ENSEMBLE METHODS: BAGGING -INTRODUCTION TO DATA SCIENCE-

ISL 5.3.4, 8.2

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NOTATION

REMINDER: For either classification or regression, we produce predictions for a given feature vector X

That is, we form

$$\hat{Y} = \hat{f}(X) \in \mathbb{R}$$
 Regression

or

$$\hat{Y} = \hat{g}(X) \in \mathcal{G}$$
 Classification

where

- \hat{f} or \hat{g} is some procedure formed with the training data (EXAMPLE: $\hat{\beta}$ formed by least squares)
- The prediction \hat{Y} formed at a desired feature vector X (Example: $\hat{Y} = X^{\top}\hat{\beta}$ formed by least squares)



BAGGING

Many methods (trees included) tend to be designed to have lower bias but high variance

HEURISTICALLY: If we split the training data into two parts at random and fit a decision tree to each part, the results could be quite different

A low variance estimator would yield similar results if applied repeatedly to distinct data sets (consider $\hat{f}(X) = 0$ for all X)

Bagging, also known as Bootstrap AGgregation, is a general purpose procedure for reducing variance.

We'll use it specifically in the context of trees, but it can be applied more broadly.

Bagging for regression

BAGGING: THE MAIN IDEA

Suppose we have n uncorrelated observations Z_1, \ldots, Z_n , each with variance σ^2 .

What is the variance of

$$\overline{Z} = \frac{1}{n} \sum_{i=1}^{n} Z_i?$$

BAGGING: THE MAIN IDEA

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What is the variance of

$$\overline{Z} = \frac{1}{n} \sum_{i=1}^{n} Z_i?$$

Answer: σ^2/n .

More generally, if we have B separate (uncorrelated) training sets, we could form B separate model fits,

$$\hat{f}^1(X),\ldots,\hat{f}^B(X)$$

Then average them:

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

BAGGING: THE BOOTSTRAP PART

Of course, this isn't practical as having access to many training sets is unlikely.

We therefore turn to the bootstrap to simulate having many training sets.

The bootstrap is a widely applicable statistical tool that can be used to quantify uncertainty without Gaussian approximations.

Let's look at an example.

Bootstrap detour

Suppose we are looking to invest in two financial instruments, X and Y. The return on these investments is random, but we still want to allocate our money in a risk minimizing way.

That is, for some $\alpha \in (0,1)$, we want to minimize

$$Var(\alpha X + (1 - \alpha)Y)$$

The minimizing α is:

$$\alpha_* = \frac{\sigma_Y^2 - \sigma_{XY}}{\sigma_X^2 + \sigma_Y^2 - 2\sigma_{XY}}$$

(Here, σ_{XY} is the covariance between X and Y)

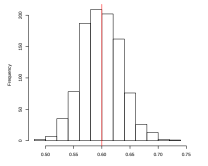
We can estimate α_* via a plug-in estimator

$$\hat{\alpha} = \frac{\hat{\sigma}_Y^2 - \hat{\sigma}_{XY}^2}{\hat{\sigma}_X^2 + \hat{\sigma}_Y^2 - 2\hat{\sigma}_{XY}^2}$$

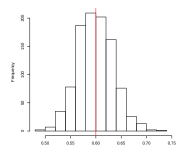
Now that we have an estimator of α_* , it would be nice to have an estimator of its variability.

In this case, computing a standard error is difficult.

Suppose for a moment that we can simulate a large number of draws (say 1000) of the data, which has actual value $\alpha = 0.6$. Then we could get estimates $\hat{\alpha}_1, \ldots, \hat{\alpha}_{1000}$:



This is the sampling distribution of $\hat{\alpha}$



The mean of all of these is:

$$\overline{\alpha} = \frac{1}{1000} \sum_{r=1}^{1000} \hat{\alpha}_r = 0.599,$$

which is very close to 0.6 (red line), and the standard error is

$$\sqrt{\frac{1}{1000 - 1} \sum_{r=1}^{1000} (\hat{\alpha}_r - \overline{\alpha})^2} = 0.079$$

The standard error of 0.035 gives a very good idea of the accuracy of $\hat{\alpha}$ for a single sample.

Roughly speaking, for a new random sample, we expect

$$\hat{\alpha} \in (\alpha - 2 * 0.079, \alpha + 2 * 0.079) = (0.442, 0.758)$$

In practice, of course, we cannot use this procedure as it relies on being able to draw a large number of (independent) samples from the same distribution as our data.

This is where the bootstrap comes in.

We instead draw a large number of samples directly from our observed data. This sampling is done with replacement, which means that the same data point can be drawn multiple times.

