Cover Sheet

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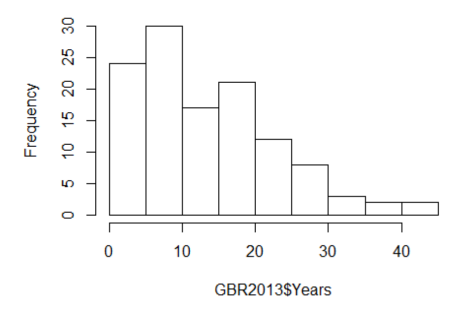
Name	Student Number	Contribution (%)
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Vinith Gunawardena	19620392	33.33

Part 1

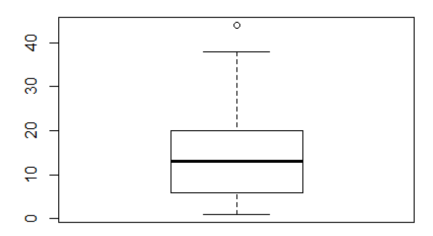
Part 1(A)

```
GBR2013 =read.csv(file.choose())
GBR2017 =read.csv(file.choose())
mean.Years2013 =mean(GBR2013$Years)
summary(GBR2013$Years, na.rm=TRUE)
##
      Min. 1st Qu. Median Mean 3rd Qu.
                                           Max.
                    13.00
##
      1.00
             6.00
                           14.43
                                    20.00 44.00
sd(GBR2013$Years, na.rm=TRUE)
## [1] 9.670479
hist(GBR2013$Years)
```

Histogram of GBR2013\$Years



boxplot(GBR2013\$Years)



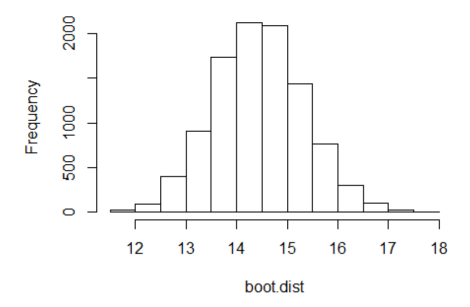
#The data is right-skewed when we observe this when we examine this in the histogram. We can also examine this type of spread when we look at the box plot. We have also identified one outlier; this and the spread has had some effect on the calculation of the mean(13, 14.43). The standard deviation is relatively large(9.67), and this can also be seen with the histogram and the boxplot. We can see from the box plot, that the data is also spread a little unevenly, more so towards the end of the plot[which confirms the spread we see in the histogram].

Part 1(B)

```
mean.Years2013 =mean(GBR2013$Years)
mean.Years2013
## [1] 14.42857

b =10000
boot.dist =rep(NA, b)
for (i in 1:b){
boot.sample =sample(119, replace=TRUE)
boot.dist[i] =mean(GBR2013$Years[boot.sample])
}
hist(boot.dist)
```

Histogram of boot.dist



```
SE.year2013 =sd(boot.dist)
SE.year2013
## [1] 0.8871243

CI.99.Bootstrap.Upper =mean.Years2013 +2.576*(SE.year2013)
CI.99.Bootstrap.Upper
## [1] 16.7138

CI.99.Bootstrap.Lower =mean.Years2013 -2.576*(SE.year2013)
CI.99.Bootstrap.Lower
## [1] 12.14334

#The 99% CI is between 12.14 and 16.71. We are 99% sure that population parameter for the mean years is between this interval. [Note: I know that each bootstrap distribution will change the result of the Confidence interval, the one uses the standard error from one of the simulations]
```

