Aprendizagem Estatística em Altas Dimensões [MAE0501/MAE5904/IBI5904]

Ícaro Maia Santos de Castro¹ Rayssa de Carvalho Roberto² Rodrigo Aoyama Nakahara³ Rodrigo Araujo⁴ Vitor Hugo Vieira de Lima⁵

Novembro de 2020

Sumário

Leitura dos Dados	. 2
Análise Descritiva	. 2
Análise Discriminante	. 11
Modelagem	. 11
Testes	
Regressão Logística	. 28
Modelagem	
Teste	
Random Forest	. 40
Modelagem	
Árvores de decisão geradas	
Teste	
Support Vector Machine	. 51
Modelagem	
Testes	

 $^{^1\}mathrm{N\'umero}$ USP: 11866921

 $^{^2\}mathrm{N\'umero}$ USP: 10940828

 $^{^3\}mathrm{N\'umero}$ USP: 3510922

⁴Número USP: 9299208

⁵Número USP: 10263886

Leitura dos Dados

```
diabetes <- read_csv("diabetes.csv")</pre>
## Parsed with column specification:
## cols(
    Pregnancies = col_double(),
##
     Glucose = col double(),
##
     BloodPressure = col_double(),
##
     SkinThickness = col_double(),
##
     Insulin = col_double(),
##
     BMI = col_double(),
##
##
     DiabetesPedigreeFunction = col_double(),
##
     Age = col_double(),
     Outcome = col_double()
##
## )
colnames(diabetes)[9] <- "diabetes"</pre>
diabetes$diabetes <- as.factor(diabetes$diabetes)</pre>
levels(diabetes$diabetes) <- c("No","Yes")</pre>
# Missings
diabetes[, 2:6] [diabetes[, 2:6] == 0] <- NA
head(diabetes) %>% kable(caption="Dados.")
```

Tabela 1: Dados.

Pregnancies	Glucose	BloodPressure	SkinThickness	Insulin	BMI	DiabetesPedigreeFunction	Age	diabetes
6	148	72	35	NA	33,6	0,63	50	Yes
1	85	66	29	NA	26,6	0,35	31	No
8	183	64	NA	NA	23,3	0,67	32	Yes
1	89	66	23	94	28,1	0,17	21	No
0	137	40	35	168	43,1	2,29	33	Yes
5	116	74	NA	NA	25,6	0,20	30	No

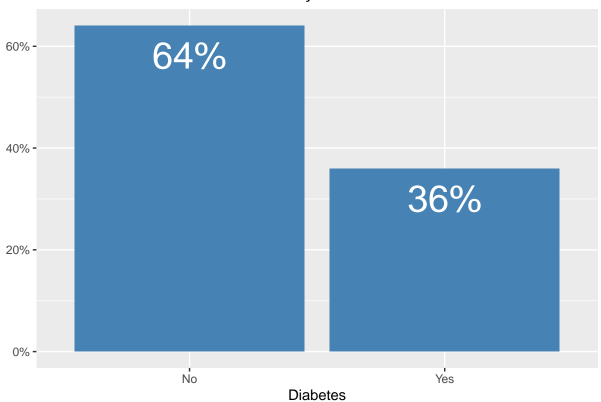
Análise Descritiva

```
# OBS: análise descritiva com base no conjunto de treinamento para evitar data snooping
library(dplyr)
library(caret)
library(lattice)
library(ggplot2)
library(GGally)
```

```
## Registered S3 method overwritten by 'GGally':
##
     method from
##
     +.gg
           ggplot2
library(ggcorrplot)
## Warning: package 'ggcorrplot' was built under R version 4.0.2
library(scales)
##
## Attaching package: 'scales'
## The following objects are masked from 'package:psych':
##
##
       alpha, rescale
## The following object is masked from 'package:purrr':
##
##
       discard
## The following object is masked from 'package:readr':
##
##
       col_factor
library(pROC)
## Warning: package 'pROC' was built under R version 4.0.2
## Type 'citation("pROC")' for a citation.
## Attaching package: 'pROC'
## The following object is masked from 'package:colorspace':
##
##
       coords
## The following objects are masked from 'package:stats':
##
##
       cov, smooth, var
# Distribuição da variável resposta
train %>% count(Diabetes = factor(diabetes)) %>% mutate(pct = prop.table(n)) %>%
  ggplot(aes(x = Diabetes, y = pct, fill = pct, label = scales::percent(pct))) +
```

```
geom_col(position = 'dodge', fill="steelblue") +
labs(title = "Classificação Diabetes", x = "Diabetes", y = "") +
geom_text(aes(label=scales::percent(pct)), vjust=1.6, color="white", size=10) +
scale_y_continuous(labels = scales::percent) +
theme(plot.title = element_text(hjust = 0.5), legend.title = element_blank())
```

Classificação Diabetes



```
# Matriz completa com dispersão, densidades, correlações, retas de regressão (com IC) e LOESS (com IC)

# Função auxiliar para curvas de regressão linear e LOESS nos gráficos abaixo
curvas <- function(data, mapping, ...){
    p <- ggplot(data = data, mapping = mapping) +
        geom_point(size = 0.5) +
        geom_smooth(method = loess, fill = "red", color = "red", ...) +
        geom_smooth(method = lm, fill = "blue", color = "blue", ...)
    p
}

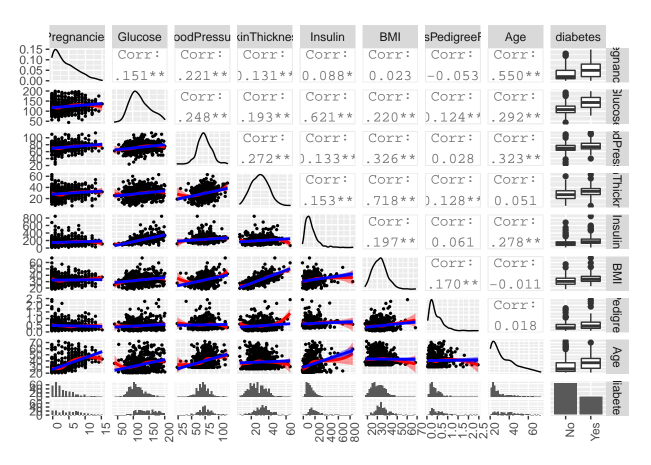
ggpairs(train, columns = 1:9, lower = list(continuous = curvas)) + # Obs: pode demorar para montar o gr
    theme(axis.text.x = element_text(angle = 90, hjust = 1))</pre>
```

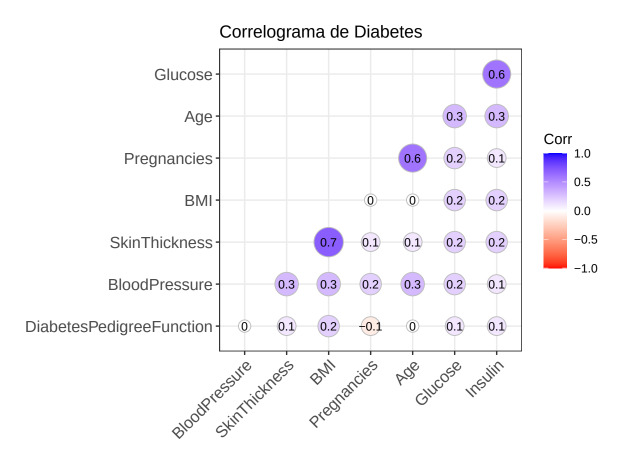
'geom_smooth()' using formula 'y ~ x'

'geom_smooth()' using formula 'y ~ x'

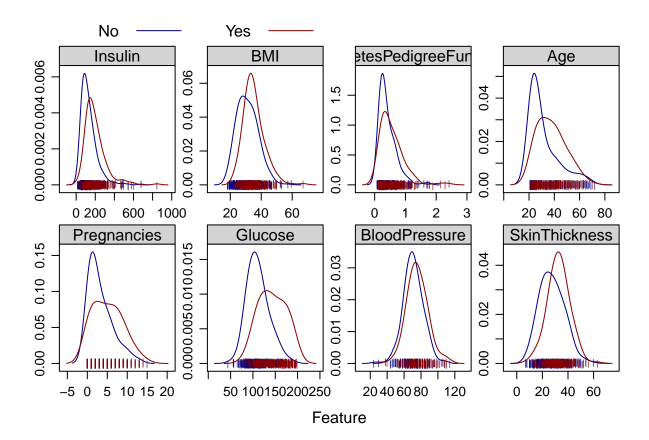
```
## 'geom_smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
## 'geom smooth()' using formula 'y ~ x'
## 'geom_smooth()' using formula 'y ~ x'
```

```
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
## 'stat_bin()' using 'bins = 30'. Pick better value with 'binwidth'.
```

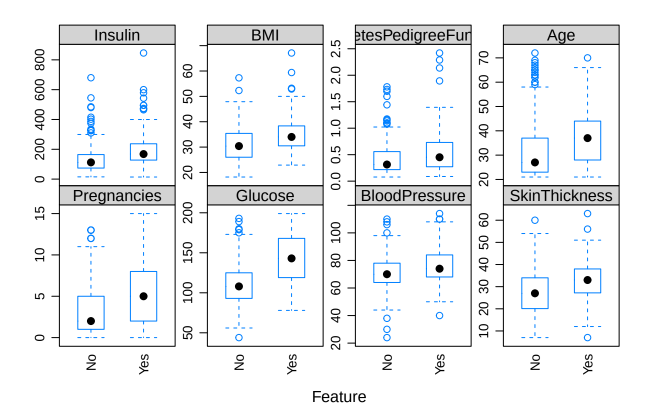




Warning in draw.key(simpleKey(...), draw = FALSE): not enough rows for columns



```
featurePlot(x = train[, 1:8], # Boxplots
    y = train$diabetes,
    plot = "box",
    scales = list(y = list(relation="free"), x = list(rot = 90)),
    layout = c(4,2),
    auto.key = list(columns = 4),
    par.settings = list(strip.background=list(col="lightgrey")))
```



Funções auxiliares

```
library(dplyr)
library(ggraph)

## Warning: package 'ggraph' was built under R version 4.0.3

library(igraph)

## Warning: package 'igraph' was built under R version 4.0.2

## Attaching package: 'igraph'

## The following objects are masked from 'package:dplyr':

## as_data_frame, groups, union

## The following objects are masked from 'package:purrr':

## compose, simplify
```

```
## The following object is masked from 'package:tidyr':
##
##
       crossing
## The following object is masked from 'package:tibble':
##
       as_data_frame
## The following objects are masked from 'package:stats':
##
       decompose, spectrum
## The following object is masked from 'package:base':
##
##
       union
plotaroc <- function(rocobj, titulo = "Curva ROC"){</pre>
  # Função que plota as curvas roc para os modelos ajustados
  b <- which.max(rocobj$sensitivities + rocobj$specificities)</pre>
  best <- round(c(rocobj$thresholds[b],rocobj$specificities[b],rocobj$sensitivities[b]), 3)
  pROC::ggroc(rocobj, col = "red", alpha = 0.5, size = 0.5) +
    theme_gray() +
    ggtitle(titulo) +
    geom_abline(intercept = 1, slope=1, linetype = "dashed") +
    labs(x="Especificidade", y = "Sensibilidade") +
    geom_point(data = tibble(Sensibilidade = best[2],
                              Especificidade = best[3]),
               mapping = aes(x=Sensibilidade, y=Especificidade),
               col = "black") +
    geom_text(mapping = aes(x = best[2] - 0.15,
                              y = best[3] - 0.05),
              label = paste( best[1], "(", best[2], ",", best[3], ")")) +
    geom text(mapping = aes(x = 0.5,
                             y = 0.01),
              label = paste("AUC: ", round(rocobj$auc,3)))
}
tree_func <- function(final_model,</pre>
                      tree num) {
  # get tree by index
  tree <- randomForest::getTree(final_model,</pre>
                                 k = tree_num,
                                 labelVar = TRUE) %>%
    tibble::rownames_to_column() %>%
    # make leaf split points to NA, so the Os won't get plotted
    mutate('split point' = ifelse(is.na(prediction), 'split point', NA))
  # prepare data frame for graph
  graph frame <- data.frame(from = rep(tree$rowname, 2),</pre>
                             to = c(tree$'left daughter', tree$'right daughter'))
```

```
# convert to graph and delete the last node that we don't want to plot
graph <- graph_from_data_frame(graph_frame) %>%
 delete vertices("0")
# set node labels
V(graph)$node_label <- gsub("_", " ", as.character(tree$'split var'))</pre>
V(graph)$leaf_label <- as.character(tree$prediction)</pre>
V(graph) $split <- as.character(round(tree $'split point', digits = 2))
# plot
plot <- ggraph(graph, 'dendrogram') +</pre>
 theme_bw() +
 geom_edge_link() +
 geom_node_point() +
  geom_node_text(aes(label = node_label), na.rm = TRUE, repel = TRUE) +
 geom_node_label(aes(label = split), vjust = 2, na.rm = TRUE, fill = "white") +
  geom_node_label(aes(label = leaf_label, fill = leaf_label), na.rm = TRUE,
                  repel = TRUE, colour = "white", fontface = "bold", show.legend = FALSE) +
 theme(panel.grid.minor = element_blank(),
        panel.grid.major = element_blank(),
        panel.background = element_blank(),
        plot.background = element_rect(fill = "white"),
       panel.border = element blank(),
        axis.line = element_blank(),
        axis.text.x = element_blank(),
        axis.text.y = element_blank(),
        axis.ticks = element_blank(),
        axis.title.x = element_blank(),
        axis.title.y = element_blank(),
        plot.title = element_text(size = 14))
return(plot)
```

