# Webinar: Building meaningful machine learning models for disease prediction

Dr. Shirin Glander
March 31, 2017

Webinar for ISDS R Group: http://www.syndromic.org/cop/r

Description: Dr Shirin Glander will go over her work on building machine-learning models to predict the course of different diseases. She will go over building a model, evaluating its performance, and answering or addressing different disease related questions using machine learning. Her talk will cover the theory of machine learning as it is applied using R.

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https://shiring.github.io

https://github.com/ShirinG

Slides and code will be available on Github: https://github.com/ShirinG/Webinar\_ML\_for\_disease

Code will also be on my website: https://shiring.github.io

Can we predict flu deaths with Machine Learning and R?: https://shiring.github.io/machine\_learning/2016/11/27/flu\_outcome\_ML\_post Extreme Gradient Boosting and Preprocessing in Machine Learning - Addendum to predicting flu outcome with R: https://shiring.github.io/machine\_learning/2016/12/02/flu\_outcome\_ML\_2\_post Feature Selection in Machine Learning (Breast Cancer Datasets): https://shiring.github.io/machine\_learning/2017/01/15/rfe\_ga\_post Predicting food preferences with sparklyr (machine learning): https://shiring.github.io/machine\_learning/2017/02/19/food\_spark Building deep neural nets with h2o and rsparkling that predict arrhythmia of the heart: https://shiring.github.io/machine\_learning/2017/02/27/h2o

#### **Dataset**

### Breast Cancer Wisconsin (Diagnostic) Dataset

The data was downloaded from the UC Irvine Machine Learning Repository. The features in these datasets characterise cell nucleus properties and were generated from image analysis of fine needle aspirates (FNA) of breast masses.

The first dataset looks at the predictor classes:

- malignant or
- benign breast mass.

The phenotypes for characterisation are:

- Sample ID (code number)
- Clump thickness
- Uniformity of cell size
- Uniformity of cell shape
- Marginal adhesion
- Single epithelial cell size
- Number of bare nuclei

- Bland chromatin
- Number of normal nuclei
- Mitosis
- Classes, i.e. diagnosis

Missing values are imputed with the *mice* package.

```
bc_data <- read.table("datasets/breast-cancer-wisconsin.data.txt", header = FALSE, sep = ",")</pre>
colnames(bc_data) <- c("sample_code_number",</pre>
                        "clump_thickness",
                        "uniformity_of_cell_size",
                        "uniformity_of_cell_shape",
                        "marginal adhesion",
                        "single_epithelial_cell_size",
                        "bare nuclei",
                        "bland_chromatin",
                        "normal_nucleoli",
                        "mitosis",
                        "classes")
bc_data$classes <- ifelse(bc_data$classes == "2", "benign",</pre>
                           ifelse(bc_data$classes == "4", "malignant", NA))
bc_data[bc_data == "?"] <- NA</pre>
# how many NAs are in the data
length(which(is.na(bc_data)))
# impute missing data
library(mice)
bc data[,2:10] <- apply(bc data[, 2:10], 2, function(x) as.numeric(as.character(x)))
dataset_impute <- mice(bc_data[, 2:10], print = FALSE)</pre>
bc_data <- cbind(bc_data[, 11, drop = FALSE], mice::complete(dataset_impute, 1))</pre>
bc_data$classes <- as.factor(bc_data$classes)</pre>
# how many benign and malignant cases are there?
summary(bc_data$classes)
```

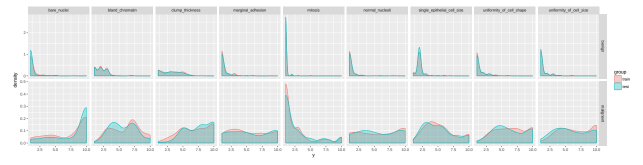
## Machine Learning packages for R

caret

```
library(caret)
```

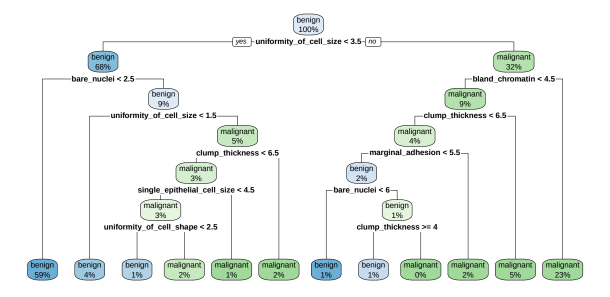
# Training, validation and test data

```
set.seed(42)
index <- createDataPartition(bc_data$classes, p = 0.7, list = FALSE)
train_data <- bc_data[index, ]
test_data <- bc_data[-index, ]</pre>
```



## Classification

#### Decision trees



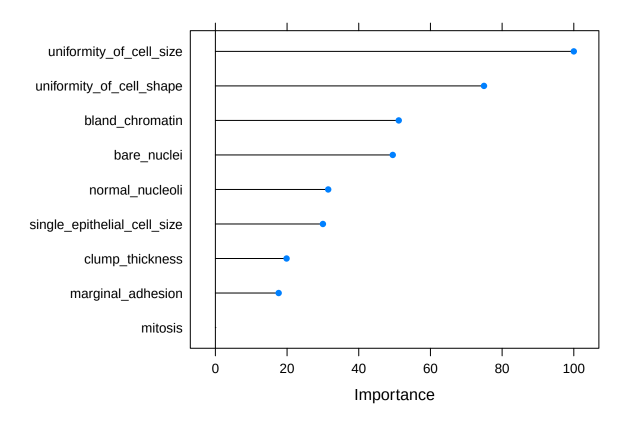
#### **Random Forests**

Can be used for classification and regression tasks. Here, I show a classification task.

When you specify savePredictions = TRUE, you can access the cross-validation resuls with model\_rf\$pred.

• Feature Importance

```
# estimate variable importance
importance <- varImp(model_rf, scale = TRUE)
plot(importance)</pre>
```



• predicting test data

```
confusionMatrix(predict(model_rf, test_data), test_data$classes)
```

```
## Confusion Matrix and Statistics
##
              Reference
##
  Prediction benign malignant
##
     benign
                  133
##
##
     malignant
                             70
##
                  Accuracy : 0.9713
##
                    95% CI: (0.9386, 0.9894)
##
##
       No Information Rate: 0.6555
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.9369
##
    Mcnemar's Test P-Value: 0.6831
##
               Sensitivity: 0.9708
##
               Specificity: 0.9722
##
            Pos Pred Value: 0.9852
##
            Neg Pred Value: 0.9459
##
                Prevalence: 0.6555
##
            Detection Rate: 0.6364
##
##
      Detection Prevalence: 0.6459
##
         Balanced Accuracy: 0.9715
```

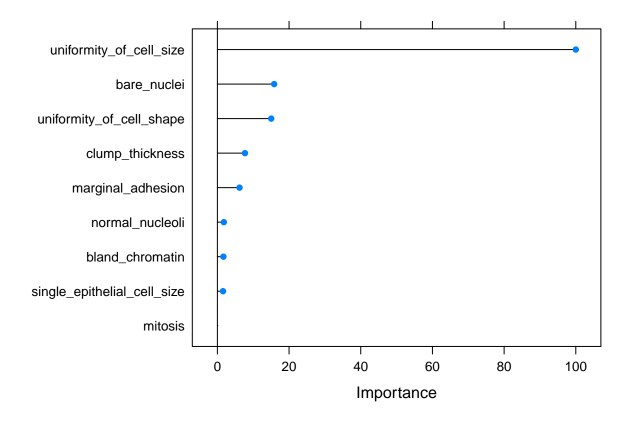
```
##
## 'Positive' Class : benign
##
```

## Extreme gradient boosting trees

Can be used for classification and regression tasks. Here, I show a classification task.