<u>Part 1(c)</u>

```
\#CLT states that SE using the formula SE = s / sqrt(n) applies when the size
is greater or equal to 30. This would suggest we can apply CLT to this case
since n = 119.
mean.Years2013 =mean(GBR2013$Years)
mean. Years 2013
## [1] 14.42857
sd.Years2013 =sd(GBR2013$Years)
sd.Years2013
## [1] 9.670479
SE.CLT.Years2013 =sd.Years2013 /sqrt(119)
SE.CLT.Years2013
## [1] 0.8864913
tvalue.Years2013 = qt(0.995, 119)
tvalue.Years2013
## [1] 2.617776
CI.99t.Upper =mean.Years2013 +tvalue.Years2013*SE.CLT.Years2013
CI.99t.Upper
## [1] 16.74921
```

```
CI.99t.Lower =mean.Years2013 -tvalue.Years2013*SE.CLT.Years2013
CI.99t.Lower
## [1] 12.10794
```

#We notice that our SE is almost identical to using the bootstrap method; We can infer that because the SE is almost identical, the CI is going to be very similar. We know that when we are dealing with means, it follows a t-distribution; so our value for z-score would be replaced by our t-value. The 99% CI for both of them are almost identical. CI using Bootstrapping method = 12.14 and 16.71. CI using CLT method was between 12.10762 and 16.74953. This gives us an indication that the formula methodology that we are using is a valid representation of a simulation; and that it can be used as a suitable alternative, if it meets the conditions for CLT.

Part 2

Part 2(A)

```
#(a) Hypothesis Test
##0: p = 0.7
##A: p < 0.7

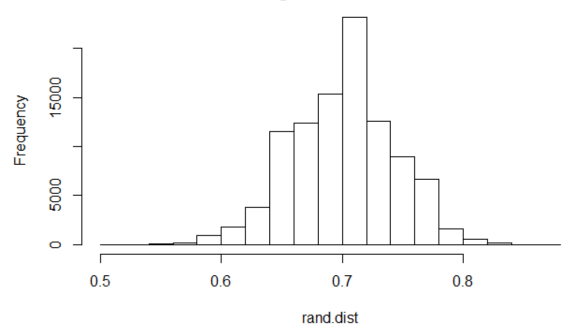
#Note: P is representing the population parameter(for proportions).
#Left-tailed test.
----
#Part (i)
----

#two values are missing and therefore we are using n = 117

#With and without Continuity Correction.
prop.test(72, 117, conf.level=0.95, correct =FALSE, alternative ="less", 0.7)
prop.test(72, 117, conf.level=0.95, correct =TRUE, alternative ="less", 0.7)</pre>
```

```
## 1-sample proportions test with continuity correction
## data: 72 out of 117, null probability 0.7
## X-squared = 3.5963, df = 1, p-value = 0.02895
## alternative hypothesis: true p is less than 0.7
## 95 percent confidence interval:
## 0.000000 0.1668754
## sample estimates:
##
     р
## 0.6153846
#p-value without correction = 0.0229
#p-value with correction = 0.02895
#Doing some extra-tests to see if the prop.test is consistent with CLT
method.
#The test is also done with the intention of confirming our SE through
randomization.
se.ztest = sqrt(0.7*0.3/117)
se.ztest
prop.Optimistic =72/117
ztest = (prop.Optimistic -0.7) / sqrt(0.7*0.3/117)
ztest
pnorm(ztest, 0 , 1)
\#p-value Z-test = 0.0229
#Part (ii)
mean(GBR2013$Optimistic >=6, na.rm=TRUE)
#Method 1 [Centered at the null hypothesis]
-----
r = 100000
n = 117
rand.dist = rep(0,r)
for (a in 1:r) {
s =sample(c(TRUE, FALSE), n, replace =TRUE, prob =c(0.7,0.3))
rand.dist[a] =mean(s ==TRUE)
}
```

```
hist(rand.dist)
sd(rand.dist)
pvalue.randomization1 =mean(rand.dist <=prop.Optimistic )
pvalue.randomization1</pre>
```



