# Chapter 9: Interactive Notebook for Instructors

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## Load functions to compute p-value

```
library(ggplot2)
p_rtail = function(sampdist,tstat)
{
  temp = density(sampdist)
  df = data.frame(temp$x, temp$y)
  formula1 = df$temp.x<tstat
  df1 = df[formula1,]
  plot(df, col = "red", type = "h")
  points(df1, col = "green", type = "h")
  pvalue = length(sampdist[sampdist>tstat])/(length(sampdist))
  return(pvalue)
  }

p_ltail = function(sampdist,tstat)
  {
  temp = density(sampdist)
   df = data.frame(temp$x, temp$y)
  formula1 = df$temp.x>tstat
```

```
df1 = df[formula1,]
  plot(df, col = "red", type = "h")
  points(df1, col = "green", type = "h")
  pvalue = length(sampdist[sampdist<tstat])/(length(sampdist))</pre>
  return(pvalue)
p_2tail = function(sampdist,tstat)
hyp = mean(sampdist)
cutoff1 = hyp - abs(tstat-hyp)
  cutoff2 = hyp + abs(tstat-hyp)
  temp = density(sampdist)
  df = data.frame(temp$x, temp$y)
  formula1 = df$temp.x<cutoff1 | df$temp.x>cutoff2
  df1 = df[formula1,]
  plot(df, col = "green", type = "h")
  points(df1, col = "red", type = "h")
  pvalue = length(sampdist[sampdist<cutoff1 | sampdist>cutoff2])/(length(sampdist))
  return(pvalue)
  }
```

## **Basic Concepts**

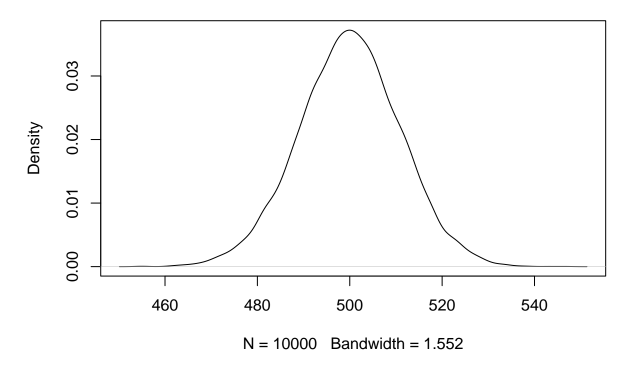
In this lesson we will explore Nonparametric statistical tests, which rely on no or few assumptions about the population distribution from which the sample was drawn.

How is this done? Remember that we need a sampling distribution to conduct a statistical test. Creating a sampling distribution requires us to specify the population distribution. Therefore, if you cannot specify a population distribution, how can you then create the sampling distribution and conduct the test? In certain situations, if you are creative, you can. Let us consider an example.

### Median Test

In a previous chapter, we conducted a test where the null hypothesis was that the median GMAT score was 510. Let us replicate that code here.

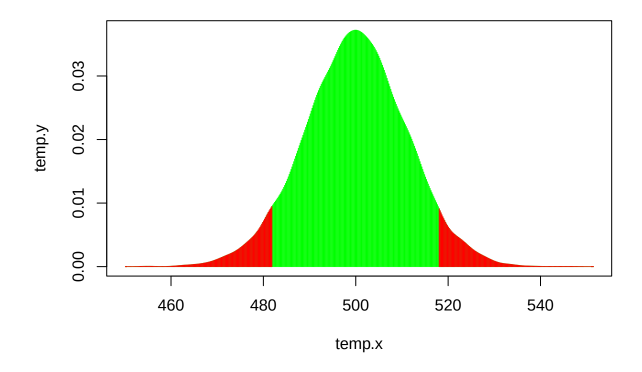
```
admission <- read.csv("../../data/admission.csv", stringsAsFactors=TRUE)
set.seed(87654321)
sampsize = nrow(admission)
f1 = function(){
   s1 = rnorm(sampsize, mean = 500, sd = sd(admission$GMAT))
   return(median(s1))
}
sampdist = replicate(10000, f1())
plot(density(sampdist))</pre>
```



```
tstat = median(admission$GMAT)
tstat
```

