# Random Forests™

Ensembles of bootstrapped, weakened trees

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# Bias / Variance (again)

Expected prediction error at  $x_0$  $E[(Y - \hat{f}_k(x_0))^2 | X = x_0]$ 

- For a single test vector  $x_0$ , **bias** "is the squared difference between the true mean  $f(x_0)$  and the expected value of the estimate" [ESL book p37] (at least under MSE error) (How far off is the expected prediction from true answer?)
- Variance is the variance of the estimates for  $x_0$  from models trained using tweaked training data (most common terminology)
- But high variance also implies model parameters (tree structure) vary a lot if we tweak the training data

Leo Breiman (1996) introduced bagging then Random Forests (2001)

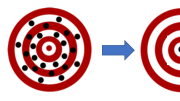
https://www.stat.berkeley.edu/~breiman/randomforest2001.pdf

#### RF motivation

- Decision trees can often get training errors close to zero because we can grow very large trees to partition the feature space into tiny regions with 1 or just a few observations / samples; trees are very accurate on the training set and have low bias
- The downside is that decision trees overfit like mad: decision trees have high variance and don't generalize well



# How can we increases generality?



- Goal: keep the high accuracy, but increase the generality
- Recall: simplifying models often increases generality at cost of some bias
- So, let's weaken our decision tree model but in a way that makes predictions noisier though with same prediction expected value
- To compensate for the noise and claw back some accuracy, make an ensemble of such weakened trees; ensemble predicts average or majority vote of trees
- Averaging predictions reduces variance without changing bias so ensemble is accurate on average
- The expected value of full strength model is same as expectation of weakened model's prediction



# The key trick is amnesia



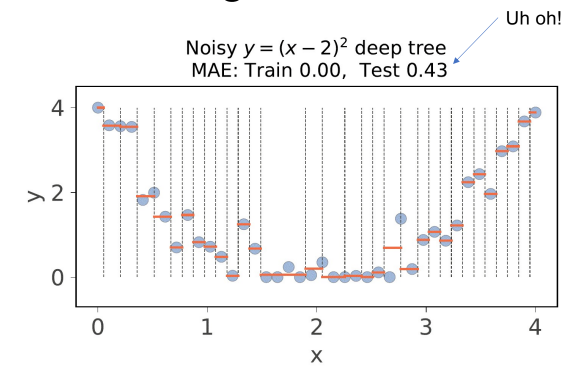
- Random forests are all about adding a bit of amnesia to the training process
- We will weaken trees by training each on a randomly selected subset of the training data: bagged trees
- Further, we will have training purposely forget about some features as we create decision nodes: random forests

# Analogy: Crowdsourcing SF house prices

- Recruit multiple real estate agents to build house price models in their heads by visiting lots of houses; then each agent can estimate prices of unvisited houses
- Agents choose and examine house subsets independently
- There will be some overlap in visited houses sets but the subsets will be independent and identically distributed (i.i.d.)
- An agent trained on large house set is very accurate
- An agent trained on an i.i.d. subset is not biased (they have same expectation) but is less accurate—a prediction for one house might be too low but a prediction for another house might be too high
- The variance of the ensemble average will be much tighter than the variance of an individual tree's prediction
- Averaging all agents' predictions reduces variance and is unbiased

# Ex: Overfit decision trees regressors

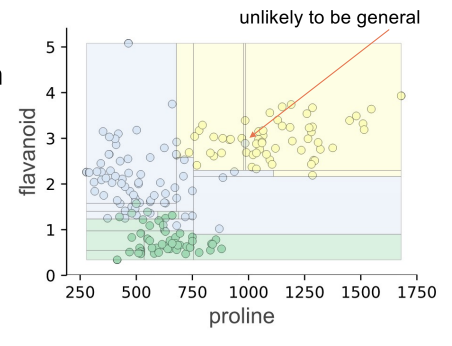
- Animation shows 1D feature space partitioning of i.i.d. sample sets
- Slightly different training data sets can yield very different decision trees
- Clearly the trees have gotten way too specific to the data set
- Notice how the training error is 0 but (20% hold out) test error is terrible!





