## Model Selection - readmission 30d

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### **Imports**

```
library(tidyverse)
library(yaml)
library(tidymodels)
library(usemodels)
library(vip)
library(bonsai)
library(lightgbm)
library(caret)
```

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))</pre>
```

Minutes to run: 0.006

#### Eligible features

```
} else {
  features = base::intersect(eligible_features, features_list)
gluedown::md_order(features, seq = TRUE, pad = TRUE)
## 01. education_level
## 02. underlying_heart_disease
## 03. heart disease
## 04. nyha_basal
## 05. prior_mi
## 06. heart_failure
## 07. transplant
## 08. endocardites
## 09. hemodialysis
## 10. comorbidities_count
## 11. procedure_type_1
## 12. reop_type_1
## 13. procedure_type_new
## 14. cied_final_1
## 15. cied_final_group_1
## 16. admission_pre_t0_count
## 17. admission_pre_t0_180d
## 18. icu_t0
## 19. dialysis_t0
## 20. admission_t0_emergency
## 21. aco
## 22. antiarritmico
## 23. betabloqueador
## 24. ieca_bra
## 25. dva
## 26. digoxina
## 27. estatina
## 28. diuretico
## 29. vasodilatador
## 30. insuf_cardiaca
## 31. espironolactona
## 32. bloq_calcio
## 33. antiplaquetario_ev
## 34. insulina
## 35. anticonvulsivante
## 36. psicofarmacos
## 37. antifungico
## 38. antiviral
## 39. classe_meds_qtde
## 40. meds_cardiovasc_qtde
## 41. meds_antimicrobianos
## 42. cec
## 43. transplante_cardiaco
## 44. outros_proced_cirurgicos
## 45. icp
## 46. intervencao_cv
## 47. cateterismo
## 48. eletrofisiologia
## 49. cateter_venoso_central
## 50. proced_invasivos_qtde
## 51. cve_desf
```

## 52. transfusao
## 53. equipe\_multiprof

## 54. ecg ## 55. holter

```
## 56. metodos_graficos_qtde
## 57. laboratorio
## 58. cultura
## 59. analises_clinicas_qtde
## 60. citologia
## 61. biopsia
## 62. histopatologia_qtde
## 63. angio_rm
## 64. angio_tc
## 65. cintilografia
## 66. ecocardiograma
## 67. endoscopia
## 68. flebografia
## 69. pet_ct
## 70. ultrassom
## 71. tomografia
## 72. radiografia
## 73. ressonancia
## 74. exames_imagem_qtde
## 75. bic
## 76. mpp
Minutes to run: 0
```

## Train test split (70%/30%)

```
set.seed(42)

if (outcome_column == 'readmission_30d') {
    df_split <- readRDS("./dataset/split_object.rds")
} else {
    df_split <- initial_split(df, prop = .7, strata = all_of(outcome_column))
}

df_train <- training(df_split) %>% dplyr::select(all_of(c(features, outcome_column)))
df_test <- testing(df_split) %>% dplyr::select(all_of(c(features, outcome_column)))
```

Minutes to run: 0.005

## Global parameters

Minutes to run: 0

#### **Functions**

```
validation = function(model_fit, new_data, plot=TRUE) {
   library(pROC)
   library(caret)

