Super Learner for Prediction (Part 1)

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Introduction

In this practical, you will apply the SuperLearner. We will use part of the National Health and Nutrition Examination Survey (NHANES) dataset. We want to write the raw dataset to a data frame in our environment, and can do this with the following code:

```
nhanes_installed <- require(NHANES)
if(!nhanes_installed) install.packages("NHANES", repos = "http://cran.us.r-project.org")
library(NHANES)

data("NHANES")
df <- NHANESraw</pre>
```

Data Processing

Inspect the data

dim(df)

[1] 20293 78

names(df)

```
"SurveyYr"
    [1] "ID"
                                                 "Gender"
##
    [4] "Age"
                            "AgeMonths"
                                                 "Race1"
    [7]
        "Race3"
                            "Education"
                                                 "MaritalStatus"
## [10] "HHIncome"
                            "HHIncomeMid"
                                                "Poverty"
## [13] "HomeRooms"
                            "HomeOwn"
                                                "Work"
## [16] "Weight"
                            "Length"
                                                "HeadCirc"
                                                "BMICatUnder20yrs"
## [19] "Height"
                            "BMI"
## [22] "BMI WHO"
                            "Pulse"
                                                "BPSysAve"
        "BPDiaAve"
                            "BPSys1"
                                                "BPDia1"
## [25]
                            "BPDia2"
                                                 "BPSys3"
## [28] "BPSys2"
## [31] "BPDia3"
                            "Testosterone"
                                                "DirectChol"
                            "UrineVol1"
## [34] "TotChol"
                                                "UrineFlow1"
  [37] "UrineVol2"
                            "UrineFlow2"
                                                 "Diabetes"
   [40] "DiabetesAge"
                            "HealthGen"
                                                 "DaysPhysHlthBad"
  [43]
       "DaysMentHlthBad"
                            "LittleInterest"
                                                "Depressed"
                                                 "Age1stBaby"
## [46] "nPregnancies"
                            "nBabies"
## [49] "SleepHrsNight"
                            "SleepTrouble"
                                                 "PhysActive"
```

```
## [52] "PhysActiveDays"
                            "TVHrsDay"
                                                "CompHrsDay"
  [55] "TVHrsDayChild"
##
                            "CompHrsDayChild"
                                               "Alcohol12PlusYr"
## [58] "AlcoholDay"
                            "AlcoholYear"
                                                "SmokeNow"
## [61] "Smoke100"
                            "SmokeAge"
                                                "Marijuana"
##
  [64] "AgeFirstMarij"
                            "RegularMarij"
                                                "AgeRegMarij"
                            "SexEver"
  [67] "HardDrugs"
                                                "SexAge"
                            "SexNumPartYear"
                                                "SameSex"
## [70] "SexNumPartnLife"
## [73] "SexOrientation"
                            "WTINT2YR"
                                                "WTMEC2YR"
## [76] "SDMVPSU"
                            "SDMVSTRA"
                                                "PregnantNow"
table(df$SmokeNow, df$Smoke100, useNA='always')
```

We will be interested in smoking as an exposure. The dataset contains two smoking variables Smoke100 which is a binary indicator of whether a person has smoked at least 100 cigarettes in their lifetime (but is not a current smoker), and SmokeNow which indicates if the person is a current smoker.

Combine these into a single factor variable that indicates whether the individual is an ex, current or never smoker.

A systolic blood pressure reading (BPSysAve), a continuous outcome, will be a primary outcome. If there is time, you can also consider diabetes status (Diabetes), a binary outcome.