#### Ex: Overfit decision tree classifiers

- Here is a previous example where partitioning trapped a lonely blue in a sea of yellow
- In practice, we're given just one data set so let's do some sampling to get some i.i.d. "copies"
- Then see how different data sets give different partitioning (from different trees)



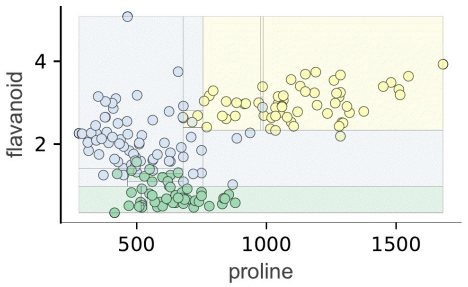
# Partitioning from bootstrapped data

 Bootstrap (X, y) to simulate multiple i.i.d. data sets<sup>†</sup>

• Each set gets ~63% of unique (X, y) data (sample n records with replacement)

- Animation shows 2D feature space partitions from various bootstraps
- Partitioning clearly varies a lot between bootstraps
- OOB == "out of bag" (more later)

Bagged tree classifier partitioning Accuracy: train 100.0% OOB 85.7% Showing all (X, y) data points



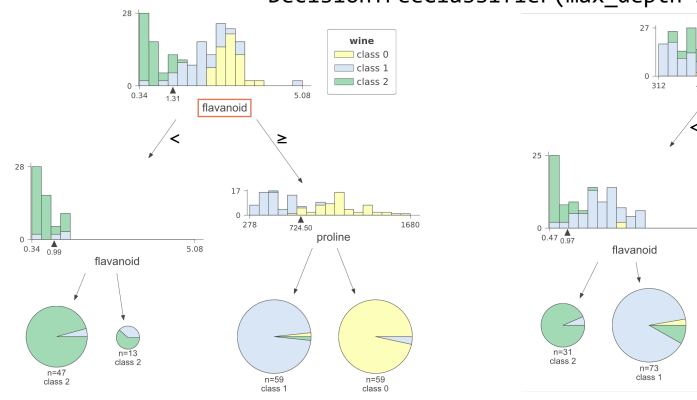
<sup>†</sup>Bootstraps are technically *conditionally independent*, conditioned on nature of original *X*, which could be weird by chance or by the way it was collected/obtained.

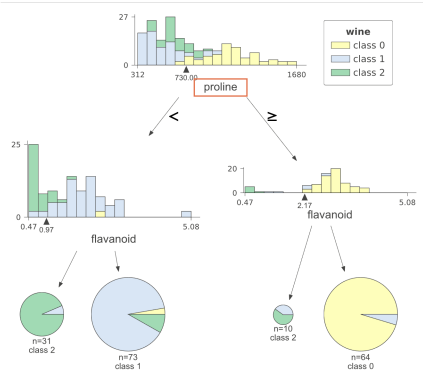


Uh oh! Varies

# Bootstrapping gives slightly different trees

DecisionTreeClassifier(max\_depth=2)





Same tree construction algorithm running on slightly different bootstraps



# Aside: Code for bootstrapping

#### NumPy

```
# Bootstrap: sample with replacement
n = len(y)
idx = np.random.randint(0,n,size=n)
X_train = X[idx]
y_train = y[idx]
```

```
# get OOB (out-of-bag) samples
mask = np.ones(n, dtype=bool)
mask[idx] = False
X_test = X[mask]
y_test = y[mask]
```

#### **Pandas**

```
# If data in dataframe
df = df.sample(len(df), replace=True)
```

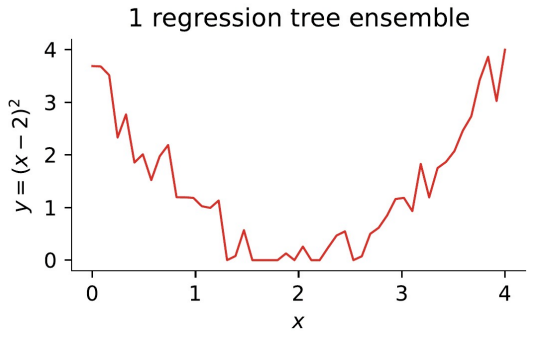


# Bagged trees

Training trees on bootstrapped samples and aggregating predictions

# Ensemble of high-variance regression trees

- Animation shows how averaging the prediction of an ensemble of overfit trees actually produces a reasonable combined prediction
- As we add trees, the average prediction (red line) smooths out to reveal the underlying quadratic distribution from which we draw noisy samples
- Note: variance of individual tree predictions stays high regardless of number of trees, but the variance of the ensemble average tightens (the magic of C.L.T.)

