Unit 4 Lecture 4: Boosting

November 11, 2021

Today, we will learn how to train and tune boosting models using the gbm package.

First, let's load some libraries:

```
library(gbm) # install.packages("gbm")
library(tidyverse)
```

Boosting models for regression

We will continue using the Hitters data from the ISLR package, splitting into training and testing:

```
Hitters = ISLR2::Hitters %>%
  as_tibble() %>%
  filter(!is.na(Salary)) %>%
  mutate(Salary = log(Salary)) # log-transform the salary
Hitters
```

```
## # A tibble: 263 x 20
##
      AtBat Hits HmRun
                                   RBI Walks Years CAtBat CHits CHmRun CRuns
                           Runs
##
       <int> <int> <int> <int> <int>
                                       <int>
                                              <int>
                                                      <int> <int>
                                                                     <int> <int> <int>
##
        315
                                           39
                                                       3449
                                                               835
                                                                        69
                                                                              321
                                                                                    414
    1
                81
                        7
                              24
                                    38
                                                  14
    2
                       18
                                    72
                                           76
                                                   3
                                                       1624
                                                               457
                                                                        63
                                                                              224
##
        479
               130
                              66
                                                                                    266
                                                       5628
                                                              1575
                                                                              828
##
    3
        496
               141
                       20
                              65
                                    78
                                           37
                                                  11
                                                                       225
                                                                                    838
##
    4
        321
                87
                       10
                              39
                                    42
                                           30
                                                   2
                                                        396
                                                               101
                                                                        12
                                                                               48
                                                                                     46
##
    5
        594
                        4
                              74
                                    51
                                           35
                                                       4408
                                                              1133
                                                                              501
                                                                                    336
               169
                                                  11
                                                                        19
##
    6
        185
                37
                        1
                              23
                                     8
                                           21
                                                   2
                                                        214
                                                                42
                                                                               30
                                                                                      9
                                                                         1
    7
        298
                                    24
                                            7
                                                   3
                                                        509
                                                                                     37
##
                73
                        0
                              24
                                                               108
                                                                         0
                                                                               41
##
    8
        323
                81
                        6
                              26
                                    32
                                            8
                                                   2
                                                        341
                                                                86
                                                                         6
                                                                               32
                                                                                     34
##
    9
        401
                92
                       17
                              49
                                    66
                                           65
                                                  13
                                                       5206
                                                             1332
                                                                       253
                                                                              784
                                                                                    890
        574
               159
                       21
                             107
                                    75
                                           59
                                                  10
                                                       4631
                                                              1300
                                                                        90
                                                                             702
                                                                                    504
   # ... with 253 more rows, and 8 more variables: CWalks <int>, League <fct>,
       Division <fct>, PutOuts <int>, Assists <int>, Errors <int>, Salary <dbl>,
       NewLeague <fct>
```

```
set.seed(1) # set seed for reproducibility
train_samples = sample(1:nrow(Hitters), round(0.8*nrow(Hitters)))
Hitters_train = Hitters %>% filter(row_number() %in% train_samples)
Hitters_test = Hitters %>% filter(!(row_number() %in% train_samples))
```

Training a gradient boosting model

Arguments:

- distribution: "gaussian" for continuous responses; "bernoulli" for binary responses
- n.trees: maximum number of trees to try; defaults to 100 but this is normally not enough trees
- interaction.depth: interaction depth; defaults to 1
- shrinkage: shrinkage parameter lambda: defaults to 0.1

- bag.fraction: subsampling fraction pi; defaults to 0.5
- cv.folds: number of CV folds to use; defaults to 0 (i.e. no CV)
- train.fraction: fraction of data to use as training; rest used as validation set

```
# read more about the inputs and outputs, bells and whistles of gbm
?gbm
```

Training the model:

We can visualize the CV error using gbm.perf, which both makes a plot and outputs the optimal number of trees:

```
opt_num_trees = gbm.perf(gbm_fit)
        0.7
                                                                                                           ı
        9
        Ö
Squared error loss
        S
        Ö.
        0.4
        0.3
        0.2
                0
                                 20
                                                   40
                                                                     60
                                                                                       80
                                                                                                        100
                                                         Iteration
```

