Tree-based Models and Ensembles

Lecture 11

Supervised Learning Techniques

Linear Regression

K-Nearest Neighbors

Perceptron

Logistic Regression

Fisher's Linear Discriminant

Linear Discriminant Analysis

Quadratic Discriminant Analysis

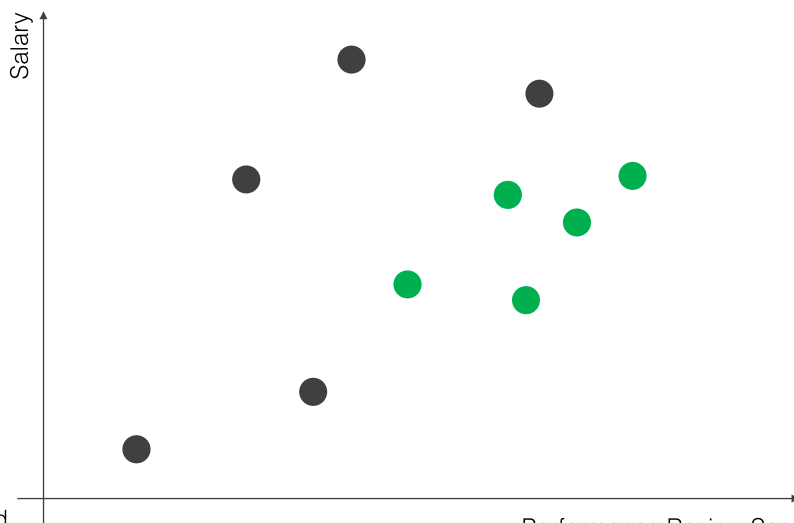
Naïve Bayes

Decision Trees

Ensemble methods (bagging and boosting)

Classification trees = decision trees

Predicting promotions of salaried employees



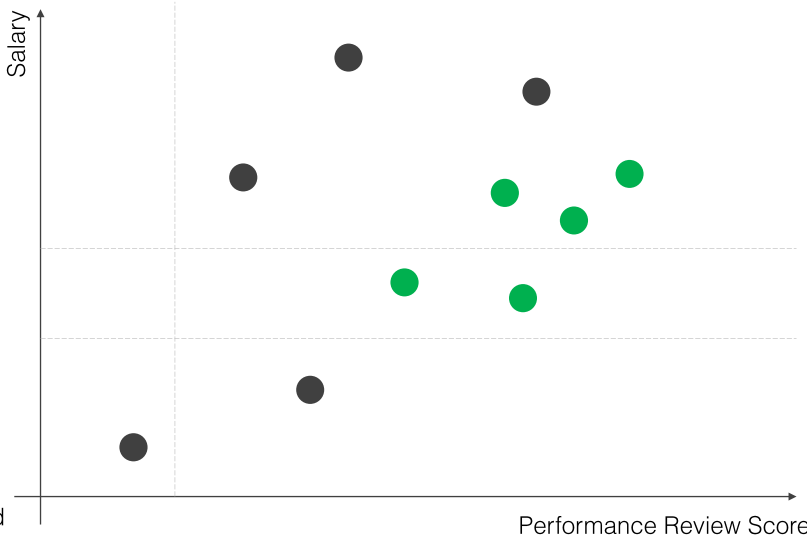
Promoted

Not promoted

Performance Review Score

Predicting promotions of salaried employees

Find the best "split" in any one feature (that best classifies the data) that divides the region in two