Now we trim the variables in the data set, keeping only the ones relevant to the practical:

```
df <- df %>% dplyr::select(one_of(
   "BPSysAve","BMI", "Age", "SleepHrsNight", "PhysActive","Smoke",
   "Gender", "Race1", "Poverty", "Diabetes", "TotChol"))
```

Inspect our outcome variables, and do the same for key covariates BMI and Age.

```
table(df$Diabetes,useNA = 'always')
##
##
                <NA>
      Nο
           Yes
         1706
## 17754
                  833
summary(df$BPSysAve)
##
      Min. 1st Qu. Median
                                Mean 3rd Qu.
                                                 Max.
                                                         NA's
##
             105.0
                      115.0
                                       127.0
                                                233.0
                                                         5426
      74.0
                               118.1
table(df$Smoke, useNA = 'always')
##
## Current
                 Ex
                      Never
                                <NA>
      2454
               2779
                      15060
                                   0
summary(df$BMI)
```

```
## 12.40 19.79 24.92 25.65 30.10 84.87 2279

summary(df$Age)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
```

For this illustration, we want to keep only the complete cases in the dataset. We do this using a single line of code.

80.00

53.00

```
df <- df[complete.cases(df),]</pre>
```

Now split the data into training and test sets. The split should be 50/50. Call the sets df_train and df holdout.

```
set.seed(101)
train_obs <- sample(nrow(df), size = nrow(df)*0.5)

df_train <- df[train_obs, ]

# Create a holdout set for evaluating model performance.
# Note: cross-validation is even better than a single holdout sample.
df_holdout <- df[-train_obs, ]</pre>
```

We want the outcome (BPSysAve) in vector form (also split into train and holdout) and kept separate from the main data frame. Do this now.

```
# The continuous outcome will be the average of the 3 systolic blood pressure measurrement
# BPSysAve
y_train <- df$BPSysAve[train_obs]
y_holdout <- df$BPSysAve[-train_obs]</pre>
```

Fitting Individual Algorithms

Linear Regression

##

0.00

10.00

28.00

32.02

We begin by trying different learners individually. For example: a linear regression, lasso (glmnet), random-Forest, XGBoost. These should ideally be tested with multiple hyperparameter settings for each algorithm. (You will get to do this later!)

Carry out a simple linear regression for the systolic blood pressure measurement, using BMI, Age, SleepHrsNight, PhysActive, Smoke, Gender, Race1, Poverty, Diabetes and TotChol as covariates.

Using the generic predict() function to make predictions, manually calculate the sum of square errors for the test set.

Boosting

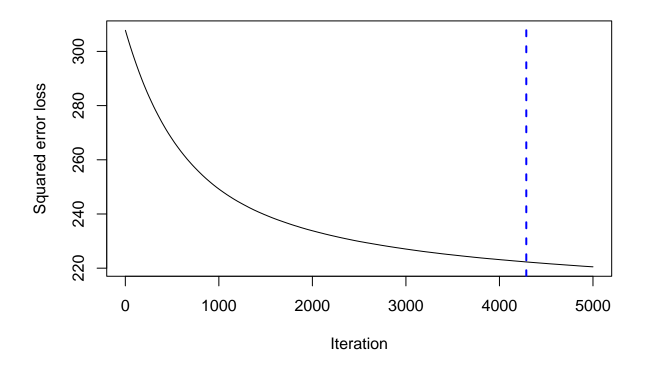
Now we run a gradient boosting algorithm on the data. We specify the formula with only the main terms, but recall that trees automatically include interactions up to the specified depth.

Use the gbm() function to fit a gradient boosting model to the data.

We can use the gbm.perf() function to check the performance of this model, and by specifying the option method, we can do this in multiple ways. Check the performance using out-of-bag validation (method = "OOB") and cross-validation (method = "CV").

How do you interpret the plot that results from running gbm.perf()?

```
#' Checks performance using the out-of-bag (00B) error
best.iter <- gbm.perf(boost, method = "00B")</pre>
```



```
print(best.iter)

## [1] 4287

## attr(,"smoother")

## Call:

## loess(formula = object$oobag.improve ~ x, enp.target = min(max(4,

## length(x)/10), 50))

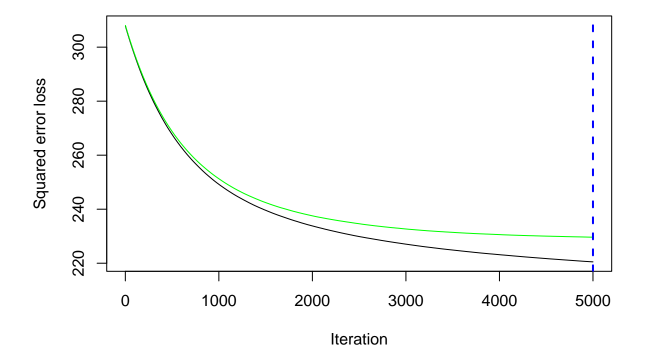
##

## Number of Observations: 5000

## Equivalent Number of Parameters: 39.99

## Residual Standard Error: 0.001578

#' Checks performance using 5-fold cross-validation
best.iter2 <- gbm.perf(boost, method = "cv")</pre>
```



```
print(best.iter2)
```

