STATS 205: Homework Assignment 3

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Solution to Problem 1

(i)

```
library(bootstrap); data(law)
t(law)
##
## LSAT 576.00 635.0 558.00 578.00 666.00 580.00 555 661.00 651.00 605.00
          3.39
                 3.3
                        2.81
                               3.03
                                      3.44
                                              3.07
## GPA
                                                     3
                                                         3.43
                                                                3.36
##
                    12
                           13
                                  14
            11
                                         15
## LSAT 653.00 575.00 545.00 572.00 594.00
## GPA
          3.12
                 2.74
                         2.76
                                2.88
theta.hat = cor(law$LSAT, law$GPA); theta.hat
## [1] 0.7763745
library(partitions)
n = 15
allCompositions = compositions(n, n);allCompositions[,1:5]
         [,1] [,2] [,3] [,4] [,5]
##
   [1,]
           15
                14
                     13
                           12
                                11
##
   [2,]
            0
                            3
## [3,]
                 0
                       0
                            0
                                 0
            0
## [4,]
            0
                 0
                       0
                            0
                                 0
##
  [5,]
            0
                 0
                      0
                            0
                                 0
  [6,]
##
            0
                 0
                       0
                            0
                                 0
## [7,]
            0
                 0
                      0
                                 0
                            0
## [8,]
            0
                      0
                 0
                            0
                                 0
## [9,]
            0
                 0
                      0
                            0
                                 0
## [10,]
            0
                 0
                      0
                            0
                                 0
## [11,]
            0
                 0
                      0
                            0
                                 0
## [12,]
            0
                 0
                      0
                            0
                                 0
## [13,]
            0
                       0
                            0
                                 0
## [14,]
            0
                 0
                      0
                                 0
                            0
## [15,]
allCompositions.sub = allCompositions[, sample(1:dim(allCompositions)[2], size=10000, replace=FALSE)]
draw.bootstrap.samples = function(df){
  n = dim(df)[1]
  ind = sample(n, replace = TRUE)
  cor.bootstrap.replicate = cor(df[ind, "LSAT"], df[ind, "GPA"])
  return(cor.bootstrap.replicate)
}
R = 10000
```

```
theta.hat.star = replicate(R, draw.bootstrap.samples(law))
# make a gaplot
library(ggplot2)
## Registered S3 methods overwritten by 'ggplot2':
##
     method
                    from
##
     [.quosures
                    rlang
##
     c.quosures
                    rlang
##
     print.quosures rlang
theta.hat.star.df = data.frame(theta.hat.star = theta.hat.star)
ggplot(theta.hat.star.df) +
  geom_density(aes(x = theta.hat.star, y = ..scaled..),
    fill = "lightblue") +
  geom_hline(yintercept=0, colour="white", size=1) +
  theme_bw() +
  ylab("density") +
  xlab(bquote(hat(theta))) +
  geom_vline(xintercept = theta.hat, col = "red")+
  scale_y_continuous(expand = c(0,0))
   1.00
   0.75
density
0.50
```

(ii)

0.25

0.00

```
sd(theta.hat.star)
```

0.50

θ

0.75

1.00

[1] 0.1326684

0.25

Solution to Problem 2

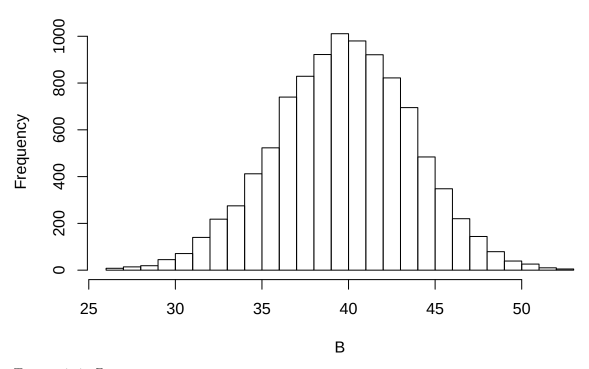
```
(i)
```

```
67 runs resulting in swallowing attempts
58 successful
9 failed

H_0: p = 0.6
H_A: p > 0.6
n = 67
successes = 58
pbar = successes / n; pbar

## [1] 0.8656716
p0 = 0.6; nsim = 10000
B = rbinom(nsim, size = n, prob = p0)
hist(B, breaks = 30)
```

Histogram of B



Test statistic Z:

$$Z_0 = \frac{B - 67(0.6)}{(67(0.6)(0.4))^{\frac{1}{2}}}$$

```
qnorm((1-0.05), mean = 0, sd = 1)
```

