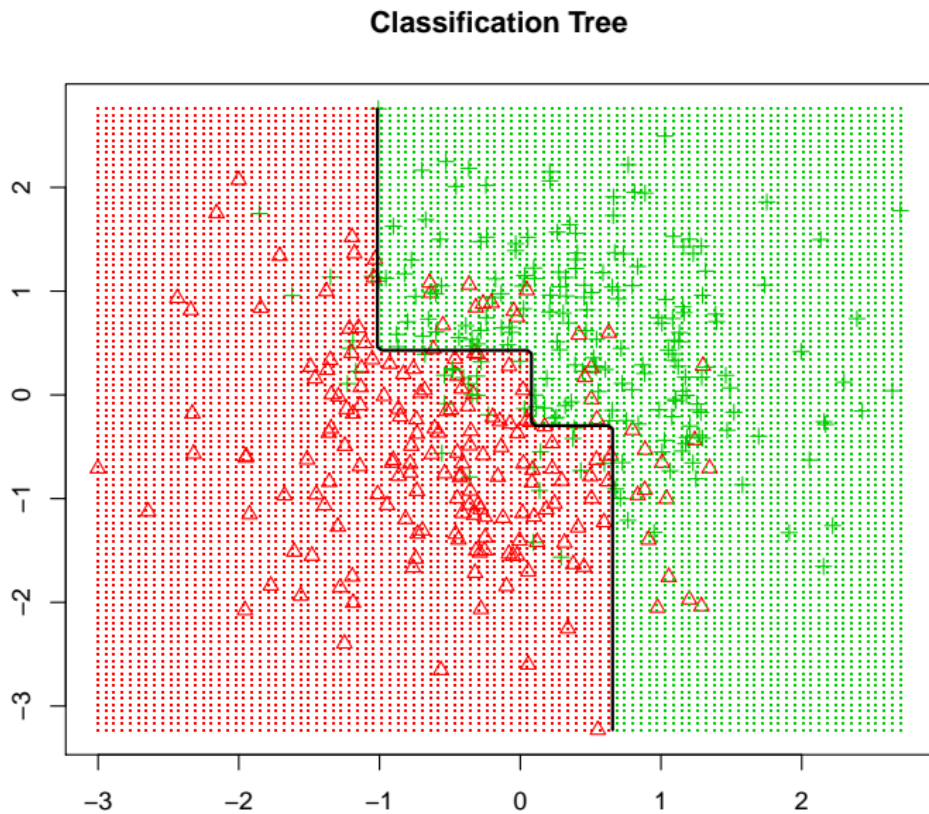
```
#####
# Classification Tree #
#####

require(rpart)
fit0 = rpart(y~.,data_train,control=rpart.control(cp=0))
fit = prune(fit0,cp=fit0$cptable[which.min(fit0$cptable[, "xerror"]),"CP"])

# test err
yhat = predict(fit,data_test,type="class")
err = 1-mean(yhat==ytrue)
err

## [1] 0.22
```

Simulation 1



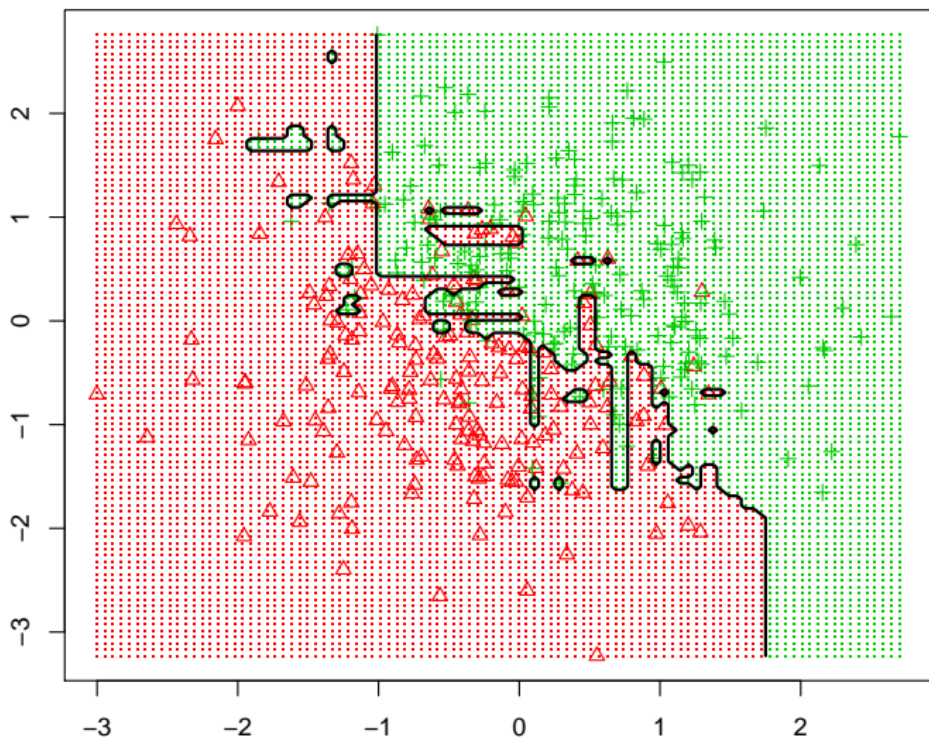
Simulation 1

```
#####
# Bagging #
#####
require(randomForest)
# mtry: number of predictors considered for each split
# bagging is a special case of a random forest with mtry = all predictors
fit = randomForest(y~.,data_train,mtry=2)
fit

##
## Call:
##   randomForest(formula = y ~ ., data = data_train, mtry = 2)
##           Type of random forest: classification
##                   Number of trees: 500
## No. of variables tried at each split: 2
##
##           OOB estimate of  error rate: 21.75%
## Confusion matrix:
##   FALSE TRUE class.error
## FALSE  140   47   0.2513369
## TRUE    40  173   0.1877934
```

Simulation 1

Bagging (overfit)



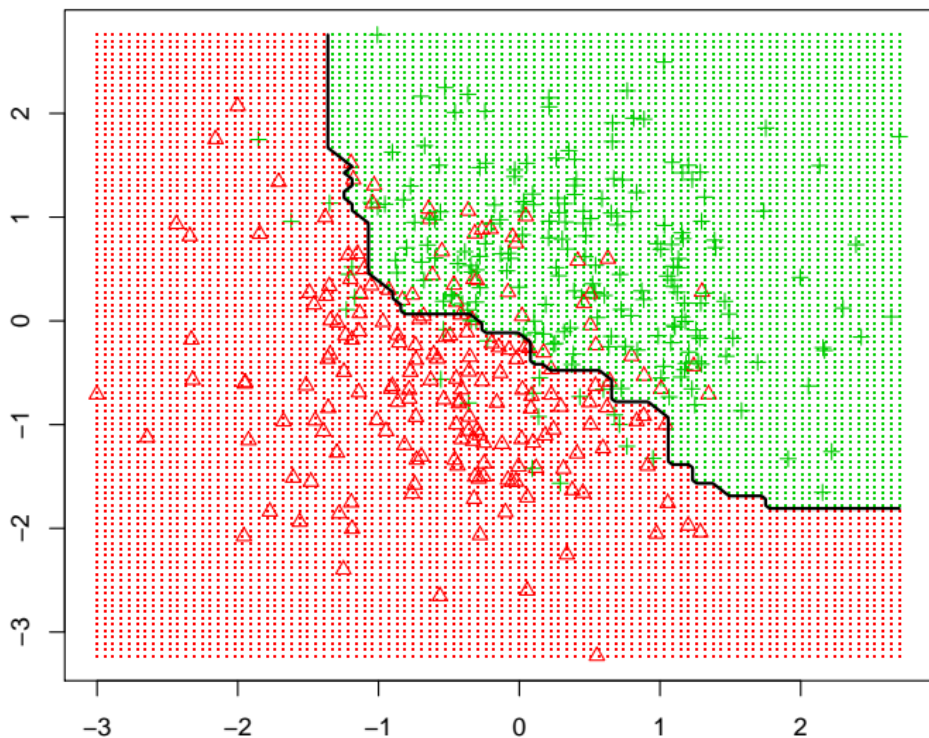
Simulation 1

```
#####
# Bagging #
#####
fit = randomForest(y~.,data_train,mtry=2,maxnodes=8)
#
# test err
yhat = predict(fit,data_test)
err = 1-mean(yhat==ytrue)
err

## [1] 0.2066667
```