test_predictions_prob <-
    predict(model_fit, new_data = new_data, type = "prob") %>%
    rename_at(vars(starts_with(".pred_")), ~ str_remove(., ".pred_")) %>%
```

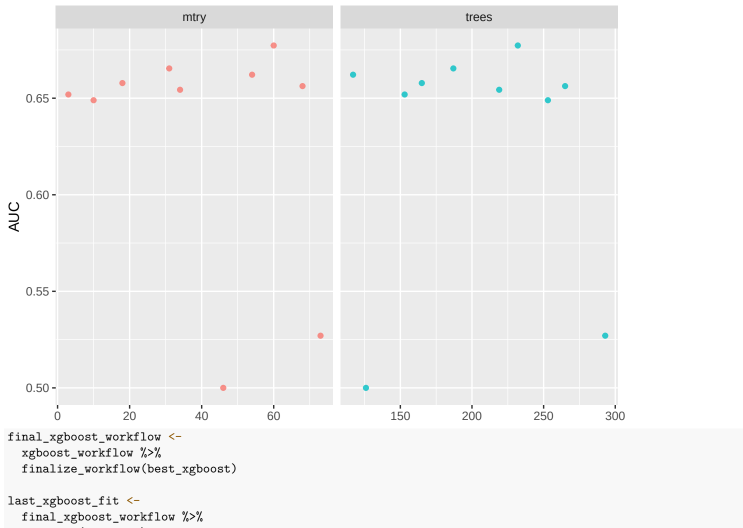
```
.$`1`
  pROC obj <- roc(
    new_data[[outcome_column]],
    test_predictions_prob,
    direction = "<",</pre>
    levels = c(0, 1),
    smoothed = TRUE,
    ci = TRUE,
    ci.alpha = 0.9,
    stratified = FALSE,
    plot = plot,
    auc.polygon = TRUE,
    max.auc.polygon = TRUE,
    grid = TRUE,
    print.auc = TRUE,
    show.thres = TRUE
  test_predictions_class <-</pre>
    predict(model_fit, new_data = new_data, type = "class") %>%
    rename_at(vars(starts_with(".pred_")), ~ str_remove(., ".pred_")) %>%
    .$class
  conf_matrix <- table(test_predictions_class, new_data[[outcome_column]])</pre>
  if (plot) {
    sens.ci <- ci.se(pROC_obj)</pre>
    plot(sens.ci, type = "shape", col = "lightblue")
    plot(sens.ci, type = "bars")
    confusionMatrix(conf_matrix) %>% print
  return(pROC_obj)
}
```

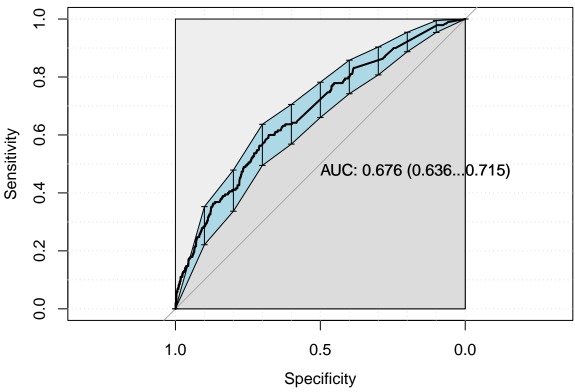
Minutes to run: 0

## Boosted Tree (XGBoost)

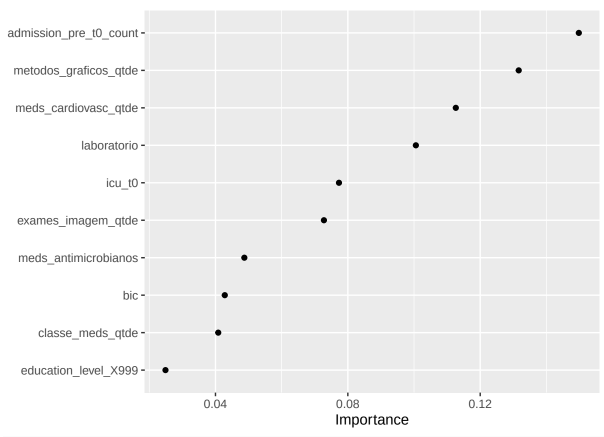
```
xgboost_recipe <-</pre>
  recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
  step_novel(all_nominal_predictors()) %>%
  step_unknown(all_nominal_predictors()) %>%
  step_other(all_nominal_predictors(), threshold = 0.05, other=".merged") %>%
  step_dummy(all_nominal_predictors(),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

```
xgboost_grid <- grid_latin_hypercube(</pre>
  finalize(mtry(), df_train),
  dials::trees(range = c(100L, 300L)),
 min_n(),
 tree_depth(),
 learn_rate(),
 loss_reduction(),
  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
     mtry trees min_n tree_depth learn_rate loss_reduction .metric .estimator mean
                                                                                     n std_err .config
##
    <int> <int> <int>
                         <int>
                                    <dbl>
                                                  <dbl> <chr>
                                                                 <chr>
                                                                           <dbl> <int>
                                                                                       <dbl> <chr>
       60 232 14
## 1
                             3 0.0108
                                            0.000746 roc_auc binary
                                                                           0.677 4 0.0106 Preprocessor