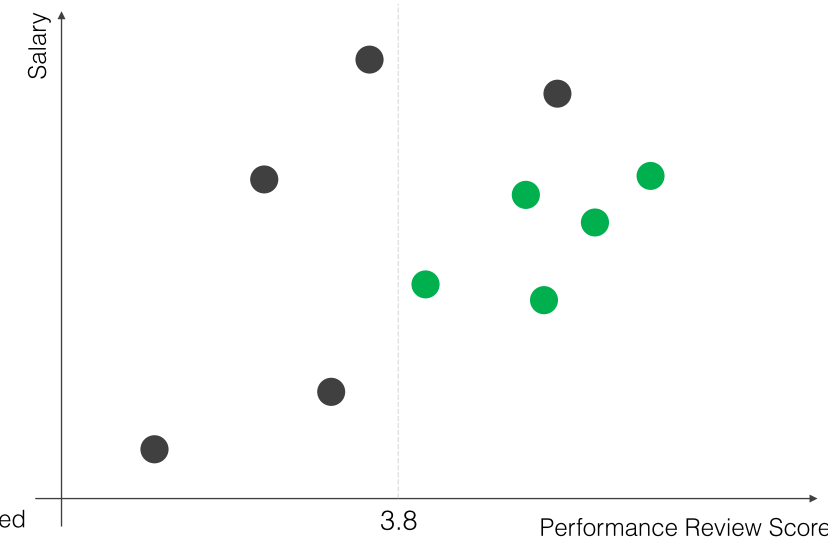


Promoted

Not promoted

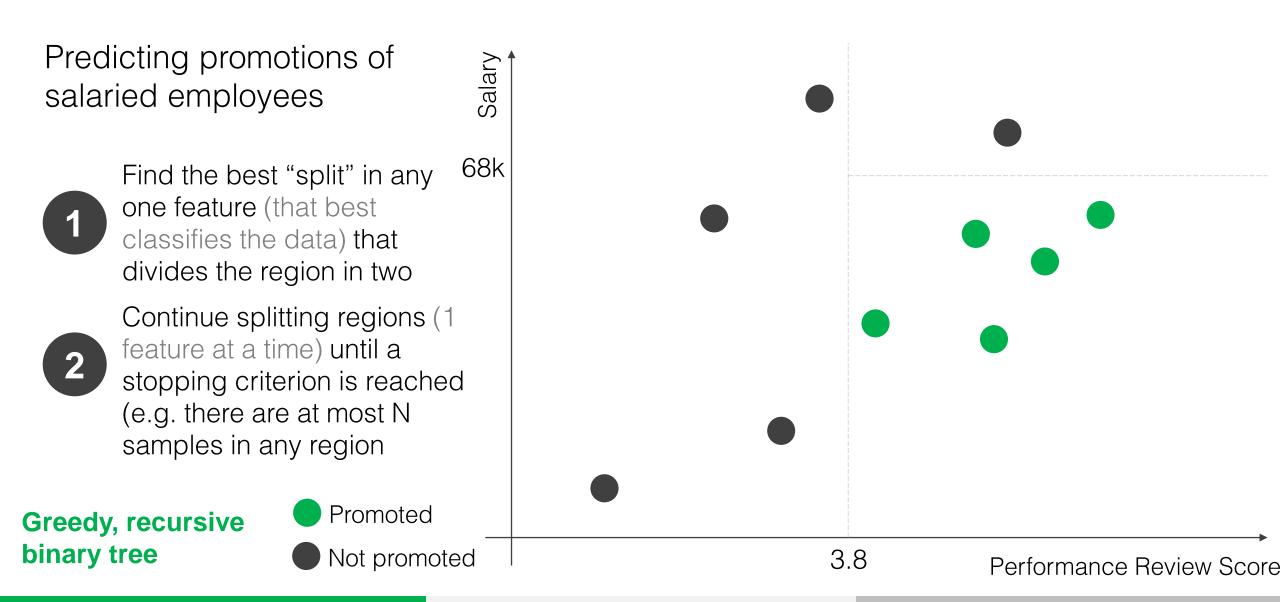
Predicting promotions of salaried employees

Find the best "split" in any one feature (that best classifies the data) that divides the region in two

