Building meaningful machine learning models for disease prediction

Dr. Shirin Glander March 31, 2017

Webinar for the ISDS R Group

This document presents the code used to produce the example analysis and figures shown in my webinar on building meaningful machine learning models for disease prediction.

My webinar slides are available on Github

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.

Setup

All analyses are done in R using RStudio. For detailed session information including R version, operating system and package versions, see the sessionInfo() output at the end of this document.

All figures are produced with ggplot2.

The dataset

The dataset I am using in these example analyses, is the **Breast Cancer Wisconsin (Diagnostic) Dataset**. The data was downloaded from the UC Irvine Machine Learning Repository.

The first dataset looks at the predictor classes:

- malignant or
- benign breast mass.

The features characterise cell nucleus properties and were generated from image analysis of fine needle aspirates (FNA) of breast masses:

- Sample ID (code number)
- Clump thickness
- Uniformity of cell size
- Uniformity of cell shape
- $\bullet \quad {\rm Marginal \ adhesion}$
- Single epithelial cell size
- Number of bare nuclei
- Bland chromatin
- Number of normal nuclei
- Mitosis
- Classes, i.e. diagnosis

```
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))
```

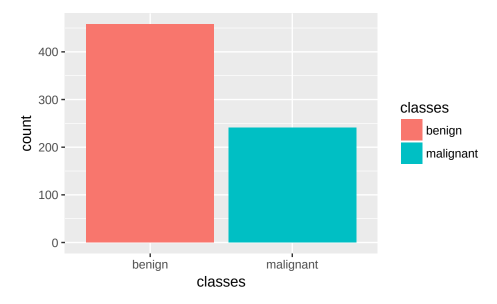
Missing data

```
bc_data[bc_data == "?"] <- NA</pre>
# how many NAs are in the data
length(which(is.na(bc_data)))
## [1] 16
# how many samples would we loose, if we removed them?
nrow(bc_data)
## [1] 699
nrow(bc_data[is.na(bc_data), ])
## [1] 16
Missing values are imputed with the mice package.
# impute missing data
library(mice)
bc_data[,2:10] <- apply(bc_data[, 2:10], 2, function(x) as.numeric(as.character(x)))</pre>
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)
```

Data exploration

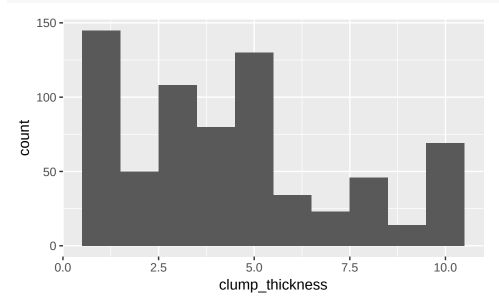
• Response variable for classification

```
library(ggplot2)
ggplot(bc_data, aes(x = classes, fill = classes)) +
  geom_bar()
```



• Response variable for regression

```
ggplot(bc_data, aes(x = clump_thickness)) +
  geom_histogram(bins = 10)
```



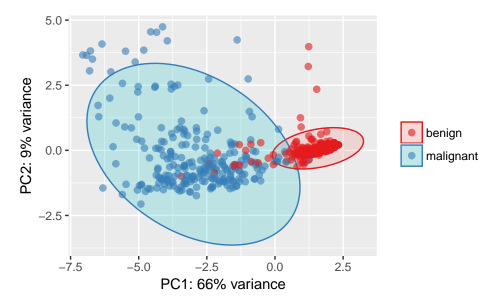
• Principal Component Analysis

```
library(pcaGoPromoter)
library(ellipse)

# perform pca and extract scores
pcaOutput <- pca(t(bc_data[, -1]), printDropped = FALSE, scale = TRUE, center = TRUE)
pcaOutput2 <- as.data.frame(pcaOutput$scores)

# define groups for plotting
pcaOutput2$groups <- bc_data$classes

centroids <- aggregate(cbind(PC1, PC2) ~ groups, pcaOutput2, mean)</pre>
```

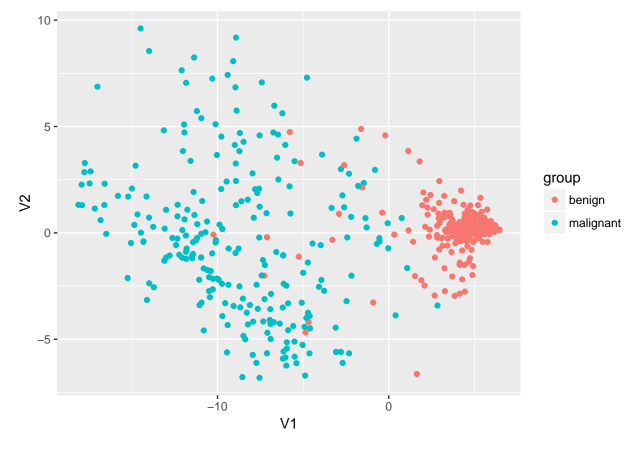


• Multidimensional Scaling

library(dplyr)

```
##
## Attaching package: 'dplyr'
##
  The following objects are masked from 'package:Biostrings':
##
##
       collapse, intersect, setdiff, setequal, union
## The following object is masked from 'package:XVector':
##
##
       slice
## The following objects are masked from 'package: IRanges':
##
       collapse, desc, intersect, regroup, setdiff, slice, union
##
## The following objects are masked from 'package:S4Vectors':
```

```
first, intersect, rename, setdiff, setequal, union
##
## The following objects are masked from 'package:BiocGenerics':
##
       combine, intersect, setdiff, union
##
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
       intersect, setdiff, setequal, union
##
select(bc_data, -1) %>%
  dist() %>%
  cmdscale %>%
  as.data.frame() %>%
  mutate(group = bc_data$classes) %>%
  ggplot(aes(x = V1, y = V2, color = group)) +
    geom_point()
```