### ## [1] 482

```
gap = abs(tstat-500)
cutoff1 = 500 - gap
cutoff2 = 500 + gap
temp = density(sampdist)
df = data.frame(temp$x, temp$y)
formula1 = df$temp.x<cutoff1 | df$temp.x>cutoff2
df1 = df[formula1,]
plot(df, col = "green", type = "h")
points(df1, col = "red", type = "h")
```



pvalue = length(sampdist[sampdist<cutoff1 | sampdist>cutoff2])/(length(sampdist))
print(pvalue)

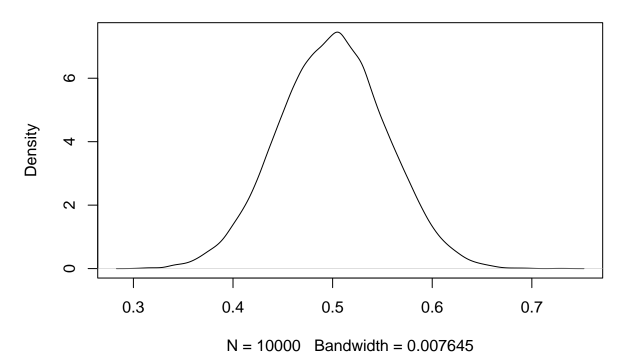
Can we conduct this test without having to assume that the population distribution for GMAT scores is normal?

We know that, regardless of what the population distribution is, by definition, half of the values should lie below the median and half should lie above the median. This is true for a normal distribution and also true for any other distribution. We will use this basic logic to design our test.

Parametric Population Description	Nonparametric Population Description
Normal Distribution	Population consists of positives (1) which are above the median and negatives (0) which are below the median with equal probabilities

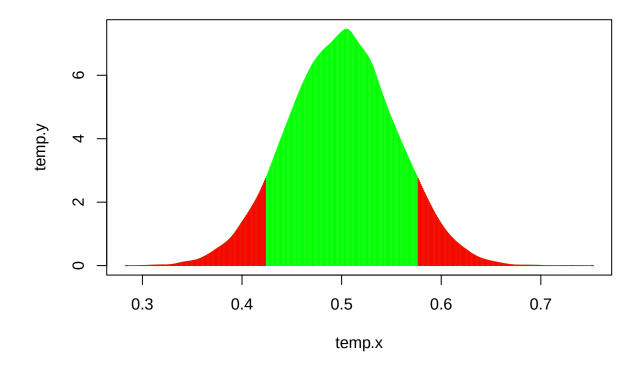
Now we can conduct the test.

```
set.seed(87654321)
sampsize = nrow(admission)
f1 = function(){
   s1 = sample(x = c(0,1), size = sampsize, replace = T, prob = c(.5,.5))
   return(sum(s1)/sampsize)
}
sampdist = replicate(10000, f1())
plot(density(sampdist))
```



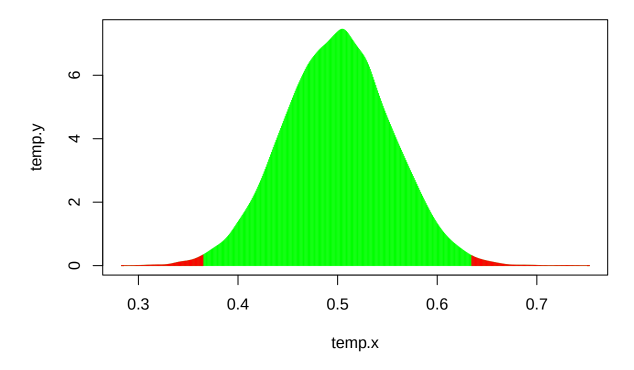
```
tstat = length(admission$GMAT[admission$GMAT>500])/(sampsize)

gap = abs(tstat-.5)
cutoff1 = .5 - gap
cutoff2 = .5 + gap
temp = density(sampdist)
df = data.frame(temp$x, temp$y)
formula1 = df$temp.x<cutoff1 | df$temp.x>cutoff2
df1 = df[formula1,]
plot(df, col = "green", type = "h")
points(df1, col = "red", type = "h")
```



pvalue = length(sampdist[sampdist<cutoff1 | sampdist>cutoff2])/(length(sampdist))
print(pvalue)

