



Machine learning with H2O

Dr. Shirin Glander
Data Scientist



What is H2O?

```
library(h2o)
h2o.init()
H2O is not running yet, starting it now...
java version "1.8.0 131"
Java (TM) SE Runtime Environment (build 1.8.0 131-b11)
Java HotSpot(TM) 64-Bit Server VM (build 25.131-b11, mixed mode)
Starting H2O JVM and connecting: ... Connection successful!
R is connected to the H2O cluster:
   H2O cluster uptime: 2 seconds 124 milliseconds
   H2O cluster version: 3.20.0.8
   H2O cluster total nodes: 1
   H2O cluster total memory: 3.56 GB
   H2O cluster total cores:
                         localhost
   H2O Connection ip:
   H2O Connection port: 54321
   H2O API Extensions:
                              XGBoost, Algos, AutoML, Core V3, Core V4
   R Version:
                              R version 3.5.1 (2018-07-02)
```



New dataset: seeds data

```
glimpse(seeds data)
Observations: 150
Variables: 8
$ area
           <dbl> 15.26, 14.88, 14.29, 13.84, 16.14, 14.38, 14.69, ...
$ perimeter <dbl> 14.84, 14.57, 14.09, 13.94, 14.99, 14.21, 14.49, ...
$ compactness <dbl> 0.8710, 0.8811, 0.9050, 0.8955, 0.9034, 0.8951, ...
$ kernel length <dbl> 5.763, 5.554, 5.291, 5.324, 5.658, 5.386, 5.563, ...
$ kernel width <dbl> 3.312, 3.333, 3.337, 3.379, 3.562, 3.312, 3.259, ...
$ asymmetry
           <dbl> 2.2210, 1.0180, 2.6990, 2.2590, 1.3550, 2.4620, ...
$ kernel groove <dbl> 5.220, 4.956, 4.825, 4.805, 5.175, 4.956, 5.219, ...
seeds data %>%
 count (seed type)
# A tibble: 3 x 2
 seed type
     <int> <int>
             50
            50
             50
```

Preparing the data for modeling with H2O

Data as H2O frame

```
seeds_data_hf <- as.h2o(seeds_data)
```

• Define **features and target** variable

```
y <- "seed_type"
x <- setdiff(colnames(seeds_data_hf), y)
```

For classification target should be a factor

```
seeds_data_hf[, y] <- as.factor(seeds_data_hf[, y])
```



Training, validation and test sets

```
sframe <- h2o.splitFrame(data = seeds data hf,</pre>
                            ratios = c(0.7, 0.15),
                            seed = 42)
train <- sframe[[1]]</pre>
valid <- sframe[[2]]</pre>
test <- sframe[[3]]</pre>
summary(train$seed type, exact quantiles = TRUE)
seed type
1:36
 2:36
 3:35
summary(test$seed type, exact quantiles = TRUE)
 seed type
 1:8
 2:8
 3:5
```

Model training with H2O

- Gradient Boosted models with h2o.gbm() & h2o.xgboost()
- Generalized linear models with h2o.glm()
- Random Forest models with h2o.randomForest()
- Neural Networks with h2o.deeplearning()



Evaluate model performance with H2O

Model performance

Predict new data

```
h2o.predict(gbm_model, test)
```





Let's practice!





Grid and random search with H2O

Dr. Shirin Glander
Data Scientist

Hyperparameters in H2O models

Hyperparameters for Gradient Boosting:

?h2o.gbm

- ntrees: Number of trees. Defaults to 50.
- max depth: Maximum tree depth. Defaults to 5.
- min rows: Fewest allowed (weighted) observations in a leaf. Defaults to 10.
- learn_rate: Learning rate (from 0.0 to 1.0) Defaults to 0.1.
- learn_rate_annealing: Scale the learning rate by this factor after each tree (e.g., 0.99 or 0.999) Defaults to 1.



Preparing our data for modeling with H2O

Convert to H2O frame

```
seeds_data_hf <- as.h2o(seeds_data)
```

Identify features and target

```
y <- "seed_type"
x <- setdiff(colnames(seeds_data_hf), y)
```

• Split data into train, test & validation set



Defining a hyperparameter grid

GBM hyperparamters

• h2o.grid function

Examine results with h2o.getGrid



Examining a grid object

- Examine results for our model gbm grid with h2o.getGrid function.
- Get the grid results sorted by validation accuracy



Extracting the best model from a grid

Top GBM model chosen by validation accuracy has id position 1

```
best_gbm <- h2o.getModel(gbm_gridperf@model_ids[[1]])</pre>
```

These are the hyperparameters for the best model:

best_gbm is a regular H2O model object and can be treated as such!

```
h2o.performance(best_gbm, test)

MSE: (Extract with `h2o.mse`) 0.04761904

RMSE: (Extract with `h2o.rmse`) 0.2182179

Logloss: (Extract with `h2o.loglos
```



Random search with H2O

• In addition to hyperparameter grid, add search criteria:



Stopping criteria

```
search criteria <- list(strategy = "RandomDiscrete",</pre>
                         stopping_metric = "mean_per_class_error",
                        stopping tolerance = 0.0001,
                        stopping rounds = 6)
gbm grid <- h2o.grid("gbm",</pre>
                     X = X
                     y = y
                     training frame = train,
                     validation frame = valid,
                     seed = 42,
                     hyper params = gbm params,
                     search criteria = search criteria)
H2O Grid Details
==========
Grid ID: gbm grid
Used hyper parameters:
    learn rate
    max depth
    ntrees
Number of models: 30
Number of failed models: 0
```





Time to practise!