Simulation 1

Bagging

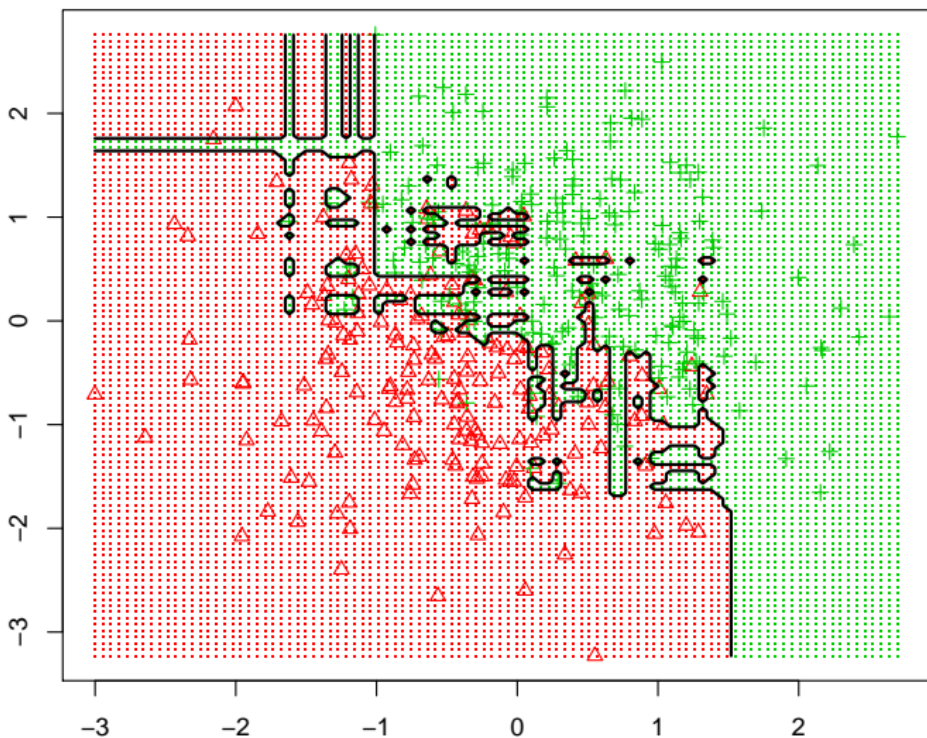


Simulation 1

```
#####
# Boosting #
#####
require(gbm)
data_boost = transform(data_train,y=as.numeric(y)-1) # requires y={0,1}
fit = gbm(y~.,data_boost,distribution="adaboost", # use AdaBoost
           n.trees=5000,
           interaction.depth=10,
           shrinkage=1)
```

Simulation 1

Boosting (overfit)



Simulation 1

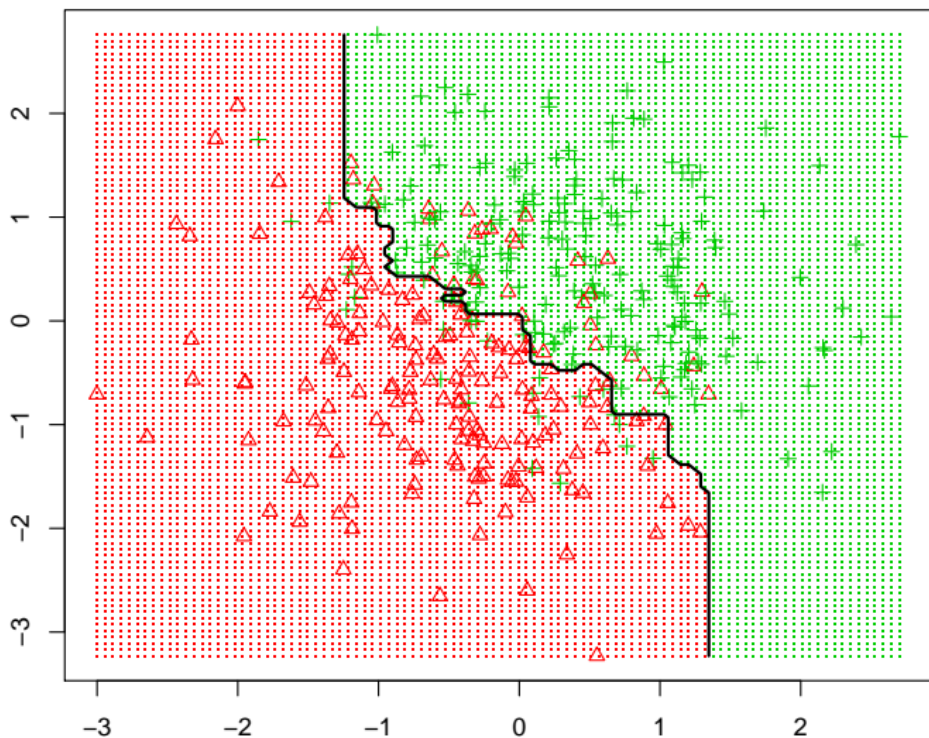
```
#####
# Boosting #
#####
require(gbm)
data_boost = transform(data_train,y=as.numeric(y)-1)
fit = gbm(y~.,data_boost,distribution="adaboost",
           n.trees=4000,interaction.depth=2,shrinkage=0.001)

# test err
phat = predict(fit,data_test,n.trees=4000,type="response")
yhat = (phat>0.5)
err = 1-mean(yhat==ytrue)
err

## [1] 0.1933333
```

