## Class 5: Data Visualization

### Loreen A17059289

### Base R graphics vs ggplot2

There are many graphics systems available in R, including so-called "base" R graphics and the very popular ggplot2 package.

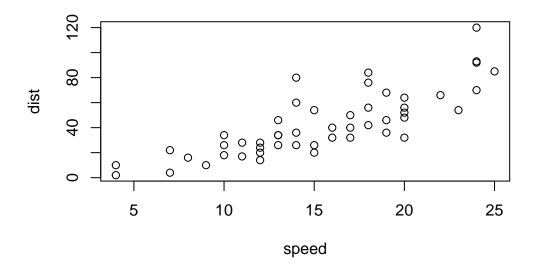
To compare these let's play with the inbuilt cars dataset.

head(cars)

#### 

To use "base" R I can simply call the plot() function:

```
plot(cars)
```



To use ggplot2 package I first need to install it with the function install.packages("ggplot2").

I will run this in my R console (the R brain) as I don't want to re-install it every time I render my report.

The main function in this package is called <code>ggplot()</code>. Can I just call it

```
library(ggplot2)
ggplot()
```

To make a figure with ggplot I need always at least three things:

- data (what I want to plot)
- ${\bf aes}$  the aesthetic mapping of the data to the plot I want
- the **geoms** (How I want to plot the data)

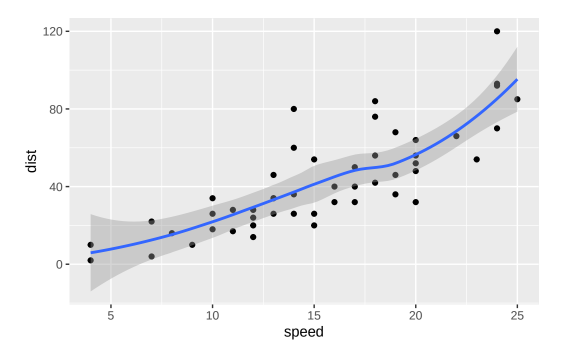
```
ggplot(data=cars) +
  aes(x=speed, y=dist) +
  geom_point()
```



If I want to add more stuff I can just keep adding layers:

```
ggplot(data=cars) +
  aes(x=speed, y=dist) +
  geom_point() +
  geom_smooth()
```

 $\ensuremath{\mbox{`geom\_smooth()`}}\ \mbox{using method} = \ensuremath{\mbox{'loess'}}\ \mbox{and formula} = \ensuremath{\mbox{'y}}\ \sim \ensuremath{\mbox{x'}}\ \mbox{'}$ 



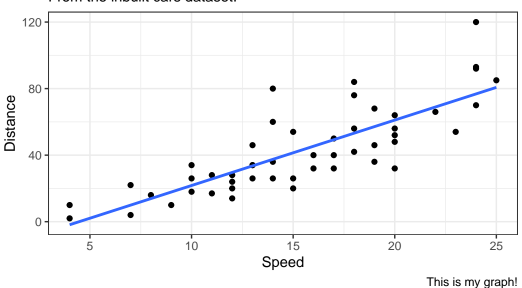
ggplot is much more verbose than base R plots for standard plots but it has a consistent layer system that I can use to make just about any plot.

Making the line straight and removing outline/highlight:

```
ggplot(data=cars) +
  aes(x=speed, y=dist) +
  geom_point() +
  geom_smooth(se=FALSE, method="lm") +
  labs(title = "Cars", caption = "This is my graph!", subtitle="From the inbuilt cars data
  theme_bw()
```

<sup>`</sup>geom\_smooth()` using formula = 'y ~ x'

# Cars From the inbuilt cars dataset:



### A more complicated plot

Let's plot some gene expression data. The code below reads the results of a differential expression analysis where a new anti-viral drug is being tested.

```
url <- "https://bioboot.github.io/bimm143_S20/class-material/up_down_expression.txt"
genes <- read.delim(url)
head(genes)</pre>
```

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

Q. How many genes are in this dataset?

```
nrow(genes)
```

### [1] 5196

Q How can we summarize that last column - the "State column?

```
table(genes$State)
```

```
down unchanging up
72 4997 127
```

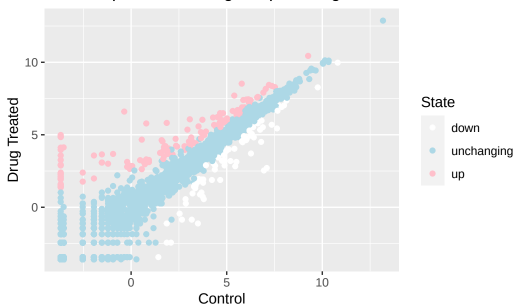
Plot this data:

```
p <- ggplot(genes) +
  aes(x=Condition1, y=Condition2, color=State) +
  geom_point()</pre>
```

I can now just call p when I want to plot or add to it.

```
p + labs(title="Gene Expression Changes Upon Drug Treatment", x="Control", y="Drug Treated
scale_colour_manual( values=c("white","lightblue","pink") )
```

### Gene Expression Changes Upon Drug Treatment



##Going Further

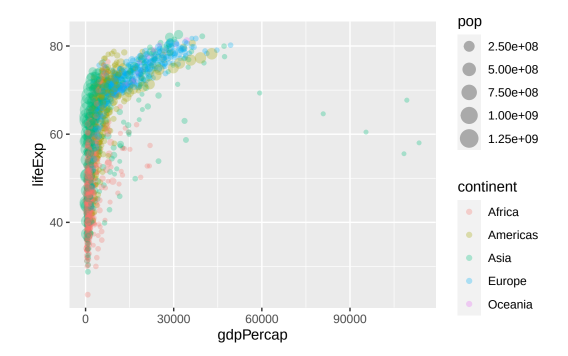
Here I read a slightly larger dataset:

```
# File location online
url <- "https://raw.githubusercontent.com/jennybc/gapminder/master/inst/extdata/gapminder.
gapminder <- read.delim(url)
head(gapminder)</pre>
```

```
countrycontinentyearlifeExppopgdpPercap1AfghanistanAsia195228.8018425333779.44532AfghanistanAsia195730.3329240934820.85303AfghanistanAsia196231.99710267083853.10074AfghanistanAsia196734.02011537966836.19715AfghanistanAsia197236.08813079460739.98116AfghanistanAsia197738.43814880372786.1134
```

Plot of data above:

```
ggplot(gapminder) +
  aes(x=gdpPercap, y=lifeExp, col=continent, size=pop) +
  geom_point(alpha=0.3)
```



A very useful layer to add sometimes is for "faceting".

```
ggplot(gapminder) +
  aes(x=gdpPercap, y=lifeExp, col=continent, size=pop) +
  geom_point(alpha=0.3) +
  facet_wrap(~continent)
```

