# PCOS modeling

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# Contents

Creating the classificators	1
Setup	 2
Load packages and data	 2
Data splitting	 2
Imputation on training set	 9
Define the models	 10
Define cross-validation parameters	 10
Modeling	 10
Logistic regression modeling	 10
Elastic net regression	 13
Random Forest	 19
Model comparison	 22
Test the performance in the validation set $\dots \dots \dots \dots \dots \dots \dots$ .	 22
Explore EN model 1	 24
Variable importance	 24
Effect of class imbalance on the best model (if time allows)	 25

# Creating the classificators

This report contains the details of the generation and comparison of several classifiers whose aim is to classify people with and without polycystic ovary syndrome (PCOS). This report is part of a larger project with a goal to develop and validate a model that takes easily collected variables, and tests its performance against models using variables obtained with increasingly invasive procedures. For a summary of the results and a clearer picture of the project, please read the Final\_report.pdf file in this repository.

### Setup

#### Load packages and data

#### Data splitting

## 364 177

After loading the data and packages, we will proceed to split the data into training and validation sets using a 70:30 random split.

```
data = data %>%
  column_to_rownames("id")
# Make the code reproducible
set.seed(504)
# Create the data split
train.index <- caret::createDataPartition(data$pcos, p = .7, list=FALSE)</pre>
# Define the training and validation data sets
train <- data[ train.index,]</pre>
valid <- data[-train.index,]</pre>
# Check the ratio of cases with and without the main outcome measure (i.e. PCOS diagnosis) in the train
table(train$pcos)
##
## No Yes
## 255 124
table(train$pcos)[1]/ table(train$pcos)[2]
##
         No
## 2.056452
# Check the ratio of cases with and without the main outcome measure (i.e. PCOS diagnosis) in the valid
table(valid$pcos)
##
## No Yes
## 109 53
table(valid$pcos)[1]/ table(valid$pcos)[2]
##
         No
## 2.056604
# Check the ratio of cases with and without the main outcome measure (i.e. PCOS diagnosis) in the origi
table(data$pcos)
##
## No Yes
```

#### table(data\$pcos)[1]/ table(data\$pcos)[2]

```
## No
## 2.056497
```

The ratio of cases with and without the main outcome measure (i.e. PCOS diagnosis) is similar between the original (2.0565), validation (2.0566) and training (2.0565) data.

#### Imputation on training set

Following the data splitting, we will impute missing values for the training set. As observed in the exploratory data analysis, the proportion of missing data is very low overall, which makes variables with missingness suitable for imputation.

```
# Explore misingness in training set
sapply(train, function(x) sum(is.na(x)))
```

##	200	0.00	rroimh+	hoimh+	hmi
	pcos	age	weight	height	bmi
##	0	0	0	0	0
##	blood_group	<pre>pulse_rate</pre>	rr	hb	cycle
##	0	2	0	0	0
##	cycle_length	marriage_status	pregnant	no_of_abortions	i_betahcg
##	0	0	0	0	0
##	<pre>ii_betahcg</pre>	fsh	lh	fsh_lh_ratio	hip
##	0	0	1	1	0
##	waist	waist_hip_ratio	tsh	amh	prl
##	0	0	0	2	0
##	vitd3	prg	rbs	weight_gain	hair_growth
##	1	0	0	0	0
##	skin_darkening	hair_loss	pimples	fast_food	reg_exercise
##	0	0	0	1	0
##	<pre>bp_systolic</pre>	<pre>bp_diastolic</pre>	follicle_no_l	follicle_no_r	avg_f_size_l
##	0	0	0	0	0
##	avg_f_size_r	endometrium			
##	0	0			

For imputation, we will use the mice R package, a widely used software to handle missing data. We will use the default options, which use the most appropriate methodology depending on the class of the data to be imputed, which are the following: "By default, the method uses pmm, predictive mean matching (numeric data) logreg, logistic regression imputation (binary data, factor with 2 levels) polyreg, polytomous regression imputation for un-ordered categorical data (factor > 2 levels) polr, proportional odds model for (ordered, > 2 levels)."

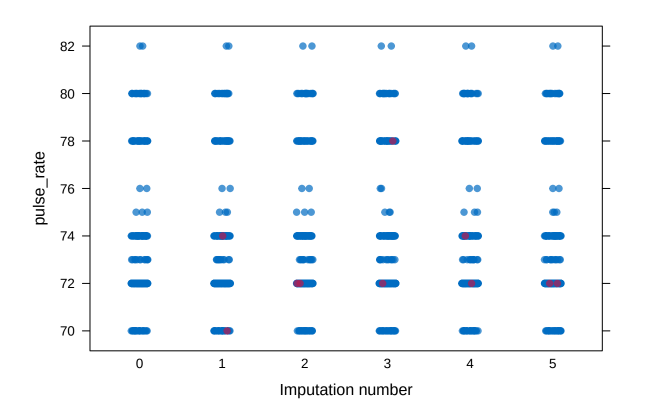
```
chained_train = mice::mice(train)
```

