Lecture 7: Supervised Learning Pt. 1

Non-Linear Models, Bootstrapping, and Bagging

INFO 1998: Introduction to Machine Learning



Agenda

- 1. Decision Trees
- 2. Logistic Regression
- 3. Validation Techniques
 - Bootstrapping & Bagging



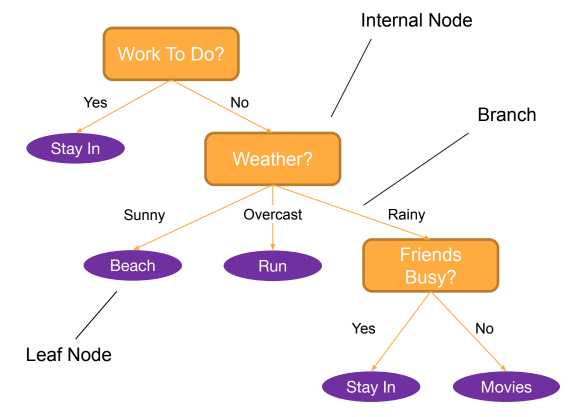
Decision Trees



How Should I Spend My Weekends

Decision Tree

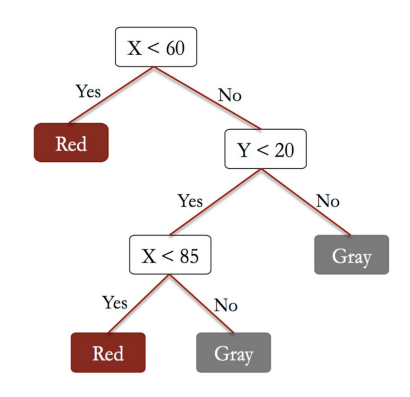
- supervised machine learning model
- breaking down our data by making a decision based on asking a series of questions based on features





CART (Classification and Regression Trees)

- Used for Classification and Regression
- At each node, split on variables
- Each split minimizes error/impurity function
- Very interpretable
- Models a non-linear relationship!



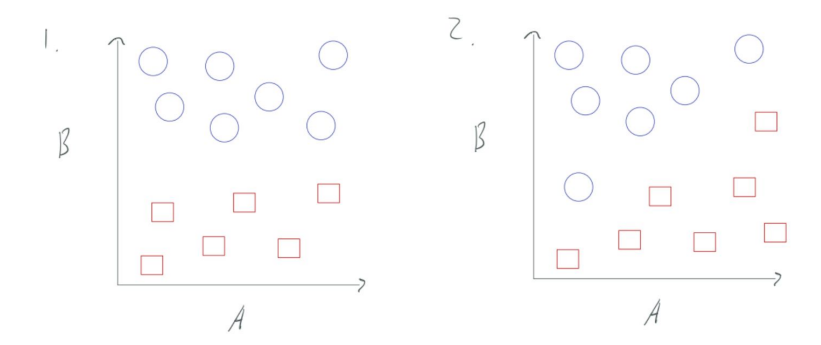


Pros and Cons of Using Decision Trees

| Pros | Cons |
|---|--|
| Easy to interpret | Overfitting 😕 |
| Requires little data preparation (robust to missing data) | Requires parameter tuning (max depth) |
| Can use a lot of features | Can only make horizontal/vertical splits (solvable with feat. eng. / ensembling) |
| Can capture non-linear relationships | |



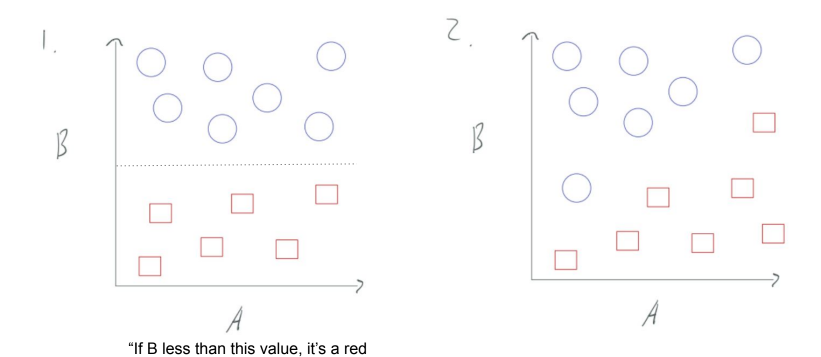
What would these decision boundaries look like?





