Model Selection - readmission_1year

Eduardo Yuki Yada

Global parameters

```
k <- params$k # Number of folds for cross validation
grid_size <- params$grid_size # Number of parameter combination to tune on each model
repeats <- params$repeats
```

Minutes to run: 0

Imports

```
library(tidyverse)
library(yaml)
library(tidymodels)
library(usemodels)
library(vip)
library(bonsai)
library(lightgbm)
library(aret)
library(pROC)
```

Minutes to run: 0

Loading data

```
load('dataset/processed_data.RData')
load('dataset/processed_dictionary.RData')

columns_list <- yaml.load_file("./auxiliar/columns_list.yaml")

outcome_column <- params$outcome_column
features_list <- params$features_list

df <- mutate(df, across(where(is.character), as.factor))

Minutes to run: 0.005</pre>
```

Minutes to run: 0

Eligible features

```
cat_features_list = readRDS(sprintf(
  "./auxiliar/significant_columns/categorical_%s.rds",
  outcome_column
))
num_features_list = readRDS(sprintf(
  "./auxiliar/significant_columns/numerical_%s.rds",
 outcome_column
))
features_list = c(cat_features_list, num_features_list)
Minutes to run: 0
eligible_columns = df_names %>%
  filter(momento.aquisicao == 'Admissão t0') %>%
  .$variable.name
exception_columns = c('death_intraop', 'death_intraop_1', 'disch_outcomes_t0')
correlated_columns = c('year_procedure_1', # com year_adm_t0
                       'age_surgery_1', # com age
                       'admission_t0', # com admission_pre_t0_count
                       'atb', # com meds_antimicrobianos
                       'classe_meds_cardio_qtde', # com classe_meds_qtde
                       'suporte_hemod', # com proced_invasivos_qtde,
                       'radiografia', # com exames_imagem_qtde
                       'ecg' # com metodos_graficos_gtde
                       )
eligible_features = eligible_columns %>%
  base::intersect(c(columns_list$categorical_columns, columns_list$numerical_columns)) %>%
  setdiff(c(exception_columns, correlated_columns))
if (is.null(features_list)) {
  features = eligible_features
} else {
  features = base::intersect(eligible_features, features_list)
}
gluedown::md_order(features, seq = TRUE, pad = TRUE)
## 01. sex
## 02. age
## 03. race
## 04. education_level
## 05. patient_state
## 06. underlying_heart_disease
## 07. heart_disease
## 08. nyha_basal
## 09. prior_mi
## 10. heart_failure
## 11. af
## 12. cardiac_arrest
## 13. transplant
## 14. valvopathy
## 15. endocardites
## 16. diabetes
## 17. renal_failure
## 18. hemodialysis
```

- ## 19. copd
- ## 20. comorbidities_count
- ## 21. procedure_type_1
- ## 22. reop_type_1
- ## 23. procedure_type_new
- ## 24. cied_final_1
- ## 25. cied_final_group_1
- ## 26. admission_pre_t0_count
- ## 27. admission_pre_t0_180d
- ## 28. year_adm_t0
- ## 29. icu_t0
- ## 30. dialysis_t0
- ## 31. admission_t0_emergency
- ## 32. aco
- ## 33. antiarritmico
- ## 34. betabloqueador
- ## 35. ieca_bra
- ## 36. dva
- ## 37. digoxina
- ## 38. estatina
- ## 39. diuretico
- ## 40. vasodilatador
- ## 41. insuf_cardiaca
- ## 42. espironolactona
- ## 43. bloq_calcio
- ## 44. antiplaquetario_ev
- ## 45. insulina
- ## 46. anticonvulsivante
- ## 47. psicofarmacos
- ## 48. antifungico
- ## 49. antiviral
- ## 50. antiretroviral
- ## 51. classe_meds_qtde
- ## 52. meds_cardiovasc_qtde
- ## 53. meds_antimicrobianos
- ## 54. ventilacao_mecanica
- ## 55. cec
- ## 56. transplante_cardiaco
- ## 57. cir_toracica
- ## 58. outros_proced_cirurgicos
- ## 59. icp
- ## 60. intervencao_cv
- ## 61. angioplastia
- ## 62. cateterismo
- ## 63. eletrofisiologia
- ## 64. cateter_venoso_central
- ## 65. proced_invasivos_qtde
- ## 66. cve_desf
- ## 67. transfusao
- ## 68. interconsulta
- ## 69. equipe_multiprof
- ## 70. holter
- ## 71. teste_esforco
- ## 72. espiro_ergoespiro
- ## 73. tilt_teste
- ## 74. metodos_graficos_qtde
- ## 75. laboratorio
- ## 76. cultura
- ## 77. analises_clinicas_qtde
- ## 78. citologia
- ## 79. biopsia

```
## 80. histopatologia_qtde
## 81. angio_rm
## 82. angio_tc
## 83. aortografia
## 84. arteriografia
## 85. cintilografia
## 86. ecocardiograma
## 87. endoscopia
## 88. flebografia
## 89. pet_ct
## 90. ultrassom
## 91. tomografia
## 92. ressonancia
## 93. exames_imagem_qtde
## 94. dieta_parenteral
## 95. bic
## 96. mpp
## 97. hospital_stay
Minutes to run: 0
```