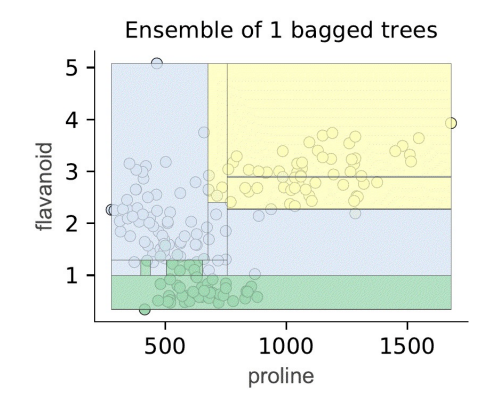


Repeatedly drawing new noisy training sets



# Ensemble of high-var. classification trees

- Animation shows overlapping prediction regions from multiple classifier trees
- Training data for each tree is bootstrapped from the original (X, y) data
- As we add trees, the averaged prediction regions become more stable and the decision boundaries more complex
- "Bag" is bootstrap aggregation

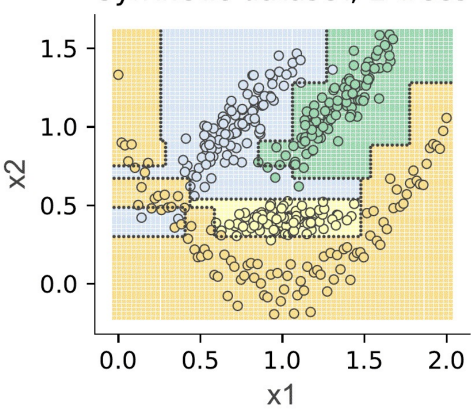




# Ensemble classifier on synthetic data set

- Animation shows prediction regions from multiple bagged classifier trees
- Colored tiles indicate the probabilities of the various classes; e.g., yellow-orange color indicates uncertainty between those two classes
- What's prob. of class k at tile?
  - proportion of trees that predict k

Synthetic dataset, 1 trees



Animation uses probability space visualization I added to dtreeviz

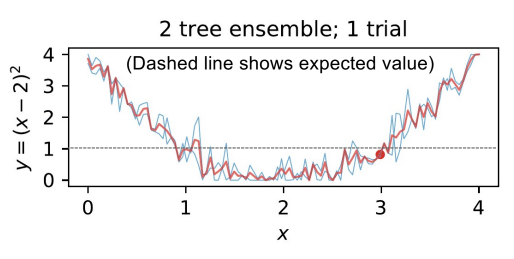
#### Ensemble's effect on bias and variance

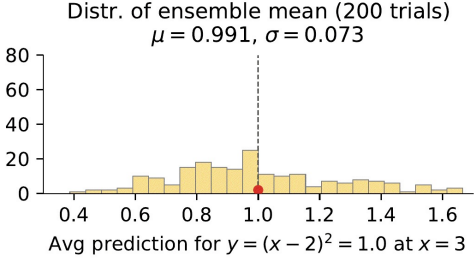
- Train T trees on T i.i.d. X data sets
- Central limit theorem says that if variance of an i.i.d. random variable is  $\sigma^2$ , the variance of the average of T such vars is  $\sigma^2/T$
- So, as we add trees, the variance of the ensemble prediction will shrink, which means better generality
- After, say, 100 trees though we're not going to get a more general model, but will get better estimates of the true prediction for a single test record (squeezing out some more noise)
- The average of the tree predictions is the same as the <u>expected</u> <u>prediction</u> from any tree trained on one of the *X* sets (since i.d.)
- If weakened trees had different expectations, adding trees would increase bias



### Ex: variance of ensemble prediction

- Animation shows tree and ensemble predictions on left for T trees;
   variance of predictions in blue tree predictions doesn't change with the number of trees but red line get tighter / less noisy with more trees
- At x = 3, expected value of ensemble is 1.0; create 200 separate ensembles of size T and compute variance of ensemble predictions at x = 3; distribution of ensemble average shown on the right





### Problem: trees are not independent thinkers

- With real estate agent analogy, we implicitly assumed agents were independent thinkers, and not clones
- But, decision trees are like robot clones and, given the same bit of data, yield the exact same bit of tree
- Imagine worst case: bootstrapping yields *T* identical sets so ensemble gives exactly the same prediction as any single tree
- In practice, if there is one strongly predictive var out of p, then all trees would be similar; initial root splits, and many others, would likely be same

