# Big Data: MLearning Classifiers Grid Search with Sparklyr and H2O-SWater

## WHAT IS THIS ALL ABOUT?

H2O is a product created by the H2O.ai company to combine the main algorithms of machine learning and statistical learning with **Big Data**. Thanks to its way of compressing and storing data, H2O can work with millions of records in a single computer (it uses all its cores) or in a cluster of many computers. Internally, H2O is written in Java and follows the Key / Value paradigm for storing data and Map / Reduce for its algorithms. Thanks to its APIs, it is possible to access all its functions from R, Python, or Scala, as well as through a web interface called Flow.

Although H2O's main advantage over other tools is its scalability, its algorithms are equally useful when working with a small volume of data. H2O is an open source, distributed machine learning platform for Everyone. H2O is an alternative open-source cross-platform for machine learning that supports the most widely used statistical & machine learning algorithms including gradient boosted machines, generalized linear models, deep learning and more.

The rsparkling extension package provides bindings to H2O's distributed machine learning algorithms via sparklyr. In particular, rsparkling allows you to access the machine learning routines provided by the H2O Sparkling Water Spark package. The main purpose of rsparkling is to provide a connector between sparklyr and H2O's machine learning algorithms. The rsparkling package uses sparklyr for Spark job deployment and initialization of Sparkling Water. After that, user can use the regular h2o R package for modeling.

This present work is about an application and practical exposition of how to apply two classic Grid Search methods for tunning four (Glm, RandomForest, Gbm, Deeplearning) of the most frequently used machine learning models for classification, by using Sparklyr, rsparkling and H2O-Sparkling Water. The main results obtained are presented here, providing a clear guideline that identifies the classic steps in the modeling process, possible to adapt to any other equivalent database. The key functions and scripts used are proporcionated, allowing the reader to consult the appropriate bibliography and tutorial examples clearly and directly.

You can get more specific information about the H2O machine learning models for classification used in this project in the following links:

Generalized Linear Model (Glm) H2O documentation R documentation Random Forest (RandomForest) H2O documentation R documentation

Gradient Boosting Machine (Gbm) H2O documentation R documentation
Deep Learning - Neural Networks (Deeplearning)
H2O documentation
R documentation

An excellent clear practical explanation of the Grid Search methods used in this project can be found here. Even though in this video Kevin (the author) use examples based on Python, his concepts explanations are so clearly exposed that I decided to include it here without a doubt.

While in this project models are run on a small data set in a local spark cluster, these methods can be scaled, for the most part, for data analysis in a distributed Apache Spark cluster.

This project has been developed using sparklyr considering a local Spark cluster environment. The excellent online publications from the University of Chicago "MACS 305001 - Computing for the Social Sciences" in its derived chapter "Spark and sparklyr" and the great "Documentation About Sparkling Water" are both used as bibliographic source of support and consultation base.

## DATA SOURCE

In this project I use a prostate cancer dataset which is included in the H2O package. The data was collected by Dr. Donn Young at the Ohio State University Comprehensive Cancer Center for a study of patients with varying degrees of prostate cancer. You can download the dataset here. The classification goal is to predict the incidence of penetration of the prostatic capsule (CAPSULE) (variable y)

Note that, the dataset is not significant [380 rows x 8 columns] and you may think that the computation takes a long time. Spark is designed to process a considerable amount of data. Spark's performances increase relative to other machine learning libraries when the dataset processed grows larger.

## LOADING THE DATA

```
# Clean memory and remove all fies
rm(list=ls())
# set working directory location for our project
mypath <- "path"</pre>
# set working directory
setwd(mypath)
# get current working directory
getwd()
# Load required minimum packages
library(rsparkling)
library(sparklyr)
library(tidyverse) # to get the whole tidyverse: dplyr, ggplot2 tibble, readr, ti
dyr, purrr
library(reshape2)
library(gridExtra) #viewing multiple plots together
library(grid) # To use with tables
library(h2o)
h2o.init(ip = "localhost",
# Number of threads -1 means use all cores on your machine.
nthreads = -1)
# max mem size = "12G")  #max mem size is the maximum memory to allocate to H2O
#Cluster data is deleted in case it has already been started.
h2o.removeAll()
# Spark connection
sc <- spark_connect(master = "local")</pre>
```

```
# Read data from web
path <- "http://h2o-public-test-data.s3.amazonaws.com/smalldata/prostate/prostate</pre>
df raw <- read.csv(path)</pre>
glimpse(df raw)
Observations: 380
Variables: 9
          1, 2, 3, 4, 5, 6, 7, 8, 9, 10, 11, 12, 13, 14, 15, 16, 17, ...
$ CAPSULE 0, 0, 0, 0, 0, 1, 0, 0, 0, 1, 1, 1, 1, 1, 0, 0, 0, 0, 1,...
$ AGE
          65, 72, 70, 76, 69, 71, 68, 61, 69, 68, 68, 72, 72, 65, 75,...
          1, 1, 1, 2, 1, 1, 2, 2, 1, 2, 2, 1, 1, 1, 1, 1, 1, 2, 1, 1, 2,...
$ RACE
$ DPROS 2, 3, 1, 2, 1, 3, 4, 4, 1, 1, 4, 2, 4, 4, 1, 2, 1, 2, 1, 3,...
$ DCAPS 1, 2, 2, 1, 1, 2, 2, 2, 1, 2, 2, 2, 2, 2, 1, 1, 1, 1, 1, 2, 2,...
          1.4, 6.7, 4.9, 51.2, 12.3, 3.3, 31.9, 66.7, 3.9, 13.0, 4.0,...
$ PSA
          0.0, 0.0, 0.0, 20.0, 55.9, 0.0, 0.0, 27.2, 24.0, 0.0, 0.0, ...
$ VOL
$ GLEASON 6, 7, 6, 7, 6, 8, 7, 7, 7, 6, 7, 7, 9, 7, 5, 5, 5, 5, 6, 7,...
# copy our R data frame onto spark
df <- copy to(sc, df raw, name = "df")</pre>
# to see tables in "sc"
src tbls(sc)
Output:
[1] "df"
```

## READJUSTMENT DATASET

#### GETTING DATASET INTO H2O FORMAT

```
# Getting mydata tbl into H2O format.
h2o df <- as.h2o(df, destination frame = "h2o df")
h2o df
Output:
 ID CAPSULE AGE RACE DPROS DCAPS PSA VOL GLEASON
        0 65 1
                    2
                        1 1.4 0.0
        0 72 1
                         2 6.7 0.0
                   1 2 4.9 0.0
        0 70 1
       0 76 2 2 1 51.2 20.0
                                      7
       0 69 1 1 1 12.3 55.9
        1 71 1
                   3 2 3.3 0.0
[380 rows x 9 columns]
```

#### TRANSFORM CATEGORICAL FEATURES INTO FACTORS

Now we have readjusted mydata\_tbl dataset to be incorporated into the H2o format for machine learning models. We isolate the model's dependent variable (y) and turn all the categorical features into numbers that represent factors.

