Model

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Imports

```
library(tidyverse)
library(yaml)
library(tidymodels)
library(usemodels)
library(vip)
```

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</pre>
```

Filtering eligible pacients

```
df = df %>%
  filter(disch_outcomes_t0 == 0)

df %>% dim
## [1] 15766 236
```

Eligible features

Train test split (70%/30%)

Imputation

Functions

```
validation = function(model_fit, new_data) {
   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`

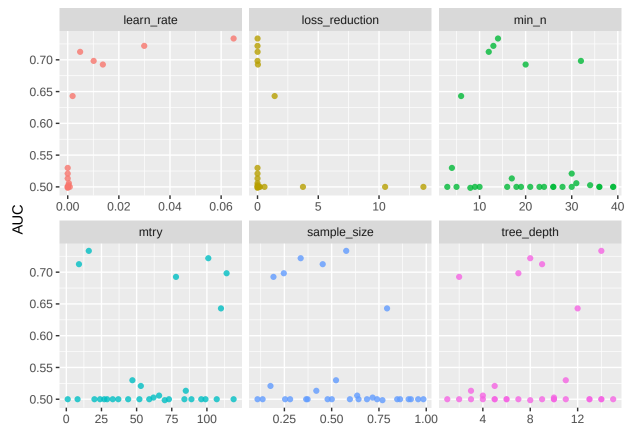
pROC_obj <- roc(
   new_data[[outcome_column]],
   test_predictions_prob,
   smoothed = TRUE,
   # arguments for ci</pre>
```

```
ci = TRUE,
  ci.alpha = 0.9,
  stratified = FALSE,
  # arguments for plot
  plot = TRUE,
  auc.polygon = TRUE,
  max.auc.polygon = TRUE,
  grid = TRUE,
  print.auc = TRUE,
  show.thres = TRUE
)
sens.ci <- ci.se(pROC_obj)</pre>
plot(sens.ci, type = "shape", col = "lightblue")
plot(sens.ci, type = "bars")
test_predictions_class <-</pre>
  predict(model_fit, new_data = new_data, type = "class") %>%
  rename_at(vars(starts_with(".pred_")), ~ str_remove(., ".pred_")) %%
conf_matrix = table(test_predictions_class, new_data[[outcome_column]])
confusionMatrix(conf_matrix) %>% print
return(pROC_obj)
```

$\mathbf{XGBoost}$

```
xgboost_recipe <-
  recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
  step_string2factor(one_of("sex", "race", "education_level", "patient_state",
    "underlying_heart_disease", "heart_disease", "nyha_basal", "hypertension",
    "prior_mi", "heart_failure", "af", "cardiac_arrest", "transplant", "valvopathy",
    "endocardites", "diabetes", "renal_failure", "hemodialysis", "stroke",
    "copd", "cancer", "surgery_count", "procedure_type_1", "reop_type_1", "cied_final_1",
    "death_intraop_1", "dialysis_hosp", "icu_hosp", "admission_pre_t0_180d", "ventilacao_mecanica")) %>%
  step_novel(all_nominal_predictors()) %>%
  step_dummy(all_nominal_predictors(), one_hot = TRUE) %>%
  step_zv(all_predictors())
xgboost_spec <- boost_tree(</pre>
 trees = 500,
 tree_depth = tune(),
 min_n = tune(),
 loss_reduction = tune(),
 sample_size = tune(),
 mtry = tune(),
  learn_rate = tune()
  set_engine("xgboost") %>%
  set_mode("classification")
xgboost_grid <- grid_latin_hypercube(</pre>
  tree_depth(),
 min_n(),
 loss_reduction(),
  sample_size = sample_prop(),
  finalize(mtry(), df_train),
```

