Addition of Emphysema-CNN score improves upon prescreening risk calculated by LCRAT

Load packages

```
# List packages to be loaded (and installed if needed)
packages <-
    c(
        "readr",
        "here",
        "dplyr",
        "psych",
        "pROC"
    )
# List packages that are not installed
not_installed <-</pre>
    packages[!(packages %in% installed.packages()[, "Package"])]
# Install packages that are not installed
if (length(not_installed))
    install.packages(not_installed)
# Load all packages
lapply(packages, require, character.only = TRUE)
```

Merge in Wes' T0 data

```
nlst_emp <- read_csv(here('data/T0_data.csv'))
data_screen_abn_neg <- readRDS(here('data/data_screen_abn_neg.rds'))
data_screen_abn_neg_emp <-
    merge(data_screen_abn_neg, nlst_emp, by = "pid")</pre>
```

Fit models

Without physician-annotated features

```
# Prescreening risk only
glm_screen_neg <-
    glm(case ~ log1yrisk - 1,
    data = data_screen_abn_neg_emp,
    family = binomial(link = 'log'))
summary(glm_screen_neg)</pre>
```

```
## Call:
## glm(formula = case ~ log1yrisk - 1, family = binomial(link = "log"),
      data = data_screen_abn_neg_emp)
##
## Deviance Residuals:
           1Q Median
                                 3Q
##
      Min
                                          Max
## -0.7549 -0.4153 -0.3584 -0.3115
##
## Coefficients:
##
            Estimate Std. Error z value Pr(>|z|)
## log1yrisk 0.49427
                       0.01639
                                30.16 <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance:
                        Inf on 1750 degrees of freedom
## Residual deviance: 879.18 on 1749 degrees of freedom
## AIC: 881.18
## Number of Fisher Scoring iterations: 6
# Prescreening risk + p_emph
glm_screen_neg_pemph <-</pre>
   glm(case ~ log1yrisk + p_emph - 1,
       data = data_screen_abn_neg_emp,
       family = binomial(link = 'log'),
       na.action = na.exclude)
summary(glm_screen_neg_pemph)
##
## Call:
## glm(formula = case ~ log1yrisk + p_emph - 1, family = binomial(link = "log"),
      data = data_screen_abn_neg_emp, na.action = na.exclude)
##
## Deviance Residuals:
      Min
           1Q
                    Median
                                  3Q
## -0.8022 -0.4161 -0.3543 -0.3010
                                      2.6001
##
## Coefficients:
            Estimate Std. Error z value Pr(>|z|)
                        0.02082 24.790 <2e-16 ***
## log1yrisk 0.51614
## p_emph
             0.62740
                        0.32944
                                 1.904
                                         0.0569 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance:
                        Inf on 1750 degrees of freedom
## Residual deviance: 875.93 on 1748 degrees of freedom
## AIC: 879.93
##
## Number of Fisher Scoring iterations: 6
```

```
# Prescreening risk + logit p_emph
glm_screen_neg_logitpemph <-</pre>
    glm(case ~ log1yrisk + I(logit(p_emph)) - 1,
        data = data_screen_abn_neg_emp,
       family = binomial(link = 'log'),
       na.action = na.exclude)
summary(glm_screen_neg_logitpemph)
##
## Call:
## glm(formula = case ~ log1yrisk + I(logit(p_emph)) - 1, family = binomial(link = "log"),
       data = data_screen_abn_neg_emp, na.action = na.exclude)
## Deviance Residuals:
      Min
           10
                    Median
                                   30
                                           Max
## -0.7680 -0.4150 -0.3548 -0.3048
                                        2.5849
##
## Coefficients:
                    Estimate Std. Error z value Pr(>|z|)
                                0.02319 19.652
## log1yrisk
                     0.45573
                                                  <2e-16 ***
## I(logit(p_emph)) 0.10048
                                0.04679 2.148
                                                  0.0317 *
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance:
                         Inf on 1750 degrees of freedom
## Residual deviance: 874.93 on 1748 degrees of freedom
## AIC: 878.93
## Number of Fisher Scoring iterations: 6
# Prescreening risk + log p_emph
glm_screen_neg_logpemph <-</pre>
   glm(case ~ log1yrisk + I(log(p_emph)) - 1,
        data = data_screen_abn_neg_emp,
        family = binomial(link = 'log'),
       na.action = na.exclude)
summary(glm_screen_neg_logpemph)
##
## glm(formula = case ~ log1yrisk + I(log(p_emph)) - 1, family = binomial(link = "log"),
##
       data = data_screen_abn_neg_emp, na.action = na.exclude)
##
## Deviance Residuals:
                    Median
##
      Min
                1Q
                                   3Q
                                           Max
## -0.7482 -0.4162 -0.3571 -0.2987
##
## Coefficients:
##
                  Estimate Std. Error z value Pr(>|z|)
                  0.41882 0.03509 11.936
## log1yrisk
## I(log(p_emph)) 0.17159
                              0.07477
                                        2.295
                                                0.0217 *
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
## Null deviance: Inf on 1750 degrees of freedom
## Residual deviance: 873.91 on 1748 degrees of freedom
## AIC: 877.91
##
## Number of Fisher Scoring iterations: 6
```

