Peer Assessment on ToothGrowth data - Data Science Specialization

Michael Karp

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Investigating the tooth growth dataset

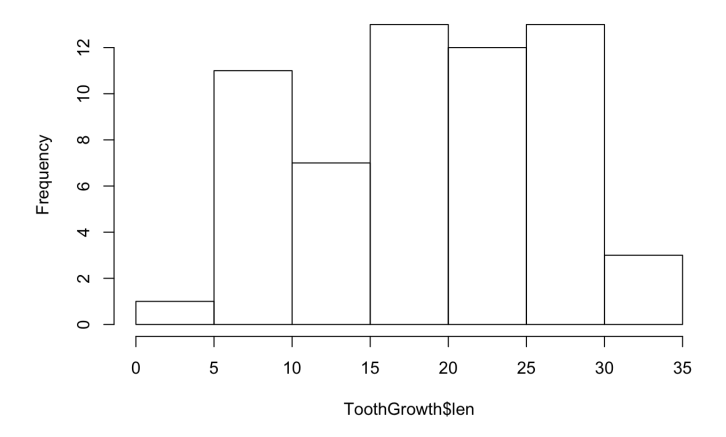
set seed to make research reproducible
set.seed(1000)

Load the ToothGrowth data and perform some basic exploratory data analyses

library(datasets)

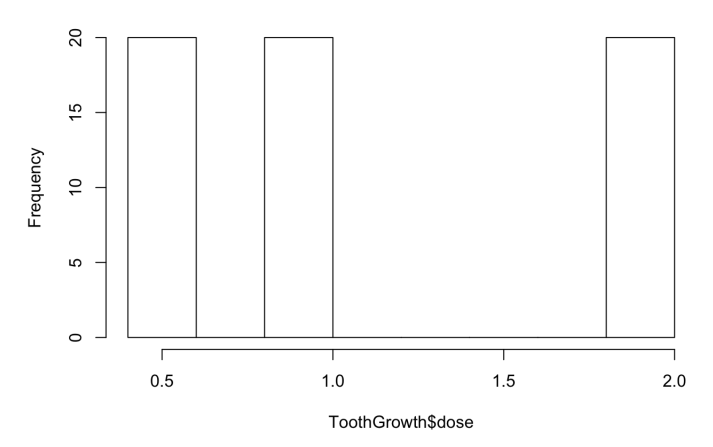
hist(ToothGrowth\$len)

Histogram of ToothGrowth\$len



hist(ToothGrowth\$dose)

Histogram of ToothGrowth\$dose



table(ToothGrowth\$supp == "VC", ToothGrowth\$len > 18.81)

FALSE TRUE

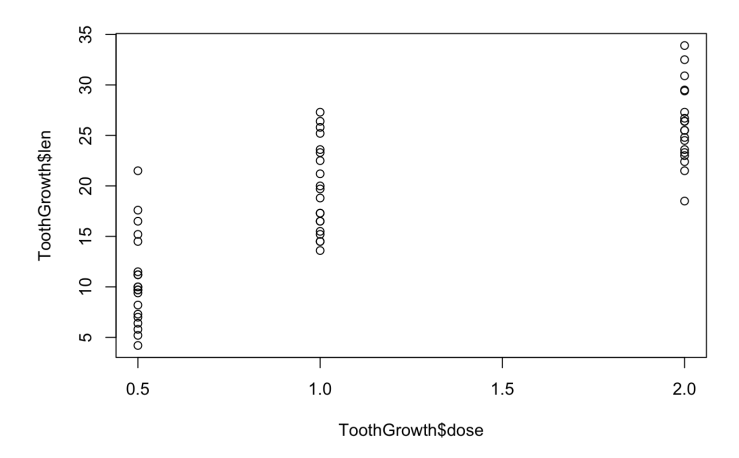
FALSE 10 20 TRUE 20 10

table(ToothGrowth\$supp == "OJ", ToothGrowth\$len > 18.81)

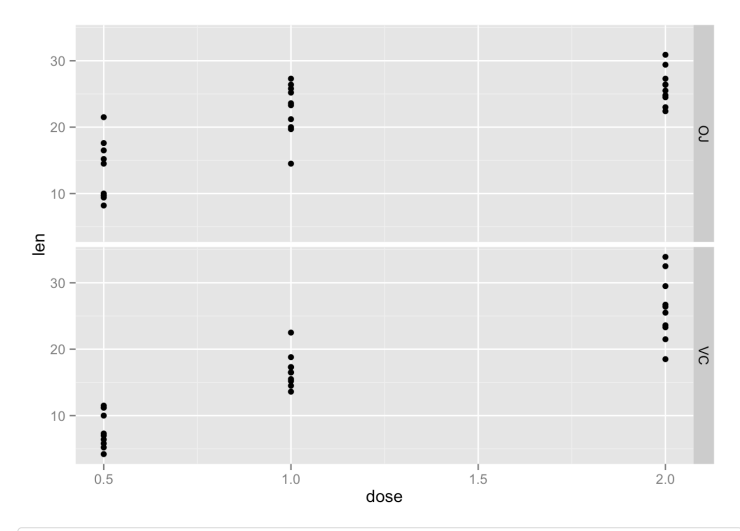
FALSE TRUE

FALSE 20 10 TRUE 10 20

plot(ToothGrowth\$len~ToothGrowth\$dose)
library(ggplot2)



```
exp_plot <- ggplot(data = ToothGrowth, aes(x = dose, y = len)) + geom_point() + facet_grid(
supp~.)
exp_plot</pre>
```



Provide a basic summary of the data.
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 ... \$ dose: num 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 0.5 ...

summary(ToothGrowth)

len supp dose

Min.: 4.20 OJ:30 Min.: 0.500 1st Qu::13.07 VC:30 1st Qu::0.500 Median:19.25 Median:1.000

Mean :18.81 Mean :1.167 3rd Qu.:25.27 3rd Qu.:2.000 Max. :33.90 Max. :2.000

```
# Use confidence intervals and/or hypothesis tests to compare tooth growth by supp and dose
. (Only use the techniques
# from class, even if there's other approaches worth considering)
# H_0 <- mean(dose) == 1.167
# H_a <- mean(dose) > 30
# State your conclusions and the assumptions needed for your conclusions.
```