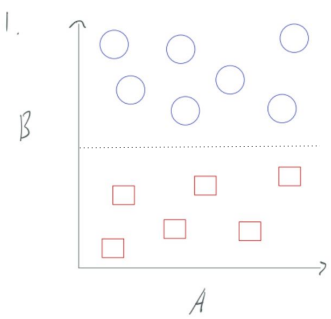
What would these decision boundaries look like?

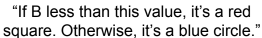
square. Otherwise, it's a blue circle."

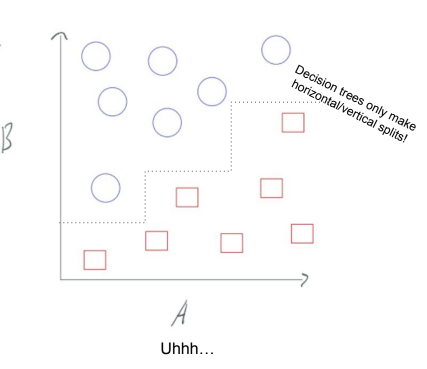




What would these decision boundaries look like?

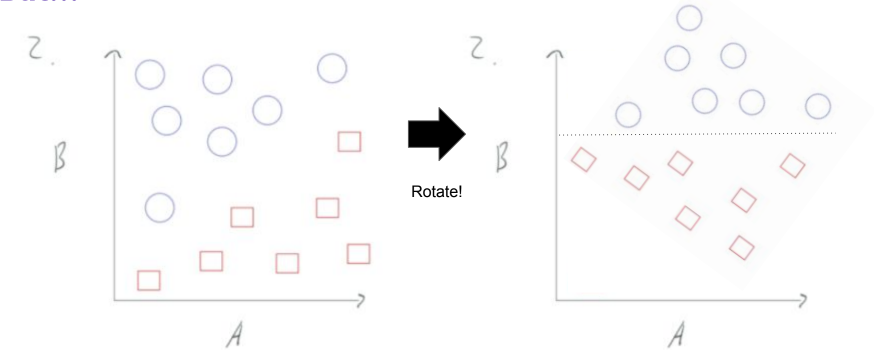








But...





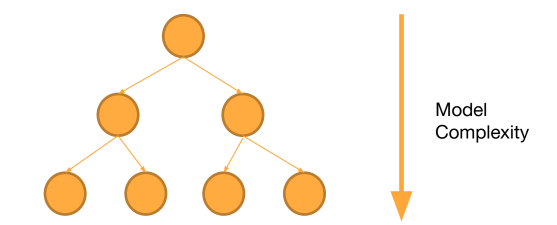
How to Reduce Overfitting

1. Limit the max depth of the tree

Depth = 0

Depth = 1

Depth = 2

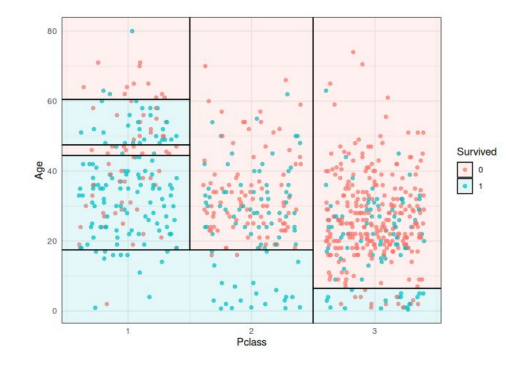


When training a decision tree, we have to specify the maximum depth a constructed tree can have



How to Reduce Overfitting

- There are no "curves" for each decision tree boundary line
- Limiting the depth of the tree limits the number of lines you are splitting on