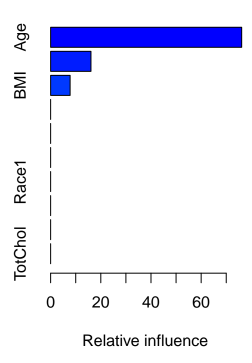
[1] 5000

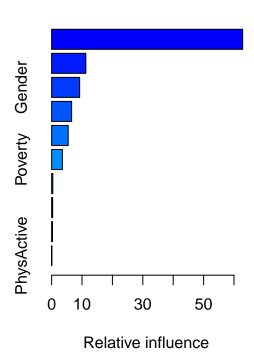
The OOB error typically underestimates the optimal number of iterations.

Using the summary() function with the n.trees options, generate plots that show the relative influence of each variable after (a) a single tree and (b) with the optimal number of iterations as determined in the last question.

```
#' Plots relative influence of each variable
par(mfrow = c(1, 2))
summary(boost, n.trees = 1)  # using first tree
```

```
##
                                 rel.inf
                           var
                           Age 76.152577
## Age
## Gender
                        Gender 16.076925
## BMI
                           BMI
                                7.770499
## SleepHrsNight SleepHrsNight
                                0.000000
## PhysActive
                    PhysActive
                               0.000000
## Smoke
                         Smoke
                               0.000000
## Race1
                               0.000000
                         Race1
## Poverty
                       Poverty
                                0.000000
## Diabetes
                      Diabetes
                                0.000000
## TotChol
                       TotChol
                                0.000000
summary(boost, n.trees = best.iter) # using estimated best number of trees
```





##		var	rel.inf
##	Age	Age	62.85775470
##	BMI	BMI	11.25197645
##	Gender	Gender	9.22223588
##	TotChol	TotChol	6.57923313
##	Race1	Race1	5.46377909
##	Poverty	Poverty	3.55175178
##	Diabetes	Diabetes	0.38909641
##	${\tt SleepHrsNight}$	${\tt SleepHrsNight}$	0.33934926
##	Smoke	Smoke	0.28974810
##	PhysActive	PhysActive	0.05507517

We see that in both (first and best tree) Age has the largest influence. Gender and BMI have second and third

largest influence, using the first tree, while the best tree has BMI as higher than Gender.

Now using the generic predict() function, make predictions for the holdout set. Don't forget to stipulate how many iterations of the gradient boosting model should be used. This can be done using the n.trees option.

Also generate the sum of square errors that results from applying this model to the holdout set.

```
#' predictions will be on the link scale
Yhat.boost <- predict(boost, newdata = df_holdout, n.trees = best.iter, type = "link")

#' Sum of Squared Errors SSE
SSE.boost <- sum((y_holdout - Yhat.boost)^2)</pre>
```

LASSO via SuperLearner

We will continue to fit single learners, but in order to get familiar it we do so using the SuperLearner package