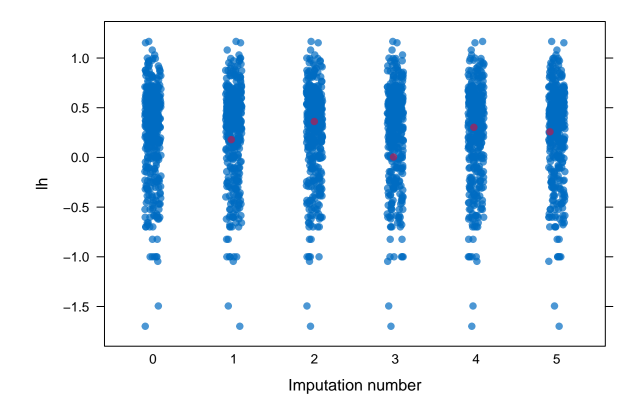
```
##
##
   iter imp variable
##
        1 pulse rate lh fsh lh ratio
                                            vitd3 fast food
                                       amh
                                            vitd3 fast_food
##
    1
        2 pulse_rate lh fsh_lh_ratio
                                       amh
##
        3 pulse_rate lh fsh_lh_ratio
                                       amh vitd3 fast_food
    1
          pulse rate lh fsh lh ratio
                                       amh vitd3 fast food
##
    1
```

```
##
     1
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     2
         1
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
            pulse_rate
                                                          fast_food
     2
                                                          fast food
##
            pulse_rate
                          lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
     2
##
            pulse_rate
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
                         lh
##
     2
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     2
                              fsh_lh_ratio
                                                  vitd3
                                                          fast food
            pulse_rate
                         lh
                                             amh
##
     3
                              fsh lh ratio
                                                  vitd3
                                                          fast food
            pulse_rate
                         lh
                                             amh
     3
                                                          fast_food
##
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
##
     3
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     3
                                             amh
                                                  vitd3
            pulse_rate
                         lh
                              fsh_lh_ratio
                                                          fast_food
##
     3
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     4
                                                  vitd3
                                                          fast_food
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
     4
##
         2
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     4
                              fsh_lh_ratio
                                                  vitd3
             pulse_rate
                                             amh
                                                          fast_food
##
     4
                              fsh_lh_ratio
                                                  vitd3
                                                          fast_food
            pulse_rate
                         lh
                                             amh
##
     4
         5
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     5
         1
                                                  vitd3
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                          fast_food
     5
##
            pulse rate
                         lh
                              fsh lh ratio
                                             amh
                                                  vitd3
                                                          fast food
##
     5
                                                  vitd3
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                          fast_food
     5
##
            pulse_rate
                         lh
                              fsh lh ratio
                                             amh
                                                  vitd3
                                                          fast_food
##
     5
            pulse_rate
                         lh
                              fsh_lh_ratio
                                             amh
                                                  vitd3
                                                          fast_food
```

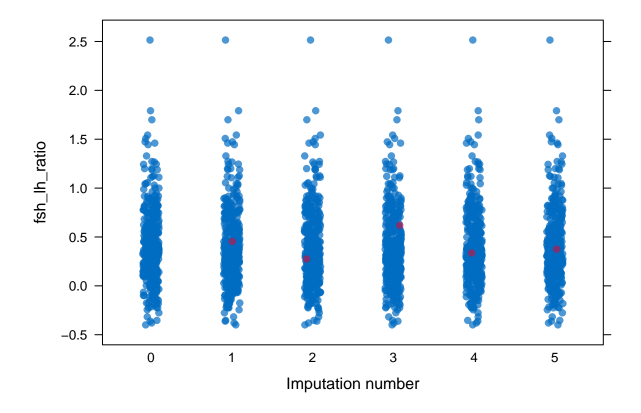
## Warning: Number of logged events: 98

# Explore the imputed data and check that the generated value are plausible
stripplot(chained\_train, pulse\_rate, pch = 19, xlab = "Imputation number")

