Load the packages

```
rm(list = ls())
setwd('C:/Users/Tao/Desktop/KNN')

pkgs <- c('dplyr', 'ggplot2', 'class', 'caret', 'pROC', 'Hmisc')
# install.packages(pkgs)
suppressMessages(for (i in pkgs) library(i, character.only = TRUE))</pre>
```

Load the data

```
load(file = 'simulated_dataset.R')
str(dataset_final)
'data.frame': 8000 obs. of 21 variables:
$ ID : num 1 2 3 4 5 6 7 8 9 10 ...
$ X_1 : num 56 63 67 50 64 65 59 59 59 58 ...
$ X_3 : num 15.4 14.8 28.5 23.3 17.9 18.9 20.8 14.6 25.6 17.9 ...
       : num 127 122 130 85 109 82 139 97 66 88 ...
$ X_4
$ X_5
        : num 0000000000...
      : num 1010001011...
$ X_6
$ X 7
      : num 1010011000...
$ X_8 : num 0010001000...
$ X_9 : num 1 1 1 1 1 2 1 2 1 1 ...
$ X_10 : num 3.85 3.96 2.31 3.59 4.38 5.5 4.27 4.77 2.66 3.1 ...
$ X_11 : num 4.2 3.72 2.57 3.1 2.61 2.57 2.09 3.48 2.78 2.85 ...
$ X_12
        : num 1000010010 ...
$ X_13
        : num 0000101321...
$ X_14 : num 0000000001...
$ X_15 : num 0000000000...
$ X_16 : num 3 3 4 2 1 3 2 2 3 1 ...
$ Y
       : num 1.51 2.85 1.61 5.21 4.71 ...
$ delta : num 1000011100...
$ Y_binary: num 0 0 1 0 0 0 0 0 0 0 ...
       : chr "raw" "raw" "raw" "raw" ...
```

Step 1: Clean and feature scaling

Clean the data

```
names(dataset_final) <-
    c('ID', 'age', 'male', 'BMI', 'SBP', 'MI', 'HF', 'COPD',
    'cancer', 'albuminuria', 'TC', 'LDLC', 'No_outpatient', 'No_inpatient',
    'liver_disease', 'hypoglycemia', 'CKD_stage',
    'AKI_time', 'AKI_status', 'AKI_binary', 'type')</pre>
head(dataset_final)
```

	ID	age	male	BMI	SBP	MI	HF	COPD	cancer	albuminuria	 LDLC	N
	<dbl></dbl>	 <dbl></dbl>	<(
1	1	56	0	15.4	127	0	1	1	0	1	 4.20	1
2	2	63	0	14.8	122	0	0	0	0	1	 3.72	0
3	3	67	1	28.5	130	0	1	1	1	1	 2.57	0
4	4	50	1	23.3	85	0	0	0	0	1	 3.10	0
5	5	64	0	17.9	109	0	0	0	0	1	 2.61	0
6	6	65	0	18.9	82	0	0	1	0	2	 2.57	1

```
dataset_final <- dataset_final %>%
        mutate(
        albuminuria = case_when( ## relabel of categorical variable
            albuminuria == 1 ~ "normal to mild",
            albuminuria == 2 ~ "moderate",
            albuminuria == 3 ~ "severe"),
        albuminuria = factor(albuminuria, levels = c("normal to mild", "moderate", "severe")),
        CKD_stage = case_when(
            CKD_stage == 1 ~ "G1-2",
            CKD_stage == 2 ~ "G3a",
            CKD_stage == 3 ~ "G3b",
            CKD_stage == 4 ~ "G4"),
         \label{eq:ckd_stage} {\tt CKD\_stage = factor(CKD\_stage, levels = c("G1-2", "G3a", "G3b", "G4")))} 
data <- dataset_final %>% filter(type == "raw")
data <- data[, c('AKI_binary', 'age', 'male', 'BMI', 'SBP', 'MI', 'HF', 'COPD',</pre>
                  'cancer', 'albuminuria', 'TC', 'LDLC', 'No_outpatient', 'No_inpatient',
                 'liver_disease', 'hypoglycemia', 'CKD_stage')]
data$AKI_binary <- as.factor(data$AKI_binary)</pre>
data_external <- dataset_final %>% filter(type == "external")
data_external <- data_external[, c('AKI_binary', 'age', 'male', 'BMI', 'SBP', 'MI', 'HF',</pre>
'COPD',
                           'cancer', 'albuminuria', 'TC', 'LDLC', 'No_outpatient',
'No_inpatient',
                           'liver_disease', 'hypoglycemia', 'CKD_stage')]
data_external$AKI_binary <- as.factor(data_external$AKI_binary)</pre>
```

