Machine learning

Random forests and gradient-boosted trees

July 14th, 2021

Decision trees review

Decision trees partition training data into **homogenous nodes** / **subgroups** with similar response values

Pros

- Decision trees are **very easy to explain** to non-statisticians.
- Easy to visualize and thus easy to interpret without assuming a parametric form

Cons

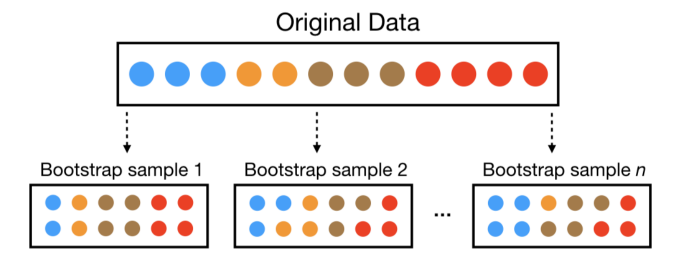
- High variance, i.e. split a dataset in half and grow tress in each half, the result will be very different
- Related note they generalize poorly resulting in higher test set error rates

But there are several ways we can overcome this via **ensemble models**

Bagging

Bootstrap aggregation (aka bagging) is a general approach for overcoming high variance

• **Bootstrap**: sample the training data *with replacement*



• **Aggregation**: Combine the results from many trees together, each constructed with a different bootstrapped sample of the data

Bagging algorithm

Start with a **specified number of trees** *B*:

- For each tree b in $1, \ldots, B$:
 - Construct a bootstrap sample from the training data
 - o Grow a deep, unpruned, complicated (aka really overfit!) tree

To generate a prediction for a new point:

- **Regression**: take the **average** across the B trees
- Classification: take the majority vote across the B trees
 - assuming each tree predicts a single class (could use probabilities instead...)

Improves prediction accuracy via **wisdom of the crowds** - but at the expense of interpretability

• Easy to read one tree, but how do you read B=500?

But we can still use the measures of **variable importance** and **partial dependence** to summarize our models

Random forests algorithm

Random forests are an extension of bagging

- For each tree b in $1, \ldots, B$:
 - Construct a bootstrap sample from the training data
 - Grow a deep, unpruned, complicated (aka really overfit!) tree **but with a twist**
 - \circ **At each split**: limit the variables considered to a **random subset** m_{try} of original p variables

Predictions are made the same way as bagging:

- **Regression**: take the **average** across the B trees
- ullet Classification: take the majority vote across the B trees

Split-variable randomization adds more randomness to make each tree more independent of each other

Introduce m_{try} as a tuning parameter: typically use p/3 (regression) or \sqrt{p} (classification)

 $ullet m_{try}=p$ is bagging

Example data: MLB 2021 batting statistics

Downloaded MLB 2021 batting statistics leaderboard from Fangraphs

```
library(tidvverse)
mlb_data <- read_csv("http://www.stat.cmu.edu/cmsac/sure/2021/materials/data/fg_batting_2021.csv")</pre>
  janitor::clean_names() %>%
  mutate at(vars(bb percent:k percent), parse number)
model mlb data <- mlb data %>%
  dplyr::select(-name, -team, -playerid)
head(model mlb data)
## # A tibble: 6 × 20
                                    sb bb percent k per...¹ iso babip
##
                   hr
                          r rbi
             ра
                                                                       avg
##
    <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
                                            <dbl>
                                                    <dbl> <dbl> <dbl> <dbl> <dbl> <dbl>
## 1
       82
            354
                   27
                         66
                              69
                                             14.4 17.2 0.336 0.346 0.336 0.438
## 2
                                                     28.1 0.395 0.333 0.302 0.385
       68
            288
                   27
                         66
                               58
                                    18
                                             12.5
## 3
       79 347
                   16
                               52
                                    0
                                             13.5
                                                          0.231 0.324 0.298 0.398
                         61
## 4
       82
            372
                         63
                               54
                                            8.9
                                                     23.9 0.256 0.329 0.286 0.349
                   21
                                    10
## 5
       78 342
                   23
                         67
                               51
                                    16
                                             13.2
                                                     24.3 0.313 0.306 0.278 0.386
## 6
            322
                               67
                                    12
                                             11.2
       82
                   31
                         60
                                                     28
                                                          0.418 0.29 0.277 0.363
## # ... with 8 more variables: slg <dbl>, w oba <dbl>, xw oba <dbl>, w rc <dbl>,
## #
      bs r <dbl>, off <dbl>, def <dbl>, war <dbl>, and abbreviated variable name
       <sup>1</sup>k percent
## #
```