BOOTSTRAP DETOUR: SMALL EXAMPLE

Suppose we have data $\mathcal{D} = (4.3, 3, 7.2, 6.9, 5.5)$.

Then we can draw bootstrap samples, which might look like:

$$\mathcal{D}_1^* = (7.2, 4.3, 7.2, 5.5, 6.9)$$
 $\mathcal{D}_2^* = (6.9, 4.3, 3.0, 4.3, 6.9)$
 \vdots
 $\mathcal{D}_B^* = (4.3, 3.0, 3.0, 5.5, 6.9)$

It turns out each of these \mathcal{D}_b^* have very similar properties as \mathcal{D}

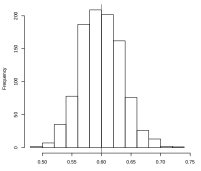
BOOTSTRAP DETOUR: SMALL EXAMPLE

Now, we form the bootstrap mean:

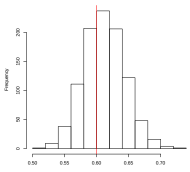
$$\operatorname{mean}_{B} = \frac{1}{B} \sum_{b=1}^{B} \hat{\alpha}_{b}^{*}$$

The bootstrap estimator of the standard error is:

$$SE_B = \sqrt{\frac{1}{B} \sum_{b=1}^{B} (\hat{\alpha}_b^* - \text{mean}_B)^2}$$



Sampling distribution of $\hat{\alpha}$ (impossible to form)



Bootstrap distribution of $\hat{\alpha}$ (possible to form)

BOOTSTRAP: END DETOUR

SUMMARY:

Suppose we want to get an idea of the sampling distribution of some statistic \hat{f} trained on \mathcal{D} .

Then we do the following: Fix a large number B (B could be, say, 1000)

Then for each $b = 1, \ldots, B$

- 1. Form a new bootstrap draw from \mathcal{D} , call it \mathcal{D}^*
- 2. Compute \hat{f}_b^* from \mathcal{D}^*

Now, we can estimate the distribution of \hat{f} trained on \mathcal{D} by looking at the distribution of the B draws, \hat{f}_b^*

End detour

BAGGING: THE BOOTSTRAP PART

Now, instead of having B separate training sets, we train on B bootstrap draws:

$$\hat{f}_1^*(X),\ldots,\hat{f}_B^*(X)$$

and then average (i.e. aggregate) them:

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

This process is known as Bagging

Bagging trees



BAGGING TREES

The procedure for trees is the following

- 1. Choose a large number B.
- 2. For each b = 1, ..., B, grow an unpruned tree on the b^{th} bootstrap draw from the data.
- 3. Average all these trees together.

Each tree, since it is unpruned, will have (low/high) variance and (low/high) bias

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Therefore averaging many trees results in an estimator that has lower variance and still low bias.

Bagging for classification

BAGGING TREES IN CLASSIFICATION

For classification, there are a few sensible methods for aggregation

For each test observation X,

- record the length B vector $[\hat{g}_1^*(X), \dots, \hat{g}_B^*(X)]^{\top}$ and classify X via majority vote
- Average the length G probability vectors from each tree and choose the argmax

WARNING: One thing you definitely do not want to do is estimate probabilities via taking proportions of times X was classified to each class across the B trees

Additional tree bagging topics

BAGGING TREES

Now that we are growing a large number (B) of random trees, we can't directly look at the dendrogram

(we have sacrificed some interpretability for better performance)

However, we do get some helpful information instead

- Mean decrease variable importance
- Out-of-Bag error estimation (OOB)
- Permutation variable importance
- Proximity

(these ideas apply to bagging any low bias procedure, not just unpruned trees)

MEAN DECREASE VARIABLE IMPORTANCE

Observation: Every time a split of a node is made on a feature, the loss function is not increased

Hence, adding up the loss decreases for each feature over all trees gives an indication of feature importance

Intuitively an important feature is one that if split upon, it leads to a large reduction in the loss

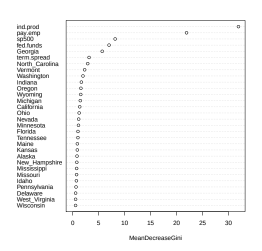
Mean decrease variable importance

To recover some information, we can do the following:

- 1. For each of the *B* trees and each of the *p* features, we record the amount that the Gini index (or cross-entropy) is reduced by splitting on that feature
- 2. Report the average reduction over all B trees

This gives us an indication of the importance of a feature

MEAN DECREASE VARIABLE IMPORTANCE



OUT-OF-BAG SAMPLES (OOB)

One can show that, on average, drawing n samples from n observations with replacement results in about 2/3 of the observations being selected.