# Random Forests

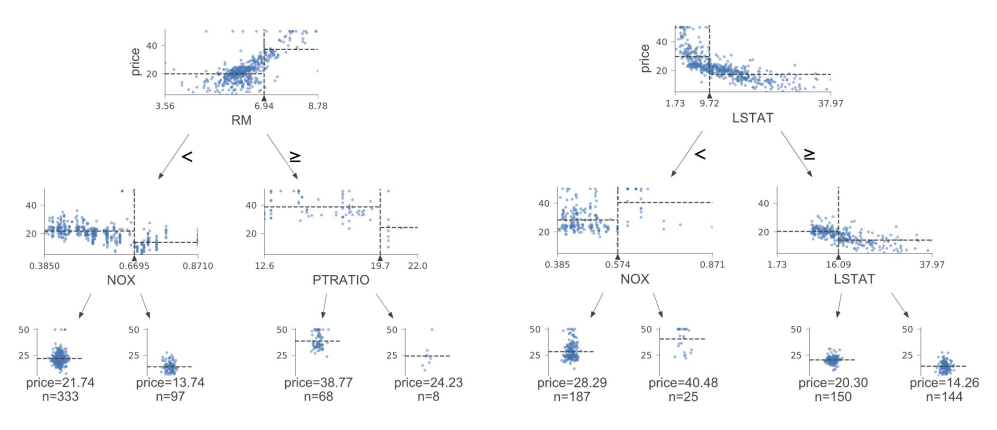
Ensembles of de-correlated bagged trees

# Making trees independent thinkers

- Bagging overcomes most of the overfitting, but we can improve generality a little by further weakening the tree training process itself in an effort to make trees think more independently
- Restrict the available features when searching for a decision node split; choose from m randomly selected features (amnesia again!)
- Choose max features per split,  $m \le p$ , such as  $m = \operatorname{sqrt}(p)$
- Make sure chance of selecting predictive variables (m/p) is high enough to find predictive variables (See ESLII p596)
- Let validation error be your guide to choosing m
- A random forest is then just an ensemble of decision trees trained on bootstraps and whose feature selection strategy has a bit of amnesia

#### Ex: Effect of limiting feature set but using full data set

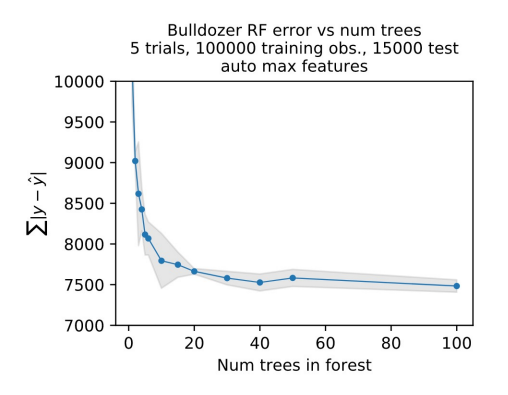
2 trees trained on **entire** Boston set with m=5 (of 13)

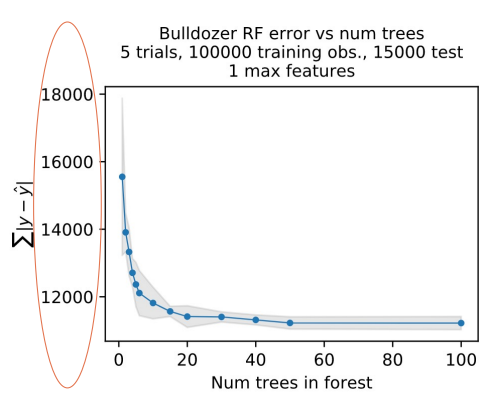


Choose from 5 randomly selected features during EACH split



# If max\_features too low, bad accuracy

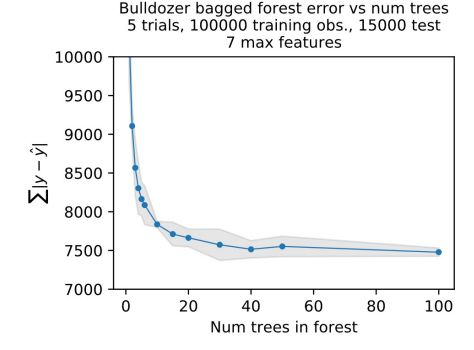




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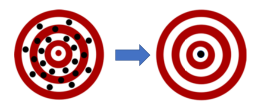
# Effect of forest size on accuracy