                            12 0.0000653 0.0000000541 roc_auc binary
## 2
       31 187
                  6
                                                                           0.665
                                                                                    4 0.0111 Preprocessor
## 3
       54 117 13
                             8 0.0789 2.62
                                                        roc_auc binary
                                                                           0.662 4 0.0151 Preprocessor
       18 165
                             2 0.0000134 0.141
## 4
                   31
                                                        roc_auc binary
                                                                           0.658
                                                                                     4 0.00477 Preprocessor1
## 5
       68 265
                  20
                             15 0.00108
                                           0.0000314
                                                        roc_auc binary
                                                                           0.656
                                                                                     4 0.0113 Preprocessor
best_xgboost <- xgboost_tune %>%
 select_best("roc_auc")
xgboost_tune %>%
  collect_metrics() %>%
 filter(.metric == "roc_auc") %>%
  select(mean, mtry:trees) %>%
 pivot_longer(mtry:trees,
              values_to = "value",
              names_to = "parameter"
  ) %>%
  ggplot(aes(value, mean, color = parameter)) +
  geom_point(alpha = 0.8, show.legend = FALSE) +
  facet_wrap(~parameter, scales = "free_x") +
  labs(x = NULL, y = "AUC")
```





```
##
   Confusion Matrix and Statistics
##
##
##
   test_predictions_class
##
                        0 4540
                                 190
##
                        1
                             0
##
##
                  Accuracy : 0.9598
                    95% CI : (0.9538, 0.9652)
##
##
       No Information Rate: 0.9598
       P-Value [Acc > NIR] : 0.5193
##
##
                     Kappa: 0
##
##
##
    Mcnemar's Test P-Value : <2e-16
##
##
               Sensitivity: 1.0000
##
               Specificity: 0.0000
##
            Pos Pred Value : 0.9598
##
            Neg Pred Value :
##
                Prevalence: 0.9598
##
            Detection Rate: 0.9598
##
      Detection Prevalence : 1.0000
##
         Balanced Accuracy: 0.5000
##
##
          'Positive' Class : 0
##
final_xgboost_fit %>%
  fit(data = df_train) %>%
  extract_fit_parsnip() %>%
  vip(geom = "point")
```