Example using ranger

ranger package is a popular / fast implementation (see randomForest for the original)

```
library(ranger)
init_mlb_rf <- ranger(war ~ ., data = model_mlb_data, num.trees = 50, importance = "impurity")</pre>
init mlb rf
## Ranger result
##
## Call:
## ranger(war ~ ., data = model_mlb_data, num.trees = 50, importance = "impurity")
##
## Type:
                                      Regression
## Number of trees:
                                      50
## Sample size:
                                      135
## Number of independent variables:
                                      19
## Mtry:
## Target node size:
## Variable importance mode:
                                      impurity
## Splitrule:
                                      variance
## 00B prediction error (MSE):
                                      0.1919574
## R squared (00B):
                                      0.8314733
```

Out-of-bag estimate

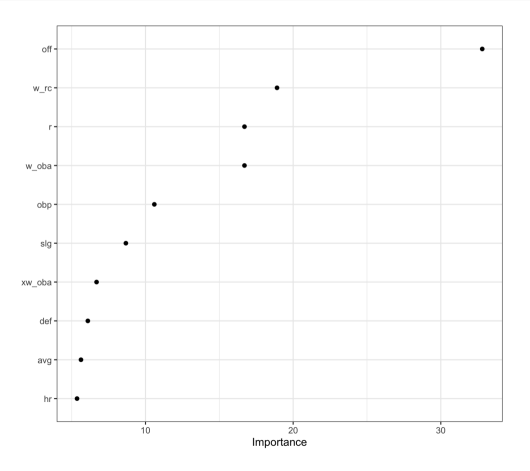
Since the trees are constructed via bootstrapped data (samples with replacements) - each sample is likely to have duplicate observations / rows

Out-of-bag (OOB) - original observations not contained in a single bootstrap sample

- Can use the OOB samples to estimate predictive performance (OOB becomes better with larger datasets)
- On average pprox 63% of original data ends up in any particular bootstrap sample

Variable importance

```
library(vip)
vip(init_mlb_rf, geom = "point") + theme_bw()
```

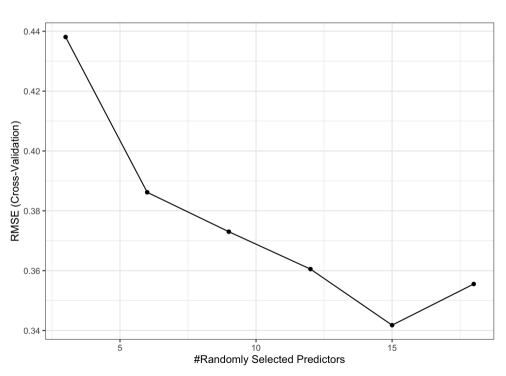


Tuning random forests

Unfortunately caret does not let you know tune number of trees - typically the error goes down with more (Exercise: check out CV performance as a function of the number trees on your own, compare with OOB error)

- Important: m_{try}
- Marginal: tree complexity, splitting rule, sampling scheme

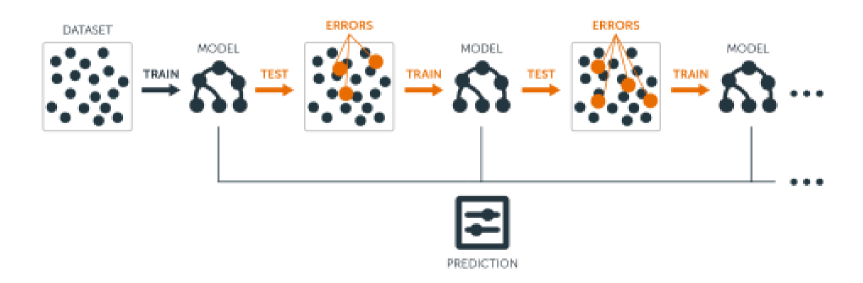
```
ggplot(caret_mlb_rf) + theme_bw()
```