• Feature Importance

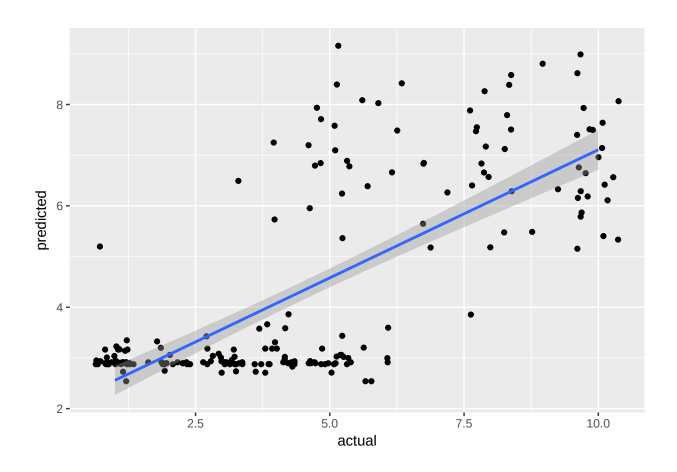
```
# estimate variable importance
importance <- varImp(model_xgb, scale = TRUE)
plot(importance)</pre>
```



• predicting test data

```
## Confusion Matrix and Statistics
##
              Reference
##
## Prediction benign malignant
##
     benign
                  132
                              2
                             70
     malignant
                    5
##
##
                  Accuracy : 0.9665
##
##
                    95% CI: (0.9322, 0.9864)
##
       No Information Rate: 0.6555
##
       P-Value [Acc > NIR] : <2e-16
##
##
                     Kappa: 0.9266
   Mcnemar's Test P-Value : 0.4497
##
##
##
               Sensitivity: 0.9635
##
               Specificity: 0.9722
##
            Pos Pred Value: 0.9851
            Neg Pred Value: 0.9333
##
##
                Prevalence: 0.6555
##
            Detection Rate: 0.6316
##
      Detection Prevalence : 0.6411
         Balanced Accuracy: 0.9679
##
##
##
          'Positive' Class : benign
##
Regression
set.seed(42)
model_glm <- caret::train(clump_thickness ~ .,</pre>
                          data = train_data,
                          method = "glm",
                          preProcess = c("scale", "center"),
                          trControl = trainControl(method = "repeatedcv",
                                                   number = 10,
                                                   repeats = 10,
                                                   savePredictions = TRUE,
                                                   verboseIter = FALSE))
data.frame(actual = test_data$clump_thickness,
           predicted = predict(model_glm, test_data)) %>%
  ggplot(aes(x = actual, y = predicted)) +
   geom_jitter() +
   geom_smooth(method = "lm")
```

confusionMatrix(predict(model\_xgb, test\_data), test\_data\$classes)



## Grid search with h2o

```
library(h2o)
h2o.init()
    Connection successful!
##
##
\mbox{\tt \#\#} R is connected to the H2O cluster:
       H2O cluster uptime:
                                     21 minutes 15 seconds
##
                                     3.10.3.6
##
       H2O cluster version:
       H2O cluster version age:
                                     13 days
##
       H20 cluster name:
                                     H20_started_from_R_s_glan02_tbd690
##
       H2O cluster total nodes:
##
       H2O cluster total memory:
                                     3.40 GB
##
       H2O cluster total cores:
                                     8
                                     2
##
       H2O cluster allowed cores:
##
       H2O cluster healthy:
                                     TRUE
       H20 Connection ip:
##
                                     localhost
##
       H20 Connection port:
                                     54321
       H2O Connection proxy:
##
       R Version:
                                     R version 3.3.2 (2016-10-31)
bc_data_hf <- as.h2o(bc_data)</pre>
##
```

```
0%
                                                     =======| 100%
library(tidyr)
h2o.describe(bc_data_hf) %>%
  gather(x, y, Zeros:Sigma) %>%
  mutate(group = ifelse(x %in% c("Min", "Max", "Mean"), "min, mean, max",
                             ifelse(x %in% c("NegInf", "PosInf"), "Inf", "sigma, zeros"))) %>%
  ggplot(aes(x = Label, y = as.numeric(y), color = x)) +
     geom_point(size = 4, alpha = 0.6) +
    scale_color_brewer(palette = "Set1") +
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust = 1)) +
    facet_grid(group ~ ., scales = "free") +
    labs(x = "Feature",
          y = "Value",
          color = "")
         0.50 -
         0.25 -
                                                                                   ΞŢ
         0.00
        -0.25 -
        -0.50 -
         10.0 -
                                                                                              Max
                                                                                              Mean
                                                                                   min, mean, max
          7.5 -
                                                                                              Min
           5.0 -
                                                                                              NegInf
           2.5 -
                                                                                              PosInf
                                                                                              Sigma
           0.0 -
                                                                                              Zeros
          400 -
                                                                                   sigma, zeros
          300 -
          200 -
          100 -
                                               rijosis nucedi el ste stape di site riordia di cell site sirde epitrelial cell site unitornità di cell site
            0 -
```

Feature

## Training, validation and test data

```
splits <- h2o.splitFrame(bc_data_hf,</pre>
                          ratios = c(0.7, 0.15),
                          seed = 1)
train <- splits[[1]]</pre>
valid <- splits[[2]]</pre>
test <- splits[[3]]</pre>
response <- "classes"
features <- setdiff(colnames(train), response)</pre>
summary(train$classes, exact_quantiles = TRUE)
## classes
## benign
             :317
## malignant:174
summary(valid$classes, exact_quantiles = TRUE)
## classes
## benign
             :71
## malignant:35
summary(test$classes, exact_quantiles = TRUE)
## classes
## benign
             :70
## malignant:32
```