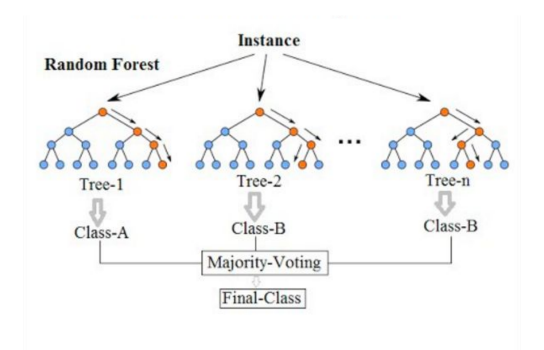




How to Reduce Overfitting

2) Train multiple decision trees and determine final output based on output of each decision tree

This is called a **Random Forest Classifier**





Demo



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Logistic Regression



Logistic Regression

Used for Binary Classification:

$$Y = \begin{cases} 1 \\ 0 \end{cases}$$

- Fits a linear relationship between the variables
- Transforms the linear relationship of probability that the outcome is 1 by using the sigmoid function

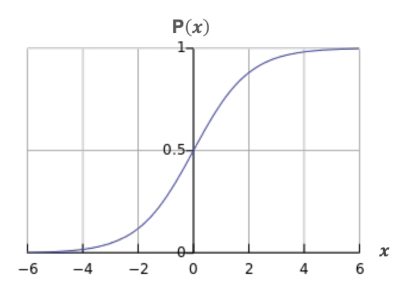
Formula:

$$P(x) = \frac{1}{1 + e^{-(\beta_0 + \beta_1 x_1 + \dots + \beta_k x_k)}} \longrightarrow \ln\left(\frac{P}{1 - P}\right) = \beta_0 + \beta_1 x_1 + \dots + \beta_k x_k$$



Logistic Function

$$P(x) = \frac{1}{1 + e^{-x}}$$



The Logistic Function "squeezes" numbers to be between 0 and 1



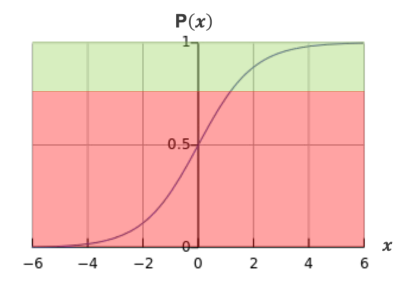
Allows us to interpret our prediction as a "probability" that something is true



Threshold

At what point do we differentiate between our classifications?

- f(x) below threshold: predict 0
- f(x) above threshold: predict 1





Pros and Cons of Using Logistic Regression

| Pros | Cons |
|--------------------------------------|---|
| Easy to interpret (probability) | Only Capable of Binary Classification |
| Computationally efficient to compute | No closed form solution (requires use of optimization algorithms) |
| Does not require parameter tuning | |

Logistic Regression is a simple model, therefore, oftentimes it is used as a good "baseline" to compare more complex models to



Validation Techniques



Review: Regression vs. Classification

Regression

- Predict Continuous Data
- "On average, how wrong are we?"

Classification

- Predict Discrete or Categorical data
- "How many points do we get wrong?"

Numbers



Continuous



Leave-P-out

Let **D** be our whole dataset

Choose a P

For every combination of **P** points in **D**:

Use a train/test split with those **P** points as test, the rest as train



Leave-P-out

Let's say **D** has a size of 4. There are four data points: *a, b, c,* and *d*. Leave-P-out:

- P = 2.
- We have 6 possible test sets:

```
{a,b}, {a,c}, {a,d}, {b,c}, {b,d}, and {c,d}
```



Leave-P-out

Pros:

- Dependable (not random)
- Representative checks all combinations

Cons:

- Slow!
 - Runtime <u>increases</u> with larger datasets
 - Runtime <u>explodes</u> with larger P



Monte Carlo Cross Validation

- Getting accuracy 1 time doesn't tell us much
- Getting accuracy 2 times tells us a bit
- Getting accuracy 3 times tells us a bit more
- ...
- Getting accuracy N times might be good enough!

