## P8106\_group2\_COVID-19\_recoveryanalysis

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## Import and data manipulation

```
# Load recovery.RData environment
load("./recovery.Rdata")
dat %>% na.omit()
# dat1 draw a random sample of 2000 participants Uni:3307
set.seed(3307)
dat1 = dat[sample(1:10000, 2000),]
dat1 =
 dat1[, -1] %>%
  mutate(
   gender = as.factor(gender),
   race = as.factor(race),
   smoking = as.factor(smoking),
   hypertension = as.factor(hypertension),
   diabetes = as.factor(diabetes),
   vaccine = as.factor(vaccine),
   severity = as.factor(severity),
    study = as.factor(
      case_when(study == "A" ~ 1, study == "B" ~ 2, study == "C" ~ 3)
    )
# dat2 draw a random sample of 2000 participants Uni:2493
set.seed(2493)
dat2 = dat[sample(1:10000, 2000),]
dat2 =
  dat2[, -1] %>%
 mutate(
   gender = as.factor(gender),
```

```
race = as.factor(race),
    smoking = as.factor(smoking),
    hypertension = as.factor(hypertension),
    diabetes = as.factor(diabetes),
    vaccine = as.factor(vaccine),
    severity = as.factor(severity),
    study = as.factor(
      case when(study == ^{"}A" \sim 1, study == ^{"}B" \sim 2, study == ^{"}C" \sim 3)
    )
# Merged dataset with unique observation
covid_dat = rbind(dat1, dat2) %>%
  unique()
covid_dat2 = model.matrix(recovery_time ~ ., covid_dat)[, -1]
# Partition dataset into two parts: training data (70%) and test data (30%)
rowTrain = createDataPartition(y = covid_dat$recovery_time, p = 0.7, list = FALSE)
trainData = covid_dat[rowTrain, ]
testData = covid_dat[-rowTrain, ]
# matrix of predictors
x1 = covid_dat2[rowTrain,]
# vector of response
y1 = covid_dat$recovery_time[rowTrain]
# matrix of predictors
x2 = covid_dat2[-rowTrain,]
# vector of response
y2 = covid_dat$recovery_time[-rowTrain]
ctrl1 = trainControl(method = "repeatedcv", number = 10, repeats = 5)
```

## Data visualization