#### Tree-Based Methods

Chapter 8

November 16, 2016

1 8.1 The basics of decision trees.

2 8.2 Bagging, random forests and boosting

#### About this chapter

- Decisions trees: splitting each variable sequentially, creating rectugular regions.
- Making fitting/prediction locally at each region.
- It is intuitive and easy to implement, may have good interpretaion.
- Generally of lower prediction accuracy.
- Bagging, random forests and boosting ... make fitting/prediction based on a number of trees.
- Bagging and Boosting are general methodologies, not just limited to trees.

### Regression trees

- Trees can be applied to both regression and classification.
- CART refers to classification and regression trees.
- We first consider regression trees through an example of predicting Baseball players' salaries.

#### The Hitters data

- Response/outputs: Salary.
- Covarites/Inputs:
   Years (the number of years that he has played in the major leagues)
  Hits (the number of hits that he made in the previous year).
- preparing data: remove the observations with missing data and log-transformed the Salary (preventing heavy right-skewness)

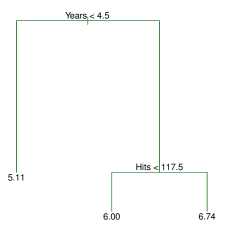


Figure: 8.1. Next page

Figure 8.1. For the Hitters data, a regression tree for predicting the log salary of a baseball player, based on the number of years that he has played in the major leagues and the number of hits that he made in the previous year. At a given internal node, the label (of the form  $X_i < t_k$ ) indicates the left-hand branch emanating from that split, and the right-hand branch corresponds to  $X_i \geq t_k$ . For instance, the split at the top of the tree results in two large branches. The left-hand branch corresponds to Years < 4.5, and the right-hand branch corresponds to Years  $\geq 4.5$ . The tree has two internal nodes and three terminal nodes, or leaves. The number in each leaf is the mean of the response for the observations that fall there.

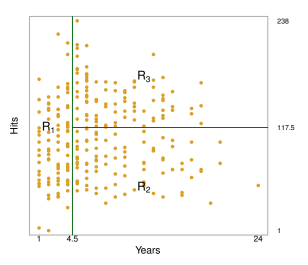


Figure: 8.2. The three-region partition for the Hitters data set from the regression tree illustrated in Figure 8.1.

# Estimation/prediction

- On Regions  $R_1$ ,  $R_2$ ,  $R_3$ , the mean-log-salary is 5.107, and 6.74.
- Our prediction for any players in  $R_1$ ,  $R_2$  and  $R_3$  are, respectively  $1000 \times e^{5.107} = \$165, 174, 1,000 \times e^{5.999} = \$402, 834,$  and  $1.000 \times e^{6.740} = \$845, 346.$

# Estimation/prediction

- Trees involve a series of splittings of the data, each time by one variable.
- The series of actions taken place sequentially creates a tree-like results.
- As in Figure 8.1, the terminal nodes are the three indexed by the numbers, which represent the regions  $R_1$ ,  $R_2$  and  $R_3$ . These regions constitute he final partiation of the data.
- Terminal nodes are also called leaves.
- Each internal node represents a splitting,
- In Figure 8.1, the two internal nodes are indexed by Y < 4.5 and Hits < 117.5.
- The lines connecting nodes are called branches.
- Trees are typically drawn upside down.

### Two step towards prediction

- Run the splitting according to input values sequentially, and obtain final partition of the data in regions  $R_1, ..., R_J$ .
- For any new observation with covariates in region  $R_k$ , we predict its response by the average of the reponses of the data points in region  $R_k$ .

#### How to split

- Suppose we wish to partition a region R. In other words, we wish to separate the data in region R into two parts, day  $R_1$  and  $R_2$ , according to one input values.
- What would be the optimal or efficient split in some sense?
- Only two flexibility in the split: 1. Choice of the input variable to split, 2. the cutpoint of the split of that chose input.
- Imagine that this is the final split of R:  $R_1$  and  $R_2$  would be leaves.
  - And we would use the mean response of data in  $R_1$  and  $R_2$  to predict the response of any new/old observations.
  - We wish our choice of  $R_1$  and  $R_2$  would be optimal in the sense of achieving miminum prediction error on the training data in region R.

