Stats Lab Bird

Joel Alder, Connor Charlton, Samir Hauser

2023-05-02

Data pre processing

We first load in the data and preprocess it.

We transform our response variable values to 0 and 1.

```
d.bird$sex_genetics[d.bird$sex_genetics == "M"] <- "m"
d.bird$sex_genetics[d.bird$sex_genetics == "F"] <- "f"
d.bird$sex_genetics[d.bird$sex_genetics == "m"] <- 0
d.bird$sex_genetics[d.bird$sex_genetics == "f"] <- 1</pre>
```

Next we define a new variable season, which indicates the time of the ringing. We do this because some of the variables depend on the time of year; for example the weight is different in summer and winter.

We have two variables for the feather length, P1 and P8. We merge them into one variable and where this variable takes the value of P8 if P8 has a value, and takes the value of P1 otherwise. There are only 32 instances for which P1 has a value and P8 does not.

```
if (sum(!is.na(d.bird$P1)) > 0) {
  d.bird$P8[is.na(d.bird$P8)] <- d.bird$P1[is.na(d.bird$P8)]
} else {
  print("NO impuatation possible as P1 is NA")
}</pre>
```

There are some duplicates in the dataset which we remove. Additionally, there is a possibility that one bird is captured more than once. If this is the case we only use the latest observation. Afterwards, we select only the adult birds and keep the ones for which we have the labels. In the end we select the columns we want to use for training. These are all the different morphological traits.

```
# Remove duplicates
d.bird <- d.bird[!duplicated(d.bird, fromLast = TRUE), ]

# check multiple captures
freq_table <- table(d.bird$ringnr)</pre>
```

```
freq_table <- freq_table[freq_table > 1]
freq_table <- sort(freq_table, decreasing = T)</pre>
double_caputre = as.data.frame(freq_table)
d.bird <- d.bird[!duplicated(d.bird$ringnr, fromLast = TRUE), ]</pre>
# select only the adult birds
df_adult <- d.bird[d.bird$Age>1,]
# select only rows with no NA in the response
df adult <- df adult[complete.cases(df adult["sex genetics"]),]</pre>
# select only the needed columns
cols <- c("season", "Age", "Wing", "P8", "Tarsus", "weight", "Fat", "Muscle",</pre>
          "Bill_length", "sex_genetics")
df_adult_sub = data.frame(df_adult[cols])
# chaning type of variables
df_adult_sub$Fat <- as.factor(df_adult_sub$Fat)</pre>
df_adult_sub$Muscle <- as.factor(df_adult_sub$Muscle)</pre>
df_adult_sub$sex_genetics <- as.factor(df_adult_sub$sex_genetics)</pre>
str(df_adult_sub)
## 'data.frame': 981 obs. of 10 variables:
               : Factor w/ 4 levels "0","1","2","3": 3 3 2 3 2 2 2 2 3 3 ...
## $ season
## $ Age
                 : num 5 4 4 5 4 4 4 6 5 5 ...
## $ Wing
                 : num 123 116 123 123 116 ...
## $ P8
                 : num 98 90 97 94 90 96.5 89 89.5 89.5 95 ...
                : num 23.4 22.7 22.9 23.4 22.5 24.2 22.6 23.2 22.8 22.7 ...
## $ Tarsus
## $ weight
                : num 38 40 37.5 34 32.5 35 33 34 38 35.5 ...
```

We now train three different models on this data - a decision tree, a random forest and lastly a logistic regression model with missing values imputed via using MICE. We use the accuracy of our predictions as our evaluation metric.

: Factor w/ 3 levels "1","2","3": 2 NA 2 2 3 1 2 3 1 1 ...

\$ Bill_length : num NA NA 10.8 14.1 NA NA NA NA NA NA ...

\$ sex_genetics: Factor w/ 2 levels "0","1": 1 2 1 1 2 1 2 2 2 1 ...