Automatic machine learning & hyperparameter tuning with H2O

Dr. Shirin Glander
Data Scientist

Automatic Machine Learning (AutoML)

• Automatic tuning of algorithms, in addition to hyperparameters

AutoML makes model tuning and optimization much faster and easier

AutoML only needs a dataset, a target variable and a time or model number
 limit for training

AutoML in H2O

AutoML compares

- Generalized Linear Model (GLM)
- (Distributed) Random Forest (DRF)
- Extremely Randomized Trees (XRT)
- Extreme Gradient Boosting (XGBoost)
- Gradient Boosting Machines (GBM)
- Deep Learning (fully-connected multi-layer artificial neural network)
- Stacked **Ensembles** (of all models & of best of family)



Hyperparameter tuning in H2O's AutoML

GBM Hyperparameters

- histogram type
- ntrees
- max depth
- min rows
- learn rate
- sample_rate
- col sample rate
- col_sample_rate_per_tree

Deep Learning Hyperparameters

- epochs
- adaptivate_rate
- activation
- rho
- epsilon
- input_dropout_ratio
- hidden
- hidden dropout ratios

min colit improvement



Using AutoML with H2O

• h2o.automl function

• returns a leaderboard of all models, ranked by the chosen metric (here "logloss")



Viewing the AutoML leaderboard

```
| model_id mean_per_class_error | model_id mean_per_class_error | GBM_grid_0_AutoML_20181029_144443_model_6 | 0.01851852 | 0.02777778 | 0.02777778 | 0.02777778 | 0.02777778 | 0.02777778 | 0.02777778 | 0.02777778 | 0.03703704 | GBM_grid_0_AutoML_20181029_144443_model_9 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703704 | 0.03703
```

Per default, the leaderboard is calculated on 5-fold cross-validation.

http://docs.h2o.ai/h2o/latest-stable/h2o-docs/automl.html



Extracting models from AutoML leaderboard

List all models by model id

```
model_ids <- as.data.frame(lb)$model_id

[1] "GBM_grid_0_AutoML_20181029_144443_model_6"

[3] "GBM_grid_0_AutoML_20181029_144443_model_18"

[19] "XRT_0_AutoML_20181029_144443"

[20] "DRF_0_AutoML_20181029_144443"

[24] "DeepLearning_0_AutoML_20181029_144443"

[41] "StackedEnsemble_BestOfFamily_0_AutoML_20181029_144443"

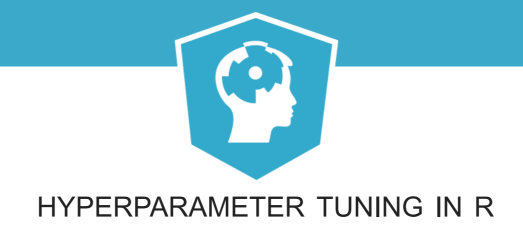
[42] "StackedEnsemble_AllModels_0_AutoML_20181029_144443"</pre>
```

Get the best model

```
aml_leader <- automl_model@leader
```

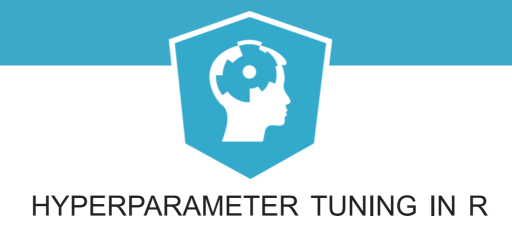
• aml leader is again a regular H2O model object and can be treated as such!





Get ready for your last round of exercises!





Congratulations!

Dr. Shirin Glander
Data Scientist



What you've learned in this course

- What hyperparameters are
- How they are different from model parameters
- And why to tune them
- **How** tuning works in three R packages:
- caret
- mlr
- h2o

Terms you can understand and apply

- Cartesian Grid Search
- Random Search
- Adaptive Resampling
- Automatic Machine Learning
- Evaluating tuning results with performance metrics
- Stopping criteria



How you can use this knowledge

- Find best hyperparameter set for your models
- Compare and contrast R packages => favorite

Where to go from here?

- Package manuals & vignettes
- Try it out!
- UC Irvine Machine Learning Repository
- Kaggle





Thank you and have fun!