With physician-annotated features

```
glm_screen_neg_cons <-
    glm(
        case ~ log1yrisk + log1yrisk:consolidation - 1,
        data = data_screen_abn_neg_emp,
        family = binomial(link = 'log'),
        na.action = na.exclude
    )
summary(glm_screen_neg_cons)</pre>
```

```
##
## glm(formula = case ~ log1yrisk + log1yrisk:consolidation - 1,
##
      family = binomial(link = "log"), data = data_screen_abn_neg_emp,
##
      na.action = na.exclude)
##
## Deviance Residuals:
      Min
                1Q
                    Median
                                  3Q
                                          Max
## -0.7233 -0.4049 -0.3462 -0.3003
                                       2.6045
##
## Coefficients:
                          Estimate Std. Error z value Pr(>|z|)
##
## log1yrisk
                           0.50851
                                      0.01842 27.602
                                                        <2e-16 ***
## log1yrisk:consolidation -0.10453
                                      0.09195 -1.137
                                                         0.256
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
                       Inf on 1536 degrees of freedom
##
      Null deviance:
## Residual deviance: 740.4 on 1534 degrees of freedom
     (214 observations deleted due to missingness)
## AIC: 744.4
##
## Number of Fisher Scoring iterations: 6
```

```
glm_screen_neg_abn_emph <-
    glm(
        case ~ log1yrisk + log1yrisk:emphysema - 1,
        data = data_screen_abn_neg_emp,</pre>
```

```
family = binomial(link = 'log'),
       na.action = na.exclude
   )
summary(glm_screen_neg_abn_emph)
##
## Call:
## glm(formula = case ~ log1yrisk + log1yrisk:emphysema - 1, family = binomial(link = "log"),
       data = data_screen_abn_neg_emp, na.action = na.exclude)
##
## Deviance Residuals:
      Min
                1Q
                     Median
                                   3Q
                                           Max
## -0.7806 -0.4157 -0.3479 -0.2724
                                        2.6343
##
## Coefficients:
##
                      Estimate Std. Error z value Pr(>|z|)
                       0.55480
                                  0.02842 19.521
## log1yrisk
                                                    <2e-16 ***
## log1yrisk:emphysema -0.09376
                                  0.03682 -2.547
                                                     0.0109 *
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance:
                        Inf on 1536 degrees of freedom
## Residual deviance: 734.85 on 1534 degrees of freedom
     (214 observations deleted due to missingness)
## AIC: 738.85
## Number of Fisher Scoring iterations: 6
glm_screen_neg_abn_cons_emph <-</pre>
   glm(
        case ~ log1yrisk + log1yrisk:consolidation + log1yrisk:emphysema - 1,
        data = data_screen_abn_neg_emp,
       family = binomial(link = 'log'),
       na.action = na.exclude
summary(glm_screen_neg_abn_cons_emph)
##
## Call:
## glm(formula = case ~ log1yrisk + log1yrisk:consolidation + log1yrisk:emphysema -
       1, family = binomial(link = "log"), data = data_screen_abn_neg_emp,
##
      na.action = na.exclude)
##
## Deviance Residuals:
                1Q
                     Median
                                   3Q
                                           Max
## -0.8406 -0.4164 -0.3482 -0.2679
                                        2.6487
## Coefficients:
                           Estimate Std. Error z value Pr(>|z|)
##
                            0.56087
                                      0.02897 19.360 < 2e-16 ***
## log1yrisk
```

```
## log1yrisk:consolidation -0.12984
                                      0.08974 -1.447 0.14792
                                      0.03691 -2.649 0.00808 **
## log1yrisk:emphysema
                          -0.09777
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance:
                        Inf on 1536 degrees of freedom
## Residual deviance: 733.27 on 1533 degrees of freedom
     (214 observations deleted due to missingness)
## AIC: 739.27
## Number of Fisher Scoring iterations: 7
glm_screen_neg_cons_emph_pemph <-</pre>
    glm(
        case ~ log1yrisk + log1yrisk:consolidation + log1yrisk:emphysema + I(log(p_emph)) - 1,
        data = data_screen_abn_neg_emp,
        family = binomial(link = 'log'),
       na.action = na.exclude
    )
summary(glm_screen_neg_cons_emph_pemph)
##
## Call:
  glm(formula = case ~ log1yrisk + log1yrisk:consolidation + log1yrisk:emphysema +
       I(log(p_emph)) - 1, family = binomial(link = "log"), data = data_screen_abn_neg_emp,
##
##
       na.action = na.exclude)
##
## Deviance Residuals:
      Min
##
              1Q
                     Median
                                  3Q
                                          Max
## -0.7769 -0.4153 -0.3389 -0.2597
                                       2.6374
##
## Coefficients:
##
                          Estimate Std. Error z value Pr(>|z|)
## log1yrisk
                           0.45693
                                      0.04953 9.225 <2e-16 ***
## I(log(p_emph))
                           0.21134
                                      0.08587
                                                2.461
                                                        0.0139 *
## log1yrisk:consolidation -0.14210
                                      0.08549 -1.662
                                                        0.0965 .
## log1yrisk:emphysema
                          -0.07085
                                      0.03846 -1.842
                                                        0.0655 .
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance:
                        Inf on 1536 degrees of freedom
## Residual deviance: 727.13 on 1532 degrees of freedom
     (214 observations deleted due to missingness)
## AIC: 735.13
## Number of Fisher Scoring iterations: 7
```