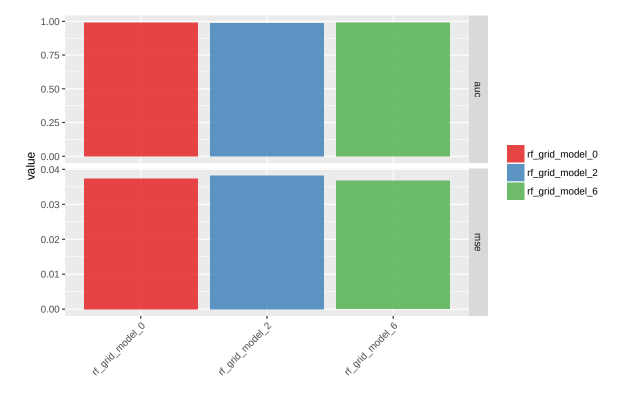
#### Classification

### Random Forest

x = features,

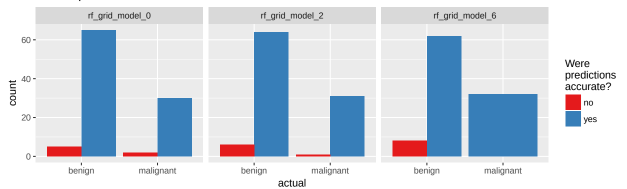
```
Can be used for classification and regression tasks. Here, I show a classification task.
hyper_params <- list(</pre>
                      ntrees = c(25, 50, 75, 100),
                      \max_{depth} = c(10, 20, 30),
                      min_rows = c(1, 3, 5)
search_criteria <- list(</pre>
                          strategy = "RandomDiscrete",
                          max_models = 50,
                          max runtime secs = 360,
                          stopping_rounds = 5,
                          stopping_metric = "AUC",
                          stopping_tolerance = 0.0005,
                          seed = 42
                          )
rf_grid <- h2o.grid(algorithm = "randomForest", # h2o.randomForest,
                                                    # alternatively h2o.gbm for Gradient boosting trees
```

```
y = response,
                     grid_id = "rf_grid",
                     training_frame = train,
                     validation_frame = valid,
                     nfolds = 25,
                     fold_assignment = "Stratified",
                     hyper_params = hyper_params,
                     search_criteria = search_criteria,
                     seed = 42
# performance metrics where smaller is better -> order with decreasing = FALSE
sort_options_1 <- c("mean_per_class_error", "mse", "err", "logloss")</pre>
for (sort_by_1 in sort_options_1) {
  grid <- h2o.getGrid("rf_grid", sort_by = sort_by_1, decreasing = FALSE)</pre>
 model_ids <- grid@model_ids</pre>
 best_model <- h2o.getModel(model_ids[[1]])</pre>
 h2o.saveModel(best model, path="models", force = TRUE)
}
# performance metrics where bigger is better -> order with decreasing = TRUE
sort_options_2 <- c("auc", "precision", "accuracy", "recall", "specificity")</pre>
for (sort_by_2 in sort_options_2) {
  grid <- h2o.getGrid("rf_grid", sort_by = sort_by_2, decreasing = TRUE)</pre>
 model ids <- grid@model ids
 best_model <- h2o.getModel(model_ids[[1]])</pre>
 h2o.saveModel(best_model, path="models", force = TRUE)
files <- list.files(path = "models")
rf_models <- files[grep("rf_grid_model", files)]</pre>
for (model_id in rf_models) {
 path <- paste0("U:\\Github_blog\\Webinar\\Webinar_ML_for_disease\\models\\", model_id)</pre>
  best_model <- h2o.loadModel(path)</pre>
  mse_auc_test <- data.frame(model_id = model_id,</pre>
                              mse = h2o.mse(h2o.performance(best_model, test)),
                              auc = h2o.auc(h2o.performance(best_model, test)))
  if (model_id == rf_models[[1]]) {
    mse_auc_test_comb <- mse_auc_test</pre>
```



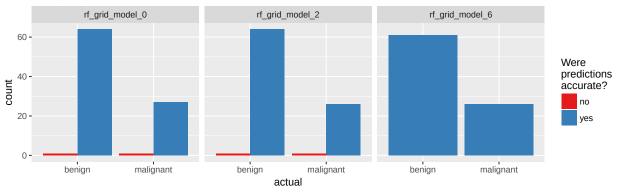
```
"yes", "no")
 finalRf_predictions$predict_stringent <- ifelse(finalRf_predictions$benign > 0.8,
                                                "benign",
                                                ifelse(finalRf_predictions$malignant > 0.8,
                                                       "malignant", "uncertain"))
 finalRf_predictions$accurate_stringent <- ifelse(finalRf_predictions$actual == finalRf_predictions$pr
                                                 "yes",
                                        ifelse(finalRf_predictions$predict_stringent == "uncertain",
                                              "na", "no"))
 if (model id == rf models[[1]]) {
   finalRf_predictions_comb <- finalRf_predictions</pre>
 } else {
   finalRf_predictions_comb <- rbind(finalRf_predictions_comb, finalRf_predictions)</pre>
 }
##
                                                                      0%
            _____| 100%
##
                                                                      0%
                                                                 =| 100%
##
                                                                      0%
                                                                ==| 100%
finalRf_predictions_comb %>%
 ggplot(aes(x = actual, fill = accurate)) +
   geom_bar(position = "dodge") +
   scale_fill_brewer(palette = "Set1") +
   facet_wrap(~ model_id, ncol = 3) +
   labs(fill = "Were\npredictions\naccurate?",
        title = "Default predictions")
```

#### Default predictions



```
finalRf_predictions_comb %>%
  subset(accurate_stringent != "na") %>%
  ggplot(aes(x = actual, fill = accurate_stringent)) +
    geom_bar(position = "dodge") +
    scale_fill_brewer(palette = "Set1") +
    facet_wrap(~ model_id, ncol = 3) +
    labs(fill = "Were\npredictions\naccurate?",
        title = "Stringent predictions")
```