Train test split (70%/30%)

Minutes to run: 0.001

Boosted Tree (XGBoost)

```
xgboost_recipe <-</pre>
  recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
  step_novel(all_nominal_predictors()) %>%
  step_unknown(all_nominal_predictors()) %>%
  step_other(all_nominal_predictors(), threshold = 0.05, other = ".merged") %>%
  step_dummy(all_nominal_predictors())
xgboost_spec <- boost_tree(</pre>
 trees = tune(),
 min_n = tune(),
 tree_depth = tune(),
 learn_rate = tune(),
 loss_reduction = tune(),
 sample_size = tune()
) %>%
  set_engine("xgboost",
            nthread = 8) %>%
  set_mode("classification")
```

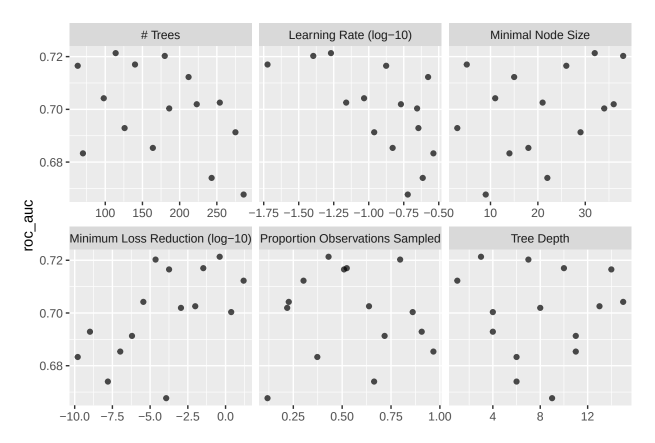
```
xgboost_grid <- grid_latin_hypercube(</pre>
  trees(range = c(50L, 300L)),
 min n(),
 tree_depth(),
 learn_rate(range = c(0.01, 0.3), trans = NULL),
 loss_reduction(),
  sample_prop(range = c(1/10, 1), trans = NULL),
  size = grid_size
xgboost_workflow <-</pre>
  workflow() %>%
  add_recipe(xgboost_recipe) %>%
  add_model(xgboost_spec)
xgboost_tune <-
  xgboost_workflow %>%
  tune_grid(resamples = df_folds,
            grid = xgboost_grid)
xgboost_tune %>%
 show_best("roc_auc")
## # A tibble: 5 x 12
## trees min_n tree_depth learn_rate loss_reduction sample~1 .metric .esti~2 mean
                                                                                            n \ std\_err .config
##
    <int> <int>
                  <int>
                                 <dbl>
                                             <dbl> <dbl> <chr>
                                                                        <chr>
                                                                                  <dbl> <int> <dbl> <chr>
     114
           32
                        3
                                 0.0537
                                             0.410
                                                           0.430 roc_auc binary 0.721 5 0.00599 Prepro~
## 1
## 2 180 38
                         7
                               0.0402

      0.524 roc_auc binary
      0.717
      5 0.00496 Prepro~

      0.509 roc_auc binary
      0.717
      5 0.00707 Prepro~

      0.302 roc_auc binary
      0.712
      5 0.00714 Prepro~

## 3 140 5
                        10
                                 0.0189
                                             0.0336
## 4
       63
              26
                         14
                                 0.133
                                             0.000181
## 5
      212
            15
                          1
                                 0.265
                                            15.8
## # ... with abbreviated variable names 1: sample_size, 2: .estimator
best_xgboost <- xgboost_tune %>%
 select_best("roc_auc")
autoplot(xgboost_tune, metric = "roc_auc")
```

