Identify the relationship between referral rate and potential predictors using logistic regression

Elisa Zhang, Shicong Wang

4/8/2022

Visulization

1. Cancer Stage

The figure below illustrates the relationship between referral rate and cancer stages. There are many overlaps among error bars. The cancer stage might not be related to the referral rate.

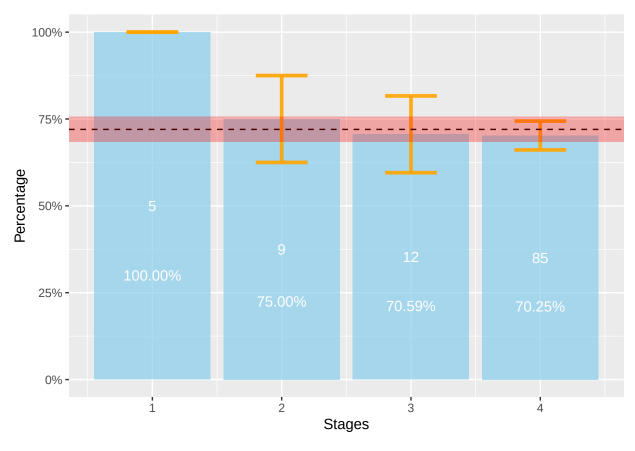


Figure 1: Proportion of Patients Who got referrals in each Cancer Stage

2. ECOG level

The figure below shows the relationship between ECOG status and referral rate. We can conclude that there is no obvious relationship between ECOG status and the referral rate.

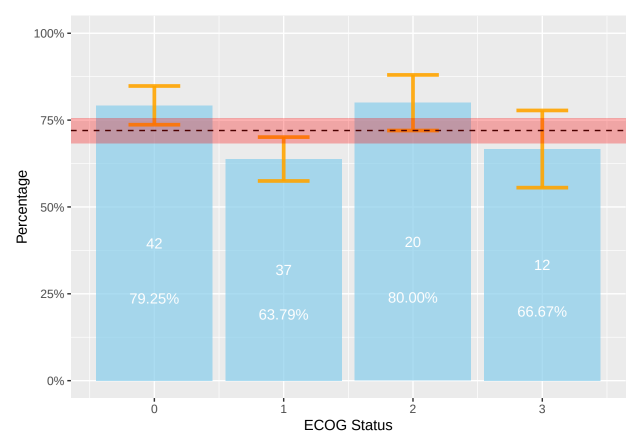
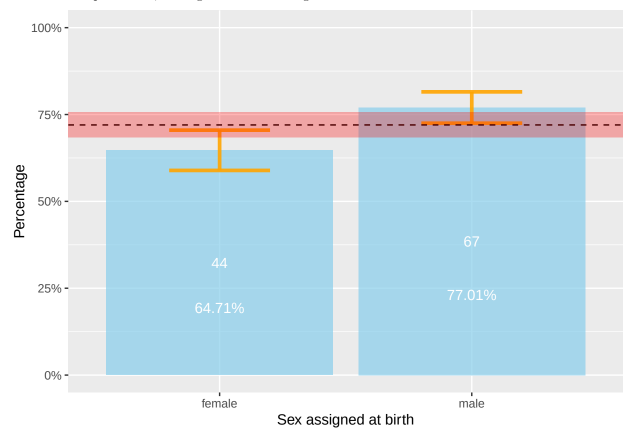


Figure 2: Proportions of Patients Who got referral rate in each ECOG Level

3. Sex Assigned at Birth

From the the plot below, we might conclude that gender is related to the referral rate.



4. Ethnicity

Since some ethnicity only contain few observations, we divide the race into white and not white and use as the new variable. And the figure below does not show that there is an obvious relationship between race and referral rate.

