## p8131\_hw3\_jsg2145

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## Problem 1

```
case_df = tibble(
 age = seq(from = 25, to = 75, by = 10),
 unexposed = c(0, 5, 21, 34, 36, 8),
  exposed = c(1, 4, 25, 42, 19, 5)
) %>%
 mutate(case = 1)
control_df = tibble(
  age = seq(from = 25, to = 75, by = 10),
  unexposed = c(106, 164, 138, 139, 88, 31),
 exposed = c(9, 26, 29, 27, 18, 0)
) %>%
 mutate(case = 0)
df = union(case_df, control_df) %>%
  pivot_longer(cols = c(exposed, unexposed), names_to = "status") %>%
  mutate(status = recode(status,
                         exposed = 1,
                         unexposed = 0))
fit = glm(cbind(value*(case == 1), value*(case == 0)) ~ status + age, family = binomial(link = 'logit')
summary(fit)
##
## Call:
## glm(formula = cbind(value * (case == 1), value * (case == 0)) ~
       status + age, family = binomial(link = "logit"), data = df)
##
## Deviance Residuals:
     \mathtt{Min}
           1Q Median
                               3Q
                                      Max
## -7.359 -5.265 0.000
                            4.223 11.108
##
## Coefficients:
              Estimate Std. Error z value Pr(>|z|)
##
## (Intercept) -5.02345
                           0.41821 -12.012 <2e-16 ***
               1.78000
                           0.18708
                                   9.514
                                             <2e-16 ***
## status
                0.06158
                           0.00729
                                     8.446
                                             <2e-16 ***
## age
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
```

```
## Null deviance: 989.49 on 21 degrees of freedom
## Residual deviance: 809.81 on 19 degrees of freedom
## AIC: 815.81
##
## Number of Fisher Scoring iterations: 4

fit = fit %>%
   broom::tidy()
```

The difference in log odds of developing cancer is 1.78 for those with the exposure (>80 grams of alcohol per day) compared to those without, holding age constant.

The log odds ratio of developing cancer is 0.06 for each 1 unit increase in age, holding exposure constant.

## Problem 2

```
bean 75 = tibble(
  y = c(10, 23, 23, 26, 17),
  m = c(39, 62, 81, 51, 39)) \%
  mutate(extract = "bean",
         seed = "75")
cuc_{75} = tibble(
  y = c(5, 53, 55, 32, 46, 10),
  m = c(6, 74, 72, 51, 79, 13)) \%
  mutate(extract = "cuc",
         seed = "75")
bean_73 = tibble(
  y = c(8, 10, 8, 23, 0),
  m = c(16, 30, 28, 45, 4)) \%
  mutate(extract = "bean",
         seed = "73")
cuc_73 = tibble(
  y = c(3, 22, 15, 32, 3),
  m = c(12, 41, 30, 51, 7)) \%
  mutate(extract = "cuc",
         seed = "73")
df_o = union_all(bean_75, cuc_75) %>%
  union_all(bean_73) %>%
  union_all(cuc_73) %>%
  mutate(y_m = m - y,
         extract = recode(extract,
                          "bean" = 0,
                          "cuc" = 1),
         seed = recode(seed,
                       "75" = 0,
                       "73" = 1))
fit_o = glm(cbind(y, y_m) ~ seed + extract, family = binomial(link = 'logit'), df_o)
```

```
summary(fit_o)
```

```
##
## Call:
## glm(formula = cbind(y, y_m) ~ seed + extract, family = binomial(link = "logit"),
##
       data = df_o)
##
## Deviance Residuals:
##
      Min
                10
                     Median
                                  3Q
                                           Max
## -2.3919 -0.9949 -0.3744
                              0.9831
                                        2.4766
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.4300
                            0.1137 -3.781 0.000156 ***
               -0.2705
                            0.1547 -1.748 0.080435 .
## seed
## extract
                1.0647
                            0.1442
                                    7.383 1.55e-13 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 98.719 on 20 degrees of freedom
## Residual deviance: 39.686 on 18 degrees of freedom
## AIC: 122.28
##
## Number of Fisher Scoring iterations: 4
```

The log odds ratio for germination is reduced by -0.27 for species Orobanche aegyptiaca 73 compared to Orobanche aegyptiaca 75, holding the extract media type constant.

The log odds ratio for germination is increased by 1.06 for cucumber extract media compared to bean extract media, holding the species constant.

There is a log odds of -0.43 for plants that are species Orobanche aegyptiaca 75 grown in bean extract media.

```
# Evaluate dispersion with deviance

dev_o = deviance(fit_o)

pval = 1 - pchisq(dev_o, 19)
pval
```

```
## [1] 0.003597422
```

A low p-value (<0.05) indicates that there is over dispersion

```
# Estimate phi with the G statistic

G.stat = sum(residuals(fit_o, type = "pearson")^2)
G.stat
```

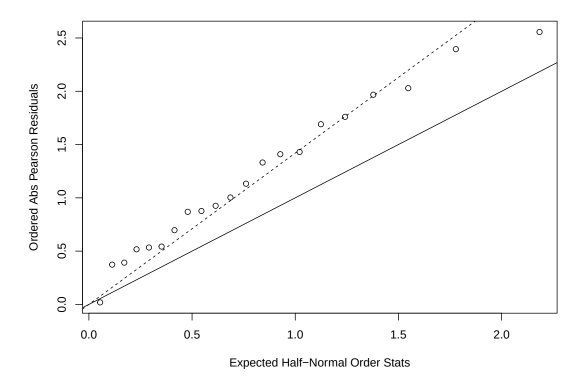
```
## [1] 38.31062
```

```
phi = G.stat/(21 - 2)
phi
## [1] 2.016348
```

```
# Estimate phi with the deviance
dev_o/(21 - 2)
```

```
## [1] 2.088731
```

```
res_o = residuals(fit_o, type = "pearson")
plot(qnorm((21 + 1:21 + 0.5) / (2 * 21 + 1.125)), sort(abs(res_o)), xlab = 'Expected Half-Normal Order abline(a = 0, b = 1)
abline(a = 0 , b = sqrt(phi), lty = 2)
```



```
summary(fit_o, dispersion = phi)
```

```
##
## Call:
## glm(formula = cbind(y, y_m) ~ seed + extract, family = binomial(link = "logit"),
## data = df_o)
##
## Deviance Residuals:
```

```
##
       Min
                10
                     Median
                                  3Q
                                          Max
## -2.3919 -0.9949 -0.3744
                              0.9831
                                       2.4766
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.4300
                           0.1615 -2.663 0.00775 **
## seed
                -0.2705
                            0.2197 -1.231 0.21828
                1.0647
                            0.2048
                                             2e-07 ***
## extract
                                    5.199
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 2.016348)
##
       Null deviance: 98.719 on 20 degrees of freedom
##
## Residual deviance: 39.686 on 18 degrees of freedom
## AIC: 122.28
##
## Number of Fisher Scoring iterations: 4
```

The type of seed is no longer significant in the model when a dispersion factor is included.

The reason for the over dispersion may be correlation within each treatment group.