### Recursive binary splitting

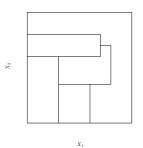
• A greedy algorithm (geedy means it is optimal at the current step): For j=1,...,p and all real value s, let  $R_1(j,s)=\{i\in R: X_j< s\}$  and  $R_2(j,s)=\{i\in R: X_j\geq s\}$ . And let  $\hat{y}_1$  and  $\hat{y}_2$  be the mean response of all observations in  $R_1(j,s)$  and  $R_2(j,s)$ , respectively. Consider the following prediction error:

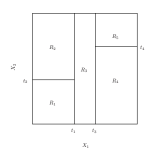
$$RSS_{new} = \sum_{i \in R_1(j,s)} (y_i - \hat{y}_1)^2 + \sum_{i \in R_2(j,s)} (y_i - \hat{y}_2)^2$$

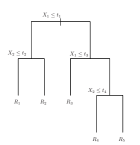
Choose the split which has the smallest prediction error. This split is the optimal one, denoted as  $R_1$  and  $R_2$ .

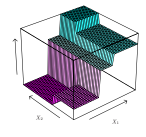
• Continue the split till the final partition.

#### 8.1 The basics of decision trees.









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Figure 8.3. Top Left: A partition of two-dimensional feature space that could not result from recursive binary splitting. Top Right: The output of recursive binary splitting on a two-dimensional example. Bottom Left: A tree corresponding to the partition in the top right panel. Bottom Right: A perspective plot of the prediction surface corresponding to that tree.

# When to stop split

- The problem of when to stop.
- If too many steps of splitting: many leaves, too complex model, small bias but large variance, may overfit.
- If too few steps of splitting: few leaves, too simple model, large bias but small variance, may underfit.

#### One natural idea

• When splitting R into  $R_1$  and  $R_2$ , consider the RSS before the split

$$RSS_{old} = \sum_{i \in R} (y_i - \hat{y})^2$$

where  $\hat{y}$  is the average of the response of data in R. With the optimal split, the reduction of RSS is

$$RSS_{old} - RSS_{new}$$

- We can pre-choose a threshold, h, and decide the worthiness of the split.
- If the reduction is smaller than h, we do not do it, and stop right there; then R is one terminal node (a leave).
- If the reduction is greater than h, we make the split, and continue with next step.

#### One natural idea

- The idea is seemingly reasonable, but is too near-sighted.
- Only look at the effect of the current split.
- It is possible that even if the current split is not effective, the future splits could be effective and, maybe, very effective.

### Tree pruning

- Grow a very large tree.
- Prune the true back to obtain a subtree.
- Objective: find the subtree that has the best test error.
- Cannot use cross-validation to examine the test errors for all possible subtrees, since there are just too many.
- Even if we can, this would probably be overfitting, since model space is too large.

# Cost complexity pruning

- Let  $T_0$  be the original (large) tree. Let T be any subtree. Use  $|T_0|$  and |T| to denote their numbers of teminal nodes, which represent complexity.
- Consider "Loss + Penalty":

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

where  $R_m$  are the terminal nodes of the subtree T, and the mean response of  $R_m$  is  $\hat{y}_m$ ;  $\alpha$  is tuning parameter.

- Denote the minimized subtree as  $T_{\alpha}$ .
- If  $\alpha = 0$ , no penalty the optimal tree is the original  $T_0$ .
- If  $\alpha = \infty$ , the tree has no split at all. The predictor is just  $\bar{y}$ .
- The larger the  $\alpha$ , the more penalty for model complexity.

### Cost complexity pruning

- Just like Lasso, there exists efficient computation algorithm to compute the entire sequence of  $T_{\alpha}$  for all  $\alpha$ .
- Use cross-validation to find the best  $\alpha$  to minimize the test error.

### The algorithm

- 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  $\alpha$ .

#### The algorithm

- 3. Use K-fold cross-validation to determine best  $\alpha$ . That is, divide the training observations into K folds. For each k=1,...,K
  - (a) Repeat Steps 1 and 2 on all but the kth 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  $\alpha$ .
  - (c) Average the results for each value of  $\alpha$ , and pick  $\alpha$  to minimize the average error.
- 4. Return the subtree from Step 2 that corresponds to the chosen value of  $\alpha$ .

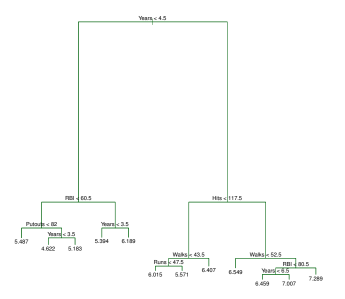


Figure: 8.4. Regression tree analysis for the Hitters data. The unpruned tree that results from top-down greedy splitting on the training data is shown.