The remaining one-third of the observations not used are referred to as out-of-bag (OOB)

OUT-OF-BAG SAMPLES (OOB)

We can think of it as a for-free cross-validation

The observations that aren't included serve as test data

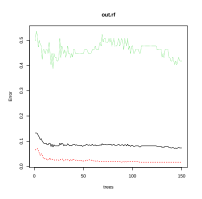
This provides a free estimate of prediction risk for each tree

We can therefore get an overall estimate of prediction risk by averaging these estimates over all bootstrapped trees (Same idea applies here as for getting a prediction at a test X. For each X_i , we can

take a majority vote of all the classifications when (X_i, Y_i) is OOB)

OUT-OF-BAG SAMPLES (OOB)

We can use the OOB samples to choose the number of trees ${\it B}$ to consider



As we are taking an average, we can iteratively compute small batches, stopping when OOB error rate stabilizes (EXERCISE)

PERMUTATION VARIABLE IMPORTANCE

Consider the b^{th} bootstrap sample

- 1. The OOB prediction accuracy is recorded
- 2. Then, the j^{th} feature is randomly permuted in the OOB samples
- The prediction error is recomputed and the change in prediction error is recorded

INTUITION: If a feature is highly important, then the OOB prediction error should increase substantially after permuting the OOB values for that feature

PROXIMITY

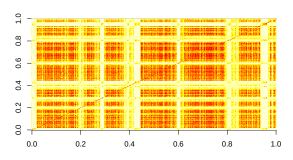
Choose any two observations on the training data: i, i'

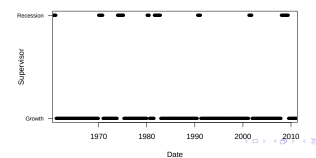
We can record

$$\operatorname{proximity}(i,i') = \frac{\# \text{ times } i,i' \text{ are in the same leaf}}{\# \text{ times } i,i' \text{ occur in same tree}}$$

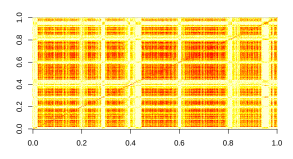
Values near 1 indicate "close" observations and values near 0 indicate "far" observations

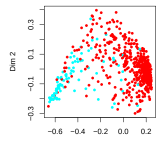
PROXIMITY





PROXIMITY





PROXIMITY

```
#First image
         = randomForest(X,Y,proximity=TRUE)
par(mfrow=c(2,1), mar=c(4,4,1.2,4))
image(1 - out.rf$proximity)
plot(dates, Y,
     xlab='Date', ylab='Supervisor')
dev.off()
#Second image
out.rf = randomForest(X,Y,proximity=TRUE)
par(mfrow=c(2,1), mar=c(4,4,1.2,4))
image(1 - out.rf$proximity)
MDSplot(out.rf,Y,palette=rainbow(2))
```

Random Forest

Random Forest is a small extension of Bagging, in which we attempt to decorrelate the bootstrap trees

IDEA: Draw a bootstrap sample and start to build a tree

- At each split, we randomly select m of the possible p features as candidates for the split.
- A new sample of size m of the features is taken at each split.

Usually $m=\sqrt{p}$ for classification and p/3 for regression (this would be 7 out of 56 features for GDP data)

In other words, at each split, we aren't even allowed to consider the majority of possible features!

What is going on here?

Suppose there is 1 really strong feature and many mediocre ones.

- Then each tree will have this one feature in it.
- Therefore, each tree will look very similar (i.e. highly correlated).
- Averaging highly correlated things leads to much less variance reduction than if they were uncorrelated.

If we don't allow some trees/splits to use this important feature, each of the trees will be much less similar and hence much less correlated.

Bagging is Random Forest when m = p

(That is, when we can consider all the features at each split)



An average of B uncorrelated random variables has variance

$$\frac{\sigma^2}{B}$$

An average of B random variables has variance

$$\rho\sigma^2 + \frac{(1-\rho)\sigma^2}{B}$$

for correlation ρ

As $B \to \infty$, the second term goes to zero, but the first term remains

Hence, correlation of the trees limits the benefit of averaging

Another way to decorrelate the trees is by introducing noise features

Generate a few new features (say 0.01p) that are not related to the supervisor

In some bootstrap samples, this feature will be included in the tree, adding a decorrelating effect