When fitting a model, H2O automatically identifies which variables are categorical and internally creates the corresponding dummy variables. It is highly recommended to allow H2O to carry out this process instead of doing it externally since its implementation is highly optimized. The behavior of the encoding of categorical variables can be controlled with the argument categorical\_encoding, by default, its value is "AUTO".

By default, H2O standardizes numerical predictors before adjusting the models so that they all have mean zero and variance one. This behavior can be controlled with the "standardize" argument

```
y <- "CAPSULE"

x <- setdiff(names(h2o_df), y)

h2o_df[,y] <- as.factor(h2o_df[,y])

h2o_df[,"RACE" ] <- as.factor(h2o_df[,"RACE" ])

h2o_df[,"DCAPS" ] <- as.factor(h2o_df[,"DCAPS" ])</pre>
```

```
h2o_df[,"DPROS" ] <- as.factor(h2o_df[,"DPROS" ])
```

Although the data set used in this project is small enough to load everything in memory and directly use the magnificent possibilities of exploratory analysis that R offers, to better represent the way to proceed with large volumes of data, now, by way of example, we use some of H2O's own functions.

#### EXPLORING THE DATA

Although the data set used in this example is small enough to load everything into memory and directly use the magnificent possibilities of exploratory analysis that R offers, to better represent the way to proceed with large volumes of data, H2O's functions are used.

```
# Data set dimensions
h2o.dim(h2o_df)
Output:
[1] 380  9
# Columns name
h2o.colnames(h2o_df)
Output:
[1] "ID"          "CAPSULE" "AGE"          "PSA"
[8] "VOL"          "GLEASON"
```

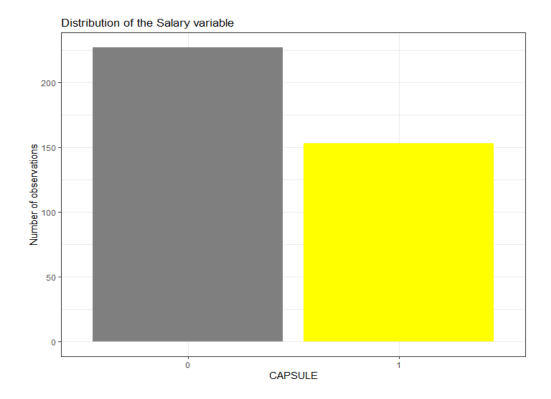
We can get a quick overview of the data and its structure. The "h2o.describe ()" function is very useful to obtain a quick analysis that shows the data type, the number of missing values, the minimum, maximum, mean value, standard deviation, and the number of categories (Cardinality) of each one of variables. H2O uses the name "enum" for data of type factor or character.

```
h2o.describe(h2o df)
Output:
   Label Type Missing Zeros PosInf NegInf Min Max
                                                        Sigma Car
                                               Mean
                               0 1.0 380.0 190.5000000 109.8407939
    ID int
                         0
                  227
                                0 0.0 1.0 0.4026316 0.4910743
2 CAPSULE enum
                          0
                                0 43.0 79.0 66.0394737 6.5270713
3
                0
                    0
                          0
    AGE int
NA
4
                0
                    3
                          0
                                0 0.0 2.0
                                                 NA
  RACE enum
                                                           NA
3
                          0 0.0 3.0
5
  DPROS enum 0 99
                                                 NA
                                                           NA
```

```
DCAPS enum
                 0 339
                            0
                                  0 0.0 1.0 0.1078947 0.3106564
2
    PSA real
                     0
                                 0 0.3 139.7 15.4086316 19.9975727
NA
                                0 0.0 97.6 15.8129211 18.3476200
8
    VOL real
                 0 167
                            0
NA
9 GLEASON int
                 0
                     2
                            0
                                 0 0.0 9.0 6.3842105 1.0919534
NA
```

To count the number of observations of each class in a categorical variable, as in this case the response variable CAPSULE, the h2o.table () function is used.

```
# A table is created with the number of observations of each type.
tabla CAPSULE <- as.data.frame(h2o.table(h2o df$CAPSULE))</pre>
tabla CAPSULE
Output:
 CAPSULE Count
      0 227
       1 153
# Once the table is created, it can be loaded into the R environment to be able t
o graph.
ggplot(data = tabla_CAPSULE,
        aes(x = as.factor(CAPSULE), y = Count, fill = as.factor(CAPSULE))) +
        geom col() +
        scale fill manual(values = c("gray50", "yellow")) +
        theme bw() +
        labs(x = "CAPSULE", y = "Number of observations", title = "Distribution o
f the Salary variable") +
        theme(legend.position = "none")
```



## GETTING THE TRAIN AND TEST DATASET

#### GETTING TRAIN AND TEST DATASETS

The h2o.splitFrame () function performs random partitions, but it does not allow them to do them in a stratified way, so it does not ensure that the distribution of response variable classes is the same in all partitions. This can be problematic with very unbalanced data (some of the groups are very minority).

In the case of unbalanced data, the base should be adjusted in advance according to traditional methods that allow more effective modeling.

```
# split our data set into 'training', 'test'
partitions <- h2o.splitFrame(h2o_df, 0.7, seed = 2020)
# Create table references
train <-partitions[[1]]
test <-partitions[[2]]</pre>
```

## GRID SEARCH GENERALITIES

#### INTRODUCTION

Grid-searching is the process of scanning the data to configure optimal parameters for a given model. Grid-searching can be applied across machine learning to calculate the best parameters to use for any given model. Grid-Search will build a model on each parameter combination possible. It iterates through every parameter combination and stores a model for each combination.