- Why does accuracy improve greatly (initially) as we add trees?
  - Each tree sees only 2/3 of data so adding bootstrapped trees increases use of training data
  - Variance is tightening quickly as we average even just a few trees
- Why does accuracy asymptotically approach a minimum instead of continual improvement?
  - With enough trees, ensemble sees 100% of the training data; it's approaching the accuracy of single decision tree in ideal world
  - We've squeezed out all bias and variance that we can with this model





# Properties (see Breiman 2001)



- p4 "Random forests do not overfit as more trees are added" Why?
  - New trees get averaged in so each additional tree has less individual effect
  - New trees <u>balance each other out</u>, one might be too high, another too low
- p7 "It's relatively robust to y outliers and X noise" **Why**?
  - y outliers get shunted to their own leaf since doing so reducing loss function, particularly if squared-error is used
  - Noise X variables aren't predictive so not chosen as split vars
- p10 Bagging helps more, the more unstable the model. Why?
  - Averaging is a smoothing operator, squeezing predictions to true value
  - If model is low variance already, there is no point in bagging

# Properties continued

- RFs are scale and range insensitive in features and target y Why?
  - Comparing feature values in decision nodes, not doing math on them
  - Computing mean or mode of y to predict
- ESLII p596 "Classifiers are less sensitive to variance [than regressors]" Why?
  - (not sure haha) I believe it has something to do with mode vs mean (mode is same until a threshold whereas mean is influenced by any value added, unless it is also the mean)

# Bootstrapping vs subsampling

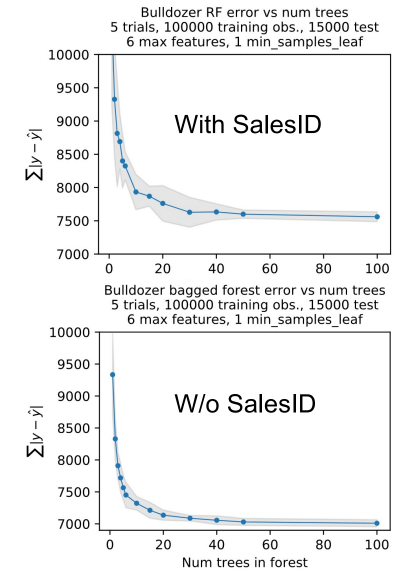
- Bootstrapping is sampling with replacement vs subsampling w/o replacement
- Friedman and Hall (2000): subsampling also works, showing that training trees with n/2 subsamples is similar in bias/variance to bagging <a href="http://statweb.stanford.edu/~jhf/ftp/bag.pdf">http://statweb.stanford.edu/~jhf/ftp/bag.pdf</a>
  - Smaller training set is a big win in terms of speed
  - Using even smaller fractions of n improve generality (reduce variance) because trees are less correlated (they work on different data chunks); note that each tree would become less accurate as n subsample size decreases

# RF Tuning strategy

- Good news: very little tuning needed
- Goal: minimize validation error
- Start with maybe 20 trees and work upwards til validation error stops getting better; or just pick 100
- Sklearn uses max\_features= sqrt(p) by default; try dropping this to log(p), or similar; ESLII suggests p/3 for regression and sqrt(p) for classification
- Try adjusting min samples per leaf: 1, 3, 5, 10, 25, 100
- Can also try grid search, but I never bother;
   Start with num trees, then tune the others

# Feature engineering beats model tuning

- SalesID: unique record ID, and is never seen again in future predictions
- Is that useful for prediction? No
- Does the model think it's useful? Yes
- Model is overfit not on noise but on falsely-predictive feature
  - Could be that sales ID correlates with inflation or change in type of models sold in auction creates "trend" in sale prices
- A case where using LESS data improves the model a lot (\$500 diff)
- Dropping useless features also often gives a small bump



# The RF algorithms



# Fitting RFs

```
Algorithm: RFfit(X, y, loss, ntrees, max\_features, min\_samples\_leaf)

for i = 1..ntrees do

X', y' = bootstrap(X, y, size = |X|)

T_i = RFdtreefit(X', y', loss, max\_features, min\_samples\_leaf)

end
```

