# R語言與資料視覺化

ALL HOLENIUM

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### 環境資訊頁面

- ■所有課程補充資料、投影片皆位於
  - https://github.com/ywchiu/cdc\_course

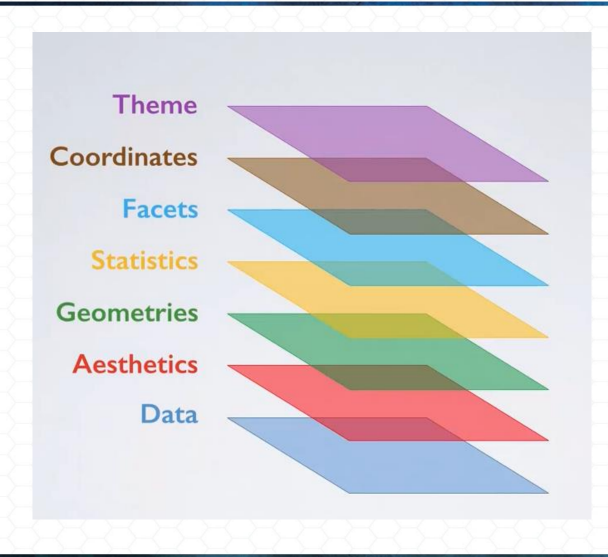
# GGPlot2

# 繪圖語法 (Grammar of Graphics)

- Wilkinson 用Grammar of Graphics 解釋統計圖形 (statistical graphic)的概念
- Wickham 則是以該語法為基礎發展出 ggplot 繪圖系統

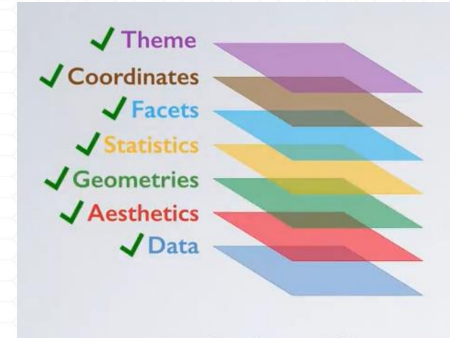


### GGPlot繪圖架構



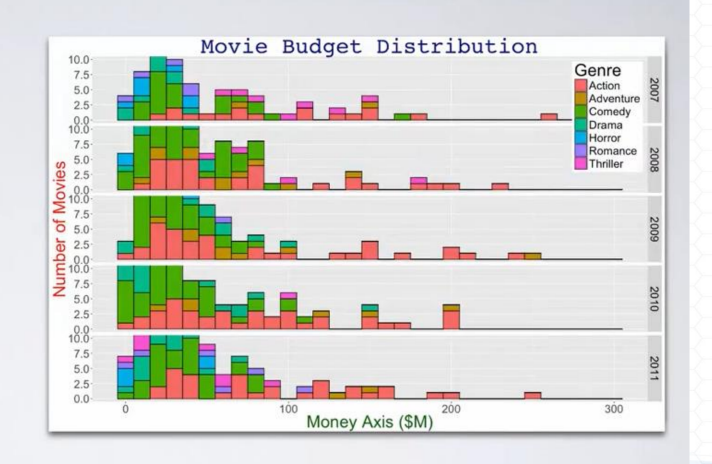
- 主題 (Theme)
  - □ 控制資料以外的繪圖組件
- 座標系統 (Coordinates)
  - □ 指定繪圖時所使用的座標系統
- 繪圖面 (Facets)
  - □ 指定如何將資料分散在多張子圖形中繪製
- 統計轉換 (Statistics)
  - □ 指定如何以將資料轉換為各種統計量。
- 幾何圖案 (Geometries)
  - □ 繪製資料用的幾何圖形
- 美學對應 (Aesthetic)
  - □ 指定原始資料與圖形之間的對應關係
- 資料來源 ( Data )
  - □ 原始資料來源

#### 繪圖範例



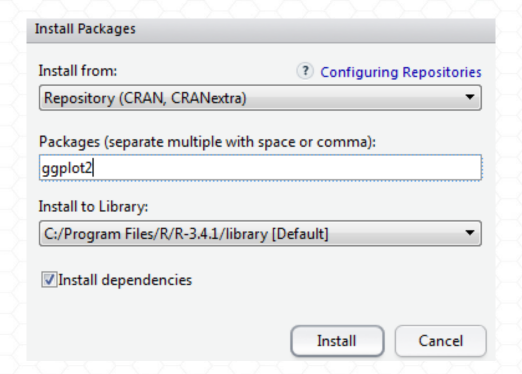
Aesthetics: x, Fill

Movie Name	Budget	Genre	
ABC	\$5 M	Action	
DEF	\$50 M	Action	
GHI	\$12 M	Comedy	



#### 安裝GGPlot2

- ■安裝ggplot2
  - □install.packages("ggplot2")
- 使用ggplot2
  - □ library(ggplot2)
- ■觀看說明頁
  - □help(package='ggplot2')