Even though grid-search is used to find the optimal hyperparameters of a model which results in the most 'accurate' predictions, It is important to note that it can be extremely computationally expensive and may take your machine quite a long time to run.

There are many hyperparameter optimization/tuning algorithms, but the two most common strategies are: cartesian grid search (or grid search) and random gid search.

## CARTESIAN AND RANDOM GRID SEARCHING OF HYPERPARAMETERS

The cartesian grid search (or grid search, by short) methodically build and evaluate a model for each combination of algorithm parameters specified in a grid.

For instance, if you have three hyperparameters H1, H2, and H3, and each of them can take on 10, 20, and 10 values, respectively, your grid will contain a total of 10 \* 20 \* 10 = 2000 models.

Cartesian Grid Search, the default grid search method in H2o, exhaustively searches over all possible combinations of the hyperparameters. *If the search space is small, this should be your method of choice*.

Random search differs from a cartesian search in that you no longer provide a discrete set of values to explore for each hyperparameter; rather, you provide a statistical distribution for each hyperparameter from which values may be randomly sampled. That is to say, Random Search replaces the exhaustive enumeration of all combinations by selecting them randomly.

The idea of random searching of hyperparameters was proposed by James Bergstra & Yoshua Bengio. You can check the original paper here. [http://www.jmlr.org/papers/volume13/bergstra12a/bergstra12a.pdf]

In a random grid search a random combination of hyperparameters (H2O will sample uniformly from the set of all possible hyperparameter value combinations) are tested instead of exhaustively testing all possible combinations. Also, the user specifies a stopping criterion, which controls when the random grid search is completed. The user can tell the random grid search to stop by specifying a maximum number of models or the maximum number of seconds allowed for the search. The user may also specify a performance-metric-based stopping criterion, which will stop the random grid search when the performance stops improving by a specified amount. *If your search space is large, this should be your method of choice*.

## GRID SEARCH MODELING

## MODEL: LOGISTIC REGRESSION (LR\_model)

#### CARTESIAN GRID SEARCH

#### Grid Search Train / Test dataset procedure

#### **GLM** hyperparameters

```
alpha_opts = list(list(0), list(.25), list(.5), list(.75), list(1))
lambda_opts = list(list(1), list(.5), list(.1), list(.01), list(.001), list(.0001), list(.00001), list(0))
glm_params <- list(alpha = alpha_opts, lambda = lambda_opts)</pre>
```

#### Train and validate a grid of GLMs

```
glm_grid <- h2o.grid("glm", y = "CAPSULE", x = c("AGE", "RACE", "PSA", "GLEASON")
,

grid_id = "glm_grid1",
 training_frame= train,
 family = "binomial",
 ignore_const_cols = TRUE,
 seed = 2020,
 hyper_params = glm_params)</pre>
```

```
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
  alpha lambda
                       model ids
1 [0.0] [0.01] glm grid1 model 16 0.7852855477855478
2 [0.75] [0.001] glm grid1 model 24 0.7852272727272728
3 [1.0] [0.001] glm grid1 model 25 0.78522727272728
4 [0.5] [0.001] glm_grid1_model_23 0.7851689976689976
5 [0.5] [0.01] glm_grid1_model_18 0.7848776223776223
   alpha lambda model ids auc
35 [0.25] [1.0] glm_grid1_model_2 0.5
36 [0.5] [1.0] glm_grid1_model_3 0.5
37 [0.75] [1.0] glm grid1 model 4 0.5
38 [1.0] [1.0] glm_grid1_model_5 0.5
39 [0.5] [0.5] glm grid1 model 8 0.5
40 [0.75] [0.5] glm grid1 model 9 0.5
```

## Grid Search KFold Cross Validation procedure

### Train and validate a grid of GLMs

```
glm_grid <- h2o.grid("glm", y = "CAPSULE", x = c("AGE", "RACE", "PSA", "GLEASON")

grid_id = "glm_grid2",

training_frame= h2o_df,

family = "binomial",

seed = 2020,

nfolds=10,

fold_assignment="Stratified", # can be "AUTO", "Modulo", "R

andom" or "Stratified"

hyper_params = glm_params)</pre>
```

```
Output:
H2O Grid Details
_____
Grid ID: glm grid2
Used hyper parameters:
 - alpha
 - lambda
Number of models: 40
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
  alpha lambda model ids
1 [0.0] [0.01] glm grid2 model 16 0.7859549105986007
2 [0.0] [0.1] glm grid2 model 11 0.7848895799141977
3 [0.75] [0.001] glm grid2 model 24  0.784659238144597
4 [0.5] [0.001] glm grid2 model 23 0.7845728599809968
5 [1.0] [0.001] glm_grid2_model_25 0.7845440672597966
   alpha lambda model ids
35 [0.25] [1.0] glm grid2 model 2 0.43675678788402295
36 [0.5] [1.0] glm_grid2_model_3 0.43675678788402295
37 [0.75] [1.0] glm grid2 model 4 0.43675678788402295
38 [1.0] [1.0] glm grid2 model 5 0.43675678788402295
39 [0.5] [0.5] glm_grid2_model_8 0.43675678788402295
40 [0.75] [0.5] glm grid2 model 9 0.43675678788402295
```

#### RANDOM GRID SEARCH

## Random Grid Search Train / Test dataset procedure

## **GLM** hyperparameters

```
search_criteria <- list(strategy = "RandomDiscrete",</pre>
```

```
max_models = 50)
```

#### Train and validate a grid of GLMs

```
# Get the grid results, sorted by validation auc
glm gridperf3 <- h2o.getGrid(grid id = "glm grid3",</pre>
                            sort_by = "auc",
                            decreasing = TRUE)
glm_gridperf3
# Output
H2O Grid Details
_____
Grid ID: glm grid3
Used hyper parameters:
 - alpha
 - lambda
Number of models: 40
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
                       model ids
  alpha lambda
1 [0.0] [0.01] glm_grid3_model_12 0.7852855477855478
2 [0.75] [0.001] glm_grid3_model_22 0.78522727272728
3 [1.0] [0.001] glm_grid3_model_35 0.78522727272728
4 [0.5] [0.001] glm grid3 model 1 0.7851689976689976
```

```
5 [0.5] [0.01] glm_grid3_model_28 0.7848776223776223

---
alpha lambda model_ids auc

35 [0.5] [1.0] glm_grid3_model_19 0.5

36 [1.0] [1.0] glm_grid3_model_24 0.5

37 [0.75] [1.0] glm_grid3_model_32 0.5

38 [0.75] [0.5] glm_grid3_model_34 0.5

39 [1.0] [0.5] glm_grid3_model_4 0.5

40 [0.5] [0.5] glm_grid3_model_5 0.5
```