[1] 1.644854

Rejection region: $Z \ge z_{0.05} = 1.645$

Observed test statistic Z_o :

$$Z_o = \frac{58 - 67(0.6)}{(67(0.6)(0.4))^{\frac{1}{2}}} = 4.44$$

```
numerator = successes - (n * p0)
denominator = sqrt(n * p0 * (1.0 - p0))
Z.obs = numerator / denominator; Z.obs
```

[1] 4.438917

The large sample approximation value $Z_o = 2.5 > 1.645$ and thus we reject $H_0: p = 0.6$ in favor of p > 0.6 at the approximate $\alpha = 0.05$ level. Thus there is evidence that the success rate of swallowing attempts is greater than 0.6.

(ii)

Power is the probability of rejecting H_0 when H_A is true. We found that test reject H_0 is $Z \ge z_{0.05} = 1.645$. Therefore, if p = 0.7,

$$Z_o = \frac{58 - 67(0.6)}{(67(0.6)(0.4))^{\frac{1}{2}}} = 4.44$$

is no longer standard normal.

We have

$$Z_{o7} = \frac{58 - 67(0.7)}{(67(0.7)(0.3))^{\frac{1}{2}}} = 2.96$$

```
p1 = 0.7
numerator = successes - (n * p1)
denominator = sqrt(n * p1 * (1.0 - p1))
Z.obs.seven = numerator / denominator; Z.obs.seven
```

[1] 2.959211

$$Power = P(Z \ge 1.645 | p = 0.7)$$

$$= P_{p=0.7} \left(\frac{B - 67(0.6)}{(67(0.6)(0.4))^{\frac{1}{2}}} \ge 1.645 \right)$$

$$= P_{p=0.7}(B \ge 1.645(67(0.6)(0.4))^{\frac{1}{2}} + 67(0.6))$$

$$=P_{p=0.7}\bigg(\frac{B-67(0.7)}{(67(0.7)(0.3))^{\frac{1}{2}}}\geq \frac{1.645(67(0.6)(0.4))^{\frac{1}{2}}+67(0.6)-67(0.7)}{(67(0.7)(0.3))^{\frac{1}{2}}}\bigg)$$

```
triple_product = n * p0 * (1.0 - p0)
first_term = 1.645 * sqrt(triple_product)
second_term = n * p0
third_term = n * p1
bottom_term = n * p1 * (1.0 - p1)
```

```
p7_numerator = first_term + second_term - third_term
p7_denominator = sqrt(bottom_term)
Pp_7_zvalue = p7_numerator / p7_denominator; Pp_7_zvalue
```

[1] -0.02761144

$$P(Z^* \ge -0.0276) = 0.4890$$

```
power7 = pnorm(Pp_7_zvalue); power7
```

[1] 0.488986

The power of the test (the probability of rejecting H_0 when H_A is true) when p = 0.7 is 0.4890. If p = 0.8,

$$Power = P(Z \ge 1.645 | p = 0.8)$$

$$= P_{p=0.8} \left(\frac{B - 67(0.8)}{(67(0.8)(0.2))^{\frac{1}{2}}} \ge \frac{1.645(67(0.6)(0.4))^{\frac{1}{2}} + 67(0.6) - 67(0.8)}{(67(0.8)(0.2))^{\frac{1}{2}}} \right)$$

```
p2 = 0.8
triple_product = n * p0 * (1.0 - p0)
first_term = 1.645 * sqrt(triple_product)
second_term = n * p0
third_term = n * p2
bottom_term = n * p2 * (1.0 - p2)
p8_numerator = first_term + second_term - third_term
p8_denominator = sqrt(bottom_term)
Pp_8_zvalue = p8_numerator / p8_denominator; Pp_8_zvalue
```

[1] -2.077971

$$P(Z^* > -2.078) = 0.01886$$

```
power8 = pnorm(Pp_8_zvalue); power8
```

[1] 0.01885601

The power of the test (the probability of rejecting H_0 when H_A is true) when p = 0.8 is 0.01886.

Solution to Problem 3

Summary: Estimate for $\hat{p} = 0.8615$ and estimate for standard deviation of $\hat{p} = 0.04284$.