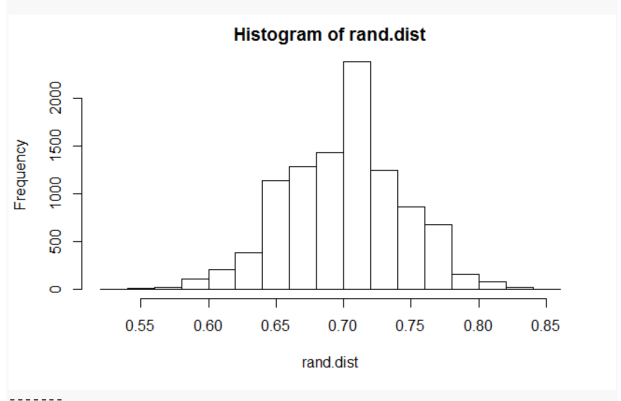
```
#pvalue1 = 0.031

#Method 2 [Done for extra Confirmation]

b =10000
rand.dist =rep(NA, b)
for (i in 1:b) {
    rand.dist[i] =(rbinom(1, size =117, prob =0.7)/117)
}
hist(rand.dist)
sd(rand.dist)
pvalue.randomization2 =mean(rand.dist <=prop.Optimistic )
pvalue.randomization2

#pvalue2 = 0.0302</pre>
```

#Two randomization methods with the intention of observing consistency. Standard error is similar to the one produced through the formula.



#Part (iii)

#All three tests give p-values which are around 0.03 which is lower than the alpha = 0.05. We notice some difference when we use the CLT with the z-score; but this is corrected when continuity is on. We would reject the Null hypothesis with all of the tests [randomization and prop.test]. There is evidence that the mean proportion for those optimistic is less than 0.7.

Part 2(B)

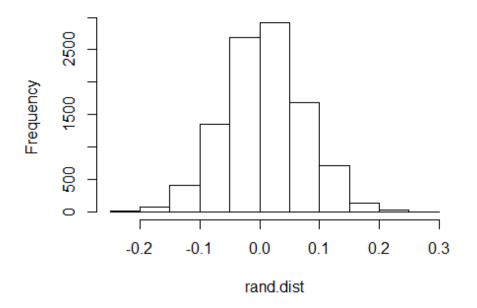
```
mean.2017.6 =mean(GBR2017$Optimistic >=6, na.rm=TRUE)
difference.mean =mean.2013.6 -mean.2017.6
difference.mean
## [1] 0.02398677
#Part (i)
----
#Note: There are two missing values for both 2013 and 2017, and we are using
n(2013) = 117 and n(2017) = 93
#H0 : p(2013) - p(2017) = 0
#HA: p(2013) - p(2017) != 0
#p is representative of the population parameter [proportions]
#Prop.test [With Correct False and True]
prop.test(c(72, 55), c(117, 93), correct =FALSE)
prop.test(c(72, 55), c(117, 93), correct =TRUE)
##
##
  2-sample test for equality of proportions with continuity
## correction
##
## data: c(72, 55) out of c(117, 93)
## X-squared = 0.044557, df = 1, p-value = 0.8328
## alternative hypothesis: two.sided
## 95 percent confidence interval:
## -0.1189019 0.1668754
## sample estimates:
##
      prop 1
               prop 2
## 0.6153846 0.5913978
#pvalue[correction] = 0.8328
#pvalue[without correction] = 0.724
#We notice that the correction for continuity has had a big influence on the
p-value; change it from 0.724 to 0.8328.
#Prop-test has its own unique method, and I had thought to add a third(z-
test) to make some clarifications. The test is valuable when we want to
examine whether or not our SE in our randomization is the same value as the
formula. The formula was very useful in our investigation, because it has
given us some hints about how to produce our randomization. The formula gave
us some indication that it was imperative that we need to split the values in
```

```
an uneven fashion (117, 93).
table(GBR2017$Optimistic)
##
## 1 2 3 4 5 6 7 8 9 10
## 11 3 3 6 15 9 18 13 3 12
table(GBR2013$Optimistic)
##
   1 2 3 4 5 6 7 8 9 10
##
##
  6 3 9 9 18 10 12 20 2 28
pooledtable =table(GBR2017$Optimistic) +table(GBR2013$Optimistic)
pooledtable
##
## 1 2 3 4 5 6 7 8 9 10
## 17 6 12 15 33 19 30 33 5 40
\#Total\ n = 210\ (missing\ two\ values\ in\ each\ year)
pooled.proportion =127/210
pooled.proportion
## [1] 0.6047619
p.1minusp =(pooled.proportion)*(1-pooled.proportion)
p.1minusp
## [1] 0.2390249
se.pooled.proprotion =sqrt(p.1minusp/119 +p.1minusp/95 )
se.pooled.proprotion
## [1] 0.06726563
zscore.pooled =(mean.2013.6 -mean.2017.6) /se.pooled.proprotion
zscore.pooled
## [1] 0.3565977
pvalue =2*(1 -pnorm(zscore.pooled, 0, 1))
pvalue
## [1] 0.721393
#Part (ii)
_ _ _ _ _
```