## Grid Search KFold Cross Validation procedure

Train and validate a grid of GLMs

```
- alpha
 - lambda
Number of models: 40
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
  alpha lambda
                       model ids
1 [0.0] [0.01] glm_grid4_model_13 0.7859549105986007
2 [0.0] [0.1] glm_grid4_model_8 0.7848895799141977
3 [0.75] [0.001] glm grid4 model 36  0.784659238144597
4 [0.5] [0.001] glm grid4 model 33 0.7845728599809968
5 [1.0] [0.001] glm grid4 model 30 0.7845440672597966
   alpha lambda model ids
35 [0.5] [1.0] glm_grid4_model_17 0.43675678788402295
36 [0.75] [1.0] glm grid4 model 25 0.43675678788402295
37 [0.5] [0.5] glm_grid4_model_3 0.43675678788402295
38 [0.75] [0.5] glm grid4 model 31 0.43675678788402295
39 [0.25] [1.0] glm grid4 model 32 0.43675678788402295
40 [1.0] [0.5] glm grid4 model 37 0.43675678788402295
```

#### MODEL: RANDOM FOREST (RForest\_model)

#### CARTESIAN GRID SEARCH

## Grid Search Train / Test dataset procedure

#### RForest hyperparameters

#### Train and validate a grid of RFs

```
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 mtries ntrees sample rate
                                   model ids
                                                            auc
                      0.55 rf grid1 model 2 0.6836247086247086
1
            50
           100
                     0.55 rf grid1 model 5 0.6831876456876457
                     0.632 rf grid1 model 16 0.6830710955710956
            100
                     0.632 rf grid1 model 13 0.6821095571095571
           50
            200
                     0.55 rf_grid1_model_11 0.6809440559440559
  mtries ntrees sample rate
                                   model ids
                                                             auc
31
            100
                        0.75 rf grid1 model 30 0.6580419580419581
                        0.55 rf grid1 model 1 0.6574009324009323
32
            50
                        0.75 rf grid1 model 32 0.6561771561771562
        3
33
            150
                        0.75 rf grid1 model 36 0.6552738927738928
34
        4
            200
35
            150
                        0.75 rf grid1 model 33 0.6528846153846154
                        0.75 rf grid1 model 25 0.6500874125874126
36
             50
```

## Grid Search KFold Cross Validation procedure

#### Train and validate a grid of RFs

```
decreasing = TRUE)
rf gridperf2
Output:
H2O Grid Details
_____
Grid ID: rf grid2
Used hyper parameters:
 - mtries
 - ntrees
 - sample rate
Number of models: 36
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 mtries ntrees sample_rate model_ids
                 0.55 rf grid2 model 10 0.7319973510696496
      2 200
      2 150 0.55 rf_grid2_model_7 0.7283694681984395
                  0.55 rf grid2 model 4 0.7271025884656359
         100
          50
                  0.55 rf grid2 model 1 0.7248279634908296
               0.55 rf grid2 model 11 0.7229420402522242
 3 200
 mtries ntrees sample rate model ids
                                                       auc
                    0.75 rf grid2 model 35 0.7043563387175722
31
     3 200
      3 150
                     0.75 rf_grid2_model_32 0.7037372952117704
32
33
     4 50
                     0.75 rf grid2 model 27 0.701793786530765
                     0.75 rf_grid2_model_30 0.7011315539431632
     4 100
34
                     0.75 rf grid2 model 33 0.7003973395525611
35
      4 150
                     0.75 rf grid2 model 36 0.6989577034925571
```

#### RANDOM GRID SEARCH

#### Train and validate a grid of RFs

```
# Get the grid results, sorted by validation auc
rf gridperf3 <- h2o.getGrid(grid id = "rf grid3",</pre>
                            sort_by = "auc",
                            decreasing = TRUE)
rf_gridperf3
# Output
H2O Grid Details
_____
Grid ID: rf grid3
Used hyper parameters:
 - mtries
  - ntrees
 - sample rate
Number of models: 36
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 mtries ntrees sample_rate
                                  model ids
                    0.55 rf grid3 model 18 0.6836247086247086
           50
                    0.55 rf_grid3_model_4 0.6831876456876457
          100
3
      2
           100
                     0.632 rf grid3 model 30 0.6830710955710956
```

```
0.632 rf grid3 model 8 0.6821095571095571
            50
      3
           200
                      0.55 rf grid3 model 15 0.6809440559440559
  mtries ntrees sample rate
                             model ids
                                                             auc
31
            100
                       0.75 rf grid3 model 12 0.6580419580419581
                       0.55 rf grid3 model 26 0.6574009324009323
32
            50
       3
                       0.75 rf grid3 model 16 0.6561771561771562
33
            150
34
            200
                       0.75 rf_grid3_model_13 0.6552738927738928
                       0.75 rf grid3 model 6 0.6528846153846154
35
            150
                       0.75 rf grid3 model 5 0.6500874125874126
36
       2
            50
```

## Grid Search KFold Cross Validation procedure

Train and validate a grid of RFs

```
Grid ID: rf grid4
Used hyper parameters:
 - mtries
 - ntrees
 - sample rate
Number of models: 36
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
mtries ntrees sample_rate model ids
                                                   auc
     2 200 0.55 rf_grid4_model_14 0.7319973510696496
     2 150 0.55 rf_grid4_model_35 0.7283694681984395
         100
                 0.55 rf_grid4_model_28 0.7271025884656359
               0.55 rf_grid4_model_30 0.7248279634908296
    2 50
5 3 200 0.55 rf grid4 model 36 0.7229420402522242
 mtries ntrees sample_rate model_ids
                                                   auc
                   0.75 rf grid4 model 15 0.7043563387175722
31
    3 200
32
     3 150
                   0.75 rf grid4 model 32 0.7037372952117704
33 4 50
                   0.75 rf grid4 model 2 0.701793786530765
                   0.75 rf grid4 model 19 0.7011315539431632
34
     4 100
35
     4 150
                    0.75 rf_grid4_model_26 0.7003973395525611
36 4 200
                    0.75 rf grid4 model 7 0.6989577034925571
```

