

non parametric
supervised methods.

Chapter 8: Tree-Based Methods

quantitative
categorical

We will introduce *tree-based* methods for regression and classification.

These involve segmenting the predictor space into a number of simple regions

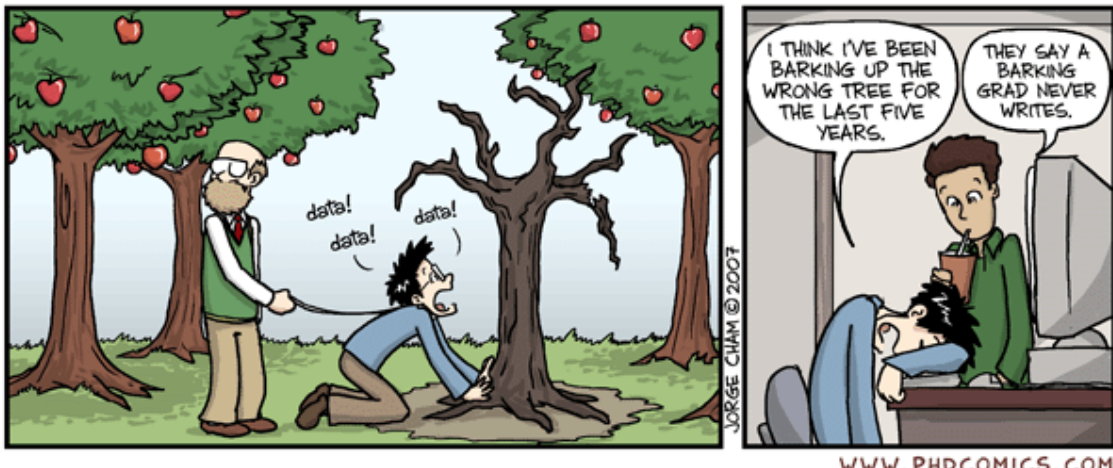
To make a prediction for an observation, we will use the mean or mode of the training observations in the regions to which it belongs.

The set of splitting rules can be summarized in a tree ⇒ “decision trees”.

- simple and useful for interpretation
- not competitive w/ other supervised approaches (eg. lasso) for prediction.

bagging, random forests,
boosting.

Combining a large number of trees can often result in dramatic improvements in prediction accuracy at the expense of interpretation.



Credit: <http://phdcomics.com/comics.php?f=852>

Decision trees can be applied to both regression and classification problems. We will start with regression.