Feature scaling

```
data <-
   data %>%
    mutate(
        AKI_binary = as.factor(AKI_binary),
        male = as.numeric(male),
        MI = as.numeric(MI),
        HF = as.numeric(HF),
        COPD = as.numeric(COPD),
        cancer = as.numeric(cancer),
        albuminuria = as.numeric(albuminuria) - 1,
        liver_disease = as.numeric(liver_disease),
        hypoglycemia = as.numeric(hypoglycemia),
        CKD_stage = as.numeric(CKD_stage) - 1
    )
data_scaled <- as.data.frame(scale(data[, -1]))</pre>
data_scaled <- cbind(AKI_binary = data$AKI_binary, data_scaled)</pre>
str(data_scaled)
levels(data_scaled$AKI_binary) <- c('no', 'yes')</pre>
```

```
'data.frame': 5000 obs. of 17 variables:
$ age
              : num -1.2 0.208 1.013 -2.408 0.41 ...
            : num -0.983 -0.983 1.017 1.017 -0.983 ...
$ male
$ BMI
             : num -1.472 -1.602 1.366 0.239 -0.931 ...
$ SBP
            : num 0.9398 0.6594 1.108 -1.4151 -0.0695 ...
$ MI
             : num -0.4 -0.4 -0.4 -0.4 -0.4 ...
             : num 1.711 -0.584 1.711 -0.584 -0.584 ...
$ HF
$ COPD : num 1.556 -0.642 1.556 -0.642 -0.642 ...
$ cancer : num -0.518 -0.518 1.931 -0.518 -0.518 ...
$ albuminuria : num -0.639 -0.639 -0.639 -0.639 ...
$ TC : num -0.5067 -0.4083 -1.8839 -0.7392 -0.0327 ...
          : num 2.31 1.484 -0.495 0.417 -0.427 ...
$ LDLC
$ No_outpatient: num 1.299 -0.534 -0.534 -0.534 -0.534 ...
$ No_inpatient : num -0.812 -0.812 -0.812 -0.812 0.379 ...
$ liver_disease: num -0.289 -0.289 -0.289 -0.289 -0.289 ...
$ hypoglycemia : num -0.136 -0.136 -0.136 -0.136 -0.136 ...
$ CKD_stage : num 1.009 1.009 2.267 -0.249 -1.507 ...
```

```
# normalizing
normalized <- function(x){</pre>
   return((x - min(x)) / (max(x) - min(x)))
data_normalized <- as.data.frame(lapply(data[, -1], normalized))</pre>
data normalized <- cbind(AKI binary = data$AKI binary, data normalized)</pre>
str(data normalized)
levels(data_normalized$AKI_binary) <- c('no', 'yes')</pre>
'data.frame': 5000 obs. of 17 variables:
\ \ AKI_binary \ \ : Factor w/ 2 levels "0","1": 1 1 2 1 1 1 1 1 1 1 ...
             : num 0.333 0.545 0.667 0.152 0.576 ...
 $ age
$ male
              : num 0011000000...
$ BMI
              : num 0.0839 0.0629 0.542 0.3601 0.1713 ...
 $ SBP
             : num 0.614 0.575 0.638 0.283 0.472 ...
             : num 00000000000...
 $ MI
 $ HF
             : num 1010001011...
 $ COPD
             : num 1010011000...
$ cancer
              : num 0010001000...
 $ albuminuria : num 0 0 0 0 0 0.5 0 0.5 0 0 ...
$ TC
              : num 0.443 0.456 0.256 0.411 0.507 ...
$ LDLC
              : num 0.636 0.531 0.279 0.395 0.287 ...
```