: Factor w/ 7 levels "0","1","2","3",...: 3 NA 2 2 2 2 2 3 2 ...

Decision Tree

\$ Fat

\$ Muscle

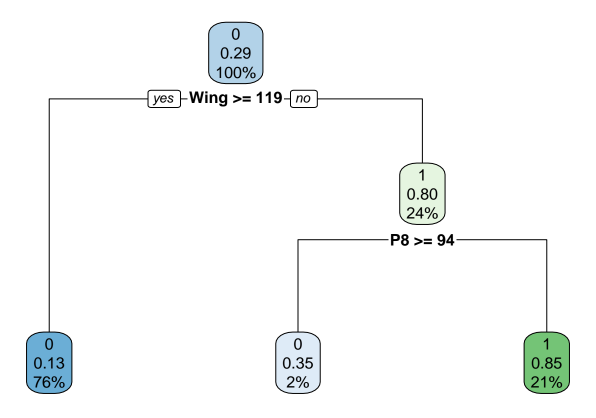
We use a ten fold cross-validation to get the accuracy score of the decision tree. The advantage of this method is that it can handle the missing values.

```
n <- dim(df_adult_sub)[1]
K <- 10
folds <- sample(cut(seq(1,n),breaks=K,labels=FALSE), replace = FALSE)
Fold.error <- numeric(K)
for (i in 1:K) {
   test.ind <- which(folds == i)
   tree_model.i <- rpart(sex_genetics ~ ., data = df_adult_sub[-test.ind,], method = "class")
   test_predictions.i <- predict(tree_model.i, df_adult_sub[test.ind,], type = "class")
   Fold.error[i] <- mean(test_predictions.i == df_adult_sub[test.ind,]$sex_genetics)</pre>
```

```
}
cat("Accuracy Decision Tree:", round(mean(Fold.error), digits = 5))
```

Accuracy Decision Tree: 0.84714

We get an accuracy around 84.7 %. One question that is raised is whether these two errors should be weighted differently. Maybe the misclassification of a female bird as a male bird should have a higher price than the opposite. We should keep in mind that the sex distribution at the population level is unbalanced, with two thirds being male. Now we plot the decision tree, to see which variables are important.



This produces a rather simple tree. The 0 values represent male birds and the 1 values represent female birds. As can be seen in the descriptive analysis of our data, the wing variable is the most informative for the prediction of genetic sex. There are only two variables in our tree.

Random Forest

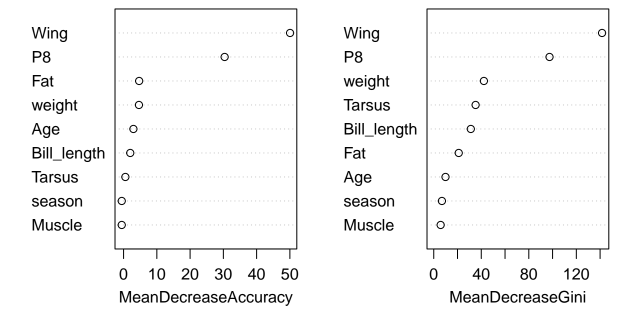
We again use ten fold cross-validation to get the accuracy score of the random forest. The random forest can also handle missing values by specifying the na.action parameter. This will start by imputing the missing values using the column-wise median or mode. Then it grows a forest and computes proximities, before iterating through and constructing a new forest using these newly imputed values and repeating this process on repeat.

```
## Tuning of parameter done via this code
```

Accuracy Random Forest: 0.84472

We get an accuracy around 84.5 %. We tuned the parameter ntree and mtry and with this combiantion we get the best result. In the plot below we see the importance of the different variables. Like in the decision tree, wing and P8 are the most important ones.