Análise Discriminante

Modelagem

```
# Obs: possibilidades de modelos de AD: rda, lda, pda (acurácias iguais) e qda (pior)

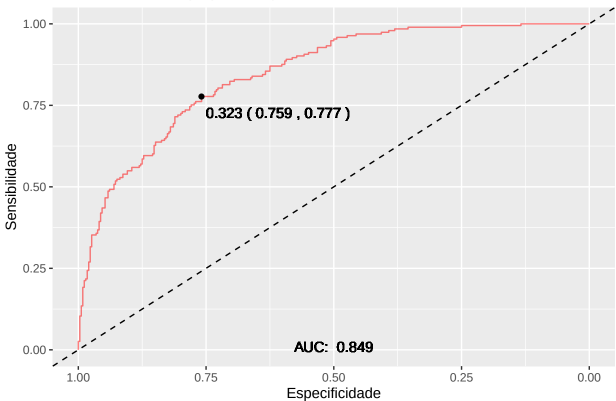
train.control <- caret::trainControl(method = "cv", number = 15, classProbs = TRUE) # Cross-validation

## ADL com dados imputados
set.seed(23)
modeloAD1 <- caret::train(diabetes ~ ., data = train, trControl = train.control, method = "lda")
print(modeloAD1)

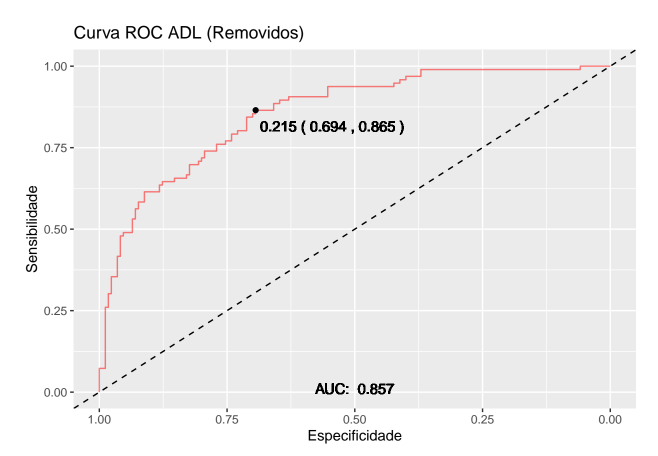
## Linear Discriminant Analysis</pre>
```

```
##
## 537 samples
     8 predictor
##
     2 classes: 'No', 'Yes'
##
##
## No pre-processing
## Resampling: Cross-Validated (15 fold)
## Summary of sample sizes: 501, 501, 501, 501, 501, 501, ...
## Resampling results:
##
##
     Accuracy
                Kappa
##
     0.7636508 0.4650271
rocAD1 <- roc(response = train$diabetes, predictor = predict(modeloAD1, train, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocAD1, titulo = "Curva ROC ADL (Imputados)")
```





```
## ADL sem missing
set.seed(23)
modeloAD2 <- caret::train(diabetes ~ ., data = train_without_NAs, trControl = train.control, method = "</pre>
print(modeloAD2)
## Linear Discriminant Analysis
## 266 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (15 fold)
## Summary of sample sizes: 248, 248, 249, 248, 247, 249, ...
## Resampling results:
##
    Accuracy Kappa
   0.7784199 0.5053595
##
rocAD2 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloAD2, train_without_NAs,</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocAD2, titulo = "Curva ROC ADL (Removidos)")
```



```
## AD Flexivel com dados imputados
set.seed(23)
modeloAD3 <- caret::train(diabetes ~ ., data = train, trControl = train.control, method = "fda")

## Loading required package: earth

## Warning: package 'earth' was built under R version 4.0.3

## Loading required package: plotmo

## Warning: package 'plotmo' was built under R version 4.0.3

## Loading required package: plotrix

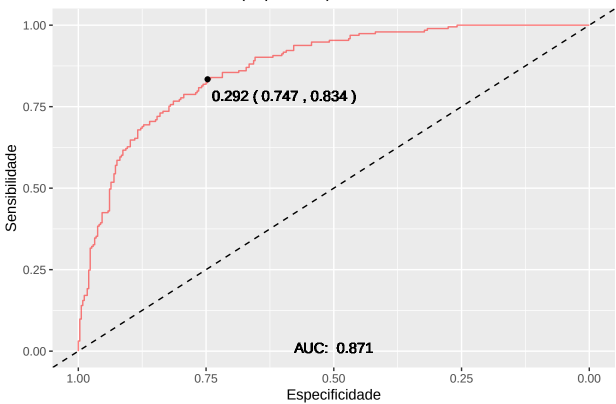
## Attaching package: 'plotrix'

## The following object is masked from 'package:scales':

## rescale</pre>
```

```
## The following object is masked from 'package:psych':
##
##
       rescale
## Loading required package: TeachingDemos
## Warning: package 'TeachingDemos' was built under R version 4.0.3
##
## Attaching package: 'TeachingDemos'
## The following objects are masked from 'package:Hmisc':
##
       cnvrt.coords, subplot
##
print(modeloAD3)
## Flexible Discriminant Analysis
##
## 537 samples
    8 predictor
##
     2 classes: 'No', 'Yes'
##
##
## No pre-processing
## Resampling: Cross-Validated (15 fold)
## Summary of sample sizes: 501, 501, 501, 501, 501, 501, ...
## Resampling results across tuning parameters:
##
##
    nprune Accuracy
                        Kappa
##
     2
            0.7450794 0.4129759
##
             0.7823280 0.5204850
     8
##
     15
             0.7823280 0.5192155
##
## Tuning parameter 'degree' was held constant at a value of 1
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were degree = 1 and nprune = 8.
rocAD3 <- roc(response = train$diabetes, predictor = predict(modeloAD3, train, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocAD3, titulo = "Curva ROC AD Flexível (Imputados)")
```

Curva ROC AD Flexível (Imputados)



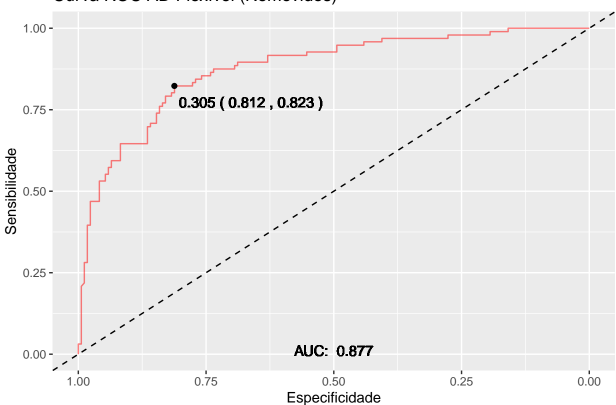
```
## AD Flexivel sem missing
set.seed(23)
modeloAD4 <- caret::train(diabetes ~ ., data = train_without_NAs, trControl = train.control, method = ".
print(modeloAD4)</pre>
```

```
## Flexible Discriminant Analysis
##
## 266 samples
##
     8 predictor
     2 classes: 'No', 'Yes'
##
##
## No pre-processing
## Resampling: Cross-Validated (15 fold)
## Summary of sample sizes: 248, 248, 249, 248, 247, 249, ...
## Resampling results across tuning parameters:
##
##
     nprune Accuracy
                        Kappa
     2
##
             0.7481137
                        0.4233678
##
             0.7516455
                        0.4422724
##
     15
             0.7446279
                        0.4240573
## Tuning parameter 'degree' was held constant at a value of 1
## Accuracy was used to select the optimal model using the largest value.
\#\# The final values used for the model were degree = 1 and nprune = 8.
```

```
rocAD4 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloAD4, train_without_NAs,
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases</pre>
```

Curva ROC AD Flexível (Removidos)

plotaroc(rocAD4, titulo = "Curva ROC AD Flexível (Removidos)")



```
## AD Quadrática com dados imputados
set.seed(23)
modeloAD5 <- caret::train(diabetes ~ ., data = train, trControl = train.control, method = "qda")
print(modeloAD5)</pre>
```

```
## Quadratic Discriminant Analysis
##
## 537 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (15 fold)
## Summary of sample sizes: 501, 501, 501, 501, 501, ...
## Resampling results:
##
```

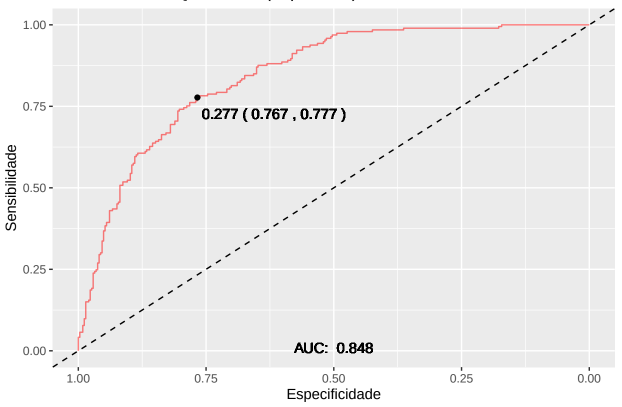
```
## Accuracy Kappa
## 0.7431217 0.4201363

rocAD5 <- roc(response = train$diabetes, predictor = predict(modeloAD5, train, type = "prob")[,2])

## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocAD5, titulo = "Curva ROC AD Quadrática (Imputados)")</pre>
```

Curva ROC AD Quadrática (Imputados)



```
## AD Quadrática sem missing
set.seed(23)
modeloAD6 <- caret::train(diabetes ~ ., data = train_without_NAs, trControl = train.control, method = "
print(modeloAD6)</pre>
```

```
## Quadratic Discriminant Analysis
##
## 266 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (15 fold)
```

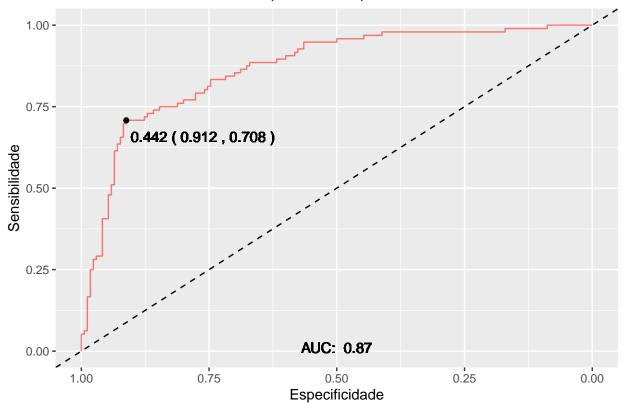
```
## Summary of sample sizes: 248, 248, 249, 248, 247, 249, ...
## Resampling results:
##
## Accuracy Kappa
## 0.7860681 0.5158632

rocAD6 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloAD6, train_without_NAs,

## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocAD6, titulo = "Curva ROC AD Quadrática (Removidos)")</pre>
```