Simulation 1

Boosting



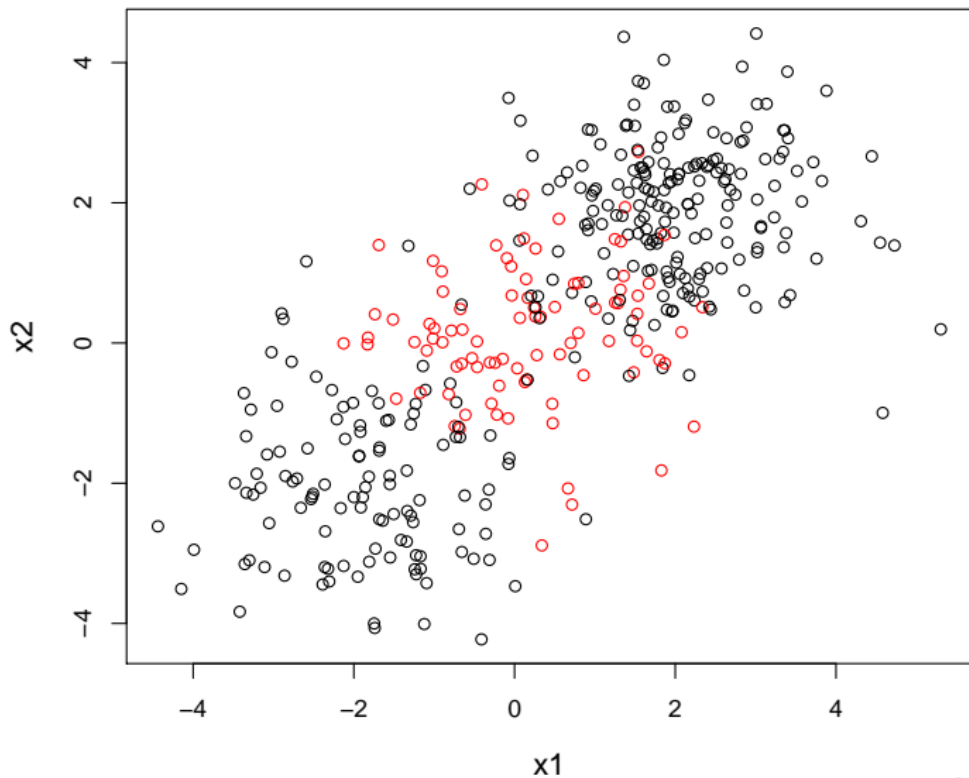
Simulation 2

In Simulation 1, we see tree-based methods struggle to capture linear decision boundaries. Now let's generate some data for binary classification with a nonlinear decision boundary:

```
# Simulation
n = 1000
x = matrix(rnorm(n*2), ncol=2)
x[1:n/2,] = x[1:n/2,] + 2
x[(n/2+1):(n/4*3),] = x[(n/2+1):(n/4*3),] - 2
y = c(rep(1, (n/4*3)), rep(2, (n/4)))

# Create training and test sets
data = data.frame(x1=x[,1], x2=x[,2], y=as.factor(y))
train = sample(n, n*0.4)
data_train = data[train,]
data_test = data[-train,]
```

Simulation 2



Simulation 2

In this case, linear logistic regression (log odds linear in x) would fail to classify $y = 2$ completely:

```
#####
# Logistic Regression: linear #
#####
fit = multinom(y~.,data_train)

# test err
ytrue = data_test[, "y"]
yhat = predict(fit,data_test)
table(ytrue,yhat)

##      yhat
## ytrue  1   2
##      1 440  0
##      2 160  0
```

Simulation 2

Instead, we can try:

```
#####
# Logistic Regression: quadratic #
#####
fit = multinom(y ~ poly(x1,2)+poly(x2,2),data_train)

coeftest(fit)

##
## z test of coefficients:
##
##           Estimate Std. Error z value Pr(>|z|)
## (Intercept) -3.36059   0.39246 -8.5629 < 2.2e-16 ***
## poly(x1, 2)1 -10.50099  5.74763 -1.8270   0.0677 .
## poly(x1, 2)2 -52.04420  8.64979 -6.0168 1.779e-09 ***
## poly(x2, 2)1 -3.28473  6.23134 -0.5271   0.5981
## poly(x2, 2)2 -47.40449  7.72478 -6.1367 8.426e-10 ***
## ---
## Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
```

Simulation 2

```
# test err
yhat = predict(fit,data_test)
table(ytrue,yhat)

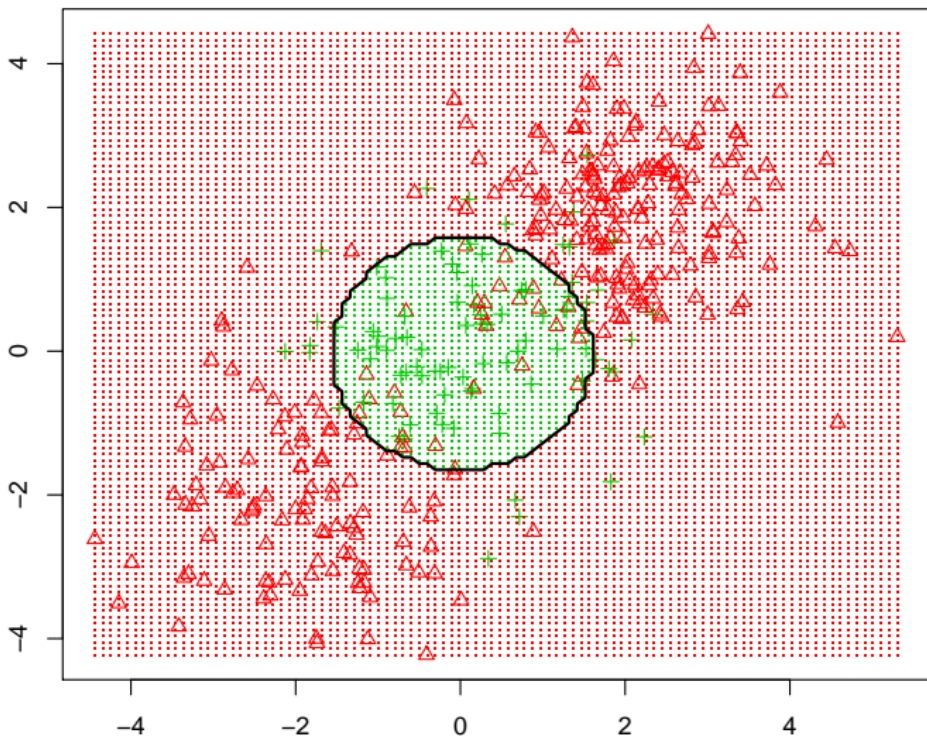
##      yhat
## ytrue   1   2
##   1 410 30
##   2  46 114

err = 1-mean(yhat==ytrue)
err

## [1] 0.1266667
```

Simulation 2

Quadratic Logistic Regression



Simulation 2

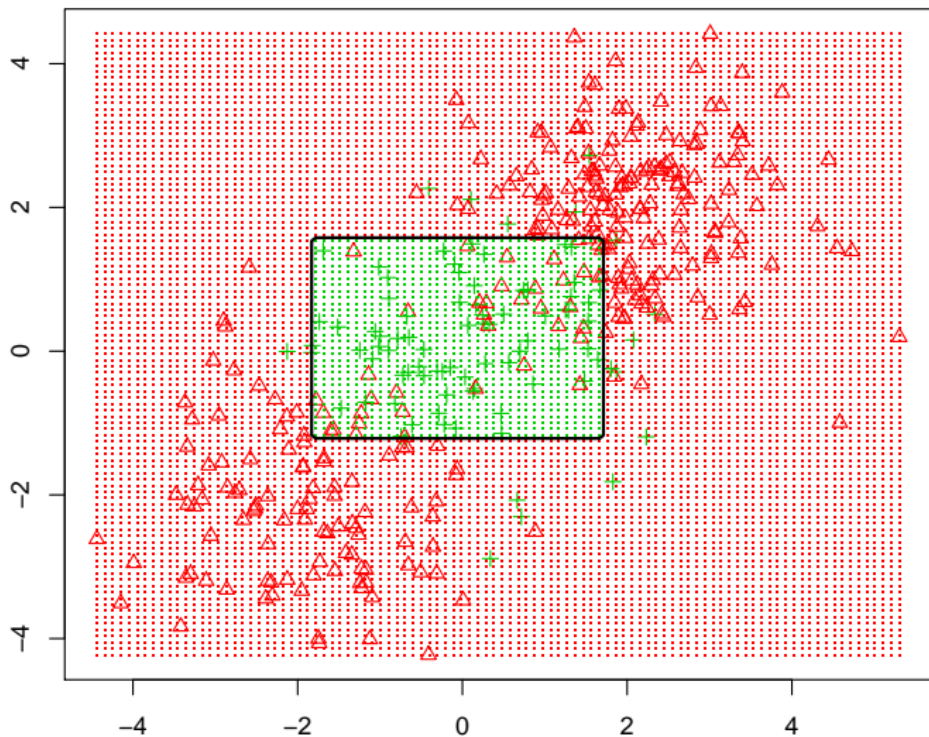
```
#####
# Classification Tree #
#####

fit0 = rpart(y~.,data_train,control=rpart.control(cp=0))
fit = prune(fit0,cp=fit0$cptable[which.min(fit0$cptable[, "xerror"]), "CP"])
#
# test err
yhat = predict(fit,data_test,type="class")
err = 1-mean(yhat==ytrue)
err

## [1] 0.1416667
```