### 資料來源(Data)

■ 從疫情中心的開放資料網站蒐集到「地區年齡性別統計表-麻疹」(2003年起各地區、各年齡層、性別之病例數統計表(疾病名稱:麻疹,日期種類:發病日,病例種類:確定病例,感染來源:本土、境外移入)

#### ■讀取資料

library(readr)

measles <- read\_csv("https://raw.githubusercontent.com/ywchiu/cdc\_course/master/data/measles.csv")
head(measles)</pre>

確定病名 <chr></chr>	發病年份 ⊴int>	發病月份 <int></int>	縣市 <chr></chr>	<b>鄉鎮</b> <chr></chr>	性別 <chr></chr>	是否為境外移入 <chr></chr>	年齢層 <chr></chr>	確定病例數
麻疹	2009	4	高雄市	小港區	M	否	20-24	2
麻疹	2009	5	基隆市	中山區	M	否	30-34	1
麻疹	2011	6	新北市	蘆洲區	F	是	25-29	1
麻疹	2014	1	高雄市	<b>副</b> 男三	F	是	0	1
麻疹	2017	3	台北市	松山區	F	是	0	1
麻疹	2018	4	桃園市	蘆竹區	F	否	30-34	1

#### 日期資料轉換

- Date
  - □為簡單版的時間資料格式

- POSIXct, POSIXIt
  - □POSIXct: 以整數形式存放時間資料 (1970/01/01至今的秒數)
  - ■POSIXIt: 以年、月、週、日等有效訊息存放資料

### 日期函式

```
x <- as.Date("2018-05-08")
class(x)
unclass(x)

y <- as.Date("1970-01-01")
unclass(y)</pre>
```

# 時間處理 (1)

```
■印出系統時間
```

```
x <- Sys.time()
```

X

■時間轉換

```
p <- as.POSIXIt(x)
```

unclass(p)

p\$sec

p1 <- as.POSIXct(x) unclass(p1)

### 時間處理 (2)

■利用strptime 轉換時間

```
ds <- c("May 8, 2018 12:00")
x <- strptime(ds, "%B %d, %Y %H:%M")</pre>
```

X

#### ■時間運算

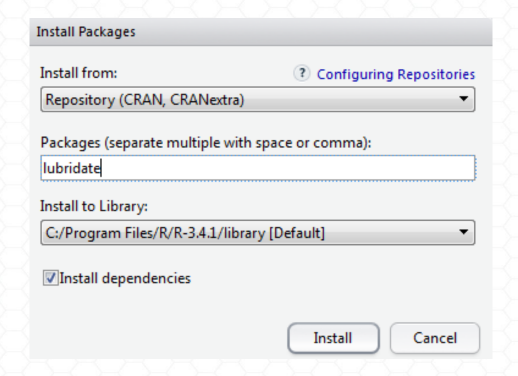
x <- strptime(ds, "%B %d, %Y %H:%M")</p>

x1 <- as.POSIXIt(as.Date('2018-05-08'))

x - x1

#### 使用Lubridate 轉換時間資料

- ■安裝lubridate
  - □install.packages("lubridate")
- ■使用lubridate
  - □ library(lubridate)
- ■觀看說明頁
  - help(package='lubridate')



#### 使用Lubridate 轉換時間資料

```
library(lubridate)
ymd("20180605")
ymd("2018/06/05")
dmy("05/06/2011")
ymd_hms('2018-05-06 08:00:00')
```

#### 取得年月等資訊

```
bday <- dmy("15/10/1988")
month(bday)
```

wday(bday, label = TRUE)

year(bday) <- 2018 wday(bday, label = TRUE)

#### 轉換時間資料

measles\$發病時間 <- as.Date(with(measles, paste(發病年份, 發病月

份, '01',sep="-")), "%Y-%m-%d")

With 語法可指名發病年份與發病月份 資料來自measles 資料集

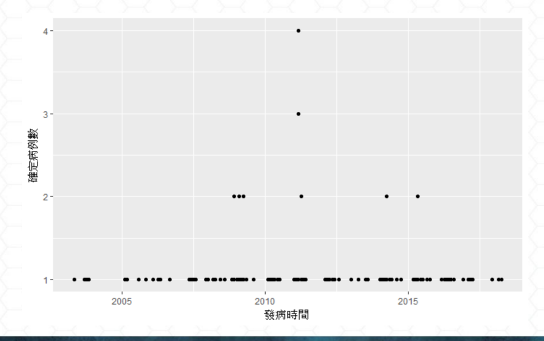
確定病名	發病年份 發	病月份 縣市	鄉鎮 <chr></chr>	性別 <chr></chr>	是否為境外移力 <chr></chr>	年齢層 <chr></chr>	確定病例數	發病時間 <date></date>
麻疹	2009	4 高	小	М	否	20-24	2	2009-04-01
麻疹	2009	5 基	中	M	否	30-34	1	2009-05-01
麻疹	2011	6 新	蘆	F	是	25-29	1	2011-06-01
麻疹	2014	1 高	三	F	是	0	1	2014-01-01
麻疹	2017	3 台	松	F	是	0	1	2017-03-01
麻疹	2018	4 桃	蘆	F	否	30-34	1	2018-04-01