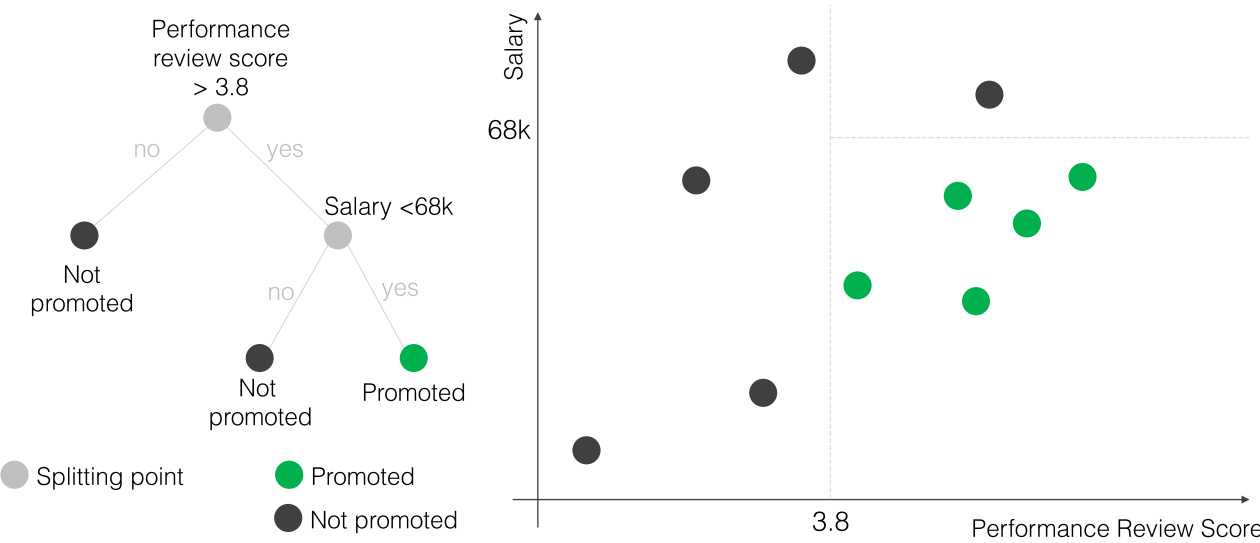


Promoted

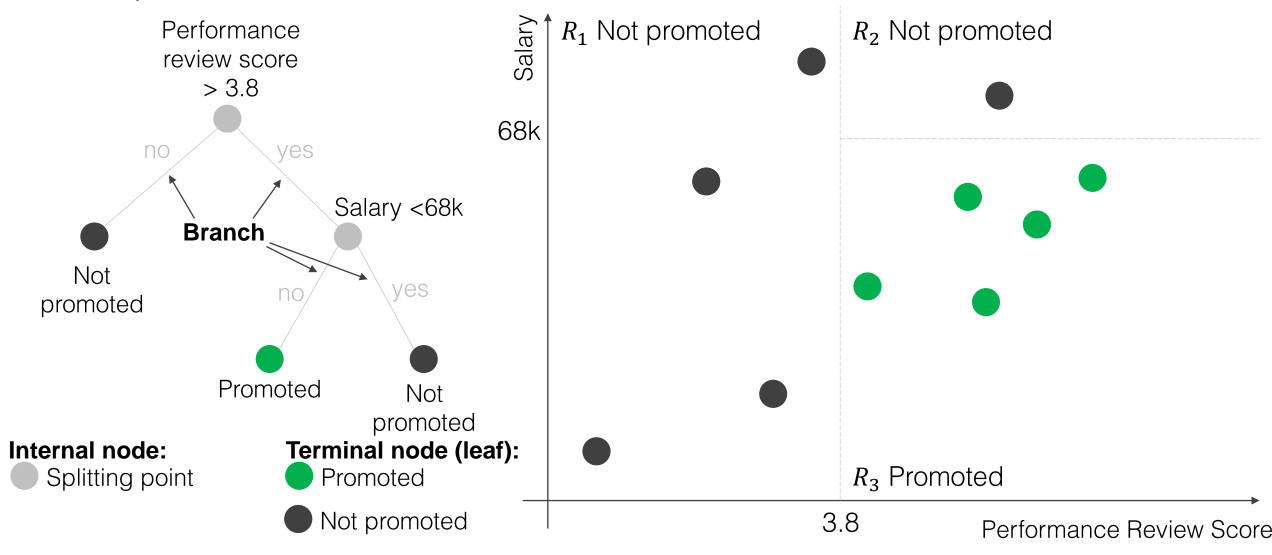
Not promoted



Tree representation:

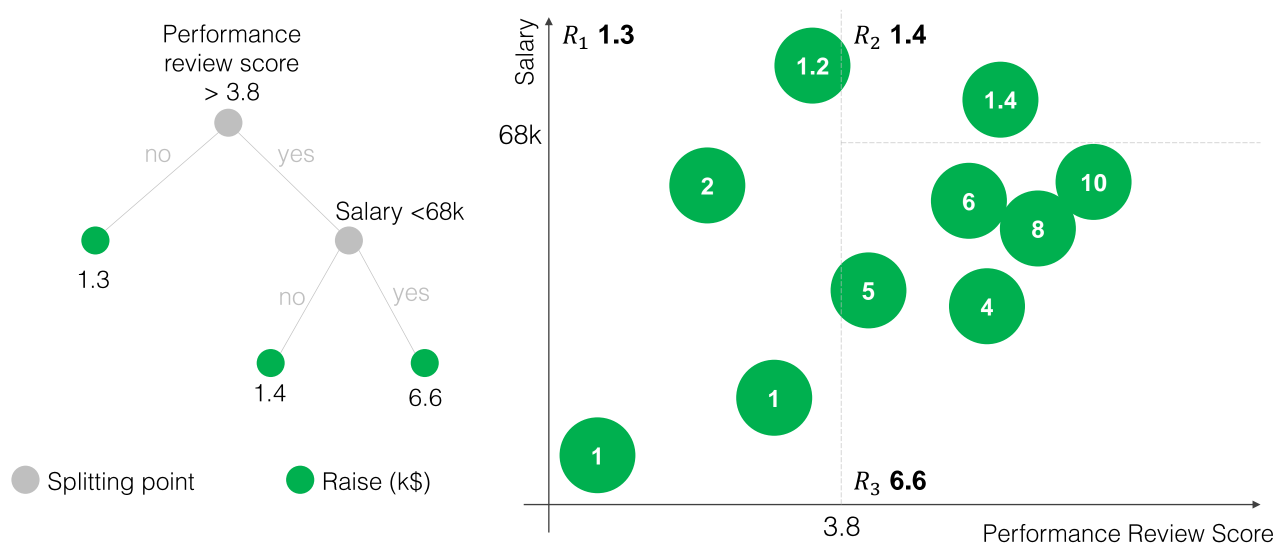


Tree representation:



The Regression Setting

In this case, each region is represented by an average of the values it contains



How do we determine which split to make?

Pick the split that reduces the error/cost criterion most after the split

Splitting criterion

$$C = \sum_{r=1}^{R_{tot}} Q(r)$$

Regression

Kyle Bradbury

Mean square error

$$Q_{MSE}(r) = \sum_{i \in R_r} (y_i - \hat{y}_{R_r})^2$$

 y_i = training data response i

 \hat{y}_{R_r} = mean value in region r, (where R_{tot} is the total # of regions)

Classification

Misclassification rate

$$Q_{Misclass} = 1 - \max_{k} (\hat{p}_{rk})$$

Gini impurity

$$Q_{Gini} = \sum_{k=1}^{K} \hat{p}_{rk} (1 - \hat{p}_{rk})$$

Cross-entropy $Q_{entropy} = -\sum_{k=1}^{K} \hat{p}_{rk} \log \hat{p}_{rk}$

 \hat{p}_{rk} = proportion of training observations in the r^{th} region from the k^{th} class

Example for the two-category case

Chindhod Range of the two-category case

O 0.5 1 p

Duda, Hart, and Stork., Pattern Classification

How to measure quality of split for classification?

Class 1

Class 2

 \hat{p}_{rk} = proportion of training observations in the r^{th} region from the k^{th} class

For each region:

Misclassification rate

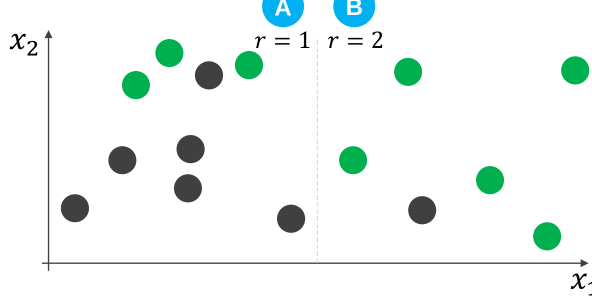
$$Q_{Misclass} = 1 - \max_{k} (\hat{p}_{rk})$$







