# Stat 502 HW3

### Dongvang Wang

10/25/2021

1

(a)

```
\#rm(list = ls())
sleep <- sleep
sleep1 <- sleep[sleep$group == 1,]</pre>
sleep2 <- sleep[sleep$group == 2,]</pre>
t.test(sleep1$extra, sleep2$extra, var.equal = T)
##
##
    Two Sample t-test
##
## data: sleep1$extra and sleep2$extra
## t = -1.8608, df = 18, p-value = 0.07919
## alternative hypothesis: true difference in means is not equal to 0
```

-3.363874 0.203874

## sample estimates:

## mean of x mean of y

## 0.75 2.33

The confidence interval is [-3.363874, 0.203874].

## 95 percent confidence interval:

Since  $Y_A, Y_B$  are normally distributed,  $Y_A - Y_B \sim N(\delta, \frac{\sigma^2}{n_A} + \frac{\sigma^2}{n_B})$ . To form a standard normal, we have  $\frac{Y_A - Y_B - \delta}{\sqrt{\frac{\sigma^2}{n_A} + \frac{\sigma^2}{n_B}}}$ . Since we also know  $\frac{n_A + n_B - 2}{\sigma^2} s_P^2$  follows chi-squared distribution with  $n_A + n_B - 2$  as the dof, and

X and Z are independent. So, by definition of t-statistic, we have  $t(Y_A, Y_B) = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{n_A} + \frac{n_B^2}{n_B}}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}} = \frac{Y_A - Y_B}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}} = \frac{\frac{Y_A - Y_B - \delta}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}{\sqrt{\frac{n_A + n_B - 2}{\sigma^2} s_P^2/(n_A + n_B - 2)}}}$  $\frac{Y_A - Y_B}{s_p \sqrt{\frac{1}{n_A} + \frac{1}{n_B}}} - \frac{\delta}{s_p \sqrt{\frac{1}{n_A} + \frac{1}{n_B}}}$ . As we can observe, this is a non-central t-distribution.

The components are (i)  $Z = \frac{Y_A - Y_B - \delta}{\sqrt{\frac{\sigma^2}{n_A} + \frac{\sigma^2}{n_B}}}$ . (ii)  $X = \frac{n_A + n_B - 2}{\sigma^2} s_P^2$ . (iii) The non-centrality parameter is  $\gamma = \frac{-\delta}{\sqrt{\frac{\sigma^2}{n_A} + \frac{\sigma^2}{n_B}}}.$ 

(c) Since power is an increasing function of absolute value of the non centrality parameter  $\gamma = |\frac{\delta}{\sqrt{\frac{\sigma^2}{n_A} + \frac{\sigma^2}{n_B}}}|$ , we want to maximize  $\gamma$ . Rewriting with  $n_B = N - n_A$ , we have  $\gamma = |\frac{\delta}{\sigma\sqrt{\frac{1}{n_A} + \frac{1}{N - n_A}}}| = |\frac{\delta}{\sigma\sqrt{\frac{N}{n_A(N - n_A)}}}| = |\frac{\delta}{\sigma\sqrt{\frac{N}{n_A(N - n_A)}}}|$  $\left|\frac{\delta\sqrt{n_A(N-n_A)}}{\sigma\sqrt{N}}\right|$ . To maximize this, we let  $N-n_A=n_A$ , that is  $n_A=\frac{1}{2}N$ . Hence, we maximized power. 3