# 美學對應 (Aesthetic)

#### 繪製散佈圖

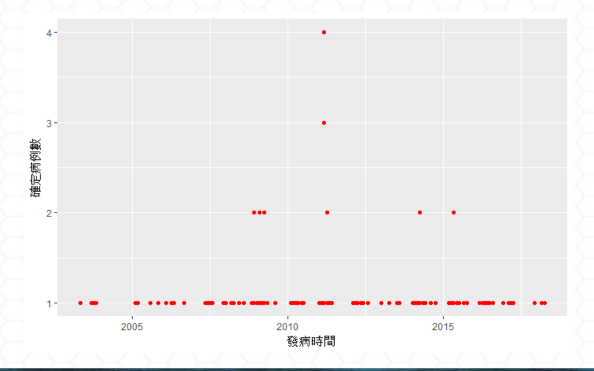
```
library(ggplot2)
```

p1 <- ggplot(measles, aes(x = 發病時間, y = 確定病例數))
p1 + geom\_point()



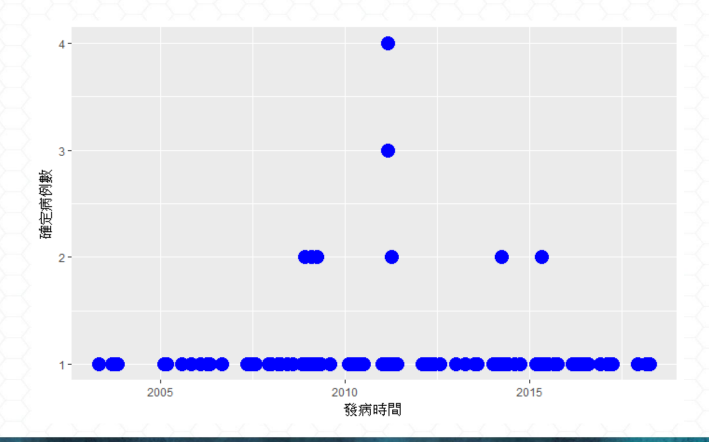
#### 修改顏色

p1 <- ggplot(measles, aes(x = 發病時間, y = 確定病例數))
p1 + geom\_point(color='red')



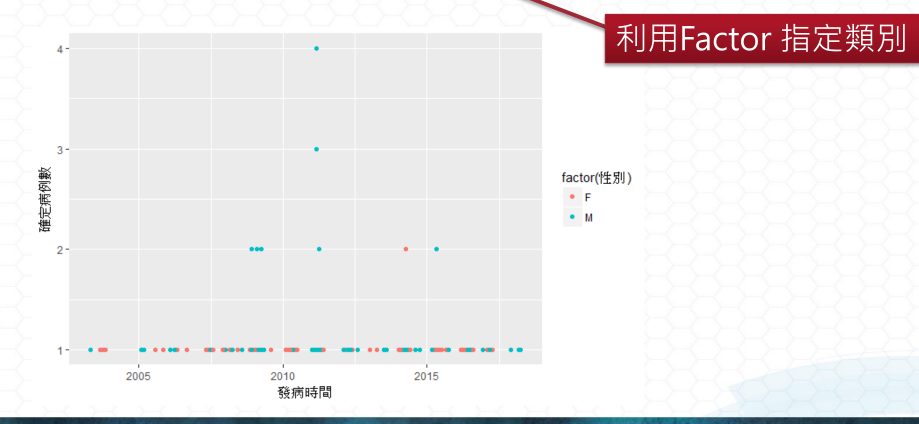
### 修改點的大小、顏色、形狀

p1 + geom\_point(color='blue', size = 5, shape = 19)



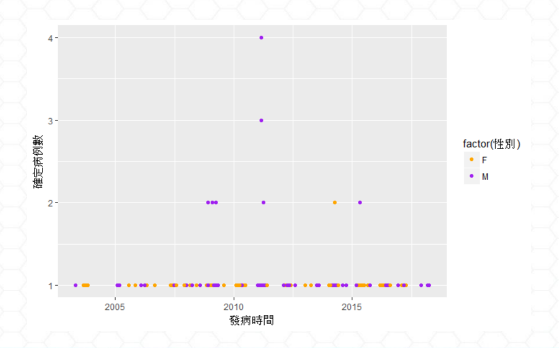
# 根據類別 (factor) 修改

p1 + geom\_point(aes(color=factor(性別))