$$Q_{Gini} = \sum_{k=1}^{K} \hat{p}_{rk} (1 - \hat{p}_{rk})$$



$$\hat{p}_{11} = 3/9$$

$$\hat{p}_{12} = 6/9$$

$$\hat{p}_{21} = 5/6$$

$$\hat{p}_{22} = 1/6$$

Cross-entropy

$$Q_{entropy} = -\sum_{k=1}^{K} \hat{p}_{rk} \log \hat{p}_{rk} \qquad 0.637$$

Tree Pruning

Trees have the tendency to overfit the data

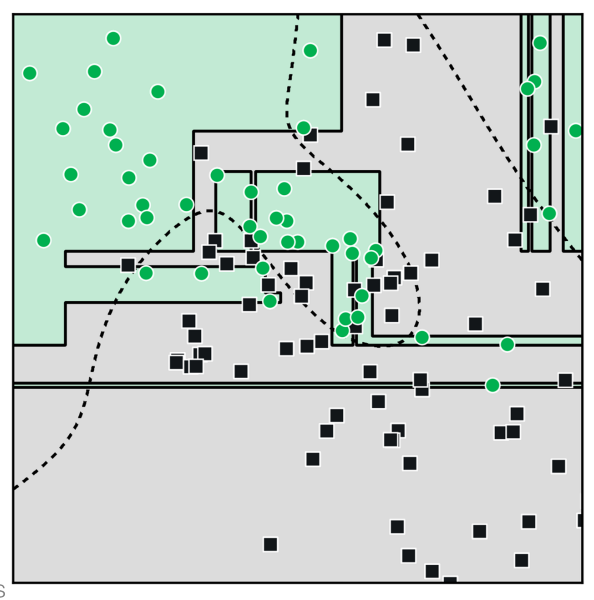
Consider the stopping rule: stop splitting once there is only 1 observation in each region (leads to complete overfit)

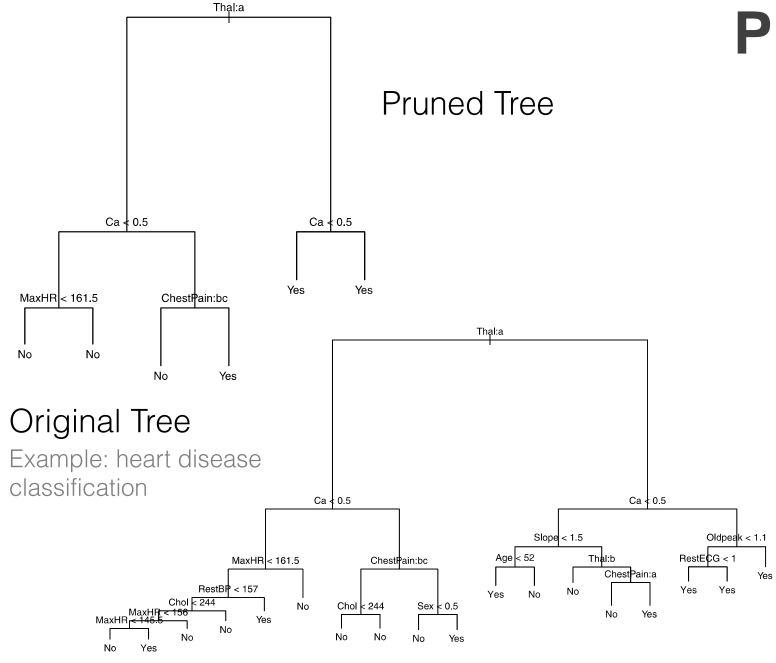
Pruning the tree back reduces this overfit (removing splits after the tree is formed)

Pruning can be optimized through a penalty on the number of terminal nodes:

$$C_{Prune} = \sum_{j=1}^{T} \sum_{i \in R_j} \left(y_i - \hat{y}_{R_j} \right)^2 + \alpha T$$
penalty on number of terminal nodes