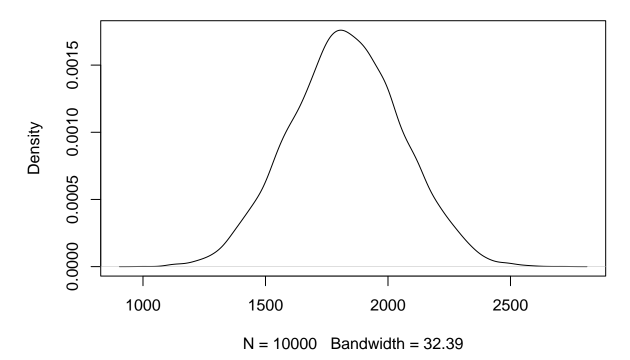
tstat = length(admission\$GMAT[admission\$GMAT>510])/(sampsize)
p\_2tail(sampdist,tstat)



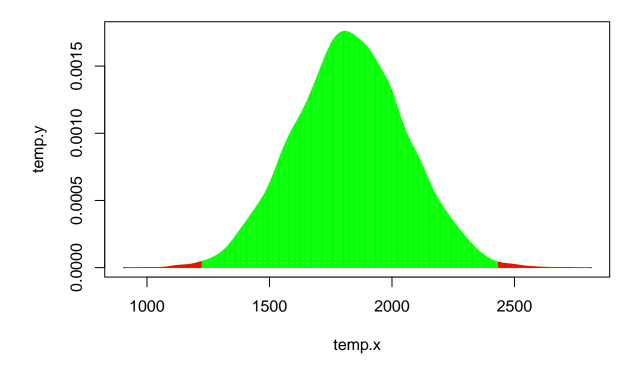
## [1] 0.0108

Since the p value is 0.0076, we reject the null hypothesis. Conducting this test did not require us to specify the population distribution for the GMAT scores!

```
set.seed(87654321)
sampsize = nrow(admission)
f1 = function(){
    s1 = sample(x = c(0,1), size = sampsize, replace = T, prob = c(.5,.5))
    r1 = seq(1:sampsize)
    x = sum(r1[s1==1])
    return(x)
}
sampdist = replicate(10000, f1())
plot(density(sampdist))
```



```
sign_gmat = ifelse(admission$GMAT<510,0,1)
dev_gmat = abs(admission$GMAT-510)
df = data.frame(sign_gmat,dev_gmat)
df = df[order(df$dev_gmat),]
df$rank_gmat = seq(1:sampsize)
tstat = sum(df[df$sign_gmat==1,]$rank_gmat)
p_2tail(sampdist,tstat)</pre>
```



```
## [1] 0.0064
print(tstat)
```

```
## [1] 1223
```

```
sum(df[df$sign_gmat==1,]$rank_gmat)
```

## [1] 1223

```
wilcox.test(x=admission$GMAT,mu = 510)
```

```
##
## Wilcoxon signed rank test with continuity correction
##
## data: admission$GMAT
## V = 1224, p-value = 0.008
## alternative hypothesis: true location is not equal to 510
```

Let us do another one.

## Two Sample Test

Suppose you have the following data on reading scores of third graders. The treatment group has been subjected to a special reading program. You want to test to see if the reading program is effective in improving the reading scores of third graders.

Treatment Group			Control Group				
24	61	59	46	42	33	46	37
43	44	52	43	43	41	10	42
58	67	62	57	55	19	17	55
71	49	54		26	54	60	28
43	53	57		62	20	53	48
49	56	33		37	85	42	

If our null hypothesis is that the treatment has no effect, then it implies that there is no difference between the treatment and the control group. Thus, we can combine all the values into one pool, shuffle them, and randomly reassign them into two groups again. The difference between the averages of each group in each shuffle is only due to sampling error, and not because there is any real and substantive difference between them.