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

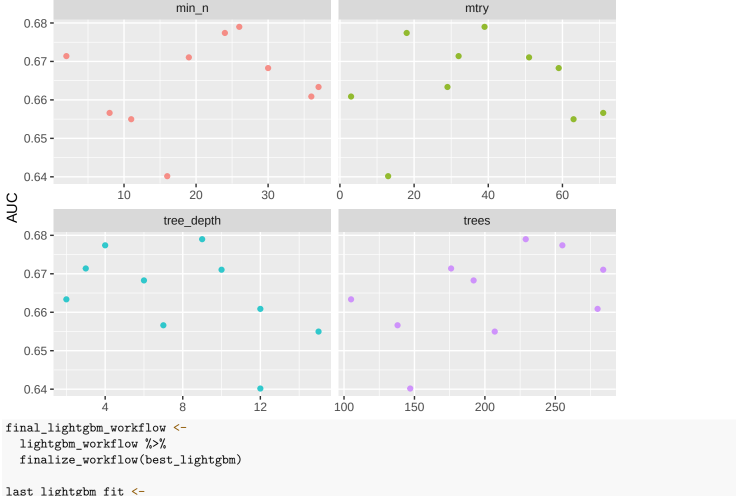
saveRDS(
    xgboost_parameters,
    file = sprintf(
        "./auxiliar/model_selection/hyperparameters/xgboost_%s.rds",
        outcome_column
    )
)
```

Minutes to run: 2.737

## 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_impute_mean(all_numeric_predictors()) %>%
  step_zv(all_predictors())
lightgbm_spec <- boost_tree(</pre>
 mtry = tune(),
  trees = tune(),
 min_n = tune(),
  tree_depth = tune(),
  learn_rate = tune(),
  loss_reduction = tune(),
  sample_size = 1
) %>%
```

```
set_engine("lightgbm") %>%
  set_mode("classification")
lightgbm_grid <- grid_latin_hypercube(</pre>
  finalize(mtry(), df_train),
  dials::trees(range = c(100L, 300L)),
 min_n(),
 tree_depth(),
 learn_rate(),
 loss_reduction(),
 size = grid_size
)
lightgbm_workflow <-</pre>
  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 12
##
     mtry trees min_n tree_depth
                                    learn_rate loss_reduction .metric .estimator mean
                                                                                          n std_err .config
##
    <int> <int> <int> <int>
                                         <dbl>
                                                      <dbl> <chr>
                                                                    <chr>
                                                                                 <dbl> <int> <dbl> <chr>
                             9 0.0000000168 6.67
       39 229 26
## 1
                                                              roc_auc binary
                                                                                 0.679 4 0.00702 Preprocess
## 2
       18 255
                   24
                              4 0.0000395
                                                0.985
                                                             roc_auc binary
                                                                                0.677
                                                                                          4 0.00875 Preprocess
       32 176
                   2
                              3 0.00139
                                                0.00000000245 roc_auc binary
## 3
                                                                                 0.671
                                                                                          4 0.00902 Preprocess
## 4
       51
            284
                   19
                             10 0.00000000965 0.0000000430 roc_auc binary
                                                                                 0.671
                                                                                           4 0.00701 Preprocess
                                                              roc_auc binary
## 5
       59
           192
                   30
                               6 0.00000104
                                                0.150
                                                                                 0.668
                                                                                           4 0.00578 Preprocess
best_lightgbm <- lightgbm_tune %>%
  select_best("roc_auc")
lightgbm_tune %>%
  collect_metrics() %>%
 filter(.metric == "roc_auc") %>%
  select(mean, mtry:tree_depth) %>%
 pivot_longer(mtry:tree_depth,
              values_to = "value",
              names_to = "parameter"
  ) %>%
  ggplot(aes(value, mean, color = parameter)) +
  geom_point(alpha = 0.8, show.legend = FALSE) +
  facet_wrap(~parameter, scales = "free_x") +
 labs(x = NULL, y = "AUC")
```



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

last_lightgbm_fit <-
    final_lightgbm_workflow %>%
    last_fit(df_split)

final_lightgbm_fit <- extract_workflow(last_lightgbm_fit)

lightgbm_auc <- validation(final_lightgbm_fit, df_test)</pre>
```

```
Sensitivity

AUC: 0.683 (0.644...0.722)

1.0 0.5 0.0

Specificity
```

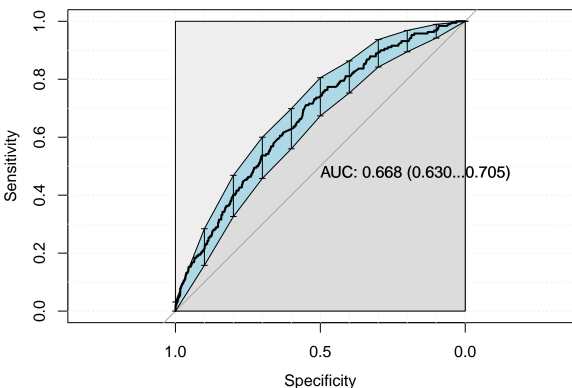
```
##
   Confusion Matrix and Statistics
##
##
##
   test_predictions_class
##
                        0 4540
                                190
##
                        1
                             0
##
##
                  Accuracy: 0.9598
                    95% CI : (0.9538, 0.9652)
##
##
       No Information Rate : 0.9598
##
       P-Value [Acc > NIR] : 0.5193
##
##
                     Kappa: 0
##
##
    Mcnemar's Test P-Value : <2e-16
##
##
               Sensitivity: 1.0000
##
               Specificity: 0.0000
##
            Pos Pred Value : 0.9598
##
            Neg Pred Value :
##
                Prevalence: 0.9598
##
            Detection Rate: 0.9598
##
      Detection Prevalence : 1.0000
##
         Balanced Accuracy: 0.5000
##
##
          'Positive' Class : 0
##
lightgbm_parameters <- lightgbm_tune %>%
  show_best("roc_auc", n=1) %>%
  select(trees, mtry, min_n, tree_depth, learn_rate, loss_reduction) %>%
  as.list
saveRDS(
  lightgbm_parameters,
  file = sprintf(
```

