## Homework 0 Answer Sketch

## Thomas E. Love

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```
hw0 <- read_csv(here("data", "hw0.csv")) %>%
    type.convert() # converts all characters to factors
```

## 1 Looking Over the Data Set

I will start with a quick summary to be sure things are imported properly in the hw0 data set...

## summary(hw0)

subject	treatment	cov1	cov2
Min. :101.0	Not Treated:95	Min. :27.03	Min. :29.49
1st Qu.:134.5	Treated :40	1st Qu.:41.74	1st Qu.:46.48
Median :228.0		Median :47.70	Median :53.48
Mean :210.2		Mean :48.36	Mean :52.68
3rd Qu.:261.5		3rd Qu.:56.62	3rd Qu.:59.99
Max. :295.0		Max. :73.28	Max. :73.25
cov3	cov4	cov5	
Min. : 8.00	Min. : 9.00		
MIII 0.00	MIII. : 9.00	Min. :0.0000	
1st Qu.:17.00	1st Qu.:17.00	Min. :0.0000 1st Qu.:0.0000	
1st Qu.:17.00	1st Qu.:17.00	1st Qu.:0.0000	
1st Qu.:17.00 Median :20.00	1st Qu.:17.00 Median :20.00	1st Qu.:0.0000 Median :0.0000	
1st Qu.:17.00 Median :20.00 Mean :20.28	1st Qu.:17.00 Median :20.00 Mean :20.06	1st Qu.:0.0000 Median :0.0000 Mean :0.4296	

#### 1.1 Using the describe function

Alternatively, we could use the describe function, which is part of the Hmisc package...

Hmisc::describe(hw0)

hwO

7	Variables	135	Observations

subject missing distinct Info . 05 Mean Gmd . 10 107.7 0 135 1 210.2 70.54 114.4 135 .75 .25 .50 .90 .95 134.5 228.0 261.5 281.6 288.3

lowest: 101 102 103 104 105, highest: 291 292 293 294 295

treatment n missing distinct 135 0 Value Not Treated Treated Frequency Proportion 0.704 0.296 cov1 Info . 05 n missing distinct Mean Gmd . 10 48.36 11.83 30.70 34.36 0 133 1 48.36 .50 .75 .90 .95 135 . 25 41.75 47.70 56.62 61.84 64.94 lowest: 27.03 28.30 28.39 28.73 29.60, highest: 66.37 67.30 67.89 68.39 73.28 n missing distinct Info Mean Gmd . 05 . 10 1 52.68 11.08 36.81 39.15 0 133 .75 .90 . 25 .50 . 95 53.48 59.99 65.39 67.05 46.48 lowest: 29.49 31.93 32.24 34.24 34.25, highest: 68.90 69.55 70.10 72.03 73.25 \_\_\_\_\_\_ cov3 Mean 20.28 .10 n missing distinct Info Gmd .05 135 0 24 0.995 5.87 12 14 .90 . 95 .50 .75 . 25 20 24 27 17 29 lowest: 8 9 11 12 13, highest: 28 29 30 32 33 \_\_\_\_\_\_ cov4 Mean Gmd .05 . 10 n missing distinct Info 135 23 0.994 20.06 4.929 13 15 0 .75 .90 .95 . 25 .50 17 20 23 26 28 lowest: 9 11 12 13 14, highest: 28 29 31 32 33 \_\_\_\_\_\_

\_\_\_\_\_

 $\operatorname{\mathtt{Sum}}$ 

58

Mean

0.4296

Gmd

0.4938

Info

0.735

2

cov5

135

n missing distinct

0

#### 1.2 Glimpsing the data's structure

Or, perhaps we just want to see the structure of the data and some of the first few values in each variable, in which case, the **str** command would help, or we could use the **dplyr** package's **glimpse** function...