Let us read and take a look at the data.

```
twosample <- read.csv("../../data/twosample.csv")
table(twosample$group)</pre>
```

```
## ## Control Treatment
## 23 21
```

**Step 1**: State the hypothesis.

There is no difference in the mean of the treatment and control groups.

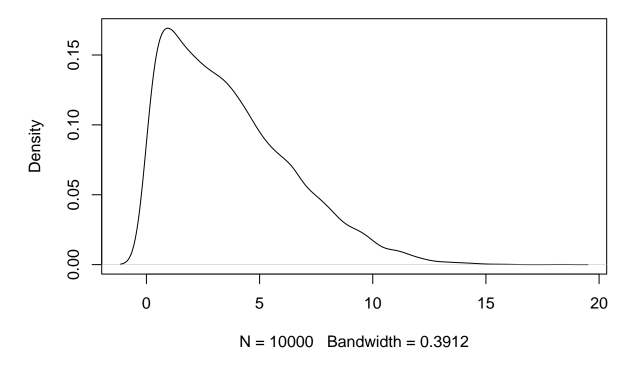
Step 2: Describe the data generation process and the population.

There is no difference between the two groups. We can pool them all into one large group.

Step 3: Create a sampling distribution.

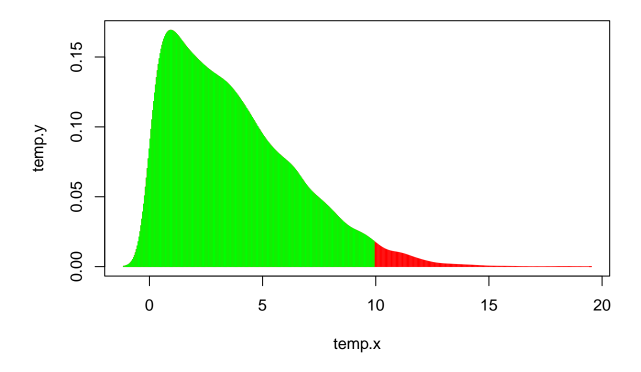
The sampling distribution is created with the following code.

```
set.seed(87654321)
f1 = function(){
  pool = twosample$score
  s1 = sample(pool)
  control1 = s1[1:23]
  treatment1 = s1[24:44]
  return(abs(mean(treatment1)-mean(control1)))
}
sampdist = replicate(10000, f1())
plot(density(sampdist))
```



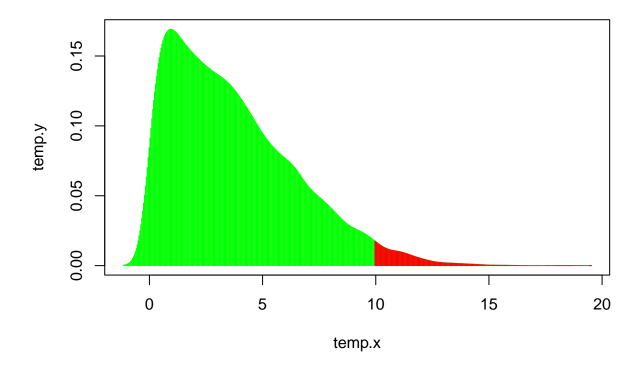
**Step 4**: Get the actual sample and compute the statistic.

```
tstat = abs(mean(twosample$score[1:21]) - mean(twosample$score[22:44]))
p_rtail(sampdist,tstat)
```



**Step 5**: Plot and compute the p value.

```
cutoff = tstat
temp = density(sampdist)
df = data.frame(temp$x, temp$y)
formula1 = df$temp.x>cutoff
df1 = df[formula1,]
plot(df, col = "green", type = "h")
points(df1, col = "red", type = "h")
```



pvalue = length(sampdist[sampdist>tstat])/(length(sampdist))
print(pvalue)

Since the p value is 0.0262, we reject the null hypothesis that there is no difference between the treatment and control groups. Remember again that we made no assumption about the distribution of the student scores. The same basic approach can be used to evaluate if the median or the 75% percentile scores improve with the special reading program.