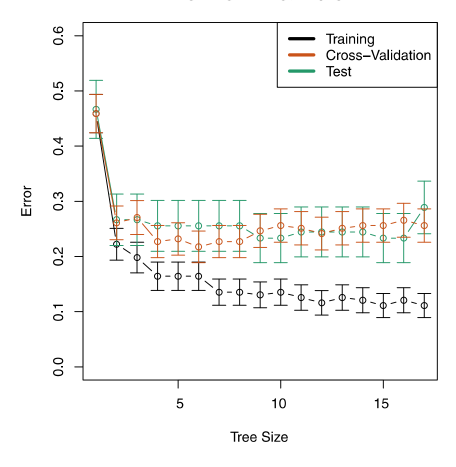
Decision Tree





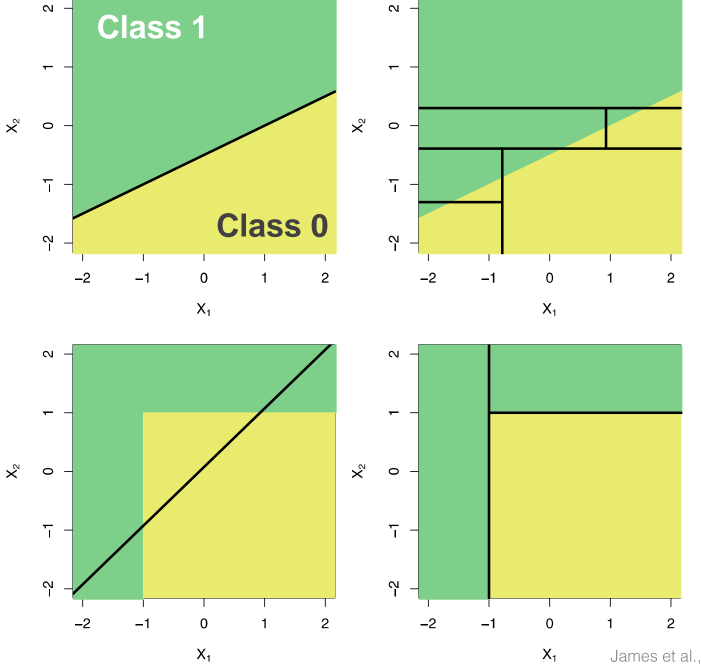
Pruning example

Performance



James et al., An Introduction to Statistical Learning

Linear model



Classification Tree

Struggle when the boundary is not parallel to an axis

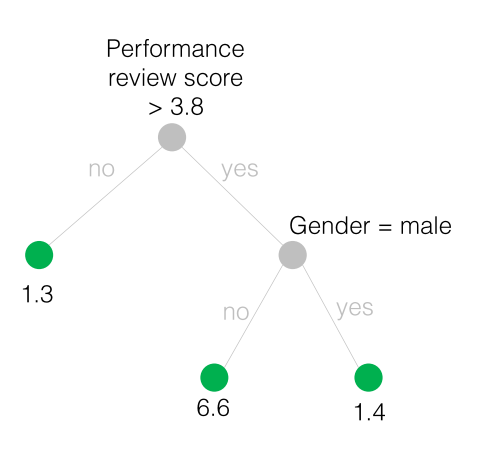
...nonlinear feature transforms could help...

James et al., An Introduction to Statistical Learning

Pros/Cons

Numerical data

Categorical data



Pros:

Trees easily handle multiple types of data

Trees are easy to interpret

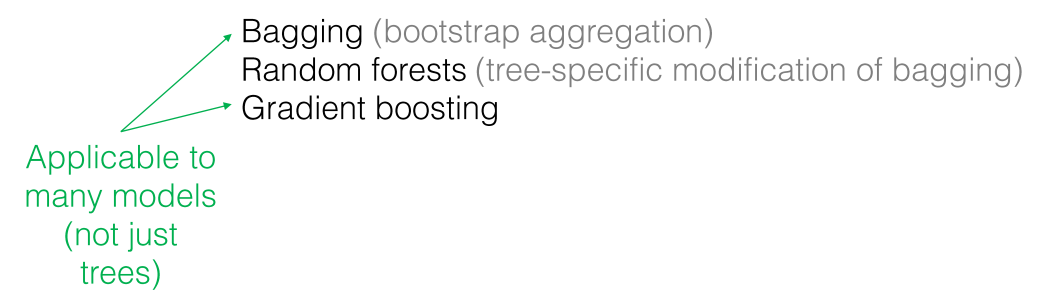
Cons:

Trees do not typically have the same level of predictive accuracy of many methods

Tend to overfit (have high variance)

Ensemble learning

How can we combine models to improve performance?