# Stringent predictions



rf\_model <- h2o.loadModel("U:\\Github\_blog\\Webinar\\Webinar\_ML\_for\_disease\\models\\rf\_grid\_model\_6")
rf model</pre>

```
## Model Details:
## =======
##
## H2OBinomialModel: drf
## Model ID: rf_grid_model_6
## Model Summary:
##
     number_of_trees number_of_internal_trees model_size_in_bytes min_depth
## 1
                                          100
                                                             24848
##
     max_depth mean_depth min_leaves max_leaves mean_leaves
## 1
                  5.77000
                                   8
                                                   12.94000
                                             17
##
##
## H20BinomialMetrics: drf
## ** Reported on training data. **
## ** Metrics reported on Out-Of-Bag training samples **
```

```
##
## MSE: 0.03055597
## RMSE: 0.1748027
## LogLoss: 0.1140213
## Mean Per-Class Error: 0.02467457
## AUC: 0.989521
## Gini: 0.979042
##
## Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
##
             benign malignant
                                 Error
                                           Rate
## benign
                305
                          12 0.037855
                                       =12/317
                  2
                          172 0.011494
                                         =2/174
## malignant
## Totals
                307
                          184 0.028513 =14/491
##
## Maximum Metrics: Maximum metrics at their respective thresholds
##
                           metric threshold
                                               value idx
## 1
                           max f1 0.445310 0.960894 172
## 2
                           max f2 0.214464 0.977528 182
## 3
                    max f0point5 0.552688 0.946712 165
## 4
                    max accuracy 0.445310 0.971487 172
## 5
                   max precision 1.000000 1.000000
## 6
                      max recall 0.214464 1.000000 182
                 max specificity 1.000000 1.000000
## 7
                max absolute_mcc 0.445310 0.939393 172
## 8
      max min_per_class_accuracy 0.487375 0.965300 167
## 10 max mean_per_class_accuracy   0.445310   0.975325   172
## Gains/Lift Table: Extract with `h2o.gainsLift(<model>, <data>)` or `h2o.gainsLift(<model>, valid=<T/
## H20BinomialMetrics: drf
## ** Reported on validation data. **
##
## MSE: 0.01590167
## RMSE: 0.1261018
## LogLoss: 0.07067446
## Mean Per-Class Error: 0.007042254
## AUC: 0.9995976
## Gini: 0.9991952
##
## Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
##
             benign malignant
                                 Error
                                          Rate
                 70
                           1 0.014085
                                         =1/71
## benign
## malignant
                 0
                           35 0.000000
                                        =0/35
                 70
                           36 0.009434 =1/106
## Totals
##
## Maximum Metrics: Maximum metrics at their respective thresholds
##
                           metric threshold
                                               value idx
## 1
                           max f1 0.293126 0.985915
## 2
                           max f2 0.293126 0.994318 34
## 3
                    max f0point5 0.555841 0.994152
## 4
                    max accuracy 0.555841 0.990566
## 5
                    max precision 1.000000 1.000000
## 6
                      max recall 0.293126 1.000000 34
## 7
                 max specificity 1.000000 1.000000
                                                      0
                max absolute_mcc 0.293126 0.979045 34
## 8
```