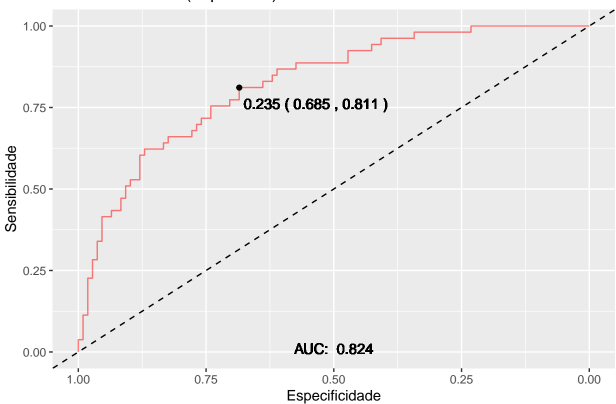
Curva ROC AD Quadrática (Removidos)



Testes

```
## ADL com dados imputados
rocAD1 <- roc(response = test$diabetes, predictor = predict(modeloAD1, test, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases</pre>
```

Curva ROC ADL (Imputados)

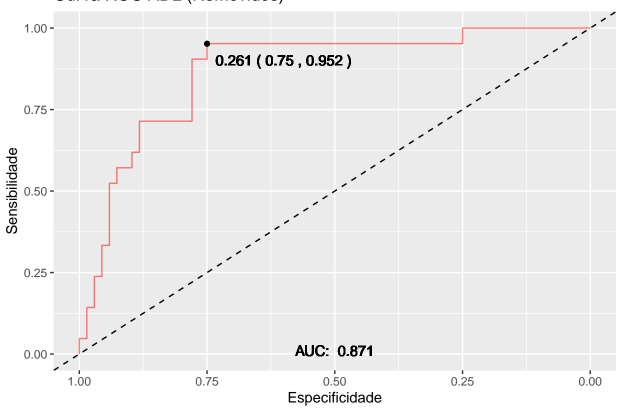


```
predictAD1 <- predict(modeloAD1, newdata = test)
confusionMatrix(predictAD1, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
          No 99
                  28
##
                  25
##
          Yes 9
##
                  Accuracy : 0.7702
##
##
                    95% CI: (0.6974, 0.8327)
       No Information Rate: 0.6708
##
       P-Value [Acc > NIR] : 0.003815
##
##
##
                     Kappa : 0.4274
##
##
    Mcnemar's Test P-Value: 0.003085
##
##
               Sensitivity: 0.9167
##
               Specificity: 0.4717
            Pos Pred Value: 0.7795
##
```

```
Neg Pred Value : 0.7353
##
                Prevalence: 0.6708
##
            Detection Rate: 0.6149
##
##
      Detection Prevalence: 0.7888
##
         Balanced Accuracy: 0.6942
##
##
          'Positive' Class: No
##
## ADL sem missing
rocAD2 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloAD2, test_without_NAs, ty</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocAD2, titulo = "Curva ROC ADL (Removidos)")
```

Curva ROC ADL (Removidos)

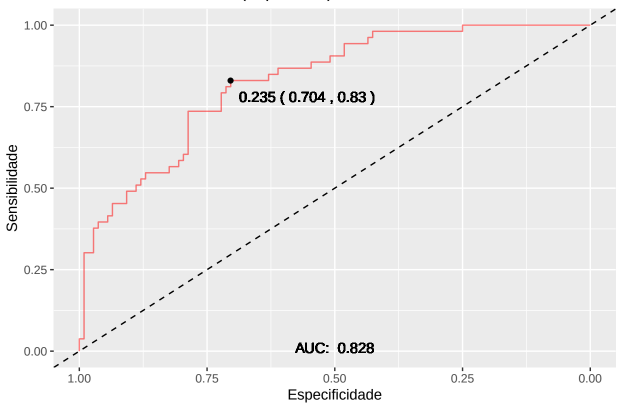


```
predictAD2 <- predict(modeloAD2, newdata = test_without_NAs)
confusionMatrix(predictAD2, test_without_NAs$diabetes)</pre>
```

Confusion Matrix and Statistics
##
Reference

```
## Prediction No Yes
         No 64 12
##
##
         Yes 4
##
##
                  Accuracy : 0.8202
##
                    95% CI: (0.7245, 0.8936)
##
      No Information Rate: 0.764
##
       P-Value [Acc > NIR] : 0.12910
##
##
                     Kappa: 0.4258
##
   Mcnemar's Test P-Value : 0.08012
##
##
##
               Sensitivity: 0.9412
##
               Specificity: 0.4286
##
            Pos Pred Value: 0.8421
##
            Neg Pred Value: 0.6923
##
                Prevalence: 0.7640
##
            Detection Rate: 0.7191
##
     Detection Prevalence: 0.8539
##
        Balanced Accuracy: 0.6849
##
##
          'Positive' Class : No
##
## ADL Flexivel com dados imputados
rocAD3 <- roc(response = test$diabetes, predictor = predict(modeloAD3, test, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocAD3, titulo = "Curva ROC AD Flexível (Imputados)")
```

Curva ROC AD Flexível (Imputados)



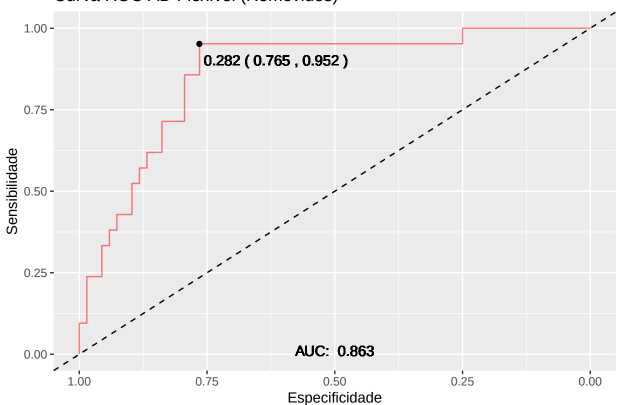
```
predictAD3 <- predict(modeloAD3, newdata = test)
confusionMatrix(predictAD3, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
          No 95 26
##
##
          Yes 13 27
##
                  Accuracy : 0.7578
##
                    95% CI : (0.6841, 0.8217)
##
##
       No Information Rate: 0.6708
       P-Value [Acc > NIR] : 0.01039
##
##
##
                     Kappa : 0.415
##
##
    Mcnemar's Test P-Value: 0.05466
##
               Sensitivity: 0.8796
##
##
               Specificity: 0.5094
##
            Pos Pred Value: 0.7851
            Neg Pred Value : 0.6750
##
##
                Prevalence: 0.6708
            Detection Rate: 0.5901
##
```

```
## Detection Prevalence : 0.7516
## Balanced Accuracy : 0.6945
##
## 'Positive' Class : No
##
## ADL Flexível sem missing
rocAD4 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloAD4, test_without_NAs, ty
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocAD4, titulo = "Curva ROC AD Flexível (Removidos)")</pre>
```

Curva ROC AD Flexível (Removidos)

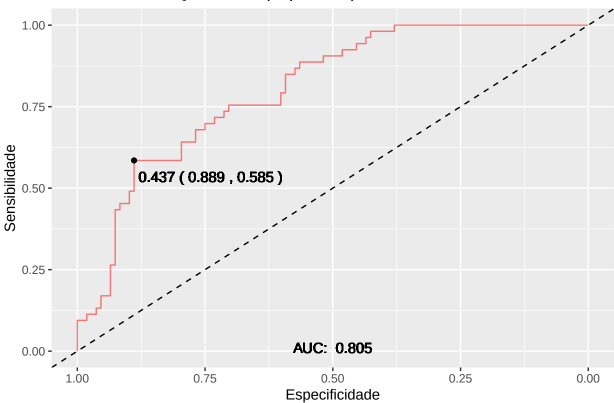


```
predictAD4 <- predict(modeloAD4, newdata = test_without_NAs)
confusionMatrix(predictAD4, test_without_NAs$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction No Yes
## No 61 11
## Yes 7 10
```

```
##
                  Accuracy : 0.7978
##
                    95% CI: (0.6993, 0.8755)
##
##
       No Information Rate: 0.764
       P-Value [Acc > NIR] : 0.2709
##
##
##
                     Kappa: 0.3996
##
##
    Mcnemar's Test P-Value: 0.4795
##
##
               Sensitivity: 0.8971
##
               Specificity: 0.4762
##
            Pos Pred Value: 0.8472
            Neg Pred Value: 0.5882
##
##
                Prevalence: 0.7640
            Detection Rate: 0.6854
##
##
      Detection Prevalence: 0.8090
         Balanced Accuracy: 0.6866
##
##
##
          'Positive' Class : No
##
## AD Quadrática com dados imputados
rocAD5 <- roc(response = test$diabetes, predictor = predict(modeloAD5, test, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases</pre>
plotaroc(rocAD5, titulo = "Curva ROC AD Quadrática (Imputados)")
```

Curva ROC AD Quadrática (Imputados)



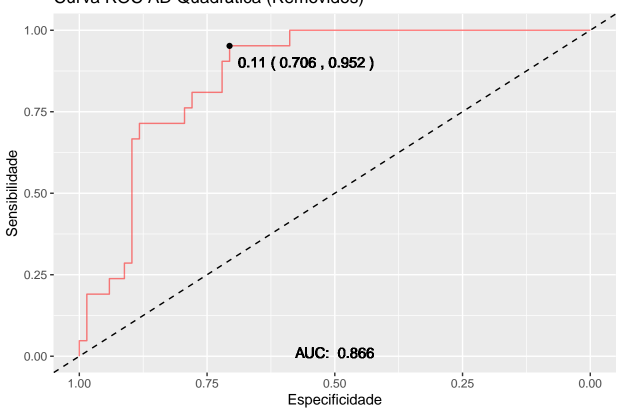
```
predictAD5 <- predict(modeloAD5, newdata = test)
confusionMatrix(predictAD5, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
             Reference
##
## Prediction No Yes
          No 96 25
##
##
          Yes 12 28
##
                  Accuracy : 0.7702
##
                    95% CI: (0.6974, 0.8327)
##
##
       No Information Rate: 0.6708
       P-Value [Acc > NIR] : 0.003815
##
##
##
                     Kappa : 0.445
##
##
    Mcnemar's Test P-Value: 0.048520
##
               Sensitivity: 0.8889
##
##
               Specificity: 0.5283
##
            Pos Pred Value: 0.7934
            Neg Pred Value : 0.7000
##
##
                Prevalence: 0.6708
            Detection Rate: 0.5963
##
```

```
## Detection Prevalence : 0.7516
## Balanced Accuracy : 0.7086
##
## 'Positive' Class : No
##
## AD Quadrática sem missing
rocAD6 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloAD6, test_without_NAs, ty
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocAD6, titulo = "Curva ROC AD Quadrática (Removidos)")</pre>
```

Curva ROC AD Quadrática (Removidos)



```
predictAD6 <- predict(modeloAD6, newdata = test_without_NAs)
confusionMatrix(predictAD6, test_without_NAs$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction No Yes
## No 60 6
## Yes 8 15
```

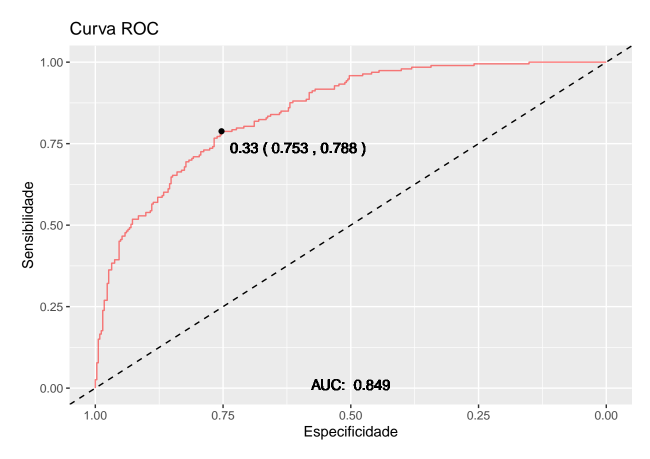
```
##
##
                  Accuracy : 0.8427
                    95% CI : (0.7502, 0.9112)
##
##
       No Information Rate: 0.764
##
       P-Value [Acc > NIR] : 0.04778
##
##
                     Kappa: 0.5776
##
##
   Mcnemar's Test P-Value: 0.78927
##
##
               Sensitivity: 0.8824
##
               Specificity: 0.7143
            Pos Pred Value: 0.9091
##
##
            Neg Pred Value: 0.6522
##
                Prevalence: 0.7640
##
            Detection Rate: 0.6742
##
     Detection Prevalence: 0.7416
##
         Balanced Accuracy: 0.7983
##
          'Positive' Class : No
##
##
```