model



Logistic regression model

Logistic regression with complete cases

Before we use MICE to impute the missing values, we only use the complete cases with 741 observation. Because there might be some interactions we build a model which includes them. We do model selection with AIC and then we fit the best model with ten fold cross-validation to compute the accuracy score.

```
# use only the complete cases
df_adult_sub_comp <- na.omit(df_adult_sub)</pre>
# define the full model and fit it
fit <- glm(sex_genetics ~ season + Age + Wing + P8 + Tarsus + weight + Fat + Muscle
           + Bill_length + Wing*P8 + Wing*Bill_length + Wing*Tarsus + Wing*weight
           + P8*Bill_length + P8*Tarsus + P8*weight + Tarsus*weight + Tarsus*Bill_length
           + weight*Bill_length, data = df_adult_sub_comp, family = "binomial")
# use the step function to select the best model. It uses AIC criterion
step(fit, direction = "both", trace = 0)
##
## Call: glm(formula = sex_genetics ~ season + Wing + P8 + Tarsus + Muscle +
       Bill_length, family = "binomial", data = df_adult_sub_comp)
##
## Coefficients:
## (Intercept)
                                                                  P8
                                                                            Tarsus
                    season1
                                  season2
                                                   Wing
       77.8480
                                                                            0.3041
##
                     0.2269
                                  -3.6361
                                                -0.5272
                                                             -0.3373
       Muscle2
                    Muscle3 Bill length
##
                                   0.7169
##
        1.5315
                     2.7487
##
## Degrees of Freedom: 740 Total (i.e. Null); 732 Residual
## Null Deviance:
                        885.3
## Residual Deviance: 464.1
                                 AIC: 482.1
# we get this model as our best
best_model <- formula(sex_genetics ~ season + Wing + P8 + Tarsus + Muscle + Bill_length)</pre>
#do 10 fold cross validation with the best model
set.seed(123)
n <- dim(df_adult_sub_comp)[1]</pre>
folds <- sample(cut(seq(1,n),breaks=K,labels=FALSE), replace = FALSE)</pre>
Fold.error <- numeric(K)</pre>
for (i in 1:K) {
  test.ind <- which(folds == i)</pre>
  glm.i <- glm(best_model, data = df_adult_sub_comp[-test.ind,], family = binomial)</pre>
 prob_test_predictions.i <- predict(glm.i, df_adult_sub_comp[test.ind,], type = "response")</pre>
  test_predictions.i <- ifelse(prob_test_predictions.i > 0.5, 1, 0)
  Fold.error[i] <- mean(test_predictions.i == df_adult_sub_comp[test.ind,]$sex_genetics)
cat("Accuracy Logistic Model Complete Cases:", round(mean(Fold.error), digits = 5))
```

Accuracy Logistic Model Complete Cases: 0.86099

This yields an accuracy of 86.1 %. Now we look at the estimated coefficients of the model and there significance.

```
##
                 Estimate Std. Error
                                        z value
                                                    Pr(>|z|)
## (Intercept) 77.8479749 7.27304118 10.7036346 9.786274e-27
               0.2269047 0.25391932 0.8936094 3.715309e-01
## season1
## season2
              -3.6361072 2.30854415 -1.5750651 1.152414e-01
              -0.5271909 0.06626136 -7.9562352 1.773531e-15
## Wing
              -0.3372846 0.06968261 -4.8402973 1.296450e-06
## P8
## Tarsus
               0.3040828 0.16593143 1.8325811 6.686489e-02
## Muscle2
               1.5315476 0.91143754 1.6803648 9.288636e-02
## Muscle3
               2.7487494 1.02567619 2.6799387 7.363563e-03
## Bill_length 0.7169340 0.26025062 2.7547829 5.873109e-03
```