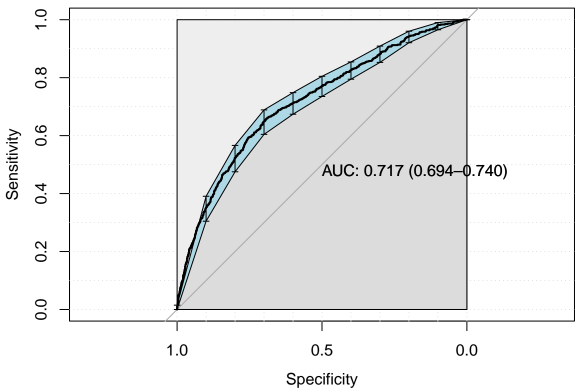


```
final_xgboost_workflow <-
    xgboost_workflow %>%
    finalize_workflow(best_xgboost)

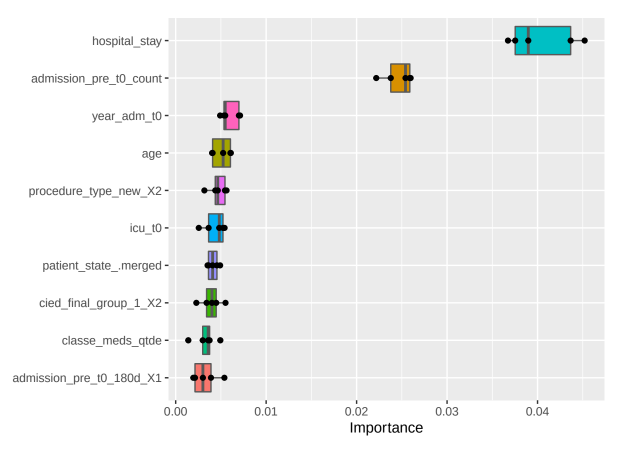
last_xgboost_fit <-
    final_xgboost_workflow %>%
    last_fit(df_split)

final_xgboost_fit <- extract_workflow(last_xgboost_fit)

xgboost_auc <- validation(final_xgboost_fit, df_test)</pre>
```



```
## [1] "Optimal Threshold: 0.12"
##
  Confusion Matrix and Statistics
##
       reference
##
  data
           0
##
      0 2863
              207
##
      1 1264 397
##
##
                  Accuracy : 0.6891
                    95% CI: (0.6757, 0.7022)
##
##
      No Information Rate: 0.8723
      P-Value [Acc > NIR] : 1
##
##
##
                     Kappa : 0.2009
##
##
   Mcnemar's Test P-Value : <2e-16
##
##
               Sensitivity: 0.6937
##
               Specificity: 0.6573
##
            Pos Pred Value : 0.9326
            Neg Pred Value: 0.2390
##
##
                Prevalence: 0.8723
##
            Detection Rate: 0.6052
##
      Detection Prevalence: 0.6489
##
         Balanced Accuracy : 0.6755
##
##
          'Positive' Class: 0
##
extract_vip(final_xgboost_fit, pred_wrapper = predict,
            reference_class = "0")
```

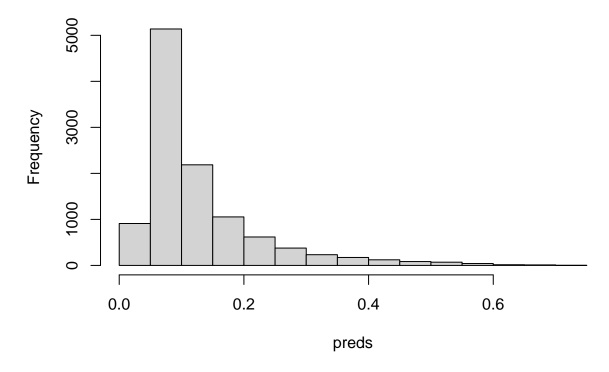


```
xgboost_parameters <- xgboost_tune %>%
    show_best("roc_auc", n = 1) %>%
    select(trees, min_n, tree_depth, learn_rate, loss_reduction) %>%
    as.list

saveRDS(
    xgboost_parameters,
    file = sprintf(
        "./auxiliar/model_selection/hyperparameters/xgboost_%s.rds",
        outcome_column
    )
)
```