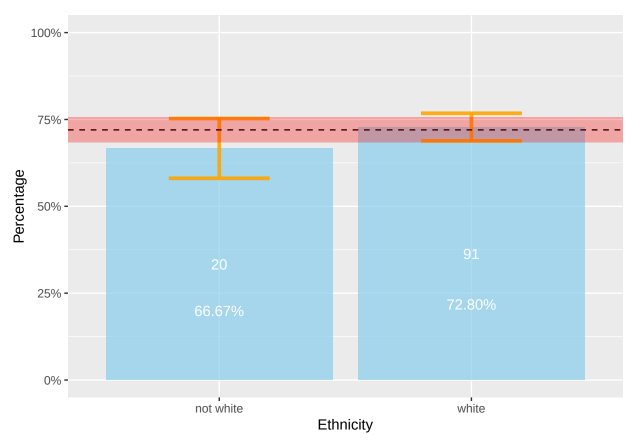


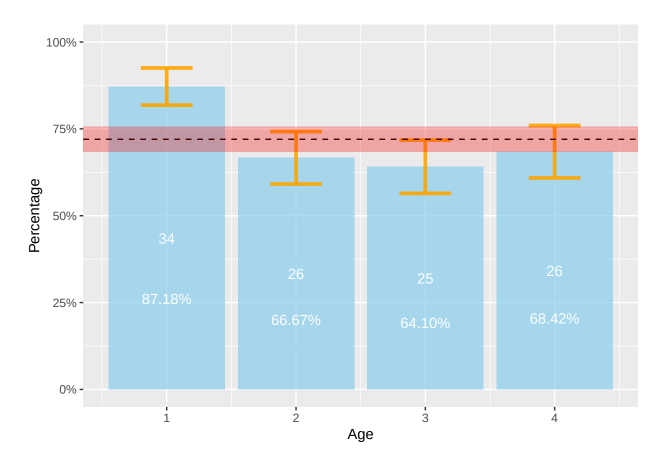
Figure 3: Proportions of Patients Who got referral rate among races

5. Age

We divide Age into groups using quantiles. The first group contains patients from 32 to 62 years old. And the second group contains patients aged from 62 to 68. And the third group include patients from 68 to 75.5 years old. And the last group include patients whose age are in the 4th quantile.

```
## 0% 25% 50% 75% 100%
## 32.0 62.0 68.0 75.5 96.0
```

From the plot below, there is no much difference for the referral rate among 2nd, 3rd and 4th groups.