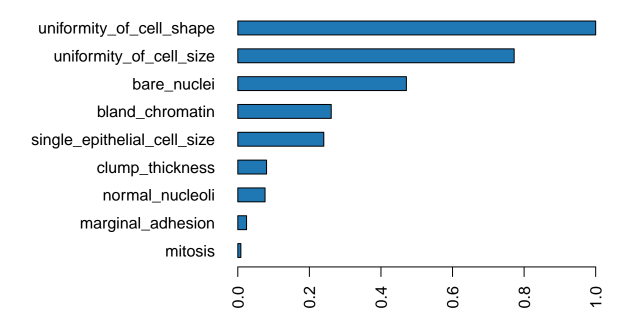
```
max min_per_class_accuracy 0.293126 0.985915 34
## 10 max mean_per_class_accuracy 0.293126 0.992958 34
## Gains/Lift Table: Extract with `h2o.gainsLift(<model>, <data>)` or `h2o.gainsLift(<model>, valid=<T/
## H20BinomialMetrics: drf
## ** Reported on cross-validation data. **
## ** 25-fold cross-validation on training data (Metrics computed for combined holdout predictions) **
## MSE: 0.03141201
## RMSE: 0.1772343
## LogLoss: 0.1163767
## Mean Per-Class Error: 0.02309728
## AUC: 0.9890496
## Gini: 0.9780993
##
## Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
##
            benign malignant
                                Error
                                          Rate
                306
                          11 0.034700
                                       =11/317
## benign
                          172 0.011494
                                        =2/174
## malignant
                 2
## Totals
                308
                         183 0.026477 =13/491
## Maximum Metrics: Maximum metrics at their respective thresholds
##
                          metric threshold
                                              value idx
## 1
                          max f1 0.458037 0.963585 178
## 2
                          max f2 0.458037 0.978385 178
                    max f0point5 0.514579 0.952116 176
## 4
                    max accuracy 0.514579 0.973523 176
                   max precision 1.000000 1.000000
## 6
                      max recall 0.182569 1.000000 196
## 7
                 max specificity 1.000000 1.000000
                max absolute_mcc 0.458037 0.943546 178
## 8
      max min_per_class_accuracy 0.530369 0.968454 174
## 10 max mean_per_class_accuracy   0.458037   0.976903   178
## Gains/Lift Table: Extract with `h2o.gainsLift(<model>, <data>)` or `h2o.gainsLift(<model>, valid=<T/
## Cross-Validation Metrics Summary:
                                 mean
                                                sd cv_1_valid cv_2_valid
                           0.97824687 0.022163384
## accuracy
                                                           1.0
                                                                       1.0
## auc
                           0.98896265 0.0137583455
                                                           1.0
                                                                       1.0
## err
                          0.021753136 0.022163384
                                                           0.0
                                                                       0.0
## err count
                                                           0.0
                                  0.4
                                               0.4
## f0point5
                           0.94340783 0.060686026
                                                           1.0
                                                                       1.0
## f1
                           0.96193475
                                       0.04148564
                                                           1.0
                                                                       1.0
## f2
                            0.9834832
                                       0.01836118
                                                           1.0
                                                                       1.0
## lift_top_group
                             3.146508
                                       0.86633277
                                                           3.0
                                                                2.6666667
                           0.12211874
                                       0.06134508 0.051629476 0.09666276
## logloss
## max_per_class_error
                          0.031400856 0.032273438
                                                           0.0
                                                                       0.0
                            0.9496154 0.051578306
                                                           1.0
                                                                       1.0
## mean_per_class_accuracy
                            0.9842996 0.016136719
                                                           1.0
                                                                       1.0
## mean_per_class_error
                          0.0
                                                                       0.0
                                       0.02053207 0.008940321 0.022683756
## mse
                          0.033157483
## precision
                            0.9323954
                                       0.07182843
                                                           1.0
## r2
                            0.8243174
                                         0.1447473 0.95976853 0.90321594
## recall
                                  1.0
                                               0.0
                                                           1.0
```

```
## rmse
## specificity
                            0.96859914 0.032273438
                                                            1.0
                                                                         1.0
                            cv 3 valid
##
                                         cv_4_valid cv_5_valid
                                                                cv 6 valid
## accuracy
                            0.95454544
                                                1.0
                                                            1.0
                                                                  0.9285714
## auc
                             0.9764706
                                                1.0
                                                            1.0
                                                                  0.9583333
## err
                           0.045454547
                                                0.0
                                                            0.0 0.071428575
                                                0.0
                                                            0.0
## err count
                                   1.0
## f0point5
                            0.86206895
                                                1.0
                                                            1.0 0.71428573
## f1
                            0.90909094
                                                1.0
                                                            1.0
                                                                         0.8
## f2
                                                                 0.90909094
                            0.96153843
                                                1.0
                                                            1.0
## lift_top_group
                                   4.4
                                                2.5
                                                      2.777777
                                                                        7.0
                                       0.032063406 0.07835095
                            0.14851098
                                                                0.38401476
## logloss
## max_per_class_error
                            0.05882353
                                                0.0
                                                            0.0 0.083333336
## mcc
                             0.8856149
                                                1.0
                                                            1.0 0.78173596
## mean_per_class_accuracy
                             0.9705882
                                                            1.0
                                                                  0.9583333
                                                1.0
## mean_per_class_error
                           0.029411765
                                                0.0
                                                            0.0 0.04166668
                           0.048121125 0.0026525913 0.017376833 0.12076553
## mse
## precision
                             0.8333333
                                                1.0
                                                            1.0
## r2
                             0.7259927
                                          0.9889475
                                                    0.92457974 0.013748176
## recall
                                   1.0
                                                1.0
                                                            1.0
## rmse
                            0.21936527
                                        0.051503316  0.13182122  0.34751335
## specificity
                             0.9411765
                                                                  0.9166667
                                       cv_8_valid cv_9_valid cv_10_valid
                            cv_7_valid
##
                            0.94736844
                                         0.9411765
                                                     0.888889
## accuracy
                                                                       1.0
## auc
                                         0.9848485
                                                     0.9230769
                                                                       1.0
                            0.98571426
## err
                            0.05263158
                                       0.05882353
                                                    0.11111111
                                                                       0.0
## err_count
                                   1.0
                                               1.0
                                                           2.0
                                                                       0.0
                            0.86206895
                                        0.88235295
## f0point5
                                                    0.75757575
                                                                       1.0
## f1
                            0.90909094
                                         0.9230769
                                                     0.8333333
                                                                       1.0
## f2
                            0.96153843
                                         0.9677419
                                                     0.9259259
                                                                       1.0
## lift_top_group
                                   3.8
                                         2.8333333
                                                           3.6
                                                                 1.7142857
## logloss
                            0.12703204
                                         0.1548704
                                                    0.36086315
                                                                0.10401911
## max_per_class_error
                           0.071428575
                                       0.09090909
                                                    0.15384616
                                                                       0.0
                             0.8796644 0.88273484
                                                    0.77742887
                                                                       1.0
## mcc
## mean_per_class_accuracy 0.96428573 0.95454544
                                                     0.9230769
                                                                       1.0
                           0.035714287 0.045454547
## mean_per_class_error
                                                    0.07692308
                                                                       0.0
## mse
                            0.04287585 0.051242344 0.104545906 0.019419663
## precision
                             0.8333333 0.85714287
                                                    0.71428573
## r2
                            0.77888316 0.77562064
                                                    0.47887886
                                                                0.92010194
## recall
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## rmse
                            0.20706484 0.22636773
                                                     0.3233356
                                                                0.13935445
## specificity
                             0.9285714 0.90909094 0.84615386
                                                                       1.0
                           cv 11 valid cv 12 valid cv 13 valid
                                                                cv 14 valid
                                         0.944444
                                                     0.9411765
## accuracy
                                  0.96
                                                                        1.0
                            0.99264705
                                            0.9625
                                                    0.95238096
## auc
                                                                        1.0
                                                    0.05882353
                                                                        0.0
                                  0.04 0.05555556
## err
## err_count
                                   1.0
                                               1.0
                                                           1.0
                                                                        0.0
                            0.90909094
                                         0.9259259
## f0point5
                                                     0.7894737
                                                                         1.0
## f1
                             0.9411765
                                       0.95238096
                                                    0.85714287
                                                                        1.0
## f2
                             0.9756098
                                        0.98039216
                                                        0.9375
                                                                         1.0
                                                     5.666665
                                                                      2,125
## lift_top_group
                                 3.125
                                               1.8
                                        0.21119072 0.18838081
                                                                0.066318005
## logloss
                            0.13761193
## max_per_class_error
                            0.05882353
                                             0.125 0.071428575
                                                                        0.0
                                         0.8918826 0.83452296
## mcc
                            0.91465914
                                                                         1.0
```

```
0.9375 0.96428573
                                                                        1.0
## mean_per_class_accuracy
                             0.9705882
## mean_per_class_error
                           0.029411765
                                            0.0625 0.035714287
                                                                        0.0
## mse
                           0.8888889 0.90909094
## precision
                                                          0.75
                                                                        1.0
## r2
                             0.8193236
                                          0.737646
                                                     0.6090993
                                                                  0.9610937
## recall
                                   1.0
                                               1.0
                                                           1.0
                                                                        1.0
## rmse
                            0.19828056 0.25451672 0.23834658
                             0.9411765
## specificity
                                             0.875
                                                     0.9285714
##
                           cv_15_valid cv_16_valid cv_17_valid cv_18_valid
## accuracy
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## auc
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
## err
                                   0.0
                                               0.0
                                                           0.0
## err_count
                                                                       0.0
## f0point5
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## f1
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## f2
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## lift_top_group
                             3.5714285
                                         3.8333333
                                                           3.5
                                                                      2.25
                            0.10126504 0.050743338 0.08231668 0.06105531
## logloss
## max_per_class_error
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
                                               1.0
                                                           1.0
                                                                       1.0
## mcc
                                   1.0
## mean_per_class_accuracy
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## mean_per_class_error
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
                           0.030016925 0.013950677 0.015176254 0.010579813
## mse
## precision
                                               1.0
                                                           1.0
## r2
                             0.8511065 0.92764795 0.92563635
                                                                 0.9571518
## recall
                                   1.0
                                               1.0
                                                           1.0
## rmse
                            0.17325394 0.11811298
                                                   0.12319194 0.102858216
## specificity
                                   1.0
                                               1.0
                                                           1.0
                           cv_19_valid cv_20_valid cv_21_valid cv_22_valid
## accuracy
                                   1.0
                                               1.0
                                                           1.0
## auc
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## err
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
## err_count
## f0point5
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## f1
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## f2
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## lift_top_group
                                                      2.142857
                             1.8333334
                                         2.1111112
                                                                 2.4285715
## logloss
                            0.12709582 0.041414153 0.11962002 0.102114245
## max_per_class_error
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
                                               1.0
                                                           1.0
                                   1.0
                                                                       1.0
## mean per class accuracy
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## mean_per_class_error
                                   0.0
                                               0.0
                                                           0.0
                                                                       0.0
                           0.026209245 0.007021444
                                                   0.03179515
## mse
                                                                0.02507944
## precision
                                   1.0
                                               1.0
                                                           1.0
                                                                       1.0
## r2
                             0.8942894
                                         0.9718362
                                                    0.87225163
                                                                0.89645773
## recall
                                   1.0
                                               1.0
                                                           1.0
                             0.1618927 0.08379406 0.17831194
                                                                0.15836489
## rmse
## specificity
                                               1.0
                                                                       1.0
                                   1.0
                                                           1.0
                           cv_23_valid cv_24_valid cv_25_valid
## accuracy
                                   1.0
                                               1.0
                                                          0.95
## auc
                                   1.0
                                               1.0
                                                     0.9880952
## err
                                   0.0
                                               0.0
                                                          0.05
## err count
                                   0.0
                                               0.0
                                                           1.0
## f0point5
                                               1.0 0.88235295
                                   1.0
```

```
## f1
                                    1.0
                                                 1.0
                                                       0.9230769
                                                 1.0
## f2
                                    1.0
                                                       0.9677419
## lift_top_group
                                   2.25
                                                 4.4
                                                       3.3333333
                             0.05492981 0.03994816 0.13094744
## logloss
## max_per_class_error
                                    0.0
                                                0.0 0.071428575
                                    1.0
## mcc
                                                 1.0
                                                       0.8921426
## mean_per_class_accuracy
                                    1.0
                                                1.0 0.96428573
## mean_per_class_error
                                    0.0
                                                0.0 0.035714287
## mse
                            0.008537599 0.006692548
                                                      0.04465809
## precision
                                    1.0
                                                 1.0
                                                      0.85714287
## r2
                             0.96542275
                                         0.96189183
                                                       0.7873424
## recall
                                    1.0
                                                 1.0
                                                             1.0
                                         0.08180799
## rmse
                             0.09239913
                                                      0.21132462
## specificity
                                                       0.9285714
                                    1.0
                                                 1.0
h2o.varimp_plot(rf_model)
```