Estimate for p using binomial confidence interval, binom.confint():

```
library(binom)
binom.confint(x=56, n=65, conf.level=.95, methods = "asymptotic")
```

```
## method x n mean lower upper ## 1 asymptotic 56 65 0.8615385 0.7775744 0.9455025
```

```
\hat{p} = (0.7776, 0.9455)
```

```
Estimate for p using 1-sample proportions test without continuity correction, prop.test():
```

```
prop.test(x=56, n=65, p = 0.6, conf.level=0.95, alternative = c("greater"))
```

```
##
## 1-sample proportions test with continuity correction
##
## data: 56 out of 65, null probability 0.6
## X-squared = 17.452, df = 1, p-value = 1.473e-05
## alternative hypothesis: true p is greater than 0.6
## 95 percent confidence interval:
## 0.7676875 1.0000000
## sample estimates:
## p
## 0.8615385
```

$$p = 0.8615$$

Estimate for p using Exact binomial test:

```
binom.test(x=56, n=65, p = 0.6, alternative = c("greater"), conf.level = 0.95)
```

```
##
## Exact binomial test
##
## data: 56 and 65
## number of successes = 56, number of trials = 65, p-value =
## 4.096e-06
## alternative hypothesis: true probability of success is greater than 0.6
## 95 percent confidence interval:
## 0.7708174 1.0000000
## sample estimates:
## probability of success
## 0.8615385
```

Standard error of \hat{p} is:

$$\sqrt{\frac{p(1-p)}{n}}$$

$$= \sqrt{\frac{(0.6)(0.4)}{65}}$$

$$= 0.06076$$

```
p = 0.6
n = 65
numerator = p * (1 - p)
denominator = n
answer = sqrt(numerator/denominator); answer
```

[1] 0.06076436

and estimate is:

$$\sqrt{\frac{\hat{p}(1-\hat{p})}{n}}$$

$$= \sqrt{\frac{(0.8615)(1-0.8615)}{65}}$$

$$= 0.04284$$

```
p.hat = 0.8615
n = 65
numerator = p.hat * (1 - p.hat)
denominator = n
answer = sqrt(numerator/denominator); answer
## [1] 0.04284458
```

Solution to Problem 4

```
binom.confint(x = 56, n = 65, conf.level = 0.96, methods = "all")
##
             method x n
                               mean
                                        lower
                                                   upper
## 1
     agresti-coull 56 65 0.8615385 0.7488973 0.9301180
## 2
         asymptotic 56 65 0.8615385 0.7735567 0.9495202
## 3
              bayes 56 65 0.8560606 0.7655984 0.9375798
## 4
            cloglog 56 65 0.8615385 0.7439982 0.9276404
## 5
              exact 56 65 0.8615385 0.7480632 0.9371740
## 6
              logit 56 65 0.8615385 0.7484912 0.9286194
## 7
             probit 56 65 0.8615385 0.7545841 0.9312980
## 8
            profile 56 65 0.8615385 0.7589798 0.9334925
## 9
                lrt 56 65 0.8615385 0.7589836 0.9335307
## 10
          prop.test 56 65 0.8615385 0.7483484 0.9308913
## 11
             wilson 56 65 0.8615385 0.7514483 0.9275670
Here are the rows relevant to our problem:
##
             method x n
                               mean
                                        lower
                                                   upper
## 1
     agresti-coull 56 65 0.8615385 0.7488973 0.9301180
## 2
         asymptotic 56 65 0.8615385 0.7735567 0.9495202
              exact 56 65 0.8615385 0.7480632 0.9371740
## 5
## 11
             wilson 56 65 0.8615385 0.7514483 0.9275670
```

where asymptotic is Laplace-Wald, agresti-coull is Agresti-Coull, exact is Clopper-Pearson, and wilson is Wilson.

It looks like agresti-coull and exact are fairly similar in terms of the location of the interval, while asymptotic is skewed towards the "right" side of the intervals, and wilson has the smallest range.

Solution to Problem 5

```
plant_vector <- c(926, 288, 293, 104)
expected = c(9/16, 3/16, 3/16, 1/16)
```

```
# goodness-of-fit test
res <- chisq.test(x = plant_vector, p = expected); res
##
   Chi-squared test for given probabilities
##
##
## data: plant_vector
## X-squared = 1.4687, df = 3, p-value = 0.6895
res$expected
## [1] 906.1875 302.0625 302.0625 100.6875
res$observed
## [1] 926 288 293 104
```

The **p-value** of the test is p = 0.6895, which is more than the significance level $\alpha = 0.05$. We cannot conclude that the data does not support Mendelian theory at the $\alpha = 0.05$ level.