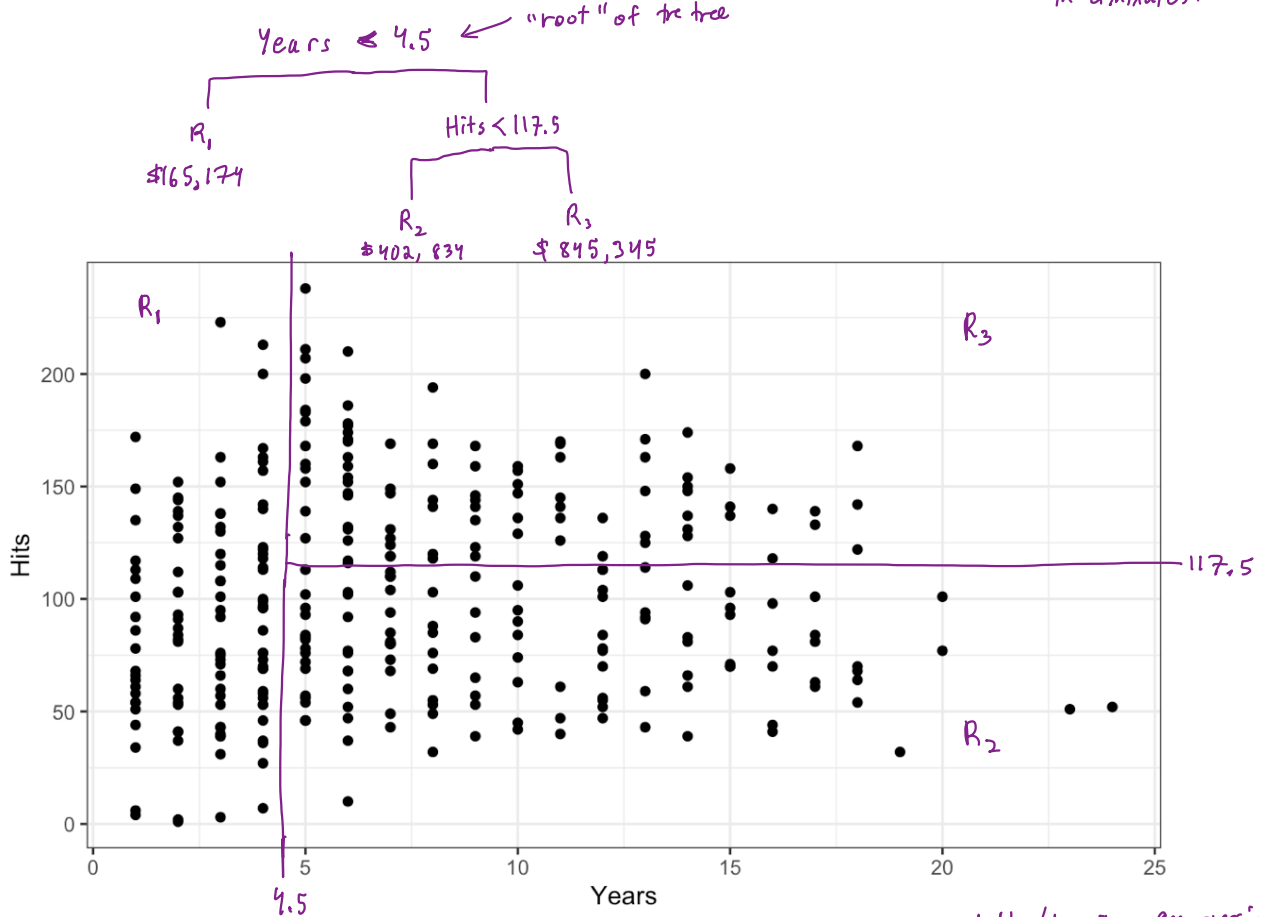
1 Regression Trees

Start with

Example: We want to predict baseball salaries using the `Hitters` data set based on `Years` (the number of years that a player has been in the major leagues) and `Hits` (the number of hits he made the previous year).

We could make a series of splitting rules to create regions and predict salary as the mean in each region.

according to a tree fit to this data (more on how in a minute).



The predicted salary for players is given by the mean response value for the players in that box. Overall, the tree segments the players into 3 regions of predictor space.

probably / definitely an oversimplification
but easy to interpret and has a nice graphical representation.

terminology: R_1, R_2, R_3 = terminal nodes or leaves of the tree

points along the tree where space is split = internal nodes

segments of tree that connect nodes = branches.

interpretation: Years is the most important factor in determining salary

2

↳ given that a player is experienced, # hits in previous year plays a role in his salary: $\text{Hits} \uparrow \text{Salary}$.

↳ given that a player is not experienced, # hits does not affect your salary.

quantitative y

\nearrow

We now discuss the process of building a regression tree. There are two steps:

\nearrow set of possible values for X_1, \dots, X_p

1. Divide predictor space

into J distinct and non-overlapping regions R_1, \dots, R_J

2. Predict

For every observation that falls into region R_j we make the same prediction,
the mean of the response Y for training values in R_j

How do we construct the regions R_1, \dots, R_J ? How to divide the predictor space?

regions could have any shape: but that is too hard (to do & interpret)

\Rightarrow divide predictor space into high dimensional rectangles or boxes

The goal is to find boxes R_1, \dots, R_J that minimize the RSS. $= \sum_{j=1}^J \sum_{i \in R_j} (y_i - \hat{y}_{R_j})^2$ where \hat{y}_{R_j} = mean response
of training data
in box R_j

Unfortunately it is computationally infeasible to consider
every possible partition.

\Rightarrow take top-down, greedy approach called recursive binary splitting

The approach is *top-down* because

We start at the top of the tree (where all observations belong to a single region)
and successively split the predictor space.

\hookrightarrow each split is indicated via two new branches in the tree.

The approach is *greedy* because

at each step of the building process, the best split is made at that particular step.

\hookrightarrow not looking ahead to make a split that will lead to a better tree later.

In order to perform recursive binary splitting,

① Select predictor and cut point s.t. splitting the predictor space in to regions

$\{X: X_j < s\}$ and $\{X: X_j \geq s\}$ leads to greatest possible reduction in RSS.
 ↗ region of predictor space where X_j takes values $< s$

↳ We consider all possible X_1, \dots, X_p and all possible cut points then choose predictor & cutpoint so tree has lowest RSS.

i.e. consider all possible half planes $R_1(j, s) = \{X | X_j < s\}$, $R_2(j, s) = \{X | X_j \geq s\}$.
 we seek j and s that minimize $\sum_{i: x_i \in R_1(j, s)} (y_i - \hat{y}_{R_1})^2 + \sum_{i: x_i \in R_2(j, s)} (y_i - \hat{y}_{R_2})^2$ ← can be quickly done if p not too large.

② Repeat process looking for next best j, s combo, but instead of splitting entire space, we will split $R_1(j, s)$ and $R_2(j, s)$ to minimize RSS.

③ Continue until stopping criteria is met (i.e. no region contains more than 5 observations).

The process described above may produce good predictions on the training set, but is likely to overfit the data.

because our resulting tree will be too complex.
 ↗ less regions.