```
glimpse(hw0)
```

The two treatment options are named "Treated" and "Not Treated", as opposed to "Treated" and "Untreated". Anything so that the thing I wanted to evaluate probabilities for (i.e. Treated as compared to Not Treated) came second alphabetically is appealing, because R, by default, treats the first level in a binary categorical variable as unsuccessful and the second level as successful and generally orders levels of binary variables alphabetically.

# 2 Fitting a Logistic Regression Model using the glm function

We are fitting a model to predict the probability of "Treated" here. If we want to see what's in m1, we can type it in, and see what we get, or ask for a summary, and get some additional details.

```
m1
```

#### Coefficients:

```
(Intercept) cov1 cov2 cov3 cov4 cov5 0.23905 0.04159 -0.02512 -0.18594 0.06993 0.79492
```

```
Null Deviance:
                    164.1
Residual Deviance: 135 AIC: 147
summary(m1)
Call:
glm(formula = treatment == "Treated" ~ cov1 + cov2 + cov3 + cov4 +
    cov5, family = binomial(), data = hw0)
Deviance Residuals:
   Min
              10
                   Median
                                3Q
                                        Max
-1.9216 -0.7418 -0.4914
                            0.8746
                                     2.4184
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
                                  0.119
(Intercept)
            0.23905
                        2.00561
                                          0.9051
                                  1.892
                                          0.0585 .
cov1
             0.04159
                        0.02198
cov2
            -0.02512
                        0.02303 -1.091
                                          0.2754
cov3
            -0.18594
                        0.04735 -3.927 8.6e-05 ***
             0.06993
                        0.05053
                                  1.384
                                          0.1664
cov4
             0.79492
                        0.44205
                                  1.798
                                          0.0721 .
cov5
Signif. codes:
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 164.08
                           on 134
                                   degrees of freedom
Residual deviance: 135.02
                           on 129
                                   degrees of freedom
AIC: 147.02
Number of Fisher Scoring iterations: 4
```

Degrees of Freedom: 134 Total (i.e. Null); 129 Residual

## 2.1 Storing Probabilities and Linear Predictions

To store the linear predictions (i.e. log odds of the estimated probabilities,) and the estimated probabilities themselves as part of the hw0 data file, I'll use the following commands:

```
hw0$linpred <- m1$linear.predictors
hw0$prob <- m1$fitted.values
```

The remaining tasks in the assignment essentially require you to obtain some numerical (perhaps) and graphical (mandatory) summaries of the estimated probabilities broken down

# 3 Some Numerical Summaries of the Fitted Probabilities by Treatment Group

#### 3.1 Using dplyr and summarise

The dplyr library can be used to compare the probs across the two treatment groups, along with some piping commands, to create a little data frame of the summaries you're interested in, as follows:

#### 3.2 Using the by command

Some of you may be more familiar with the by command - that works, as well...

```
by(hw0$prob, hw0$treatment, describe)
hw0$treatment: Not Treated
dd[x,]
          missing distinct
                                Info
                                         Mean
                                                    Gmd
                                                              .05
                                                                       .10
      95
                0
                         95
                                   1
                                       0.2343
                                                 0.1891 0.04319 0.05634
     .25
               .50
                        .75
                                 .90
                                           . 95
 0.10597
          0.18805 0.31990 0.47906
                                      0.55551
lowest: 0.01772653 0.02853116 0.03021174 0.03724054 0.04076704
highest: 0.60114577 0.60689182 0.71653717 0.82781846 0.84216789
hw0$treatment: Treated
dd[x,]
                                                              .05
          missing distinct
                                Info
                                         Mean
                                                    Gmd
                                                                       .10
       n
                0
                         40
                                       0.4435
                                                 0.2411
                                                          0.1284
                                                                   0.1543
      40
                                   1
     . 25
              .50
                        .75
                                 .90
                                           .95
```