#### 指定各類別的顏色

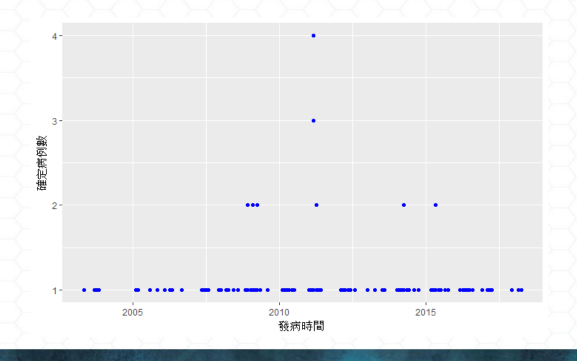
```
p1 +
geom_point(aes(color=factor(性別))) +
scale_color_manual(values = c("orange", "purple"))
```



指定類別顏色

#### 覆載原本的點樣式

```
p1 +
geom_point(color='red') +
geom_point(color='blue')
```



修改原本點的顏色

#### 修改顏色與性別

#### source('multiplot.R')

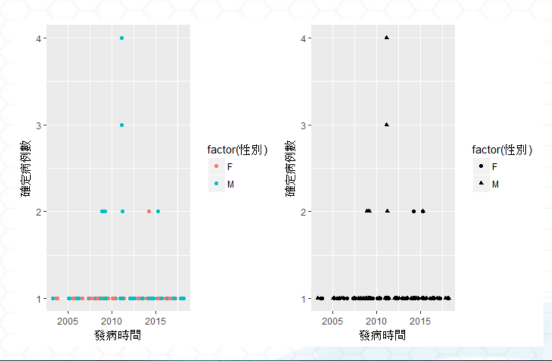
p2 <- p1 + geom\_point(aes(color=factor(性別)))

p3 <- p1 + geom\_point(aes(shape=factor(性別)))

multiplot(p2, p3, cols=2)

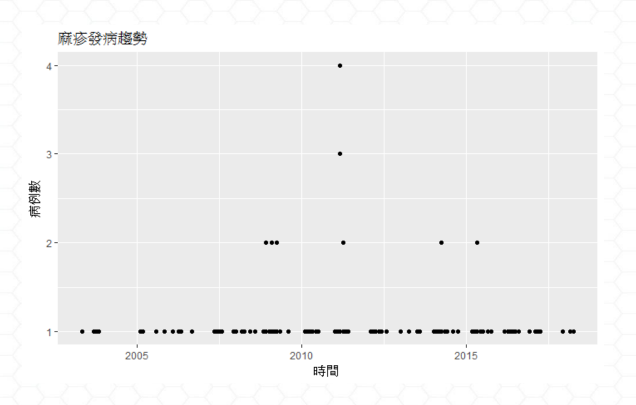
#### multiplot.R 可載自

https://raw.githubusercontent.com/ywchiu/cdc\_course/master/script/multiplot.R



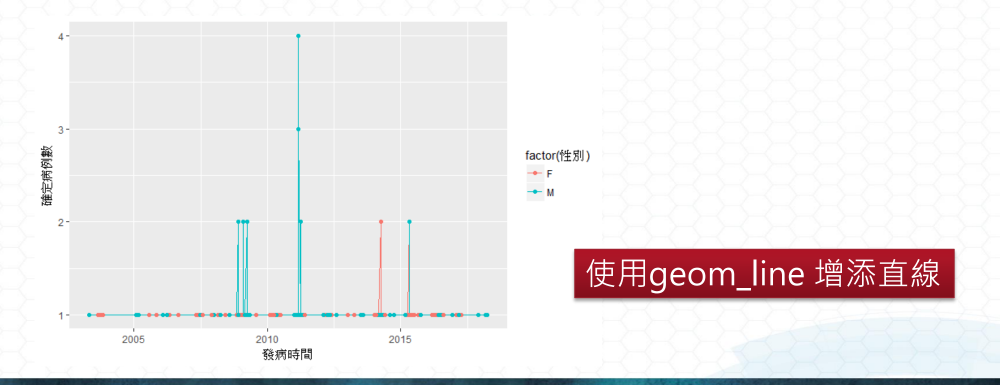
#### 修改標籤名稱與標題名稱

```
p1 +
geom_point() +
xlab("時間") +
ylab("病例數") +
ggtitle("麻疹發病趨勢")
```



#### 圖層疊加

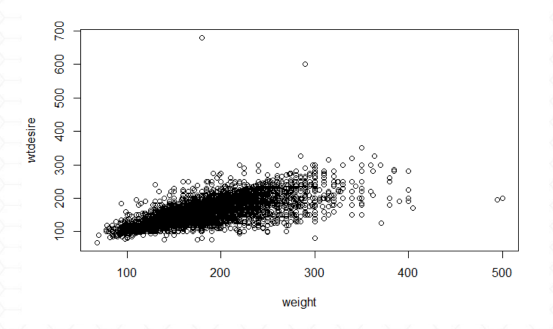
```
p1 + geom_point(aes(color=factor(性別))) + geom_line(aes(color=factor(性別))) p1 + geom_line(aes(color=factor(性別))) + geom_point(aes(color=factor(性別)))
```