```
learn_rate(),
  size = 30
set.seed(234)
df_folds <- vfold_cv(df_train, v = 5, strata = all_of(outcome_column))</pre>
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 min_n tree_depth learn_rate
##
    <int> <int>
                 <int>
                                <dbl>
## 1
                       14
                              0.0649
       16 14
## 2
      101 13
                        8 0.0299
## 3
       9
           12
                        9 0.00485
## 4
     114
             32
                         7
                              0.0101
## 5
      78
             20
                         2
                              0.0137
## # ... with 8 more variables:
## #
      loss_reduction <dbl>,
## #
      sample_size <dbl>, .metric <chr>,
## #
       .estimator <chr>, mean <dbl>,
## #
      n <int>, std_err <dbl>, .config <chr>
best_xgboost <- xgboost_tune %>%
  select_best("roc_auc")
xgboost_tune %>%
  collect_metrics() %>%
  filter(.metric == "roc_auc") %>%
  select(mean, mtry:sample_size) %>%
 pivot_longer(mtry:sample_size,
              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")
```



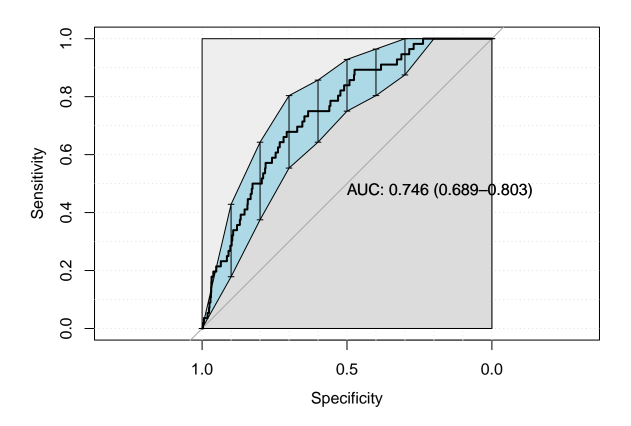
```
final_xgboost_workflow <-
    xgboost_workflow %>%
    finalize_workflow(best_xgboost)

last_xgboost_fit <-
    final_xgboost_workflow %>%
    last_fit(df_split)

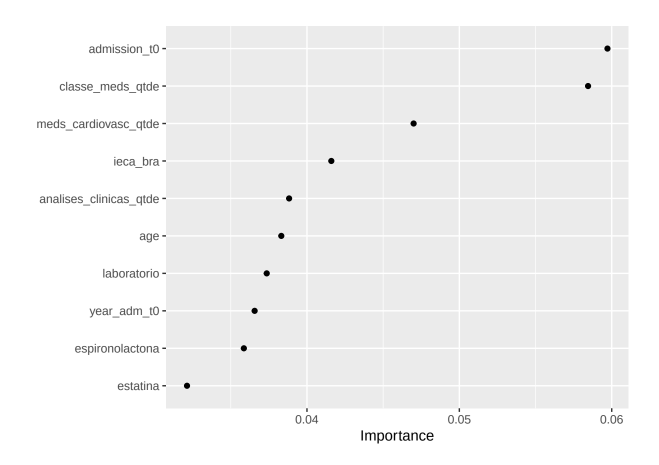
final_xgboost_fit <- extract_workflow(last_xgboost_fit)

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

Setting levels: control = 0, case = 1
Setting direction: controls < cases</pre>



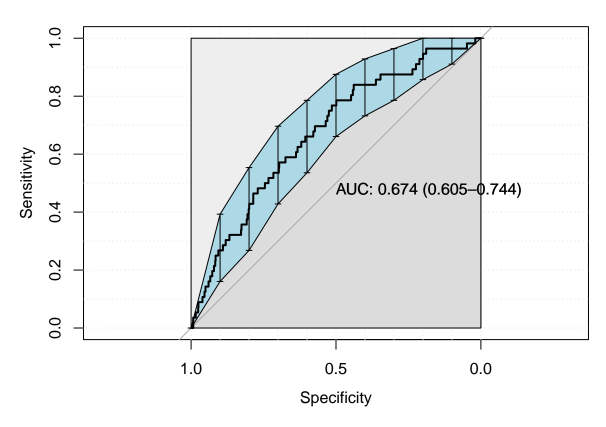
```
Confusion Matrix and Statistics
##
##
##
   test_predictions_class
                                   1
##
                        0 4674
                                  56
##
                        1
                                   0
##
                  Accuracy : 0.9882
##
##
                    95% CI: (0.9847, 0.991)
       No Information Rate: 0.9882
##
       P-Value [Acc > NIR] : 0.5355
##
##
##
                     Kappa: 0
##
##
    Mcnemar's Test P-Value : 1.987e-13
##
##
               Sensitivity: 1.0000
##
               Specificity: 0.0000
##
            Pos Pred Value: 0.9882
##
            Neg Pred Value :
                Prevalence: 0.9882
##
##
            Detection Rate: 0.9882
##
      Detection Prevalence: 1.0000
##
         Balanced Accuracy : 0.5000
##
##
          'Positive' Class : 0
##
final_xgboost_fit %>%
  fit(data = df_train) %>%
  extract_fit_parsnip() %>%
  vip(geom = "point")
```