Solution to Problem 6

```
df = data.frame(Low = c(0, 2), High = c(5, 1))
rownames(df) = c("Multiple attack", "Primary attack"); df
                   Low High
## Multiple attack
                     0
                          5
## Primary attack
                     2
fisher.test(df, alternative = "less")
##
   Fisher's Exact Test for Count Data
##
##
## data: df
## p-value = 0.1071
## alternative hypothesis: true odds ratio is less than 1
## 95 percent confidence interval:
## 0.00000 1.75251
## sample estimates:
## odds ratio
##
```

There is not enough evidence that the unknown probability that a multiple-attack patient will have low reactivity is less than the unknown probability that a primary-attack patient will have low reactivity at the $\alpha = 0.05$ level. The P-value achieved by these data if we use Fisher's exact test of H_0 against the alternative $p_1 < p_2$ is 0.1071.

```
df = data.frame(Low = c(0, 2), High = c(5, 1))
rownames(df) = c("Multiple attack", "Primary attack"); df
```

```
##
                    Low High
## Multiple attack
                      0
                           5
## Primary attack
                      2
```

fisher.test(df, alternative = "two.sided") ## ## Fisher's Exact Test for Count Data ## ## data: df ## p-value = 0.1071 ## alternative hypothesis: true odds ratio is not equal to 1 ## 95 percent confidence interval: ## 0.00000 2.75184 ## sample estimates: ## odds ratio

The P-value achieved when testing the null against a two-sided hypothesis is the same, 0.1071, and therefore, there is still not enough evidence that the unknown probability that a multiple-attack patient will have low reactivity is not equal to the unknown probability that a primary-attack patient will have low reactivity.

Solution to Problem 7

Since we want to find out if having tonsils reduces the rate of Hodgkin's, we want to see if there is a higher rate of tonsillectomy among Hodgkin's cases than among non-Hodgkin's cases. Therefore, we will test the null hypothesis that there is no difference in the rate of tonsillectomy between the two populations of Hodgkin's and non-Hodgkin's cases against the alternative hypothesis that there is higher rate of tonsillectomy among Hodgkin's cases.

$$H_0: p_1 = p_2$$

$$H_A: p_1 > p_2$$

```
x = c(67, 43)
n = c(101, 107)
prop.test(x, n, alternative = c("greater"), conf.level = 0.99)
##
   2-sample test for equality of proportions with continuity
##
##
   correction
##
## data: x out of n
## X-squared = 13.229, df = 1, p-value = 0.0001379
## alternative hypothesis: greater
## 99 percent confidence interval:
  0.09655707 1.00000000
## sample estimates:
##
      prop 1
                prop 2
## 0.6633663 0.4018692
```

The 99% confidence interval for difference between rate of tonsillectomy in Hodgkin's cases and rate of tonsillectomy in non-Hodgkin's cases is between 9.66% and 100%.

The P-value achieved by these data if we use a 2-sample test for equality of proportions with continuity correction is 0.0001379. There is $strong\ evidence$ at the $\alpha=0.05$ level that there is higher rate of tonsillectomy among Hodgkin's cases.

Solution to Problem 8

```
elisa_matrix <- matrix(c(82, 13, 6, 0), nrow=2, ncol=2, byrow=TRUE, dimnames = list("ABC-ELISA" = c("+"
##
            Standard ELISA
## ABC-ELISA + -
##
           + 82 13
##
           - 6 0
# prop.test(x = elisa\_matrix) < -- this produced a p value not less than 0.005, so I may be using it wro
mcnemar.test(x = elisa_matrix)
##
##
   McNemar's Chi-squared test with continuity correction
##
## data: elisa_matrix
## McNemar's chi-squared = 1.8947, df = 1, p-value = 0.1687
```

The *P*-value for McNemar's test is 0.1687. What we can conclude concerning the hypothesis of equal proportions positive for both ELISA tests is that equal proportions may not be positive for both ELISA tests.

Solution to Problem 9

```
ptsd_matrix <- matrix(c(29, 7, 21, 30), nrow = 2, ncol = 2, byrow = TRUE, dimnames = list(c("Yes", "No"
##
       Battered women
## Yes
                   29
## No
                   21
##
       Maritally distressed women who had not experienced battering
## Yes
## No
                                                                  30
prop.test(ptsd_matrix, correct = F, alternative = "two.sided")
##
   2-sample test for equality of proportions without continuity
##
##
   correction
##
## data: ptsd_matrix
## X-squared = 13.389, df = 1, p-value = 0.0002531
## alternative hypothesis: two.sided
## 95 percent confidence interval:
## 0.2068191 0.5807626
## sample estimates:
      prop 1
                prop 2
## 0.8055556 0.4117647
```

The P-value for 2-sample test for equality of proportions without continuity correction is 0.0002531, which is significant at the $\alpha = 0.05$ level. There is strong evidence that there is a significant difference in the PTSD rates for battered women versus maritally distressed women (who had not experienced battering).