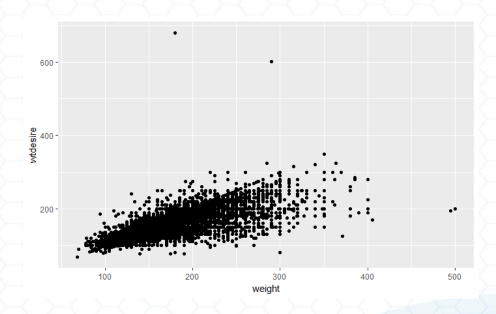
# 幾何圖案 (Geometries)

# 散佈圖 (Scatter Plot)

load('cdc.RData')
plot(wtdesire ~ weight, data = cdc)

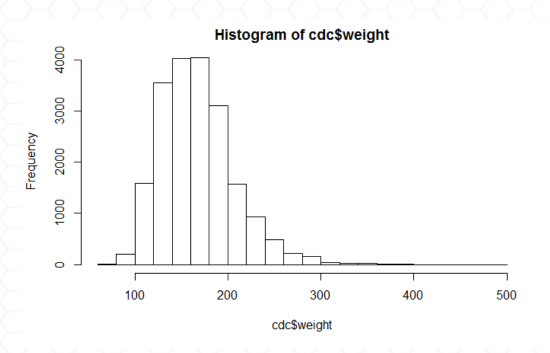


load('cdc.RData')
ggplot(data=cdc, aes(x=weight, y =
wtdesire)) + geom\_point()

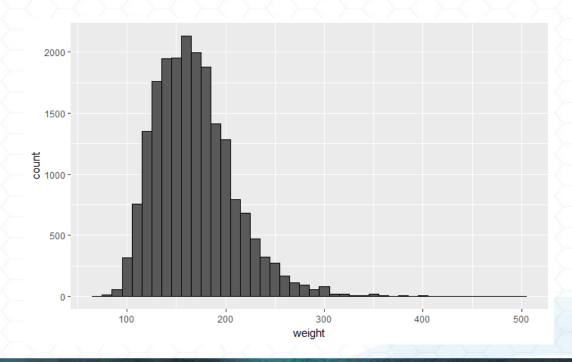


# 直方圖 (Histogram)

hist(cdc\$weight)

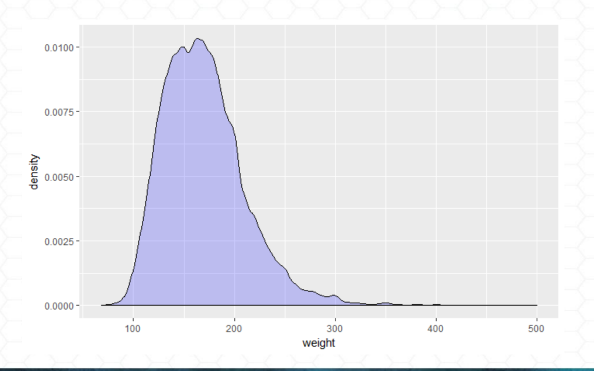


histogram <- ggplot(data=cdc, aes(x=weight))
histogram + geom\_histogram(binwidth=10,
color="black")



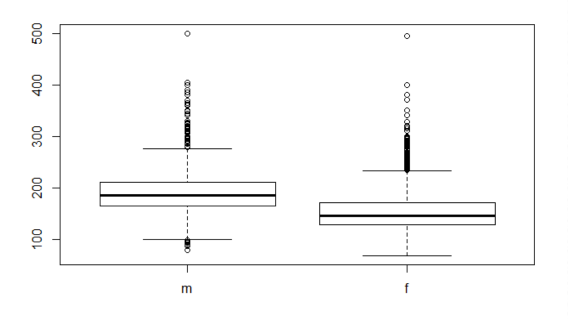
# 密度圖 (Density Plot)

```
density <- ggplot(data=cdc, aes(x=weight))
density + geom_density(stat="density", alpha=I(0.2), fill='blue')</pre>
```

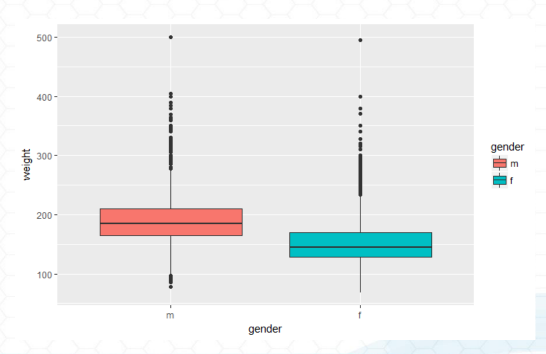


# 箱型圖 (Box Plot)

boxplot(cdc\$weight~ cdc\$gender)