GLM

```
glmnet_recipe <-</pre>
  recipe(formula = sprintf("%s ~ .", outcome_column) %% as.formula, data = df_train) %>%
  step_string2factor(one_of("sex", "race", "education_level", "patient_state",
    "underlying_heart_disease", "heart_disease", "nyha_basal", "hypertension",
    "prior_mi", "heart_failure", "af", "cardiac_arrest", "transplant", "valvopathy",
    "endocardites", "diabetes", "renal_failure", "hemodialysis", "stroke",
    "copd", "cancer", "surgery_count", "procedure_type_1", "reop_type_1", "cied_final_1",
"death_intraop_1", "dialysis_hosp", "icu_hosp", "admission_pre_t0_180d", "ventilacao_mecanica")) %>%
  step_novel(all_nominal_predictors()) %>%
  step_dummy(all_nominal_predictors()) %>%
  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_fit %>%
   pull_workflow_fit() %>%
    tidy()
```

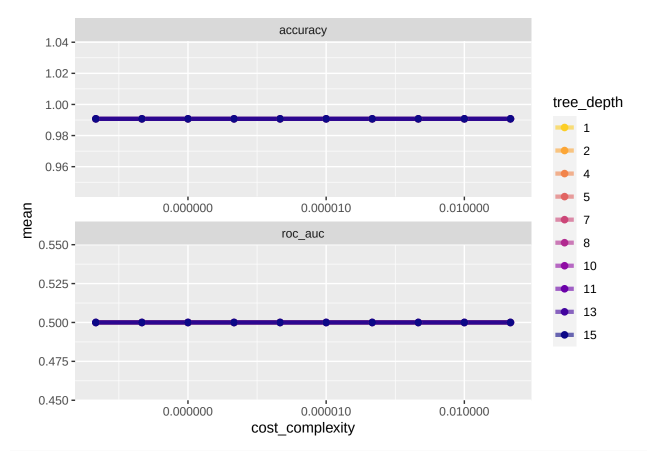
```
## Setting direction: controls < cases
```



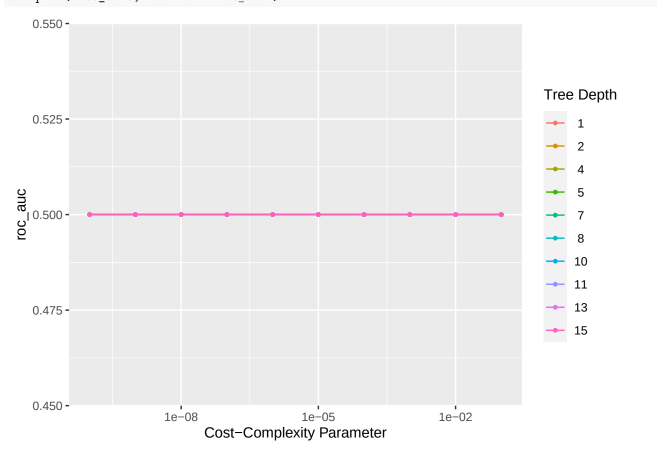
```
## Confusion Matrix and Statistics
##
##
##
  test_predictions_class
##
                        0 4674
                                  56
##
                        1
                              0
##
##
                  Accuracy : 0.9882
                    95% CI: (0.9847, 0.991)
##
##
       No Information Rate : 0.9882
##
       P-Value [Acc > NIR] : 0.5355
##
                     Kappa: 0
##
##
    Mcnemar's Test P-Value : 1.987e-13
##
##
##
               Sensitivity: 1.0000
##
               Specificity: 0.0000
            Pos Pred Value: 0.9882
##
##
            Neg Pred Value :
                Prevalence: 0.9882
##
##
            Detection Rate: 0.9882
##
      Detection Prevalence: 1.0000
##
         Balanced Accuracy: 0.5000
##
##
          'Positive' Class : 0
##
```