Reducing Variance or Bias through ensembles

Bagging

Boosting

Models in ensemble:

high variance, low bias

high bias, low variance

Effect of aggregating:

Reduce variance through averaging output

Reduce bias through sequentially fitting models to previous model errors

Bagging

Bootstrap aggregation

Trees overfit (have high variance). Averaging over observations reduces variance

Recall bootstrap sampling (sampling with replacement):

Original Data:

















Bootstrapped sample 1:



Bootstrapped sample 3:





















Can be applied to many machine learning techniques!

Bootstrap aggregation

- 1 Create a random bootstrap sample from the training data
- Train a model on that bootstrap sample and call it $\hat{f}_i(x)$
- Repeat 1 and 2 until we have B models trained on different bootstrap samples
- Take the average of the output for our new model estimate:

$$\hat{f}_{bag}(\mathbf{x}) = \frac{1}{B} \sum_{i=1}^{B} \hat{f}_{i}(\mathbf{x})$$

(for classification models we can take a majority vote instead)

Bagging

Tree Number:









Observations

Included: (out of 1-9)

[1,2,3,3,8]

[1,2,4,7,7]

[1,5,6,8,9]

[2,2,2,4,9]

Features list:

[A, B, C, D]

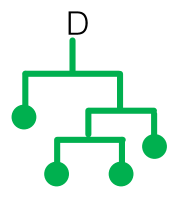
[A, B, C, D]

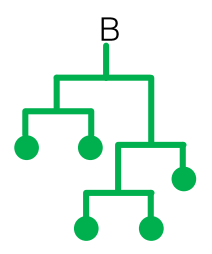
[A, B, C, D]

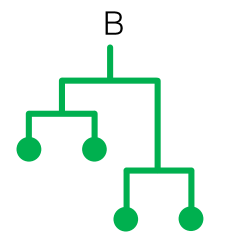
[A, B, C, D]

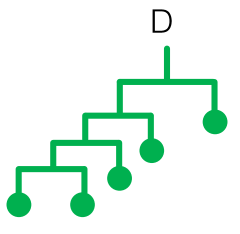
First split:

Trees:









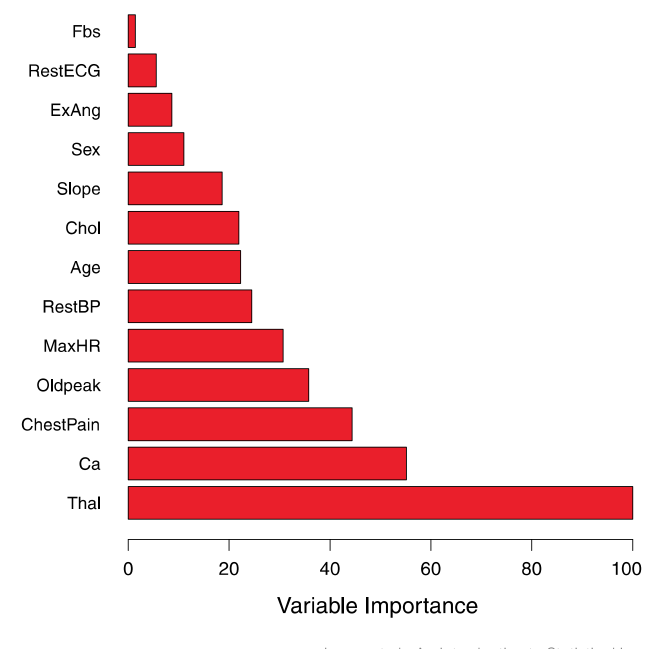
Variable Importance

Decision trees are very interpretable, but this is lost with bagging

We can construct another measure called "variable importance" to compare feature contributions

Calculate the total amount the error (or impurity) decreased by splitting on each feature.

Average over all the trees resulting from bagging



James et al., An Introduction to Statistical Learning

Random Forests

A small tweak on bagging

Random forests decorrelate the bagged trees

Decision trees are constructed greedily

This can lead to highly correlated trees

"Strong" features will typically be split before moderately strong predictors.

