## assess 1

## Jodie

## 2024-04-17

General question: In this randomized study, what is the effect of the drug on elevated potassium levels (i.e. hyperkalemia)?

Data sets: 1. t05xc: Adverse event data 2. t016: Lab test data 3. master\_data: Master database for clinical trial

Notes: 1. Hyperkalemia is defined as any potassium value >=5.5 2. Potassium values are recorded in the column labeled "k" in both the adverse event data set and the lab test data set. 3. Each patient may have multiple potassium values recorded. We are interested in whether each patient EVER experienced hyperkalemia. Assume that if no potassium values are available, then hyperkalemia was not present. 4. Patients are identified by "master\_id" in each of the data sets. 5. Each patient's treatment assignment is identified by "treat" (0 for placebo; 1 for drug) in the master data set. 6. Geographic region is identified by "region" in the master data set.

Questions: 1. Is there evidence that the drug is associated with hyperkalemia? 2. Does the drug effect depend on geographic region?

Further questions: after concluding there is an association, how to identify the relationship—try doing regression analysis? What test to answer—does the drug effect depend on geographic region?

Is there evidence that the drug is associated with hyperkalemia? Primary approach: Do hypothesis test to see if there significant difference between drug group and placebo group -> chi sq test of independence (essentially a log linear model) Do logistic regression on the drug, with a random intercept terms for the patient and for the geographic region. Regress against hyperkalemia as label In other words, response variable is hyperkalemia and feature

```
# join dataframe by master_id

# test that looks at
# assuming all other variables are held constant
# do hypothesis test to see if there significant difference between drug group and placebo group -> chi
## HO: no difference in proportion between groups; Ha: there is difference between groups; if p-val < 0
# there hyperkalmeia is associated with presence of drug

# create table
# 1. create two subsets of data--> 1 for drug and 0 for no drug
# 2. calculate num of patients who experienced hyperkalemia in each group
# manipulating dataframe
```

library(dplyr)

Create a new column-overall\_hyperk and join dataframes to create a single dataframe that has all variables of interest "treat", "region", "master\_id", "overall\_hyperk"

```
## Warning: package 'dplyr' was built under R version 4.2.3
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
       filter, lag
## The following objects are masked from 'package:base':
##
##
       intersect, setdiff, setequal, union
library(tidyr)
## Warning: package 'tidyr' was built under R version 4.2.3
master <- read.csv("/Users/jojoc/Downloads/Data specialist assessment/Assignment No 1/master_data.csv")
adverse.df <- read.csv("/Users/jojoc/Downloads/Data specialist assessment/Assignment No 1/t016.csv")
lab.df <- read.csv("/Users/jojoc/Downloads/Data specialist assessment/Assignment No 1/t05xc.csv")
#head(master)
#head(adverse.df)
#tail(lab.df)
#table(lab.df$test_type)
lab.df$test_type <- tolower(lab.df$test_type)</pre>
#lab.df
# revise lab df
# select only rows with "pota" or "k" in test_type col for lab.df using regular expression --> to inclu
sel_lab_ <- lab.df[grep1("pota|k", lab.df$test_type, ignore.case = TRUE), ]</pre>
#sel_lab_# updated lab df to contain only potassium tests, need to aggregate by patient later to assign
# aggregate by master_id to check if at least one k is >= 5.5; if so assign 1 (true) to has_hyperk col
sel_lab_2 <- sel_lab_ %>%
 mutate(k = replace_na(k, 0)) %>%
  group_by(master_id) %>%
  summarize(has_hyperk = any(k >= 5.5)) %>%
  mutate(has_hyperk_num = as.numeric(has_hyperk))
#sel_lab_2 # most updated lab
#sel_lab_
# revise master to contain treat and drug
rev_master <- master[, c("master_id", "treat", "region")] # only one occurrence for each patient</pre>
                                                           # as determined doing table()
#rev master # most updated master
unique(rev_master$region) # region 1 or 2
```