Regressão Logística

Modelagem

```
library(caret)
train.control <- caret::trainControl(method = "cv", number = 10) # Cross-validation com k=10
## Logística simples com ligação logit / dados imputados
set.seed(23)
modeloRL1 <- caret::train(diabetes ~ ., data = train, trControl = train.control, method = "glm", family
print(modeloRL1)
## Generalized Linear Model
##
## 537 samples
##
    8 predictor
##
     2 classes: 'No', 'Yes'
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 483, 483, 484, 483, 482, 484, ...
## Resampling results:
##
##
     Accuracy
                Kappa
    0.7595623 0.4586088
##
```

```
rocRL1 <- roc(response = train$diabetes, predictor = predict(modeloRL1, train, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRL1)</pre>
```



```
## Logística simples com ligação logit / dados sem missing
set.seed(23)
modeloRL2 <- caret::train(diabetes ~ ., data = train_without_NAs, trControl = train.control, method = ",
print(modeloRL2)</pre>
```

```
## Generalized Linear Model
##
## 266 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 240, 240, 239, 239, 239, 239, ...
```

```
## Resampling results:
##
## Accuracy Kappa
## 0.7673789 0.4828079

rocRL2 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloRL2, train_without_NAs,

## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

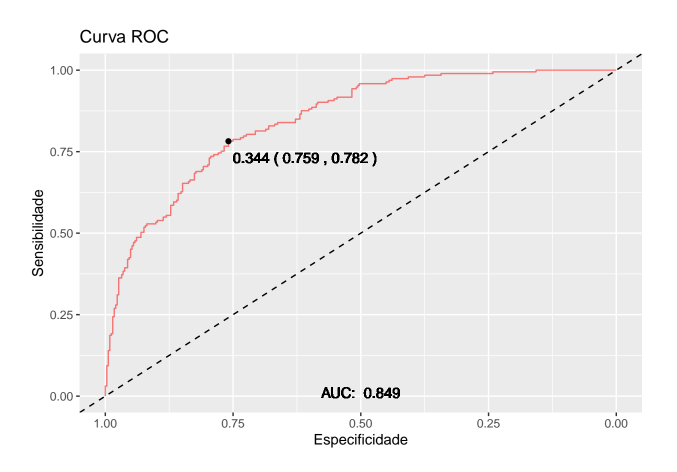
plotaroc(rocRL2)</pre>
```

Curva ROC 1.00 0.75 0.275 (0.706, 0.844) 0.25 0.00 AUC: 0.858 1.00 0.75 0.50 Especificidade

```
## Logistica regularizada / dados imputados
set.seed(23)
modeloRL3 <- caret::train(diabetes ~ ., data = train, trControl = train.control, method = "regLogistic"
print(modeloRL3)</pre>
```

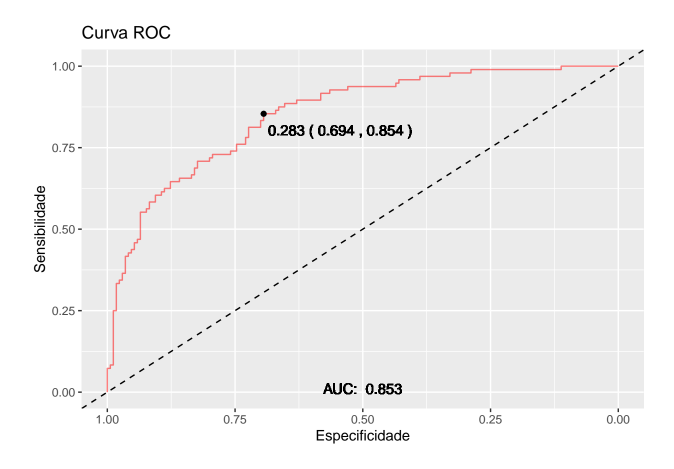
```
## Regularized Logistic Regression
##
## 537 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
```

```
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 483, 483, 484, 483, 482, 484, ...
## Resampling results across tuning parameters:
##
                     epsilon Accuracy
    cost loss
                                         Kappa
##
    0.5
                     0.001
                              0.7482765 0.4294592
          L1
##
                     0.010
    0.5
          L1
                              0.7426847 0.4156713
##
    0.5
          L1
                     0.100
                              0.7482053 0.4254680
##
    0.5
                     0.001
          L2_dual
                              0.6961737 0.2507155
##
    0.5
          L2_dual
                     0.010
                              0.6813925 0.2298081
##
    0.5
          L2_dual
                     0.100
                              0.6217813 0.1749702
##
    0.5
          L2_primal 0.001
                              0.7445715 0.4210244
##
    0.5
          L2_primal
                     0.010
                              0.6998126 0.3193385
##
    0.5
          L2_primal
                     0.100
                              0.6964195 0.3018360
##
    1.0
          L1
                     0.001
                              0.7558236 0.4469887
##
    1.0
         L1
                     0.010
                              0.7539369 0.4431172
##
    1.0
          L1
                     0.100
                              0.7426161 0.4055951
##
    1.0
          L2_dual
                     0.001
                              0.6039839 0.1296527
                              0.6144038 0.2065002
##
    1.0
          L2 dual
                     0.010
##
    1.0
         L2_dual
                     0.100
                              0.6311416 0.1155712
##
    1.0
          L2_primal 0.001
                              0.7464583 0.4229977
##
    1.0
          L2_primal 0.010
                              0.6998126 0.3193385
##
    1.0
          L2_primal 0.100
                              0.6964195 0.3018360
##
    2.0
                     0.001
          L1
                              0.7595623 0.4554919
##
    2.0 L1
                     0.010
                              0.7558236 0.4468964
##
    2.0
          L1
                     0.100
                              0.7538670 0.4388071
##
         L2_dual
                     0.001
    2.0
                              0.6149863 0.1897020
##
    2.0
         L2_dual
                     0.010
                              0.5702999 0.1900379
                     0.100
##
    2.0
          L2_dual
                              0.5902649 0.1230814
##
    2.0
          L2_primal 0.001
                              0.7500934 0.4318992
##
    2.0
          L2_primal 0.010
                              0.7070802 0.3426133
##
    2.0
          L2_primal 0.100
                              0.6964195 0.3018360
##
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were cost = 2, loss = L1 and epsilon
## = 0.001.
rocRL3 <- roc(response = train$diabetes, predictor = predict(modeloRL3, train, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRL3)
```



```
## Logística regularizada / dados sem missing
set.seed(23)
modeloRL4 <- caret::train(diabetes ~ ., data = train_without_NAs, trControl = train.control, method = "
print(modeloRL4)
## Regularized Logistic Regression
##
## 266 samples
##
    8 predictor
     2 classes: 'No', 'Yes'
##
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 240, 240, 239, 239, 239, 239, ...
## Resampling results across tuning parameters:
##
##
     cost loss
                      epsilon Accuracy
                                          Kappa
##
     0.5
           L1
                      0.001
                               0.7598291 0.4596795
##
    0.5
          L1
                      0.010
                               0.7636752 0.4669602
##
     0.5
                      0.100
                               0.7488604 0.4299137
##
    0.5
          L2_dual
                      0.001
                               0.6394587 0.1554546
##
     0.5
          L2_dual
                      0.010
                               0.6065527 0.1369186
##
    0.5
          L2_dual
                      0.100
                               0.6501425 0.1759711
##
    0.5
          L2_primal 0.001
                               0.7263533 0.3848223
           L2_primal
##
     0.5
                      0.010
                               0.7149573 0.3616567
```

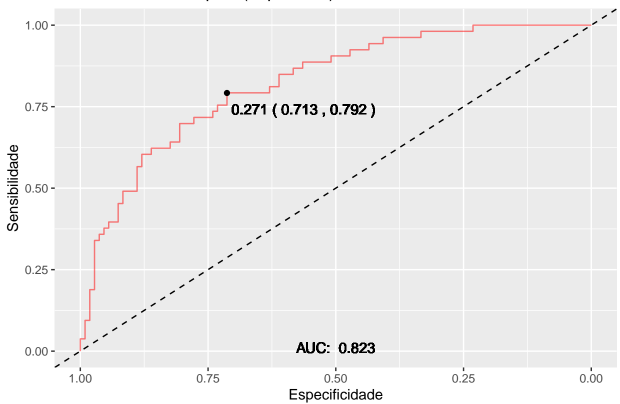
```
##
    0.5
          L2_primal 0.100
                            0.6807692 0.2740010
                    0.001
##
    1.0 L1
                            0.7709402 0.4834174
    1.0 L1
                    0.010
                            0.7747863 0.4903988
##
##
                    0.100
    1.0
          L1
                            0.7522792 0.4352616
                    0.001
##
    1.0
         L2_dual
                            0.6249288 0.1842344
##
    1.0 L2_dual
                   0.010
                            0.5915954 0.1786622
                            0.6173789 0.1770171
##
    1.0 L2 dual
                    0.100
          L2_primal 0.001
##
    1.0
                            0.7565527 0.4527102
                            0.7339031 0.4037206
##
    1.0
          L2_primal 0.010
##
    1.0
          L2_primal 0.100
                            0.6807692 0.2740010
##
    2.0
                    0.001
                            0.7710826 0.4902196
##
    2.0
                            0.7747863 0.4970394
          L1
                    0.010
##
                            0.7712251 0.4816945
    2.0
         L1
                    0.100
##
    2.0 L2_dual
                    0.001
                            0.5992877 0.1660801
##
    2.0
          L2_dual
                    0.010
                            0.6019943 0.2116899
##
    2.0
          L2_dual
                    0.100
                            0.6427350 0.1862283
##
    2.0
          L2_primal 0.001
                            0.7601140 0.4623563
##
    2.0
          L2_primal 0.010
                             0.7487179 0.4343284
##
    2.0
          L2_primal 0.100
                             0.6807692 0.2740010
##
## Accuracy was used to select the optimal model using the largest value.
## The final values used for the model were cost = 1, loss = L1 and epsilon = 0.01.
rocRL4 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloRL4, train_without_NAs,</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRL4)
```



Teste

```
## Regressão Logística Simples com dados imputados
rocRL1 <- roc(response = test$diabetes, predictor = predict(modeloRL1, test, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRL1, titulo = "Curva ROC RL Simples (Imputados)")</pre>
```

Curva ROC RL Simples (Imputados)



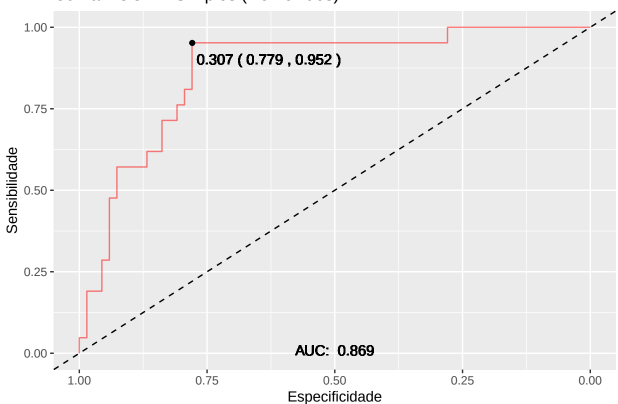
```
predictRL1 <- predict(modeloRL1, newdata = test)
confusionMatrix(predictRL1, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
          No 99
                  27
##
##
          Yes 9
                  26
##
##
                  Accuracy : 0.7764
                    95% CI : (0.7041, 0.8382)
##
##
       No Information Rate: 0.6708
       P-Value [Acc > NIR] : 0.002203
##
##
##
                     Kappa : 0.4458
##
##
    Mcnemar's Test P-Value: 0.004607
##
               Sensitivity: 0.9167
##
               Specificity: 0.4906
##
##
            Pos Pred Value: 0.7857
            Neg Pred Value : 0.7429
##
##
                Prevalence: 0.6708
            Detection Rate: 0.6149
##
```

```
## Detection Prevalence : 0.7826
## Balanced Accuracy : 0.7036
##
## 'Positive' Class : No
##
## Regressão Logística Simples sem missing
rocRL2 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloRL2, test_without_NAs, ty
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocRL2, titulo = "Curva ROC RL Simples (Removidos)")</pre>
```

Curva ROC RL Simples (Removidos)