\$ liver_disease: num 0 0 0 0 0 0 0 0 1 ...
\$ hypoglycemia : num 0 0 0 0 0 0 0 0 ...
\$ CKD_stage : num 0.667 0.667 1 0.333 0 ...

```
data_external <-
    data_external %>%
    mutate(
        AKI_binary = as.factor(AKI_binary),
        male = as.numeric(male),
        MI = as.numeric(MI),
        HF = as.numeric(HF),
        COPD = as.numeric(COPD),
        cancer = as.numeric(cancer),
        albuminuria = as.numeric(albuminuria) - 1,
        liver disease = as.numeric(liver disease),
        hypoglycemia = as.numeric(hypoglycemia),
        CKD_stage = as.numeric(CKD_stage) - 1
    )
# scaling
data_external_scaled <- as.data.frame(scale(data_external[, -1]))</pre>
data_external_scaled <- cbind(AKI_binary = data_external$AKI_binary, data_external_scaled)</pre>
str(data_external_scaled)
levels(data_external_scaled$AKI_binary) <- c('no', 'yes')</pre>
```

```
'data.frame': 3000 obs. of 17 variables:
$ AKI_binary : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 ...
$ age
              : num -0.401 0.855 -2.076 1.413 0.436 ...
$ male
             : num -0.995 1.005 1.005 1.005 1.005 ...
$ BMI
             : num -0.668 -1.025 2.245 1.274 1.987 ...
$ SBP
             : num -2.349 1.119 0.868 0.767 0.315 ...
$ MI
             : num -0.418 -0.418 2.389 -0.418 -0.418 ...
              : num -0.603 1.658 -0.603 -0.603 1.658 ...
$ HF
$ COPD
              : num -0.638 1.567 1.567 -0.638 -0.638 ...
$ cancer
               : num -0.519 -0.519 1.928 -0.519 -0.519 ...
$ albuminuria : num -0.846 0.49 -0.846 1.825 -0.846 ...
$ TC : num -1.68 -0.616 -0.698 0.852 -0.185 ...
$ LDLC : num -0.9545 0.0344 0.5219 -0.2581 0.4244 ...
$ No_outpatient: num 1.194 -0.579 -0.579 -0.579 -0.579 ...
$ No_inpatient : num 1.897 -0.739 -0.739 0.579 -0.739 ...
$ liver_disease: num -0.295 -0.295 -0.295 -0.295 ...
$ hypoglycemia : num -0.171 -0.171 -0.171 -0.171 -0.171 ...
$ CKD_stage : num 1.05 -1.43 1.05 -1.43 -0.19 ...
```

```
# normalizing
normalized <- function(x){
    return((x - min(x)) / (max(x) - min(x)))
}
data_external_normalized <- as.data.frame(lapply(data_external[, -1], normalized))
data_external_normalized <- cbind(AKI_binary = data_external$AKI_binary,
data_external_normalized)
str(data_external_normalized)
levels(data_external_normalized$AKI_binary) <- c('no', 'yes')</pre>
```

```
'data.frame': 3000 obs. of 17 variables:
$ AKI_binary : Factor w/ 2 levels "0","1": 1 1 1 1 1 1 1 1 1 1 ...
$ age
             : num 0.48 0.66 0.24 0.74 0.6 0.58 0.64 0.56 0.54 0.5 ...
$ male
            : num 0111110000 ...
$ BMI
            : num 0.237 0.168 0.798 0.611 0.748 ...
$ SBP
            : num 0.146 0.625 0.59 0.576 0.514 ...
             : num 0010000000...
$ MI
$ HF
             : num 0100100001...
$ COPD
             : num 0110000110...
$ cancer : num 0 0 1 0 0 1 0 0 1 0 ...
$ albuminuria : num 0 0.5 0 1 0 0 1 0.5 0 0 ...
$ TC : num 0.272 0.427 0.415 0.64 0.489 ...
          : num 0.203 0.348 0.419 0.305 0.404 ...
$ LDLC
$ No_outpatient: num 0.333 0 0 0 0 ...
$ No_inpatient : num   0.4 0 0 0.2 0 0.2 0 0 0 0.2 ...
$ liver_disease: num 0 0 0 0 0 1 0 1 0 0 ...
\ hypoglycemia : num \ 0\ 0\ 0\ 0\ 0\ 0\ 0\ 0\ 0\ \dots
$ CKD_stage : num 0.667 0 0.667 0 0.333 ...
```