Take the average of those **N** times



Monte Carlo CV

- Need to use new, random train/test split each time
 - If you use the same train/test split each time, you're not getting any new information!
- Pros:
 - easy to implement
 - easy to make faster/slower by changing number of iterations
- Cons:
 - random -> train/test splits not guaranteed to be representative of dataset (might overlap, or miss some data)
 - harder to calculate how many iterations you need



The Bootstrap

What if we don't have enough data?

- Use **bootstrap datasets** to approximate the test error
- Sample with replacement from the original training dataset (with n samples) to generate bootstrap datasets of size n
 - Some data points may appear more than once in the generated data
 - Some data points may not appear
- Estimate of test error = average error among bootstrap datasets



Demo



Pipeline of the Bootstrap

Algorithm 1 Estimating prediction error via the bootstrap

- 1: **Input**: dataset $\{(x_i, y_i)\}_{i=1}^n$, loss function ℓ .
- 2: **for** $b = 1, 2, \dots, B$ **do**
- 3: generate set S_b by sampling n items with replacement from $\{(x_i, y_i)\}_{i=1}^n$.
- 4: form prediction \hat{f}^b by fitting the logistic regression model with training data \mathcal{S}_b .
- 5: compute $\widehat{\text{err}}_b := \frac{1}{n} \sum_{i=1}^n \ell(y_i, \hat{f}^b(x_i))$
- 6: end for
- 7: **return** $\widehat{\text{err}}_{\text{boot}} := \frac{1}{B} \sum_{b=1}^{B} \widehat{\text{err}}_{b}$
- Does this give very good estimate of the prediction error?
- ❖ If **no**, why not? Does it **underestimate** or **overestimate** the prediction error?



Pipeline of the Bootstrap

No.

Since the bootstrap is sampling with replacement for B validation folds, each fold would have significant overlap with the original data used for training. Approximately, **2/3** of the training data would appear in each validation fold.

This leads to significant **underestimation** of the prediction error.

❖ Why 2/3?



Proof

Suppose there are n samples (yi, xi) in the training set data, then for each of the bootstrap validation fold b, every sample has probability 1/n of being selected into b. The probability of each data sample (yi, xi) not being in fold b is (1 - 1/n), respectively.

Probability of avoid selecting all n data points in fold b: (1 - 1/n)^n.

When n gets large, we have this probability converges to 1/e.

(Why this converges? Probably should review Calc I, or see:

https://math.stackexchange.com/questions/882741/limit-of-1-x-nn-when-n-tends-to-infinity.)

Therefore, the fraction of overlapping data points in each fold is (1-1/e), which is about 2/3.



How we fix the problem

- Requires some thoughts when generating validation set, especially for real-world complex data.
- Can partly fix this problem by only using predictions for those observations that did not (by chance) occur in the current bootstrap sample.
- But the method gets complicated.



Bootstrap vs. k-fold

In K-fold validation, each of the K folds is distinct from the other (K - 1) folds used for training: there is **no overlap**.

This is crucial for its success in estimating prediction error.



Why do we still use Bootstrap?

- Bootstrap allows us to use a computer to mimic the process of obtaining new data sets.
- Can be used to quantify the uncertainty associated with a given estimator or statistical learning method.
- Provides an estimate of the standard error of a coefficient, or a confidence interval for that coefficient.
 - i.e. the variability of the model!



Bagging (Bootstrap Aggregating)

What if we don't have enough data?

- Bagging is a common technique that builds on Bootstrapping
- Main Idea: Do Bootstrapping a bunch and make a classifier for each bootstrap, then choose majority prediction.
- Many weak learners aggregated typically outperform a single learner over the entire set, and overfits less.
 - Principle behind Random Forests ("forest" of decision trees)



Coming Up

- Last day for Mid-Semester Check-Ins!
- **Assignment 6:** Due tonight at 11:59pm
- **Assignment 7**: Due November 5th, 2025
- Final Project: Due December 3rd, 2025
- **Next Lecture**: Supervised Learning Part 2