## MODEL: GRADIENT BOOSTING MACHINE (GBM\_model)

#### CARTESIAN GRID SEARCH

## Grid Search Train / Test dataset procedure

#### **GBM** hyperparameters

## Train and validate a grid of GBMs

```
Grid ID: gbm grid1
Used hyper parameters:
 - learn rate
 - max depth
 - sample rate
Number of models: 24
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 learn rate max depth sample rate
                                        model ids
                                                                  auc
                             1.0 gbm grid1 model 24 0.9996794871794872
                             1.0 gbm grid1 model 21 0.9994463869463869
        0.1
                  15
                            1.0 gbm grid1 model 18 0.999067599067599
        0.1
                 10
        0.1
                 20
                            0.8 gbm grid1 model 12 0.995425407925408
        0.1
                  15
                             0.8 gbm grid1 model 9 0.995425407925408
  learn rate max depth sample rate
                                         model ids
       0.001
19
                   15
                            0.8 gbm grid1 model 7 0.8703088578088579
20
       0.001
                    5
                              0.8 gbm grid1 model 1 0.867511655011655
21
      0.001
                    5
                              1.0 gbm_grid1_model_13 0.8423368298368299
                              1.0 gbm grid1 model 16 0.8423368298368299
22
      0.001
                    10
                             1.0 gbm grid1 model 19 0.8423368298368299
      0.001
23
                   15
       0.001
                              1.0 gbm grid1 model 22 0.8423368298368299
24
                    20
```

## Grid Search KFold Cross Validation procedure

## Train and validate a grid of GBMs

```
fold_assignment="Stratified", # can be "AUTO", "Modulo", "R

# Early stop

score_tree_interval = 100,

stopping_rounds = 3,

stopping_tolerance = 0.001,

# Fixed hyperparameters

ntrees = 100,

# Optimized hyperparameters

hyper_params = gbm_params,

# Search type

search_criteria = list(strategy = "Cartesian"),

seed = 2020)
```

```
gbm_gridperf2 <- h2o.getGrid(grid_id = "gbm_grid2",</pre>
                           sort by = "auc",
                           decreasing = TRUE)
gbm gridperf2
Output
H2O Grid Details
_____
Grid ID: gbm_grid2
Used hyper parameters:
 - learn_rate
 - max_depth
 - sample rate
Number of models: 24
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 learn rate max depth sample rate
                                        model ids
                        1.0 gbm_grid2_model_14  0.741081454608275
      0.05
                            0.8 gbm grid2 model 1 0.7371224554432639
      0.001
3
       0.05
                  5
                            0.8 gbm_grid2_model_2 0.7367193573464629
```

```
0.8 gbm grid2 model 5 0.7359995393164609
        0.05
                    10
        0.05
                    20
                               0.8 gbm grid2 model 11 0.7358843684316605
   learn rate max depth sample rate
                                            model ids
                                                                      auc
19
        0.001
                     15
                                1.0 gbm grid2 model 19 0.7140882784831994
                                1.0 gbm grid2 model_22 0.7140882784831994
20
        0.001
                     20
                                1.0 gbm grid2 model 18 0.7123607152111947
         0.1
21
                     10
22
         0.1
                     20
                                1.0 gbm_grid2_model_24 0.7085888687339841
                                0.8 gbm grid2 model 6 0.7085888687339841
23
                     10
                                1.0 gbm grid2 model 21 0.7068037200195791
24
         0.1
                     15
```

#### RANDOM GRID SEARCH

## Random Grid Search Train / Test dataset procedure

#### Train and validate a grid of GBMs

```
hyper_params = gbm_params,

# Search type
search_criteria = search_criteria,
seed = 2020)
```

```
gbm_gridperf3 <- h2o.getGrid(grid_id = "gbm_grid3",</pre>
                          sort by = "auc",
                         decreasing = TRUE)
gbm gridperf3
Output
H2O Grid Details
_____
Grid ID: gbm grid3
Used hyper parameters:
 - learn rate
 - max depth
 - sample rate
Number of models: 24
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 1.0 gbm grid3 model 13 0.9996794871794872
       0.1
                20
       0.1
                15
                          1.0 gbm grid3 model 9 0.9994463869463869
       0.1
                 10
                           1.0 gbm grid3 model 10 0.999067599067599
                           0.8 gbm grid3 model 14 0.995425407925408
       0.1
                15
       0.1
                20
                          0.8 gbm_grid3_model_15  0.995425407925408
 learn rate max depth sample rate model ids
                                                              auc
19
       0.001
                  10
                           0.8 gbm grid3 model 6 0.8703088578088579
                            0.8 gbm_grid3_model_3 0.867511655011655
       0.001
                           1.0 gbm grid3 model 11 0.8423368298368299
      0.001
                  5
21
                            1.0 gbm grid3 model 12 0.8423368298368299
22
       0.001
                  10
```

```
23 0.001 15 1.0 gbm_grid3_model_16 0.8423368298368299
24 0.001 20 1.0 gbm_grid3_model_23 0.8423368298368299
```