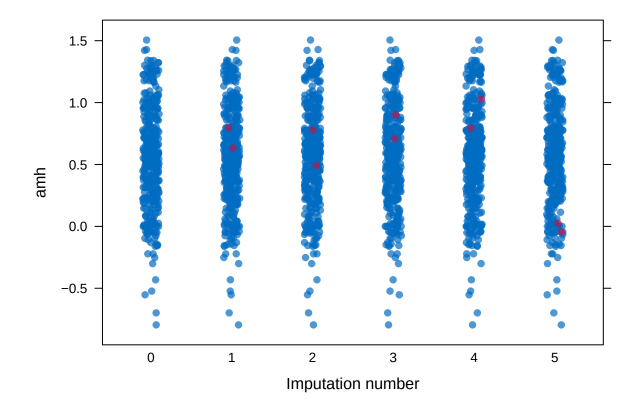




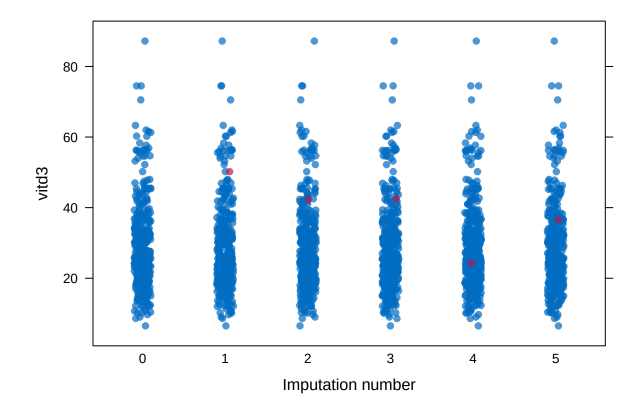
stripplot(chained\_train, fsh\_lh\_ratio, pch = 19, xlab = "Imputation number")



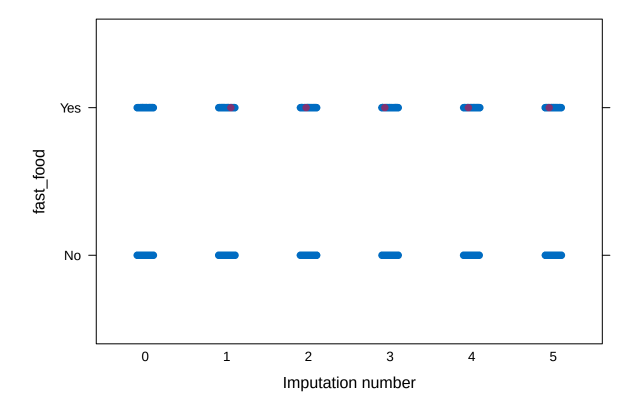
stripplot(chained\_train, amh, pch = 19, xlab = "Imputation number")



stripplot(chained\_train, vitd3, pch = 19, xlab = "Imputation number")



stripplot(chained\_train, fast\_food, pch = 19, xlab = "Imputation number")



The values appear to be plausible, therefore, we will proceed with extracting the imputed data

```
chained_train = complete(data=chained_train)
sapply(chained_train, function(x) sum(is.na(x)))
```

##	pcos	age	weight	height	bmi
##	0	0	0	0	0
##	blood_group	<pre>pulse_rate</pre>	rr	hb	cycle
##	0	0	0	0	0
##	cycle_length	marriage_status	pregnant	no_of_abortions	i_betahcg
##	0	0	0	0	0
##	<pre>ii_betahcg</pre>	fsh	lh	fsh_lh_ratio	hip
##	0	0	0	0	0
##	waist	waist_hip_ratio	tsh	amh	prl
##	0	0	0	0	0
##	vitd3	prg	rbs	weight_gain	hair_growth
##	0	0	0	0	0
##	skin_darkening	hair_loss	pimples	fast_food	reg_exercise
##	0	0	0	0	0
##	<pre>bp_systolic</pre>	<pre>bp_diastolic</pre>	follicle_no_l	follicle_no_r	avg_f_size_l
##	0	0	0	0	0
##	avg_f_size_r	endometrium			
##	0	0			

#### Define the models

For the aim of this project, we will develop 4 different models with an increasing number of variables according to how easy they are to collect. Model 1 will contain only variables that are collected through patient history, model 2 will add variables collected through clinical examination, model 3 adds results of blood tests and model 4 will include all the relevant variables available in the data, including results from the transvaginal ultrasound.

It is worth mentioning that we removed some non-relevant variables from the dataset based on the scientific literature of the field. A more detailed information about this selection can be found in the final report available in this repository.

#### Define cross-validation parameters

```
# Set the cross-validation parameters for all models
fitControl <- trainControl(
   method = 'cv',
   number = 5,
   savePredictions = 'final',
   classProbs = TRUE,
   summaryFunction=twoClassSummary)</pre>
```

### Modeling

#### Logistic regression modeling

First, we will fit a logistic regression model with the features mentioned above.

```
# Set the seed to ensure reproducibility
set.seed(504)

#Since we have few samples, we will use bootstrapping instead of cross fold validation
models_logreg = list()
```

```
for (model in list(model1_vars, model2_vars, model3_vars, model4_vars)){
  cv_model = caret::train(
   pcos ~ .,
   data = chained train %>%
     dplyr::select(all_of(model)),
   method = "glm",
   family = "binomial",
   trControl = fitControl)
  models_logreg = append(models_logreg, list(cv_model))
## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not
## in the result set. ROC will be used instead.
## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not
## in the result set. ROC will be used instead.
## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not
## in the result set. ROC will be used instead.
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## Warning in train.default(x, y, weights = w, ...): The metric "Accuracy" was not
## in the result set. ROC will be used instead.
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
## Warning: glm.fit: fitted probabilities numerically 0 or 1 occurred
names(models_logreg) = c("model1", "model2", "model3", "model4")
models_logreg
## $model1
## Generalized Linear Model
##
## 379 samples
## 11 predictor
   2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 304, 303, 303, 303, 303
```

```
## Resampling results:
##
##
     ROC
                Sens
                           Spec
     0.9050523 0.9019608 0.718
##
##
##
## $model2
## Generalized Linear Model
##
## 379 samples
  19 predictor
     2 classes: 'No', 'Yes'
##
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 304, 303, 303, 303
## Resampling results:
##
##
    ROC
                Sens
                           Spec
##
     0.8943791 0.9019608 0.6946667
##
##
## $model3
## Generalized Linear Model
##
## 379 samples
## 27 predictor
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 304, 303, 303, 303
## Resampling results:
##
##
     ROC
                Sens
                           Spec
     0.8718627 0.8941176 0.6943333
##
##
##
## $model4
## Generalized Linear Model
##
## 379 samples
   32 predictor
##
##
     2 classes: 'No', 'Yes'
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 304, 303, 303, 303
## Resampling results:
##
##
     ROC
                           Spec
                Sens
##
    0.9350327 0.9137255 0.815
```