Randome split the data into train and test

```
# set.seed(123)
# id <- sample(seq_len(nrow(data_normalized)), size = 0.7 * nrow(data_normalized))
# train <- data_normalized[id, ]
# test <- data_normalized[-id, ]

set.seed(123)
id <- sample(seq_len(nrow(data_normalized)), size = 0.7 * nrow(data_normalized))
train <- data_normalized[id, ]
test <- data_normalized[-id, ]</pre>
```

```
KNN_pred <-
knn(
    train = train[, -1],
    test = test[, -1],
    cl = train$AKI_binary,
    k = ceiling(sqrt(nrow(train))),
    prob = TRUE
)

pred_prob <- attr(KNN_pred, 'prob')
pred_prob <- ifelse(KNN_pred == 'yes', pred_prob, 1-pred_prob)

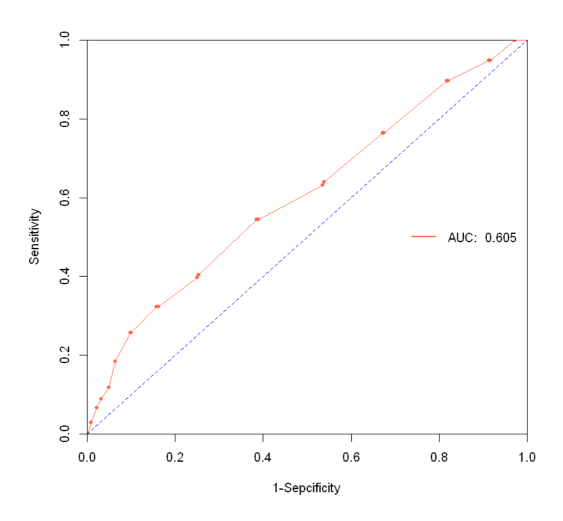
# confusion matrix
cm <- table(actual = test$AKI_binary, predicted = KNN_pred)
cm

predicted
actual no yes
    no 1364    0
    yes 136    0</pre>
```

```
# AUC
roc <- roc(as.factor(as.numeric(test$AKI_binary) - 1), pred_prob)
AUC <- auc(roc)
round(AUC, 3)

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

```
plot(1 - roc$specificities,
    roc$sensitivities,
    pch = 20,
    col = "tomato",
    xlim = c(0, 1), ylim = c(0, 1),
    xlab = '1-Sepcificity',
    ylab = 'Sensitivity',
    xaxs = 'i',
    yaxs = 'i')
lines(1- roc$specificities,
     roc$sensitivities,
     col = "tomato")
abline(0, 1, lty = 2, col = 'blue')
legend("right",
      legend = paste("AUC: ", round(AUC, 3)),
      col = "tomato",
      lwd = 2,
      bty = "n")
```



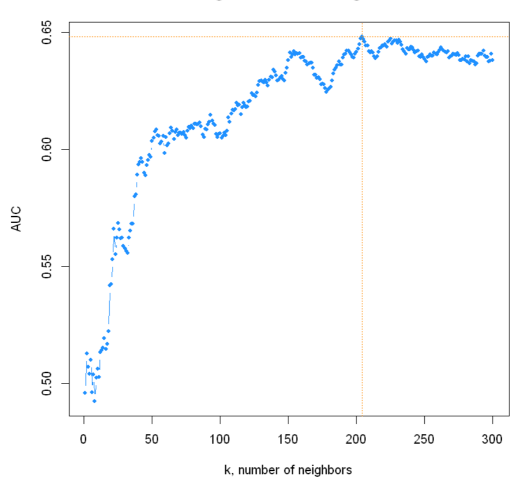
Step 3: Model tuning

Random split