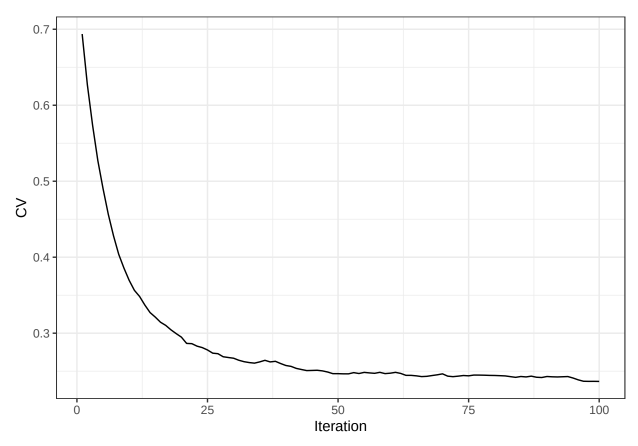
```
opt_num_trees
```

[1] 100

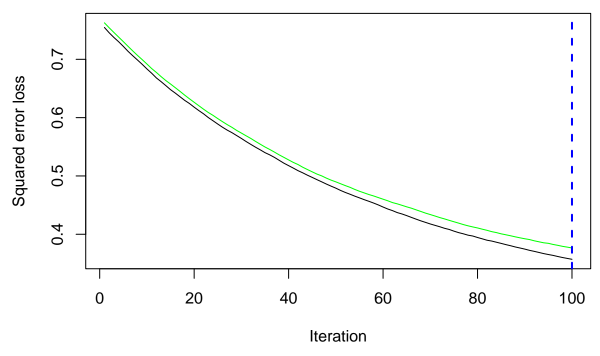
The green curve is the CV error; the black curve is the training error. The dashed blue line indicates the minimum of the CV error.

Note that gbm_fit\$cv.error also contains the CV errors, so these can be plotted manually as well:

```
ntrees = 100
tibble(Iteration = 1:ntrees, CV = gbm_fit$cv.error) %>%
    ggplot(aes(x = Iteration, y = CV)) + geom_line() +
    theme_bw()
```

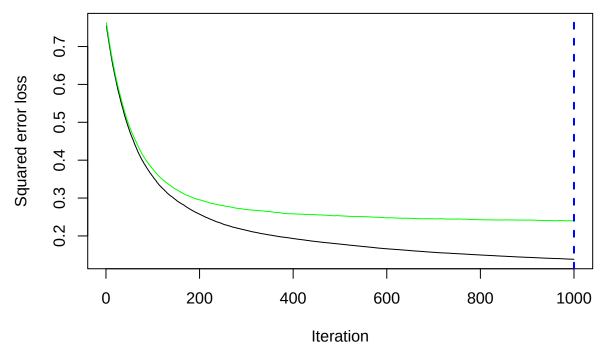


We want to make sure there are enough trees that the CV curve has reached its minimum. For example, suppose we had chosen a smaller shrinkage parameter, e.g. 0.01:



[1] 100

We see that 100 is not enough trees for lambda = 0.01. In this case, we would need to increase the number of trees:



[1] 1000

Tuning the interaction depth

The quick way to tune the interaction depth is to try out a few different values:

```
set.seed(1)
gbm_fit_1 = gbm(Salary ~ .,
              distribution = "gaussian",
              n.trees = 100,
              interaction.depth = 1,
              shrinkage = 0.1,
              cv.folds = 5,
              data = Hitters_train)
gbm_fit_2 = gbm(Salary ~ .,
              distribution = "gaussian",
              n.trees = 100,
              interaction.depth = 2,
              shrinkage = 0.1,
              cv.folds = 5,
              data = Hitters_train)
gbm_fit_3 = gbm(Salary ~.,
              distribution = "gaussian",
              n.trees = 100,
              interaction.depth = 3,
              shrinkage = 0.1,
              cv.folds = 5,
              data = Hitters_train)
```

We can extract the CV errors from each of these objects by using the cv.error field:

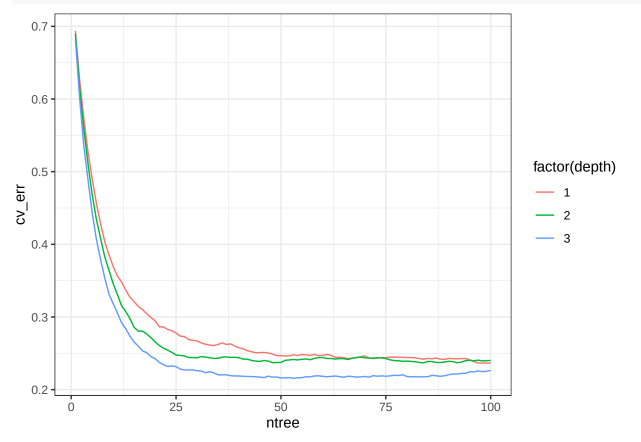
```
ntrees = 100
cv_errors = bind_rows(
  tibble(ntree = 1:ntrees, cv_err = gbm_fit_1$cv.error, depth = 1),
```

```
tibble(ntree = 1:ntrees, cv_err = gbm_fit_2$cv.error, depth = 2),
tibble(ntree = 1:ntrees, cv_err = gbm_fit_3$cv.error, depth = 3)
)
cv_errors
```

```
## # A tibble: 300 x 3
##
      ntree cv_err depth
      <int> <dbl> <dbl>
##
##
          1 0.694
   1
          2 0.627
##
    2
                       1
##
    3
          3 0.574
                       1
##
   4
          4 0.527
                       1
##
   5
          5 0.490
          6 0.457
##
   6
         7 0.428
##
   7
                       1
##
   8
         8 0.404
##
         9 0.386
   9
                       1
## 10
         10 0.370
## # ... with 290 more rows
```

We can then plot these as follows:

```
cv_errors %>%
ggplot(aes(x = ntree, y = cv_err, colour = factor(depth))) +
geom_line() + theme_bw()
```



Which value of interaction.depth seems to work the best here?

Let's save the optimal model and optimal number of trees (note plot.it = FALSE in gbm.perf to extract

the optimal number of trees without making the CV plot again):

```
gbm_fit_optimal = gbm_fit_3
optimal_num_trees = gbm.perf(gbm_fit_3, plot.it = FALSE)
optimal_num_trees
```

[1] 53

Model interpretation

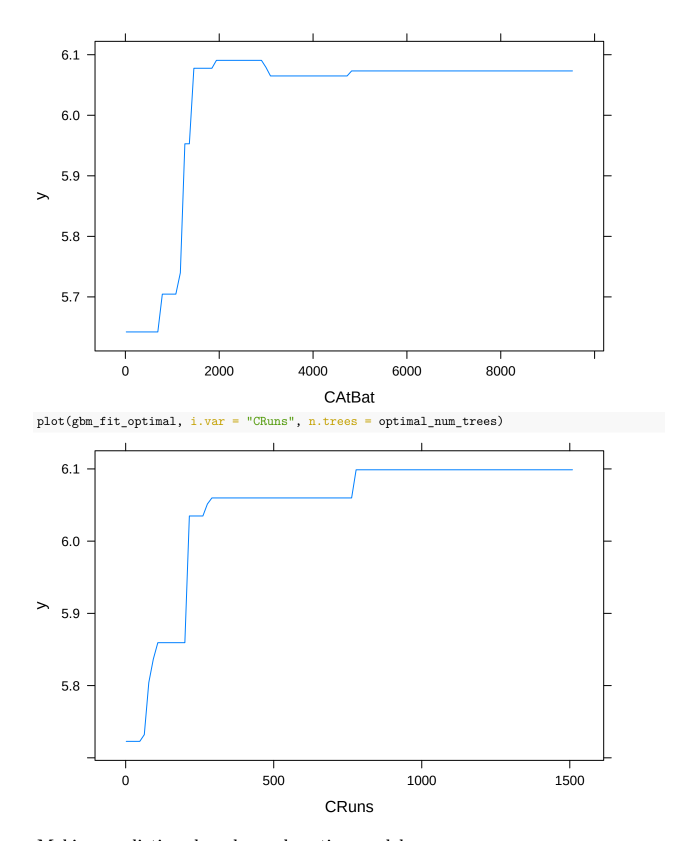
Let's now interpret our tuned model. To get the variable importance measures, we use summary, specifying the number of trees via the n.trees argument:

```
summary(gbm_fit_optimal, n.trees = optimal_num_trees, plotit = FALSE)
```

```
##
                         rel.inf
                   var
## CAtBat
               CAtBat 24.5153770
## CRBI
                 CRBI 16.6073144
                CRuns 11.5190958
## CRuns
## CWalks
               CWalks 9.7032929
## CHits
                CHits 6.6707587
## AtBat
                AtBat 4.4347641
## PutOuts
              PutOuts 4.3226905
## Hits
                 Hits 4.2193568
## Years
                Years 3.1751279
## Walks
                Walks 3.0915187
## CHmRun
                CHmRun 3.0041789
## RBI
                  RBI 2.9560862
## Runs
                 Runs 2.4589581
## HmRun
                HmRun 0.9662435
## Errors
               Errors
                       0.8391951
## League
               League
                       0.6814164
## Division
                       0.4754207
             Division
## Assists
               Assists
                       0.3592044
## NewLeague NewLeague 0.0000000
```