• t-SNE dimensionality reduction

```
library(tsne)
select(bc_data, -1) %>%
  dist() %>%
  tsne() %>%
```

```
as.data.frame() %>%
mutate(group = bc_data$classes) %>%
ggplot(aes(x = V1, y = V2, color = group)) +
   geom_point()
```

```
## sigma summary: Min. : 0.2945 | 1st Qu. : 0.5325 | Median : 0.5978 | Mean : 0.7045 | 3rd Qu. : 0.9128 | Ma.

## Epoch: Iteration #100 error is: 12.9051118607517

## Epoch: Iteration #200 error is: 0.55100578564023

## Epoch: Iteration #300 error is: 0.507964548605455

## Epoch: Iteration #400 error is: 0.497858645047617

## Epoch: Iteration #500 error is: 0.494929061688897

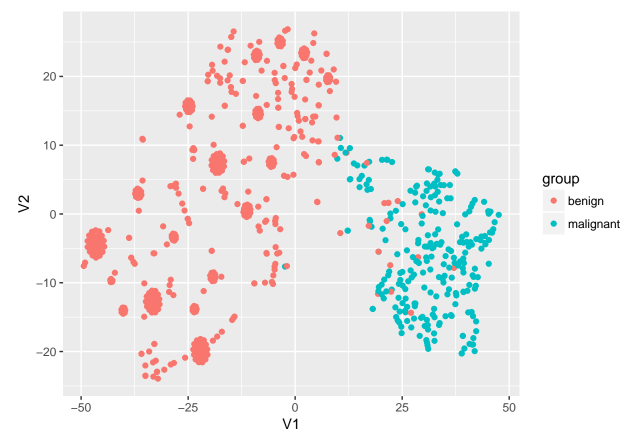
## Epoch: Iteration #600 error is: 0.493629390821366

## Epoch: Iteration #700 error is: 0.492881739116146

## Epoch: Iteration #800 error is: 0.492432292261577

## Epoch: Iteration #900 error is: 0.492109439434399

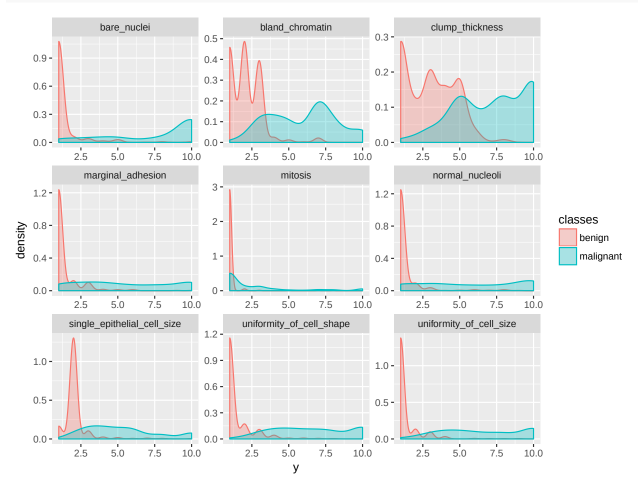
## Epoch: Iteration #1000 error is: 0.491872988560014
```



• Features

```
library(tidyr)
gather(bc_data, x, y, clump_thickness:mitosis) %>%
```

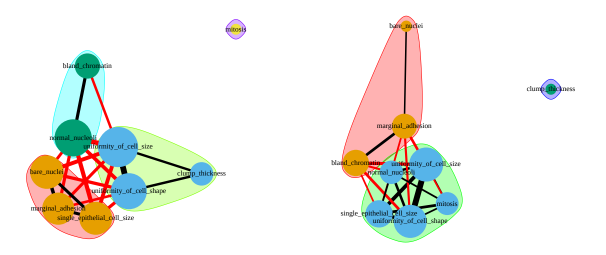
```
ggplot(aes(x = y, color = classes, fill = classes)) +
geom_density(alpha = 0.3) +
facet_wrap( ~ x, scales = "free", ncol = 3)
```



• Correlation graphs

```
mode = "upper")
# http://kateto.net/networks-r-igraph
cut.off_b <- mean(E(g_benign)$weight)</pre>
cut.off_m <- mean(E(g_malignant)$weight)</pre>
g_benign_2 <- delete_edges(g_benign, E(g_benign)[weight < cut.off_b])</pre>
g_malignant_2 <- delete_edges(g_malignant, E(g_malignant)[weight < cut.off_m])</pre>
c_g_benign_2 <- cluster_fast_greedy(g_benign_2)</pre>
c_g_malignant_2 <- cluster_fast_greedy(g_malignant_2)</pre>
par(mfrow=c(1,2))
plot(c_g_benign_2, g_benign_2,
     vertex.size = colSums(co_mat_benign) * 10,
     vertex.frame.color = NA,
     vertex.label.color = "black",
     vertex.label.cex = 0.8,
     edge.width = E(g_benign_2)$weight * 15,
     layout = layout_with_fr(g_benign_2),
     main = "Benign tumors")
plot(c_g_malignant_2, g_malignant_2,
     vertex.size = colSums(co_mat_malignant) * 10,
     vertex.frame.color = NA,
     vertex.label.color = "black",
     vertex.label.cex = 0.8,
     edge.width = E(g_malignant_2)$weight * 15,
     layout = layout_with_fr(g_malignant_2),
     main = "Malignant tumors")
```