## Random Grid Search KFold Cross Validation procedure

## Train and validate a grid of GBMs

```
gbm grid <- h2o.grid("gbm", y = "CAPSULE", x = c("AGE", "RACE", "PSA", "GLEASON")</pre>
                      grid_id = "gbm_grid4",
                      training frame= h2o df,
                      ignore const cols = TRUE,
                      nfolds = 10,
                      fold assignment="Stratified", # can be "AUTO", "Modulo", "R
andom" or "Stratified"
                      # Early stop
                      score tree interval = 100,
                      stopping rounds = 3,
                      stopping tolerance = 0.001,
                      # Fixed hyperparameters
                      ntrees = 100,
                      # Optimized hyperparameters
                      hyper params = gbm params,
                      # Search type
                      search_criteria = search_criteria,
                      seed = 2020)
```

```
______
Grid ID: gbm gri4
Used hyper parameters:
 - learn rate
 - max depth
 - sample rate
Number of models: 24
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
 learn rate max depth sample rate
                                       model ids
                                                                auc
                            1.0 gbm_grid4_model_15  0.741081454608275
      0.05
                            0.8 gbm grid4 model 23 0.7371224554432639
2
      0.001
                   5
                            0.8 gbm grid4 model 9 0.7367193573464629
      0.05
                  5
      0.05
                 10
                            0.8 gbm grid4 model 6 0.7359995393164609
                            0.8 gbm grid4 model 17 0.7358843684316605
      0.05
                 20
  learn rate max depth sample rate model ids
                            1.0 gbm grid4 model 18 0.7140882784831994
19
       0.001
                   15
       0.001
                             1.0 gbm grid4 model 22 0.7140882784831994
20
                   20
21
        0.1
                   10
                             1.0 gbm_grid4_model_5 0.7123607152111947
                             1.0 gbm_grid4_model_19 0.7085888687339841
22
        0.1
                   20
                            0.8 gbm grid4 model 7 0.7085888687339841
23
        0.1
                   10
                             1.0 gbm grid4 model 14 0.7068037200195791
24
         0.1
                   15
```

## MODEL: DEEP LEARNING - NEURAL NETWORKS (DL\_model)

#### CARTESIAN GRID SEARCH

## Grid Search Train / Test dataset procedure

## DL hyperparameters

## Train and validate a grid of DLs

```
dl grid <- h2o.grid(</pre>
                       # Algorithm
                      algorithm="deeplearning",
                      activation=c("RectifierWithDropout"),
                      epochs=500,
                      #adaptive rate = FALSE,
                      # Response variable and predictors
                      y = "CAPSULE",
                      x = c("AGE", "RACE", "PSA", "GLEASON"),
                      grid_id = "dl_grid1",
                      # Training data
                      training frame= train,
                      shuffle_training_data = FALSE,
                       # Preprocessed
                      standardize = TRUE,
                      missing_values_handling = "Skip",
                      ignore_const_cols = TRUE,
                      # Early stop
                      stopping metric="AUC",
                      stopping_tolerance= 0.01,
                      stopping rounds = 3,
                      # Regularization
                      11=1e-5,
                      12=1e-5,
                      # Optimized hyperparameters
```

```
hyper_params = dl_params,

# Search type
search_criteria = list(strategy = "Cartesian"),
seed = 2020)
```

## Get the grid results, sorted by validation AUC

```
dl gridperf1 <- h2o.getGrid(grid id = "dl grid1",</pre>
                            sort by = "auc",
                            decreasing = TRUE)
dl gridperf1
Output
H2O Grid Details
_____
Grid ID: dl grid1
Used hyper parameters:
 - hidden
Number of models: 9
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
     hidden model ids
    [1024] dl grid6 model 5 0.81931818181818
      [512] dl grid6 model 4 0.8131410256410256
     [256] dl grid6 model 3 0.8083041958041959
4 [256, 256] dl grid6 model 8 0.8052156177156178
      [128] dl grid6 model 2 0.8037004662004663
6 [512, 512] dl grid6 model 9 0.8035256410256411
7 [128, 128] dl_grid6_model_7 0.8032342657342657
       [64] dl grid6 model 1 0.8026515151515151
  [64, 64] dl grid6 model 6 0.8010780885780886
```

## Grid Search KFold Cross Validation procedure

Train and validate a grid of GBMs

```
dl grid <- h2o.grid(</pre>
                      # Algorithm
                      algorithm="deeplearning",
                      activation=c("RectifierWithDropout"),
                      epochs=500,
                      #adaptive rate = FALSE,
                      # Response variable and predictors
                      y = "CAPSULE",
                      x = c("AGE", "RACE", "PSA", "GLEASON"),
                      grid id = "dl grid2",
                      # Training data
                      training frame= h2o df,
                      shuffle_training_data = FALSE,
                      # Cross Validation
                      nfolds = 10,
                      fold assignment="Stratified", # can be "AUTO", "Modulo", "R
andom" or "Stratified"
                      # Preprocessed
                      standardize = TRUE,
                      missing_values_handling = "Skip",
                      ignore const cols = TRUE,
                      # Early stop
                      stopping_metric="AUC",
                      stopping tolerance= 0.01,
                      stopping rounds = 3,
                      # Regularization
                      11=1e-5,
                      12=1e-5,
                      # Optimized hyperparameters
                      hyper params = dl params,
                      # Search type
                      search criteria = list(strategy = "Cartesian"),
                      seed = 2020)
```

```
dl_gridperf2 <- h2o.getGrid(grid_id = "dl_grid2",</pre>
```

```
sort by = "auc",
                            decreasing = TRUE)
dl gridperf2
Output
H2O Grid Details
_____
Grid ID: dl_grid2
Used hyper parameters:
 - hidden
Number of models: 9
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
     hidden model ids
   [64, 64] dl grid2 model 6 0.7866171431862025
      [64] dl grid2 model 1 0.7848319944717975
2
      [256] dl_grid2_model_3 0.7813192824853876
      [128] dl grid2 model 2 0.7697446085629552
      [512] dl grid2 model 4 0.765022602286142
6 [128, 128] dl_grid2_model_7 0.7602718032881288
7 [512, 512] dl grid2 model 9 0.7557513460597161
    [1024] dl_grid2_model_5 0.7545996372117129
9 [256, 256] dl grid2 model 8 0.7353373067288589
```