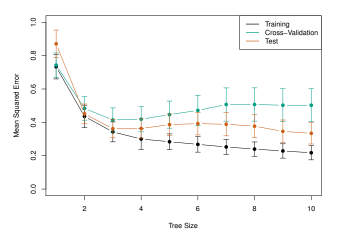


Figure: 8.5. Regression tree analysis for the Hitters data. The training, cross-validation, and test MSE are shown as a function of the number of terminal nodes in the pruned tree. Standard error bands are displayed. The minimum cross-validation error occurs at a tree size of three.

#### Classification trees

- Regression has numerical responses; and classification has qualitative responses.
- Recall that for regression trees, we chose to obtain the greatest reduction of RSS.
  - RSS is using sum of squares to measure the error.
- For classification trees, one can follow the same line of procedure as that of regression trees, but using error measurements that are more appropriate for classification.

#### Classification error rates

- For a region R, let  $\hat{p}_k$  be the percentage of observations in this region that belong to class k.
- We assign any new observation in region R as from the class with largest  $\hat{p}_k$ , which is the so-called most commonly occurring class in training data.

#### The impurity measure

• The classification error rate (for this region R) is

$$E = 1 - \max_k \hat{p}_k.$$

• The Gini index is

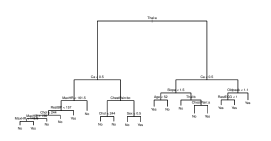
$$G = \sum_{k=1}^{K} \hat{p}_k (1 - \hat{p}_k)$$

• The cross-entropy is

$$D = -\sum_{k=1}^{K} \hat{p}_k \log(\hat{p}_k)$$

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- If R is nearly pure, most of the observations are from one class, then the Gini-index and cross-entropy would take smaller values than classification error rate.
- Gini-index and cross-entropy are more sentive to node purity.
- To evaluate the quality of a particluar split, the Gini-index and cross-entropy are more popularly used as error measurement crietria than classification error rate.
- Any of these three approaches might be used when pruning the tree.
- The classification error rate is preferable if prediction accuracy of the final pruned tree is the goal.



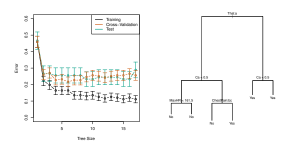


Figure 8.6. Heart data. Top: The unpruned tree. Bottom Left: Cross-validation error, training, and test error, for different sizes of the pruned tree. Bottom Right: The pruned tree corresponding to the minimal cross-validation error.

#### Trees vs. Linear models

• For regression model:

$$Y = f(X) + \epsilon$$

Linear model assumes

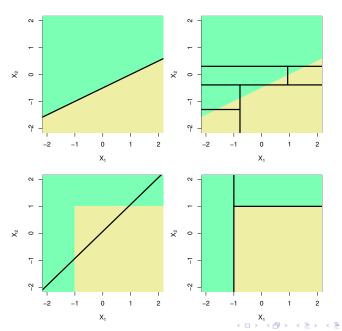
$$f(X) = \beta_0 + \sum_{j=1}^{p} X_j \beta_j$$

Regression trees assume

$$f(X) = \sum_{j=1}^{M} c_m 1(X \in R_m)$$

where  $R_1, ..., R_M$  are rectagular partitions of the input space.

• If the underlying realation is close to linear, linear model is better. Otherwise, regression trees are generally better. (Useless comments)



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Figure 8.7. Top Row: A two-dimensional classification example in which the true decision boundary is linear, and is indicated by the shaded regions. A classical approach that assumes a linear boundary (left) will outperform a decision tree that performs splits parallel to the axes (right). Bottom Row: Here the true decision boundary is non-linear. Here a linear model is unable to capture the true decision boundary (left), whereas a decision tree is successful (right).

# Advantages of Trees

- Trees are very easy to explain to people. In fact, they are even easier to explain than linear regression!
- Some people believe that decision trees more closely mirror human decision-making than do the regression and classification approaches seen in previous chapters.
- Trees can be displayed graphically, and are easily interpreted even by a non-expert (especially if they are small).
- Trees can easily handle qualitative predictors without the need to create dummy variables.

### Disadvantages of Trees

- Trees generally do not have the same level of predictive accuracy as some of the other regression and classification approaches seen in this book.
- Trees can be very non-robust. In other words, a small change in the data can cause a large change in the final estimated tree.
- However, by aggregating many decision trees, using methods like bagging, random forests, and boosting, the predictive performance of trees can be substantially improved. We introduce these concepts in the next section.