```
0.2972 0.4299 0.6032 0.6858 0.7658
```

lowest: 0.05370511 0.10947830 0.12944077 0.15246158 0.15452007 highest: 0.68436074 0.69878775 0.76469008 0.78683381 0.86569552

#### 3.3 Using the tableone library

Or, you could use the tableone library to produce a summarized Table 1 describing our results...

#### Stratified by treatment Not Treated Treated test р 95 40 prob (mean (SD)) 0.23 (0.18) 0.44 (0.21) < 0.001 linpred (mean (SD)) -1.44 (1.09) -0.29 (1.01) < 0.001 cov1 (mean (SD)) 47.07 (10.20) 51.42 (10.01) 0.025 cov2 (mean (SD)) 53.40 (9.79) 50.98 (9.30) 0.187 cov3 (mean (SD)) 21.45 (4.98) 17.50 (4.74) <0.001 cov4 (mean (SD)) 19.66 (4.17) 21.00 (4.83) 0.107 cov5 = 1 (%)38 (40.0) 0.378 20 (50.0)

You could even use non-parametric tests, and report quartiles for the continuous covariates...

```
print(tab1, nonnorm=c("prob", "linpred", "cov1", "cov2", "cov3", "cov4"))
```

```
Stratified by treatment
                       Not Treated
                                             Treated
                                                                   p
                           95
                                                 40
n
prob (median [IQR])
                         0.19 [0.11, 0.32]
                                              0.43 [0.30, 0.60]
                                                                   <0.001
linpred (median [IQR]) -1.46 [-2.13, -0.75] -0.28 [-0.86, 0.42]
                                                                   < 0.001
cov1 (median [IQR])
                       46.42 [39.97, 53.94] 51.07 [43.32, 60.79]
                                                                   0.046
cov2 (median [IQR])
                       54.08 [47.08, 60.36] 51.66 [44.02, 55.49]
                                                                    0.122
                       21.00 [18.00, 24.00] 18.00 [14.75, 21.00] < 0.001
cov3 (median [IQR])
                       20.00 [17.00, 23.00] 19.50 [18.00, 24.00]
cov4 (median [IQR])
                                                                   0.267
cov5 = 1 (\%)
                           38 (40.0)
                                                20 (50.0)
                                                                    0.378
                      Stratified by treatment
n
prob (median [IQR])
                       nonnorm
```

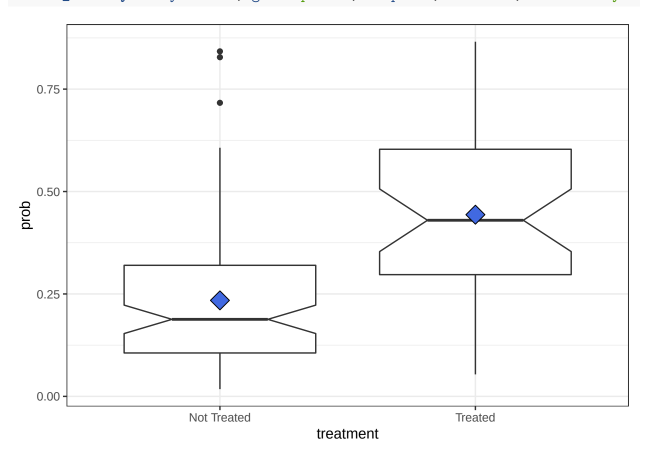
```
linpred (median [IQR]) nonnorm cov1 (median [IQR]) nonnorm cov2 (median [IQR]) nonnorm cov3 (median [IQR]) nonnorm cov4 (median [IQR]) nonnorm cov5 = 1 (%)
```

## 4 Plotting the Fitted Probabilities by Treatment Group

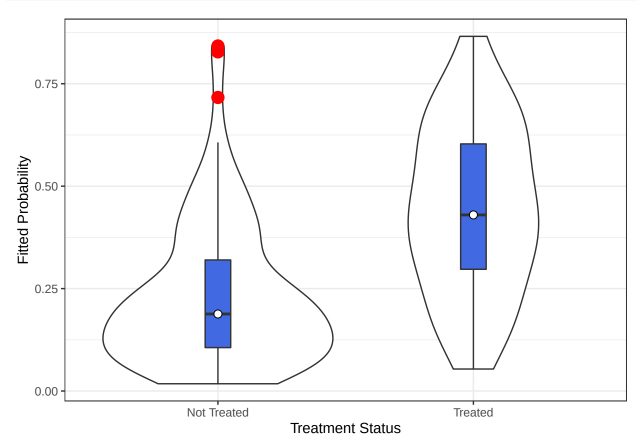
OK. So we've seen a numerical summary - let's focus on the important issue - a plot.

#### 4.1 A Boxplot using ggplot2, with Notches and Means Indicated