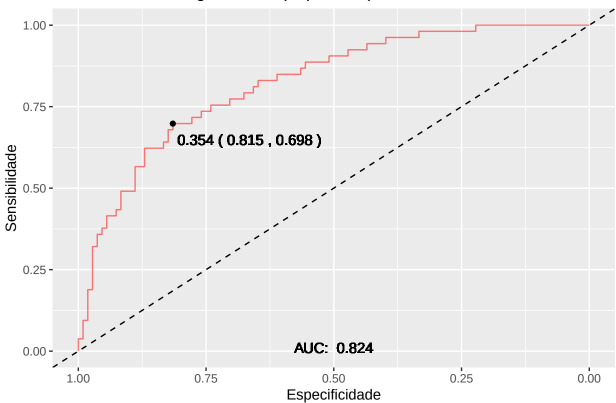


```
predictRL2 <- predict(modeloRL2, newdata = test_without_NAs)
confusionMatrix(predictRL2, test_without_NAs$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction No Yes
## No 64 11
## Yes 4 10
```

```
##
##
                  Accuracy : 0.8315
                    95% CI: (0.7373, 0.9025)
##
##
       No Information Rate: 0.764
       P-Value [Acc > NIR] : 0.08127
##
##
##
                     Kappa: 0.4717
##
##
    Mcnemar's Test P-Value: 0.12134
##
##
               Sensitivity: 0.9412
##
               Specificity: 0.4762
##
            Pos Pred Value: 0.8533
            Neg Pred Value: 0.7143
##
##
                Prevalence: 0.7640
            Detection Rate: 0.7191
##
##
      Detection Prevalence: 0.8427
         Balanced Accuracy: 0.7087
##
##
          'Positive' Class : No
##
##
## Regressão Logística Regularizada com dados imputados
rocRL3 <- roc(response = test$diabetes, predictor = predict(modeloRL3, test, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRL3, titulo = "Curva ROC RL Regularizada (Imputados)")
```

Curva ROC RL Regularizada (Imputados)



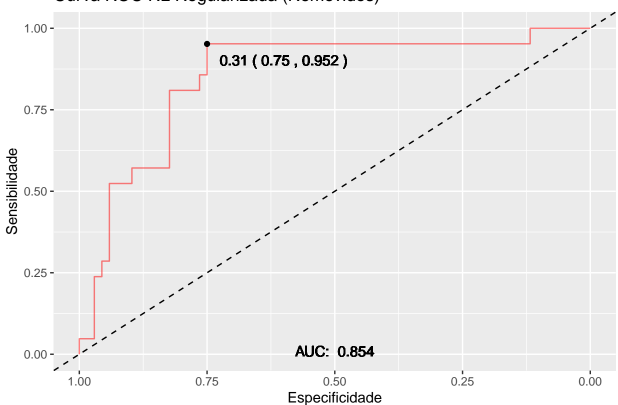
```
predictRL3 <- predict(modeloRL3, newdata = test)
confusionMatrix(predictRL3, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
          No 99
                  28
##
##
          Yes 9
##
                  Accuracy : 0.7702
##
                    95% CI: (0.6974, 0.8327)
##
##
       No Information Rate: 0.6708
       P-Value [Acc > NIR] : 0.003815
##
##
##
                     Kappa : 0.4274
##
##
    Mcnemar's Test P-Value: 0.003085
##
               Sensitivity: 0.9167
##
##
               Specificity: 0.4717
##
            Pos Pred Value: 0.7795
            Neg Pred Value : 0.7353
##
##
                Prevalence: 0.6708
            Detection Rate: 0.6149
##
```

```
## Detection Prevalence : 0.7888
## Balanced Accuracy : 0.6942
##
## 'Positive' Class : No
##
## Regressão Logística Regularizada sem missing
rocRL4 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloRL4, test_without_NAs, ty
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocRL4, titulo = "Curva ROC RL Regularizada (Removidos)")</pre>
```

Curva ROC RL Regularizada (Removidos)



```
predictRL4 <- predict(modeloRL4, newdata = test_without_NAs)
confusionMatrix(predictRL4, test_without_NAs$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction No Yes
## No 64 11
## Yes 4 10
```

```
##
##
                  Accuracy: 0.8315
##
                    95% CI: (0.7373, 0.9025)
##
      No Information Rate: 0.764
##
      P-Value [Acc > NIR] : 0.08127
##
##
                     Kappa: 0.4717
##
   Mcnemar's Test P-Value: 0.12134
##
##
##
              Sensitivity: 0.9412
##
               Specificity: 0.4762
            Pos Pred Value: 0.8533
##
##
            Neg Pred Value: 0.7143
##
               Prevalence: 0.7640
##
            Detection Rate: 0.7191
##
     Detection Prevalence: 0.8427
##
         Balanced Accuracy: 0.7087
##
##
          'Positive' Class : No
##
```

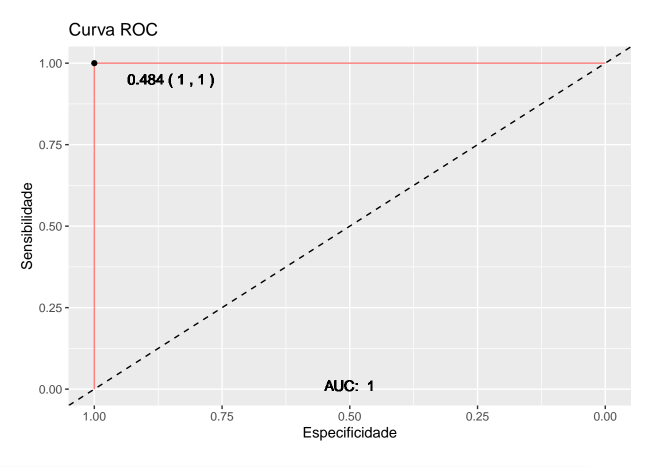
Random Forest

Modelagem

```
library(caret)
train.control <- caret::trainControl(method = "cv", number = 10) # Cross-validation com k=10
## Random Forest / dados imputados
set.seed(23)
tuneGrid <- expand.grid(.mtry = c(1: 10))</pre>
modeloRF1 <- train(diabetes~.,</pre>
    train,
    method = "rf",
    metric = "Accuracy",
   tuneGrid = tuneGrid,
   trControl = train.control,
    importance = TRUE,
   ntree = 1000)
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
```

```
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
print(modeloRF1)
```

```
## Random Forest
##
## 537 samples
    8 predictor
##
##
     2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 483, 483, 484, 483, 482, 484, ...
## Resampling results across tuning parameters:
##
##
     mtry Accuracy
                      Kappa
##
           0.7506099 0.4361204
     1
     2
##
           0.7562004 0.4559871
##
      3
           0.7468001 0.4344252
##
      4
           0.7486869 0.4382093
##
      5
           0.7487218 0.4409065
##
      6
           0.7505387 0.4457852
##
      7
           0.7523569 0.4506149
           0.7468001 0.4377380
##
     8
##
      9
           0.7431300 0.4295050
##
     10
           0.7487567 0.4392488
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 2.
rocRF1 <- roc(response = train$diabetes, predictor = predict(modeloRF1, train, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRF1)
```



```
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range

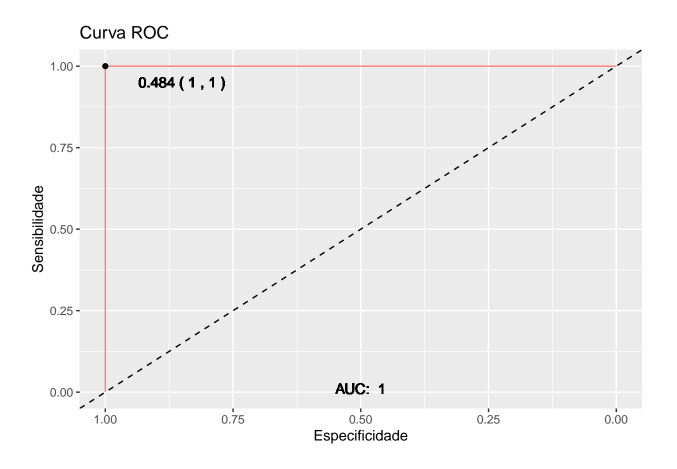
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range

## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range

## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
```

```
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
## Warning in randomForest.default(x, y, mtry = param$mtry, ...): invalid mtry:
## reset to within valid range
print(modeloRF2)
## Random Forest
## 266 samples
   8 predictor
```

```
2 classes: 'No', 'Yes'
##
## No pre-processing
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 240, 240, 239, 239, 239, 239, ...
## Resampling results across tuning parameters:
##
##
     mtry Accuracy
                      Kappa
##
     1
           0.7750712 0.4927747
##
      2
           0.7787749 0.5063822
##
      3
           0.7903134 0.5363471
##
          0.7938746 0.5468630
      4
##
      5
          0.7826211 0.5182975
##
      6
          0.7827635 0.5166230
##
     7
          0.7864672 0.5255357
##
     8
          0.7827635 0.5201542
##
     9
          0.7904558 0.5372170
           0.7903134 0.5347168
##
    10
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was mtry = 4.
rocRF2 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloRF1, train_without_NAs,</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRF2)
```



Árvores de decisão geradas

```
tree_num <- which(modeloRF1$finalModel$forest$ndbigtree == min(modeloRF1$finalModel$forest$ndbigtree))
print(paste("Menor arvore com imputação",tree_num))

## [1] "Menor arvore com imputação 876"

tree_func(final_model = modeloRF1$finalModel, tree_num[1])

## Warning: Duplicated aesthetics after name standardisation: na.rm

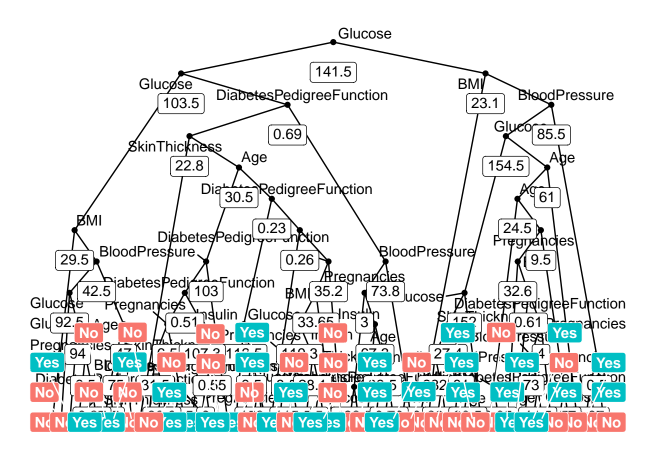
## Warning: Duplicated aesthetics after name standardisation: na.rm

## Warning: Duplicated aesthetics after name standardisation: na.rm

## Warning: Removed 64 rows containing missing values (geom_text_repel).

## Warning: Removed 64 rows containing missing values (geom_label).

## Warning: Removed 63 rows containing missing values (geom_label_repel).</pre>
```

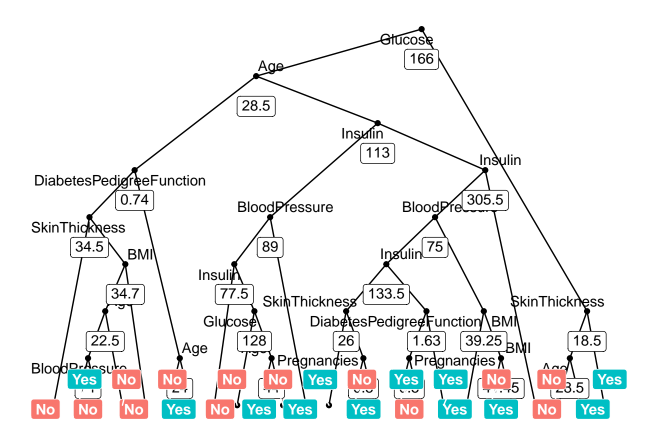


```
tree_num <- which(modeloRF2$finalModel$forest$ndbigtree == min(modeloRF2$finalModel$forest$ndbigtree))
print(paste("Menor árvore sem imputação:",tree_num))</pre>
```

[1] "Menor árvore sem imputação: 250"

```
tree_func(final_model = modeloRF2$finalModel, tree_num[1])
```

- ## Warning: Duplicated aesthetics after name standardisation: na.rm
- ## Warning: Duplicated aesthetics after name standardisation: na.rm
- ## Warning: Duplicated aesthetics after name standardisation: na.rm
- ## Warning: Removed 25 rows containing missing values (geom_text_repel).
- ## Warning: Removed 25 rows containing missing values (geom_label).
- ## Warning: Removed 24 rows containing missing values (geom_label_repel).



