## Class05.R

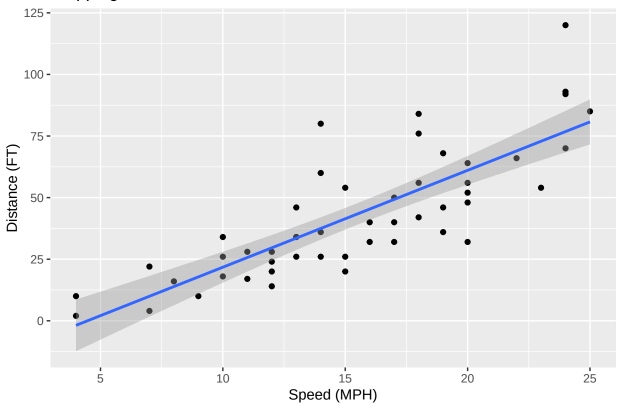
julianapoli

2021-10-13

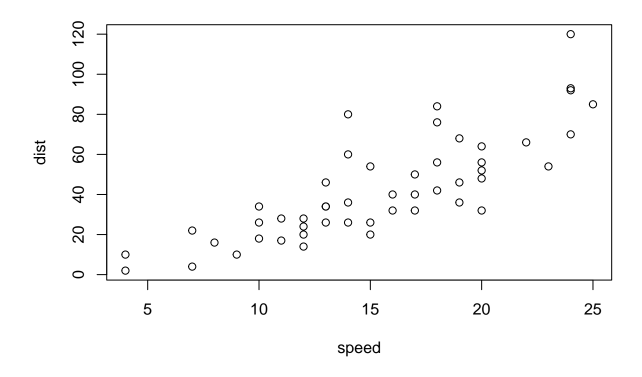
```
# Class 05: Data Visualization
# Today we are going to use ggplot2 package
# Need to install packages, but we only do this once
# install.packages("ggplot2")
library(ggplot2)
\textit{\# We will use this built-in "cars" dataset first}
head(cars)
    speed dist
##
## 1
       4 2
       4 10
## 2
## 3
       7 4
## 4
     7 22
## 5
      8 16
## 6
# All ggplots have at least 3 layers,
\# data + aes + geom
ggplot(data=cars) +
 aes(x=speed, y = dist) +
 geom_point() +
 geom_smooth(method = "lm") +
 labs(title="Stopping Distance of Old Cars",
      x = "Speed (MPH)",
      y = "Distance (FT)")
```

## 'geom\_smooth()' using formula 'y ~ x'

## Stopping Distance of Old Cars



# Side-note: ggplot is not the only graphics system
# a very popular one is good old "base" R graphics
plot(cars)



```
# Doing the adding more plot aesthetics part of lab worksheet
url <- "https://bioboot.github.io/bimm143_S20/class-material/up_down_expression.txt"
genes <- read.delim(url)
head(genes)</pre>
```

```
## Gene Condition1 Condition2 State
## 1 A4GNT -3.6808610 -3.4401355 unchanging
## 2 AAAS 4.5479580 4.3864126 unchanging
## 3 AASDH 3.7190695 3.4787276 unchanging
## 4 AATF 5.0784720 5.0151916 unchanging
## 5 AATK 0.4711421 0.5598642 unchanging
## 6 AB015752.4 -3.6808610 -3.5921390 unchanging
```

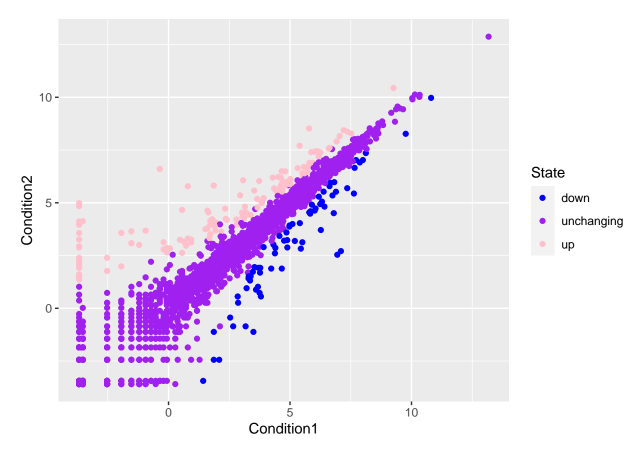
```
# Q: What are the column names? colnames(genes)
```

```
## [1] "Gene" "Condition1" "Condition2" "State"
```

```
# Q: How many genes are in the dataset?
nrow(genes)
```

## [1] 5196

```
# How many columns are there in the dataset?
ncol(genes)
## [1] 4
# Q: How many upregulated genes are there?
table(genes$State)
##
##
         down unchanging
                                up
           72
                    4997
                               127
127/5196
## [1] 0.02444188
round(table(genes$State)/nrow(genes) * 100,2)
##
##
         down unchanging
                               up
         1.39 96.17
                               2.44
##
# Let's make a figure
p <- ggplot(genes) +</pre>
 aes(x=Condition1, y=Condition2, col = State) +
 geom_point()
p + scale_color_manual(values= c("blue", "purple", "pink"))
```



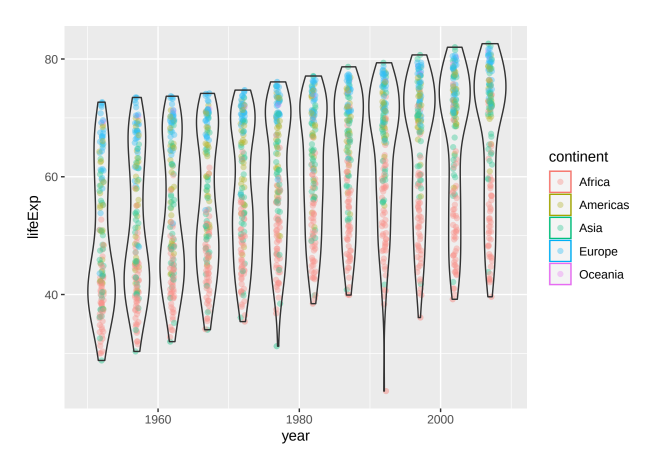
```
# Let's do some optional gapminder stuff
#install.packages("gapminder")
library(gapminder)
head(gapminder)
```

```
## # A tibble: 6 x 6
##
                 continent year lifeExp
     country
                                               pop gdpPercap
     <fct>
                 <fct>
                            <int>
                                    <dbl>
                                             <int>
                                                        <dbl>
## 1 Afghanistan Asia
                             1952
                                     28.8 8425333
                                                         779.
## 2 Afghanistan Asia
                             1957
                                     30.3 9240934
                                                         821.
## 3 Afghanistan Asia
                             1962
                                     32.0 10267083
                                                         853.
## 4 Afghanistan Asia
                             1967
                                     34.0 11537966
                                                         836.
## 5 Afghanistan Asia
                             1972
                                     36.1 13079460
                                                         740.
## 6 Afghanistan Asia
                             1977
                                     38.4 14880372
                                                         786.
```

## # Number of countries nrow(gapminder)

## ## [1] 1704

```
# Let's make a new plot of year vs. life expectancy
ggplot(gapminder) +
  aes(x=year, y =lifeExp, col = continent) +
  geom_jitter(width=0.3,alpha=0.4) +
  geom_violin(aes(group = year), alpha = 0.2)
```



```
# Install the plotly
#install.packages("plotly")
# library(plotly)
# ggplotly()
```