```
ggplot(hw0, aes(x = treatment, y = prob)) +
  geom_boxplot(notch=TRUE) +
  stat_summary(fun.y="mean", geom="point", shape=23, size = 5, fill = "royalblue")
```

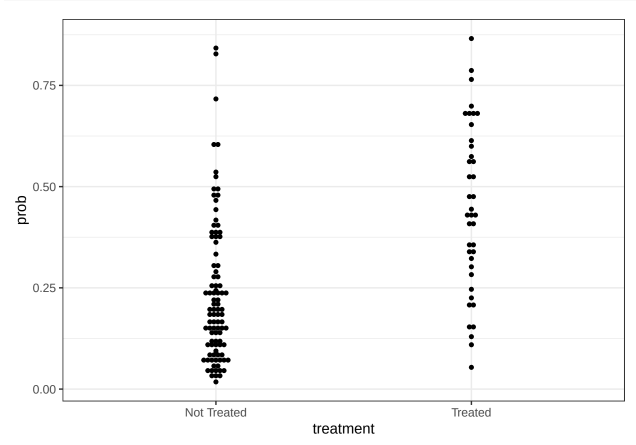


#### 4.2 A Violin Plot



## 4.3 A DotPlot to compare the probabilities, via ggplot2

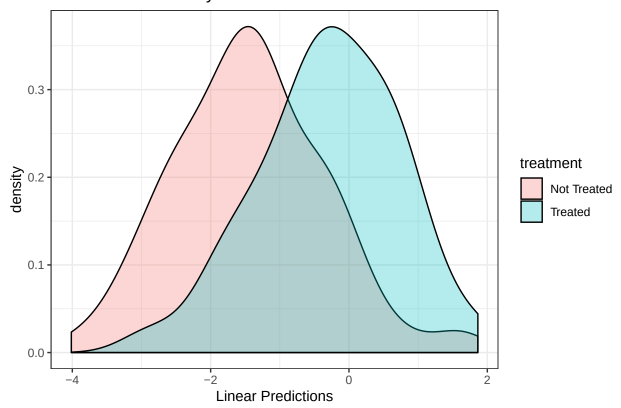
```
ggplot(hw0, aes(x = treatment, y = prob)) +
geom_dotplot(binaxis="y", binwidth=0.01, stackdir="center")
```



#### 4.4 A Density Plot, using ggplot2

A possibly more impressive picture would be a density plot. The best way to get this (here, I'll look at the linear probability (i.e. log odds of treatment) results rather than the raw probabilities on a 0-1 scale just to see if we observe something different) uses the ggplot2 library again...

#### **Linear Predictions By Treatment**

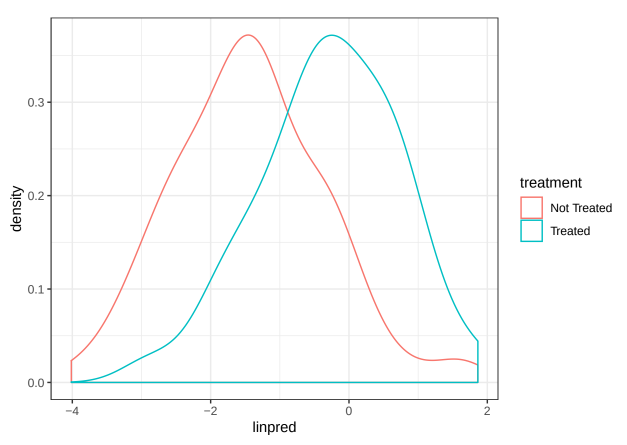


One advantage of the linear probabilities over the raw probability estimates is that the log odds results (linear probabilities) are a bit more likely to follow a normalish distribution. Again, it looks like there is fairly substantial overlap in the fitted probabilities across the treatment groups.

## 4.5 Another Density Plot, using ggplot2

We can use color instead of fill to indicate the densities.

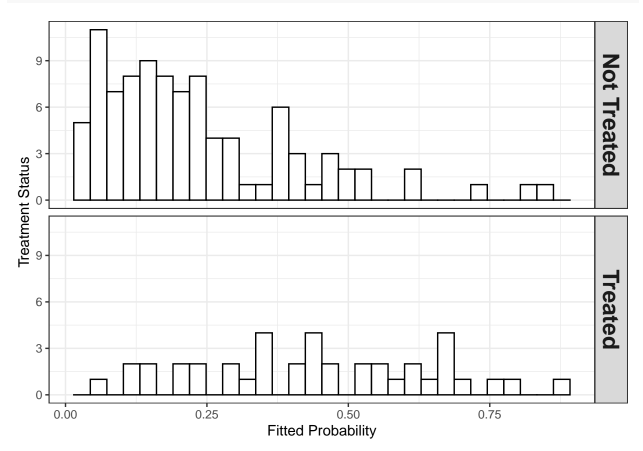
```
ggplot(hw0, aes(x=linpred, color=treatment)) +
  geom_density() +
  theme_bw()
```



### 4.6 Our Old Standby - Comparing Distributions via Histograms

The slickest approach I have here is this:

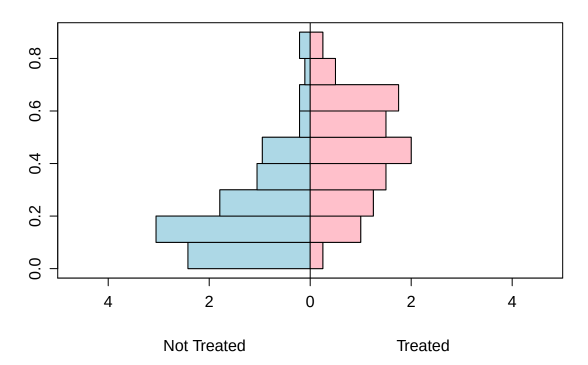
```
ggplot(hw0, aes(x = prob)) +
  geom_histogram(fill="white", color="black") +
  facet_grid(treatment ~ .) +
  theme(strip.text = element_text(face="bold", size=rel(1.5))) +
  xlab("Fitted Probability") + ylab("Treatment Status")
```



#### 4.7 A Back-to-Back Histogram

A former student suggested this approach, from the Hmisc library. There are likely better ways to get such a plot out of R.

#### **Back to Back Histogram of Fitted Probabilities**



#### 5 What About the ROC Curve and C Statistic?

Recall that our model m1 was

