# P8130 Biostatistical Methods Homework 5

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### Problem 1

```
#Read the CSV data into a dataframe
antibodies_df <- read.csv("./data/Antibodies.csv")

# Make AgeCategory, Smell, and Gender appopriate datatypes. Helps to ensure we know all unique values,
unique(antibodies_df$AgeCategory)
    antibodies_df <- antibodies_df %>%
        mutate(AgeCategory = factor(AgeCategory, labels = c("18-30", "31-50", "51+") ))
unique(antibodies_df$Smell)
    antibodies_df <- antibodies_df %>%
        mutate(Smell = factor(Smell, levels = c("Normal", "Altered", "Unanswered/Others")))
    antibodies_df = antibodies_df %>% filter(Smell != "Unanswered/Others")
unique(antibodies_df$Gender)
    antibodies_df <- antibodies_df %>%
        mutate(Gender = factor(Gender, levels = c("Male", "Female")))
```

In order to assess the difference in IgM levels between the two smell factor groups (Normal vs Altered) given non-normal distributions, we opt for a non-parametric test called the Wilcoxon Rank-Sum test. It's the nonparametric equivalent of the two-sample independent t-test, and examines if the medians of the two populations are equal versus not equal:

 $H_0$  = the medians of the two groups' IgM levels are equal, and  $H_A$  = the medians of the two groups' IgM levels are not equal. The decision rule is that we reject  $H_0$  = if  $T > z_{1-(\alpha/2)}$ .

The test statistic is computed, with a continuity correction, one of two ways. We first combine the data from the two groups, order the values from lowest to highest, assign ranks to the individual values (1 to n), and if ties, assign the average rank. Then, select a group and compute the sum of ranks  $T_1$  for the first group, and then use the appropriate test statistic formula.

With no ties (referring to two equally ranked values once the values are listed), the test statistic is

$$T = \frac{\left| T_1 - \frac{n_1(n_1 + n_2 + 1)}{2} \right| - \frac{1}{2}}{\sqrt{(n_1 n_2 / 12)(n_1 + n_2 + 1)}}$$

and with ties, the test statistic is

$$T = \frac{\left| T_1 - \frac{n_1(n_1 + n_2 + 1)}{2} \right| - \frac{1}{2}}{\sqrt{(n_1 n_2 / 12)[(n_1 + n_2 + 1) - \sum_{i=1}^g t_i (t_i^2 - 1) / (n_1 + n_2)(n_1 + n_2 - 1)]}}$$

where  $t_i$  refers to the number of observations with the same absolute value in the  $i^{th}$  group and g is the number of tied groups.

In our case, the test statistic calculated by R is slightly different, since it does not by default add the  $n_1(n_1+1)/2$  term (and is denoted by W).

The p-value under the normal approximation, with  $n_1$  and  $n_2 \ge 10$  is described by  $2 * [1 - \Phi(T)]$ .

```
antibodies_df2 =
  antibodies_df %>%
pivot_wider(
         names_from = Smell,
         values_from = Antibody_IgM
    )

wilcox.test(antibodies_df2$Normal, antibodies_df2$Altered, mu = 0)

##

## Wilcoxon rank sum test with continuity correction

##

## data: antibodies_df2$Normal and antibodies_df2$Altered

## W = 5836, p-value = 0.01406

## alternative hypothesis: true location shift is not equal to 0
```

In context, and ignoring missing values and the unanswered smell category, we find evidence to reject the null hypothesis and conclude that the true location shift between the Normal and Altered smell categories is not equal to zero (in other words, the median IgM values are different for the two groups).

# Problem 2

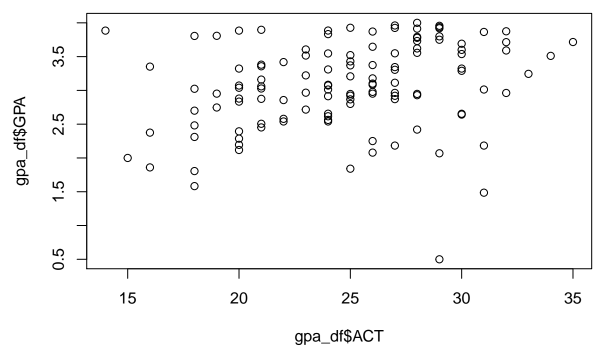
Let's come back to this.