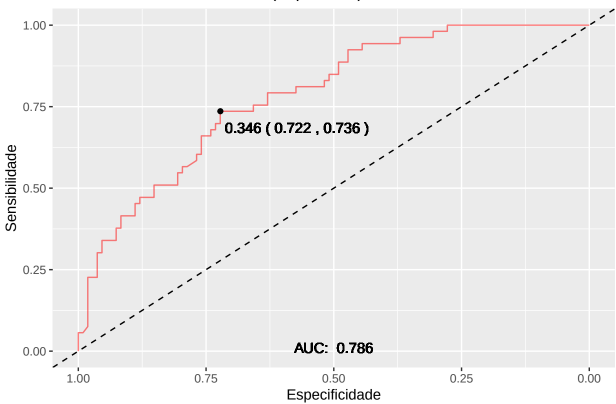
Teste

```
## Random Forest com dados imputados
rocRF1 <- roc(response = test$diabetes, predictor = predict(modeloRF1, test, type = "prob")[,2])
## Setting levels: control = No, case = Yes

## Setting direction: controls < cases

plotaroc(rocRF1, titulo = "Curva ROC Random Forest (Imputados)")</pre>
```

Curva ROC Random Forest (Imputados)



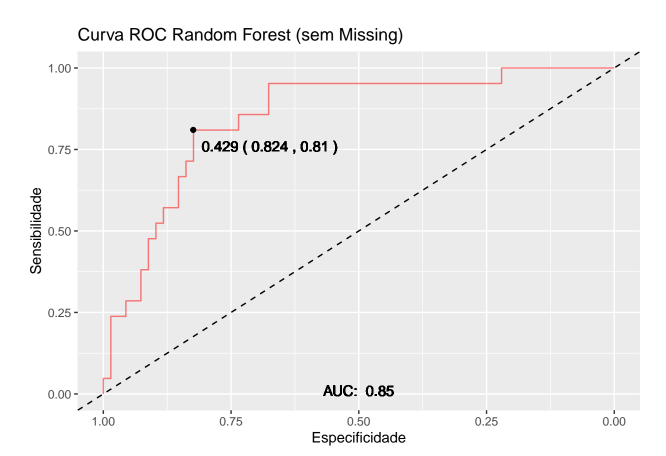
```
predictRF1 <- predict(modeloRF1, newdata = test)
print("Dados imputados:")</pre>
```

[1] "Dados imputados:"

confusionMatrix(predictRF1, test\$diabetes)

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
##
          No 93 28
          Yes 15 25
##
##
##
                  Accuracy : 0.7329
                    95% CI : (0.6576, 0.7995)
##
##
       No Information Rate: 0.6708
##
       P-Value [Acc > NIR] : 0.05368
##
##
                     Kappa : 0.355
##
    Mcnemar's Test P-Value : 0.06725
##
##
               Sensitivity: 0.8611
##
```

```
##
               Specificity: 0.4717
##
            Pos Pred Value: 0.7686
##
            Neg Pred Value: 0.6250
                Prevalence: 0.6708
##
##
            Detection Rate: 0.5776
##
      Detection Prevalence: 0.7516
##
         Balanced Accuracy: 0.6664
##
##
          'Positive' Class : No
##
## Random Forest com dados imputados
rocRF2 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloRF2, test_without_NAs, ty
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocRF2, titulo = "Curva ROC Random Forest (sem Missing)")
```



```
predictRF2 <- predict(modeloRF2, newdata = test_without_NAs)
print("Sem Missing:")</pre>
```

[1] "Sem Missing:"

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
          No 59
##
##
          Yes 9
##
                  Accuracy : 0.7978
##
                    95% CI : (0.6993, 0.8755)
##
       No Information Rate: 0.764
##
       P-Value [Acc > NIR] : 0.2709
##
##
##
                     Kappa: 0.4391
##
   Mcnemar's Test P-Value: 1.0000
##
##
##
               Sensitivity: 0.8676
##
               Specificity: 0.5714
##
            Pos Pred Value: 0.8676
##
            Neg Pred Value: 0.5714
##
                Prevalence: 0.7640
##
            Detection Rate: 0.6629
##
      Detection Prevalence: 0.7640
         Balanced Accuracy: 0.7195
##
##
##
          'Positive' Class : No
##
```

Support Vector Machine

Support Vector Machine (SVM) estabelece um limite de decisão ideal que separa os pontos de dados de diferentes classes e, em seguida, prevê a classe de novas observações com base nesse limite de separação. Os diferentes grupos podem ser separáveis por uma linha reta linear ou por uma linha limite não linear.

Modelagem

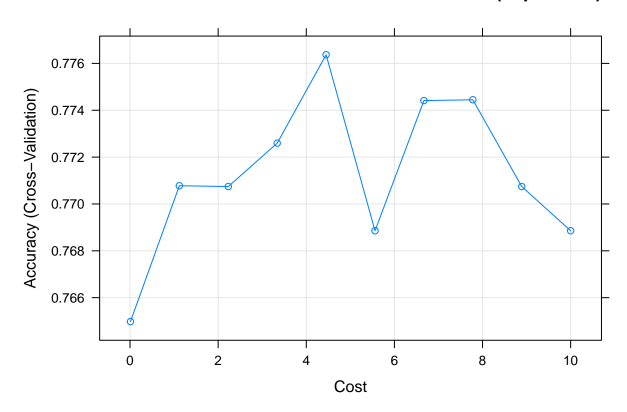
SVM Kernel Linear com dados imputados

```
train.control <- caret::trainControl(method = "cv", number = 10, savePred=T, classProb=T) # Cross-valid
#library(e1071) #SVM
## SVM Kernel Linear com dados imputados
set.seed(23)
modeloSVM1 <- caret::train(
    diabetes ~., data = train, method = "svmLinear",
    probability = T,</pre>
```

```
trControl = train.control,
  preProcess = c("center", "scale"),
  tuneGrid = expand.grid(C = seq(0.01, 10, length = 10))
print(modeloSVM1)
## Support Vector Machines with Linear Kernel
##
## 537 samples
##
    8 predictor
     2 classes: 'No', 'Yes'
##
##
## Pre-processing: centered (8), scaled (8)
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 483, 483, 484, 483, 482, 484, ...
## Resampling results across tuning parameters:
##
##
           Accuracy
                      Kappa
##
     0.01 0.7649794 0.4734524
##
      1.12 0.7707782 0.4824355
##
      2.23 0.7707433 0.4808278
##
      3.34 0.7725951 0.4843473
##
      4.45 0.7763687 0.4917503
      5.56 0.7688565 0.4769505
##
##
      6.67 0.7744133 0.4896157
     7.78 0.7744470 0.4862920
##
##
     8.89 0.7707433 0.4778763
##
     10.00 0.7688565 0.4754528
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was C = 4.45.
```

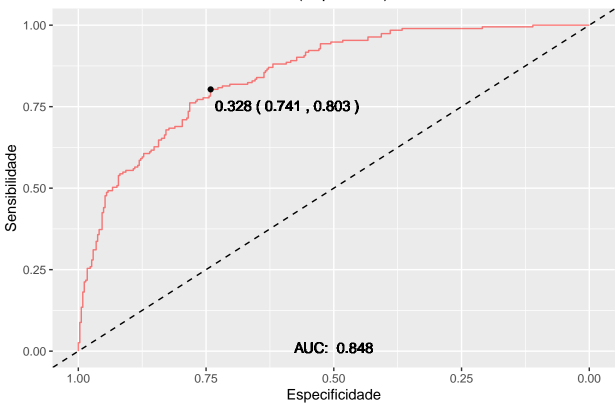
plot(modeloSVM1, main="SVM Kernel Linear - Acurácia vs Valores de Cost (Imputados)")

SVM Kernel Linear – Acurácia vs Valores de Cost (Imputados)



```
rocSVM1 <- roc(response = train$diabetes, predictor = predict(modeloSVM1, train, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocSVM1, titulo = "Curva ROC SVM Kernel Linear (Imputados)")</pre>
```

Curva ROC SVM Kernel Linear (Imputados)



SVM Kernel Linear sem missing

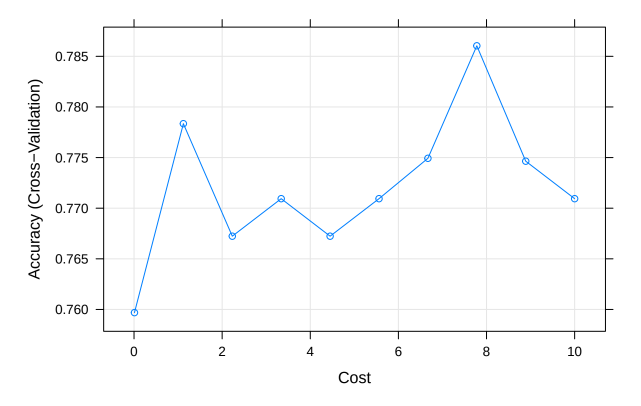
```
## SVM Kernel Linear sem missing
set.seed(23)
modeloSVM2 <- caret::train(
   diabetes ~., data = train_without_NAs, method = "svmLinear",
   trControl = train.control,
   preProcess = c("center", "scale"),
   tuneGrid = expand.grid(C = seq(0.01, 10, length = 10))
)
print(modeloSVM2)</pre>
```

```
## Support Vector Machines with Linear Kernel
##
## 266 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## Pre-processing: centered (8), scaled (8)
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 240, 240, 239, 239, 239, ...
## Resampling results across tuning parameters:
##
```

```
##
            Accuracy
                       Kappa
##
     0.01 0.7596866 0.4629441
      1.12
##
           0.7783476
                      0.4980691
      2.23
           0.7672365
                       0.4744505
##
##
      3.34
           0.7709402
                      0.4839230
      4.45
           0.7672365
                      0.4774236
##
##
      5.56
           0.7709402 0.4781789
           0.7749288
                       0.4893745
##
     6.67
##
     7.78
           0.7860399
                       0.5103509
##
     8.89
           0.7746439
                       0.4906956
##
     10.00 0.7709402
                      0.4754996
##
## Accuracy was used to select the optimal model using the largest value.
## The final value used for the model was C = 7.78.
```

plot(modeloSVM2, main="SVM Kernel Linear - Acurácia vs Valores de Cost (Removidos)")

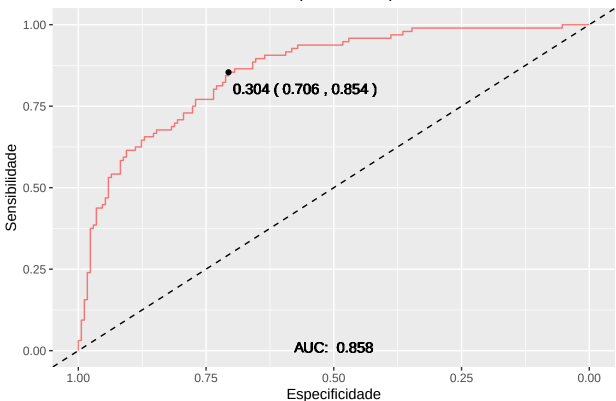
SVM Kernel Linear – Acurácia vs Valores de Cost (Removidos)



rocSVM2 <- roc(response = train_without_NAs\$diabetes, predictor = predict(modeloSVM2, train_without_NAs\$</pre>

Setting levels: control = No, case = Yes
Setting direction: controls < cases</pre>

Curva ROC SVM Kernel Linear (Removidos)



SVM Kernel Não Linear com dados imputados

```
## SVM Kernel Não Linear com dados imputados
set.seed(23)
modeloSVM3 <- caret::train(
   diabetes ~., data = train, method = "svmRadial",
   trControl = train.control,
   preProcess = c("center","scale"),
   tuneGrid = expand.grid(sigma = seq(0.01, 1, length = 10) , C = seq(1, 10, length = 10))
)
print(modeloSVM3)</pre>
```