## ## [1] 1 2

```
# revise adverse events df to also have has_hyperk col
adverse.df2 <- adverse.df %>%
  select(master_id, k) %>%
  mutate(k = replace_na(k, 0)) %>% # replace na's with 0
  group_by(master_id) %>%
  summarize(has_hyperk = any(k >= 5.5)) %>%
  mutate(has_hyperk_num = as.numeric(has_hyperk))
# most updated dataframes are sel_lab_2, rev_master, and adverse.df2
head(rev master)
##
    master_id treat region
## 1 10010013
                   1
## 2 10010025
                   0
## 3 10010037
## 4 10010049
                          1
                 1
## 5 10010051
                 1
## 6 10010063
                  0
                         1
head(sel_lab_2)
## # A tibble: 6 x 3
   master_id has_hyperk has_hyperk_num
##
         <int> <lgl>
                                   <dbl>
## 1 10010025 FALSE
                                       0
## 2 10010051 FALSE
                                       0
## 3 10010102 TRUE
                                       1
## 4 10010114 FALSE
                                       0
## 5 10010152 FALSE
                                       0
## 6 10010188 FALSE
                                       0
head(adverse.df2)
## # A tibble: 6 x 3
    master_id has_hyperk has_hyperk_num
##
         <int> <lgl>
                                 <dbl>
## 1 10010013 FALSE
                                       0
## 2 10010025 FALSE
                                       0
## 3 10010037 FALSE
                                       0
## 4 10010049 TRUE
                                       1
## 5 10010051 TRUE
                                       1
## 6 10010063 FALSE
                                       0
\# might be some overlaps between sel_lab_2 and adverse.df2--
# assuming that that there are patients in both dataframes,
# if has hyperk is true in adverse.df2 but false in sel_lab_2, mark as true overall for has_hyperk
sel_lab_adverse <- rbind(sel_lab_2, adverse.df2)</pre>
sel_lab_adverse
```

```
## # A tibble: 3,826 x 3
##
     master_id has_hyperk has_hyperk_num
          <int> <lgl>
                                    <dbl>
##
  1 10010025 FALSE
                                        0
##
##
   2 10010051 FALSE
                                        0
## 3 10010102 TRUE
                                        1
## 4 10010114 FALSE
                                        0
## 5 10010152 FALSE
                                        0
## 6 10010188 FALSE
                                        0
## 7 10010190 FALSE
                                        0
## 8 10010215 FALSE
                                        0
## 9 10010227 FALSE
                                        0
## 10 10010241 FALSE
                                        0
## # i 3,816 more rows
sel_lab_adverse2 <- sel_lab_adverse %>%
  group_by(master_id) %>%
  summarize(overall_has_hyperk = any(has_hyperk_num==1)) %>%
  mutate(overall_hyperk = as.numeric(overall_has_hyperk))
dim(sel_lab_adverse2) # final joint data frame created by joining sel_lab_2 and adverse.df2
## [1] 3437
               3
dim(rev_master)
## [1] 3445
# join sel_lab_adverse2 and rev_master by master_id
# include all=TRUE to include patients that only appear in one of the two dfs being merged
final_df <- merge(sel_lab_adverse2, rev_master, by = "master_id")</pre>
head(final_df) # 3446 entries (when all=TRUE), otherwise 3,436 when all is false
     master_id overall_has_hyperk overall_hyperk treat region
## 1 10010013
                            FALSE
                                               0
                                                     1
                                                            1
## 2 10010025
                            FALSE
                                               0
                                                     0
                                                            1
                                               0
## 3 10010037
                            FALSE
                                                     1
                                                            1
## 4 10010049
                             TRUE
                                               1
                                                     1
                                                            1
## 5 10010051
                             TRUE
                                               1
                                                     1
                                                            1
## 6 10010063
                            FALSE
                                                            1
is.factor(final_df$region) # check if this column has levels, currently not so convert
```

To answer Q1, logistic regression on the treat and region and regress against overall\_hyperk as label.