#### RANDOM GRID SEARCH

## Random Grid Search Train / Test dataset procedure

#### Train and validate a grid of DLs

```
dl_grid <- h2o.grid(
```

```
# Algorithm
algorithm="deeplearning",
activation=c("RectifierWithDropout"),
epochs=500,
#adaptive rate = FALSE,
# Response variable and predictors
y = "CAPSULE",
x = c("AGE", "RACE", "PSA", "GLEASON"),
grid_id = "dl_grid3",
# Training data
training frame= train,
shuffle_training_data = FALSE,
# Preprocessed
standardize = TRUE,
missing values handling = "Skip",
ignore_const_cols = TRUE,
# Early stop
stopping_metric="AUC",
stopping tolerance= 0.01,
stopping rounds = 3,
# Regularization
11=1e-5,
12=1e-5,
# Optimized hyperparameters
hyper params = dl params,
# Search type
search_criteria = search_criteria,
seed = 2020)
```

```
______
Grid ID: dl grid3
Used hyper parameters:
 - hidden
Number of models: 9
Number of failed models: 0
Hyper-Parameter Search Summary: ordered by decreasing auc
     hidden model ids
    [1024] dl grid6 model 5 0.81931818181818
     [512] dl grid6 model 4 0.8131410256410256
      [256] dl_grid6_model_3 0.8083041958041959
4 [256, 256] dl_grid6_model_8 0.8052156177156178
    [128] dl grid6 model 2 0.8037004662004663
6 [512, 512] dl grid6 model 9 0.8035256410256411
7 [128, 128] dl grid6 model 7 0.8032342657342657
       [64] dl_grid6_model_1 0.8026515151515151
  [64, 64] dl grid6 model 6 0.8010780885780886
```

## Random Grid Search KFold Cross Validation procedure

## Train and validate a grid of DLs

```
dl_grid <- h2o.grid(
    # Algorithm
    algorithm="deeplearning",
    activation=c("RectifierWithDropout"),
    epochs=500,
    #adaptive_rate = FALSE,
    # Response variable and predictors
    y = "CAPSULE",
    x = c("AGE", "RACE", "PSA", "GLEASON"),
    grid_id = "dl_grid4",</pre>
```

```
# Training data
                      training frame= h2o df,
                      shuffle_training_data = FALSE,
                      # Cross Validation
                      nfolds = 10,
                      fold assignment="Stratified", # can be "AUTO", "Modulo", "R
andom" or "Stratified"
                      # Preprocessed
                      standardize = TRUE,
                      missing_values_handling = "Skip",
                      ignore_const_cols = TRUE,
                      # Early stop
                      stopping_metric="AUC",
                      stopping_tolerance= 0.01,
                      stopping_rounds = 3,
                      # Regularization
                      11=1e-5,
                      12=1e-5,
                      # Optimized hyperparameters
                      hyper_params = dl_params,
                      # Search type
                      search criteria = search criteria,
                      seed = 2020)
```

# Get the grid results, sorted by validation AUC

```
Number of models: 9

Number of failed models: 0

Hyper-Parameter Search Summary: ordered by decreasing auchidden model_ids auc

1 [1024] dl_grid4_model_5 0.8020500417494457

2 [64, 64] dl_grid4_model_2 0.7836227001813941

3 [64] dl_grid4_model_3 0.7834787365753938

4 [128, 128] dl_grid4_model_9 0.7688808269269529

5 [512, 512] dl_grid4_model_8 0.7673260199821486

6 [128] dl_grid4_model_1 0.7659151766433446

7 [512] dl_grid4_model_4 0.763237453571737

8 [256] dl_grid4_model_7 0.7584578618525237

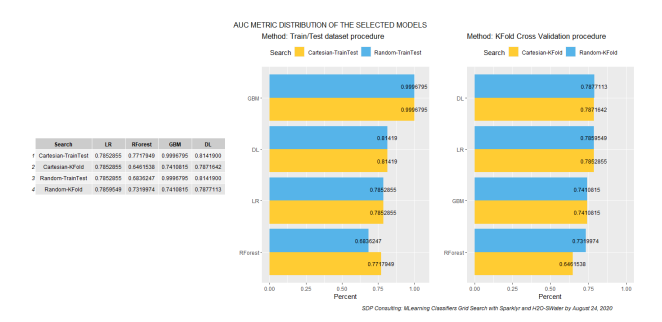
9 [256, 256] dl_grid4_model_6 0.7539661973453111
```

# SELECT BEST MODEL AND MAKE PREDICTIONS

### SELECT BEST MODEL

The models are compared based on the method (Train/Test or KFold) and the AUC metric obtained in each case.

These metric values are arranged in a table and then arranged in a possible format to process with ggplot2, generating the respective comparative visualization.



The Gradient Boosting Machine (GBM) algorithm performed the best and had a comparatively better AUC metric in the Train/Test method for both the Cartesian and Random procedures. Meantime, on the other hand, the Deep Learning-Neural Network (DL) model got a better AUC result for the KFold method in the two procedures applied.

### BEST MODELS BY TRAIN/TEST METHOD

### Selected model

The selected model is the Gradient Boosting Machine (GBM) algorithm with AUC = 0.9996794871794872 and Hyper-Parameter:

```
learn_rate = 0.1
max_depth = 20
sample_rate = 1.0
```

### Model characterization

The characterization of the model can be obtained using the print command as follows.

```
#Train your model using all data and the best known parameters
GBM model <- h2o.gbm(x = c("AGE", "RACE", "PSA", "GLEASON"),
                y = "CAPSULE",
                training_frame = train,
                learn rate = 0.1,
                max_depth = 20,
                sample rate = 1.0,
                ntrees = 100,
                seed = 2020)
print(GBM_model)
Output:
Model Details:
_____
H2OBinomialModel: gbm
Model ID: GBM model R 1598310950573 1
Model Summary:
 number_of_trees number_of_internal_trees model_size_in_bytes min_depth
            100
                                                       32264
                                      100
```

```
max depth mean depth min leaves max leaves mean leaves
1 16 10.04000 19 24 20.96000
H2OBinomialMetrics: gbm
** Reported on training data. **
MSE: 0.03192855
RMSE: 0.1786856
LogLoss: 0.1498627
Mean Per-Class Error: 0.009090909
AUC: 0.9996795
AUCPR: 0.9898817
Gini: 0.999359
R^2: 0.8653624
Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
      0 1 Error Rate
     162 3 0.018182 =3/165
     0 104 0.000000 =0/104
Totals 162 107 0.011152 =3/269
Maximum Metrics: Maximum metrics at their respective thresholds
                     metric threshold value idx
                    max f1 0.389477 0.985782 104
1
2
                    max f2 0.389477 0.994264 104
              max f0point5 0.469360 0.994094 99
              max accuracy 0.469360 0.988848 99
4
              max precision 0.999511 1.000000 0
5
                max recall 0.389477 1.000000 104
7
            max specificity 0.999511 1.000000 0
           max absolute mcc 0.389477 0.976878 104
8
   max min per class accuracy 0.430621 0.987879 102
10 max mean per class accuracy 0.389477 0.990909 104
                   max tns 0.999511 165.000000 0
11
12
                   max fns 0.999511 103.000000 0
```

### **Making Predictions**

The Train/Test method train the models using the "train" data as the training data. So, to use the model to predict we have to train the model using all data (h2o\_df) and the best-known parameters.