Decision Tree

```
tree_recipe <-
  recipe(formula = sprintf("%s ~ .", outcome_column) %% as.formula, data = df_train) %>%
  step_string2factor(one_of("sex", "race", "education_level", "patient_state",
    "underlying_heart_disease", "heart_disease", "nyha_basal", "hypertension",
    "prior_mi", "heart_failure", "af", "cardiac_arrest", "transplant", "valvopathy",
    "endocardites", "diabetes", "renal_failure", "hemodialysis", "stroke",
    "copd", "cancer", "surgery_count", "procedure_type_1", "reop_type_1", "cied_final_1",
   "death_intraop_1", "dialysis_hosp", "icu_hosp", "admission_pre_t0_180d", "ventilacao_mecanica")) %>%
  step_novel(all_nominal_predictors()) %>%
  step_zv(all_predictors())
tree_spec <-
  decision_tree(cost_complexity = tune(),
                tree_depth = tune()) %>%
  set_mode("classification") %>%
 set_engine("rpart")
tree_grid <- grid_regular(cost_complexity(),</pre>
                          tree_depth(),
                          levels = 10)
set.seed(234)
df_folds <- vfold_cv(df_train, v = 5, strata = all_of(outcome_column))</pre>
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 %>%
  collect metrics() %>%
 mutate(tree_depth = factor(tree_depth)) %>%
  ggplot(aes(cost_complexity, mean, color = tree_depth)) +
  geom_line(size = 1.5, alpha = 0.6) +
 geom_point(size = 2) +
  facet_wrap(~ .metric, scales = "free", nrow = 2) +
  scale_x_log10(labels = scales::label_number()) +
  scale_color_viridis_d(option = "plasma", begin = .9, end = 0)
```







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

best_tree <- tree_tune %>%
```

```
select_best("roc_auc")

final_tree_workflow <-
    tree_workflow %>%
    finalize_workflow(best_tree)

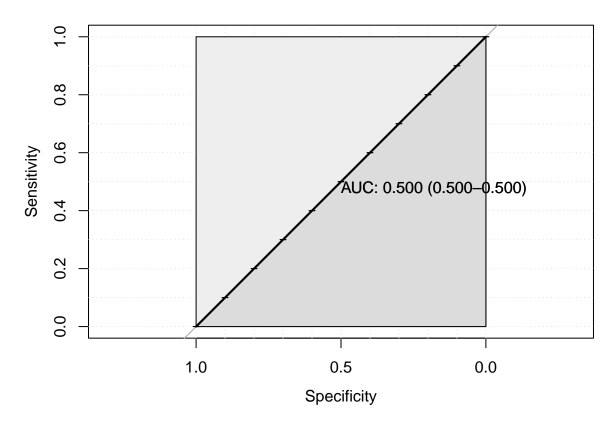
last_tree_fit <-
    final_tree_workflow %>%
    last_fit(df_split)

final_tree_fit <- extract_workflow(last_tree_fit)

tree_auc = validation(final_tree_fit, df_test)

## Setting levels: control = 0, case = 1

## Setting direction: controls < cases</pre>
```

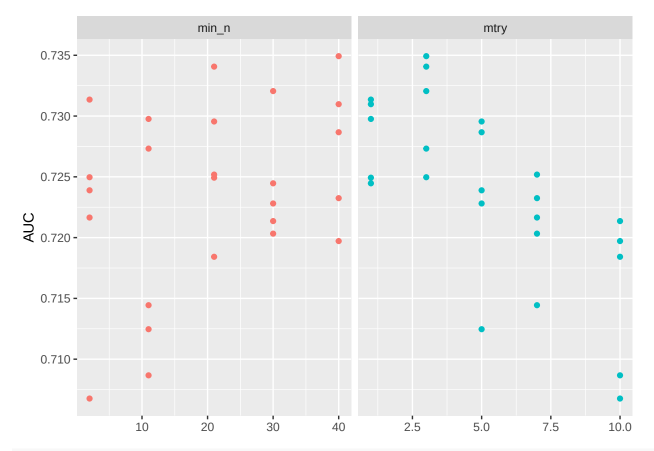


```
if (tree_auc$auc > 0.55){
  final_tree_fit %>%
    extract_fit_parsnip() %>%
    vip()
}
```

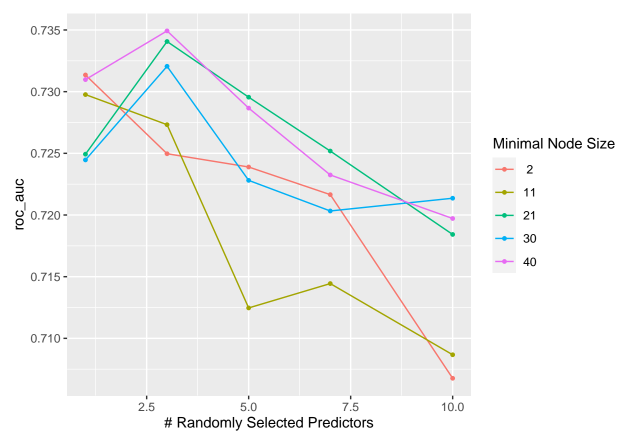
Random Forest