Benign tumors Malignant tumors

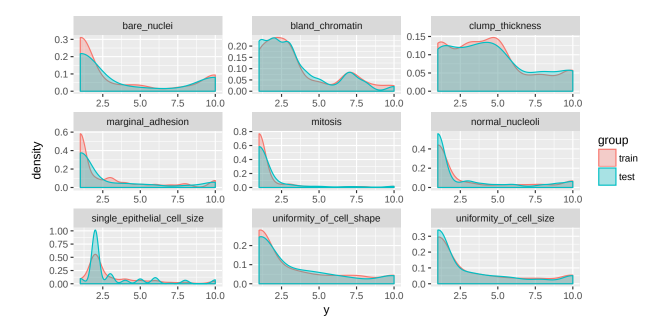


Machine Learning packages for R

caret

```
library(caret)
```

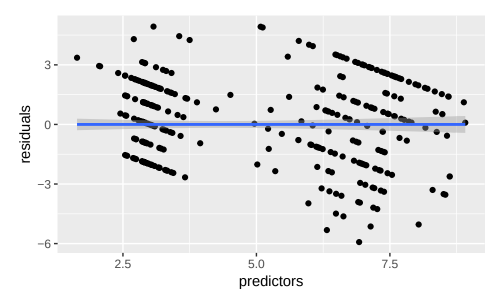
Training, validation and test data

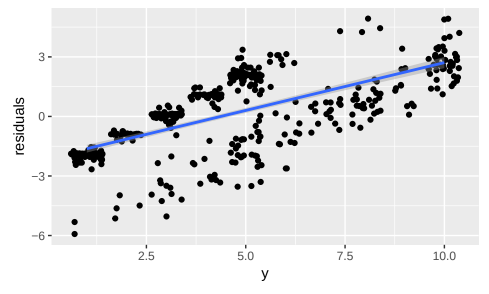


Regression

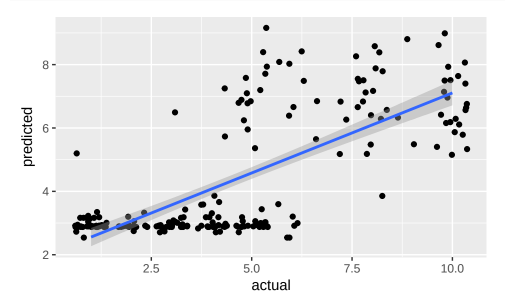
```
set.seed(42)
model_glm <- caret::train(clump_thickness ~ .,</pre>
                            data = train_data,
                            method = "glm",
                            preProcess = c("scale", "center"),
                            trControl = trainControl(method = "repeatedcv",
                                                     number = 10,
                                                      repeats = 10,
                                                     savePredictions = TRUE,
                                                      verboseIter = FALSE))
model_glm
## Generalized Linear Model
##
## 490 samples
     9 predictor
##
##
## Pre-processing: scaled (9), centered (9)
  Resampling: Cross-Validated (10 fold, repeated 10 times)
## Summary of sample sizes: 441, 441, 440, 442, 441, 440, ...
## Resampling results:
##
##
     RMSE
                Rsquared
##
     1.974296
               0.5016141
##
##
predictions <- predict(model_glm, test_data)</pre>
{\it\# model\_glm\$finalModel\$linear.predictors} \; == \; {\it model\_glm\$finalModel\$fitted.values}
data.frame(residuals = resid(model_glm),
```

```
predictors = model_glm$finalModel$linear.predictors) %>%
ggplot(aes(x = predictors, y = residuals)) +
  geom_jitter() +
  geom_smooth(method = "lm")
```





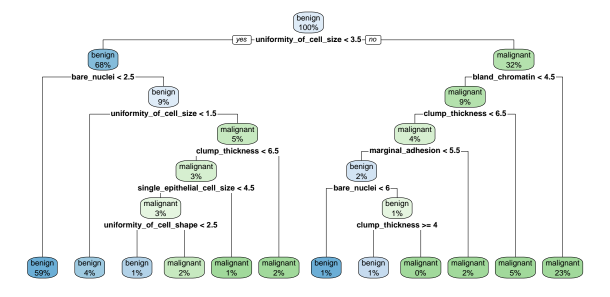




Classification

Decision trees

rpart



Random Forests

Random Forests predictions are based on the generation of multiple classification trees. They can be used for both, 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. model_rf\$finalModel\$confusion

```
## benign malignant class.error
## benign 313 8 0.02492212
## malignant 4 165 0.02366864
```

• Feature Importance

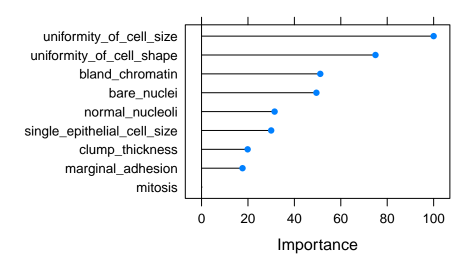
```
imp <- model_rf$finalModel$importance
imp[order(imp, decreasing = TRUE), ]</pre>
```

```
##
       uniformity_of_cell_size
                                    uniformity_of_cell_shape
##
                      54.416003
                                                    41.553022
##
               bland_chromatin
                                                 bare_nuclei
##
                      29.343027
                                                    28.483842
##
               normal_nucleoli single_epithelial_cell_size
                      19.239635
                                                    18.480155
##
```

```
## clump_thickness marginal_adhesion
## 13.276702 12.143355

## mitosis
## 3.081635

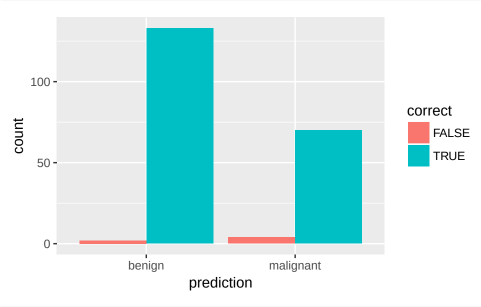
# estimate variable importance
importance <- varImp(model_rf, scale = TRUE)
plot(importance)</pre>
```



• predicting test data

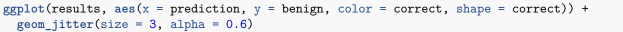
```
confusionMatrix(predict(model_rf, test_data), test_data$classes)
```

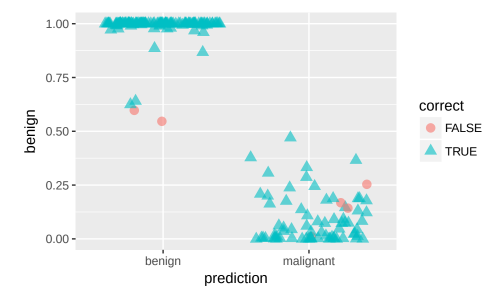
```
## Confusion Matrix and Statistics
##
##
              Reference
## Prediction benign malignant
                  133
##
     benign
                               2
##
     malignant
                             70
##
                  Accuracy : 0.9713
##
                    95% CI: (0.9386, 0.9894)
##
       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
##
```



ggplot(results, aes(x = prediction, fill = correct)) +

geom_bar(position = "dodge")





Extreme gradient boosting trees

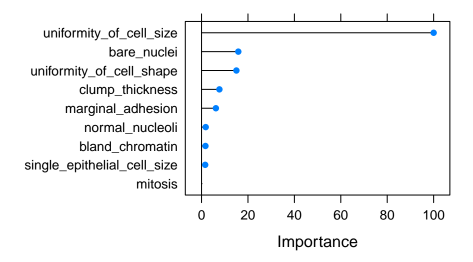
Extreme gradient boosting (XGBoost) is a faster and improved implementation of gradient boosting for supervised learning.

"XGBoost uses a more regularized model formalization to control over-fitting, which gives it better performance." Tianqi Chen, developer of xgboost

XGBoost is a tree ensemble model, which means the sum of predictions from a set of classification and regression trees (CART). In that, XGBoost is similar to Random Forests but it uses a different approach to model training. Can be used for classification and regression tasks. Here, I show a classification task.