# Variable Importance: DRF



# #h2o.varimp(rf\_model)

One performance metric we are interested in is the mean per class error for training and validation data.

```
h2o.mean_per_class_error(rf_model, train = TRUE, valid = TRUE, xval = TRUE)
```

```
## train valid xval
## 0.024674571 0.007042254 0.023097284
```

The confusion matrix tells us, how many classes have been predicted correctly and how many predictions were accurate. Here, we see the errors in predictions on validation data

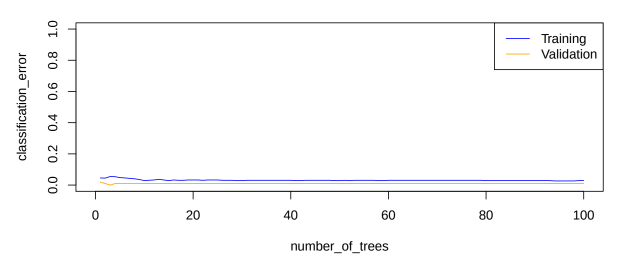
```
h2o.confusionMatrix(rf_model, valid = TRUE)
```

```
## Confusion Matrix (vertical: actual; across: predicted) for max f1 @ threshold = 0.293125896751881:
##
             benign malignant
                                           Rate
                                  Error
## benign
                 70
                             1 0.014085
                                          =1/71
## malignant
                  0
                           35 0.000000
                                          =0/35
## Totals
                 70
                           36 0.009434
                                         =1/106
```

We can also plot the classification error.

```
plot(rf_model,
    timestep = "number_of_trees",
    metric = "classification_error")
```

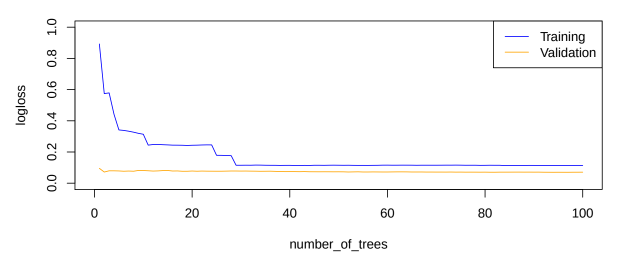
# **Scoring History**



Next to the classification error, we are usually interested in the logistic loss (negative log-likelihood or log loss). It describes the sum of errors for each sample in the training or validation data or the negative logarithm of the likelihood of error for a given prediction/ classification. Simply put, the lower the loss, the better the model (if we ignore potential overfitting).

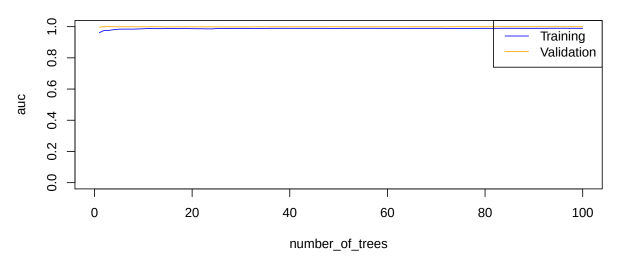
```
plot(rf_model,
    timestep = "number_of_trees",
    metric = "logloss")
```

# **Scoring History**



```
plot(rf_model,
    timestep = "number_of_trees",
    metric = "AUC")
```

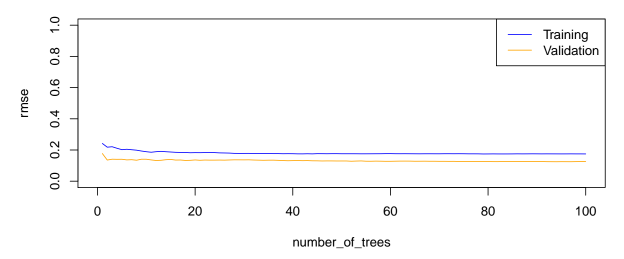
# **Scoring History**



We can also plot the mean squared error (MSE). The MSE tells us the average of the prediction errors squared, i.e. the estimator's variance and bias. The closer to zero, the better a model.

```
plot(rf_model,
    timestep = "number_of_trees",
    metric = "rmse")
```

# **Scoring History**



Next, we want to know the area under the curve (AUC). AUC is an important metric for measuring binary classification model performances. It gives the area under the curve, i.e. the integral, of true positive vs false positive rates. The closer to 1, the better a model. AUC is especially useful, when we have unbalanced datasets (meaning datasets where one class is much more common than the other), because it is independent of class labels.

```
h2o.auc(rf_model, train = TRUE)

## [1] 0.989521
h2o.auc(rf_model, valid = TRUE)

## [1] 0.9995976
h2o.auc(rf_model, xval = TRUE)
```

## [1] 0.9890496

Now that we have a good idea about model performance on validation data, we want to know how it performed on unseen test data. A good model should find an optimal balance between accuracy on training and test data. A model that has 0% error on the training data but 40% error on the test data is in effect useless. It overfit on the training data and is thus not able to generalize to unknown data.

```
perf <- h2o.performance(rf_model, test)
perf

## H20BinomialMetrics: drf
##
## MSE: 0.03673598
## RMSE: 0.1916663
## LogLoss: 0.1158835</pre>
```

```
## AUC: 0.990625
## Gini: 0.98125
##
```

## Mean Per-Class Error:

## Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
## benign malignant Error Rate