Simulation 2

Classification Tree



Simulation 2

```
#####
# Bagging #
#####
fit = randomForest(y~.,data_train,mtry=2,maxnodes=10)
#
# test err
yhat = predict(fit,data_test)
err = 1-mean(yhat==ytrue)
err

## [1] 0.1233333
```

Simulation 2

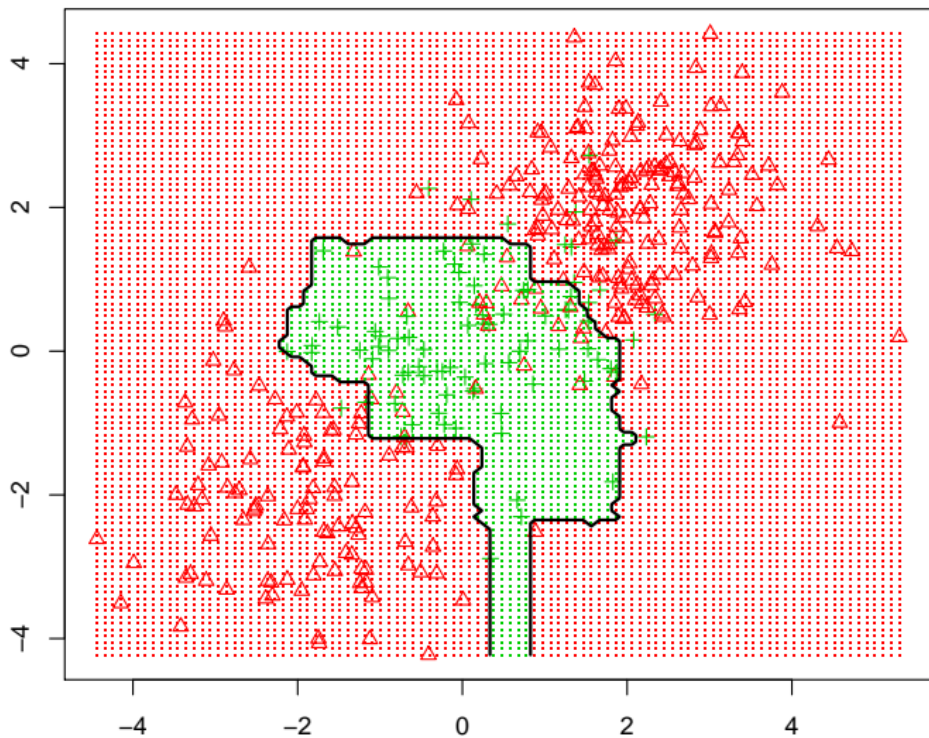
Here we can do better with random forest:

```
#####
# Random Forest #
#####
fit = randomForest(y~.,data_train,mtry=1,maxnodes=10) # use mtry=1
#
# test err
yhat = predict(fit,data_test)
err = 1-mean(yhat==ytrue)
err

## [1] 0.105
```

Simulation 2

Random Forest



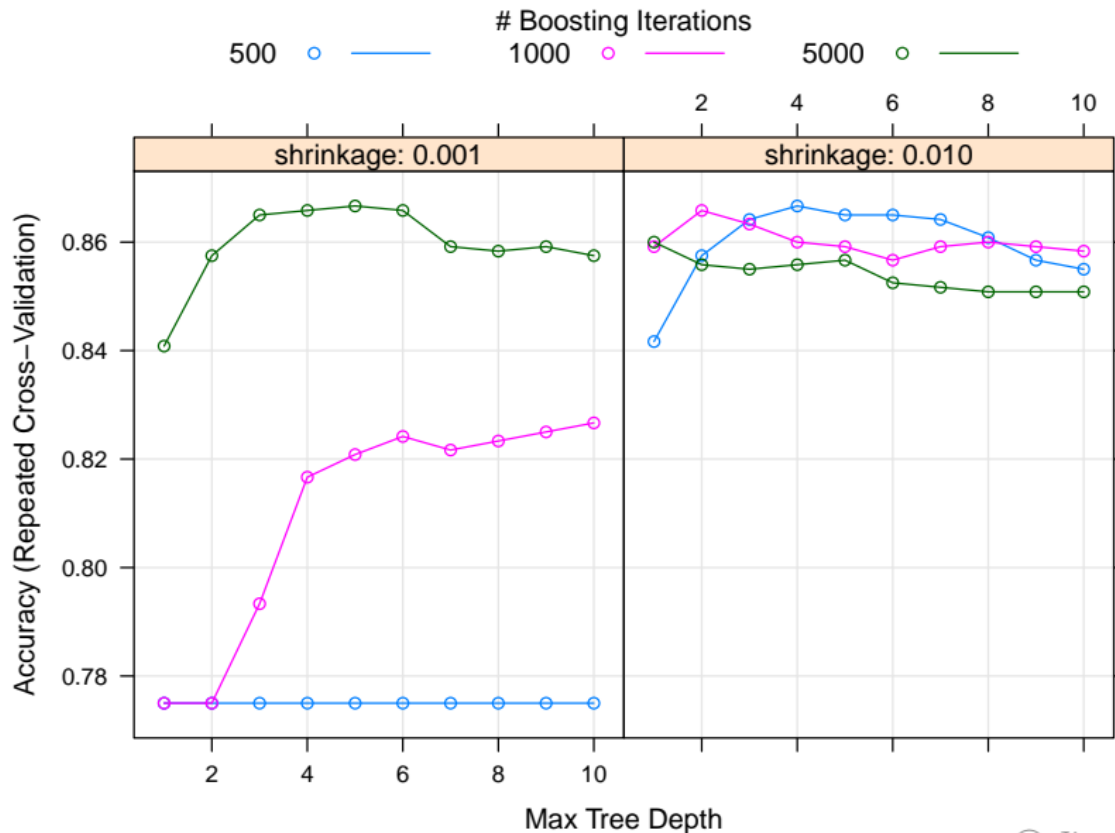
Simulation 2

```
#####
# Boosting #
#####

# We can use cross-validation to select the best number of trees
# (iterations), the size of each tree, as well as the shrinkage factor
require(caret)

fit = train(y~., data=data_train,
            method="gbm", distribution="adaboost",
            tuneGrid=expand.grid(n.trees=c(500,1000,5000),
                                interaction.depth=seq(1:10),
                                n.minobsinnode=1,
                                shrinkage=c(0.001,0.01)),
            trControl=trainControl(method="repeatedcv", repeats=3))
```