```
Call: glm(formula = treatment == "Treated" ~ cov1 + cov2 + cov3 + cov4 +
cov5, family = binomial(), data = hw0)
```

Since we're looking at a logistic regression, someone in a previous version of this class asked if I could show you how I get the C statistic (area under the ROC curve) for such a model. I usually use the Epi library . . .

Note that we need to specify the formula (abbreviated form in the ROC function) again, but that's it to get these results.

- The C statistic (area under the curve) for this logistic regression model is 0.786
- Very briefly, the ability of the model's predicted values to discriminate between patients with one outcome vs. the other is quantified by the area under the curve, also called the C statistic or concordance index, which ranges from 0.5 (discrimination is not better than chance) to 1.0 (perfect discriminating power.)
- The ROC procedure comes from signal detection theory and has been adopted into the language of diagnostic testing, essentially treating the response in the logistic regression model as the true status variable, and the set of predictors as the test to be evaluated by things like sensitivity, specificity, and positive and negative predictive values based on dichotomizing along the levels of the predictor set.
- For more on the ROC, visit Wikipedia for Receiver Operating Characteristic. Or try Google.
- A value of 0.786 would indicate a less-than-terrific model in terms of this issue. Values of 0.8 or even 0.9 are usually needed to declare the model to be reasonably accurate in this sense.