We can also make the partial dependence plots for the different features using plot:

```
plot(gbm_fit_optimal, i.var = "CAtBat", n.trees = optimal_num_trees)
```



Making predictions based on a boosting model:

We can make predictions using predict, as usual, but we need to specify the number of trees to use:

[1] 0.2845969

Boosting for classification

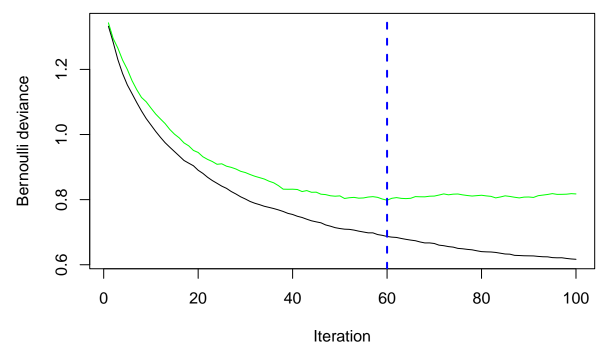
Boosting models work very similarly for classification. Let's continue with the heart disease data from last time:

Fitting a boosting model uses the same basic syntax, but with distribution = "bernoulli":

Common pitfalls when fitting a gbm:

- The binary response is coded as a character, e.g. "Yes"/"No".
- The binary response is coded as a factor.
- Any of the features are coded as strings, rather than factors.

```
gbm.perf(gbm_fit)
```



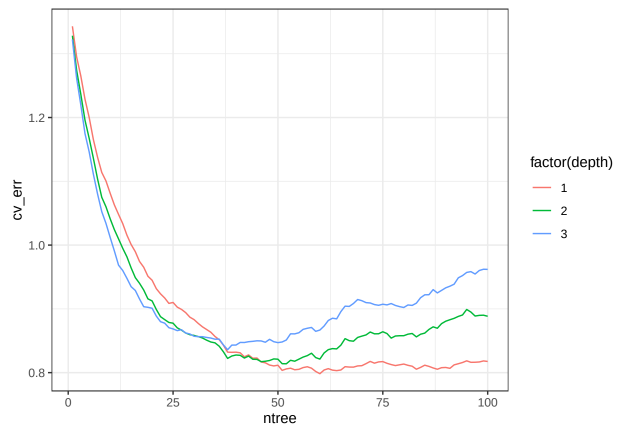
[1] 60

We can tune the interaction depth in the same way as before:

```
# try a few values
set.seed(1)
gbm_fit_1 = gbm(AHD \sim .,
              distribution = "bernoulli",
              n.trees = 100,
              interaction.depth = 1,
              shrinkage = 0.1,
              cv.folds = 5,
              data = Heart_train)
set.seed(1)
gbm_fit_2 = gbm(AHD \sim .,
              distribution = "bernoulli",
              n.trees = 100,
              interaction.depth = 2,
              shrinkage = 0.1,
              cv.folds = 5,
              data = Heart_train)
set.seed(1)
gbm_fit_3 = gbm(AHD \sim .,
              distribution = "bernoulli",
              n.trees = 100,
              interaction.depth = 3,
              shrinkage = 0.1,
              cv.folds = 5,
              data = Heart_train)
# extract CV errors
ntrees = 100
cv_errors = bind_rows(
```

```
tibble(ntree = 1:ntrees, cv_err = gbm_fit_1$cv.error, depth = 1),
tibble(ntree = 1:ntrees, cv_err = gbm_fit_2$cv.error, depth = 2),
tibble(ntree = 1:ntrees, cv_err = gbm_fit_3$cv.error, depth = 3)
)

# plot CV errors
cv_errors %>%
ggplot(aes(x = ntree, y = cv_err, colour = factor(depth))) +
geom_line() + theme_bw()
```