Simulation 2



Simulation 2

```
fit$bestTune

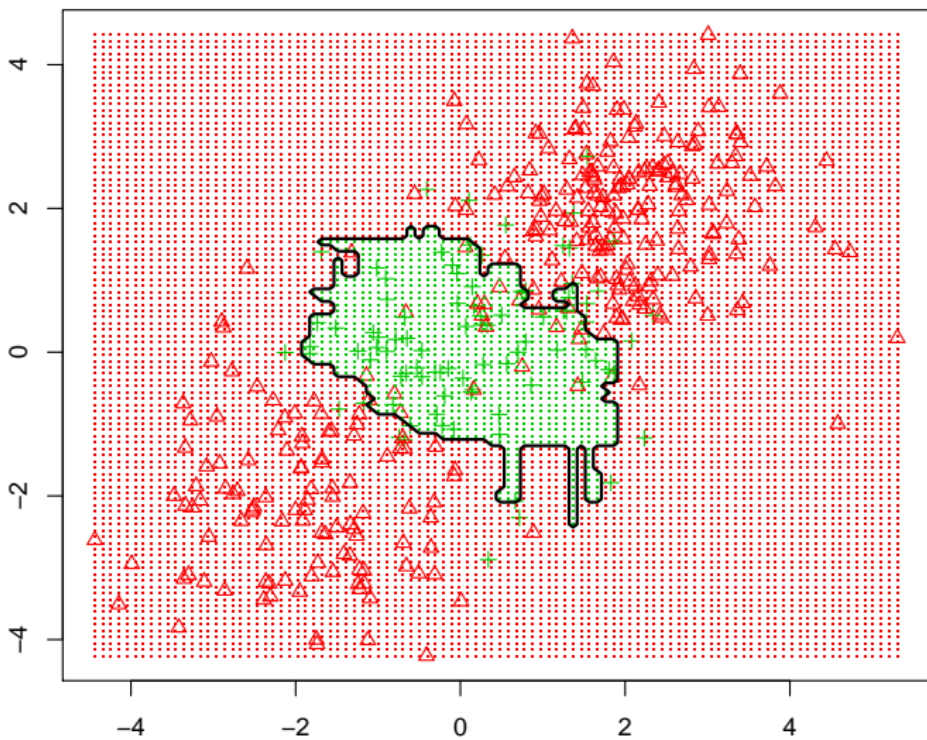
##      n.trees interaction.depth shrinkage n.minobsinnode
## 40      500                  4       0.01          1

# test error
yhat = predict(fit,data_test) # predict using fit$bestTune
err = 1-mean(yhat==ytrue)
err

## [1] 0.1216667
```

Simulation 2

Boosting



Polish Bankruptcy

Data: Polish companies' financial ratios and bankruptcy status^{8,9}.

```
data = read.csv("polish.csv")
data$bankrupt = as.factor(data$bankrupt)
dim(data)

## [1] 5630    56

sum(data$bankrupt==1)/nrow(data) # bankruptcy rate

## [1] 0.07175844
```

⁸We look at bankruptcy status one year after the financial ratios.

⁹Bankrupt companies were analyzed in the period 2000-2012, while the still operating companies were evaluated from 2007 to 2013. See [Source](#).

Polish Bankruptcy

X1 net profit / total assets
X2 total liabilities / total assets
X3 working capital / total assets
X4 current assets / short-term liabilities
X5 [(cash + short-term securities + receivables - short-term liabilities) / (operating expenses - depreciation)] * 365
X6 retained earnings / total assets
X7 EBIT / total assets
X8 book value of equity / total liabilities
X9 sales / total assets
X10 equity / total assets
X11 (gross profit + extraordinary items + financial expenses) / total assets
X12 gross profit / short-term liabilities
X13 (gross profit + depreciation) / sales
X14 (gross profit + interest) / total assets
X15 (total liabilities * 365) / (gross profit + depreciation)
X16 (gross profit + depreciation) / total liabilities
X17 total assets / total liabilities
X18 gross profit / total assets
X19 gross profit / sales
X20 (inventory * 365) / sales
X22 profit on operating activities / total assets
X23 net profit / sales
X24 gross profit (in 3 years) / total assets
X25 (equity - share capital) / total assets
X26 (net profit + depreciation) / total liabilities
X29 logarithm of total assets
X30 (total liabilities - cash) / sales

Polish Bankruptcy

X31 (gross profit + interest) / sales
X32 (current liabilities * 365) / cost of products sold
X33 operating expenses / short-term liabilities
X34 operating expenses / total liabilities
X35 profit on sales / total assets
X36 total sales / total assets
X38 constant capital / total assets
X39 profit on sales / sales
X40 (current assets - inventory - receivables) / short-term liabilities
X41 total liabilities / ((profit on operating activities + depreciation) * (12/365))
X42 profit on operating activities / sales
X43 rotation receivables + inventory turnover in days
X44 (receivables * 365) / sales
X46 (current assets - inventory) / short-term liabilities
X47 (inventory * 365) / cost of products sold
X48 EBITDA (profit on operating activities - depreciation) / total assets
X49 EBITDA (profit on operating activities - depreciation) / sales
X50 current assets / total liabilities
X51 short-term liabilities / total assets
X52 (short-term liabilities * 365) / cost of products sold
X55 working capital
X56 (sales - cost of products sold) / sales
X57 (current assets - inventory - short-term liabilities) / (sales - gross profit - depreciation)
X58 total costs /total sales
X59 long-term liabilities / equity
X61 sales / receivables
X62 (short-term liabilities *365) / sales
X63 sales / short-term liabilities

Polish Bankruptcy

```
# create training and test sets
require(caret)
train = createDataPartition(data$bankrupt,p=0.7,list=F) # 70% in training
data_train = data[train,]
data_test = data[-train,]

# training bankruptcy rate
sum(data_train$bankrupt==1)/nrow(data_train)

## [1] 0.07179097

# test bankruptcy rate
sum(data_test$bankrupt==1)/nrow(data_test)

## [1] 0.07168246
```

Polish Bankruptcy

```
#####
# Logistic Regression #
#####

require(AER)
fit = glm(bankrupt~.,data_train,family="binomial")
result = coeftest(fit)
result[1:4,]

##           Estimate Std. Error   z value Pr(>|z|)
## (Intercept) 2.387528e+15  26334094 90663008     0
## Attr1       5.607271e+14  21817873 25700357     0
## Attr2      -1.424042e+15  23449239 -60728697     0
## Attr3      -3.915047e+14  5993645 -65319962     0
```

All coefficient estimates are significant ...

```
sum(result[,4]>0.05)

## [1] NA
```

Polish Bankruptcy

```
# test err
ytrue = data_test$bankrupt
phat = predict(fit,data_test,type="response")
yhat = as.numeric(phat > 0.5)
table(ytrue,yhat)

##      yhat
## ytrue    0    1
##      0 1482   85
##      1    60   61

err = 1-mean(yhat==ytrue)
err

## [1] 0.08590047
```

Polish Bankruptcy

```
#####
# Classification Tree #
#####

require(rpart)
fit0 = rpart(bankrupt~, data=data_train, control=rpart.control(cp=0))
fit = prune(fit0, cp=fit0$cptable[which.min(fit0$cptable[, "xerror"]),"CP"])

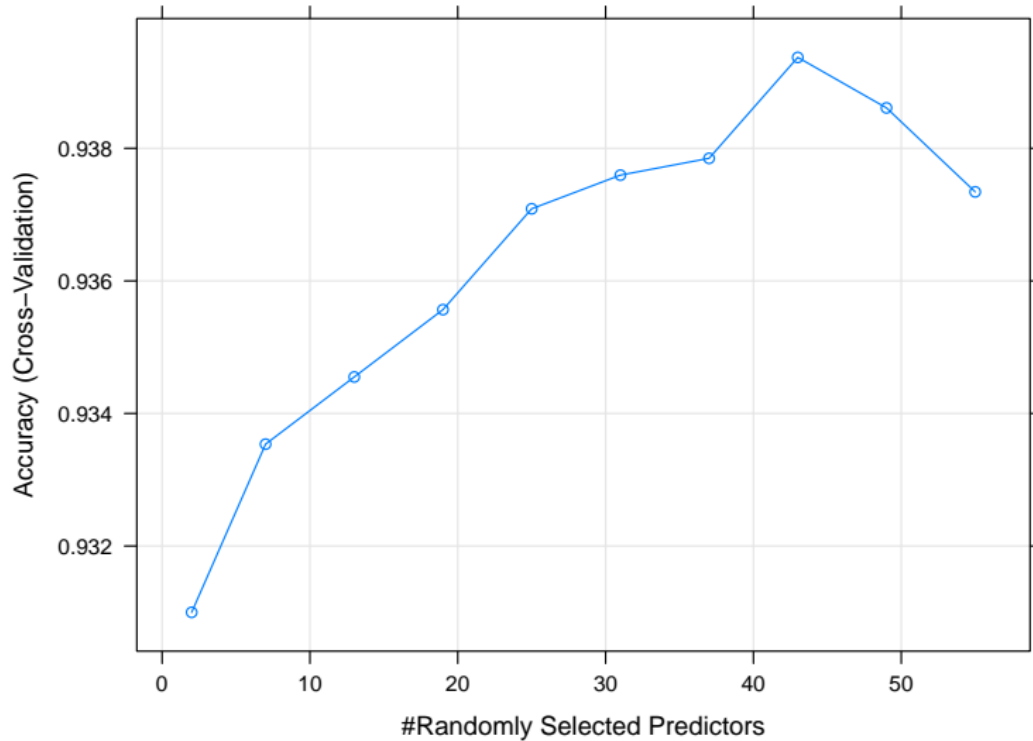
# test err
yhat = predict(fit, data_test, type="class")
err = 1-mean(yhat==ytrue)
err

## [1] 0.07227488
```