0.0625

```
## Totals
                 74
                            28 0.039216
                                         =4/102
##
## Maximum Metrics: Maximum metrics at their respective thresholds
##
                            metric threshold
                                                value idx
## 1
                                   0.735027 0.933333
                            max f1
## 2
                            max f2
                                   0.294222 0.952381
## 3
                     max f0point5
                                   0.735027 0.972222
                                                        25
                                                        25
##
                     max accuracy
                                   0.735027 0.960784
##
                    max precision
                                   1.000000 1.000000
                                                         0
                       max recall
                                   0.294222 1.000000
                                                       37
##
##
                  max specificity
                                   1.000000 1.000000
                 max absolute_mcc 0.735027 0.909782
## 8
                                                       25
## 9
       max min_per_class_accuracy
                                   0.424524 0.937500
                                                        31
## 10 max mean_per_class_accuracy
                                    0.294222 0.942857
                                                        37
##
## Gains/Lift Table: Extract with `h2o.gainsLift(<model>, <data>)` or `h2o.gainsLift(<model>, valid=<T/
Plotting the test performance's AUC plot shows us approximately how good the predictions are.
```

70

4

0 0.000000

28 0.125000

=0/70

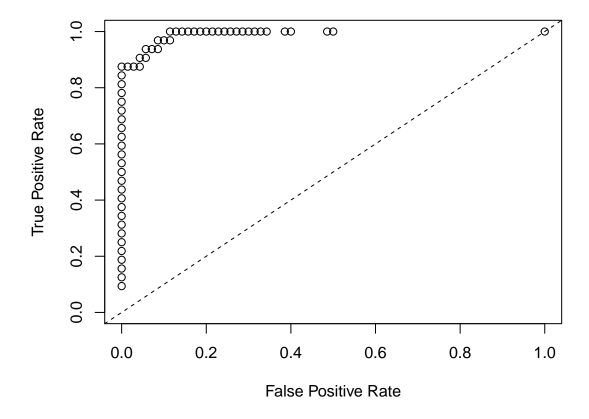
=4/32

## benign

plot(perf)

## malignant

# **True Positive Rate vs False Positive Rate**



We also want to know the log loss, MSE and AUC values, as well as other model metrics for the test data:

```
h2o.logloss(perf)
## [1] 0.1158835
h2o.mse(perf)
## [1] 0.03673598
h2o.auc(perf)
## [1] 0.990625
head(h2o.metric(perf))
## Metrics for Thresholds: Binomial metrics as a function of classification thresholds
     threshold
                     f1
                              f2 f0point5 accuracy precision
## 1 1.000000 0.171429 0.114504 0.340909 0.715686 1.000000 0.093750
## 2 0.998333 0.222222 0.151515 0.416667 0.725490 1.000000 0.125000
## 3 0.998000 0.270270 0.187970 0.480769 0.735294 1.000000 0.156250
## 4 0.997222 0.315789 0.223881 0.535714 0.745098 1.000000 0.187500
## 5 0.996210 0.358974 0.259259 0.583333 0.754902 1.000000 0.218750
## 6 0.994048 0.400000 0.294118 0.625000 0.764706 1.000000 0.250000
     specificity absolute_mcc min_per_class_accuracy mean_per_class_accuracy
## 1
        1.000000
                     0.257464
                                            0.093750
                                                                    0.546875
## 2
        1.000000
                     0.298807
                                            0.125000
                                                                    0.562500
## 3
        1.000000
                                            0.156250
                                                                    0.578125
                     0.335794
## 4
        1.000000
                     0.369755
                                            0.187500
                                                                    0.593750
## 5
        1.000000
                     0.401478
                                            0.218750
                                                                    0.609375
## 6
        1.000000
                     0.431474
                                            0.250000
                                                                     0.625000
##
    tns fns fps tps
                          tnr
                                   fnr
                                            fpr
                                                     tpr idx
     70
         29
                  3 1.000000 0.906250 0.000000 0.093750
## 1
              0
                   4 1.000000 0.875000 0.000000 0.125000
## 2
     70
         28
              0
## 3
     70
         27
              Ω
                   5 1.000000 0.843750 0.000000 0.156250
    70 26
              0 6 1.000000 0.812500 0.000000 0.187500
## 4
    70 25
              0 7 1.000000 0.781250 0.000000 0.218750
## 5
                   8 1.000000 0.750000 0.000000 0.250000
## 6 70 24
Deep learning with neural networks
hyper_params <- list(</pre>
                     activation = c("Rectifier", "Maxout", "Tanh", "RectifierWithDropout",
                                    "MaxoutWithDropout", "TanhWithDropout"),
                     hidden = list(c(5, 5, 5, 5, 5), c(10, 10, 10, 10), c(50, 50, 50)),
                     epochs = c(50, 100, 200),
                     11 = c(0, 0.00001, 0.0001),
                     12 = c(0, 0.00001, 0.0001),
                     rate = c(0, 01, 0.005, 0.001),
                     rate_annealing = c(1e-8, 1e-7, 1e-6),
                     rho = c(0.9, 0.95, 0.99, 0.999),
```

epsilon = c(1e-10,1e-8,1e-6,1e-4),
input\_dropout\_ratio = c(0, 0.1, 0.2),
max\_w2 = c(10, 100, 1000, 3.4028235e+38)

```
y = response,
grid_id = "dl_grid",
training_frame = train,
validation_frame = valid,
nfolds = 25,
fold_assignment = "Stratified",
hyper_params = hyper_params,
search_criteria = search_criteria,
seed = 42
)
grid <- h2o.getGrid("dl_grid", sort_by = "auc", decreasing = TRUE)</pre>
```

Because training can take a while, depending on how many samples, features, nodes and hidden layers you are training on, it is a good idea to save your model.

model\_ids <- grid@model\_ids</pre>

best\_model <- h2o.getModel(model\_ids[[1]])</pre>

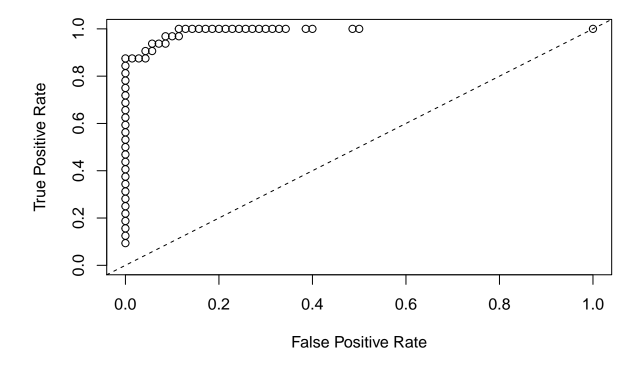
```
h2o.saveModel(best_model, path="models", force = TRUE)
```

We can then re-load the model again any time to check the model quality and make predictions on new data.

```
dl_model <- h2o.loadModel("U:\\Github_blog\\Webinar\\Webinar_ML_for_disease\\models\\dl_grid_model_8")</pre>
```

```
perf <- h2o.performance(best_model, test)
plot(perf)</pre>
```

# **True Positive Rate vs False Positive Rate**



```
h2o.confusionMatrix(best_model, test)
## Confusion Matrix (vertical: actual; across: predicted) for max f1 @ threshold = 0.735026690140367:
##
             benign malignant
                                           Rate
                                  Error
## benign
                 70
                            0 0.000000
                                          =0/70
                                          =4/32
## malignant
                  4
                           28 0.125000
## Totals
                 74
                           28 0.039216
                                        =4/102
```