Boosting

Build ensemble models sequentially

- start with a **weak learner**, e.g. small decision tree with few splits
- each model in the sequence *slightly* improves upon the predictions of the previous models **by focusing on** the observations with the largest errors / residuals



Boosted trees algorithm

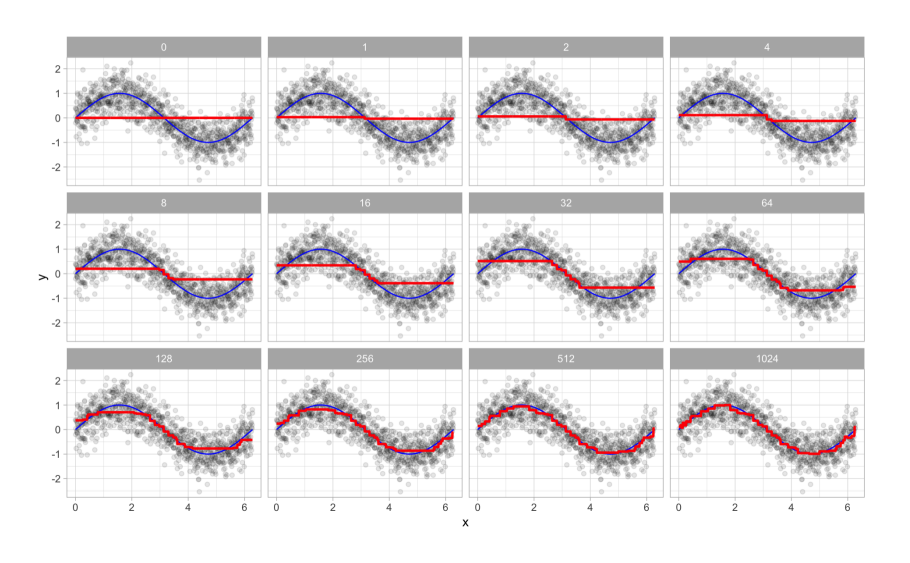
Write the prediction at step t of the search as $\hat{y}_i^{(t)}$, start with $\hat{y}_i^{(0)} = 0$

- ullet Fit the first decision tree f_1 to the data: $\hat{y}_i^{(1)} = f_1(x_i) = \hat{y}_i^{(0)} + f_1(x_i)$
- ullet Fit the next tree f_2 to the residuals of the previous: $y_i \hat{y}_i^{(1)}$
- ullet Add this to the prediction: $\hat{y}_i^{(2)}=\hat{y}_i^{(1)}+f_2(x_i)=f_1(x_i)+f_2(x_i)$
- ullet Fit the next tree f_3 to the residuals of the previous: $y_i \hat{y}_i^{(2)}$
- ullet Add this to the prediction: ${\hat y}_i^{(3)}={\hat y}_i^{(2)}+f_3(x_i)=f_1(x_i)+f_2(x_i)+f_3(x_i)$

Continue until some stopping criteria to reach final model as a sum of trees:

$$\hat{y_i} = f(x_i) = \sum_{b=1}^B f_b(x_i)$$

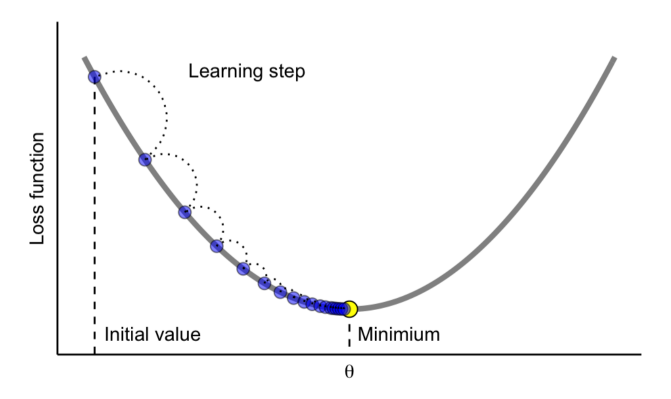
Visual example of boosting in action



Gradient boosted trees

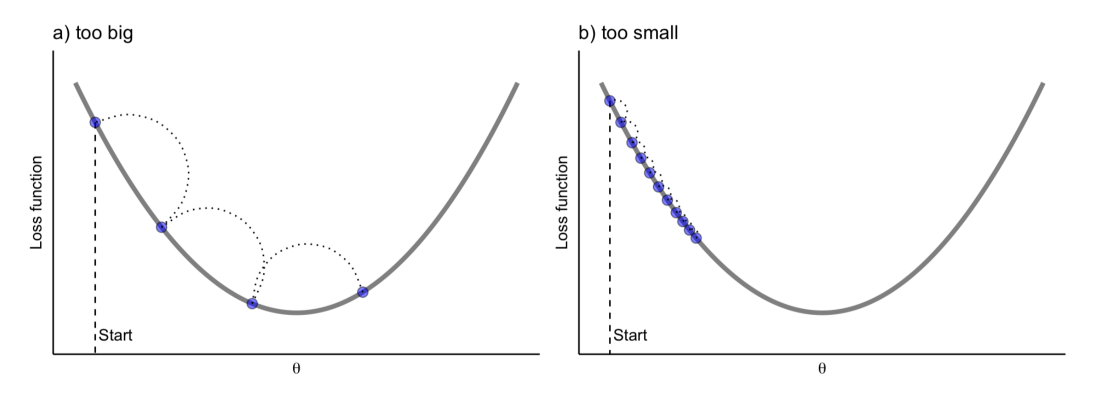
Regression boosting algorithm can be generalized to other loss functions via **gradient descent** - leading to gradient boosted trees, aka **gradient boosting machines (GBMs)**

Update the model parameters in the direction of the loss function's descending gradient



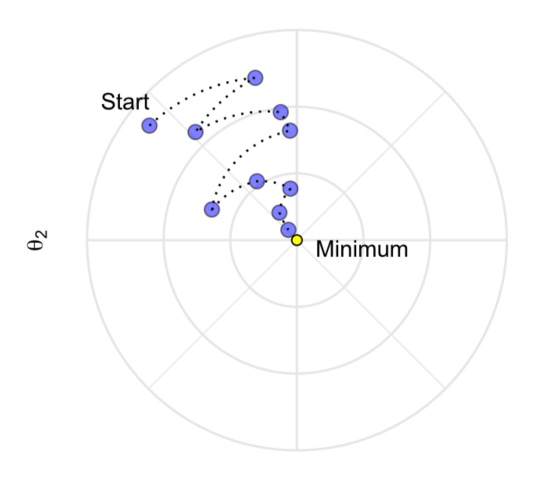
Tune the learning rate in gradient descent

We need to control how much we update by in each step - the learning rate



Stochastic gradient descent can help with complex loss functions

Can take random samples of the data when updating - makes algorithm faster and adds randomness to get closer to global minimum (no guarantees!)



eXtreme gradient boosting with XGBoost



Tuning GBMs with xgboost

XGBoost (extreme gradient boosting) is a very powerful, efficient boosting library that is available to use within R via the xgboost package

What we have to consider tuning (our **hyperparameters**):

- number of trees B (nrounds)
- learning rate (eta), i.e. how much we update in each step
- these two really have to be tuned together
- complexity of the trees (depth, number of observations in nodes)
- XGBoost also provides more **regularization** (via gamma) and early stopping

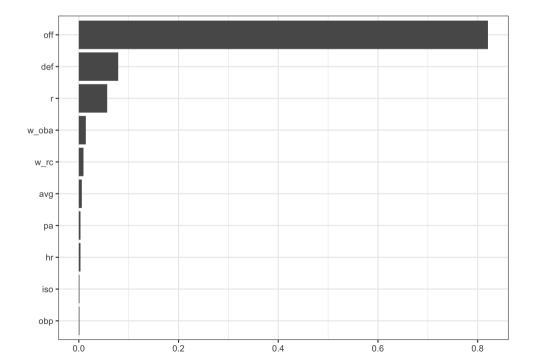
More work to tune properly as compared to random forests

- But GBMs have more flexibility in their usage for particular objective functions
- Insert with great power comes great responsibility meme

XGBoost example

```
## [16:50:58] WARNING: src/c_api/c_api.cc:935: `ntree_limit` is deprecated, use `iteration_range` instead
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```

XGBoost example



XGBoost example