```
rf_recipe <-
  recipe(formula = sprintf("%s ~ .", outcome_column) %>% as.formula, data = df_train) %>%
  step_string2factor(one_of("sex", "race", "education_level", "patient_state",
    "underlying_heart_disease", "heart_disease", "nyha_basal", "hypertension",
    "prior_mi", "heart_failure", "af", "cardiac_arrest", "transplant", "valvopathy",
    "endocardites", "diabetes", "renal_failure", "hemodialysis", "stroke",
    "copd", "cancer", "surgery_count", "procedure_type_1", "reop_type_1", "cied_final_1",
```

```
"death_intraop_1", "dialysis_hosp", "icu_hosp", "admission_pre_t0_180d", "ventilacao_mecanica")) %>%
  step_novel(all_nominal_predictors()) %>%
  step_zv(all_predictors()) %>%
  step_dummy(all_nominal_predictors()) %>%
  step_impute_mode(all_nominal_predictors()) %>%
  step_impute_mean(all_numeric_predictors())
rf_spec <-
  rand_forest(mtry = tune(),
              trees = 1000,
              min_n = tune()) %>%
  set_mode("classification") %>%
  set_engine("ranger")
rf_grid <- grid_regular(mtry(range = c(1, 10)),</pre>
                        min_n(),
                        levels = 5)
set.seed(234)
df_folds <- vfold_cv(df_train, v = 5, strata = all_of(outcome_column))</pre>
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()
rf_tune %>%
  collect_metrics() %>%
  filter(.metric == "roc_auc") %>%
  select(mean, min_n, mtry) %>%
 pivot_longer(min_n:mtry,
    values_to = "value",
   names_to = "parameter"
  ) %>%
  ggplot(aes(value, mean, color = parameter)) +
  geom_point(show.legend = FALSE) +
  facet_wrap(~parameter, scales = "free_x") +
  labs(x = NULL, y = "AUC")
```



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)

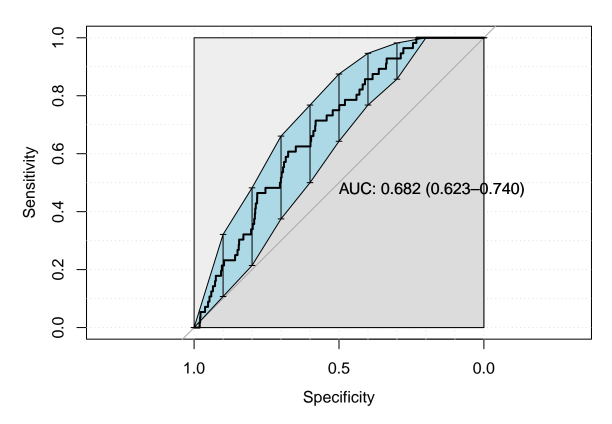
last_rf_fit <-
    final_rf_workflow %>%
    last_fit(df_split)

final_rf_fit <- extract_workflow(last_rf_fit)

rf_auc = validation(final_rf_fit, df_test)

## Setting levels: control = 0, case = 1

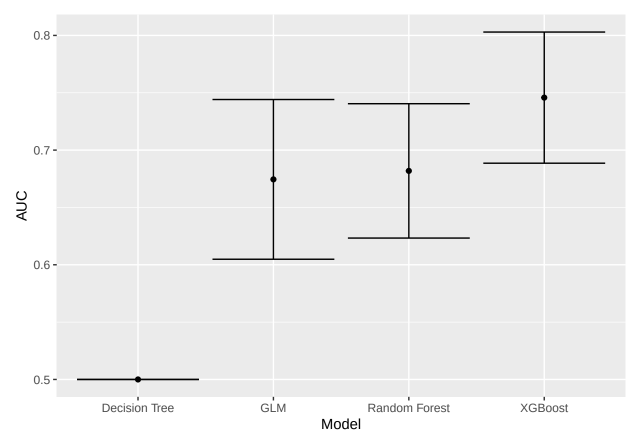
## Setting direction: controls < cases</pre>
```



```
# final_rf_fit %>%
# extract_fit_parsnip() %>%
# vip()
```

Models Comparison

```
df_auc %>%
  ggplot(aes(x = Model, y = AUC, ymin = `Lower Limit`, ymax = `Upper Limit`)) +
  geom_point() +
  geom_errorbar()
```



saveRDS(df_auc, sprintf("../EDA/auxiliar/performance/%s_auc_result.RData", outcome_column))