#### Exercises

Try to run the analyses on the following datasets:

#### Arrhythmia data

The arrhythmia dataset from the UC Irvine Machine Learning repository contains 279 features from ECG heart rhythm diagnostics and one output column. I am not going to rename the feature columns because they are too many and the descriptions are too complex. Also, we don't need to know specifically which features we are looking at for building the models. For a description of each feature, see https://archive.ics.uci.edu/ml/machine-learning-databases/arrhythmia/arrhythmia.names. The output column defines 16 classes: class 1 samples are from healthy ECGs, the remaining classes belong to different types of arrhythmia, with class 16 being all remaining arrhythmia cases that didn't fit into distinct classes.

```
arrhythmia <- read.table("datasets/arrhythmia.data.txt", sep = ",")
arrhythmia[arrhythmia == "?"] <- NA

# making sure, that all feature columns are numeric
arrhythmia[-280] <- lapply(arrhythmia[-280], as.character)
arrhythmia[-280] <- lapply(arrhythmia[-280], as.numeric)

# renaming output column and converting to factor
colnames(arrhythmia)[280] <- "class"
arrhythmia$class <- as.factor(arrhythmia$class)

arrhythmia$diagnosis <- ifelse(arrhythmia$class == 1, "healthy", "arrhythmia")
arrhythmia$diagnosis <- as.factor(arrhythmia$diagnosis)</pre>
```

#### Flu data

Among the many R packages, there is the outbreaks package. It contains datasets on epidemics, on of which is from the 2013 outbreak of influenza A H7N9 in China, as analysed by Kucharski et al. (2014):

```
library(outbreaks)
data(fluH7N9_china_2013)

# convert ? to NAs
fluH7N9_china_2013$age[which(fluH7N9_china_2013$age == "?")] <- NA

# create a new column with case ID
fluH7N9_china_2013$case.ID <- paste("case", fluH7N9_china_2013$case.ID, sep = "_")

# preparing the data frame for modeling</pre>
```

```
library(dplyr)
dataset <- fluH7N9 china 2013 %>%
  mutate(hospital = as.factor(ifelse(is.na(date_of_hospitalisation), 0, 1)),
         gender_f = as.factor(ifelse(gender == "f", 1, 0)),
         province_Jiangsu = as.factor(ifelse(province == "Jiangsu", 1, 0)),
         province_Shanghai = as.factor(ifelse(province == "Shanghai", 1, 0)),
         province_Zhejiang = as.factor(ifelse(province == "Zhejiang", 1, 0)),
         province_other = as.factor(ifelse(province == "Zhejiang" | province == "Jiangsu" | province ==
         days_onset_to_outcome = as.numeric(as.character(gsub(" days", "",
                                       as.Date(as.character(date_of_outcome), format = "%Y-%m-%d") -
                                         as.Date(as.character(date_of_onset), format = "%Y-%m-%d")))),
         days_onset_to_hospital = as.numeric(as.character(gsub(" days", "",
                                       as.Date(as.character(date_of_hospitalisation), format = "%Y-%m-%d
                                         as.Date(as.character(date_of_onset), format = "%Y-\m-\d")))),
         age = as.numeric(as.character(age)),
         early_onset = as.factor(ifelse(date_of_onset < summary(fluH7N9_china_2013$date_of_onset)[[3]],</pre>
         early_outcome = as.factor(ifelse(date_of_outcome < summary(fluH7N9_china_2013$date_of_outcome)</pre>
  subset(select = -c(2:4, 6, 8))
rownames(dataset) <- dataset$case id</pre>
dataset <- dataset[, -1]</pre>
# impute missing data
library(mice)
##
## Attaching package: 'mice'
## The following object is masked from 'package:tidyr':
##
##
       complete
dataset_impute <- mice(dataset[, -1], print = FALSE)</pre>
# recombine imputed data frame with the outcome column
dataset_complete <- merge(dataset[, 1, drop = FALSE], mice::complete(dataset_impute, 1), by = "row.name
rownames(dataset_complete) <- dataset_complete$Row.names</pre>
dataset_complete <- dataset_complete[, -1]</pre>
sessionInfo()
## R version 3.3.2 (2016-10-31)
## Platform: x86_64-w64-mingw32/x64 (64-bit)
## Running under: Windows 7 x64 (build 7601) Service Pack 1
## locale:
## [1] LC_COLLATE=English_United States.1252
## [2] LC CTYPE=English United States.1252
## [3] LC_MONETARY=English_United States.1252
## [4] LC_NUMERIC=C
## [5] LC_TIME=English_United States.1252
```

```
##
## attached base packages:
## [1] stats
                 graphics grDevices utils
                                               datasets methods
                                                                    base
##
## other attached packages:
  [1] mice_2.30
                            outbreaks_1.2.0
                                                h2o_3.10.3.6
##
   [4] plyr_1.8.4
                            xgboost 0.6-4
                                                randomForest_4.6-12
                            rpart_4.1-10
                                                tidyr_0.6.1
## [7] rpart.plot_2.1.1
## [10] dplyr_0.5.0
                            caret_6.0-73
                                                ggplot2_2.2.1.9000
## [13] lattice_0.20-34
## loaded via a namespace (and not attached):
  [1] reshape2_1.4.2
                           splines_3.3.2
                                              colorspace_1.3-2
  [4] htmltools_0.3.5
                           stats4_3.3.2
                                              yaml_2.1.14
## [7] mgcv_1.8-17
                           survival_2.40-1
                                              ModelMetrics_1.1.0
## [10] e1071_1.6-8
                           nloptr_1.0.4
                                              DBI_0.5-1
## [13] RColorBrewer_1.1-2 foreach_1.4.3
                                              stringr_1.2.0
## [16] MatrixModels 0.4-1 munsell 0.4.3
                                              gtable 0.2.0
## [19] codetools_0.2-15
                           evaluate_0.10
                                              labeling_0.3
## [22] knitr_1.15.1
                           SparseM_1.74
                                              quantreg_5.29
## [25] pbkrtest_0.4-6
                           parallel_3.3.2
                                              class_7.3-14
## [28] Rcpp_0.12.9
                           scales_0.4.1
                                              backports_1.0.5
## [31] jsonlite_1.3
                           lme4_1.1-12
                                              digest_0.6.12
## [34] stringi_1.1.2
                           grid_3.3.2
                                              rprojroot_1.2
## [37] tools_3.3.2
                           bitops_1.0-6
                                              magrittr_1.5
## [40] lazyeval_0.2.0
                           RCurl_1.95-4.8
                                              tibble_1.2
## [43] car_2.1-4
                           MASS_7.3-45
                                              Matrix_1.2-8
## [46] data.table_1.10.4
                           assertthat_0.1
                                              minqa_1.2.4
## [49] rmarkdown_1.3
                           iterators_1.0.8
                                              R6_2.2.0
## [52] nnet_7.3-12
                           nlme_3.1-131
```