First, check which learners have been integrated into the SuperLearner package. We can use any of these when we run the SuperLearner:

```
listWrappers(what = "SL")
```

```
##
    [1] "SL.bartMachine"
                               "SL.bayesglm"
                                                      "SL.biglasso"
##
   [4] "SL.caret"
                               "SL.caret.rpart"
                                                      "SL.cforest"
## [7] "SL.earth"
                               "SL.extraTrees"
                                                      "SL.gam"
## [10] "SL.gbm"
                               "SL.glm"
                                                      "SL.glm.interaction"
## [13] "SL.glmnet"
                               "SL.ipredbagg"
                                                      "SL.kernelKnn"
## [16] "SL.knn"
                               "SL.ksvm"
                                                      "SL.lda"
## [19] "SL.leekasso"
                               "SL.lm"
                                                      "SL.loess"
## [22] "SL.logreg"
                               "SL.mean"
                                                      "SL.nnet"
## [25] "SL.nnls"
                               "SL.polymars"
                                                      "SL.qda"
## [28] "SL.randomForest"
                               "SL.ranger"
                                                      "SL.ridge"
                               "SL.rpartPrune"
## [31] "SL.rpart"
                                                      "SL.speedglm"
                               "SL.step"
## [34] "SL.speedlm"
                                                      "SL.step.forward"
## [37] "SL.step.interaction" "SL.stepAIC"
                                                      "SL.svm"
## [40] "SL.template"
                               "SL.xgboost"
```

SuperLearner (SL) likes the outcome and matrix of covariates to be kept separate. Do this now.

```
X <- df %>% dplyr::select(-one_of("BPSysAve") )

# Also divide our design matrix into training and testing sets
x_train <- X[train_obs, ]
x_holdout <- X[-train_obs, ]</pre>
```

Now let's fit penalised regression LASSO, but using SL. For now using all the defaults which we will explain later:

```
# Review the elements in the SuperLearner object.
names(sl_lasso)
```

```
[1] "call"
                             "libraryNames"
                                                  "SL.library"
##
                             "coef"
   [4] "SL.predict"
                                                  "library.predict"
  [7] "Z"
                             "cvRisk"
                                                  "family"
## [10] "fitLibrary"
                             "cvFitLibrary"
                                                  "varNames"
## [13] "validRows"
                             "method"
                                                  "whichScreen"
## [16] "control"
                             "cvControl"
                                                  "errorsInCVLibrary"
## [19] "errorsInLibrary"
                             "metaOptimizer"
                                                  "env"
## [22] "times"
```

Again using the generic predict() function, calculate the sum of square errors on the test set.

```
# Predict outcome in the holdout using lasso
Yhat.lasso <- predict(sl_lasso, x_holdout, onlySL = TRUE)$pred

# calculate sum of square errors
SSE.lasso <- sum((y_holdout- Yhat.lasso)^2)</pre>
```

The SSE corresponding to LASSO is larger than the one obtained from boosting.

Random Forest via SuperLearner

Fit a random forest model using the wrapper SL.ranger. All other options as before. Also calculate the sum of square errors on the test set.

The SL.ranger wrapper fits a random forest with 500 trees, minimum node size 5, and the number of variables used to split the squared root of the number of available predictors.

With these defaults, the SSE is a bit larger than the one we obtained with boosting.

Multiple Learners Stacked in SuperLearner

Instead of fitting the models separately and looking at the performance using sum of square errors as we have been doing, we now fit them simultaneously by including them all in the SuperLearner library.

For now, we include those algorithms we tried up to now:

```
# Select candidate algorithms
my.library <-c("SL.glm","SL.glmnet", "SL.ranger")</pre>
# Set seed
set.seed(101)
# Execute the call to SuperLearner
sl0 <- SuperLearner(Y = y_train, # Y is the outcome variable
                    X = x_train, # X is a dataframe of predictor variables, in this case
                                  # everything except for outcome
                    family = gaussian(), # family will be discussed in more detail when
                                           # we see how wrappers are written.
                                           # for now gaussian (outcome is coninuous)
                                           # binomial() for 0/1 outcome
                    method = "method.NNLS",
                    # method specifies how the ensembling is done (i.e. how the optimal
                    # combination is chose)
                    # for now we will use the \sum_{k=1}^{K} \alpha_k f_{k,n} method by deafault
                    SL.library = my.library,
                    cvControl = list(shuffle = F, V = 5)
                    # cvControl specifies parameters related to cross validation,
                    # used to estimate the risk on future data
                    # the default is for V = 10-fold
```