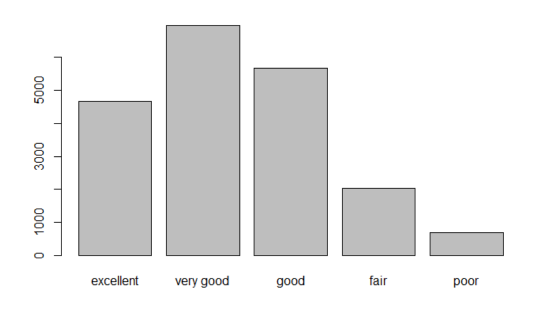


box <- ggplot(data=cdc,
aes(x=gender, y=weight))
box + geom\_boxplot(aes(fill=gender ))</pre>

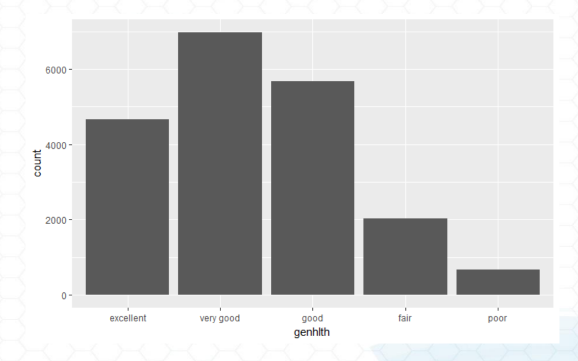


# 長條圖 (Bar Plot)

barplot(table(cdc\$genhlth))

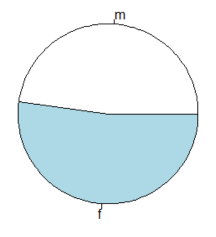


bar <- ggplot(data=cdc, aes(x=genhlth))
bar + geom\_bar()</pre>

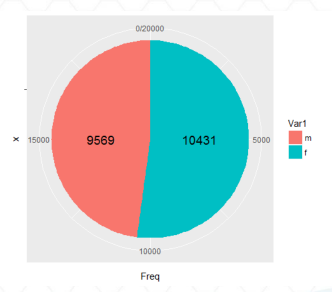


### 圓餅圖 (Pie Chart)

pie(table(cdc\$gender))



pie <- ggplot(cdc\_sex, aes(x="", y=Freq
,fill=Var1 )) + geom\_bar(width=1, stat =
"identity") + geom\_text(aes(label=Freq, y
=c(15000, 5000) ) , size=5)
pie + coord\_polar(theta="y", start = 0)</pre>



# 統計轉換(Statistics)

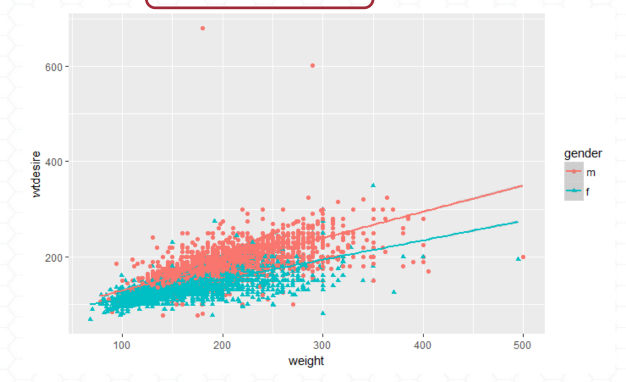
# 增添迴歸線 (Smooth)

smooth <- ggplot(data=cdc, aes(x=weight, y=wtdesire, color=gender)) +</pre>

geom\_point(aes(shape=gender), size=1.5)

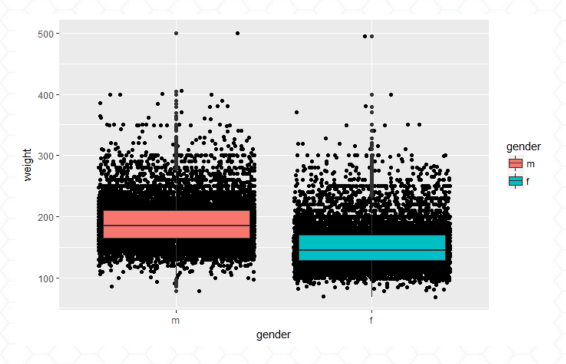
smooth + geom\_smooth(method="lm")

可替換方法為 method="loess"



#### 增添資料點到箱型圖上

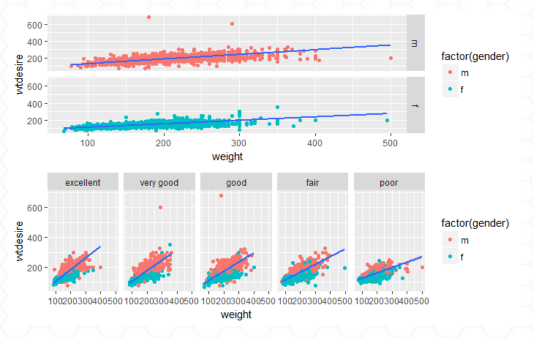
```
box <- ggplot(data=cdc, aes(x=gender, y=weight))
box + geom_jitter() + geom_boxplot(aes(fill=gender ))</pre>
```