Randomization #The formula for the SE splits the sizes of the sample into two separate calculations I would expect that splitting the samples in the same manner according to the formula will give a similar Standard error. We have two methods that are shown below which will demonstrate this phenomenon. Replacement is false, and the values has been successfully split. b =10000 rand.dist =rep(NA, b) for (i in 1:b) { unifiedSample=c(GBR2013\$Optimistic,GBR2017\$Optimistic) rand.samp=(sample(210, 117, replace =FALSE)) group.size117 =unifiedSample[rand.samp] group.size93 =unifiedSample[-rand.samp] rand.dist[i] =mean(group.size117 >=6, na.rm=TRUE) -mean(group.size93 >=6, na.rm=TRUE)

Histogram of rand.dist

hist(rand.dist)



```
sd(rand.dist)
## [1] 0.06700783
```

```
prop.difference =mean.2013.6 -mean.2017.6

pvaluernd1 =2*mean(rand.dist >=prop.difference)
pvaluernd1

## [1] 0.8266

#We notice that the prop.test function has slightly different p-value when continuity is on. Our p-value using randomization is around 0.82. Continuity is more similar to the randomization, and it seems as if the way it calculates the value is a better representation than the general CLT method.This is very similar to our p-value when we randomized 0.8266. In either case; when significance level alpha is set to 0.05, since 0.82 > 0.05, we do not reject the null hypothesis, and there is no evidence that the optimism was different between the years 2013-2017.
```

Part 3

Part 3(a)

```
GBR2013$QualityOfLife
GBR2013$WantToProtect

H0: p(correlation) = 0
HA : p (correlation) > 0

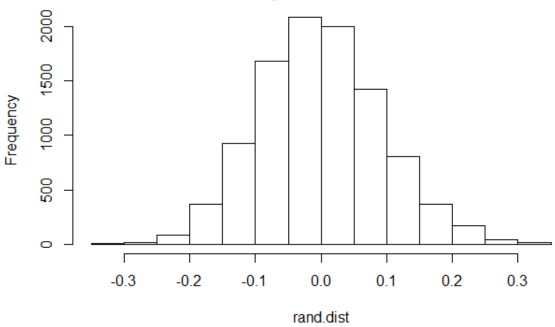
#Note: Both values are representative of the population
parameter[correlation].

Part 3(b)

cor.Value.2013 =cor(GBR2013$QualityOfLife,GBR2013$WantToProtect,
use="complete.obs")
cor.Value.2013

#We use in this section; replacement = false.

b =10000
rand.dist =rep(NA, b)
for (i in 1:b) {
    shuffle =sample(GBR2013$QualityOfLife)
```



Part 3(c)

```
Alpha = 0.10
```

#0ur p-value is 0.185, and this is larger than alpha = 0.10. We do not reject null hypothesis. This suggests that there is no evidence to support the claim that that the "The GBR contributes to my quality of life" and "I would like to do more help to protect the GBR" are positively correlated.