#### Problem 3

#### Part 1

Load the data and plot it.

```
gpa_df <- read.csv("./data/GPA.csv")
plot(gpa_df$ACT, gpa_df$GPA)</pre>
```



whether a linear association exists between ACT score (x) and GPA at the end of freshman year (Y).

Let  $\alpha = 0.05$  and let  $\beta_1$  represent the true slope to be estimated.

The null hypothesis is  $H_0: \beta_1 = \beta_{10}$  where  $\beta_{10} = 0$ . The alternative hypothesis is  $H_A: \beta_1 \neq \beta_{10}$ . In context, testing  $H_0: \beta_1 = 0$  examines whether a student's GPA at the end of freshman year can be predicted from the ACT test score.

Test

The test statistic follows the t distribution with n-2 degrees of freedom, such that

$$t = \frac{\widehat{\beta_1} - \beta_{10}}{se(\widehat{\beta_1})} \sim t_{n-2}$$
, under  $H_0 = \frac{0.03883 - 0}{0.01277} = 3.040$  using degrees of freedom  $= n = 120, df = n - 2$ 

The corresponding critical value is fixed by  $t_{n-2,1-(\alpha/2)}$  and the decision rule is that we reject  $H_0$  if  $|t| > t_{n-2,1-(\alpha/2)}$  and fail to reject  $H_0$  if  $|t| \le t_{n-2,1-(\alpha/2)}$ . As such, the critical value is  $t_{118,0.975} = 1.980272$ .

Therefore,  $|t| > t_{118,0.975}$  using the 5% significance level, we find evidence to reject the null hypothesis and conclude that there is a significant linear association between students' ACT scores and GPA at the end of freshman year.