### Correlation Test

Now let us conduct a nonparametric test for the correlation between two numeric variables. We do not want to assume any underlying population distribution for either of the two numeric variables.

The basic logic is similar to the two sample test. If there is no correlation between the two variables, then high values in one are not associated with high values in the other if they are positively correlated (high and low for negative correlation). In essence, when there is no correlation, you can view it as values in the two being randomly shuffled. The following code implements this logic for the correlation test.

```
immer = read.csv("../../data/immer.csv")
```

**Step 1**: State the hypothesis.

There is no correlation between Y1 and Y2.

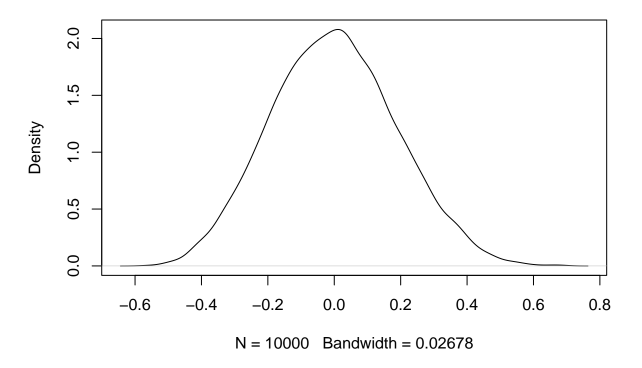
#### Step 2: Describe the data generation process and the population.

Since there is no correlation, we can shuffle the vectors and find their correlation to create a sampling distribution.

### **Step 3**: Create a sampling distribution.

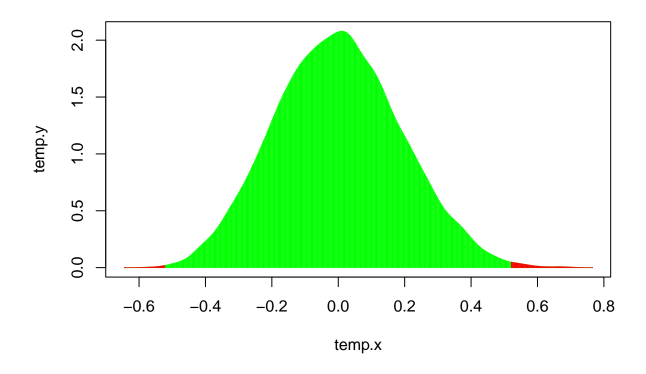
The sampling distribution is created with the following code.

```
set.seed(87654321)
f1 = function(){
  s1 = sample(immer$Y1)
  s2 = sample(immer$Y1)
  return(cor(x = s1,y = s2))
}
sampdist = replicate(10000, f1())
plot(density(sampdist))
```



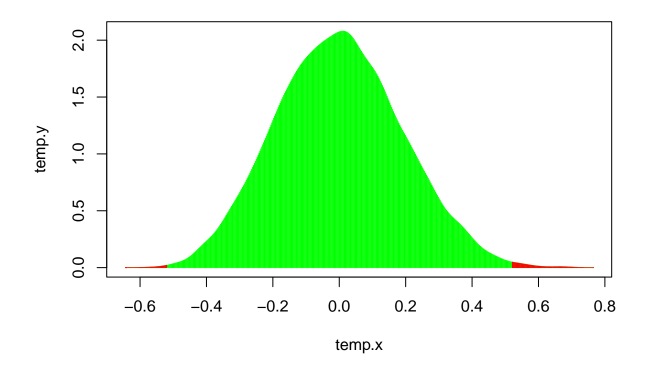
**Step 4**: Get the actual sample and compute the statistic.

```
tstat = cor(immer$Y1, immer$Y2)
p_2tail(sampdist,tstat)
```