Now check the output by simply calling the SuperLearner object.

```
##
##
## Call:
## SuperLearner(Y = y_train, X = x_train, family = gaussian(), SL.library = my.library,
## method = "method.NNLS", cvControl = list(shuffle = F, V = 5))
##
##
##
##
Risk Coef
## SL.glm_All 236.4878 0.4204597
## SL.glmnet_All 236.4112 0.0000000
## SL.ranger_All 233.7565 0.5795403
```

The output has two main components:

Firstly, the risk is a measure of model accuracy or performance, and we want our models to minimize the estimated risk (according to the specified loss function). Because we did not change it by an option, our SL has used the default mean square error, but this can be altered (see the help files for SuperLearner). In this case, the risks for each algorithm in the library are all broadly similar, with the random forest slightly outperforming the other two algorithms in the library.

The coef column tells us the importance of each algorithm in the final ensemble. By default (because we use NNLS) the weights are always greater than or equal to 0 and sum to 1 (a 'convex combination'). If a coefficient is 0, it means that the algorithm is not being used in the SuperLearner ensemble. Here we see that glm has been given no weight in the final (ensemble) predictor and so is not used.

Now see which has the lowest risk, and how long it took to run, using the following code:

```
# Let's see which is the discrete SL (i.e min risk)
sl0$cvRisk[which.min(sl0$cvRisk)]
```

```
## SL.ranger_All
## 233.7565

# Review how long it took to run the SuperLearner:
sl0$times$everything
## user system elapsed
```

Now that we have a SL ensemble predictor, make predictions on the holdout data set and review the results. This can be done using the generic predict() function, but we stipulate the option onlySL = TRUE so we do not fit algorithms in the library that had zero weight, saving computation.

```
pred.sl0 <- predict(sl0, x_holdout, onlySL = TRUE)</pre>
```

Check the structure of this prediction object using str(pred.s10). You will see that the prediction object is a list with two objects. The first is a vector of SL predictions. The second is a matrix of predictions; these are the predictions for each of the individual learners in the SL library.

We want the first object in the list to serve as our predictions. Use this to calculate the sum of square errors resulting from applying the SL predictor to the test set.

```
# Pick out SL predictions
Yhat.sl0 <- pred.sl0[[1]]

# Calculate the sum of square errors
SSE.sl0 <- sum((y_holdout - Yhat.sl0)^2)</pre>
```

The SSE is the lower than those SSEs corresponding to LASSO and RF (which were included in the SL library).

Extend the Ensemble

##

11.284

0.481 10.929

We will now add other base learners to the library You can check which ones are implemented by running listWrappers(). (You can also write your own wrapper functions, we will see later how to do this).

For now, add gbm() and glm.interaction. [Note: xgboost has deprecated a function used in the SL wrapper so warnings "reg:linear is now deprecated in favor of reg:squarederror" will be displayed, so I won't use for this exercise, though ordinarily I do!]

Run another SL predictor using the same three algorithms as before alongside the two new ones. Review the output and the time it took to run as before.

```
# Review the SL object
sl1
##
## Call:
## SuperLearner(Y = y_train, X = x_train, family = gaussian(), SL.library = my.library.2,
##
       cvControl = list(shuffle = F, V = 5))
##
##
##
                              Risk
                                         Coef
## SL.glm_All
                          236.4878 0.0000000
## SL.glm.interaction_All 234.0326 0.2662155
## SL.glmnet_All
                          236.4585 0.0000000
## SL.ranger_All
                          233.6424 0.1555134
## SL.gbm_All
                          229.3069 0.5782711
# Review times
sl1$times$everything
```

user system elapsed ## 81.447 2.045 187.981

You should see that the gradient boosted predictor (gbm()) has the most weight. The new glm with interactions is the second most important, and the others are either zero or make only small contributions to the ensemble.

As before, use the SL predictor to make predictions on the holdout set.

```
Yhat.sl1 <- predict(sl1, x_holdout, onlySL = TRUE)[[1]]
# SSE
SSE.sl1 <- sum((y_holdout - Yhat.sl1)^2)</pre>
```

We see that adding more learners slightly improves the SSE. Adding more (without removing) will result in further improvements.