# Random forests STAT 471

### Where we are

Unit 1: Intro to modern data mining



Unit 2: Tuning predictive models



Unit 3: Regression-based methods

Unit 4: Tree-based methods

Unit 5: Deep learning

Lecture 1: Growing decision trees

Lecture 2: Tree pruning and bagging

Lecture 3: Random forests

Lecture 4: Boosting

Lecture 5: Unit review and quiz in class

Homework 4 due the following Wednesday.

# Recall: Bagging

#### Bootstrap sample 1

()rio	ıinal	training	data
			JALL

Obs ID	X	Y
1	X <sub>1</sub>	Y <sub>1</sub>
2	<i>X</i> <sub>2</sub>	<b>Y</b> <sub>2</sub>
3	<b>X</b> 3	<b>Y</b> <sub>3</sub>
4	<i>X</i> <sub>4</sub>	Y <sub>4</sub>
5	<i>X</i> <sub>5</sub>	<b>Y</b> <sub>5</sub>

Obs ID	X	Y
5	$X_5$	<b>Y</b> <sub>5</sub>
3	<b>X</b> 3	<b>Y</b> 3
2	$X_2$	Y <sub>2</sub>
3	<i>X</i> <sub>3</sub>	<b>Y</b> <sub>3</sub>
1	X <sub>1</sub>	Y <sub>1</sub>

 $\longrightarrow T^{*b}$ 

#### Bootstrap sample B

Obs ID	X	Y
4	<i>X</i> <sub>4</sub>	Y <sub>4</sub>
1	X <sub>1</sub>	Y <sub>1</sub>
1	X <sub>1</sub>	Y <sub>1</sub>
5	<i>X</i> <sub>5</sub>	<b>Y</b> <sub>5</sub>
4	X <sub>4</sub>	<b>Y</b> <sub>4</sub>

Regression:

$$\hat{f}(X) = \frac{1}{B} \sum_{b=1}^{B} T^{*b}(X)$$

Classification:

$$\hat{f}(X) = \text{mode}(\{T^{*b}(X)\}_{b=1}^{B})$$

# Variance reduction of bagging

The bagging prediction is defined by  $\hat{f}(X) = \frac{1}{B} \sum_{b=1}^{B} T^{*b}(X)$ .

Suppose  $\operatorname{Corr}[T^{*b_1}(X), T^{*b_2}(X)] = \rho \in [0,1]$ . Then, we can derive that

$$\operatorname{Var}[\hat{f}(X)] = \frac{1}{B^2} \sum_{b_1 = 1}^{B} \sum_{b_2 = 1}^{B} \operatorname{Cov}[T^{*b_1}(X), T^{*b_2}(X)] \approx \left(\frac{1}{B} + \frac{B - 1}{B}\rho\right) \operatorname{Var}[T(X)] \approx \rho \cdot \operatorname{Var}[T(X)],$$

where T(X) is a single decision tree. Take-aways:

- The variance is reduced by a factor of  $\rho = \text{Corr}[T^{*b_1}(X), T^{*b_2}(X)]$ , so the less correlated the bootstrapped trees prediction are, the better.
- ullet As long as B is large enough, the variance reduction is about the same.

### Random forests: More variance reduction

Random forests are the same as bagging, but with one key modification:

At each split point of each tree:

- Randomly sample a subset of  $m \le p$  features
- Split on the best feature among this subset

Intuition: Sampling features at each split decorrelates the trees, reducing variance and therefore boosting prediction performance.

Hopefully each individual tree is still roughly unbiased even though it has access to a smaller number of features at each split.

Note that setting m = p recovers bagging.

### Random forests

#### Parameters:

- B: number of bootstrap samples
- m: number of variables to sample at each split
- criterion to stop splitting, like max number of nodes and/or min samples per node

#### Training:

- Extract B bootstrap samples from your training data
- For each bootstrap sample b = 1, ..., B,
  - Grow a decision tree based on the bootstrap sample, randomly sampling *m* candidate variable to split on at each step, until stopping criterion is met

#### Prediction:

• aggregate the decision trees using the mean (for regression) or mode (for classification)

# A bias-variance trade-off in choosing m

If m is larger, the random forest will have lower bias (it can better fit the underlying trend) but higher variance (more correlated trees).

If m is smaller, the random forest will have higher bias (it might not be able to fit the underlying trend as well) but lower variance (less correlated trees).

Default choices: m = p/3 for regression and  $m = \sqrt{p}$  for classification.

For best predictive performance, m should be tuned.

# Tuning random forests via out-of-bag error

We usually tune prediction methods via cross-validation. For random forests, there is a clever and computationally faster alternative: out-of-bag error.

The idea behind cross-validation is that we want to using parts of our training data as validation sets. By bootstrapping, random forests already do this!

For each bootstrap sample, define the "bag" to be the set of unique training observations in the sample. Then, predictions based on that tree can be made on the out-of-bag (OOB) samples.

OOB error: For each training observation, aggregate the predictions of all trees not trained on this observation. Then compute the error (mean-squared or misclassification) of these predictions.

On average, each tree will be trained on about 2/3 of the training observations. Therefore, about 1/3 of all trees will be trained without a given training observation.

### Parameters to tune (or not)

Random forests generally work pretty well even if not tuned (i.e. if default parameter choices are used).

However, parameters can be tuned using OOB error to improve performance:

- *m*: most important tuning parameter
- criteria to stop splitting: can be tuned but growing trees about as deep as possible generally works pretty well
- B: least necessary to tune; just choose a large value like 100-1000.

### Interpretability and variable importance measures

Compared to trees, main drawback of random forests is reduced interpretability.

However, variable importance measures can help improve the interpretability.

Two types of variable importance measures are used for random forests:

- purity based importance: how much improvement in node purity results from splitting on a feature
- OOB prediction based importance: how much deterioration in prediction accuracy results from scrambling a feature out of bag

### Purity-based variable importance

Consider the construction of one tree. For each split, note the feature that was split on and resulting reduction in RSS or Gini index (i.e. improvement in purity).

We can then define the importance of each feature in this single tree by summing up the improvement in purity for all splits including this feature.

For random forests, we can average this quantity over all of the trees to get a purity-based variable importance metric.

### **OOB** prediction based variable importance

Recall the OOB error introduced a few slides ago.

For each feature j and each tree, consider making predictions on the OOB data after first scrambling feature j. We can therefore get a scrambled OOB error.

For each feature j, we can define an OOB prediction based variable importance by the difference in OOB error when this feature is scrambled and when it is not.

### Summary

- Random forests are a fancier version of bagging based on random subsampling of m features at each split point.
- They improve on bagging by de-correlating the bootstrapped decision trees and therefore reducing the variance of the method.
- OOB error is a nice alternative to cross-validation error for random forests, and can be used to tune parameters such as m.
- Random forests usually give much better prediction performance than individual decision trees, but at the cost of interpretability.
- Nevertheless, there are a couple ways to measure variable importance in random forests, giving us some interpretability.

Random forests are a state-of-the-art tool for predictive modeling.