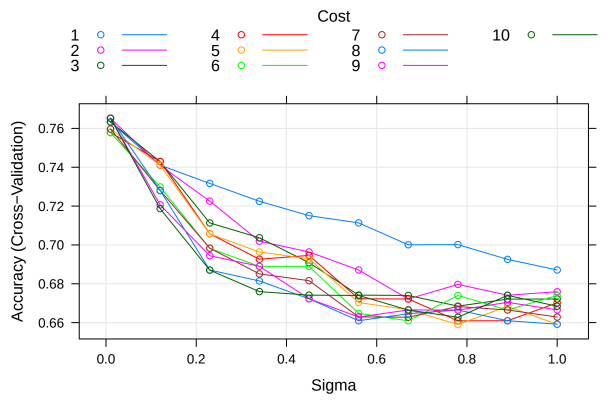
```
## Support Vector Machines with Radial Basis Function Kernel
##
## 537 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## Pre-processing: centered (8), scaled (8)
## Resampling: Cross-Validated (10 fold)
```

```
## Summary of sample sizes: 483, 483, 484, 483, 482, 484, ...
  Resampling results across tuning parameters:
##
##
     sigma C
                 Accuracy
                            Kappa
##
     0.01
             1
                0.7632323
                            0.4654255
##
     0.01
                            0.4686906
             2 0.7651541
##
     0.01
                            0.4652478
             3 0.7633371
##
     0.01
             4
                0.7578502
                            0.4499811
##
     0.01
             5
                0.7596334
                            0.4565153
##
     0.01
                0.7578153
                            0.4521902
##
     0.01
             7
                0.7597021
                            0.4544051
##
     0.01
                0.7652576
                            0.4649773
             8
##
     0.01
             9
                0.7652576
                            0.4667128
##
     0.01
                            0.4665617
                0.7652227
##
     0.12
                0.7410412
                            0.4172729
             1
##
     0.12
             2
                0.7409688
                            0.4130742
##
                0.7428905
     0.12
                            0.4189473
             3
##
     0.12
                0.7429928
                            0.4173560
##
     0.12
                0.7410724
                            0.4173790
             5
##
     0.12
             6
                0.7298901
                            0.3891193
##
     0.12
             7
                0.7280382
                            0.3836402
##
     0.12
                0.7280382
                            0.3865933
##
     0.12
             9 0.7206296
                            0.3666429
##
     0.12
                0.7186754
                            0.3627544
            10
##
     0.23
             1
                0.7316746
                            0.3933253
##
     0.23
             2
                0.7225513
                            0.3738773
##
     0.23
                0.7113354
                            0.3457537
     0.23
                0.7057449
##
             4
                            0.3323967
##
     0.23
                0.7056750
                            0.3313558
##
     0.23
                0.6982314
                            0.3163946
             6
##
     0.23
             7
                0.6983362
                            0.3096625
##
     0.23
             8
                0.6870853
                            0.2825074
     0.23
##
                0.6944578
                            0.3009454
##
     0.23
                0.6870167
                            0.2782018
            10
##
     0.34
                0.7224827
                            0.3714102
             1
##
     0.34
             2 0.7019376
                            0.3273452
##
     0.34
               0.7036872
                            0.3239199
##
     0.34
             4 0.6926421
                            0.2970354
##
     0.34
                0.6963808
                            0.3111219
             5
##
     0.34
                0.6889721
                            0.2906708
             6
                0.6850975
##
     0.34
                            0.2810966
##
     0.34
                0.6814624
                            0.2675126
             8
     0.34
##
             9
                0.6890071
                            0.2828918
##
     0.34
            10
                0.6759742
                            0.2551517
##
     0.45
                            0.3509448
                0.7150753
             1
##
     0.45
             2
                0.6963833
                            0.3062685
##
     0.45
             3
                0.6908589
                            0.2926231
##
     0.45
                0.6945988
                            0.3106269
##
     0.45
                0.6928156
                            0.3041198
             5
##
     0.45
             6
                0.6889734
                            0.2984869
##
     0.45
             7
                0.6816009
                            0.2875827
##
     0.45
                0.6722718
                            0.2590777
##
     0.45
             9
                0.6723067
                            0.2552091
##
     0.45
            10 0.6741586 0.2616750
```

```
##
     0.56
             1 0.7113379 0.3428026
             2 0.6870866
##
     0.56
                           0.2825257
     0.56
##
                0.6741573
                           0.2575748
##
     0.56
                0.6723054
                           0.2584329
##
     0.56
             5
                0.6703500
                           0.2495136
##
     0.56
             6 0.6647958
                           0.2365721
##
     0.56
                0.6629439
                           0.2385834
     0.56
                           0.2203032
##
             8 0.6609872
                0.6628740
##
     0.56
             9
                           0.2280941
##
     0.56
            10
                0.6739864
                           0.2501550
##
     0.67
                0.7000870
                           0.3084599
             1
##
     0.67
                0.6721670
                           0.2416235
##
     0.67
                0.6739864
                           0.2500257
##
     0.67
                0.6721695
                           0.2490706
##
     0.67
                0.6666463
                           0.2338014
             5
##
     0.67
                0.6609186
                           0.2177422
##
     0.67
                           0.2275974
             7
                0.6628054
##
     0.67
                0.6645887
                           0.2323034
##
     0.67
             9 0.6665428
                           0.2321039
##
     0.67
            10 0.6664754
                           0.2375644
##
     0.78
             1 0.7001569
                           0.3088335
##
     0.78
                0.6796805
                           0.2585207
##
     0.78
             3 0.6684982
                           0.2323193
##
     0.78
             4 0.6609186
                           0.2142967
##
             5 0.6590318
                           0.2111359
     0.78
##
     0.78
             6 0.6739877
                           0.2470423
##
     0.78
                0.6682924
                           0.2377308
             7
##
     0.78
               0.6664754
                           0.2266465
             8
##
     0.78
                0.6664754
                           0.2258914
                0.6627705
                           0.2125283
     0.78
##
            10
##
     0.89
             1
                0.6925761
                           0.2908961
##
     0.89
             2 0.6741236
                           0.2431205
##
     0.89
                0.6721009
                           0.2334575
##
     0.89
             4 0.6609536
                           0.2093185
##
     0.89
                0.6683610
                           0.2275698
             5
##
     0.89
             6 0.6665091
                           0.2235560
##
     0.89
                0.6665091
                           0.2188595
##
     0.89
             8 0.6609872
                           0.2072523
##
     0.89
             9
                0.6704212
                           0.2236802
##
     0.89
            10 0.6740550
                           0.2476682
##
     1.00
                0.6870891
                           0.2753881
                0.6759081
##
     1.00
                           0.2379519
     1.00
             3 0.6721695
                           0.2306436
##
##
     1.00
             4 0.6702128
                           0.2300421
##
     1.00
             5 0.6591017
                           0.2023119
##
     1.00
             6 0.6739864
                           0.2383797
     1.00
##
             7
                0.6628728
                           0.2097380
##
     1.00
                0.6592040
                           0.2043506
##
     1.00
                0.6665790
                           0.2159346
##
     1.00
            10
               0.6684308
                           0.2258624
##
```

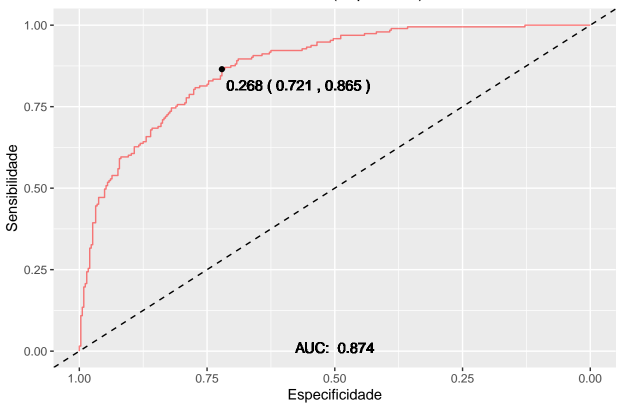
Accuracy was used to select the optimal model using the largest value. ## The final values used for the model were sigma = 0.01 and C = 9.

SVM Kernel não Linear - Acurácia vs Valores de Sigma (Imputados)



```
rocSVM3 <- roc(response = train$diabetes, predictor = predict(modeloSVM3, train, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocSVM3, titulo = "Curva ROC SVM Kernel não Linear (Imputados)")</pre>
```

Curva ROC SVM Kernel não Linear (Imputados)



SVM Kernel Não Linear sem missing

```
## SVM Kernel Não Linear sem missing
set.seed(23)
modeloSVM4 <- caret::train(
    diabetes ~., data = train_without_NAs, method = "svmRadial",
    trControl = train.control,
    preProcess = c("center", "scale"),
    tuneGrid = expand.grid(sigma = seq(0.01, 1, length = 10) , C = seq(1, 10, length = 10))
)
print(modeloSVM4)</pre>
```

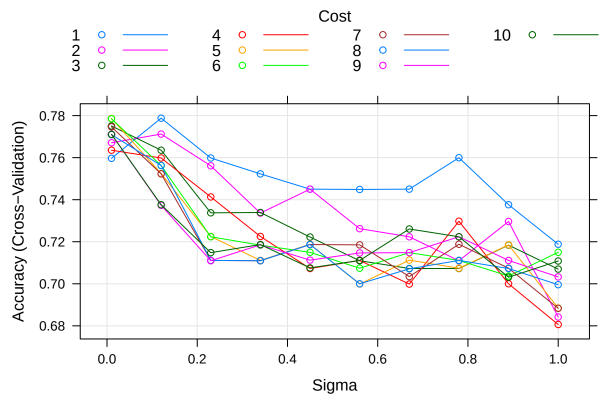
```
## Support Vector Machines with Radial Basis Function Kernel
##
## 266 samples
## 8 predictor
## 2 classes: 'No', 'Yes'
##
## Pre-processing: centered (8), scaled (8)
## Resampling: Cross-Validated (10 fold)
## Summary of sample sizes: 240, 240, 239, 239, 239, 239, ...
## Resampling results across tuning parameters:
##
```

```
Accuracy
##
     sigma C
                             Kappa
##
     0.01
                            0.4650398
             1
                 0.7596866
##
     0.01
                 0.7670940
                            0.4812458
     0.01
##
                 0.7750712
                            0.4961683
##
     0.01
                 0.7635328
                            0.4698621
##
     0.01
                            0.4972411
                 0.7786325
##
     0.01
                 0.7784900
                            0.4946139
##
     0.01
             7
                 0.7746439
                            0.4843767
##
     0.01
             8
                 0.7709402
                            0.4757028
##
     0.01
                 0.7710826
                            0.4782618
##
     0.01
             10
                 0.7710826
                            0.4782618
##
     0.12
                 0.7787749
                            0.4951781
##
     0.12
             2
                 0.7712251
                            0.4802989
##
     0.12
                 0.7635328
                            0.4566007
##
     0.12
                 0.7598291
                             0.4445875
##
     0.12
             5
                 0.7524217
                             0.4302310
##
                            0.4371691
     0.12
             6
                 0.7562678
##
     0.12
                 0.7522792
                            0.4200541
##
     0.12
                 0.7562678
                            0.4370246
             8
##
     0.12
                 0.7373219
                            0.3911532
##
     0.12
             10
                 0.7376068
                            0.3892707
##
     0.23
                 0.7598291
                            0.4523565
##
     0.23
                            0.4451085
             2
                 0.7561254
                 0.7337607
##
     0.23
                            0.4021808
##
     0.23
                 0.7413105
                            0.4199586
##
     0.23
                 0.7225071
                            0.3736789
##
     0.23
                 0.7223647
                            0.3670588
##
     0.23
             7
                 0.7111111
                            0.3417905
##
     0.23
                             0.3408347
                 0.7111111
     0.23
##
                 0.7109687
                             0.3405406
##
     0.23
             10
                 0.7148148
                            0.3367158
##
     0.34
             1
                 0.7522792
                            0.4378279
     0.34
##
                 0.7336182
                            0.4009842
##
     0.34
                 0.7339031
                            0.3950800
##
     0.34
                 0.7225071
                            0.3770402
##
     0.34
                            0.3487114
             5
                 0.7111111
##
     0.34
                 0.7183761
                            0.3638857
##
     0.34
             7
                 0.7109687
                             0.3366912
##
     0.34
             8
                 0.7109687
                            0.3441970
##
     0.34
                            0.3508626
             9
                 0.7185185
##
     0.34
                 0.7186610
                            0.3590489
             10
##
     0.45
                 0.7450142
                            0.4217840
     0.45
                            0.4322630
##
             2
                 0.7450142
##
     0.45
                 0.7222222
                            0.3712480
##
     0.45
                 0.7072650
                            0.3469918
##
     0.45
                 0.7188034
                             0.3656221
             5
##
     0.45
             6
                 0.7149573
                            0.3580174
##
     0.45
                 0.7186610
                            0.3583701
##
     0.45
             8
                 0.7186610
                             0.3619806
##
     0.45
             9
                 0.7112536
                            0.3476623
##
     0.45
                            0.3454457
             10
                 0.7075499
     0.56
##
                 0.7448718
                            0.4243368
##
     0.56
             2
                 0.7262108
                            0.3809057
     0.56
##
             3 0.7109687 0.3386173
```