Aha! We see some overfitting! For which values of interaction depth do we see more overfitting, and why? What is the optimal interaction depth?

```
gbm_fit_optimal = gbm_fit_1
optimal_num_trees = gbm.perf(gbm_fit_1, plot.it = FALSE)
```

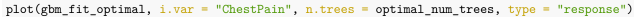
We can calculate variable importance scores as before:

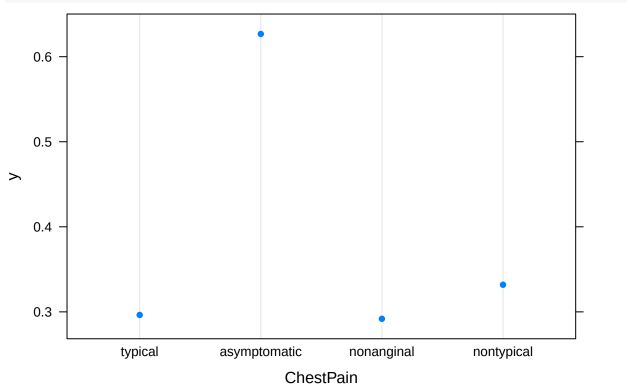
```
summary(gbm_fit_optimal, n.trees = optimal_num_trees, plotit = FALSE)
```

```
##
                   var
                         rel.inf
## ChestPain ChestPain 26.171634
## Thal
                  Thal 21.926979
## Ca
                    Ca 21.021617
## Oldpeak
               Oldpeak 7.777384
## MaxHR
                 MaxHR 6.794431
## Chol
                  Chol 3.920631
## Slope
                 Slope 3.283046
```

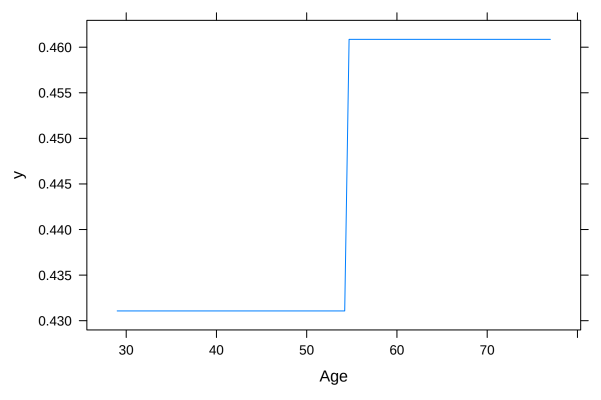
```
## RestBP RestBP 3.008152
## Sex Sex 2.769791
## ExAng ExAng 1.857264
## Age Age 1.469070
## Fbs Fbs 0.000000
## RestECG RestECG 0.000000
```

For the partial dependence plots, it's useful to specify type = "response" so we can interpret the y axis on the probability scale:





plot(gbm_fit_optimal, i.var = "Age", n.trees = optimal_num_trees, type = "response")



To make predictions, use the same syntax as before but with type = "response" to get predictions on the probability scale:

```
gbm_probabilities = predict(gbm_fit_optimal, n.trees = optimal_num_trees, type = "response", newdata = Heart_test)

gbm_probabilities

## [1] 0.95737346 0.05912815 0.33020657 0.76891179 0.47295576 0.27863385

## [7] 0.19513646 0.26055091 0.88884444 0.72941877 0.08203111 0.54234917

## [13] 0.05981156 0.92066149 0.39445784 0.36753679 0.94231197 0.97648564

## [19] 0.18233581 0.23391324 0.92945600 0.14398180 0.38773436 0.41093601

## [25] 0.05416493 0.91803106 0.23112832 0.96456990 0.85078912 0.71560082

## [31] 0.95463630 0.95027231 0.27237411 0.73277821 0.03889593 0.35264965

## [37] 0.87589252 0.86364608 0.92799472 0.33914436 0.08598292 0.92220713

## [43] 0.41009369 0.08898101 0.96951434 0.03503601 0.12429338 0.26230606

## [49] 0.93001342 0.08791087 0.69489085 0.83768148 0.04893160 0.06336597

## [55] 0.05542077 0.90798838 0.28925915 0.20442883 0.31517227 0.65976992
```

We can then threshold the probabilities at 0.5 as usual and calculate the misclassification error:

```
gbm_predictions = as.numeric(gbm_probabilities > 0.5)
mean(gbm_predictions != Heart_test$AHD)
```

[1] 0.1