• Feature Importance

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

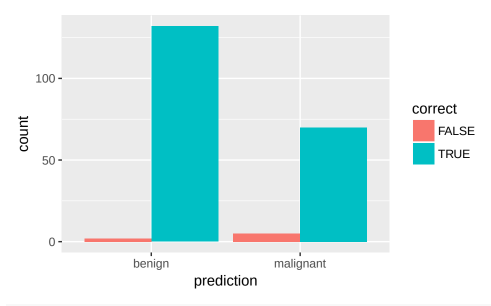


• predicting test data

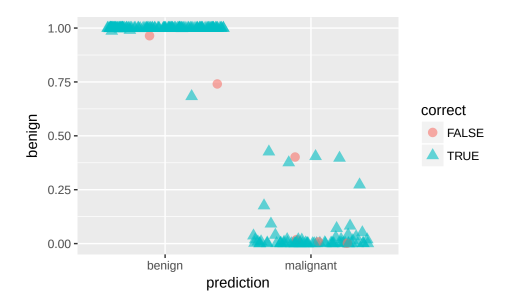
```
confusionMatrix(predict(model_xgb, test_data), test_data$classes)
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction benign malignant
## benign 132 2
## malignant 5 70
##
```

```
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
##
results <- data.frame(actual = test_data$classes,</pre>
                      predict(model_xgb, test_data, type = "prob"))
results$prediction <- ifelse(results$benign > 0.5, "benign",
                              ifelse(results$malignant > 0.5, "malignant", NA))
results$correct <- ifelse(results$actual == results$prediction, TRUE, FALSE)
ggplot(results, aes(x = prediction, fill = correct)) +
  geom_bar(position = "dodge")
```



```
ggplot(results, aes(x = prediction, y = benign, color = correct, shape = correct)) +
  geom_jitter(size = 3, alpha = 0.6)
```



Feature Selection

Machine learning uses so called features (i.e. variables or attributes) to generate predictive models. Using a suitable combination of features is essential for obtaining high precision and accuracy. Because too many (unspecific) features pose the problem of overfitting the model, we generally want to restrict the features in our models to those, that are most relevant for the response variable we want to predict. Using as few features as possible will also reduce the complexity of our models, which means it needs less time and computer power to run and is easier to understand.

Performing feature selection on the whole dataset would lead to prediction bias, we therefore need to run the whole modeling process on the training data alone!

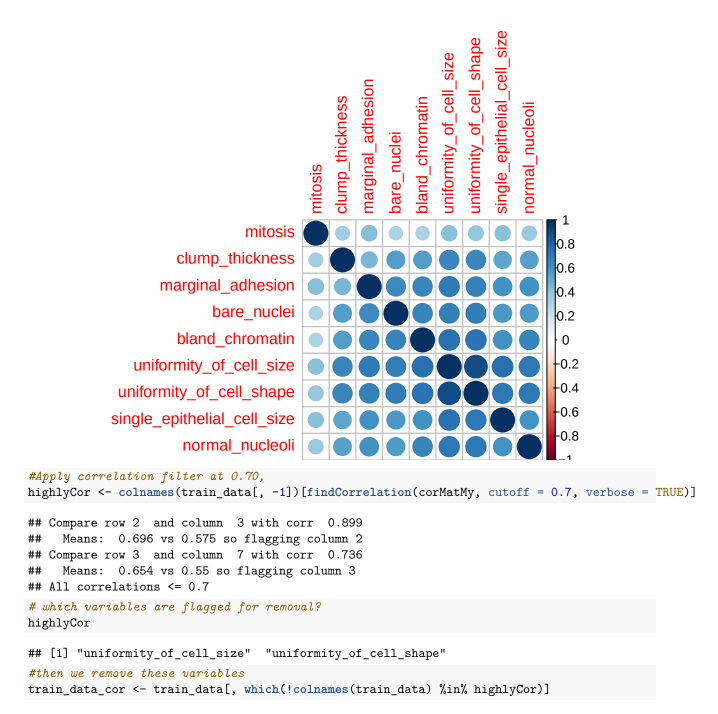
• Correlation

Often we have features that are highly correlated and thus provide redundant information. By eliminating highly correlated features we can avoid a predictive bias for the information contained in these features. This also shows us, that when we want to make statements about the biological/ medical importance of specific features, we need to keep in mind that just because they are suitable to predicting an outcome they are not necessarily causal - they could simply be correlated with causal factors.

Correlations between all features are calculated and visualised with the *corrplot* package. I am then removing all features with a correlation higher than 0.7, keeping the feature with the lower mean.

```
library(corrplot)

# calculate correlation matrix
corMatMy <- cor(train_data[, -1])
corrplot(corMatMy, order = "hclust")</pre>
```



• Recursive Feature Elimination (RFE)

Another way to choose features is with Recursive Feature Elimination. RFE uses a Random Forest algorithm to test combinations of features and rate each with an accuracy score. The combination with the highest score is usually preferential.

```
# chosen features
predictors(results_rfe)

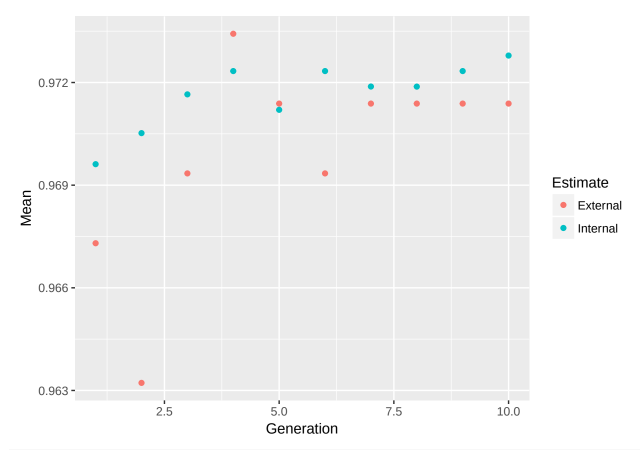
## [1] "bare_nuclei" "uniformity_of_cell_size"
## [3] "clump_thickness" "uniformity_of_cell_shape"
## [5] "bland_chromatin" "marginal_adhesion"
```

• Genetic Algorithm (GA)

The Genetic Algorithm (GA) has been developed based on evolutionary principles of natural selection: It aims to optimize a population of individuals with a given set of genotypes by modeling selection over time. In each generation (i.e. iteration), each individual's fitness is calculated based on their genotypes. Then, the fittest individuals are chosen to produce the next generation. This subsequent generation of individuals will have genotypes resulting from (re-) combinations of the parental alleles. These new genotypes will again determine each individual's fitness. This selection process is iterated for a specified number of generations and (ideally) leads to fixation of the fittest alleles in the gene pool.

This concept of optimization can be applied to non-evolutionary models as well, like feature selection processes in machine learning.