Minutes to run: 3.423

Histogram of preds



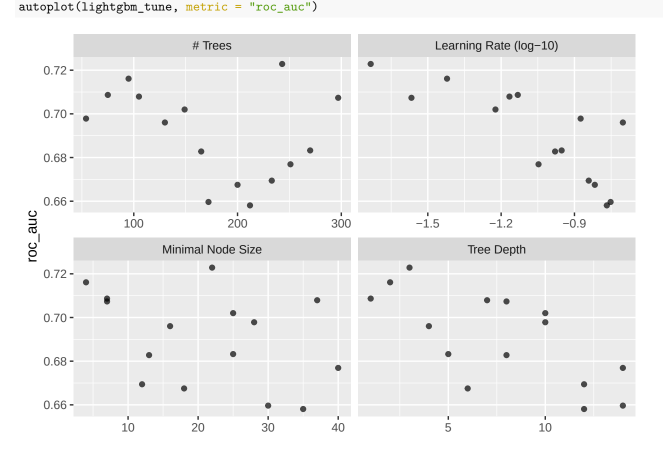
Minutes to run:

0.006

Boosted Tree (LightGBM)

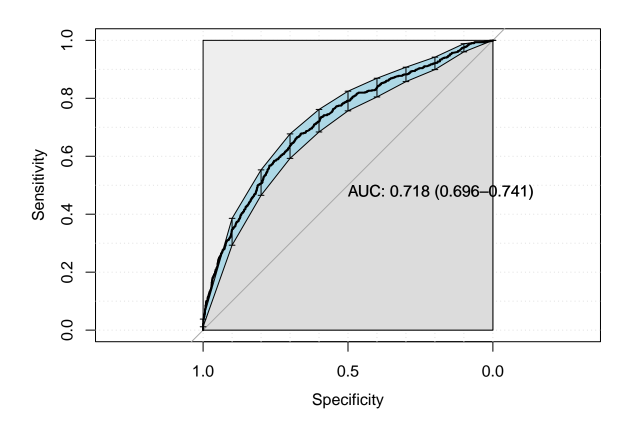
```
lightgbm_recipe <-</pre>
  recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
  step_novel(all_nominal_predictors()) %>%
  step_unknown(all_nominal_predictors()) %>%
  step_other(all_nominal_predictors(), threshold = 0.05, other = ".merged") %>%
  step_dummy(all_nominal_predictors())
lightgbm_spec <- boost_tree(</pre>
  trees = tune(),
  min_n = tune(),
  tree_depth = tune(),
  learn_rate = tune(),
  # loss_reduction = tune(),
  sample_size = 1
) %>%
  set_engine("lightgbm",
             nthread = 8) %>%
  set_mode("classification")
lightgbm_grid <- grid_latin_hypercube(</pre>
  trees(range = c(50L, 300L)),
  min_n(),
  tree_depth(),
  learn_rate(range = c(0.01, 0.2), trans = NULL),
  # loss_reduction(),
  size = grid_size
)
lightgbm_workflow <-
```

```
workflow() %>%
  add_recipe(lightgbm_recipe) %>%
  add_model(lightgbm_spec)
lightgbm_tune <-
  lightgbm_workflow %>%
  tune_grid(resamples = df_folds,
            grid = lightgbm_grid)
lightgbm_tune %>%
  show_best("roc_auc")
## # A tibble: 5 x 10
                                                                      n std_err .config
##
     trees min_n tree_depth learn_rate .metric .estimator
                                                            mean
##
     <int> <int>
                       <int>
                                  <dbl> <chr>
                                                 <chr>>
                                                            <dbl> <int>
                                                                           <dbl> <chr>
##
       243
              22
                           3
                                 0.0185 roc_auc binary
                                                            0.723
                                                                       5 0.00661 Preprocessor1_Model08
  1
##
  2
        95
               4
                           2
                                 0.0379 roc_auc binary
                                                            0.716
                                                                       5 0.00570 Preprocessor1_Model01
##
  3
        75
               7
                           1
                                                            0.709
                                                                       5 0.00431 Preprocessor1_Model02
                                 0.0738 roc_auc binary
              37
                           7
##
  4
       105
                                 0.0682 roc_auc binary
                                                            0.708
                                                                       5 0.00544 Preprocessor1_Model14
##
  5
       297
               7
                           8
                                 0.0271 roc_auc binary
                                                            0.707
                                                                       5 0.00721 Preprocessor1_Model03
best_lightgbm <- lightgbm_tune %>%
  select_best("roc_auc")
```



```
final_lightgbm_workflow <-
    lightgbm_workflow %>%
    finalize_workflow(best_lightgbm)

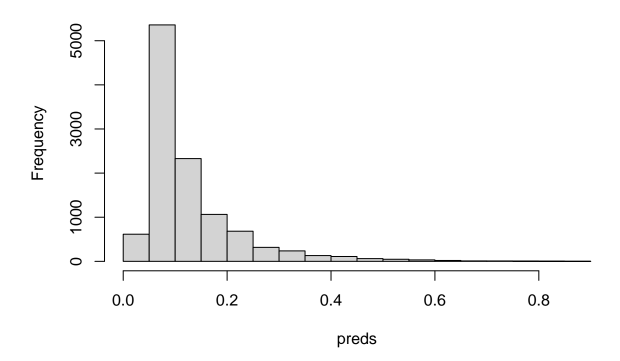
last_lightgbm_fit <-
    final_lightgbm_workflow %>%
    last_fit(df_split)
```



```
## [1] "Optimal Threshold: 0.11"
  Confusion Matrix and Statistics
##
##
       reference
##
           0
  data
##
      0 2792 202
##
      1 1335 402
##
##
                  Accuracy : 0.6751
                    95% CI: (0.6616, 0.6885)
##
      No Information Rate : 0.8723
##
##
      P-Value [Acc > NIR] : 1
##
                     Kappa : 0.19
##
##
   Mcnemar's Test P-Value : <2e-16
##
##
##
               Sensitivity: 0.6765
               Specificity: 0.6656
##
##
            Pos Pred Value : 0.9325
##
            Neg Pred Value: 0.2314
##
                Prevalence: 0.8723
##
            Detection Rate: 0.5902
##
      Detection Prevalence : 0.6328
##
         Balanced Accuracy: 0.6710
##
          'Positive' Class: 0
##
##
```

