

Class 5: Data visualization with gg plot

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Today we will have our first play with the **ggplot2** package - one of the most popular graphics on the planet.

There are many plotting systems in R. These include so called “*base*” plotting/graphics.

```
plot(cars)
```



base plot is generally rather short code and somewhat dull plots - but it is always there for you and is fast for big data sets.

If I want to use **ggplot2** it takes some more work.

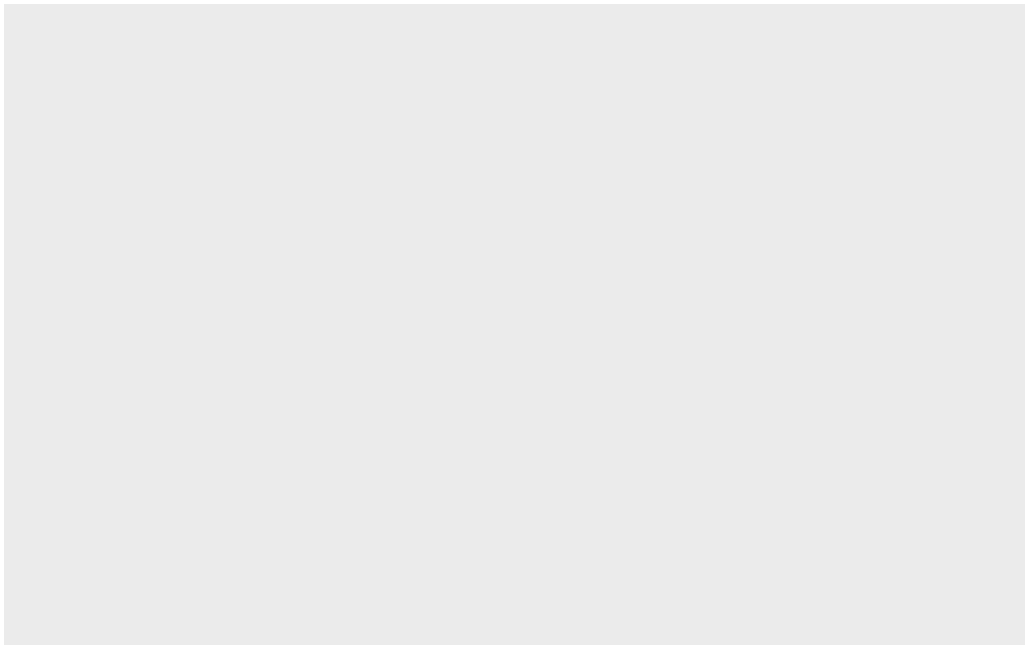
```
# ggplot(cars)
```

I need to install the package first to my computer. To do this I can use the function `install.packages("ggplot2")`

Everytime I want to use a package I need to load it up with a `library()` call.

```
library(ggplot2)
```

```
ggplot(cars)
```