```
"./auxiliar/model_selection/hyperparameters/lightgbm_%s.rds",
   outcome_column
)
```

Minutes to run: 7.706

#### GLM

```
glmnet_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(), one_hot = TRUE) %>%
  step_zv(all_predictors()) %>%
  step_normalize(all_numeric_predictors())
glmnet_spec <-</pre>
  logistic_reg(penalty = 0) %>%
  set_mode("classification") %>%
  set_engine("glmnet")
glmnet_workflow <-</pre>
  workflow() %>%
  add_recipe(glmnet_recipe) %>%
  add_model(glmnet_spec)
glm_fit <- glmnet_workflow %>%
  fit(df_train)
glm_auc = validation(glm_fit, df_test)
```



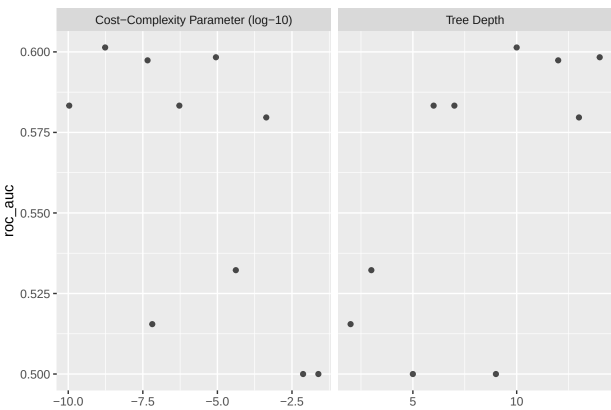
```
## |
## Confusion Matrix and Statistics
##
```

```
##
## test_predictions_class
##
                        0 4540 190
##
##
##
                  Accuracy: 0.9598
##
                    95% CI: (0.9538, 0.9652)
##
       No Information Rate: 0.9598
##
      P-Value [Acc > NIR] : 0.5193
##
##
                     Kappa: 0
##
##
   Mcnemar's Test P-Value : <2e-16
##
##
               Sensitivity: 1.0000
##
               Specificity: 0.0000
##
            Pos Pred Value: 0.9598
##
            Neg Pred Value :
##
                Prevalence: 0.9598
##
           Detection Rate: 0.9598
##
     Detection Prevalence: 1.0000
##
        Balanced Accuracy: 0.5000
##
          'Positive' Class : 0
##
##
```

Minutes to run: 0.087

#### **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(), one_hot = TRUE) %>%
  step_zv(all_predictors())
tree_spec <-
  decision_tree(cost_complexity = tune(),
                tree_depth = tune()) %>%
  set_mode("classification") %>%
  set_engine("rpart")
tree_grid <- grid_latin_hypercube(cost_complexity(),</pre>
                                  tree_depth(),
                                   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()
```



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

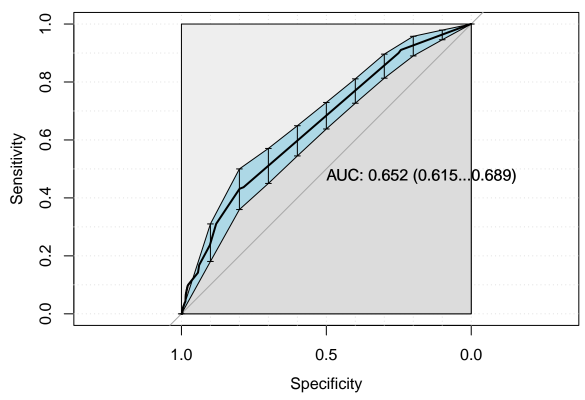
best_tree <- tree_tune %>%
    select_best("roc_auc")

final_tree_workflow <-
    tree_workflow %>%
    finalize_workflow(best_tree)