(a) The model is  $y_{ij} = \mu + \tau_i + \epsilon_{ij}$ . The meaning of  $\mu$  means the mean when no treatment is in place.  $\tau_i$  represent the within treatment variation for the four treatments. Key assumptions include  $E(\epsilon_{ij}) = 0$ ,  $Var(\epsilon_{ij}) = \sigma^2$ . We also want the sum of  $\tau_i$  is 0.

```
(b)
```

```
A <- c(62,60,63,59,64)

B <- c(65,67,73,65,66)

C <- c(69,66,71,67,67,68,62)

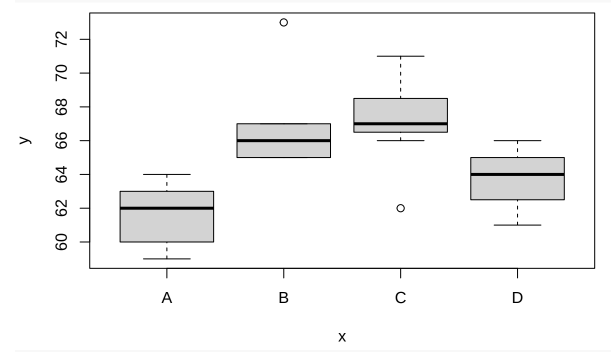
D <- c(66,62,65,61,64,65,63)

effect <- c(A,B,C,D)

diet <- c(rep('A', 5), rep('B', 5),rep('C', 7), rep('D', 7))

data <- data.frame(effect, diet)

plot(as.factor(data$diet), data$effect)
```



```
mean(A)
```

## [1] 61.6

mean(B)

## [1] 67.2

mean(C)

## [1] 67.14286

mean(D)

## [1] 63.71429

mean(data)

## Warning in mean.default(data): argument is not numeric or logical: returning NA

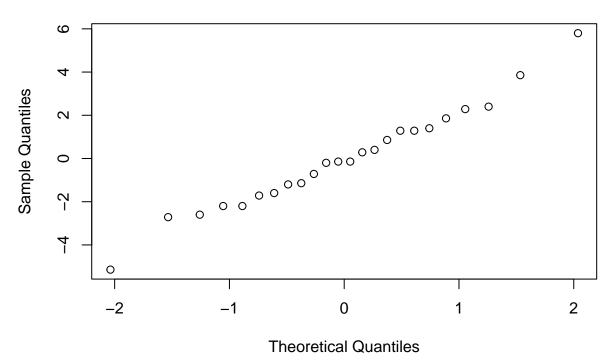
## [1] NA

There is no big difference between C and B, but among others theire is a difference.

(c)

```
#Group sample variances
sA \leftarrow var(A)
sB <- var(B)
sC <- var(C)
sD <- var(D)
sA;sB;sC;sD
## [1] 4.3
## [1] 11.2
## [1] 7.809524
## [1] 3.238095
# MSE
mse \leftarrow (4*sA+4*sB+6*sC+6*sD)/(24 - 4)
mse
## [1] 6.414286
Group sample variances are 4.3, 11.2, 7.809524, 3.238095. The MSE is 6.414286.
 (d)
mst <- 5/3*(mean(A) - mean(data\$effect))^2 + 5/3*(mean(B) - mean(data\$effect))^2
+7/3*(mean(C) - mean(data\$effect))^2 + 7/3*(mean(D) - mean(data\$effect))^2
## [1] 14.57143
mst
## [1] 27.33333
MST is 41.90476. It is way larger than MSE, so we can expect to reject the null that the diets are no different.
 (e)
anova(lm(data$effect ~ data$diet))
## Analysis of Variance Table
##
## Response: data$effect
              Df Sum Sq Mean Sq F value
## data$diet 3 125.71 41.905
                                    6.533 0.002937 **
## Residuals 20 128.29
                           6.414
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
Yes, since the p-value is 0.002937, we can safely reject the null and say that there is a difference for these four
diets.
 (f) Since the fitted values are the mean for each diet, we have
\text{data} miu.hat \leftarrow \text{c(rep(mean(A), 5), rep(mean(B), 5), rep(mean(C), 7), rep(mean(D), 7))}
data$residual <- data$effect - data$miu.hat</pre>
qqnorm(data$residual)
```

# Normal Q-Q Plot



the residuals appear to follow a normal distribution, since the points on the plot are pretty much along the diagonal.