#### Elastic net regression

## glmnet

## 379 samples

##

We will also fit an elastic net model, which is a method that performs feature selection and remove redundant terms in the models.

```
models_EN = list()
for (model in list(model1_vars, model2_vars, model3_vars, model4_vars)){
  en_model = caret::train(
    pcos ~ .,
    trControl = fitControl,
    data = chained_train %>%
      dplyr::select(all_of(model)),
    method = "glmnet",
    tuneGrid = expand.grid(alpha = seq(0.1,.2,by = 0.05),
                           lambda = seq(0.05, 0.3, by = 0.05)),
    verbose = FALSE,
    metric="ROC")
  models_EN = append(models_EN, list(en_model))
}
names(models_EN) = c("model1", "model2", "model3", "model4")
models_EN
## $model1
```

```
##
   11 predictor
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 304, 303, 303, 303
## Resampling results across tuning parameters:
##
##
     alpha lambda ROC
                               Sens
                                          Spec
##
     0.10
           0.05
                    0.9064837 0.9254902
                                         0.7253333
##
           0.10
     0.10
                    0.9051275 0.9372549
                                         0.6933333
##
     0.10
           0.15
                   0.9040359 0.9490196 0.6613333
##
     0.10
           0.20
                   0.9027745 0.9529412 0.6373333
##
     0.10
           0.25
                    0.9026307 0.9568627
                                         0.6053333
##
     0.10
           0.30
                    0.9026307 0.9568627
                                          0.5650000
##
     0.15
           0.05
                    0.9056176 0.9254902
                                         0.7253333
##
     0.15
           0.10
                    0.9037157
                              0.9411765
                                         0.6933333
##
     0.15
           0.15
                    0.9019771 0.9490196
                                         0.6533333
##
     0.15
           0.20
                    0.9018333 0.9568627
                                         0.6133333
##
           0.25
                    0.9008072 0.9568627 0.5730000
     0.15
     0.15
           0.30
                   0.8992386 0.9568627
##
                                         0.5086667
##
     0.20
           0.05
                   0.9037353 0.9215686 0.7253333
##
     0.20
           0.10
                   0.9027680 0.9411765
                                         0.6773333
##
     0.20
           0.15
                    0.9011993 0.9529412
                                         0.6533333
     0.20
##
           0.20
                    0.8990752 0.9568627
                                         0.5810000
##
     0.20
            0.25
                    0.8971928 0.9568627 0.5246667
```