```
# using the for loop
results <- data.frame()</pre>
suppressMessages(for(k in 1:300) {
    KNN_fit <-
        knn(
            train = train[, -1],
            test = test[, -1],
            cl = train$AKI_binary,
            k = k,
            prob = TRUE
    pred_prob <- attr(KNN_fit, 'prob')</pre>
    pred_prob <- ifelse(KNN_fit == 'yes', pred_prob, 1-pred_prob)</pre>
    roc <- roc(as.factor(as.numeric(test$AKI_binary) - 1), pred_prob)</pre>
    AUC <- auc(roc)
    result <- data.frame(k = k, AUC = AUC)</pre>
    results <- rbind(results, result)</pre>
})
```

```
plot(results,
    type = "b",
    col = "dodgerblue",
    cex = 1,
    pch = 20,
    xlab = "k, number of neighbors",
    ylab = "AUC",
    main = "Tuning for k: AUC vs Neighbors")
# add line for max AUC seen
abline(h = max(results$AUC), col = "darkorange", lty = 3)
abline(v = results$k[results$AUC == max(results$AUC)], col = "darkorange", lty = 3)
```

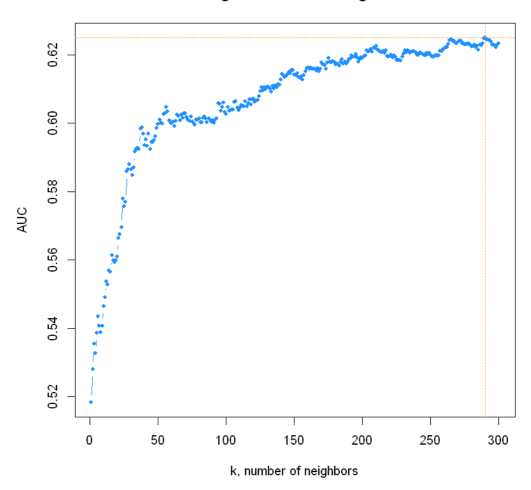
Tuning for k: AUC vs Neighbors



Cross-validation

```
type = "b",
  col = "dodgerblue",
  cex = 1,
  pch = 20,
  xlab = "k, number of neighbors",
  ylab = "AUC",
  main = "Tuning for k: AUC vs Neighbors")
# add line for max AUC seen
abline(h = max(KNN_tune$results$ROC), col = "darkorange", lty = 3)
abline(v = KNN_tune$results$k[KNN_tune$results$ROC == max(KNN_tune$results[['ROC']])], col =
"darkorange", lty = 3)
```

Tuning for k: AUC vs Neighbors



Step 4: Model refit

Random split

```
best_model_k_split <- results$k[results$AUC == max(results$AUC)]
best_model_k_split</pre>
```

204

```
best_model_split <-
    knn(
        train = data_normalized[, -1],
        test = data_normalized[, -1],</pre>
```

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

Setting direction: controls < cases
```

0.654201324309117

Cross-validation

```
best_model_k_cv <- KNN_tune$bestTune$k
best_model_k_cv</pre>
```

290

```
best_model_cv <-
   knn3(
        AKI_binary ~ .,
        data = data_normalized,
        k = best_model_k_cv
)

predictions_cv <- predict(best_model_cv, data_normalized, type = "class")
pred_prob_cv <- predict(best_model_cv, data_normalized, type = "prob")
pred_prob_cv <- pred_prob_cv[,2]</pre>
```

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

Setting direction: controls < cases
```

Random split

```
best_model_external <-
    knn(
        train = data_normalized[, -1],
        test = data_external_normalized[, -1],
        cl = data_normalized$AKI_binary,
        k = best_model_k_split,
        prob = TRUE
    )
pred_prob <- attr(best_model_external, 'prob')
pred_prob <- ifelse(best_model_external == 'yes', pred_prob, 1-pred_prob)