RANDOM FOREST: BIAS AND VARIANCE

With either approach, we are trading bias and variance again

Bagging has the same bias as the underlying procedure, but may not get much variance reduction

Random Forest is biased due to subsampling/noise features, but gets more variance reduction be decreasing ρ (recall that the variance is $\rho\sigma^2+\frac{(1-\rho)\sigma^2}{B}$)

Example: Recession data

TREE RESULTS: CONFUSION MATRICES

			Т		
			Growth	Recession	Mis-Class
Our Preds	Null	Growth	111	26	
		Recession	0	0	18.9%
	Tree	Growth	99	3	
		Recession	12	23	10.9%
	RANDOM	Growth	102	5	
	Forest	Recession	9	21	10.2%
	Bagging	Growth	104	3	
		Recession	7	23	7.3%

Tree results: Sensitivity & specificity

	Sensitivity	Specificity
NULL	0.000	1.000
TREE	0.884	0.891
RANDOM FOREST	0.807	0.918
BAGGING	0.884	0.936

OUT-OF-BAG ERROR ESTIMATION FOR BAGGING

		Т		
		Growth	Recession	Miss-Class
OOB BAGGING	Growth	401	9	
	Recession	23	44	6.71%
Test Bagging	Growth	104	3	
	Recession	7	23	7.3%

RANDOM FOREST IN R

```
require(randomForest)
out.rf = randomForest(X,Y,importance=TRUE,mtry=ncol(X))
class.rf = predict(out.rf,X_0)
```

NOTES:

- The importance statement tells it to produce the variable importance measures
- the mtry = ncol(X) tells randomForest to consider all the features at each split
 (This particular choice corresponds to bagging. Leaving this out uses the default √p)
- randomForest also supports formulae
 out.rf = randomForest(Y~.,data=X)

 However, it can take much longer to run

RANDOM FOREST IN R

```
Call:
randomForest(x = X, y = Y, mtry = ncol(X), importance=T)
              Type of random forest: classification
                     Number of trees: 500
No. of variables tried at each split: 56
        OOB estimate of error rate: 7.17%
Confusion matrix:
   0 1 class.error
0 401 9 0.02195122
1 25 42 0.37313433
```

RANDOM FOREST IN R.

```
> head(importance(out.rf,type=1))#Permutation
           MeanDecreaseAccuracy
Alabama
                      3.7277511
Alaska
                      1.7941463
Arizona
                      2.9659623
Arkansas
                     -0.8341577
California
                      7.2973572
> head(importance(out.rf,type=2))#Mean decrease
           MeanDecreaseGini
                  0.4551073
Alabama
                  1.6440170
Alaska
Arizona
                  0.7025527
Arkansas
                  0.3503138
                  1.4616203
California
#variable importance plot:
varImpPlot(out.rf,type=2)
```

Missing data

In practice, there will often be missing data

Estimating this missing data is known as imputation

RANDOM FOREST provides a method for imputation

It follows two steps:

 na.roughfix: uses either the median or mode to impute missing values

```
X \leftarrow \text{na.roughfix}(X)
```

- 2. rf.impute: Gets the proximity matrix, and re-computes the imputation
 - For numeric features, it uses weighted (with respect to proximity) average
 - For categorical features, it uses the category that maximizes the average proximity

```
require(randomForest)
x = rnorm(12)
xNA = x
xNA[sample(12,2,replace=F)] = NA
X = matrix(xNA,nrow=6)
Y = matrix(x,nrow=6) %*% c(1,.5) + rnorm(6)
XnewImp = rfImpute(X,Y)
#First column is the supervisor vector
Xnew = XnewImp[,-1]
```

```
> X
         NA -1.2589350282
  0.6202015 0.1780122216
-0.9340213 -0.6483047015
  0.1142546 0.0489260332
-1.1039581
                       NA
  0.2064204 0.0007548357
> matrix(x,nrow=6)
0.1485857 - 1.2589350282
0.6202015 0.1780122216
-0.9340213 -0.6483047015
0.1142546 0.0489260332
-1.1039581 -0.2468085986
0.2064204 0.0007548357
```

```
> Xnew
0.1287655 -1.2589350282
0.6202015 0.1780122216
-0.9340213 -0.6483047015
0.1142546 0.0489260332
-1.1039581 -0.2015763074
0.2064204 0.0007548357
```

- Data size/complexity (Does it fit in RAM?)
- Business purpose
 (Is data precious? Development time?)
- Are any observations/features missing a large fraction of values?
- Type of features
 (Any sparsity? Is multivariate normality appropriate?)
- Any atypical missing value indicators?
 (e.g. using -1000 for income to indicate a missing value)