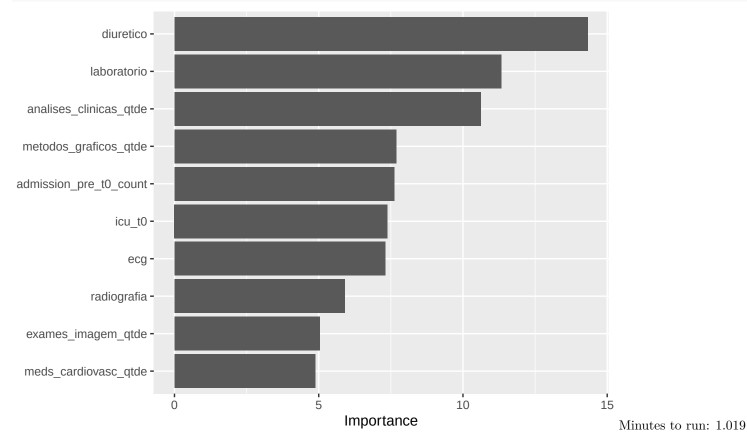
last_tree_fit <-
    final_tree_workflow %>%
    last_fit(df_split)

final_tree_fit <- extract_workflow(last_tree_fit)

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

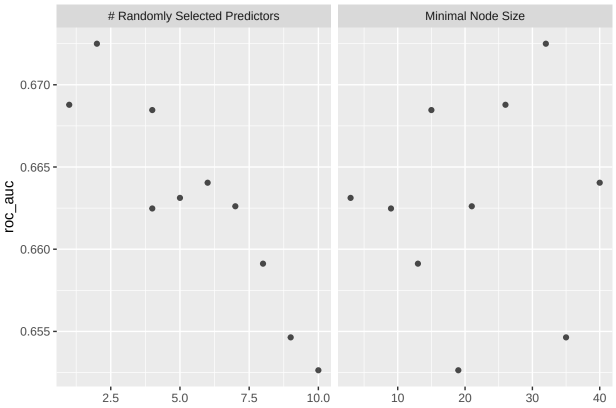


```
if (tree_auc$auc > 0.55){
  final_tree_fit %>%
    extract_fit_parsnip() %>%
    vip()
}
```



#### **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(), one_hot = TRUE) %>%
  step_zv(all_predictors()) %>%
  step_impute_mean(all_numeric_predictors())
rf_spec <-
  rand_forest(mtry = tune(),
              trees = 100,
              min_n = tune()) %>%
  set_mode("classification") %>%
  set_engine("ranger")
rf_grid <- grid_latin_hypercube(mtry(range = c(1, 10)),</pre>
                                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")

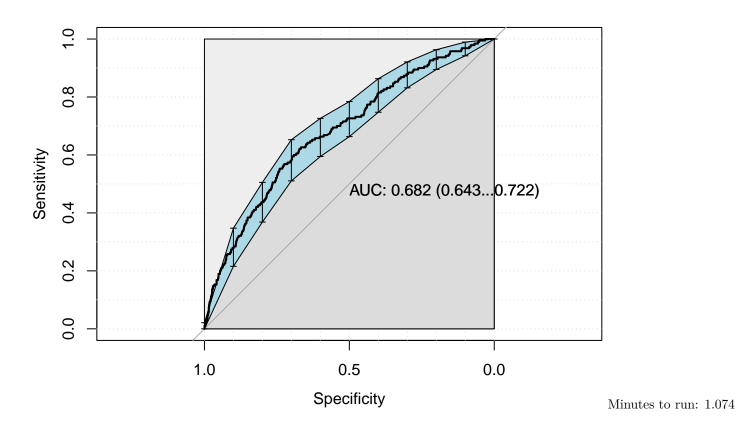
best_rf <- rf_tune %>%
    select_best("roc_auc")