For regression, pass in loss = MSE or stddev For classifier, pass in loss = gini



# Fitting a single tree in RF

Same as decision tree except we pass max\_features to RFbestsplit()

```
Algorithm: RFdtreefit(X, y, loss, max\_features, min\_samples\_leaf)

if |X| \leq min\_samples\_leaf then return Leaf(y)

col, split = RFbestsplit(X, y, loss, max\_features)

if col = -1 then return Leaf(y)

lchild = RFdtreefit(X[X_{col} \leq split], y[X_{col} \leq split], ...)

rchild = RFdtreefit(X[X_{col} > split], y[X_{col} > split], ...)

return DecisionNode(col, split, lchild, rchild)
```

# Finding best split in decision node in RF

```
Algorithm: RFbestsplit(X, y, loss, max\_features)
  best = (col = -1, split = -1, loss = loss(y))
                                                                         Only diff with decision tree
  vars = pick \ max_features \ variables \ from \ all \ p
  for col \in vars do
                                                                         Pick, say, 11 not all possible X
    candidates = randomly pick k \ll n values from X_{col}
                                                                         values. We get better generality
    foreach split \in candidates do
                                                                         and code is much faster!
       yl = y[X_{col} \leq split]
       yr = y[X_{col} > split]
       if |yl| < min\_samples\_leaf or |yr| < min\_samples\_leaf then continue
       l = \frac{|yl| \times loss(yl) + |yr| \times loss(yr)}{|y|}
                                           (weighted average of subregion losses)
       if l = 0 then return col, split
                                                                         Should pick midpoint between
       if l < best[loss] then best = (col, split, l)
                                                                         split value and next smallest X
    end
  end
  return best/col/, best/split/
                                                                                     TY OF SAN FRANCISCO
```

# Simplest RF prediction (ESLII p588)

- But doesn't use all information to make best prediction
- Should use weighted averages / votes

Regression: 
$$\hat{f}_{rf}^B(x) = \frac{1}{B} \sum_{b=1}^B T_b(x)$$
.

Classification: Let  $\hat{C}_b(x)$  be the class prediction of the bth random-forest tree. Then  $\hat{C}_{rf}^B(x) = majority\ vote\ \{\hat{C}_b(x)\}_1^B$ .

# RF prediction

Weighted average of y values among the leaves reached by running x down each tree

```
Algorithm: RFpredict_{regr}(\{T_1..T_{ntrees}\}, x)

Let leaves = \{leaf(T_t, x) \ \forall t = 1..ntrees\}

nobs = \sum_{t=1}^{ntrees} |leaves_t|

ysum = \sum_{t=1}^{ntrees} \sum_{y \in leaves_t} y

return \frac{1}{nobs}ysum Algorithm:
```

Count all y votes among the leaves reached by running x down each tree

```
Algorithm: RFpredict_{class}(\{T_1..T_{ntrees}\}, x)
counts[k] = 0 \ \forall \ classes \ k
foreach t = 1..ntrees do
leaf = leaf(T_t, x) \qquad (leaf \ reached \ by \ x)
foreach y \in leaf do
counts[y] += 1 \qquad (track \ count \ of \ leaf \ modes)
end
end
end
return argmax(counts)
```

### Extremely randomized trees (Geurts et al 2006)

- The variable/value pair is highly sensitive to the training set, and responsible for much of the error rate
- "The optimal cut-point was shown to depend very strongly on the particular learning sample used...this cut-point variance appeared to be responsible for a significant part of the error rates of tree-based methods." <a href="https://link.springer.com/article/10.1007/s10994-006-6226-1">https://link.springer.com/article/10.1007/s10994-006-6226-1</a>
- Geurts wondered if more randomness could reduce variance further
- Pick random split value in min(X[:,j]) .. max(X[:,j]), ignoring y!
- Like RF, select  $m \le p$  variables and choose var/value with lowest loss
- Fits using entire *X* training set, not bootstrap and not subsample (trying to increase accuracy of prediction/decrease noise)
- Our use of just 11 (not n) X candidate values in the project is similar (an effort to reduce variance and increase speed)

# RF advantages

- Ensemble of decision trees trained on different bootstraps that sometimes forgetting about features during training
- Prediction is ensemble average or majority vote (weighted)
- Easy to understand, efficient, excellent accuracy, interpretable
- Very little tuning is required
- Gracefully handles label-encoded categorical variables, no need to normalize numerical variables
- Robust to noise in X, y and nonpredictive variables
- Built-in out of bag validation sets
- Negative: cannot extrapolate beyond support data