## [1] FALSE

```
final_df$overall_hyperk <- as.factor(final_df$overall_hyperk)</pre>
final_df$treat <- as.factor(final_df$treat)</pre>
final_df$region <- as.factor(final_df$region) # making them factors might not make diference since each
# m1 <- glm(overall_hyperk ~ treat, data = final_df, family = binomial)</pre>
# summary(m1)
m1_2 <- glm(overall_hyperk ~ treat + region, data = final_df, family = binomial)
summary(m1_2)
##
## Call:
## glm(formula = overall_hyperk ~ treat + region, family = binomial,
       data = final df)
##
## Deviance Residuals:
##
       Min
                 1Q
                      Median
                                   3Q
                                           Max
  -0.7171 -0.5590 -0.4886 -0.3756
                                         2.3180
##
## Coefficients:
##
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -2.06542
                           0.09272 -22.276 < 2e-16 ***
                                     8.007 1.18e-15 ***
## treat1
                0.83868
                           0.10475
## region2
               -0.55049
                           0.10253 -5.369 7.92e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
##
## (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 2768.2 on 3435 degrees of freedom
## Residual deviance: 2671.1 on 3433 degrees of freedom
## AIC: 2677.1
##
## Number of Fisher Scoring iterations: 5
levels(final_df$treat) # inrepretreting the multiplicative change of odds in favor of y=1 as patient go
## [1] "0" "1"
chisq.test(table(final_df$overall_hyperk, final_df$treat))
##
##
  Pearson's Chi-squared test with Yates' continuity correction
##
## data: table(final_df$overall_hyperk, final_df$treat)
## X-squared = 65.475, df = 1, p-value = 5.885e-16
```

Drug effect is significantly associated with presence of hyperkalemia as indicated by the chi-squared test. The estimate for treat is 0.83868 (significant as it its p-value < 0.05). For patients on drugs, the odds of developing hyperkalemia is multiplied by a factor of  $\exp(0.83868) = 2.31$  compared to patients not on drugs. In other words, the odds of developing hyperkalemia when on drugs are 2.31 times the odds for when one is not on drugs. Another way of interpreting this is, the probability of developing hyperkalemia are  $(2.31 - 1) \times 100 = 131\%$  higher when on drugs compared to not being on drugs.

```
m2 <- glm(overall_hyperk ~ treat*region + treat + region, data = final_df, family = binomial)
summary(m2)</pre>
```

To answer Q2, add an interaction term (treatment\*region) to the model and compare with m1\_2 (without the interaction term)

```
##
## Call:
## glm(formula = overall_hyperk ~ treat * region + treat + region,
##
       family = binomial, data = final_df)
##
## Deviance Residuals:
##
                      Median
                                   3Q
       Min
                 1Q
                                           Max
## -0.7610 -0.5024 -0.4445
                             -0.4284
                                        2.2063
##
## Coefficients:
##
                  Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                  -2.34206
                              0.11931 -19.630
                                               < 2e-16 ***
## treat1
                   1.25098
                              0.14231
                                        8.790
                                               < 2e-16 ***
                   0.07687
## region2
                              0.16795
                                        0.458
                                                  0.647
                                      -4.641 3.47e-06 ***
## treat1:region2 -0.99190
                              0.21374
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
##
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
       Null deviance: 2768.2
                             on 3435
                                       degrees of freedom
## Residual deviance: 2649.5 on 3432
                                       degrees of freedom
## AIC: 2657.5
##
## Number of Fisher Scoring iterations: 5
```

If drug effect or treatment depends on region, we expect the model with the interaction term to fit our data better and is therefore the better model. To compare the models, we can use AIC which is included in the summary output. AIC tells you which model has better out of sample prediction accuracy. The better model will have a lower AIC. In this case, the model with the interaction term has lower AIC 2657.5 < 2677.1. Therefore, our model with the drug and region interaction term does better and drug depends on region. I considered computing the leave one cross validation to compare the two models but AIC for regression models is a sufficient approximation. It would be computationally expensive since I would need to fit the model once for each data point.

Another way to answer Q2, compute the CI for the interaction term