# Bagging (Boostrap Aggregating)

- A general purpose procedure to reduce variance of a learning method.
- A model averaging technique.
- Decision tree is generally a high variance method. (Apply the method based on different data based on same sampling scheme would lead to very different result.)
- Average of iid random variables would have a reduced variance  $\sigma^2/n$

## The procedure.

• Model

$$y_i = f(x_i) + \epsilon_i, \quad i = 1, ..., n.$$

- Suppose a statistical learning method gives  $\hat{f}(\cdot)$  based on the training data  $(y_i, x_i), i = 1, ..., n$ .
- For example,
  - 1 Linear model:  $\hat{f}(x) = \hat{\beta}_0 + \hat{\beta}^T x_i$
  - 2 KNN:  $\hat{f}(x) = \sum_{j=1}^{J} \bar{y}_{\tilde{R}_{j}}$  with least distance to K-cluster partition.
  - **3** Decision tree:  $\hat{f}(x) = \sum_{j=1}^{J} \bar{y}_{R_j}$  with rectangular partition.
  - **4** ...

### The procedure of Bagging

- Data  $(y_i, x_i), i = 1, ..., n$ ; and a learning method  $\hat{f}$
- Draw a boostrap sample from the data, and compute a  $\hat{f}_1^*$  based on this set of bootstrap sample.
- Draw another boostrap sample from the data, and compute a  $\hat{f}_2^*$  based on this set of bootstrap sample.
- ....
- Repeat M times, obtain  $\hat{f}_1^*, ...., \hat{f}_M^*$ .
- Produce the learning method with bagging as

$$\frac{1}{M} \sum_{i=1}^{M} \hat{f}_{i}^{*}$$

### The Bagging

- Bagging is general-purpose.
- It works best for high variance low bias learning methods.
- This is the case for decision trees, particularly deep trees.
- Also the case for large p.
- If the response is qualitative, we can take the majority vote (not averaging) of the predicted class based on all learning methods based on boostrap samples.

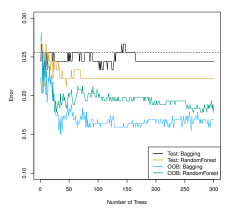


Figure: 8.8. Bagging and random forest results for the Heart data. The test error (black and orange) is shown as a function of B, the number of bootstrapped training sets used. Random forests were applied with  $m = \sqrt{p}$ . The dashed line indicates the test error resulting from a single classification tree. The green and blue traces show the OOB error, which in this case is considerably lower

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## Out-of-Bag (OOB) error estimation

- Estimation of test error for the bagged model.
- For each bootstrap sample, observation i is bootstrap sampled with probabilty  $(1 1/n)^n \approx 1/e$ .
- For each bootstrap sample, the number of observations not taken into this bootstrap sample is  $n(1-1/n)^n \approx n/e$ . These are referred to as out-of-bag (OOB) observations.
- For totally B bootstrap samples, about B/e times, the bootstrap sample does not contain observation i.
- The trees based on these bootstrap sample can be used to predict the response of observation i. Tatoally about B/e predictions.
- We average these predictions (for regression) or take majority vote (for classification) to produce the Bagged prediction for observation i, denote it as  $\hat{f}^*(x_i)$ .

### Out-of-Bag (OOB) error estimation

• The OOB MSE is

$$\sum_{i=1}^{n} (y_i - \hat{f}^*(x_i))^2$$

• The OOB classification error is

$$\sum_{i=1}^{n} I(y_i \notin \hat{f}^*(x_i))$$

- The resulting OOB error is a valid estimate of the test error for the bagged model, since the response for each observation is predicted using only the trees that were not fit using that observation.
- It can be shown that with B sufficiently large, OOB error is virtually equivalent to leave-one-out cross-validation error.

### Variable importance measures

- Bagging improves prediction accuracy at the expense of interpretability.
- An overall summary of the importance of each predictor using the RSS (for bagging regression trees) or the Gini index (for bagging classification trees).
- Bagging regression trees, we can record the total amount that the RSS is decreased due to splits over a given predictor, averaged over all B trees.
- A large value indicates an important predictor.
- Bagging classification trees, we can add up the total amount that the Gini index is decreased by splits over a given predictor, averaged over all B trees.