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>
```



### **KNN**

```
# knn_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(), one_hot = TRUE) %>%
#
    step_zv(all_predictors()) %>%
#
    step_impute_mean(all_numeric_predictors())
#
# knn_spec <-
    nearest_neighbor(neighbors = tune(),
#
                     weight_func = tune(),
#
                     dist_power = tune()) %>%
    set_mode("classification") %>%
#
    set_engine("kknn")
  knn_grid <- grid_latin_hypercube(neighbors(),</pre>
                                    weight_func(),
#
                                    dist_power(),
#
                                    size = 5)
#
# knn_workflow <-</pre>
    workflow() %>%
#
    add_recipe(knn_recipe) %>%
    add_model(knn_spec)
# knn_tune <-
#
    knn_workflow %>%
    tune_grid(resamples = df_folds,
              grid = knn_grid
# knn_tune %>%
    collect_metrics()
```

```
# autoplot(knn_tune, metric = "roc_auc")
#
# knn_tune %>%
# show_best("roc_auc")
#
# best_knn <- knn_tune %>%
# select_best("roc_auc")
#
# final_knn_workflow <-
# knn_workflow %>%
# finalize_workflow(best_knn)
#
# last_knn_fit <-
# final_knn_workflow %>%
# last_fit(df_split)
#
# final_knn_fit <- extract_workflow(last_knn_fit)
#
# knn_auc = validation(final_knn_fit, df_test)</pre>
```

Minutes to run: 0

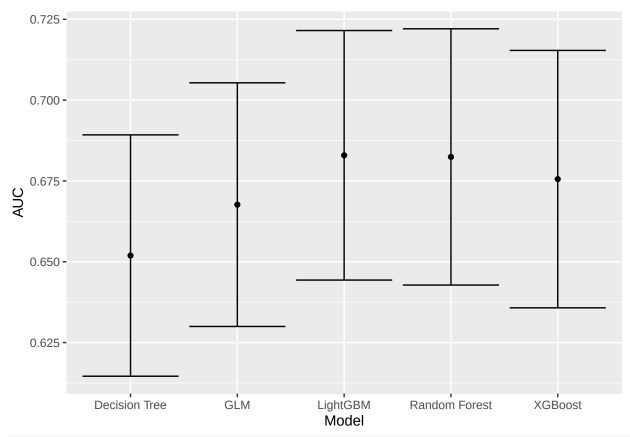
#### SVM

```
# svm_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(), one_hot = TRUE) %>%
#
   step_zv(all_predictors()) %>%
   step_impute_mean(all_numeric_predictors())
#
# svm_spec <-
#
   svm_rbf(cost = tune(), rbf_sigma = tune()) %>%
   set_mode("classification") %>%
#
   set_engine("kernlab")
# svm_grid <- grid_latin_hypercube(cost(),</pre>
#
                                    rbf\_sigma(),
#
                                    size = grid\_size)
#
# svm_workflow <-
  workflow() %>%
   add_recipe(svm_recipe) %>%
#
   add_model(svm_spec)
# svm_tune <-
   svm_workflow %>%
#
    tune\_grid(resamples = df\_folds,
              grid = 5)
#
# svm_tune %>%
#
   collect_metrics()
# autoplot(svm_tune, metric = "roc_auc")
# svm_tune %>%
   show_best("roc_auc")
```

```
# best_sum <- sum_tune %>%
# select_best("roc_auc")
#
# final_sum_workflow <-
# sum_workflow %>%
# finalize_workflow(best_sum)
#
# last_sum_fit <-
# final_sum_workflow %>%
# last_fit(df_split)
#
# final_sum_fit <- extract_workflow(last_sum_fit)
#
# sum_auc = validation(final_sum_fit, df_test)</pre>
```

Minutes to run: 0

# **Models Comparison**



saveRDS(df\_auc, sprintf("./auxiliar/model\_selection/performance/%s.RData", outcome\_column))

Minutes to run: 0.002