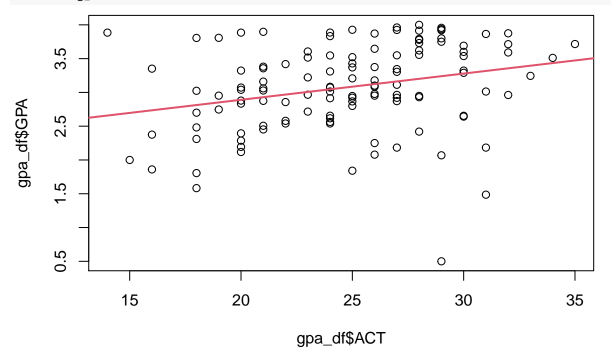
```
reg_admit<-lm(gpa_df$GPA~gpa_df$ACT)
# Summarize regression
summary(reg_admit)
##
## Call:
##
  lm(formula = gpa_df$GPA ~ gpa_df$ACT)
##
##
   Residuals:
##
        Min
                                      3Q
                                              Max
                   1Q
                        Median
##
   -2.74004 -0.33827
                       0.04062
                                0.44064
                                          1.22737
##
##
  Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                2.11405
                            0.32089
                                       6.588
                                              1.3e-09 ***
## gpa_df$ACT
                 0.03883
                            0.01277
                                       3.040
                                              0.00292 **
```

```
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.6231 on 118 degrees of freedom
## Multiple R-squared: 0.07262,
                                    Adjusted R-squared: 0.06476
## F-statistic: 9.24 on 1 and 118 DF, p-value: 0.002917
tidy(reg_admit)
## # A tibble: 2 x 5
##
    term
                 estimate std.error statistic
                                                     p.value
##
     <chr>>
                   <dbl>
                              <dbl>
                                        <dh1>
                                                       <dbl>
## 1 (Intercept)
                   2.11
                             0.321
                                         6.59 0.00000000130
## 2 gpa_df$ACT
                   0.0388
                             0.0128
                                         3.04 0.00292
glance(reg_admit)
## # A tibble: 1 x 12
   r.squared adj.r.squared sigma statistic p.value
                                                         df logLik
                                                                     AIC
                                                                           BIC
                       <dbl> <dbl>
                                       <dbl> <dbl> <dbl> <dbl> <dbl> <dbl> <
                                                          1 -113. 231.
        0.0726
                      0.0648 0.623
                                        9.24 0.00292
## 1
## # ... with 3 more variables: deviance <dbl>, df.residual <int>, nobs <int>
# Regression objects
names(reg_admit)
## [1] "coefficients" "residuals"
                                         "effects"
                                                         "rank"
## [5] "fitted.values" "assign"
                                         "qr"
                                                         "df.residual"
## [9] "xlevels"
                        "call"
                                         "terms"
                                                         "model"
# Get fitted.values
reg_admit$fitted.values
                            3
                                     4
                                              5
                   2
                                                        6
                                                                 7
## 2.929419 2.657629 3.201209 2.968246 2.929419 3.317690 3.356517 3.162382
          9
                  10
                           11
                                    12
                                             1.3
                                                       14
                                                                15
## 3.240036 3.123555 3.045900 3.278863 3.045900 3.045900 3.395344 3.162382
                                                      22
                                                                23
                  18
                           19
                                    20
                                             21
         17
## 3.084727 3.317690 3.084727 2.890592 3.045900 2.929419 3.201209 3.162382
         25
                  26
                           27
                                    28
                                             29
                                                       30
                                                                31
## 3.201209 3.123555 3.201209 2.968246 3.123555 2.929419 3.084727 2.735283
         33
                  34
                           35
                                    36
                                             37
                                                       38
                                                                39
## 3.201209 3.123555 2.968246 3.045900 2.929419 3.278863 3.162382 3.123555
                  42
                           43
                                              45
         41
                                    44
                                                       46
                                                                47
## 3.123555 3.278863 3.045900 3.123555 3.240036 3.045900 3.317690 2.696456
         49
                  50
                           51
                                    52
                                             53
                                                      54
                                                                55
## 2.851765 2.812938 3.162382 2.735283 3.162382 3.123555 3.045900 3.278863
         57
                  58
                           59
                                    60
                                             61
                                                      62
                                                                63
## 2.929419 2.890592 3.278863 3.240036 3.084727 3.007073 3.084727 3.007073
##
                  66
                           67
                                    68
                                             69
                                                       70
                                                                71
## 3.278863 2.929419 3.045900 3.356517 2.812938 3.007073 2.890592 3.007073
                  74
                           75
                                    76
                                             77
                                                      78
                                                                79
                                                                         80
## 2.812938 2.812938 3.240036 2.890592 3.007073 3.123555 3.201209 3.434172
                  82
                           83
                                    84
                                             85
                                                      86
                                                                87
## 2.890592 2.890592 3.123555 3.356517 3.084727 3.162382 3.162382 3.240036
         89
                  90
                           91
                                    92
                                             93
                                                       94
                                                                95
```

## 2.851765 2.929419 3.045900 3.162382 3.084727 2.812938 3.240036 3.045900

```
97
                   98
                            99
                                     100
                                              101
                                                        102
                                                                 103
                                                                           104
##
## 3.162382 2.929419 2.851765 2.812938 3.084727 2.812938 2.890592 3.356517
##
        105
                  106
                           107
                                              109
                                                        110
                                                                 111
                                     108
## 3.045900 3.472999 3.084727 3.201209 3.201209 3.084727 2.968246 3.278863
                                                       118
##
        113
                  114
                           115
                                     116
                                              117
                                                                 119
                                                                           120
## 2.890592 2.890592 3.317690 2.890592 3.240036 3.201209 2.735283 3.201209
```

```
# Scatterplot and regression line overlaid
plot(gpa_df$ACT, gpa_df$GPA)
abline(reg_admit,lwd=2,col=2)
```



Part 2 The basic regression model follows the form  $Y_i = \beta_0 + \beta_1 X_i + \epsilon_i$