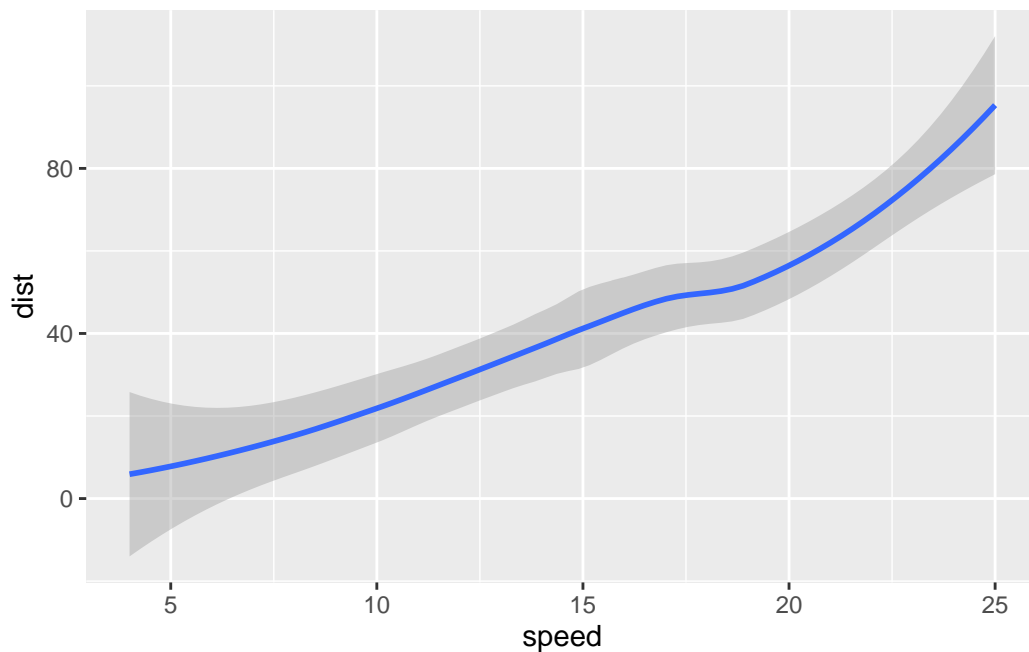


Every ggplot has at least 3 things:

- **data** (the data.frame with the data you want to plot)
- **aes** (the aesthetic mapping of the data to the plot)
- **geom** (how do you want the plot to look, points, lines, etc)

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

`geom_smooth()` using method = 'loess' and formula = 'y ~ x'

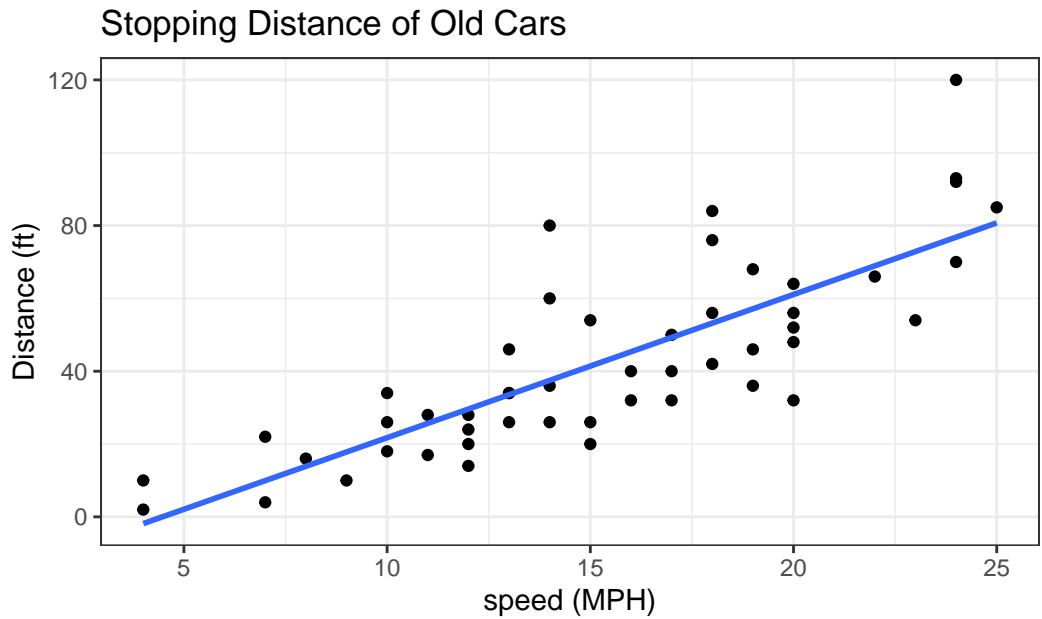


I want a linear model and no standard error bounds shown on my plot. I also want nicer axis labels a title etc.

```
bp <- ggplot(cars) + aes(x=speed, y=dist) + geom_point()
```

```
bp + geom_smooth(se=FALSE, method = "lm") + labs(title="Stopping Distance of Old Cars", x=
```

`geom_smooth()` using formula = 'y ~ x'



From the 'cars' dataset

A more complicated scatter plot

Here we make a plot of gene expression data:

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

	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

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

```
[1] 5196
```

```
colnames(genes)
```

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

```
ncol(genes)
```

```
[1] 4
```

```
url <- "https://bioboot.github.io/bimm143_S20/class-material/up_down_expression.txt"
genes <- read.delim(url)
table(genes$State)
```

down	unchanging	up
72	4997	127

```
url <- "https://bioboot.github.io/bimm143_S20/class-material/up_down_expression.txt"
genes <- read.delim(url)
round( sum(genes$State == "up")/ nrow(genes) * 100, 2)
```

```
[1] 2.44
```