Logistic regression with imputing missing values

We again use ten fold cross-validation to compute the accuracy score of the logistic regression model. Because this model type can not handle missing values, the missing values are first imputed using MICE. This works by producing 35 separate imputations on which the logistic regression model is fitted independently, and the modal classification adopted. We use again the best model select by the AIC.

```
imp <- mice(df_adult_sub, seed = 123, print = F, m = 35, maxit = 10)</pre>
imp <- mice::complete(imp, "all")</pre>
Fold.error <- numeric(K)</pre>
for (i in 1:K) {
  test.ind <- which(folds == i)</pre>
  train data <- df adult sub[-test.ind,]
  test_data <- df_adult_sub[test.ind,]</pre>
  # mice
  imp_train = mice(train_data, seed = 123, print = F, m=35, maxit = 10)
  # fit model
  fit_1 <- with(imp_train, glm(sex_genetics ~ season + Wing + P8 + Tarsus + Muscle
                                + Bill_length, family = binomial))
  pooled <- pool(fit_1)</pre>
  # hack for predict
  pooled_lm = fit_1$analyses[[1]]
  pooled_lm$coefficients = summary(pooled)$estimate
  size = nrow(imp_train[[1]][test.ind,-(10)]) # remove sex
  # loop over imputed data
  dat = matrix(nrow = size, ncol = 35)
  for (k in 1:35)
    predicted_values = predict(pooled_lm, newdata = imp[[k]][test.ind,-(10)],
                                 type="response")
    binary_predictions <- ifelse(predicted_values > 0.5, 1, 0)
    dat[,k] = binary_predictions
 }
```

```
#majority vote
test_predictions.i <- apply(dat,1,Mode)

Fold.error[i] <- mean(test_predictions.i == df_adult_sub[test.ind,]$sex_genetics)
}
cat("Accuracy Logistic Model with imputation:", round(mean(Fold.error), digits = 5))</pre>
```

Accuracy Logistic Model with imputation: 0.87852

This results in a slightly improved accuracy of 87.9 %. Now we examine the pooled estimates for the coefficients of the logistic regression model to the significance of them. We omit the fat variable because the standard error of the coefficient estimates was too high and the inclusion of the fat variable, as a predictor, explains too little variance in the regressand.

```
summary(pooled)
```

```
##
                             std.error statistic
                                                         df
                                                                 p.value
            term
                    estimate
## 1 (Intercept) 83.77497783 7.02201982 11.9303249 751.0696 3.605425e-30
## 2
                 0.08640196 0.22787975 0.3791559 843.0295 7.046676e-01
## 3
         season2 -1.72071584 0.80862323 -2.1279575 714.2446 3.368243e-02
## 4
            Wing -0.56643458 0.06287997 -9.0081885 869.7928 1.301949e-18
## 5
              P8 -0.34252105 0.06481787 -5.2843617 794.3216 1.630851e-07
## 6
                 0.29398607 0.15829457
                                        1.8572088 496.4544 6.387345e-02
## 7
         Muscle2
                 0.93197754 0.55022727
                                        1.6938047 890.7103 9.065202e-02
## 8
         Muscle3
                 2.19493298 0.70279906 3.1231302 864.6421 1.848974e-03
                 0.72407727 0.24621127 2.9408779 418.5761 3.454424e-03
## 9 Bill_length
```

Similarly, to the decision tree this suggests that both the Wing and P8 variables are highly significant and the Bill length and Muscle variables are significant on a 5 % level.

Comparison of the coefficient estimates

Here we compare the coefficients of the logistic regression only with the complete cases to the one with the imputed data.

```
##
                     [,1]
                                  [,2]
## (Intercept) 77.8479749 83.77497783
## season1
                0.2269047 0.08640196
## season2
               -3.6361072 -1.72071584
## Wing
               -0.5271909 -0.56643458
## P8
               -0.3372846 -0.34252105
## Tarsus
                0.3040828
                           0.29398607
## Muscle2
                1.5315476
                           0.93197754
## Muscle3
                2.7487494
                           2.19493298
## Bill_length 0.7169340 0.72407727
```

Conclusion

In conclusion, the imputation of missing values via MICE and subsequent classification using logistic regression is the best model, yielding around 87 % accuracy for the prediction of the genetic sex of the birds. The two most important variables to measure going forward are the Wing length and the feather length (P8).