Each time a split is considered, a **random subset of** m **features** is selected as candidates from the full set of p features

Typically chose:
$$m = \sqrt{p}$$

(If m = p, then we would be back to the bagging approach)

Bagging

Random forests

Observations Included: (out of 1-9)

[1,2,3,3,8]

[A, B, C, D]

[1,2,3,3,8]

Features list:

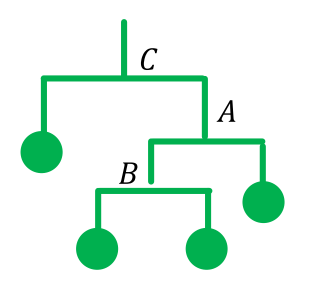
Feature options

for each split:

[A, B, C, D]

[A, B, C, D]

[A, B, C, D]



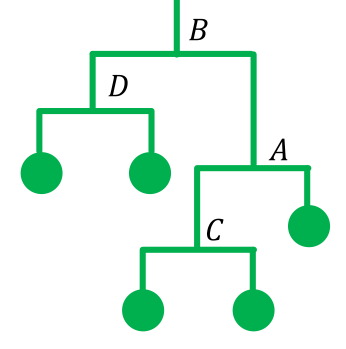
[A,B,C,D]

[C,D]

[A,B]

[A, B]

[B,C]

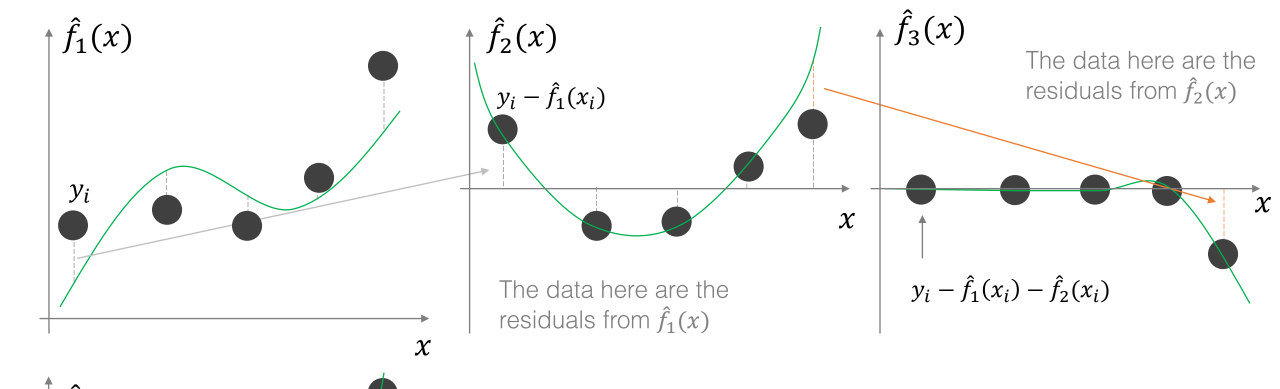


Boosting

Can be applied to many machine learning techniques!

Bagging created trees that were designed to be as independent as possible

Boosting involves building trees sequentially, each building on the errors of the last



 $\hat{f}_{hoost}(x)$ We build consecutive models, each fit to the residuals of the last model y_i

We sum models output to get the boosted prediction $\hat{f}_{boost}(x) = \hat{f}_1(x) + \hat{f}_2(x) + \hat{f}_3(x)$

Lecture 11

Boosting

Boosting for regression trees

- Select the number of models to train, B, and learning rate λ
- λ slows down the learning process to avoid overfitting

- Set $\hat{f}(x) = 0$ and $r_i = y_i$ for all the training data
- Fit a tree, $\hat{f}_i(x)$ to the residuals, r_i (with d splits)

Often this is just a small number of splits

4 Update $\hat{f}(x) = \hat{f}(x) + \lambda \hat{f}_i(x)$

Repeat B times

- 5 Update the residuals $r_i = r_i \lambda \hat{f}_i(\mathbf{x}_i)$
- Output the boosted model: $\hat{f}(x) = \sum_{i=1}^{\nu} \lambda \hat{f}_i(x)$

Model Stacking

Train multiple supervised learning techniques (could be different models)

THEN Train a supervised learning technique that includes the **outputs** of the other models as **features**

Supervised Learning Techniques

- Linear Regression
- K-Nearest Neighbors
 - Perceptron
 - Logistic Regression
 - Fisher's Linear Discriminant
 - Linear Discriminant Analysis
 - Quadratic Discriminant Analysis
 - Naïve Bayes
- Decision Trees and Random Forests
- Ensemble methods (bagging, boosting, stacking)

Appropriate for:

Classification

Regression

Can be used with many machine learning techniques