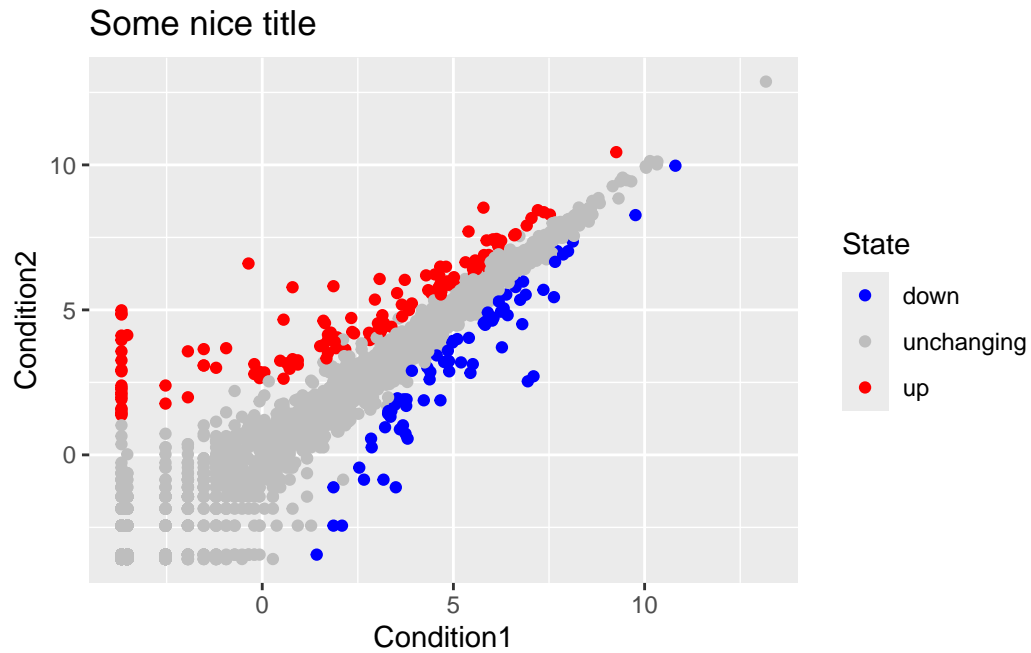
```
head(genes, 2)
```

	Gene	Condition1	Condition2	State
1	A4GNT	-3.680861	-3.440135	unchanging
2	AAAS	4.547958	4.386413	unchanging

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

Change the colors

```
p + labs(title= "Some nice title") + scale_colour_manual( values=c("blue","gray","red") )
```



Exploring the gapminder dataset

Here we will load up the gapminder dataset to get practice with different aes mappings.

```
url <- "https://raw.githubusercontent.com/jennybc/gapminder/master/inst/extdata/gapminder.  
gapminder <- read.delim(url)
```

How many entries rows are in this dataset?

```
nrow(gapminder)
```

```
[1] 1704
```

```
ncol(gapminder)
```

```
[1] 6
```

```
dim(gapminder)
```

```
[1] 1704      6
```

```
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

```
table(gapminder$year)
```

1952	1957	1962	1967	1972	1977	1982	1987	1992	1997	2002	2007
142	142	142	142	142	142	142	142	142	142	142	142

How many continents

```
table(gapminder$continent)
```

Africa	Americas	Asia	Europe	Oceania
624	300	396	360	24

I could use the ‘unique()’ function

```
length(unique(gapminder$continent))
```

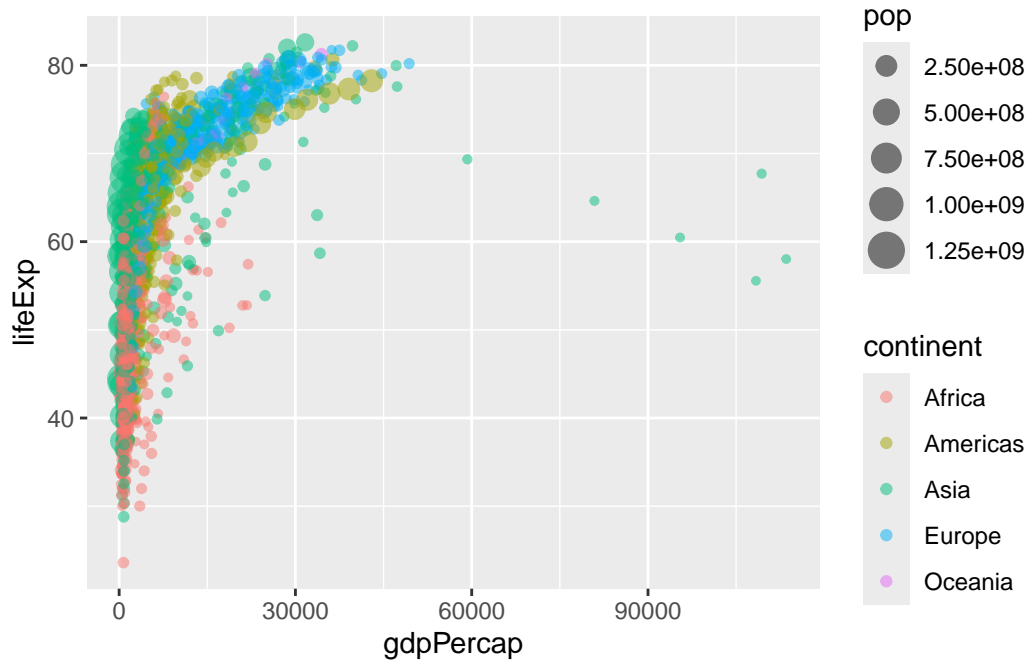
```
[1] 5
```

how many countries in data set?

