MATH 829: Introduction to Data Mining and Analysis Random forest

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The bootstrap

. We saw before that decision trees often overfit the data.

• We will now discuss techniques that can be used to mitigate that problem.

Bootstrapping: General statistical method that relies on resampling data with replacement.

Idea: Given data (y_i, x_i) , i = 1, ..., n, construct bootstrap samples by sampling n of the observations with replacement (i.e., allow repetitions):

Sample 1	Sample 2	Sample 3
(y_{i_1}, x_{i_1})	(y_{j_1}, x_{j_1})	(y_{k_1}, x_{k_1})
(y_{i_2}, x_{i_2})	(y_{j_2}, x_{j_2})	(y_{k_2}, x_{k_2})
1	:	:
(y_{i_n}, x_{i_n})	(y_{j_n}, x_{j_n})	(y_{k_n}, x_{k_n})

 Each bootstrap sample mimics the statistical properties of the original data.

• Often used to estimate parameter variability (or uncertainty).

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Bagging

Bagging: (bootstrap aggregation) Suppose we have a model $y \approx \hat{f}(x)$ for data $(y_i, x_i) \in \mathbb{R}^{p+1}$.

 $\textcircled{O} \ {\sf Construct} \ B \in \mathbb{N} \ {\sf bootstrap} \ {\sf samples}.$

- Train the method on the b th bootstrap sample to get f^{*b}(x).
- Ompute the average of the estimators:

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

- Bagging is often used with regression trees.
- Can improve estimators significantly.

Note: Each bootstrap tree will typically involve different features than the original, and might have a different number of terminal nodes.

The bagged estimate is the average prediction at \boldsymbol{x} from these \boldsymbol{B} trees.

For classification: Use a majority vote from the B trees.

Example: trees with simulated data (ESL, Example 8.7.1)

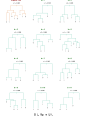
Simulation:

- N = 30 samples with p = 5 features.
- Features from a standard Gaussian distribution with pairwise correlation 0.95.
- Y generated according to
 - $P(Y = 1|X_1 \le 0.5) = 0.2$ $P(Y = 1|X_1 > 0.5) = 0.8.$
- A test sample of size 2,000 was also generated using the same model.
- The test error for the original tree and the bagged tree are reported.

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Example (cont.)

Bootstrap trees:



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Random forests

- Idea of bagging: average many noisy but approximately unbiased models, and hence reduce the variance.
- However, the bootstrap trees are generally correlated.
- Random forests improve the variance reduction of bagging by reducing the correlation between the trees.
- Achieved in the tree-growing process through random selection of the input variables.
- Popular method.

Example (cont.)

Test error:



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Out-of-bag error: Mean prediction error on each training sample x_i , using only the trees that did not have x_i in their bootstrap sample.

Can be used to approximate the prediction error.

Random forests (cont.)



Random forests: Each time a split in a tree is considered, a random selection of m predictors is chosen as split candidates from the full set of p predictors.

• Typical value for m is \sqrt{p} .

• We construct T_1, \ldots, T_B trees using that method on bootstrap samples. The random forest (regression) predictor is

$$\hat{f}_{tf}^{B}(x) = \frac{1}{B} \sum_{b=1}^{B} T_{b}(x).$$

For classification: use majority vote.

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Example (Izenman, 2013)

Diagnostic classification of four childhood tumors (Khan et al., 2001):

- Small, round, blue-cell tumors (SRBCTs) of childhood.
- Four types of SRBCTs (EWS, BL, NB, RMS).
- Tumors have a similar appearance.
- Getting the diagnosis correct impacts directly upon the type of treatment, therapy, and prognosis the patient receives.
- Currently, no single clinical test that can discriminate between these cancers.

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- 83 cases (29 EWS, 11 BL, 18 NB, 25 RMS).
- Gene expression data for 6,567 genes, reduced to 2,308 by requiring a minimum intensity.
- research.nhgri.nih.gov/microarray/Supplement.
- \bullet A random forest was applied to these data using 500 fully grown trees with m=25 variables at each split.
- Able to get a 0% Out-of-bag misclassification rate.

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Boosting

Like bagging, boosting is a general approach that can be applied to many models. Combines weak learners into a single strong learner.

 ${\bf Boosting:}\ {\rm Recursively}\ {\rm fit}\ {\rm trees}\ {\rm to}\ {\rm residuals}.\ ({\rm Compensate}\ {\rm the}\ {\rm shortcoming}\ {\rm of}\ {\rm previous}\ {\rm model}.)$

Input: $(y_i, x_i) \in \mathbb{R}^{p+1}$, i = 1, ..., n. Initialize $\hat{f}(x) = 0$, $r_i = y_i$. For b = 1, ..., B:

- Fit a tree estimator f^b with d splits to the training data.
- Update the estimator using:

$$\hat{f}(x) \leftarrow \hat{f}(x) + \lambda \cdot \hat{f}^{b}(x).$$

Opdate the residuals:

$$r_i \leftarrow r_i - \lambda \cdot \hat{f}^b(x_i)$$

Output: Boosted tree:

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

Note: $\lambda > 0$ is a learning rate.

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Boosting (cont.)

Can use many small trees (by choosing d small) and learn slowly (λ small) to avoid overfitting.

Choosing the parameters:

- O Number of trees B: choose by cross-validation.
- Number of splits: can use a small value (e.g. d = 1).
- Learning rate: can use 0.01, 0.001. Note: A small λ will generally require a larger B...

Gradient boosting: More generally, one can work with a general loss function (instead of sum of squares) and replace the residuals with the (negative) of the gradient of the loss function.

Relative importance of predictor variables

- The previous methodologies can improve decision trees considerably.
- However, we lose the nice interpretability of decision trees.
 A relative importance of each predictor can be computed to help understand a model with multiple trees.
 - Let T be a (binary) decision tree with J-1 internal nodes.
 - At each internal node t, a variable $X_{v(t)}$ is split, resulting in an improvement \hat{t}_i^2 in squared error.
 - . We define a measure of relevance of X1 by

$$I_l^2(T) := \sum_{t=1}^{J-1} \hat{\iota}_t^2 \cdot I(v(t) = l).$$

In other words, we add up the improvements at the nodes where X_l is split.

Relative importance of predictor variables (cont.)

 \bullet Similarly, in a model obtained from M trees (e.g. bagging, random forest), we use:

$$I_l^2 = \frac{1}{M} \sum_{m=1}^M I_l^2(T_m)$$

• Taking the square root of the relevance measure, we obtain the relevance of X_l .

 Typically, we do not report the actual relevance of a variable. We rather report the percentage of relevance of a given variable with respect to the variable with the largest relevance.

Relative importance of predictor for the spam data