# confusion matrix
cm <- table(actual = data_external_normalized$AKI_binary, predicted = best_model_external)
cm

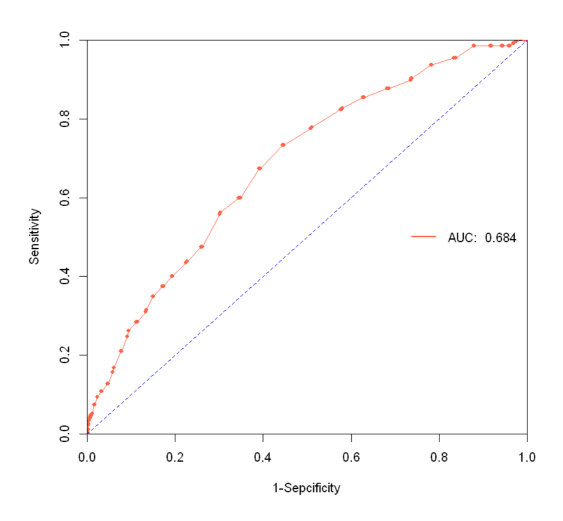
    predicted
actual no yes
    no 2733    0
    yes 267    0</pre>
```

```
# AUC
roc <- roc(data_external$AKI_binary, pred_prob)
AUC <- auc(roc)
AUC

Setting levels: control = 0, case = 1

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

```
plot(1 - roc$specificities,
    roc$sensitivities,
    pch = 20,
    col = "tomato",
    xlim = c(0, 1), ylim = c(0, 1),
    xlab = '1-Sepcificity',
    ylab = 'Sensitivity',
    xaxs = 'i',
    yaxs = 'i')
lines(1- roc$specificities,
     roc$sensitivities,
     col = "tomato")
abline(0, 1, lty = 2, col = 'blue')
legend("right",
      legend = paste("AUC: ", round(AUC, 3)),
      col = "tomato",
      lwd = 2,
      bty = "n")
```

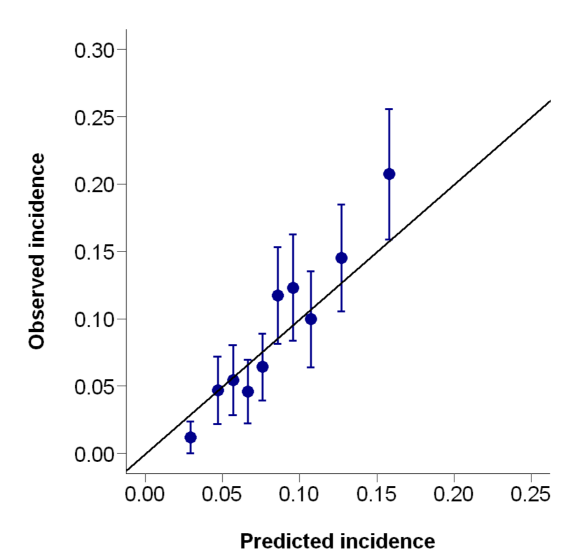


```
png(filename = 'roc_split.png', width = 1600, height = 1600, unit = 'px', res = 300)
plot(1 - roc$specificities,
    roc$sensitivities,
    pch = 20,
    col = "tomato",
    xlim = c(0, 1), ylim = c(0, 1),
    xlab = '1-Sepcificity',
    ylab = 'Sensitivity',
    xaxs = 'i',
    yaxs = 'i')
lines(1- roc$specificities,
      roc$sensitivities,
      col = "tomato")
abline(0, 1, lty = 2, col = 'blue')
legend("right",
      legend = paste("AUC: ", round(AUC, 3)),
       col = "tomato",
      1wd = 2,
      bty = "n")
dev.off()
```