Minutes to run: 2.366

Histogram of preds



Minutes to run:

0.007

GLM

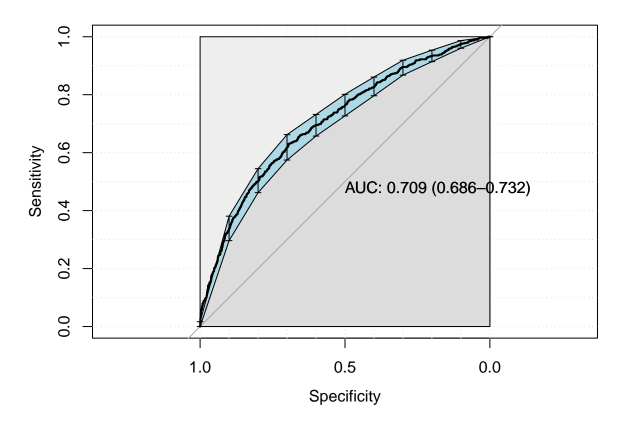
```
glmnet_recipe <-
    recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
    step_novel(all_nominal_predictors()) %>%
    step_unknown(all_nominal_predictors()) %>%
    step_other(all_nominal_predictors(), threshold = 0.05, other = ".merged") %>%
    step_dummy(all_nominal_predictors()) %>%
    step_zv(all_predictors()) %>%
    step_normalize(all_numeric_predictors())
```

```
logistic_reg(penalty = 0) %>%
set_mode("classification") %>%
set_engine("glmnet")

glmnet_workflow <-
    workflow() %>%
    add_recipe(glmnet_recipe) %>%
    add_model(glmnet_spec)

glm_fit <- glmnet_workflow %>%
    fit(df_train)

glmnet_auc <- validation(glm_fit, df_test)</pre>
```