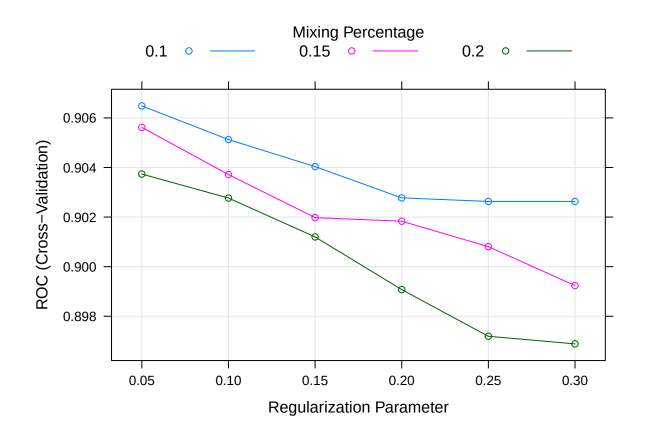
```
##
     0.20
            0.30
                    0.8968856 0.9568627 0.4446667
##
## ROC was used to select the optimal model using the largest value.
## The final values used for the model were alpha = 0.1 and lambda = 0.05.
## $model2
##
  glmnet
##
## 379 samples
##
    19 predictor
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 304, 303, 303, 303, 303
## Resampling results across tuning parameters:
##
##
            lambda
     alpha
                    ROC
                                Sens
                                           Spec
##
     0.10
            0.05
                    0.9011961
                                0.9215686
                                           0.6930000
##
     0.10
            0.10
                    0.9031307
                                0.9372549
                                           0.6770000
##
     0.10
            0.15
                    0.9031438
                                0.9490196
                                           0.6610000
##
     0.10
                    0.9033137
                                0.9490196
            0.20
                                           0.6366667
##
     0.10
            0.25
                    0.9041176
                                0.9529412
                                           0.5876667
                                0.9529412
##
     0.10
            0.30
                    0.9044379
                                           0.5636667
##
     0.15
            0.05
                    0.9012288
                                0.9254902
                                           0.6933333
##
     0.15
            0.10
                    0.9044052
                                0.9411765
                                           0.6853333
##
            0.15
                    0.9045882
     0.15
                                0.9450980
                                           0.6526667
##
     0.15
            0.20
                    0.9050850
                                0.9529412
                                           0.5880000
##
     0.15
            0.25
                    0.9044575
                               0.9529412
                                           0.5636667
##
     0.15
            0.30
                    0.9034118
                                0.9607843
                                           0.5316667
##
     0.20
            0.05
                    0.9015556
                                0.9294118
                                           0.7013333
##
     0.20
            0.10
                    0.9039477
                                0.9372549
                                           0.6770000
##
     0.20
            0.15
                    0.9052614
                                0.9411765
                                           0.6363333
     0.20
##
            0.20
                    0.9042222
                                0.9529412
                                           0.5956667
##
     0.20
            0.25
                    0.9026601
                                0.9568627
                                           0.5556667
##
     0.20
                                0.9647059
            0.30
                    0.9014706
                                           0.4590000
##
## ROC was used to select the optimal model using the largest value.
## The final values used for the model were alpha = 0.2 and lambda = 0.15.
##
## $model3
   glmnet
##
##
## 379 samples
##
    27 predictor
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 304, 303, 303, 303, 303
## Resampling results across tuning parameters:
##
##
     alpha lambda ROC
                                Sens
                                           Spec
##
     0.10
            0.05
                    0.8999869 0.9058824 0.6930000
```

```
##
     0.10
            0.10
                    0.9050327 0.9176471
                                           0.6930000
##
     0.10
            0.15
                    0.9045882 0.9294118
                                           0.6686667
##
     0.10
            0.20
                    0.9049150 0.9372549
                                           0.6366667
##
     0.10
            0.25
                    0.9041111
                               0.9450980
                                           0.6206667
##
     0.10
            0.30
                    0.9040980
                               0.9529412
                                           0.5806667
##
     0.15
            0.05
                    0.9015686 0.9058824
                                           0.6930000
##
     0.15
            0.10
                    0.9049085 0.9176471
                                           0.6930000
##
     0.15
            0.15
                    0.9038105
                               0.9294118
                                           0.6526667
##
     0.15
            0.20
                    0.9033203
                                0.9411765
                                           0.6286667
##
     0.15
            0.25
                    0.9032941
                               0.9529412
                                           0.5806667
##
     0.15
            0.30
                    0.9017190
                                0.9529412
                                           0.5323333
##
     0.20
            0.05
                    0.9031699
                               0.9098039
                                           0.6930000
##
     0.20
            0.10
                    0.9046144
                               0.9215686
                                           0.6930000
            0.15
##
     0.20
                    0.9025294
                               0.9333333
                                           0.6366667
##
     0.20
            0.20
                    0.9025359
                                0.9529412
                                           0.6126667
##
     0.20
            0.25
                    0.9009542
                                0.9529412
                                           0.5483333
##
     0.20
            0.30
                    0.9011111 0.9568627
                                           0.4603333
##
## ROC was used to select the optimal model using the largest value.
  The final values used for the model were alpha = 0.1 and lambda = 0.1.
##
## $model4
  glmnet
##
##
## 379 samples
##
    32 predictor
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 304, 303, 303, 303, 303
## Resampling results across tuning parameters:
##
##
     alpha lambda
                    ROC
                                Sens
                                           Spec
##
     0.10
            0.05
                    0.9597255
                                0.9529412
                                           0.7803333
##
     0.10
            0.10
                    0.9595621
                               0.9607843
                                           0.7403333
##
     0.10
            0.15
                    0.9592288 0.9647059
                                           0.7163333
##
     0.10
            0.20
                    0.9587451
                               0.9647059
                                           0.7083333
##
     0.10
            0.25
                    0.9590784
                                0.9725490
                                           0.7000000
##
            0.30
     0.10
                    0.9590784
                               0.9803922
                                           0.6920000
            0.05
##
     0.15
                    0.9597320
                               0.9490196
                                           0.7723333
##
     0.15
            0.10
                    0.9597190 0.9607843
                                           0.7323333
##
     0.15
            0.15
                    0.9597124
                               0.9647059
                                           0.7163333
##
            0.20
                    0.9582745 0.9686275
     0.15
                                           0.7000000
##
     0.15
            0.25
                    0.9573203
                               0.9843137
                                           0.6920000
                                0.9843137
##
     0.15
            0.30
                    0.9573268
                                           0.6513333
##
     0.20
            0.05
                    0.9587647
                                0.9490196
                                           0.7723333
##
     0.20
            0.10
                    0.9595556
                                0.9568627
                                           0.7403333
##
     0.20
            0.15
                    0.9585948
                                0.9647059
                                           0.7163333
##
     0.20
            0.20
                    0.9579673
                                0.9764706
                                           0.700000
##
     0.20
            0.25
                    0.9582876
                                0.9843137
                                           0.6430000
##
     0.20
            0.30
                    0.9594052
                               0.9882353
                                           0.6270000
##
```