png: 2

```
# calibration curve
n_group <- 10
cal_data <- data.frame(AKI_binary = as.numeric(data_external$AKI_binary) - 1, pred_prob)
cal_data <- cal_data[order(cal_data$pred_prob),]
cal_data <- cal_data %>% mutate(tiles = cut2(pred_prob, g = n_group))
```

```
cal_data <- cal_data %>%
            group_by(tiles) %>%
            summarise(
                n = n()
                obs_mean = mean(AKI_binary),
                obs_sd = sd(AKI_binary),
                # obs_mean = sum(AKI_binary) / n(),
                # obs_sd = sqrt(obs_mean * (1 - obs_mean)),
                pred_mean = mean(pred_prob),
                obs_upper = obs_mean + 1.96*obs_sd/sqrt(n()),
                obs_lower = obs_mean - 1.96*obs_sd/sqrt(n())
cal_plot <-</pre>
    ggplot(
        cal_data,
        aes(x = pred_mean, y = obs_mean)) +
    # geom_line() +
    geom_point(size = 5, colour = 'darkblue') +
    geom errorbar(
        aes(ymin = obs_lower, ymax = obs_upper),
        linewidth = 1,
        width = .005,
       colour = 'darkblue'
    geom_abline(linewidth = 1) +
    xlab("\nPredicted incidence") +
    ylab("Observed incidence\n") +
    scale_x_continuous(limits = c(0, 0.25), breaks = seq(0, 0.5, 0.05)) +
    scale_y_continuous(limits = c(0, 0.3), breaks = seq(0, 0.5, 0.05)) +
    theme(
        panel.background = element_blank(),
        panel.grid = element_blank(),
        # panel.grid.major.y = element_line(color = "gray"),
        plot.title = element_text(size = 20, face = "bold", hjust = 0),
        axis.title = element_text(size = 20, face = "bold"),
        axis.text = element_text(size = 20, color = 'black'),
        axis.ticks.length.x = unit(0.3, 'cm'),
        axis.ticks.length.y = unit(0.3, 'cm'),
        axis.line = element_line(colour = "black", linewidth = 0.5)
cal_plot
```



```
png(filename = 'calibration_curve_split.png', width = 3200, height = 3200, unit = 'px', res =
300)
cal_plot
dev.off()
```

png: 2

Cross-validation

```
best_model_external <-
    knn(
        train = data_normalized[, -1],
        test = data_external_normalized[, -1],
        cl = data_normalized$AKI_binary,
        k = best_model_k_cv,
        prob = TRUE
    )
pred_prob <- attr(best_model_external, 'prob')
pred_prob <- ifelse(best_model_external == 'yes', pred_prob, 1-pred_prob)</pre>
```

```
# confusion matrix
cm <- table(actual = data_external$AKI_binary, predicted = best_model_external)
cm</pre>
```

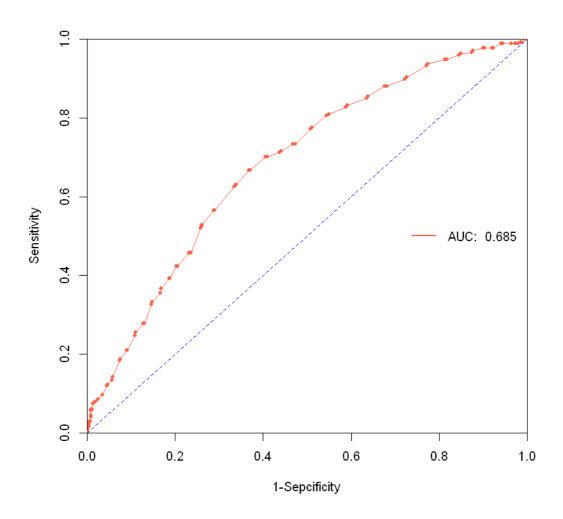
```
predicted
actual no yes
0 2733 0
1 267 0
```

```
# AUC
roc <- roc(data_external$AKI_binary, pred_prob)
AUC <- auc(roc)
AUC

Setting levels: control = 0, case = 1

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