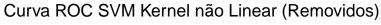
```
##
    0.56
            4 0.7111111 0.3454171
##
    0.56
            5 0.6998575 0.3195248
    0.56
##
            6 0.7074074 0.3369069
    0.56
##
            7 0.7185185
                         0.3779832
##
    0.56
            8 0.7000000 0.3153407
##
    0.56
            9 0.7146724 0.3515126
           10 0.7111111 0.3485484
##
    0.56
    0.67
##
            1
              0.7450142 0.4361749
##
    0.67
            2 0.7223647
                         0.3844141
##
    0.67
            3 0.7072650
                         0.3418878
##
    0.67
            4 0.6998575
                          0.3169579
##
    0.67
            5 0.7112536
                         0.3447735
##
    0.67
            6 0.7148148 0.3577632
##
    0.67
            7 0.7034188 0.3278498
##
    0.67
            8 0.7071225
                          0.3375983
##
    0.67
            9 0.7148148
                         0.3570661
##
    0.67
           10 0.7260684 0.3767459
##
    0.78
            1 0.7599715
                         0.4719043
##
    0.78
            2 0.7109687
                          0.3479731
##
    0.78
            3 0.7072650 0.3401029
##
    0.78
            4 0.7297721 0.3925998
##
    0.78
            5 0.7074074 0.3478802
##
    0.78
            6 0.7111111
                         0.3430724
##
    0.78
            7 0.7188034 0.3574965
##
    0.78
            8 0.7111111 0.3475439
##
    0.78
            9 0.7223647
                          0.3713868
##
    0.78
           10 0.7223647
                          0.3729232
##
    0.89
            1 0.7376068 0.4239749
##
    0.89
            2 0.7296296
                         0.3890878
    0.89
##
            3 0.7183761
                          0.3677189
##
    0.89
            4 0.7000000 0.3174928
##
    0.89
            5 0.7185185 0.3622729
    0.89
##
            6 0.7037037
                          0.3369286
##
    0.89
            7 0.7074074 0.3383008
##
    0.89
            8 0.7072650 0.3383578
##
    0.89
           9 0.7111111 0.3472359
##
    0.89
           10 0.7031339
                         0.3065973
##
    1.00
            1 0.7188034 0.3915682
##
    1.00
            2 0.6841880 0.2662962
##
    1.00
            3 0.7069801 0.3218391
##
    1.00
            4 0.6806268 0.2520008
##
    1.00
            5 0.6884615 0.2735769
    1.00
            6 0.7149573 0.3506383
##
##
    1.00
            7 0.6883191 0.2773509
##
    1.00
            8 0.6995726
                         0.2933212
    1.00
##
            9 0.7032764 0.3099552
    1.00
           10 0.7108262 0.3449889
##
##
```

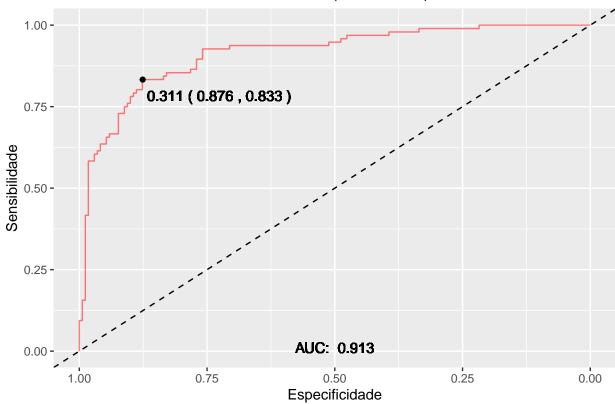
Accuracy was used to select the optimal model using the largest value. ## The final values used for the model were sigma = 0.12 and C = 1.

SVM Kernel não Linear - Acurácia vs Valores de Sigma (Removidos)



```
rocSVM4 <- roc(response = train_without_NAs$diabetes, predictor = predict(modeloSVM4, train_without_NAs
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocSVM4, titulo ="Curva ROC SVM Kernel não Linear (Removidos)")</pre>
```

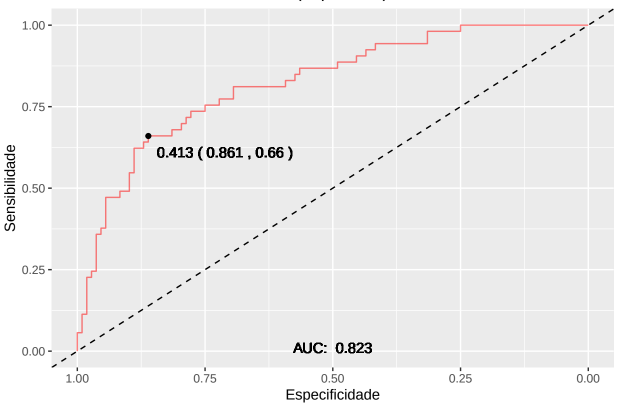




Testes

```
## SVM Kernel Linear com dados imputados
rocSVM1 <- roc(response = test$diabetes, predictor = predict(modeloSVM1, test, type = "prob")[,2])
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocSVM1, titulo = "Curva ROC SVM Kernel Linear (Imputados)")</pre>
```

Curva ROC SVM Kernel Linear (Imputados)



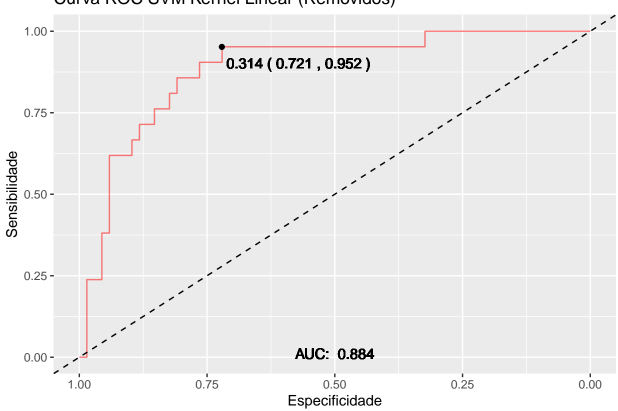
```
predictSVM1 <- predict(modeloSVM1, newdata = test)
confusionMatrix(predictSVM1, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
             Reference
##
## Prediction No Yes
          No 98 27
##
##
          Yes 10 26
##
                  Accuracy : 0.7702
##
                    95% CI: (0.6974, 0.8327)
##
##
       No Information Rate: 0.6708
       P-Value [Acc > NIR] : 0.003815
##
##
##
                     Kappa : 0.4334
##
##
    Mcnemar's Test P-Value: 0.008529
##
               Sensitivity: 0.9074
##
               Specificity: 0.4906
##
##
            Pos Pred Value: 0.7840
            Neg Pred Value : 0.7222
##
##
                Prevalence: 0.6708
            Detection Rate: 0.6087
##
```

```
## Detection Prevalence : 0.7764
## Balanced Accuracy : 0.6990
##
## 'Positive' Class : No
##
## SVM Kernel Linear sem missing
rocSVM2 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloSVM2, test_without_NAs,
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocSVM2, titulo = "Curva ROC SVM Kernel Linear (Removidos)")</pre>
```

Curva ROC SVM Kernel Linear (Removidos)

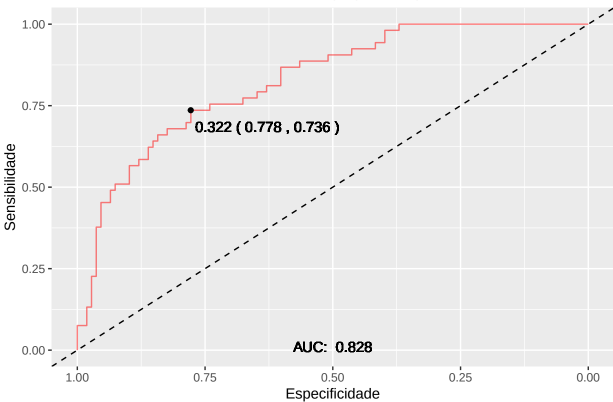


```
predictSVM2 <- predict(modeloSVM2, newdata = test_without_NAs)
confusionMatrix(predictSVM2, test_without_NAs$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction No Yes
## No 64 12
## Yes 4 9
```

```
##
##
                  Accuracy : 0.8202
                    95% CI: (0.7245, 0.8936)
##
##
       No Information Rate: 0.764
       P-Value [Acc > NIR] : 0.12910
##
##
##
                     Kappa: 0.4258
##
##
    Mcnemar's Test P-Value: 0.08012
##
##
               Sensitivity: 0.9412
##
               Specificity: 0.4286
##
            Pos Pred Value : 0.8421
            Neg Pred Value: 0.6923
##
##
                Prevalence: 0.7640
            Detection Rate: 0.7191
##
##
      Detection Prevalence: 0.8539
         Balanced Accuracy: 0.6849
##
##
##
          'Positive' Class : No
##
## SVM Kernel não Linear com dados imputados
rocSVM3 <- roc(response = test$diabetes, predictor = predict(modeloSVM3, test, type = "prob")[,2])</pre>
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases
plotaroc(rocSVM3, titulo = "Curva ROC SVM Kernel não Linear (Imputados)")
```





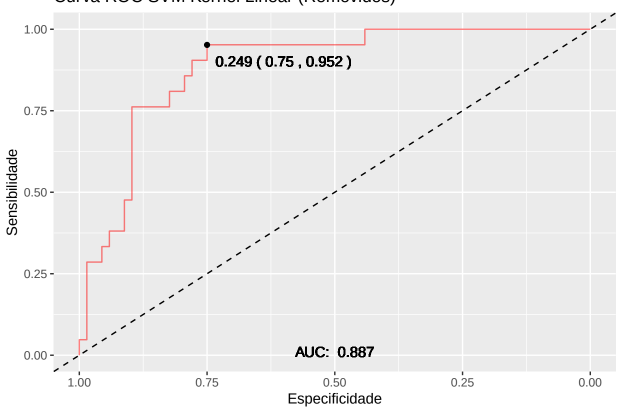
```
predictSVM3 <- predict(modeloSVM3, newdata = test)
confusionMatrix(predictSVM3, test$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
##
             Reference
## Prediction No Yes
          No 99
                  26
##
##
          Yes 9
                  27
##
                  Accuracy : 0.7826
##
                    95% CI: (0.7109, 0.8437)
##
##
       No Information Rate: 0.6708
       P-Value [Acc > NIR] : 0.001230
##
##
##
                     Kappa : 0.464
##
##
    Mcnemar's Test P-Value: 0.006841
##
               Sensitivity: 0.9167
##
##
               Specificity: 0.5094
##
            Pos Pred Value: 0.7920
            Neg Pred Value : 0.7500
##
##
                Prevalence: 0.6708
            Detection Rate: 0.6149
##
```

```
## Detection Prevalence : 0.7764
## Balanced Accuracy : 0.7131
##
## 'Positive' Class : No
##
## SVM Kernel Linear sem missing
rocSVM4 <- roc(response = test_without_NAs$diabetes, predictor = predict(modeloSVM4, test_without_NAs,
## Setting levels: control = No, case = Yes
## Setting direction: controls < cases

plotaroc(rocSVM4, titulo = "Curva ROC SVM Kernel Linear (Removidos)")</pre>
```

Curva ROC SVM Kernel Linear (Removidos)



```
predictSVM4 <- predict(modeloSVM4, newdata = test_without_NAs)
confusionMatrix(predictSVM4, test_without_NAs$diabetes)</pre>
```

```
## Confusion Matrix and Statistics
##
## Reference
## Prediction No Yes
## No 63 13
## Yes 5 8
```

Accuracy : 0.7978 95% CI: (0.6993, 0.8755) ## ## No Information Rate : 0.764 P-Value [Acc > NIR] : 0.27088 ## ## Kappa : 0.354 ## ## Mcnemar's Test P-Value : 0.09896 ## ## Sensitivity: 0.9265 ## Specificity: 0.3810 ## Pos Pred Value: 0.8289 ## Neg Pred Value: 0.6154 ## ## Prevalence: 0.7640 Detection Rate: 0.7079 ## ## Detection Prevalence: 0.8539 ## Balanced Accuracy : 0.6537 ## 'Positive' Class : No

##