## PSTAT131 - HW4

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```
library(tidyverse)
## -- Attaching packages ----- tidyverse 1.3.1 --
## v ggplot2 3.3.5 v purrr 0.3.4

## v tibble 3.1.6 v dplyr 1.0.8

## v tidyr 1.2.0 v stringr 1.4.0

## v readr 2.1.1 v forcats 0.5.1
## -- Conflicts ----- tidyverse_conflicts() --
## x dplyr::filter() masks stats::filter()
## x dplyr::lag() masks stats::lag()
library(tidymodels)
## -- Attaching packages ------ tidymodels 0.2.0 --
## v broom 0.7.12 v rsample 0.1.1
## v dials 0.1.0 v tune 0.2.0
## v infer
               1.0.0
                           v workflows 0.2.6
## v modeldata 0.1.1 v workflowsets 0.2.1
## v parsnip 0.2.1 v yardstick 0.0.9
## v recipes 0.2.0
## -- Conflicts ----- tidymodels_conflicts() --
## x scales::discard() masks purrr::discard()
## x dplyr::filter() masks stats::filter()
## x recipes::fixed() masks stringr::fixed()
## x dplyr::lag() masks stats::lag()
## x yardstick::spec() masks readr::spec()
## x recipes::step() masks stats::step()
## * Search for functions across packages at https://www.tidymodels.org/find/
library(discrim)
## Attaching package: 'discrim'
```

```
## The following object is masked from 'package:dials':
##
##
       smoothness
library(poissonreg)
library(corrr)
library(klaR) # for naive bayes
## Loading required package: MASS
## Attaching package: 'MASS'
## The following object is masked from 'package:dplyr':
##
       select
tidymodels_prefer()
Titanic <- read.csv("titanic.csv")</pre>
Titanic$survived <- factor(Titanic$survived,levels = c("Yes","No"))</pre>
Titanic$pclass <- as.character(Titanic$pclass)</pre>
Titanic$pclass <- as.factor(Titanic$pclass)</pre>
\#\#Question 1:
set.seed(2216)
titan_split <- initial_split(Titanic, prop = 0.80,</pre>
                                  strata = survived)
titan_train <- training(titan_split)</pre>
titan_test <- testing(titan_split)</pre>
titan_split
## <Analysis/Assess/Total>
## <712/179/891>
dim(titan_train)
## [1] 712 12
titan_recipe <- recipe(survived ~</pre>
                           pclass +
                           sex +
                           age +
                           sib_sp +
                           parch +
                           fare,
                         data = titan_train) %>%
  step_impute_linear(age) %>%
```

```
step_dummy(all_nominal_predictors()) %>%
  step_interact(~ starts_with("sex"):fare) %>%
  step_interact(~ age:fare)
titan_recipe
## Recipe
##
## Inputs:
##
##
         role #variables
##
      outcome
##
    predictor
##
## Operations:
##
## Linear regression imputation for age
## Dummy variables from all_nominal_predictors()
## Interactions with starts_with("sex"):fare
## Interactions with age:fare
\#\#Question 2:
titan_folds <- vfold_cv(titan_train, v = 10)</pre>
titan_folds
## # 10-fold cross-validation
## # A tibble: 10 x 2
##
      splits
                       id
##
      st>
                       <chr>
  1 <split [640/72] > Fold01
##
   2 <split [640/72] > Fold02
  3 <split [641/71] > Fold03
##
  4 <split [641/71] > Fold04
## 5 <split [641/71] > Fold05
## 6 <split [641/71] > Fold06
## 7 <split [641/71] > Fold07
## 8 <split [641/71] > Fold08
## 9 <split [641/71] > Fold09
## 10 <split [641/71]> Fold10
```

##Question 3: 1) We use k-fold cross-validation so that we can hold out a subset of the training observations from the fitting process, and apply the learned model to those held out observations.

2) We use k-fold cross-validation method so that it results in a less biased model, "because it ensures that every observation from the original dataset has the chance of appearing in training and test set." (cited from: https://towardsdatascience.com/why-and-how-to-cross-validate-a-model-d6424b45261f#:~:text=K-Folds%20Cross%20Validation%3A&text=Because%20it%20ensures%20that%20every,have%20a%20limited%20input%20data

3) If we use the entire training dataset, we would use the training-validation-test approach.

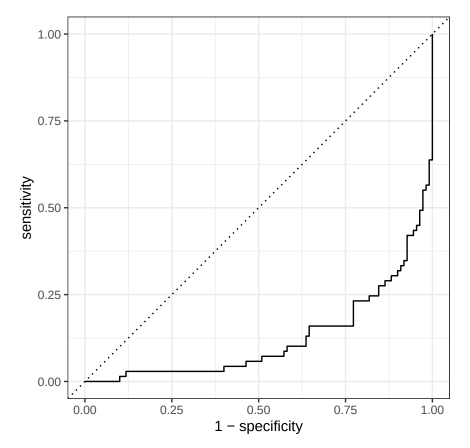
##Question 4:

```
log_reg <- logistic_reg() %>%
  set_engine("glm") %>%
  set_mode("classification")
log_wkflow <- workflow() %>%
  add_model(log_reg) %>%
  add_recipe(titan_recipe)
lda_mod <- discrim_linear() %>%
  set_mode("classification") %>%
  set_engine("MASS")
lda_wkflow <- workflow() %>%
  add_model(lda_mod) %>%
  add_recipe(titan_recipe)
qda_mod <- discrim_quad() %>%
  set_mode("classification") %>%
  set_engine("MASS")
qda_wkflow <- workflow() %>%
  add_model(qda_mod) %>%
  add_recipe(titan_recipe)
Therefore, 30 models in total will be fitted to the data, because there are 10 folds for each of 3 different
models.
\#\#Question 5:
# Fit the logistic regression model
log_fit <-</pre>
  log_wkflow %>%
  fit_resamples(titan_folds)
# Fit the LDA model
lda fit <-
  lda_wkflow %>%
  fit_resamples(titan_folds)
# Fit the QDA model
qda_fit <-
  qda_wkflow %>%
  fit_resamples(titan_folds)
\#\#Question 6:
# See the result of logistic regression model
collect_metrics(log_fit)
## # A tibble: 2 x 6
##
     .metric .estimator mean
                                 n std_err .config
##
    <chr>
             <chr> <dbl> <int>
                                       <dbl> <chr>
## 1 accuracy binary
                       0.810 10 0.0166 Preprocessor1_Model1
## 2 roc_auc binary 0.843
                                  10 0.0212 Preprocessor1_Model1
```

```
# See the result of LDA model
collect_metrics(lda_fit)
## # A tibble: 2 x 6
##
     .metric .estimator mean
                                   n std_err .config
                                       <dbl> <chr>
##
     <chr>>
              <chr>
                         <dbl> <int>
                                  10 0.0213 Preprocessor1_Model1
## 1 accuracy binary
                         0.789
                                  10 0.0212 Preprocessor1_Model1
## 2 roc_auc binary
                         0.845
# See the result of QDA model
collect_metrics(qda_fit)
## # A tibble: 2 x 6
##
     .metric .estimator mean
                                   n std_err .config
     <chr>
             <chr> <dbl> <int>
                                       <dbl> <chr>
                                  10 0.0130 Preprocessor1_Model1
## 1 accuracy binary
                         0.782
## 2 roc_auc binary
                         0.843
                                  10 0.0194 Preprocessor1_Model1
##Question 7: Since the logistic regression model has the highest accuracy, I will use the logistic regression
model as my final model.
final_fit <- fit(log_wkflow, titan_train)</pre>
\#\#Question 8:
predict(final_fit, new_data = titan_test, type = "prob")
## # A tibble: 179 x 2
##
      .pred_Yes .pred_No
##
          <dbl>
                  <dbl>
##
         0.914
                  0.0864
  1
## 2
        0.117
                  0.883
## 3
        0.0873
                  0.913
        0.568
## 4
                  0.432
## 5
        0.264
                  0.736
##
  6
        0.245
                  0.755
        0.0510
                  0.949
##
  7
## 8
        0.169
                  0.831
## 9
         0.0352
                  0.965
         0.775
                  0.225
## 10
## # ... with 169 more rows
augment(final_fit, new_data = titan_test) %>%
 conf_mat(truth = survived, estimate = .pred_class)
##
             Truth
## Prediction Yes No
##
         Yes 50 16
##
          No
               19 94
```

```
multi_metric <- metric_set(accuracy, sensitivity, specificity)
augment(final_fit, new_data = titan_test) %>%
multi_metric(truth = survived, estimate = .pred_class)
```

```
augment(final_fit, new_data = titan_test) %>%
  roc_curve(survived, .pred_No) %>%
  autoplot()
```



Therefore, as the accuracy shown above, the accuracy generated based on the entire training dataset is very close to the average accuracy generated based on the folds.

Thus, overall, the model fits well in predicting the response parameter "survived".