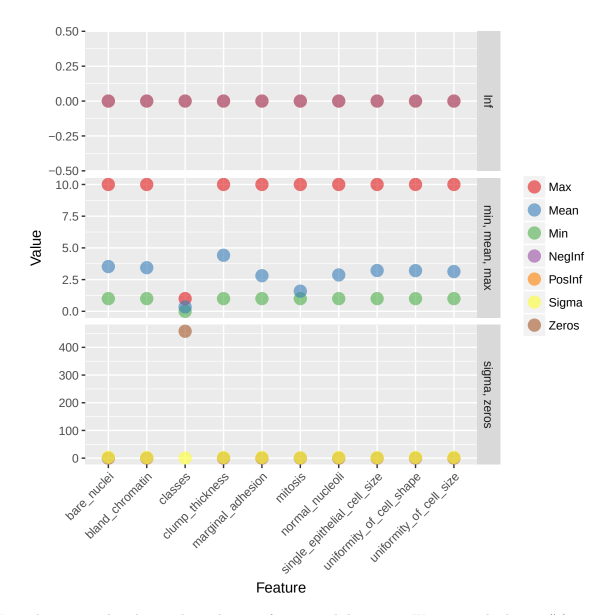
train_data_ga <- train_data[, c(1, which(colnames(train_data) %in% model_ga\$ga\$final))]</pre>

Grid search with h2o

The R package h2o provides a convenient interface to H2O, which is an open-source machine learning and deep learning platform. H2O distributes a wide range of common machine learning algorithms for classification, regression and deep learning.

```
library(h2o)
h2o.init(nthreads = -1)
## H2O is not running yet, starting it now...
##
## Note: In case of errors look at the following log files:
##
      C:\Users\s_glan02\AppData\Local\Temp\RtmpMnhd0g/h2o_s_glan02_started_from_r.out
      C:\Users\s_glan02\AppData\Local\Temp\RtmpMnhd0g/h2o_s_glan02_started_from_r.err
##
##
##
## Starting H2O JVM and connecting: . Connection successful!
## R is connected to the H2O cluster:
      H2O cluster uptime:
                                  1 seconds 832 milliseconds
##
                                  3.10.3.6
##
      H2O cluster version:
##
      H2O cluster version age:
                                 1 month and 5 days
##
      H2O cluster name:
                                 H20_started_from_R_s_glan02_tvy462
##
      H2O cluster total nodes:
                                 3.54 GB
##
      H2O cluster total memory:
##
      H2O cluster total cores:
##
      H2O cluster allowed cores:
                                 8
##
      H2O cluster healthy:
                                 TRUE
##
      H20 Connection ip:
                                  localhost
##
      H20 Connection port:
                                  54321
      H20 Connection proxy:
##
##
      R Version:
                                 R version 3.3.3 (2017-03-06)
bc_data_hf <- as.h2o(bc_data)</pre>
##
                                                                      0%
  |-----| 100%
```

We can now access all functions from the $\mathbf{h2o}$ package that are built to work on H2O Frames. A useful such function is h2o.describe(). It is similar to base R's summary() function but outputs many more descriptive measures for our data. To get a good overview about these measures, I am going to plot them.



I am also interested in the correlation between features and the output. We can use the h2o.cor() function to calculate the correlation matrix. It is again much easier to understand the data when we visualize it, so I am going to create another plot.

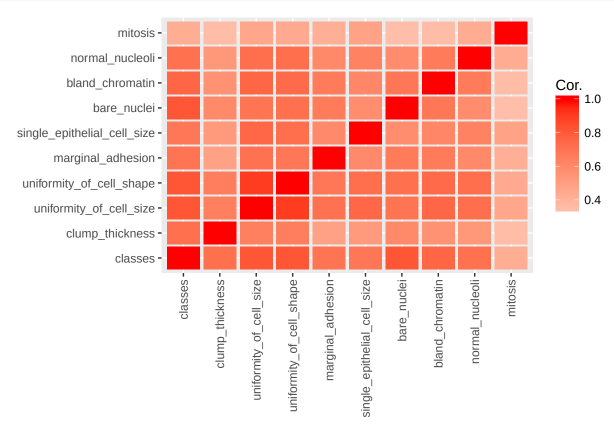
```
library(reshape2) # for melting

bc_data_hf[, 1] <- h2o.asfactor(bc_data_hf[, 1])

cor <- h2o.cor(bc_data_hf)
rownames(cor) <- colnames(cor)

melt(cor) %>%
    mutate(Var2 = rep(rownames(cor), nrow(cor))) %>%
    mutate(Var2 = factor(Var2, levels = colnames(cor))) %>%
    mutate(variable = factor(variable, levels = colnames(cor))) %>%
    ggplot(aes(x = variable, y = Var2, fill = value)) +
    geom_tile(width = 0.9, height = 0.9) +
```

```
scale_fill_gradient2(low = "white", high = "red", name = "Cor.") +
theme(axis.text.x = element_text(angle = 90, vjust = 0.5, hjust = 1)) +
labs(x = "",
    y = "")
```

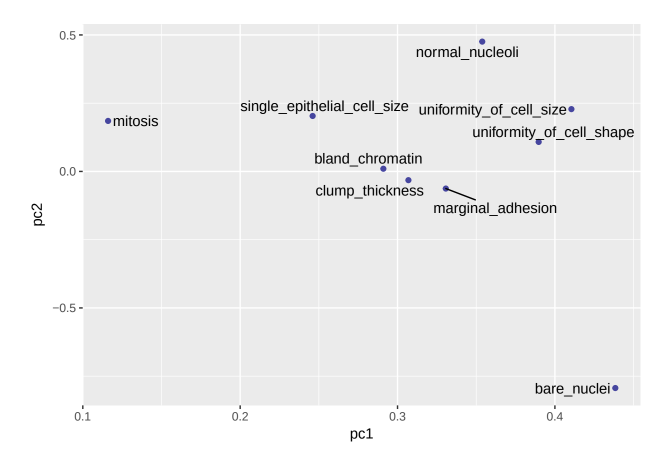


Training, validation and test data

Now we can use the h2o.splitFrame() function to split the data into training, validation and test data. Here, I am using 70% for training and 15% each for validation and testing. We could also just split the data into two sections, a training and test set but when we have sufficient samples, it is a good idea to evaluate model performance on an independent test set on top of training with a validation set. Because we can easily overfit a model, we want to get an idea about how generalizable it is - this we can only assess by looking at how well it works on previously unknown data.