Polish Bankruptcy

```
#####
# Random Forest #
#####
# here we can use cross validation to select the best mtry
require(caret)
fit = train(bankrupt~, data=data_train, method="rf",
            trControl=trainControl(method="cv"),
            tuneLength=10) #tuneLength: number of mtry to try
```

Polish Bankruptcy



Polish Bankruptcy

```
fit$bestTune

##    mtry
## 8    43

#
# test err
yhat = predict(fit,data_test) # predict using fit$bestTune
err = 1-mean(yhat==ytrue)
err

## [1] 0.06042654
```

Polish Bankruptcy

```
#####
# Boosting #
#####
require(gbm)
data_boost = transform(data_train,bankrupt=as.numeric(bankrupt)-1)
fit = gbm(bankrupt~.,data=data_boost,distribution="adaboost",
          n.trees=5000,
          interaction.depth=10,
          shrinkage=0.1) # tuning parameters selected by cv
```

```
# test error
phat = predict(fit,data_test,n.trees=5000,type="response")
yhat = as.numeric(phat>0.5)
err <- 1-mean(yhat==ytrue)
err

## [1] 0.0521327
```

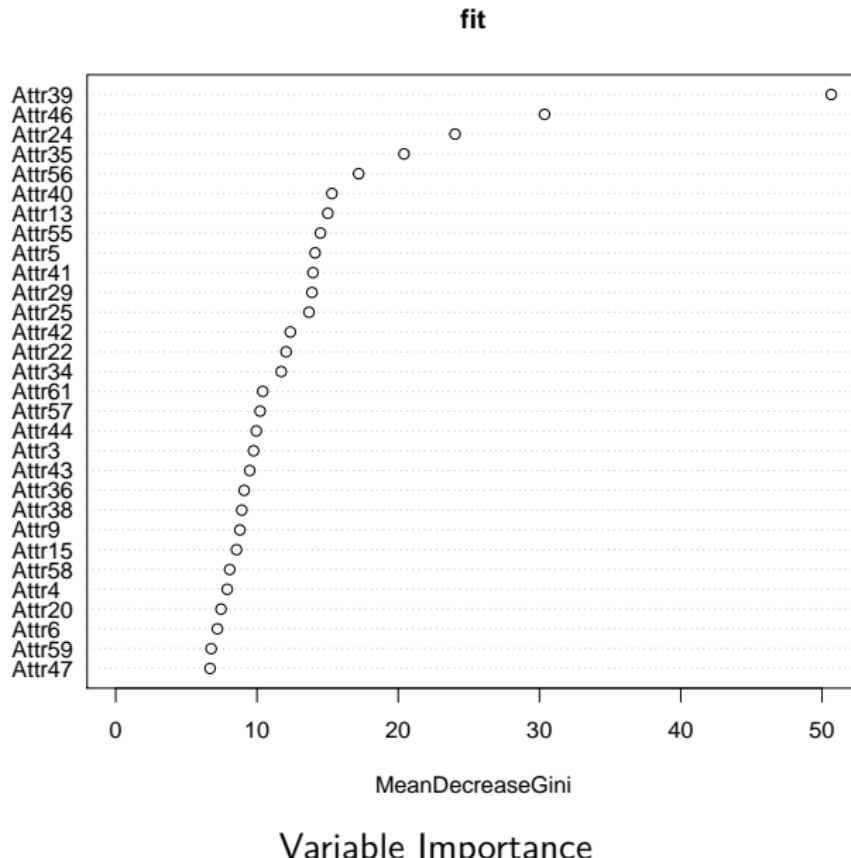
Polish Bankruptcy

Bagging (random forest) and boosting typically result in improved accuracy over prediction using a single tree, but the improved accuracy comes at the expense of interpretability.

To help better understand their results, we can calculate, for each variable in the model, its

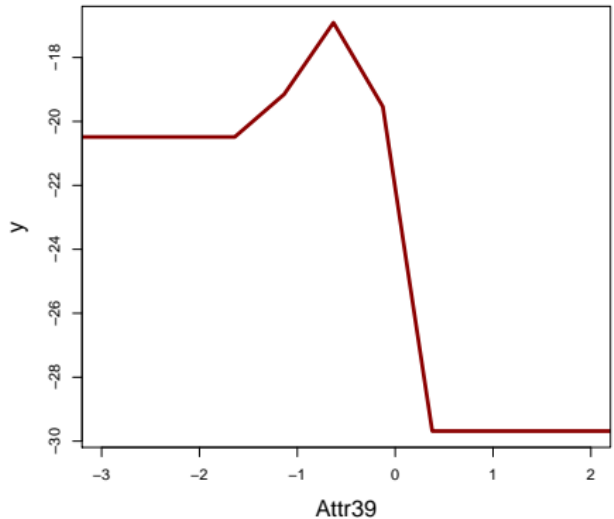
- **Variable importance:** how much in-sample error is decreased due to splits over the variable, averaged over all trees in the ensemble.
- **Partial dependence:** the *marginal* effect of the variable on y after integrating out all the other variables.

Polish Bankruptcy

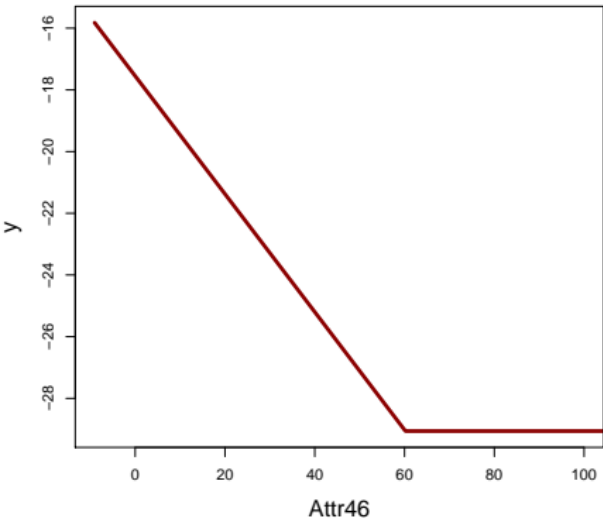


Polish Bankruptcy

Operating Profit Margin



Quick Ratio



Partial Dependence

Boston Housing Price

Now let's predict median house prices based on all variables in the data set:

```
# create training & test sets
train = sample(nrow(Boston), nrow(Boston)*0.6)
data_train = Boston[train,]
data_test = Boston[-train,]

#####
# Linear Regression #
#####

require(AER)
fit = lm(medv~.,data_train)

# test error
ytrue = data_test[, "medv"] #true medv on test data
yhat = predict(fit,data_test)
mean((yhat-ytrue)^2)

## [1] 21.89855
```

