### **Statistical Inference Course Project**

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My R Markdown file borrows heavily from https://github.com/bcaffo/courses/blob/master/06\_StatisticalInference/03\_01\_TwoGroupIntervals (https://github.com/bcaffo/courses/blob/master/06\_StatisticalInference/03\_01\_TwoGroupIntervals)

### 1. Load the ToothGrowth data and perform some basic exploratory data analyses

In RStudio run help(ToothGrowth) to see a description of the data set.

#### Description

The response is the length of odontoblasts (teeth) in each of 10 guinea pigs at each of three dose levels of Vitamin C (0.5, 1, and 2 mg) with each of two delivery methods (orange juice or ascorbic acid).

```
str(ToothGrowth)

## 'data.frame': 60 obs. of 3 variables:
```

```
## $ len : num 4.2 11.5 7.3 5.8 6.4 10 11.2 11.2 5.2 7 ...
## $ supp: Factor w/ 2 levels "OJ", "VC": 2 2 2 2 2 2 2 2 2 2 2 2 ...
## $ dose: num 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 ...
```

```
head(ToothGrowth)
```

```
## len supp dose
## 1 4.2 VC 0.5
## 2 11.5 VC 0.5
## 3 7.3 VC 0.5
## 4 5.8 VC 0.5
## 5 6.4 VC 0.5
## 6 10.0 VC 0.5
```

```
nrow(ToothGrowth)
```

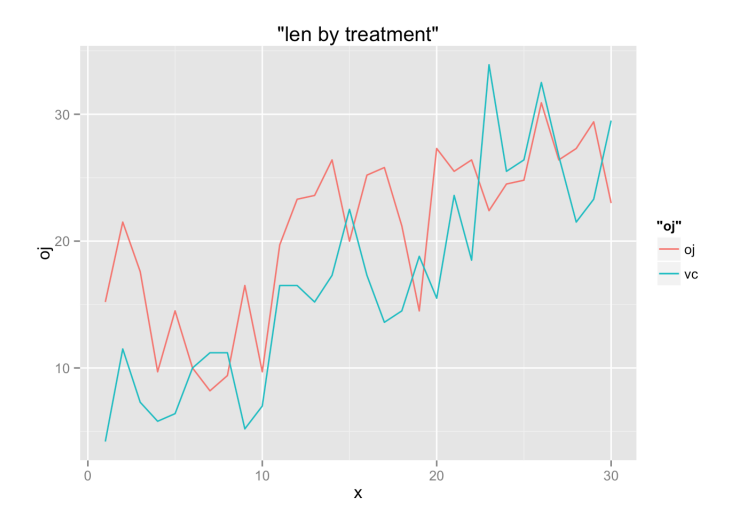
```
## [1] 60
```

```
nrow(na.omit(ToothGrowth) )
```

```
## [1] 60
```

The following graph suggest their is not much difference in tooth growth based on treatment

```
library(ggplot2)
vc <- ToothGrowth$len[ToothGrowth$supp == "VC"]
oj <- ToothGrowth$len[ToothGrowth$supp == "OJ"]
x <- 1:length(vc)
plotData <- data.frame(oj, vc, x)
pp <- ggplot(plotData, aes(x, title="len by treatment"))
pp + geom_line(aes(y=oj, colour="oj")) + geom_line(aes(y=vc, colour="vc"))</pre>
```



### 2. Provide a basic summary of the data.

summary(ToothGrowth)

```
##
       len
                             dose
                 supp
  Min. : 4.2 OJ:30 Min. :0.50
##
##
   1st Qu.:13.1 VC:30
                         1st Qu.:0.50
   Median :19.2
                        Median :1.00
##
##
   Mean :18.8
                         Mean :1.17
   3rd Qu.:25.3
                         3rd Qu.:2.00
##
   Max. :33.9
                         Max. :2.00
##
```

## 3. Use confidence intervals and/or hypothesis tests to compare tooth growth by supp and dose.

We'll use "Two Group intervals Statistical Inference"

First create 6 list, one for each combination of treament and dose

```
vc05 <- ToothGrowth$len[ToothGrowth$supp == "VC" & ToothGrowth$dose == 0.5]
vc1 <- ToothGrowth$len[ToothGrowth$supp == "VC" & ToothGrowth$dose == 1.0]
vc2 <- ToothGrowth$len[ToothGrowth$supp == "VC" & ToothGrowth$dose == 2.0]

oj05 <- ToothGrowth$len[ToothGrowth$supp == "OJ" & ToothGrowth$dose == 0.5]
oj1 <- ToothGrowth$len[ToothGrowth$supp == "OJ" & ToothGrowth$dose == 1.0]
oj2 <- ToothGrowth$len[ToothGrowth$supp == "OJ" & ToothGrowth$dose == 2.0]</pre>
```

Now for each does we'll compare the VC vs OJ

```
mean(vc05) - mean(oj05)

## [1] -5.25

t.test(vc05, oj05, paired = FALSE, var.equal = TRUE)$conf
```

```
## [1] -8.73 -1.77
## attr(,"conf.level")
## [1] 0.95
```

```
mean(vc1) - mean(oj1)
```

```
## [1] -5.93
```

```
t.test(vc1, oj1, paired = FALSE, var.equal = TRUE)$conf
```

```
## [1] -9.019 -2.841
## attr(,"conf.level")
## [1] 0.95

mean(vc2) - mean(oj2)
```

```
## [1] 0.08
```

```
t.test(vc2, oj2, paired = FALSE, var.equal = TRUE)$conf
```

```
## [1] -3.563 3.723
## attr(,"conf.level")
## [1] 0.95
```

Lets see if we find a differece between treaments when we ignore the dose

```
vc <- ToothGrowth$len[ToothGrowth$supp == "VC"]
oj <- ToothGrowth$len[ToothGrowth$supp == "OJ"]
mean(vc) - mean(oj)</pre>
```

```
## [1] -3.7
```

```
t.test(vc, oj, paired = FALSE, var.equal = TRUE)$conf
```

```
## [1] -7.567 0.167
## attr(,"conf.level")
## [1] 0.95
```

# 4. State your conclusions and the assumptions needed for your conclusions.

regardless of the size of the dose guinea pigs recevied, there does not appear to be a statistically signifigant difference in the amount of tooth growth.

The only assumption I made was groups where independent so that we could not use a paired t test