A smaller tree, with less splits might lead to lower variance and better interpretation at the cost of a little bias.

Idea: Only split tree if it results in a large enough drop in RSS.

↑
bad idea because a seemingly worthless split early in tree can be followed by a good split!

Better idea: A strategy is to grow a very large tree T_0 and then prune it back to obtain a subtree.

How to prune the tree?

goal: Select a subtree that leads to lowest test error rate → could use CV to estimate error for every possible subtree, but this is expensive (large # of possible subtrees).

Solution: "cost complexity pruning", aka "weakest link pruning"

Consider a sequence of trees indexed by a nonnegative tuning parameter α

For each value of α , \exists a corresponding subtree $T \subset T_0$ st.

$\sum_{m=1}^{|T|} \sum_{i: x_i \in R_m} (y_i - \hat{y}_{R_m})^2 + \alpha |T|$ is as small as possible
 ↗ # of terminal nodes in the tree.

$R_m = m^{\text{th}}$ terminal node region

\hat{y}_{R_m} = predicted response for R_m

α controls trade off between subtree's complexity & fit to training data

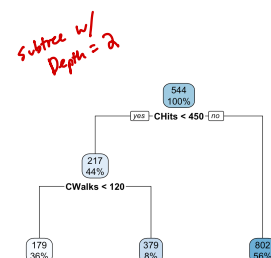
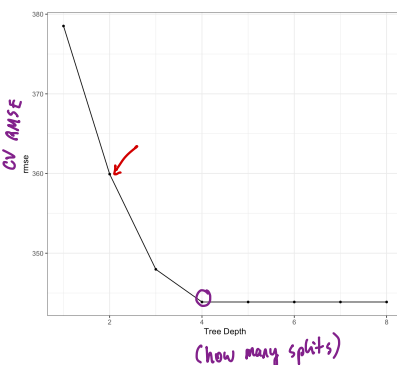
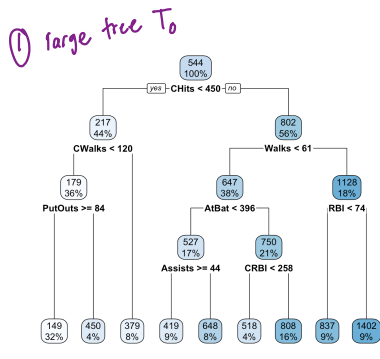
Select α via CV, then use full training dataset & chosen α to get subtree.

When $\alpha > 0$
 ↗ T = T_0
 $\alpha \uparrow \Rightarrow$ price to pay for having more terminal nodes ↑
 \Rightarrow smaller tree.

Algorithm for building a regression tree:

- ① Use recursive binary splitting to grow a large tree on training data set, stopping when each terminal node has fewer than some minimum # of observations
- ② Apply cost complexity pruning to the large tree to get a sequence of best trees as a function of α .
- ③ Use k-fold CV to choose α .
 - Divide training data into K folds, for each $k=1, \dots, K$
 - (a) Repeat steps ① and ② on all but k^{th} fold
 - (b) Evaluate MSE on data in k^{th} fold
 - Average results for each value of α and pick α to minimize CV error.
- ④ Return the subtree from ② that corresponds to α from ③.

Example: Fit regression Tree to Hitters using 9 features \rightarrow 50% train/test split.



2 Classification Trees

A *classification tree* is very similar to a regression tree, except that it is used to predict a categorical response.

Recall from regression trees, predicted response for an observation is given by the mean response of training obs. in that region.

For a classification tree, we predict that each observation belongs to the most commonly occurring class of training observation in the region to which it belongs.

the mode

We are often also interested in class proportions that fall into each terminal node.

↳ this can give us some idea of how reliable the prediction is

e.g. terminal node w/ 100% class 1 vs. 55% class 1
45% class 2.

The task of growing a classification tree is quite similar to the task of growing a regression tree.

Use binary recursive splitting to grow a classification tree
but RSS cannot be used as criterion for splitting.

Instead natural alternative is classification error rate

= fraction of training obs that do not belong to most common class

$$= 1 - \max_k (\hat{p}_{mk})$$

It turns out that classification error is not sensitive enough for tree growing.

Preferred measures:

$$\textcircled{1} \text{ Gini Index } G = \sum_{k=1}^K \hat{p}_{mk} (1 - \hat{p}_{mk}) \text{ measure of total variance across K classes.}$$

↳ takes small values if all \hat{p}_{mk} 's are close to 0 or 1. \Rightarrow measure of "node purity"
 $\downarrow G \Rightarrow$ nodes contain observations primarily from one class.

$$\textcircled{2} \text{ Entropy } D = - \sum_{k=1}^K \hat{p}_{mk} \log \hat{p}_{mk}$$

↳ will take values near zero if \hat{p}_{mk} 's close to 0 or 1. $\Rightarrow \downarrow D \Rightarrow$ nodes more "pure"

Gini and Entropy are actually quite similar numerically

When building a classification tree, either the Gini index or the entropy are typically used to evaluate the quality of a particular split.