**Step 5**: Plot and compute the p value.

```
gap = abs(tstat-0)
cutoff1 = 0 - gap
cutoff2 = 0 + gap
temp = density(sampdist)
df = data.frame(temp$x, temp$y)
formula1 = df$temp.x<cutoff1 | df$temp.x>cutoff2
df1 = df[formula1,]
plot(df, col = "green", type = "h")
points(df1, col = "red", type = "h")
```



pvalue = length(sampdist[sampdist<cutoff1 | sampdist>cutoff2])/(length(sampdist))
print(pvalue)

Therefore, we can reject the null hypothesis of no correlation between the two variables.

## Bootstrapping

Bootstrapping is a very popular nonparametric technique to define population distributions. Let us first take a look at the Merriam-Webster dictionary definition of the term. Bootstrap is defined as **to promote** or develop by initiative and effort with little or no assistance. Let us see how this works.

Suppose we have the following sample of data.

Observed number of complaints = (4, 3, 5, 13, 7, 10, 9, 9, 3, 6, 4, 3, 7, 10, 7, 6, 7, 8, 7, 7)

```
sample1 = c(4,3,5,13,7,10,9,9,3,6,4,3,7,10,7,6,7,8,7,7)
```

The bootstrapping approach works as follows.

1. Since the sample came from the population, the sample is clearly part of the population.

2. Bootstrapping assumes that the data we have are a reasonable representation of the population from which they came and that other data from the population that we did not collect will in fact look like the data we do have. If we do this repeatedly, our population will begin to emerge.

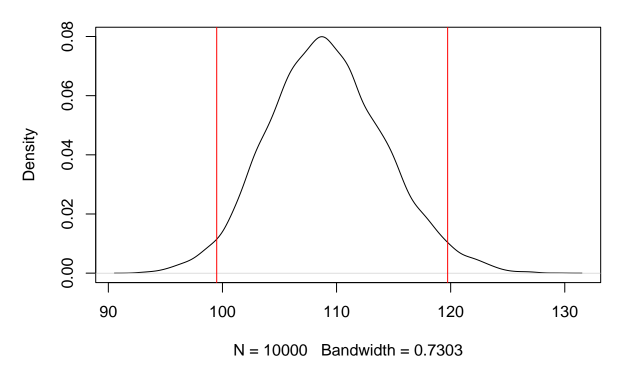
Given this, we can create the sampling distribution by 'taking samples from the one sample we have'. In other words, we sample from the sample with replacement. Once we have the bootstrap sampling distribution, we can use it to create confidence intervals and for hypothesis testing as well.

Let us create the bootstrap sampling distribution and calculate the 95% confidence interval for the variable Y1 in the immer data frame.

```
bootsampdist = replicate(10000, mean(sample(immer$Y1, replace = T)))

q2 = quantile(bootsampdist, c(.05/2,1-(.05/2)))
plot(density(bootsampdist))
abline(v = q2, col = "red")
```

## density.default(x = bootsampdist)



```
paste("95% Confidence interval = [", round(q2,2)[1],", ",round(q2,2)[2],"]")
## [1] "95% Confidence interval = [ 99.5 , 119.73 ]"
```

With the confidence interval defined, a null hypothesis which states that the population value is outside of the interval can be rejected.

## Synthetic Data

#### head(admission)

```
## GPA GMAT De
## 1 2.96 596 admit
## 2 3.14 473 admit
## 3 3.22 482 admit
## 4 3.29 527 admit
## 5 3.69 505 admit
## 6 3.46 693 admit
```

• It would be an interesting exercise for students to run a variety of statistical tests on both original and synthetic data, and compare performance.

### library(synthpop)

```
## Find out more at https://www.synthpop.org.uk/
```

```
s1 = syn(admission)
```

#### head(s1\$syn)

```
## GPA GMAT De
## 1 3.33 559 admit
## 2 2.63 414 notadmit
## 3 3.38 553 admit
## 4 2.89 496 border
## 5 3.15 440 border
## 6 3.03 626 border
```