I am also defining response and feature column names now.

```
summary(train$classes, exact_quantiles = TRUE)
##
    classes
## benign
             :317
## malignant:174
summary(valid$classes, exact_quantiles = TRUE)
##
    classes
## benign
             :71
## malignant:35
summary(test$classes, exact_quantiles = TRUE)
##
  classes
## benign
             :70
## malignant:32
We can also run a PCA on the training data, using the h2o.prcomp() function to calculate the singular value
decomposition of the Gram matrix with the power method.
pca <- h2o.prcomp(training_frame = train,</pre>
           x = features,
           validation_frame = valid,
           transform = "NORMALIZE",
           impute_missing = TRUE,
           k = 3,
           seed = 42)
##
                                                                             0%
                                                                       ==| 100%
eigenvec <- as.data.frame(pca@model$eigenvectors)</pre>
eigenvec$label <- features</pre>
library(ggrepel)
ggplot(eigenvec, aes(x = pc1, y = pc2, label = label)) +
  geom_point(color = "navy", alpha = 0.7) +
  geom_text_repel()
```



Classification

We can use the h2o.grid() function to perform a Random Grid Search (RGS). We could also test all possible combinations of parameters with Cartesian Grid or exhaustive search, but RGS is much faster when we have a large number of possible combinations and usually finds sufficiently accurate models.

For RGS, we first define a set of hyper-parameters and search criteria to fine-tune our models. Because there are many hyper-parameters, each with a range of possible values, we want to find an (ideally) optimal combination to maximize our model's accuracy. We can also specify how long we want to run the grid search for. Based on the results of each model tested in the grid, we can choose the one with the highest accuracy or best performance for the question on hand.

Random Forest

We now want to extract the best model from the grid model list. What makes a model *the best* depends on the question you want to address with it: in some cases, the model with highest AUC is the most suitable, or the one with the lowest mean squared error, etc. We first use the h2o.getGrid() function to sort all models by the quality metric we choose (depending on the metric, you want it ordered by descending or ascending values). We can then get the model that's the first in the list to work with further. This model's hyper-parameters can be found with best_model@allparameters. You can now work with your best model as with any regular model in h2o.

```
# performance metrics where smaller is better -> order with decreasing = FALSE
sort_options_1 <- c("mean_per_class_error", "mse", "err", "logloss")

for (sort_by_1 in sort_options_1) {
    grid <- h2o.getGrid("rf_grid", sort_by = sort_by_1, decreasing = FALSE)

    model_ids <- grid@model_ids
    best_model <- h2o.getModel(model_ids[[1]])
    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")

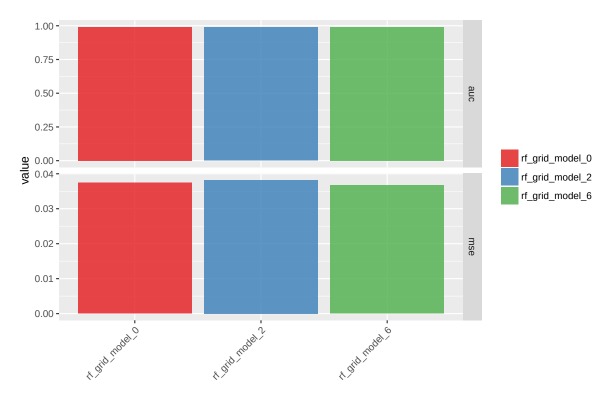
for (sort_by_2 in sort_options_2) {
    grid <- h2o.getGrid("rf_grid", sort_by = sort_by_2, decreasing = TRUE)

    model_ids <- grid@model_ids
    best_model <- h2o.getModel(model_ids[[1]])

    h2o.saveModel(best_model, path = "models", force = TRUE)
}</pre>
```

The ultimate performance test for our model will be it's prediction accuracy on the test set it hasn't seen before. Here, I will compare the AUC and mean squared error for each best model from before. You could of course look at any other quality metric that is most appropriate for your model.

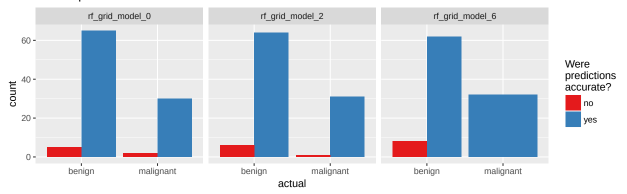
```
files <- list.files(path = "models")</pre>
rf_models <- files[grep("rf_grid_model", files)]</pre>
for (model_id in rf_models) {
  path <- paste0("U:\\Github_blog\\Webinar\\Webinar_ML_for_disease\\models\\", model_id)</pre>
  #path <- paste0("/Users/Shirin/Documents/Github/Webinar_ML_for_disease/models/", 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>
  } else {
    mse_auc_test_comb <- rbind(mse_auc_test_comb, mse_auc_test)</pre>
  }
}
mse_auc_test_comb %>%
  gather(x, y, mse:auc) %>%
  ggplot(aes(x = model_id, y = y, fill = model_id)) +
    facet_grid(x ~ ., scales = "free") +
    geom_bar(stat = "identity", alpha = 0.8, position = "dodge") +
    scale_fill_brewer(palette = "Set1") +
    theme(axis.text.x = element_text(angle = 45, vjust = 1, hjust = 1),
          plot.margin = unit(c(0.5, 0, 0, 1.5), "cm")) +
    labs(x = "", y = "value", fill = "")
```



```
for (model_id in rf_models) {
  best_model <- h2o.getModel(model_id)</pre>
 finalRf_predictions <- data.frame(model_id = rep(best_model@model_id,</pre>
                                                    nrow(test)),
                                     actual = as.vector(test$classes),
                                     as.data.frame(h2o.predict(object = best_model,
                                                                newdata = test)))
  finalRf_predictions$accurate <- ifelse(finalRf_predictions$actual == finalRf_predictions$predict,
                                          "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 {
```

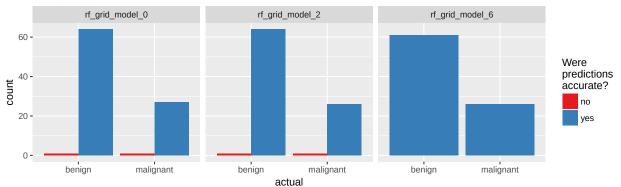
```
finalRf_predictions_comb <- rbind(finalRf_predictions_comb, finalRf_predictions)</pre>
 }
}
##
                                                                   0%
                        -----| 100%
##
                                                                   0%
##
                                                                   0%
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")
```

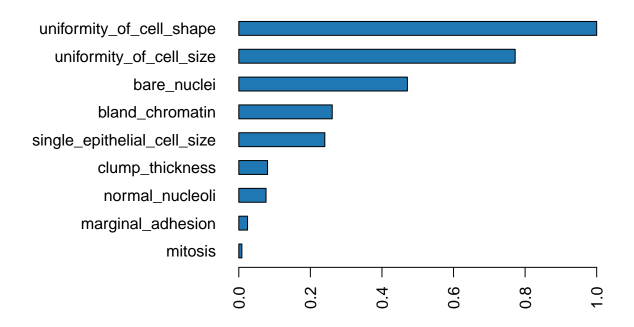
Stringent predictions



```
 \begin{tabular}{ll} rf_model &<- h2o.loadModel("U:\Github_blog\Webinar_ML_for_disease\models\rf_grid_model_6") \\ \#rf_model &<- h2o.loadModel("models/rf_grid_model_6") \\ \#summary(rf_model) \\ \#str(rf_model) \\ \end{tabular}
```

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

We can also plot the classification error.