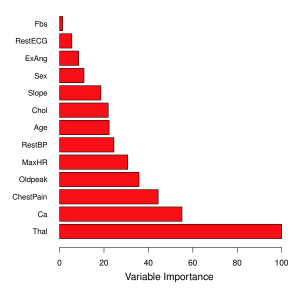


Figure: 8.9. A variable importance plot for the Heart data. Variable importance is computed using the mean decrease in Gini index, and expressed relative to the maximum. Chapter 8

#### Random forest

- Same as bagging decision trees, except ...
- When building these decision trees, each time a split in a tree is considered, a random sample of m predictors is chosen as split candidates from the full set of p predictors
- Typically  $m \approx \sqrt{p}$ .

#### Random forest

- Every step, the split is constrained on a small number m and randomly selected inputs.
- Avoid all trees are too similar to each other.
- Too similar trees are too highly correlated, average highly correlated trees cannot achieve large amount of variance reduction.
- Extreme case: If all trees are the same, average of them is still the same one.
- Averaging uncorrelated or low-correlated trees can achieve large amount of variance reduction.
- Random forest produces less correlated trees.
- Random forest reduces to bagging if m = p.

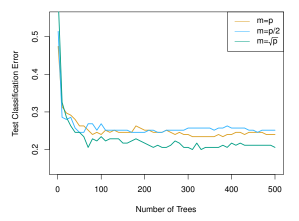


Figure: 8.10. Results from random forests for the 15-class gene expression data set with p=500 predictors. The test error is displayed as a function of the number of trees. Each colored line corresponds to a different value of m, the number of predictors available for splitting at each interior tree node. Random forests (m < p) lead to a slight improvement over bagging (m=p). A single classification tree has an error rate of 45.7%.

## Boosting

- General purpose for improving learning methods by combining many weaker learners in attempt to produce a strong learner.
- Like bagging, boosting involves combining a large number of weaker learners.
- The weaker learners are created sequentially. (no boostrap involved).
- Bagging create large variance and possibly over-fit boostrap learners and try to reduce their variance by averaging.
- Boosting create weak learners sequentially and slowly (to avoid over-fit).

## Boosting

• Suppose we have model

$$y_i = f(x_i) + \epsilon_i$$

and a learning method to produce  $\hat{f}$  based on  $(y_i, x_i), i = 1, ..., n$ .

- Start with an initial predictor  $\hat{f} = 0$ . Let  $r_i = y_i$ .
- Start loop:
  - 1 Fit the data  $(x_i, r_i), i = 1, ..., n$ , to produce  $\hat{g}$ .
  - **2** Update  $\hat{f}$  by  $\hat{f} + \lambda \hat{g}$ .
  - 3 Update  $r_i$  by  $r_i \lambda \hat{g}(x_i)$ .
- Continue the loop ... till a stop.
- Output  $\hat{f}$
- Note that the output  $\hat{f}$  is the sum of  $\lambda \hat{g}$  at each step.

## Algorithm for tree boosting

- 1. Set f(x) = 0 and  $r_i = y_i$  for all i in the training set.
- 2. For b = 1, 2, ..., B, repeat:
  - 1 Fit a tree with d splits (d+1) terminal nodes to the training data  $(x_i, r_i)$ .
  - 2 Update  $\hat{f}$  by adding in a shrunken version of the new tree:

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

3 Update the residuals,

$$r_i \leftarrow r_i - \lambda \hat{f}_b(x_i) = y_i - \hat{f}(x_i).$$

• 3. Output the boosted model  $\hat{f}$ . In fact,

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

### Tuning parameters for boosting trees

- The number of trees B. Large B leads to overfit. (not a tuning parameter for bagging)
- The learning rate  $\lambda$ .
- The number d in splits in each tree (the size of each tree). Often d = 1 works well, in which case each tree is a stump, consisting of a single split

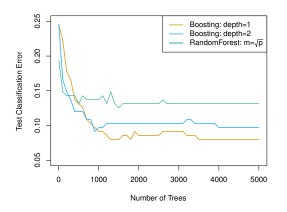


Figure: 8.11. Results from performing boosting and random forests on the 15-class gene expression data set in order to predict cancer versus normal. The test error is displayed as a function of the number of trees. For the two boosted models,  $\lambda = 0.01$ . Depth-1 trees slightly outperform depth-2 trees, and both outperform the random forest, although the standard errors are around 0.02, making none of these differences significant. The test error rate for a single tree is 24%.

#### Exercises

Run the R-Lab codes in Section \*.3 of ISLR Exercises 1-4 and 7-8 of Section 8.4 of ISLR 8.2 Bagging, random forests and boosting

End of Chapter 8.