Boston Housing Price

```
coeftest(fit)

##
## t test of coefficients:
##
##              Estimate Std. Error t value Pr(>|t|)    
## (Intercept) 36.5105043  6.7569471  5.4034 1.369e-07 ***
## crim        -0.1437336  0.0389463 -3.6906 0.0002674 ***
## zn          0.0433694  0.0209236  2.0728 0.0390814 *  
## indus       -0.0461247  0.0867671 -0.5316 0.5954170  
## chas        3.9607844  1.0902345  3.6330 0.0003315 *** 
## nox         -19.1640319 4.9886347 -3.8415 0.0001503 *** 
## rm          3.5954278  0.5718327  6.2876 1.188e-09 *** 
## age         0.0179551  0.0185882  0.9659 0.3348840  
## dis         -1.3856486  0.2662350 -5.2046 3.694e-07 *** 
## rad         0.3356831  0.0891520  3.7653 0.0002015 *** 
## tax        -0.0085282  0.0053658 -1.5894 0.1130717  
## ptratio     -0.8464195  0.1763061 -4.8009 2.538e-06 *** 
## black       0.0059278  0.0037670  1.5736 0.1166706  
## lstat      -0.6189374  0.0666922 -9.2805 < 2.2e-16 *** 
## ---
```

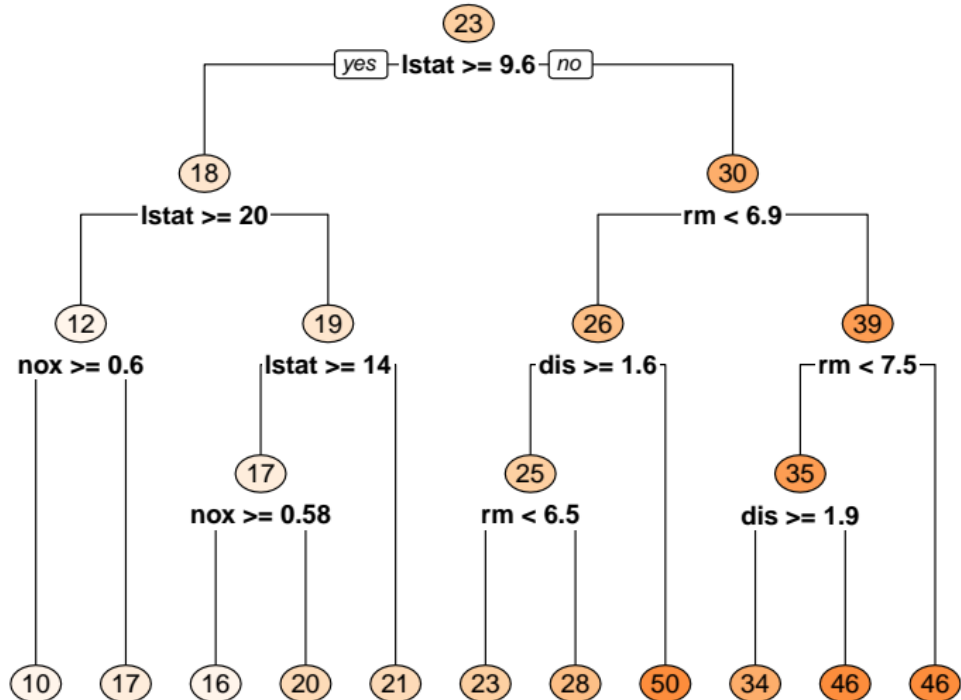
Boston Housing Price

```
#####
# Regression Tree #
#####
fit0 = rpart(medv~.,data=Boston,subset=train,
              control=rpart.control(cp=0,minbucket=2))
fit = prune(fit0,cp=fit0$cptable[which.min(fit0$cptable[, "xerror"]),"CP"])

# test error
yhat = predict(fit,data_test)
mean((yhat-ytrue)^2)

## [1] 17.34199
```

Boston Housing Price



Boston Housing Price

```
#####
# Bagging #
#####
require(randomForest)
fit = randomForest(medv~.,data_train,mtry=13,importance =TRUE)

# test error
yhat = predict(fit,data_test)
mean((yhat-ytrue)^2)

## [1] 10.99368
```

Boston Housing Price

```
#####
# Random Forest #
#####

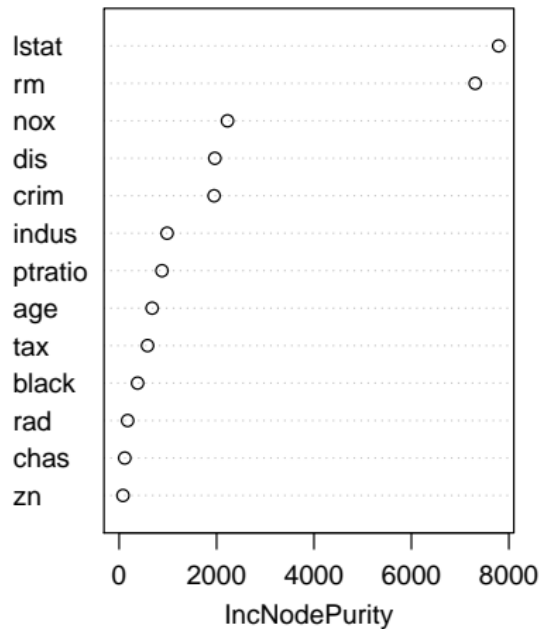
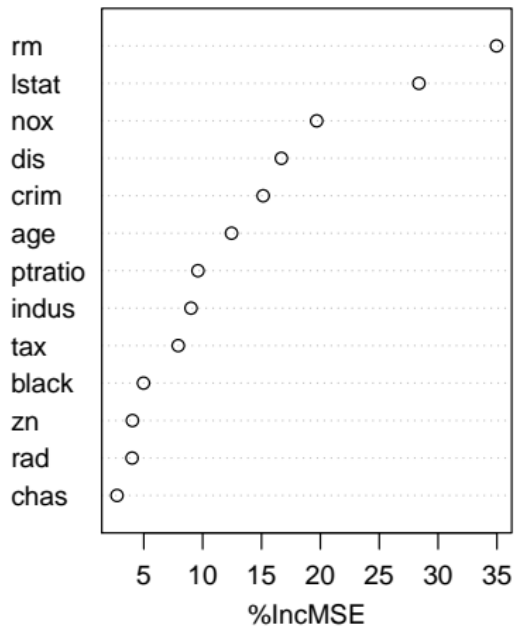
fit = randomForest(medv~.,data_train,mtry=5, # mtry selected via cv
                     importance =TRUE)

# test error
yhat = predict(fit,data_test)
mean((yhat-ytrue)^2)

## [1] 8.788307
```