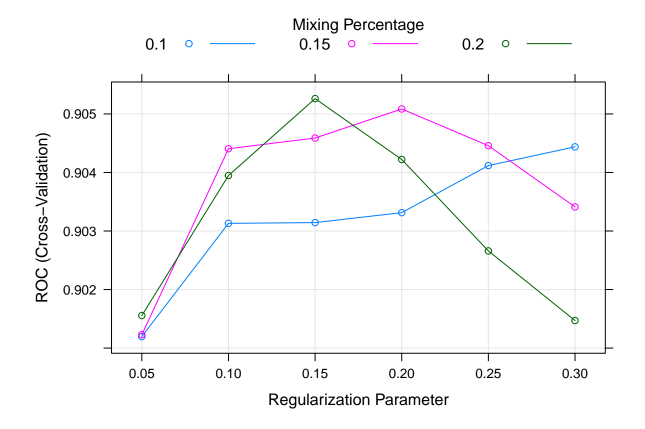
## ROC was used to select the optimal model using the largest value.

## The final values used for the model were alpha = 0.15 and lambda = 0.05.

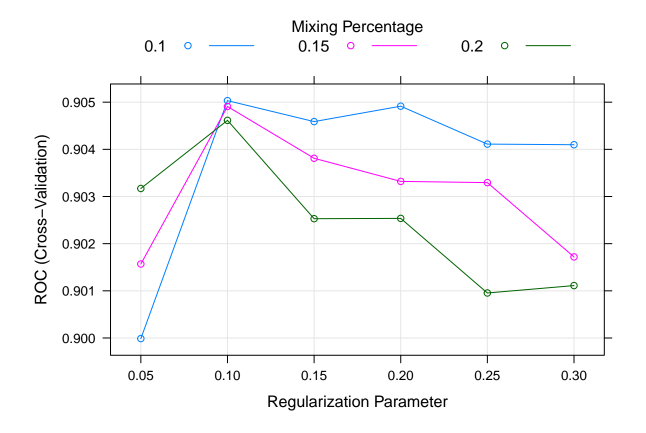
#Plot parameter tuning of model 1
plot(models\_EN[[1]])



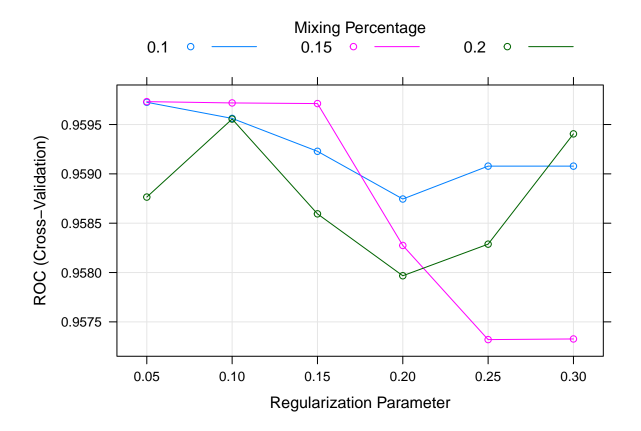
#Plot parameter tuning of model 2
plot(models\_EN[[2]])



#Plot parameter tuning of model 3
plot(models\_EN[[3]])



#Plot parameter tuning of model 4
plot(models\_EN[[4]])



#### Random Forest

Finally, we will fit a model of the CART family.

```
# Set the seed to ensure reproducibility
set.seed(504)
# Train the random forest model for all four sets of data
models_rf = list()
for (model in list(model1_vars, model2_vars, model3_vars, model4_vars)){
  rf_model = caret::train(
   pcos ~ .,
    data = chained_train %>%
      dplyr::select(all_of(model)),
    method = "ranger",
    num.trees = 500,
    tuneLength = 5,
    metric = "Sens",
    trControl = fitControl,
    importance="impurity")
  models_rf = append(models_rf, list(rf_model))
}
names(models_rf) = c("model1", "model2", "model3", "model4")
```