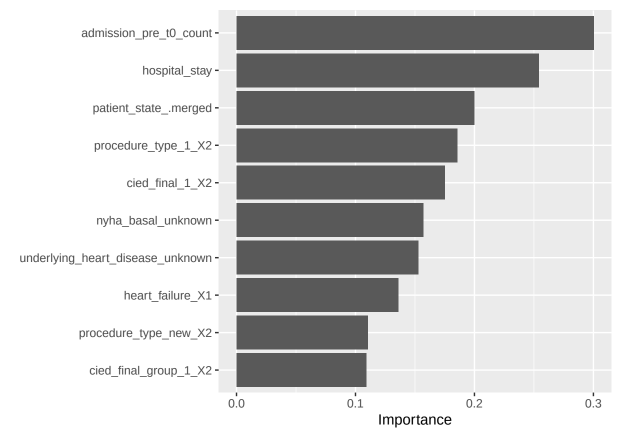


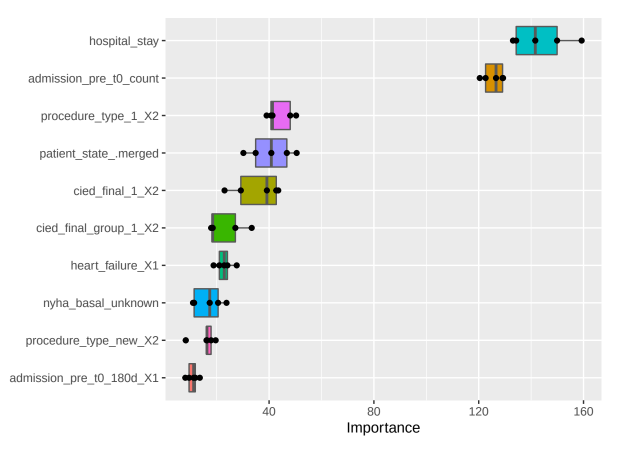
```
## [1] "Optimal Threshold: 0.12"
##
  Confusion Matrix and Statistics
##
##
       reference
##
           0
  data
                1
##
      0 2866
              223
      1 1261 381
##
##
##
                  Accuracy : 0.6863
##
                    95% CI : (0.6729, 0.6995)
##
       No Information Rate : 0.8723
##
       P-Value [Acc > NIR] : 1
##
                     Kappa : 0.1876
##
##
    Mcnemar's Test P-Value : <2e-16
##
##
               Sensitivity: 0.6945
##
##
               Specificity: 0.6308
```

```
Neg Pred Value : 0.2320
##
##
                Prevalence: 0.8723
##
            Detection Rate: 0.6058
      Detection Prevalence : 0.6529
##
##
         Balanced Accuracy: 0.6626
##
          'Positive' Class: 0
##
##
pfun_glmnet <- function(object, newdata) predict(object, newx = newdata)</pre>
extract_vip(glm_fit, pred_wrapper = pfun_glmnet,
            reference_class = "1", method = 'model')
```

Pos Pred Value : 0.9278

##





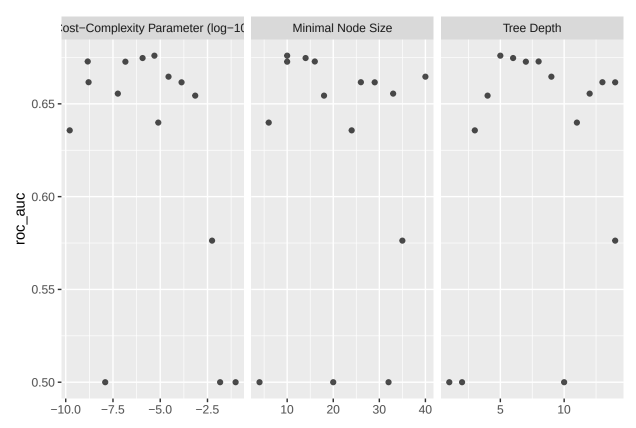
Minutes to run:

2.061

Decision Tree

```
tree_recipe <-
 recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
  step_novel(all_nominal_predictors()) %>%
  step_unknown(all_nominal_predictors()) %>%
  step_other(all_nominal_predictors(), threshold = 0.05, other = ".merged") %>%
  step_dummy(all_nominal_predictors()) %>%
  step_zv(all_predictors())
tree_spec <-
  decision_tree(cost_complexity = tune(),
                tree_depth = tune(),
                min_n = tune()) %>%
 set mode("classification") %>%
 set_engine("rpart")
tree_grid <- grid_latin_hypercube(cost_complexity(),</pre>
                                  tree_depth(),
                                  min_n(),
                                  size = grid_size)
tree_workflow <-
  workflow() %>%
  add_recipe(tree_recipe) %>%
  add_model(tree_spec)
tree_tune <-
 tree_workflow %>%
 tune_grid(resamples = df_folds,
            grid = tree_grid)
```

```
tree_tune %>%
  collect_metrics()
autoplot(tree_tune, metric = "roc_auc")
```



```
tree_tune %>%
    show_best("roc_auc")

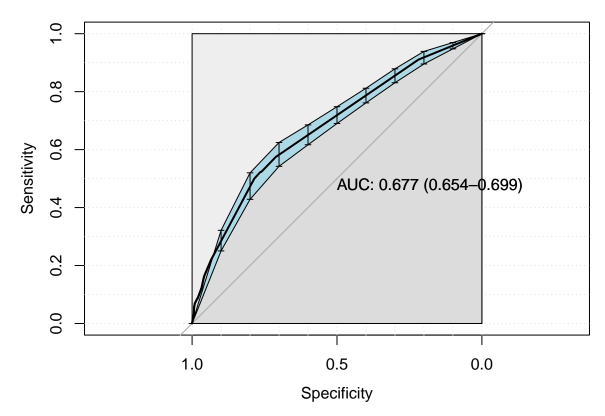
best_tree <- tree_tune %>%
    select_best("roc_auc")

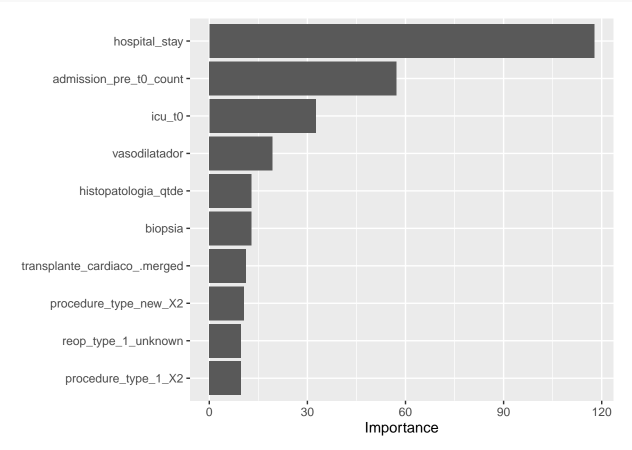
final_tree_workflow <-
    tree_workflow %>%
    finalize_workflow(best_tree)

last_tree_fit <-
    final_tree_workflow %>%
    last_fit(df_split)

final_tree_fit <- extract_workflow(last_tree_fit)

tree_auc <- validation(final_tree_fit, df_test)</pre>
```





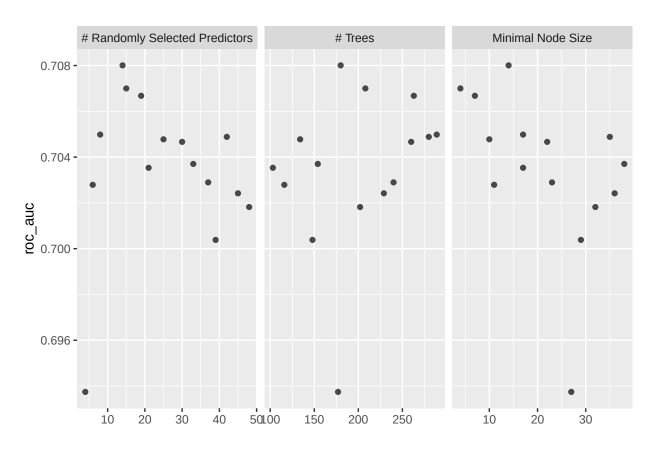
```
# extract_vip(final_tree_fit, pred_wrapper = predict,
# reference_class = "1", use_matrix = FALSE,
```

```
# method = 'permute')
```