Any of the 3 methods can also be used for pruning.

If prediction accuracy of final tree is the goal,

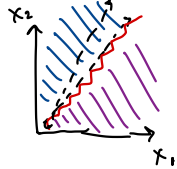
classification error rate should be used.

Note:
neither Gini nor
entropy works
well w/ unbalanced
data. There
are other options
but we'll
split on.

3 Trees vs. Linear Models

Regression and classification trees have a very different feel from the more classical approaches for regression and classification.

e.g. linear regression: $f(x) = \beta_0 + \sum_{j=1}^p x_j \beta_j$



regression trees : $f(x) = \sum_{m=1}^M c_m \mathbb{I}(x \in R_m)$. where R_1, \dots, R_m is a partition of the predictor space

Which method is better? It depends on the problem

- If the relationship b/w features and response is approximately linear, then a linear model will but perform a tree.
- If the relationship is highly non-linear, decision tree may be better.

Trees have interpretation / visualization.

3.1 Advantages and Disadvantages of Trees

Advantages

- easy to explain, even easier than linear regression.
- (?) - some people think decision trees more closely mirror human decision making.
- can be displayed graphically (good for non-experts)
- can handle categorical predictors without needing to make dummy variables.

Disadvantages

- do not have same level of predictive performance as other methods we have seen.
- Not robust: small change in data can have large effect on fitted tree.
(high variability).



We can aggregate many trees to try and improve this!

4 Bagging

"Bootstrap Aggregation"

4 Bagging

Decision trees suffer from *high variance*.

i.e. if we split data in half randomly, fit decision tree to both halves, resulting trees could be quite different.

vs. low variance will yield similar results if applied repeatedly to different samples from same population.
 ↳ linear regression when $n \gg p$.

Bootstrap aggregation or *bagging* is a general-purpose procedure for reducing the variance of a statistical learning method, particularly useful for trees.

Recall: For a given set of n independent observations Z_1, \dots, Z_n each w/ variance $\sigma^2 < \infty$.

$$\text{Var}(\bar{Z}) = \text{Var}\left(\frac{1}{n} \sum_{i=1}^n Z_i\right) \stackrel{\text{indp}}{=} \frac{1}{n^2} \sum_{i=1}^n \text{Var} Z_i = \frac{1}{n^2} \sum_{i=1}^n \sigma^2 = \frac{\sigma^2}{n}$$

i.e. averaging a set of observations reduces variance.

So a natural way to reduce the variance is to take many training sets from the population, build a separate prediction model using each training set, and average the resulting predictions.

i.e. take B training sets.

calculate $\hat{f}^1(x), \dots, \hat{f}^B(x)$.

obtain low-variance statistical learning model:

$$\hat{f}_{\text{AVG}}(x) = \frac{1}{B} \sum_{b=1}^B \hat{f}^b(x).$$

Of course, this is not practical because we generally do not have access to multiple training sets. *Collecting training data can be expensive*.

Instead we could take repeated samples (w/ replacement) from training data set.

(these are called "bootstrapped" training data sets because we are "pulling ourselves up by our bootstraps").

↳ assumes empirical dsn in sample is similar to population dsn, i.e. have representative sample.

Then we could ^{fit} our method on b^{th} bootstrapped training data set to get

$\hat{f}^{*(b)}(x)$ and average

$$\hat{f}_{\text{bag}}(x) = \frac{1}{B} \sum_{b=1}^B \hat{f}^{*(b)}(x).$$

While bagging can improve predictions for many regression methods, it's particularly useful for decision trees.

- To apply the regression trees :
- (1) construct B trees using bootstrapped data sets.
 - (2) average resulting predictions.

These trees are grown deep and not pruned.

How can bagging be extended to a classification problem?

4.1 Out-of-Bag Error

There is a very straightforward way to estimate the test error of a bagged model, without the need to perform cross-validation.

4.2 Interpretation

5 Random Forests

Random forests provide an improvement over bagged trees by a small tweak that decorrelates the trees.

As with bagged trees, we build a number of decision trees on bootstrapped training samples.

In other words, in building a random forest, at each split in the tree, the algorithm is not allowed to consider a majority of the predictors.

The main difference between bagging and random forests is the choice of predictor subset size m .

6 Boosting

Boosting is another approach for improving the prediction results from a decision tree.

While bagging involves creating multiple copies of the original training data set using the bootstrap and fitting a separate decision tree on each copy,

Boosting does not involve bootstrap sampling, instead each tree is fit on a modified version of the original data set.

Boosting has three tuning parameters:

1.

2.

3.