# Class05: Data visualization

Gen (PID: A15900230)

# 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.

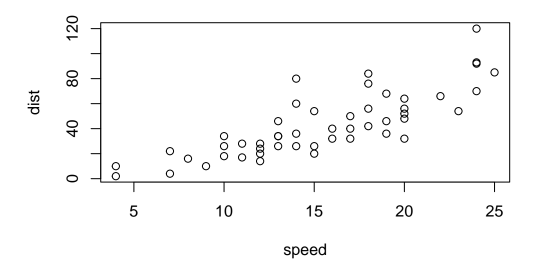
To compare these let's play with the inbuilt cars dataset

```
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 (i.e. the R brain) as I do not want to re-install it every time I render my report...

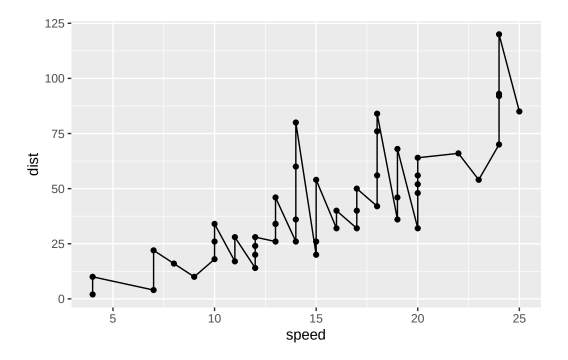
The main function in this package is called <code>ggplot()</code>. Can I just call it:

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

To make a figure with ggplot, I always need at least 3 things:

- data (i.e. what i want to plot)
- aes the aesthetic mapping of the data to the plot i want.
- $\bullet \;$  the  $\mathbf{geoms}$  i.e. How I want to plot the data.

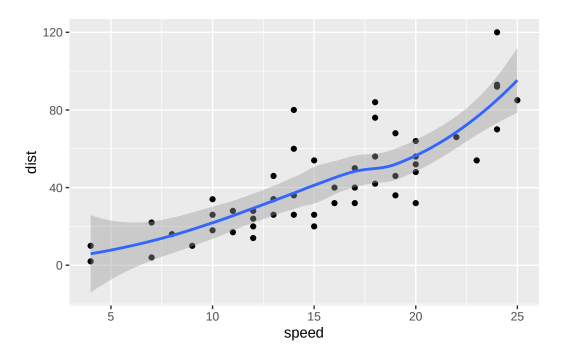
```
ggplot(data=cars) +
  aes(x=speed, y=dist) +
  geom_point() + geom_line()
```



If I want to add more stuff, I can just keep adding layers, e.g.

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

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



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

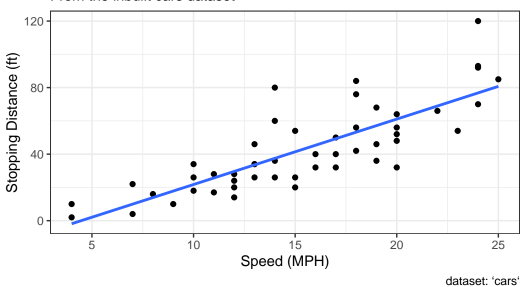
let's make a plot with a straight line fit - i.e. a linear model and no standard error shown.

```
ggplot(data=cars) +
  aes(x=speed, y=dist) +
  geom_point() +
  geom_smooth(se=FALSE, method="lm") +
  labs(title="Speed and Stopping Distances of Cars", x="Speed (MPH)", y="Stopping Distances)
```

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

# Speed and Stopping Distances of Cars

### From the inbuilt cars dataset



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
```

nrow(genes)

#### [1] 5196

Q: how many genes are in this dataset?

nrow(genes)

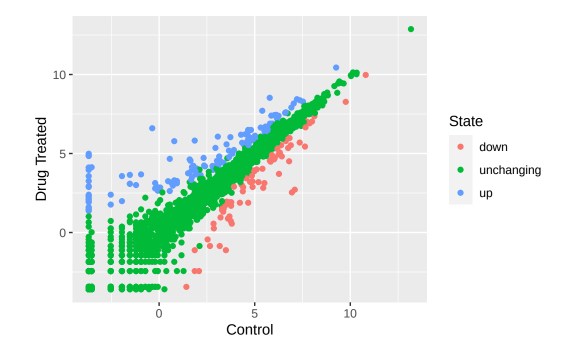
# [1] 5196

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

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

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

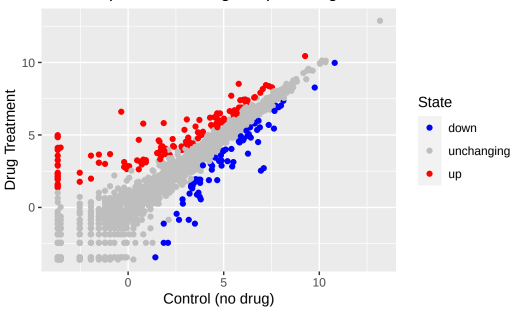
```
{\tt ggplot(genes) + aes(x=Condition1, y=Condition2, color=State) + geom\_point() + labs(x="Contor=State) + geom\_point() + geom\_point()
```



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

p + scale_colour_manual( values=c("blue", "gray", "red") ) + labs(title= "Gene Expression Ch</pre>
```

# Gene Expression Changes Upon Drug Treatment



# Going further

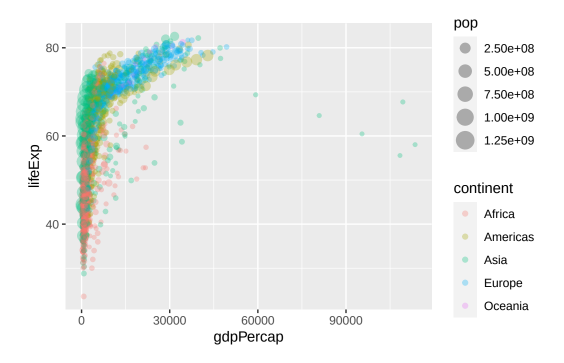
Here I read a slightly larger dataset

ggplot(gapminder) +

geom\_point(alpha=0.3)

```
# File location online
  url <- "https://raw.githubusercontent.com/jennybc/gapminder/master/inst/extdata/gapminder.
  gapminder <- read.delim(url)</pre>
  head(gapminder)
      country continent year lifeExp
                                         pop gdpPercap
1 Afghanistan
                   Asia 1952 28.801 8425333 779.4453
2 Afghanistan
                  Asia 1957 30.332 9240934 820.8530
3 Afghanistan
                  Asia 1962 31.997 10267083 853.1007
4 Afghanistan
                  Asia 1967 34.020 11537966 836.1971
5 Afghanistan
                  Asia 1972 36.088 13079460 739.9811
6 Afghanistan
                  Asia 1977 38.438 14880372 786.1134
```

aes(x=gdpPercap, y=lifeExp, color=continent, size=pop) +



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

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