70

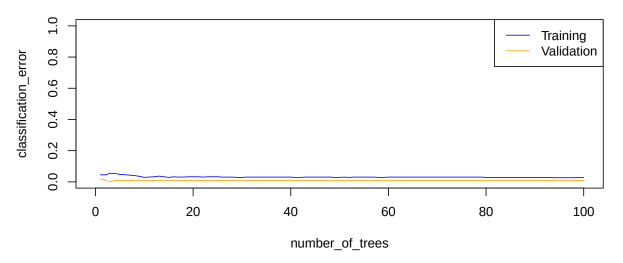
Totals

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

=1/106

36 0.009434

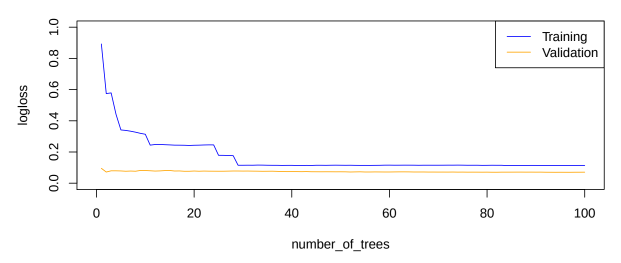
Scoring History



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).

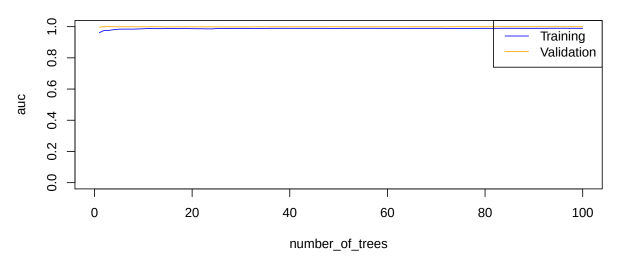
```
plot(rf_model,
    timestep = "number_of_trees",
    metric = "logloss")
```

Scoring History



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

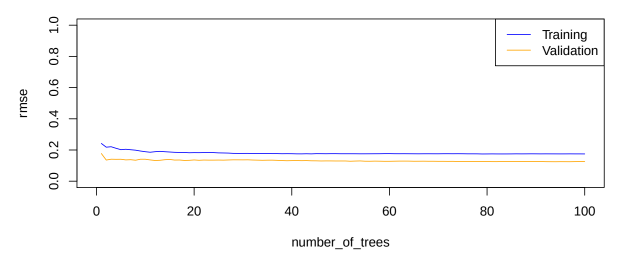
Scoring History



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

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.

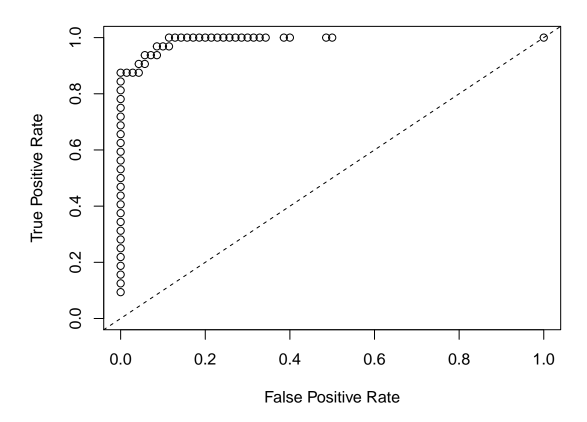
```
perf <- h2o.performance(rf_model, test)</pre>
perf
## H20BinomialMetrics: drf
##
        0.03673598
## MSE:
## RMSE: 0.1916663
## LogLoss: 0.1158835
## Mean Per-Class Error:
                          0.0625
## AUC: 0.990625
## Gini: 0.98125
##
## Confusion Matrix (vertical: actual; across: predicted) for F1-optimal threshold:
##
             benign malignant
                                  Error
                                           Rate
## benign
                 70
                             0 0.000000
                                          =0/70
```

```
## malignant
                           28 0.125000
                                          =4/32
## Totals
                 74
                           28 0.039216
                                        =4/102
##
## Maximum Metrics: Maximum metrics at their respective thresholds
##
                           metric threshold
                                                value idx
## 1
                           max f1 0.735027 0.933333
## 2
                                   0.294222 0.952381
                           max f2
## 3
                     max f0point5
                                   0.735027 0.972222
##
                     max accuracy
                                   0.735027 0.960784
                                                       25
##
                    max precision
                                   1.000000 1.000000
                                                        0
##
                       max recall
                                   0.294222 1.000000
                  max specificity
                                   1.000000 1.000000
##
                 max absolute_mcc
##
  8
                                   0.735027 0.909782
                                                       25
       max min_per_class_accuracy
## 9
                                   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.

plot(perf)

True Positive Rate vs False Positive Rate



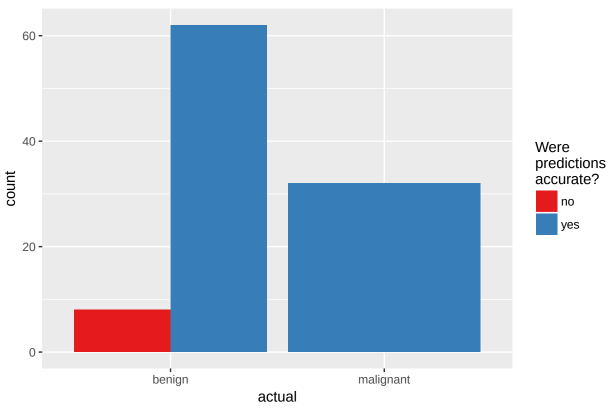
We also want to know the log loss, MSE and AUC values, as well as other model metrics for the test data:

```
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
      0.998333 0.222222 0.151515 0.416667 0.725490
                                                      1.000000 0.125000
      0.998000 0.270270 0.187970 0.480769 0.735294
                                                      1.000000 0.156250
      0.997222 0.315789 0.223881 0.535714 0.745098
                                                      1.000000 0.187500
##
      0.996210 0.358974 0.259259 0.583333 0.754902
                                                      1.000000 0.218750
##
      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
                      0.335794
                                              0.156250
                                                                       0.578125
## 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
          29
                    3 1.000000 0.906250 0.000000 0.093750
## 1
               0
      70
## 2
          28
               0
                    4 1.000000 0.875000 0.000000 0.125000
                                                             1
##
  3
      70
          27
               0
                    5 1.000000 0.843750 0.000000 0.156250
                                                             2
##
      70
          26
               0
                    6 1.000000 0.812500 0.000000 0.187500
                                                             3
  4
                    7 1.000000 0.781250 0.000000 0.218750
  5
      70
          25
               0
                                                             4
                    8 1.000000 0.750000 0.000000 0.250000
## 6
      70
          24
               0
                                                             5
```

The final predictions with probabilities can be extracted with the h2o.predict() function. Beware though, that the number of correct and wrong classifications can be slightly different from the confusion matrix above. Here, I combine the predictions with the actual test diagnoses and classes into a data frame. For plotting I also want to have a column, that tells me whether the predictions were correct. By default, a prediction probability above 0.5 will get scored as a prediction for the respective category. I find it often makes sense to be more stringent with this, though and set a higher threshold. Therefore, I am creating another column with stringent predictions, where I only count predictions that were made with more than 80% probability. Everything that does not fall within this range gets scored as "uncertain". For these stringent predictions, I am also creating a column that tells me whether they were accurate.

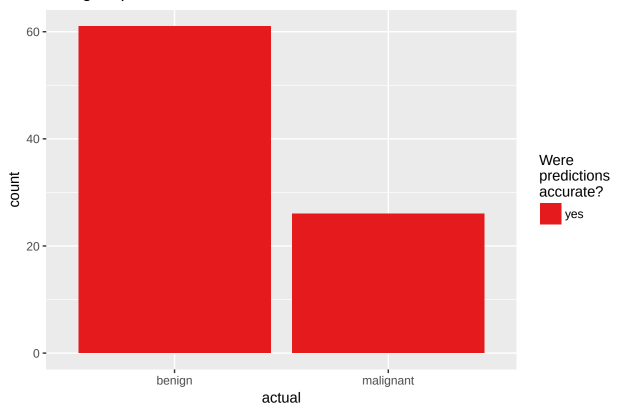
```
finalRf_predictions$accurate <- ifelse(finalRf_predictions$actual == finalRf_predictions$predict, "yes"
finalRf_predictions$predict_stringent <- ifelse(finalRf_predictions$benign > 0.8, "benign",
                                                 ifelse(finalRf_predictions$malignant > 0.8, "malignant"
finalRf_predictions$accurate_stringent <- ifelse(finalRf_predictions$actual == finalRf_predictions$pred
                                       ifelse(finalRf_predictions$predict_stringent == "uncertain", "na
finalRf_predictions %>%
  group_by(actual, predict) %>%
  dplyr::summarise(n = n())
## Source: local data frame [3 x 3]
## Groups: actual [?]
##
##
        actual
                 predict
##
        <fctr>
                 <fctr> <int>
## 1
        benign
                  benign
                            62
        benign malignant
                             8
## 2
## 3 malignant malignant
                            32
finalRf_predictions %>%
  group_by(actual, predict_stringent) %>%
  dplyr::summarise(n = n())
## Source: local data frame [4 x 3]
## Groups: actual [?]
##
##
        actual predict_stringent
                                     n
##
        <fctr>
                           <chr> <int>
## 1
        benign
                         benign
                                    61
## 2
        benign
                       uncertain
                                     9
## 3 malignant
                       malignant
                                    26
## 4 malignant
                       uncertain
finalRf_predictions %>%
  ggplot(aes(x = actual, fill = accurate)) +
    geom_bar(position = "dodge") +
    scale_fill_brewer(palette = "Set1") +
    labs(fill = "Were\npredictions\naccurate?",
         title = "Default predictions")
```

Default predictions



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

Stringent predictions



h2o.shutdown()

If you are interested in more machine learning posts, check out the category listing for **machine_learning** on my blog.

sessionInfo()

```
## R version 3.3.3 (2017-03-06)
## 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] stats4
                 parallel stats
                                     graphics grDevices utils
                                                                   datasets
## [8] methods
                 base
##
## other attached packages:
  [1] ggrepel_0.6.5
                                                  h2o_3.10.3.6
                             reshape2_1.4.2
```

```
plyr_1.8.4
    [4] corrplot_0.77
                                                   xgboost_0.6-4
  [7] randomForest_4.6-12
                                                   lattice_0.20-34
                             caret_6.0-73
## [10] igraph_1.0.1
                             tidyr_0.6.1
                                                   dplyr_0.5.0
## [13] pcaGoPromoter_1.18.0 Biostrings_2.42.1
                                                  XVector_0.14.0
## [16] IRanges_2.8.1
                             S4Vectors_0.12.1
                                                   BiocGenerics_0.20.0
## [19] ellipse 0.3-8
                             ggplot2_2.2.1.9000
## loaded via a namespace (and not attached):
   [1] Rcpp_0.12.9
                             class_7.3-14
                                                   assertthat_0.1
##
   [4] rprojroot_1.2
                             digest_0.6.12
                                                   foreach_1.4.3
                                                  MatrixModels_0.4-1
  [7] R6_2.2.0
                             backports_1.0.5
## [10] RSQLite_1.1-2
                                                   e1071_1.6-8
                             evaluate_0.10
                             lazyeval_0.2.0
## [13] zlibbioc_1.20.0
                                                  minqa_1.2.4
## [16] data.table_1.10.4
                             SparseM_1.76
                                                   car_2.1-4
## [19] nloptr_1.0.4
                             Matrix_1.2-8
                                                  rmarkdown_1.3
## [22] labeling_0.3
                             splines_3.3.3
                                                   lme4_1.1-12
## [25] stringr_1.2.0
                             RCurl_1.95-4.8
                                                  munsell_0.4.3
## [28] mgcv_1.8-17
                             htmltools 0.3.5
                                                  nnet 7.3-12
## [31] tibble_1.2
                             codetools_0.2-15
                                                  MASS_7.3-45
## [34] bitops 1.0-6
                             ModelMetrics_1.1.0
                                                  grid_3.3.3
## [37] nlme_3.1-131
                             jsonlite_1.3
                                                   gtable_0.2.0
## [40] DBI_0.6
                             magrittr_1.5
                                                   scales_0.4.1
## [43] stringi_1.1.2
                             RColorBrewer_1.1-2
                                                   iterators_1.0.8
                                                  pbkrtest_0.4-6
## [46] tools 3.3.3
                             Biobase 2.34.0
## [49] yaml_2.1.14
                             AnnotationDbi_1.36.2 colorspace_1.3-2
## [52] memoise_1.0.0
                             knitr_1.15.1
                                                  quantreg_5.29
```