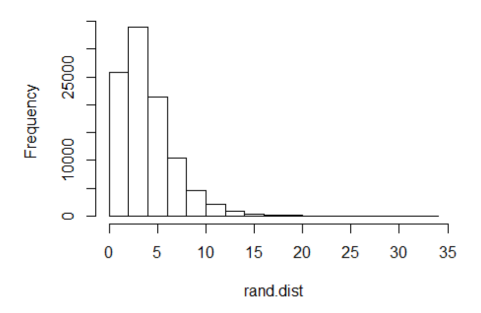
Part 4

Part 4 (a)

```
#Table:
_ _ _ _ _ _
#Note some values are missing n(2013) = 118 and n(2017) = 92
table.2013 =table(GBR2013$ClimateChangeView)
table.2013
##
## 1 2 3 4 5
## 59 21 27 5 6
table.2017 =table(GBR2017$ClimateChangeView)
table.2017
##
## 1 2 3 4 5
## 58 4 22 4 4
#Note: We must conduct randomization because three values are below 5.
#Prop Table[manual work]:
proportion.table2013 =prop.table(table(GBR2013$ClimateChangeView))
proportion.table2013
##
## 0.50000000.17796610 0.22881356 0.04237288 0.05084746
proportion.table2017 =prop.table(table(GBR2017$ClimateChangeView))
proportion.table2017
##
## 0.63043478 0.04347826 0.23913043 0.04347826 0.04347826
expectedCounts =proportion.table2013 *92
```

```
chisq.test =sum(((table.2017 -expectedCounts)^2 )/expectedCounts)
chisq.test
\#Chisq.test = 12.624
#Chi-squared goodness for fit
Note: In this situation we are testing if the proportions in 2017 are
different from the year 2013's proportions.
H0: p1 = 0.5, p2=0.178, p3=0.2288 p4=0.0424 p5=0.0508 2017 proportions are
not different from 2013's proportions [assume proportions will stay the same]
Ha: That there is some probability that is different.
#Note: We must conduct randomization because three values are below 5.
chisq.test( table(GBR2017$ClimateChangeView), p =c(0.5,0.17796610,0.22881356,
0.04237288, 0.05084746))
#pvalue[without randomization(function)] = 0.01567. We did this just for some
additional insight into the change between the values(if there are any).
#Simulated Randomization using Chis-quared function
chisq.test( table(GBR2017$ClimateChangeView), p =c(0.5,0.17796610,0.22881356,
0.04237288, 0.05084746), simulate.p.value = T, B=100000)
##
## Chi-squared test for given probabilities with simulated p-value
## (based on 1e+05 replicates)
## data: table(GBR2017$ClimateChangeView)
## X-squared = 12.624, df = NA, p-value = 0.01492
##pvalue[with randomization] = 0.01327
#Simulated Randomization; manual[three values are less than 5]
r =table(GBR2017$ClimateChangeView)
e = c(47.5, 16.906780, 21.737288, 4.025424, 4.8300508)
b = 100000
rand.dist =rep(NA, b)
for (i in 1:b) {
```

```
r =rmultinom(1,92, c(0.5,0.17796610,0.22881356, 0.04237288, 0.05084746))
rand.dist[i] =sum((r-e)^2/e)
}
hist(rand.dist)
```



```
mean(rand.dist >=12.624)
```