Below is a brief view of the predictions obtained based on the dataset.

```
# Generate the predictions on a test set (if necessary):

pred <- h2o.predict(GBM_model, newdata = test)

pred

Output:

predict p0 p1

1 1 0.5893065 0.410693521

2 1 0.3543368 0.645663223

3 1 0.5505129 0.449487099

4 0 0.9556726 0.044327425

5 0 0.9905264 0.009473623

6 0 0.8623784 0.137621574
```

[111 rows x 3 columns]

### BEST MODELS BY KFOLD METHOD

### Selected model

The selected model is the Deep Learning-Neural Network (DL) algorithm with AUC = 0.7920589674930177 and Hyper-Parameter:

```
hidden = [1024]
```

### Model characterization

The characterization of the model can be obtained using the print command as follows.

```
DL model <- h2o.deeplearning(</pre>
                       # Algorithm
                      activation=c("RectifierWithDropout"),
                      epochs=500,
                       #adaptive_rate = FALSE,
                      hidden = c(1024),
                      input dropout ratio = 0.0,
                       # Response variable and predictors
                      y = "CAPSULE",
                       x = c("AGE", "RACE", "PSA", "GLEASON"),
                       # Training data
                       training frame= h2o df,
                       # Preprocessed
                       standardize = TRUE,
                      missing values handling = "Skip",
                      ignore const cols = TRUE,
                       # Early stop
                       stopping_metric="AUC",
                       stopping tolerance= 0.01,
                       stopping rounds = 3,
                       # Regularization
                      11=1e-5,
```

```
12=1e-5,
                    seed = 2020)
print(DL model)
Output:
Model Details:
_____
H2OBinomialModel: deeplearning
Model ID: DeepLearning model R 1598718661326 672
Status of Neuron Layers: predicting CAPSULE, 2-class classification, bernoulli di
stribution, CrossEntropy loss, 10.242 weights/biases, 135,0 KB, 190.000 training
samples, mini-batch size 1
                                        11
 layer units
                      type dropout
                                                 12 mean rate rate rms
    1 7
                      Input 0.00 %
                                                 NA NA
                                         NA
    2 1024 RectifierDropout 50.00 % 0.000010 0.000010 0.193687 0.343031
                     Softmax NA 0.000010 0.000010 0.009714 0.009128
 momentum mean weight weight rms mean bias bias rms
1 NA
                 NA
                           NA
                                    NA
                                            NA
2 0.000000 -0.052502 0.262709 -0.056548 0.279210
3 0.000000 0.008416 0.148099 0.056109 0.140763
H2OBinomialMetrics: deeplearning
** Reported on training data. **
** Metrics reported on full training frame **
MSE: 0.1659942
RMSE: 0.4074238
LogLoss: 0.4883401
Mean Per-Class Error: 0.2623161
AUC: 0.825329
AUCPR: 0.7819455
Gini: 0.6506579
Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
```

```
Error
                           Rate
      145 82 0.361233 =82/227
       25 128 0.163399 =25/153
Totals 170 210 0.281579 =107/380
Maximum Metrics: Maximum metrics at their respective thresholds
                      metric threshold value idx
1
                      max f1 0.319374 0.705234 207
2
                      max f2 0.134448 0.821545 304
                max f0point5 0.737796 0.724947 77
               max accuracy 0.554822 0.771053 136
4
              max precision 0.999995 1.000000 0
5
6
                  max recall 0.036747 1.000000 339
7
              max specificity 0.999995 1.000000 0
8
             max absolute mcc 0.554822 0.518132 136
   max min per class accuracy 0.443643 0.726872 171
9
10 max mean per class accuracy 0.554822 0.754038 136
11
                     max tns 0.999995 227.000000
                     max fns 0.999995 152.000000
12
13
                     max fps 0.000000 227.000000 376
                     max tps 0.036747 153.000000 339
14
15
                     max tnr 0.999995 1.000000 0
16
                     max fnr 0.999995 0.993464
                     max fpr 0.000000 1.000000 376
17
                     max tpr 0.036747 1.000000 339
18
Gains/Lift Table: Extract with `h2o.gainsLift(, )` or `h2o.gainsLift(, valid=, xv
al=)
```

## **Making Predictions**

The KFold method trains the models using all data (h2o\_df) as the training data. So we can use directly the selected model which already considers the best-known parameters.

```
# Best model chosen by AUC:
best_model <- h2o.getModel(dl_gridperf4@model_ids[[1]])</pre>
```

Below is a brief view of the predictions obtained based on the dataset.

```
# Generate the predictions on a test set (if necessary):

pred <- h2o.predict(best_model, newdata = test)

pred

Output:

predict p0 p1

1 1 0.66265631 3.373437e-01

2 1 0.36200723 6.379928e-01

3 1 0.08907328 9.109267e-01

4 0 0.99992892 7.107996e-05

5 0 0.98164801 1.835199e-02

6 1 0.20583521 7.941648e-01
```

## FINAL WORDS

This project is about a practical exposition of how to apply two classic Grid Search methods for tunning four of the most frequently used machine learning models for classification, by using Sparklyr and H2O-SWater.

The tune processes can obviously be optimized either by modifying the current hyperparameter values or by incorporating others. However, it should be noted that incorporating more hyperparameter into the tune will require a greater amount of process time and, in some cases, will not be feasible to carry out.

Data can be loaded directly into the H2O cluster, or -as was done in this projectby first loading it into memory in the R session and then transferring it. The second option is not recommended if the data volume is very large.

While these models ran on a small data set in a local spark cluster, these methods can be scaled, for the most part, for data analysis in a distributed Apache Spark cluster.

Hector Alvaro Rojas | Data Science, Visualizations and Applied Statistics | August 30, 2020