Boston Housing Price

Variable Importance Plot



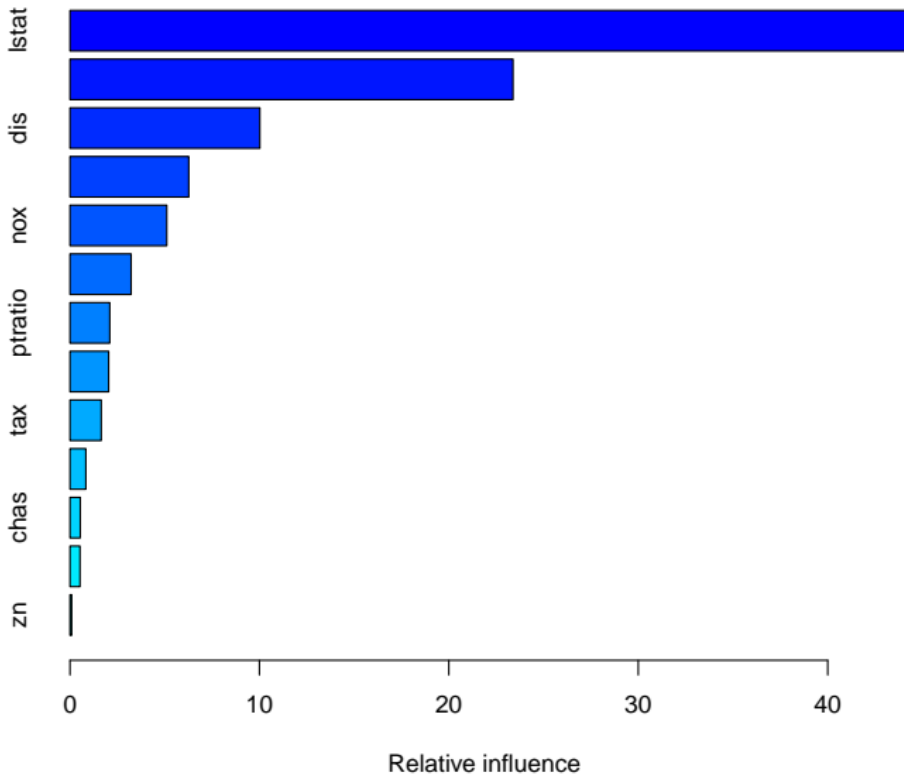
Boston Housing Price

```
#####
# Boosting #
#####
require(gbm)
fit = gbm(medv~.,data_train,distribution="gaussian",
           n.trees=10000,
           interaction.depth=5,
           shrinkage=0.001) # tuning parameters selected by cv

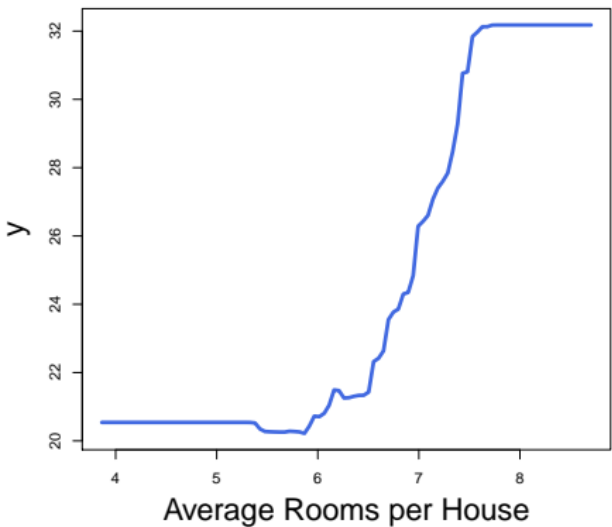
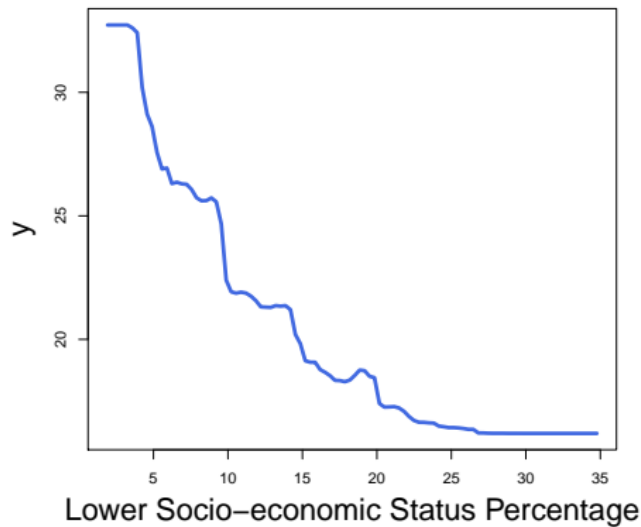
# test error
yhat = predict(fit,data_test,n.trees=10000)
mean((yhat-ytrue)^2)

## [1] 9.069948
```

Boston Housing Price



Boston Housing Price



Appendix: AdaBoost as Forward Stagewise Modeling

Let $y \in \{-1, 1\}$. Start with the forward stagewise procedure outlined on [page 75](#), using the exponential loss function $\ell(y, f(x)) = \exp(-yf(x))$.

In step m , given $\hat{f}^{(m-1)}(x)$ – the result of the previous $m - 1$ steps, the goal is find

$$(\hat{\alpha}_m, \hat{\phi}_m) = \arg \min_{\alpha, \phi} \sum_{i=1}^N \exp \left[-y_i (\hat{f}^{(m-1)}(x) + \alpha \phi(x_i)) \right] \quad (13)$$

$$= \arg \min_{\alpha, \phi} \sum_{i=1}^N w_i^{(m)} \exp [-\alpha y_i \phi(x_i)] \quad (14)$$

, where $w_i^{(m)} = \exp (-y_i \hat{f}^{(m-1)}(x))$.

Appendix: AdaBoost as Forward Stagewise Modeling

Minimizing (14) \Rightarrow

$$\hat{\alpha}_m = \frac{1}{2} \log \frac{1 - \epsilon_m}{\epsilon_m}$$

, where

$$\epsilon_m = \frac{\sum_{i=1}^N w_i^{(m)} \mathcal{I}(y_i \neq \hat{\phi}_m(x_i))}{\sum_{i=1}^N w_i^{(m)}}$$

, and

$$\hat{\phi}_m = \arg \min_{\alpha, \phi} \sum_{i=1}^N w_i^{(m)} \mathcal{I}(y_i \neq \phi(x_i))$$

, i.e. $\hat{\phi}_m$ is the function that minimizes the weighted misclassification error ϵ_m .

Appendix: AdaBoost as Forward Stagewise Modeling

We can then update $\hat{f}^{(m)}(x) = \hat{f}^{(m-1)}(x) + \hat{\alpha}_m \hat{\phi}_m(x)$, and

$$\begin{aligned} w_i^{(m+1)} &= \exp\left(-y_i \hat{f}^{(m)}(x_i)\right) \\ &= \exp\left(-y_i \left[\hat{f}^{(m-1)}(x_i) + \hat{\alpha}_m \hat{\phi}_m(x_i)\right]\right) \\ &= w_i^{(m)} \exp\left(-y_i \hat{\alpha}_m \hat{\phi}_m(x_i)\right) \\ &= w_i^{(m)} \exp(2\hat{\alpha}_m \mathcal{I}(y_i \neq \phi(x_i))) \cdot \exp(-\hat{\alpha}_m) \end{aligned}$$

, where we use the fact that $-y_i \hat{\phi}_m(x_i) = 2\mathcal{I}(y_i \neq \phi(x_i)) - 1$.

Therefore, we can multiply $\hat{\alpha}_m$ by 2, i.e., let $\hat{\alpha}_m = \log \frac{1-\epsilon_m}{\epsilon_m}$ and update w_i as

$$w_i^{(m+1)} = w_i^{(m)} \exp(\hat{\alpha}_m \mathcal{I}(y_i \neq \phi(x_i)))$$

This gives us the AdaBoost algorithm.

Acknowledgement I

Part of this lecture is adapted from the following sources:

- Géron, A. 2017. *Hands-On Machine Learning with Scikit-Learn and TensorFlow*. O'Reilly Media.
- Hastie, T., R. Tibshirani, and J. Friedman. 2008. *The Elements of Statistical Learning* (2nd ed.). Springer.
- James, G., D. Witten, T. Hastie, and R. Tibshirani. 2013. *An Introduction to Statistical Learning: with Applications in R*. Springer.
- Sontag, D. *Introduction To Machine Learning*. Lecture at NYU, retrieved on 2018.01.01. [[link](#)]
- Taddy, M. *Big Data*. Lecture at the University of Chicago Booth School of Business, retrieved on 2017.01.01. [[link](#)]
- Tibshirani, R. *Data Mining*, Lecture at Carnegie Mellon University, retrieved on 2017.01.01. [[link](#)]

Acknowledgement II

- Van der Schaar, M. and S. Flaxman. *Statistical Machine Learning*. Lecture at Oxford University, retrieved on 2018.01.01. [[link](#)]

Reference



Dietterich, T. G. 2000. "Ensemble Methods in Machine Learning," In: Multiple Classifier Systems. MCS 2000. Lecture Notes in Computer Science, vol 1857. Springer.