```
plot(1 - roc$specificities,
   roc$sensitivities,
   pch = 20,
   col = "tomato",
   xlim = c(0, 1), ylim = c(0, 1),
    xlab = '1-Sepcificity',
    ylab = 'Sensitivity',
    xaxs = 'i',
    yaxs = 'i')
lines(1- roc$specificities,
     roc$sensitivities,
     col = "tomato")
abline(0, 1, lty = 2, col = 'blue')
legend("right",
      legend = paste("AUC: ", round(AUC, 3)),
      col = "tomato",
      lwd = 2,
     bty = "n")
```

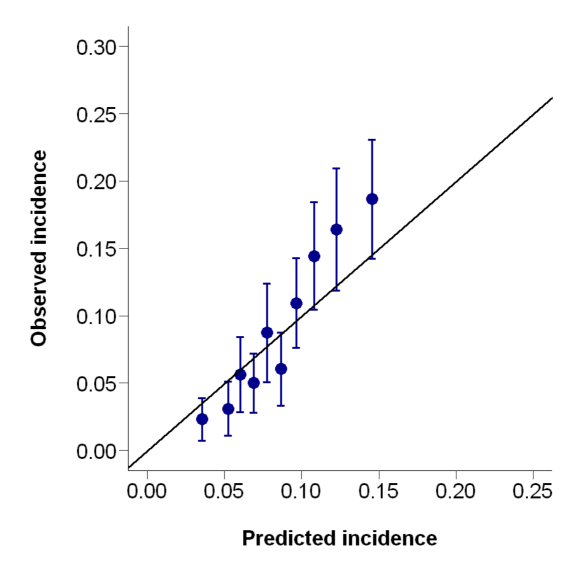


```
png(filename = 'roc_cv.png', width = 1600, height = 1600, unit = 'px', res = 300)
plot(1 - roc$specificities,
    roc$sensitivities,
    pch = 20,
    col = "tomato",
    xlim = c(0, 1), ylim = c(0, 1),
    xlab = '1-Sepcificity',
    ylab = 'Sensitivity',
    xaxs = 'i',
    yaxs = 'i')
lines(1- roc$specificities,
     roc$sensitivities,
      col = "tomato")
abline(0, 1, lty = 2, col = 'blue')
legend("right",
      legend = paste("AUC: ", round(AUC, 3)),
       col = "tomato",
      1wd = 2,
      bty = "n")
dev.off()
```

png: 2

```
# calibration curve
n_group <- 10
cal_data <- data.frame(AKI_binary = as.numeric(data_external$AKI_binary) - 1, pred_prob)
cal_data <- cal_data[order(cal_data$pred_prob),]
cal_data <- cal_data %>% mutate(tiles = cut2(pred_prob, g = n_group))
```

```
cal_data <- cal_data %>%
            group_by(tiles) %>%
            summarise(
                n = n()
                obs_mean = mean(AKI_binary),
                obs_sd = sd(AKI_binary),
                # obs_mean = sum(AKI_binary) / n(),
                # obs_sd = sqrt(obs_mean * (1 - obs_mean)),
                pred_mean = mean(pred_prob),
                obs_upper = obs_mean + 1.96*obs_sd/sqrt(n()),
                obs_lower = obs_mean - 1.96*obs_sd/sqrt(n())
cal_plot <-</pre>
    ggplot(
        cal_data,
        aes(x = pred_mean, y = obs_mean)) +
    # geom_line() +
    geom_point(size = 5, colour = 'darkblue') +
    geom errorbar(
        aes(ymin = obs_lower, ymax = obs_upper),
        linewidth = 1,
        width = .005,
       colour = 'darkblue'
    geom_abline(linewidth = 1) +
    xlab("\nPredicted incidence") +
    ylab("Observed incidence\n") +
    scale_x_continuous(limits = c(0, 0.25), breaks = seq(0, 0.5, 0.05)) +
    scale_y_continuous(limits = c(0, 0.3), breaks = seq(0, 0.5, 0.05)) +
    theme(
        panel.background = element_blank(),
        panel.grid = element_blank(),
        # panel.grid.major.y = element_line(color = "gray"),
        plot.title = element_text(size = 20, face = "bold", hjust = 0),
        axis.title = element_text(size = 20, face = "bold"),
        axis.text = element_text(size = 20, color = 'black'),
        axis.ticks.length.x = unit(0.3, 'cm'),
        axis.ticks.length.y = unit(0.3, 'cm'),
        axis.line = element_line(colour = "black", linewidth = 0.5)
cal_plot
```



```
png(filename = 'calibration_curve_cv.png', width = 3200, height = 3200, unit = 'px', res = 300)
cal_plot
dev.off()
```

png: 2