#### models\_rf

```
## $model1
## Random Forest
##
## 379 samples
##
   11 predictor
##
    2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 304, 303, 303, 303, 303
## Resampling results across tuning parameters:
##
##
    mtry
          splitrule
                      ROC
                                 Sens
                                            Spec
##
     2
          gini
                      0.9067255 0.9098039
                                            0.7020000
##
          extratrees 0.9043922 0.9137255 0.7260000
##
     4
                      gini
##
      4
          extratrees 0.8933203 0.8862745
                                            0.7176667
##
     6
          gini
                      0.8884575
                                0.8745098
                                           0.7263333
##
     6
          extratrees 0.8869673
                                0.8823529
                                           0.7256667
##
     8
                      0.8834641
                                 0.8627451
                                            0.7180000
          gini
##
     8
          extratrees 0.8850458
                                 0.8627451
                                            0.7340000
##
                      0.8799837 0.8549020
    11
          gini
                                            0.7183333
##
          extratrees 0.8803072 0.8588235
                                           0.7173333
##
## Tuning parameter 'min.node.size' was held constant at a value of 1
## Sens was used to select the optimal model using the largest value.
## The final values used for the model were mtry = 2, splitrule = extratrees
   and min.node.size = 1.
##
## $model2
## Random Forest
##
## 379 samples
   19 predictor
##
    2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 304, 303, 303, 303
## Resampling results across tuning parameters:
##
##
    mtry
          splitrule
                                 Sens
                                            Spec
##
     2
                      0.9013791 0.9215686 0.6366667
          gini
##
          extratrees 0.9013203 0.9176471 0.7016667
##
     6
          gini
                      0.8963399 0.8901961 0.6850000
##
     6
          extratrees 0.8971569 0.8980392 0.7176667
##
                      0.8883954 0.8862745
    10
          gini
                                            0.7010000
##
    10
          extratrees 0.8913497
                                0.8823529
                                            0.7420000
##
    14
                      0.8824967 0.8862745
                                           0.6853333
##
    14
          extratrees 0.8836863 0.8784314 0.7340000
##
                      0.8795000 0.8823529 0.6770000
    19
          gini
```

```
##
           extratrees 0.8864706 0.8823529 0.7423333
##
## Tuning parameter 'min.node.size' was held constant at a value of 1
## Sens was used to select the optimal model using the largest value.
## The final values used for the model were mtry = 2, splitrule = gini
   and min.node.size = 1.
##
## $model3
## Random Forest
##
## 379 samples
##
   27 predictor
##
    2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 303, 303, 304
## Resampling results across tuning parameters:
##
##
     mtry
          splitrule
                       ROC
                                  Sens
                                             Spec
##
     2
           gini
                       0.9082157 0.9294118 0.6053333
##
      2
           extratrees 0.9030458 0.9333333 0.6703333
##
     8
                       0.8923301 0.8980392 0.6943333
           gini
          extratrees 0.8944020 0.8980392
##
     8
                                             0.6783333
##
     14
           gini
                       0.8788856 0.8862745
                                            0.7103333
##
     14
           extratrees 0.8931830 0.8862745
                                             0.7100000
##
     20
                       0.8764902 0.8862745
           gini
                                             0.7023333
##
     20
           extratrees 0.8860131
                                 0.8862745
                                             0.6860000
     27
                       0.8673660 0.8627451
##
                                            0.6863333
##
     27
           extratrees 0.8798007 0.8823529
                                            0.6696667
##
## Tuning parameter 'min.node.size' was held constant at a value of 1
## Sens was used to select the optimal model using the largest value.
## The final values used for the model were mtry = 2, splitrule = extratrees
##
   and min.node.size = 1.
##
## $model4
## Random Forest
##
## 379 samples
   32 predictor
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (5 fold)
## Summary of sample sizes: 303, 303, 303, 304
## Resampling results across tuning parameters:
##
                       ROC
##
           splitrule
     mtry
                                  Sens
                                             Spec
##
     2
                       0.9608105
                                  0.9686275
                                             0.7433333
           gini
##
      2
           extratrees 0.9599477
                                 0.9490196
                                             0.7026667
##
     9
                       0.9513007 0.9490196
                                            0.8150000
##
     9
           extratrees 0.9595131 0.9333333
                                            0.7913333
##
     17
           gini
                       0.9432712 0.9372549 0.8150000
```

```
##
    17
          extratrees 0.9565098 0.9294118 0.7990000
##
    24
          gini
                      0.9419216 0.9372549 0.8070000
##
    24
          extratrees 0.9556340 0.9215686 0.8070000
##
    32
                      0.9413399 0.9294118 0.7906667
##
    32
          extratrees 0.9578758 0.9254902 0.8310000
##
## Tuning parameter 'min.node.size' was held constant at a value of 1
## Sens was used to select the optimal model using the largest value.
## The final values used for the model were mtry = 2, splitrule = gini
  and min.node.size = 1.
```

The number of trees was kept at the default of 500, as increases in the number of trees to 1000, 5000 and 10000 improved model performance by less than 1% for ROC, sensitivity and specificity. However, the processing time was increased significantly and thus the default value was not changed. Main tuning parameters were determined using the inbuilt 'tuneLength' parameter by trying different default grid values for the main parameters, which were optimized to 'sensitivity' as per the aims of this project. Further detail and justification for the metrics used in this project can be found in the Final Report in this repository.