[1] 0.01293

#pvalue[without randomization(manual)] = 0.01308

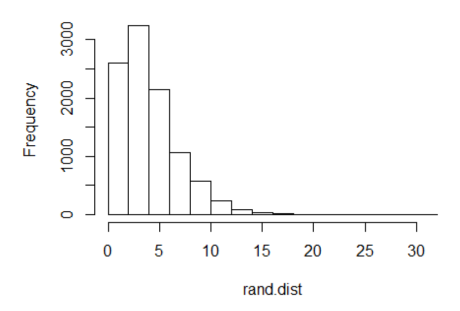
#P-Values for both types of tests(randomization and without) are around 0.013-0.015. It hadn't changed the p-value too greatly, but for formal reasons, we will stick with the value that was given from randomization methods. There wasn't much difference between the simulations, in comparison to the general test; this is probably because the values are very close to 5.We need to use randomization in this case, and this has given us a value of p=0.01327, which is lower than alpha = 0.05. This would suggest that there is evidence against the null hypothesis, and for the alternative, however; we would not be able to test this without confirmation of the significance level.

. .

Part 4(b)

```
#HO: Attitude on climate change is not associated with the year
#Ha : Attitude on climate change is associated with the year
degreeOfFreedom = (2-1)*(5-1)
F =matrix(c(59,21,27,5,6,58,4,22,4,4), nrow =2, ncol=5, byrow=TRUE)
n = sum(F)
E =rowSums(F) %o%colSums(F) /n
#Randomiztion Simulation using ChiSquared function
chisq.test(F, p = E, simulate.p.value = T, B =10000)
##
  Pearson's Chi-squared test with simulated p-value (based on 10000
##
## replicates)
##
## data: F
## X-squared = 9.5167, df = NA, p-value = 0.0483
#pvalue(randomization[function]) = 0.0447
#Simulation using randomization manually
rowSums(F)
## [1] -118 -92
colSums(F)
## [1] 117 25 49 9 10
V = sum((F-E)^2/E)
## [1] 9.516694
pval.merge =pchisq(V, df=4, lower.tail=FALSE)
pf = c(rep(1,59+58), rep(2,21+4), rep(3,27+22), rep(4,5+4), rep(5,6+4))
gr = c(rep(1,59+21+27+5+6), rep(2,58+4+22+4+4))
b = 10000
rand.dist = rep(NA, b)
for (i in 1:b) {
r =table(sample(gr), pf)
rand.dist[i] = sum((r-E)^2/E)
```

```
hist(rand.dist)
```



```
mean(rand.dist >=V)
## [1] 0.0463
#pvalue(randomization[manual]) = 0.0463
#The p-value is 0.0447[chis-squared test function] and 0.0463 for simulation(randomization). This is a relatively low p-value, and the outcome of the test would be determined about our choice of Alpha.
```

Part 4(c)

The p-values are little bit different, part(a) = 0.013, and part(b) 0.045. The difference isn't massive because the tests are very similar. However, the difference is significant enough that could cause a different outcome of a hypothesis test depending on alpha(alpha = 0.01 for example).

The goodness of fit tests whether or not there is a difference in proportions from a set of values to the other. A goodness for association(independence) determines whether or not the responses are dependent on the year. The test for association will give us insight about whether there is some property about the year which causes a change in a response or value.

It is a little difficult to determine which test is more suitable, because the question can be interpreted in many ways. From our research into the question, we found that there are certain statisticians who prefer one of the methods over the other. We had identified a similar question from the text book [Question 7.45] where part B's method was employed. To determine which test to choose is dependent on the question being asked; the question itself can have a significant impact on the way we manipulate the data and what tests to employ.

This particular question can be interpreted in many ways, firstly, on one hand it can be interpreted to see if there was a difference that has occurred between the years, and if it is interpreted in this way, then part (a) is more suitable, since we don't require a second test in the opposite direction. However, if we are interpreting the question in a different way, and we are trying to determine if there is some attribute or property about the year (dependent on time or year)which is causing this change, then the methodology in part (b) is more suitable. The good thing about the test for independence is that it will give us an understanding of what to expect, if the responses are dependent upon time (or year) then we will know that the responses will vary between the years(and therefore expect a difference, because the response is dependent on time). This test gives us more insight into answering the question; because it has the capacity to answer both interpretations of the question.