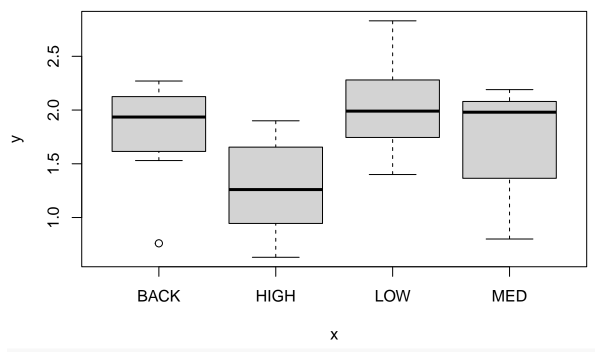
#### 4

(a)

```
library(dplyr)
```

```
##
## 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

zinc <- readRDS("zinc.RDS")
plot(as.factor(zinc$ZINC), zinc$DIVERSITY)</pre>
```



#### mean(zinc\$DIVERSITY)

```
## [1] 1.710312
```

```
# Method 1
meanZ \leftarrow zinc \%>\%
  group_by(ZINC) %>%
  summarize(mean(DIVERSITY))
aggregate(x= zinc$DIVERSITY,
         # Specify group indicator
         by = list(zinc$ZINC),
         # Specify function (i.e. mean)
         FUN = mean)
```

Group.1 ## 1 BACK 1.79750 ## 2 HIGH 1.28125 ## 3 LOW 2.03250 MED 1.73000

Yes, there appears to be a difference in biodiversity of rivers with different Zinc levels.

(b)

```
#Group sample variances
varZ <- zinc %>%
  group_by(ZINC) %>%
  summarize(var(DIVERSITY))
varZ
## # A tibble: 4 x 2
```

## ZINC `var(DIVERSITY)` ## <fct> <dbl> ## 1 BACK 0.235 ## 2 HIGH 0.208

```
## 3 LOW
                       0.198
## 4 MED
                       0.288
#count
numZ <- zinc %>%
  count(ZINC)
#MSE
mse1 <- (0.2354786+0.2081268+0.1980214 +0.2876286
)/4
mse1
## [1] 0.2323139
Group variances are BACK 0.2354786, HIGH 0.2081268, LOW 0.1980214, MED 0.2876286. MSE is 0.2323139.
 (c)
meanZ
## # A tibble: 4 x 2
     ZINC `mean(DIVERSITY)`
     <fct>
##
                        <dbl>
## 1 BACK
                          1.80
## 2 HIGH
                          1.28
## 3 LOW
                          2.03
## 4 MED
                          1.73
mst1 < -8/3*((1.79750 - mean(zinc$DIVERSITY))^2 + (1.28125 - mean(zinc$DIVERSITY))^2 +
                (2.03250 - mean(zinc\$DIVERSITY))^2 + (1.73000 - mean(zinc\$DIVERSITY))^2)
mst1
## [1] 0.7890365
fratio <- mst1 / mse1
fratio
## [1] 3.396425
#verify
anova(lm(zinc$DIVERSITY ~ zinc$ZINC))
## Analysis of Variance Table
##
## Response: zinc$DIVERSITY
             Df Sum Sq Mean Sq F value Pr(>F)
## zinc$ZINC 3 2.3671 0.78904 3.3964 0.03151 *
## Residuals 28 6.5048 0.23231
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Yes, I would have expected this F ratio – it is large such that we can reject the null that there is no difference
in biodiversity in zones with different Zinc levels. This is consistent with my observation in the plots and
early data exploration of the mean.
 (d)
zinc.level <- zinc$ZINC</pre>
fr <- c()
for(i in 1:1000){
  zinc.sim <- sample(zinc.level)</pre>
 fr <- c(fr, anova(lm(zinc$DIVERSITY ~ zinc.sim))$F[1] )</pre>
```

```
mean(fr >= 3.3964)
```

## [1] 0.027

With a p-value of 0.027, we can reject the null and say that Zinc levels do affect biodiversity.