Minutes to run: 1.879

Random Forest

```
rf_recipe <-
  recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula,
         data = df_train) %>%
  step_novel(all_nominal_predictors()) %>%
  step_unknown(all_nominal_predictors()) %>%
  step_other(all_nominal_predictors(), threshold = 0.05, other = ".merged") %>%
  step_dummy(all_nominal_predictors()) %>%
  step_zv(all_predictors()) %>%
  step_impute_mean(all_numeric_predictors())
rf_spec <-
  rand_forest(mtry = tune(),
              trees = tune(),
              min_n = tune()) %>%
  set_mode("classification") %>%
  set_engine("randomForest",
             probability = TRUE,
             nthread = 8)
rf_grid <- grid_latin_hypercube(mtry(range = c(1L, 50L)),</pre>
                                trees(range = c(100L, 300L)),
                                min_n(),
                                size = grid_size)
rf_workflow <-
  workflow() %>%
  add_recipe(rf_recipe) %>%
  add_model(rf_spec)
rf_tune <-
  rf_workflow %>%
  tune_grid(resamples = df_folds,
            grid = rf_grid)
rf_tune %>%
  collect_metrics()
autoplot(rf_tune, metric = "roc_auc")
```



```
rf_tune %>%
    show_best("roc_auc")

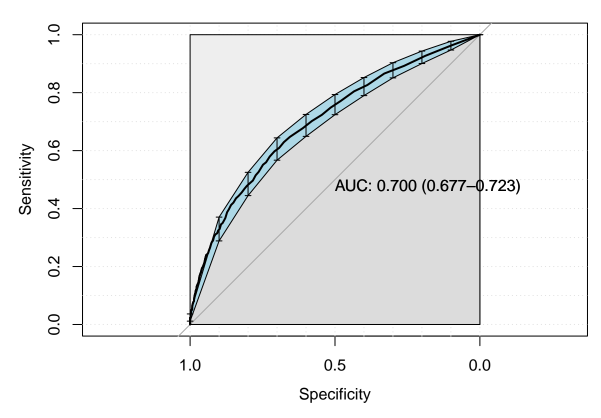
best_rf <- rf_tune %>%
    select_best("roc_auc")

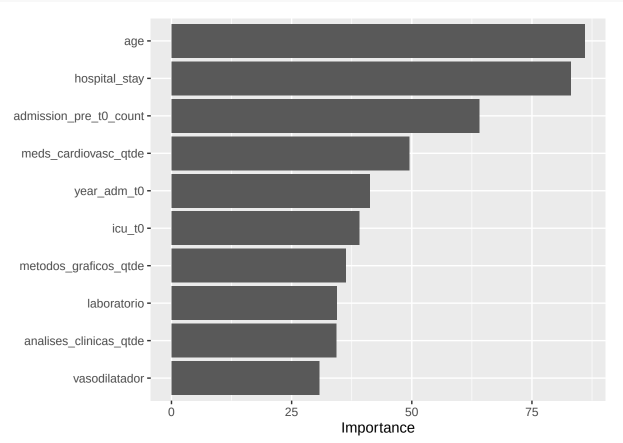
final_rf_workflow <-
    rf_workflow %>%
    finalize_workflow(best_rf)

last_rf_fit <-
    final_rf_workflow %>%
    last_fit(df_split)

final_rf_fit <- extract_workflow(last_rf_fit)

rf_auc <- validation(final_rf_fit, df_test)</pre>
```

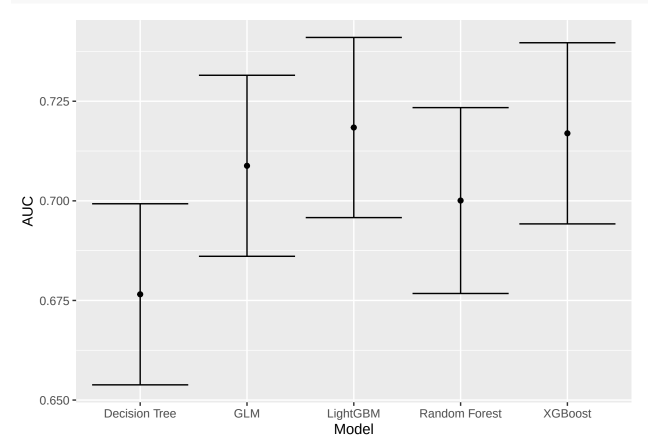




```
# extract_vip(final_rf_fit, pred_wrapper = predict,
# reference_class = "1", use_matrix = FALSE,
# method = 'permute')
```

Minutes to run: 22.712

Models Comparison



saveRDS(df_auc, sprintf("./auxiliar/model_selection/performance/%s.RData", outcome_column))

Minutes to run: 0.008