## 繪圖面 (Facets)

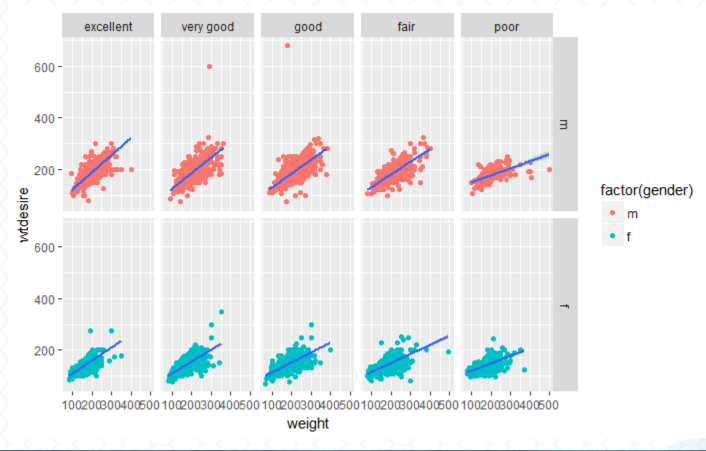
### 產生多張子圖

```
w <- ggplot(data=cdc, aes(x=weight, y = wtdesire)) + geom_point(aes(color=factor(gender))) +
geom_smooth(method = 'lm')
w1 <- w + facet_grid(gender~.)
w2 <- w + facet_grid(.~genhlth)
multiplot(w1, w2, cols = 1)</pre>
```



### 產生多張子圖 (2)

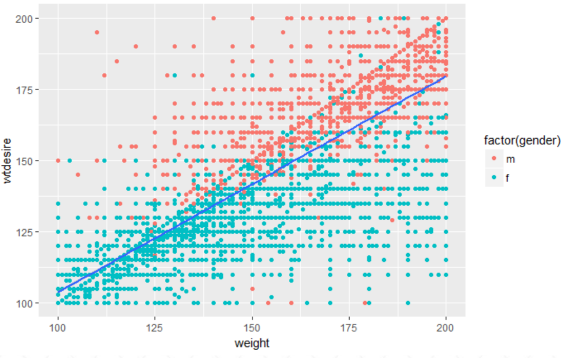
### w + facet\_grid(gender~genhlth)



### 座標系統 (Coordinates)

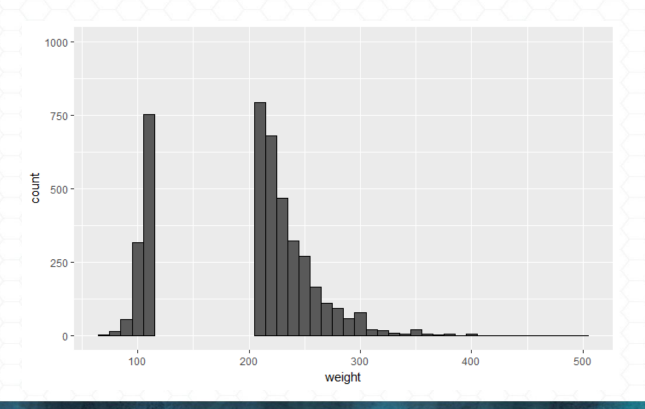
### 局部放大

```
w <- ggplot(data=cdc, aes(x=weight, y = wtdesire)) +
geom_point(aes(color=factor(gender))) + geom_smooth(method = 'lm')
w + xlim(100,200) + ylim(100,200)</pre>
```



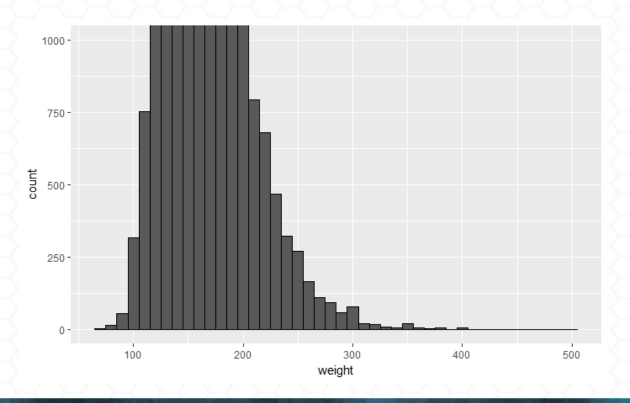
### 並非調整ylim 就可以調整所有圖表

```
histogram <- ggplot(data=cdc, aes(x=weight))
histogram + geom_histogram(binwidth=10, color="black") + ylim(0,1000)
```