Model Fit

We will use logistic regression to fit our data since we have binary outcome of whether the patients got a referral from GIM or not. We will fit three models. The full model will include both cancer stage and ECOG level. And the other two will only include either ECOG level or Cancer Stage.

```
##
## glm(formula = 'Genetics Referrals' ~ 'Stage at Dx (#0-4)' + 'ECOG at Initial (#0-4)' +
       'Age at Dx (#)' + 'Ethnicity/ Ancestry' + 'Sex Assigned\nat Birth (m/f)',
##
##
       family = "binomial", data = dat_1)
##
##
  Deviance Residuals:
                      Median
                                    3Q
##
       Min
                 1Q
                                            Max
                      0.7106
                                         1.2903
##
  -1.8614
           -1.3437
                                0.8358
##
## Coefficients:
##
                                          Estimate Std. Error z value Pr(>|z|)
```

```
## (Intercept)
                                       3.4843634 1.8067729 1.929
                                                                     0.0538 .
## 'Stage at Dx (#0-4)'
                                      -0.2976799 0.2814298 -1.058
                                                                     0.2902
                                                                     0.9984
## 'ECOG at Initial (#0-4)'
                                     -0.0003937 0.1960274 -0.002
## 'Age at Dx (#)'
                                      -0.0277649 0.0198487 -1.399
                                                                     0.1619
## 'Ethnicity/ Ancestry'white
                                       0.2280004 0.4502499
                                                            0.506
                                                                     0.6126
## 'Sex Assigned\\nat Birth (m/f)'male 0.5031592 0.3687174
                                                            1.365
                                                                     0.1724
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
      Null deviance: 184.94 on 154 degrees of freedom
## Residual deviance: 178.24 on 149 degrees of freedom
## AIC: 190.24
##
## Number of Fisher Scoring iterations: 4
Model without Cancer Stage
model2 <- glm(`Genetics Referrals` ~ `ECOG at Initial (#0-4)`</pre>
             + `Age at Dx (#)` +
               `Ethnicity/ Ancestry` +
               `Sex Assigned
at Birth (m/f),
family = "binomial", data = dat_1)
summary(model2)
##
## Call:
  glm(formula = 'Genetics Referrals' ~ 'ECOG at Initial (#0-4)' +
##
       'Age at Dx (#)' + 'Ethnicity/ Ancestry' + 'Sex Assigned\nat Birth (m/f)',
      family = "binomial", data = dat_1)
##
##
## Deviance Residuals:
##
      Min
            1Q Median
                                  3Q
                                          Max
## -1.8419 -1.3603
                    0.7124 0.8382
                                       1.2384
##
## Coefficients:
                                      Estimate Std. Error z value Pr(>|z|)
##
## (Intercept)
                                       2.29660 1.39086 1.651 0.0987 .
## 'ECOG at Initial (#0-4)'
                                      -0.04970
                                                  0.19004 - 0.262
                                                                   0.7937
## 'Age at Dx (#)'
                                      -0.02543
                                                  0.01943 - 1.309
                                                                   0.1907
## 'Ethnicity/ Ancestry'white
                                       0.19515
                                                  0.44769
                                                           0.436
                                                                   0.6629
## 'Sex Assigned\\nat Birth (m/f)'male 0.52785
                                                  0.36674
                                                           1.439
                                                                   0.1501
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 184.94 on 154 degrees of freedom
## Residual deviance: 179.46 on 150 degrees of freedom
## AIC: 189.46
## Number of Fisher Scoring iterations: 4
```

```
model3 <- glm(`Genetics Referrals` ~ `Stage at Dx (#0-4)` +</pre>
               `Age at Dx (#)` +
               `Ethnicity/ Ancestry` +
                `Sex Assigned
at Birth (m/f) , family = "binomial", data = dat_1)
summary(model3)
##
## Call:
## glm(formula = 'Genetics Referrals' ~ 'Stage at Dx (#0-4)' + 'Age at Dx (#)' +
       'Ethnicity/ Ancestry' + 'Sex Assigned\nat Birth (m/f)', family = "binomial",
##
##
       data = dat_1)
##
## Deviance Residuals:
##
      Min
                1Q
                    Median
                                  ЗQ
                                           Max
                                        1.2903
## -1.8613 -1.3438
                    0.7106 0.8357
##
## Coefficients:
##
                                       Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                                       3.48520 1.75846
                                                            1.982 0.0475 *
## 'Stage at Dx (#0-4)'
                                       -0.29780
                                                  0.27537 -1.081
                                                                   0.2795
## 'Age at Dx (#)'
                                       -0.02778
                                                  0.01862 -1.492
                                                                    0.1357
## 'Ethnicity/ Ancestry'white
                                       0.22809
                                                  0.44797
                                                             0.509
                                                                    0.6106
## 'Sex Assigned\\nat Birth (m/f)'male 0.50323
                                                  0.36703
                                                            1.371
                                                                    0.1703
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
       Null deviance: 184.94 on 154 degrees of freedom
##
## Residual deviance: 178.24 on 150 degrees of freedom
## AIC: 188.24
##
## Number of Fisher Scoring iterations: 4
```

None of the variables are associated with the processing of getting referral from GIM.

Diagnostic Plot

The diagnostic plot can be used to measure the goodness of fit of the model.

Binned residual plot

The binned residual plot can assess the overall fit of regression models for binary outcomes. Since most of points are within the grey line and there is no obvious pattern for the points, the model does not have much problem.

Binned residual plot

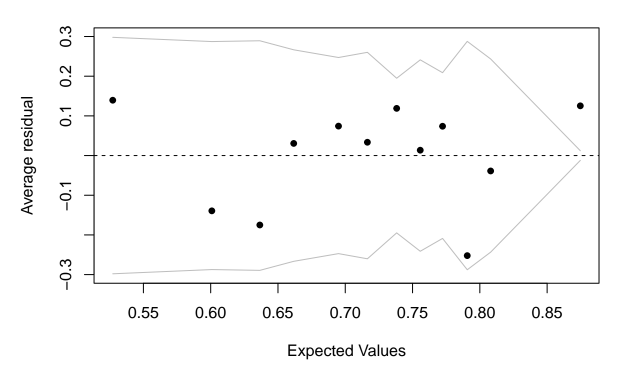


Figure 4: Binned residual plot for full model