## Model comparison

#### Test the performance in the validation set

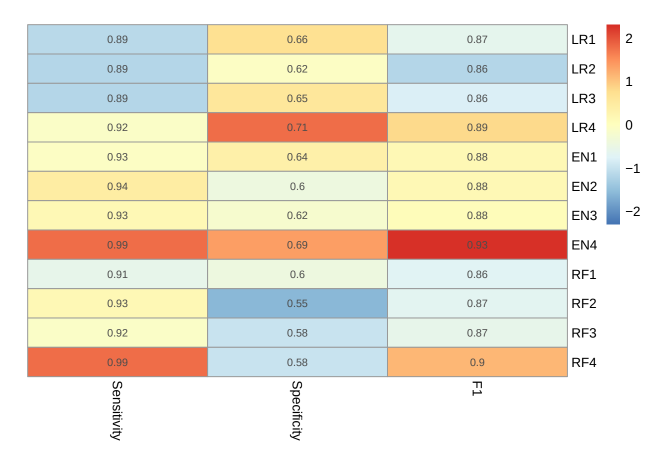
For this step, we will first fit the models in the validation data set to get the performance metrics.

We will concatenate all the models and estimate their performance in the validation set.

```
#Get the outcomes in the validation set.
outcomes = list(valid %>%
                  dplyr::select(all_of(model1_vars)) %>%
                  drop_na() %>%
                  pull(pcos),
                valid %>%
                  dplyr::select(all_of(model2_vars)) %>%
                  drop_na() %>%
                  pull(pcos),
                valid %>%
                  dplyr::select(all_of(model3_vars)) %>%
                  drop_na() %>%
                  pull(pcos),
                valid %>%
                  dplyr::select(all_of(model4_vars)) %>%
                  drop na() %>%
                  pull(pcos))
models_aggregated = c(models_logreg, models_EN, models_rf)
#Generate prediction metrics
pred_models = models_aggregated %>%
  purrr::map(\(x) predict(object = x, newdata = valid, type = "raw")) %>%
  purrr::map2(c(rep(outcomes,3)), \(x,y) caret::confusionMatrix(x,y))
#Create data frame with the metrics
(metrics = tibble(Method = c(rep("LR",4),
```

```
## # A tibble: 12 x 5
     Method Model Specificity Sensitivity
##
                                        F1
##
     <chr> <chr>
                     <dbl>
                            <dbl> <dbl>
## 1 LR
                      0.660
                               0.890 0.866
           1
## 2 LR
           2
                      0.623
                               0.888 0.856
                      0.654
                               0.887 0.862
## 3 LR
           3
                      0.712
                                0.925 0.895
## 4 LR
          4
## 5 EN
                      0.642
                                0.927 0.882
          1
## 6 EN
          2
                      0.604
                                0.944 0.882
## 7 EN
          3
                      0.615
                                0.934 0.88
## 8 EN
          4
                     0.692
                               0.991 0.925
## 9 RF
                                0.908 0.865
           1
                     0.604
## 10 RF
           2
                               0.935 0.866
                    0.547
## 11 RF
                      0.577
                                0.925 0.867
## 12 RF
           4
                      0.577
                                0.991 0.901
```

```
pheatmap(metrics %>%
          unite(name, Method, Model, sep = "") %>%
          column_to_rownames(var= "name") %>%
          select(c(Sensitivity, Specificity, F1)),
          cluster_rows = F,
          cluster_cols = F,
          scale = "column",
          display_numbers = round(metrics %>%
          select(c(Sensitivity, Specificity, F1)),2)
          )
```



In the plot above, we can observe that models 1,2 and 3 have a very similar performance in all of the methods applied. Model 4 has consistently a better performance than 1/2/3, which suggests that ultrasound variables are informative. However, getting that information usually requires a more specialized medical equipement, which is not that easily accessible.

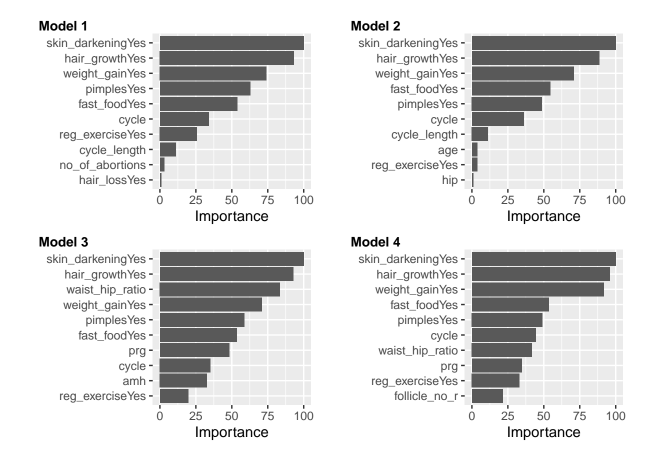
If we look at Sensitivity and F1, we can see that model 1 usually outperforms models 2 and 3. Since the aim of our project is to create a model that balances performance and easily collected variables, model 1 is the best option.

Furthermore, out of all of the models 1, the method that creates the best model is Elastic Net based on the metrics. Therefore, we will propose it as the final model in our project.

### Explore EN model 1

# Variable importance

Next, we will explore the most important variables in the elastic net models



Effect of class imbalance on the best model (if time allows)