```
length(unique(gapminder$country))
```

```
[1] 142
```

```
ggplot(gapminder) + aes(x=gdpPercap, y=lifeExp, color=continent, size=pop) + geom_point(al
```



```
library(dplyr)
```

Attaching package: 'dplyr'

The following objects are masked from 'package:stats':

filter, lag

The following objects are masked from 'package:base':

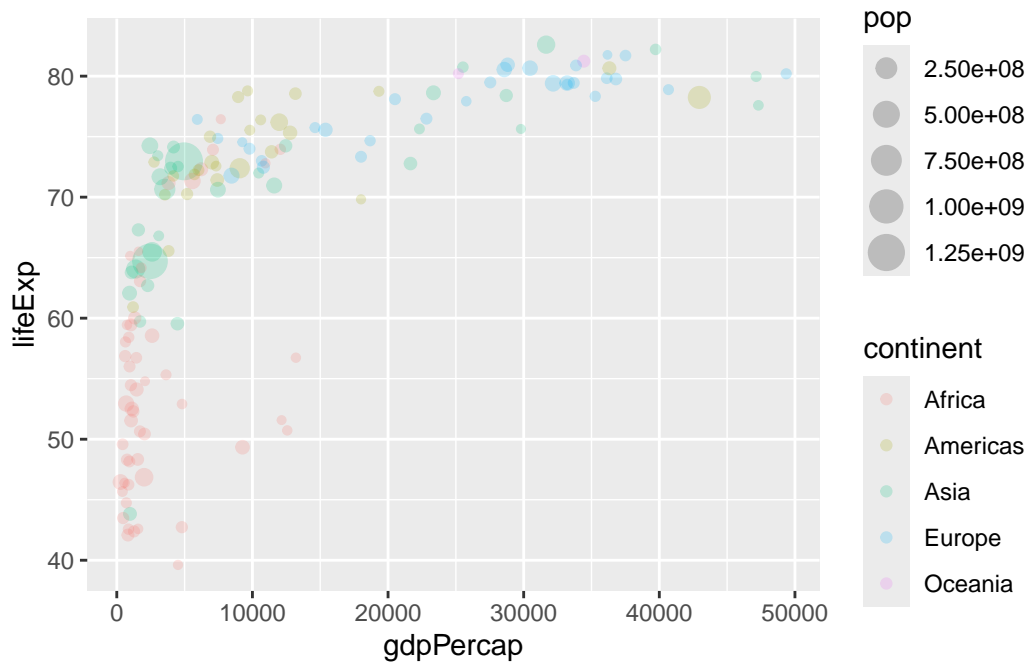
intersect, setdiff, setequal, union

```
gapminder_2007 <- filter(gapminder, year==2007)
```

```
head(gapminder_2007)
```


	country	continent	year	lifeExp	pop	gdpPercap
1	Afghanistan	Asia	2007	43.828	31889923	974.5803
2	Albania	Europe	2007	76.423	3600523	5937.0295
3	Algeria	Africa	2007	72.301	33333216	6223.3675
4	Angola	Africa	2007	42.731	12420476	4797.2313
5	Argentina	Americas	2007	75.320	40301927	12779.3796
6	Australia	Oceania	2007	81.235	20434176	34435.3674

```
ggplot(gapminder_2007) + aes(x=gdpPercap, y=lifeExp, col=continent, size=pop) + geom_point
```



```
ggplot(gapminder) + aes(x=gdpPercap, y=lifeExp) + geom_point() + facet_wrap(~continent)
```