### 調整座標

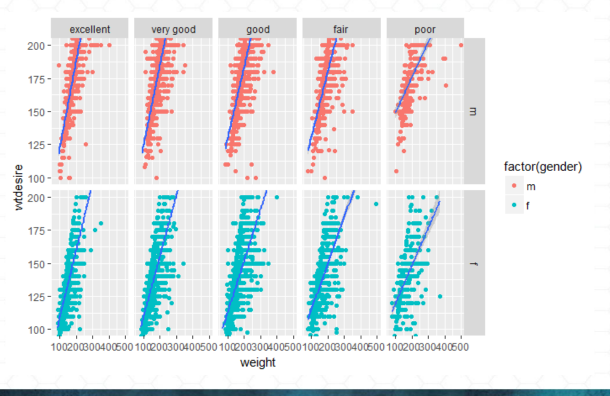
histogram + geom\_histogram(binwidth=10, color="black") + coord\_cartesian( ylim = c(0,1000) )



### 調整座標 (2)

```
w <- ggplot(data=cdc, aes(x=weight, y = wtdesire)) + geom_point(aes(color=factor(gender))) +
geom_smooth(method = 'lm')</pre>
```

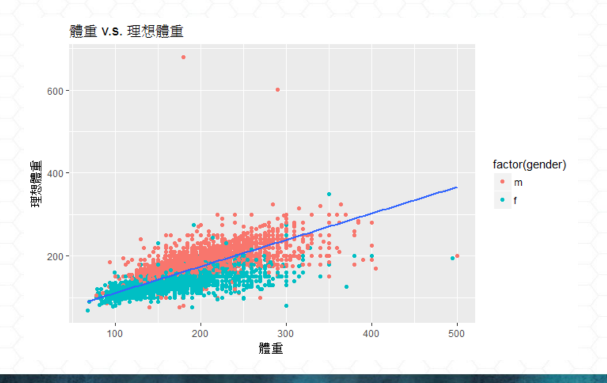
w + facet\_grid(gender~genhlth) + coord\_cartesian(ylim = c(100,200))



### 主題 (Theme)

### 增加圖表標籤敘述

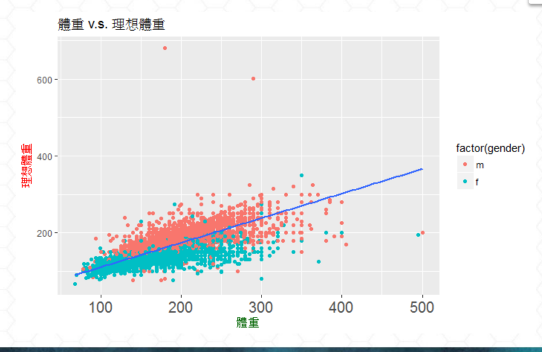
```
w <- ggplot(data=cdc, aes(x=weight, y = wtdesire)) + geom_point(aes(color=factor(gender))) + geom_smooth(method = 'lm') w + xlab('體重') + ylab('理想體重') + ggtitle('體重 v.s. 理想體重')
```



### 修改標籤樣式

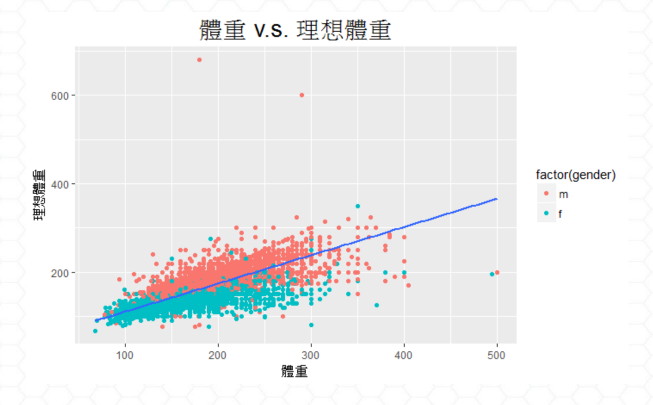
```
w + xlab('體重') + ylab('理想體重') + ggtitle('體重 v.s. 理想體重') + theme(axis.title.x = element_text(color = 'DarkGreen', size = 10), axis.title.y = element_text(color = 'Red', size = 10), axis.text.x = element_text(size = 15), axis.text.y = element_text(size = 15))
```

#### 善用?theme 觀看選項



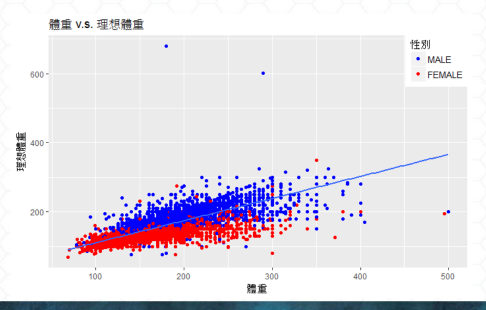
### 調整標題

w + xlab('體重') + ylab('理想體重') + ggtitle('體重 v.s. 理想體重') + theme(plot.title = element\_text(size = 20, hjust = 0.5))



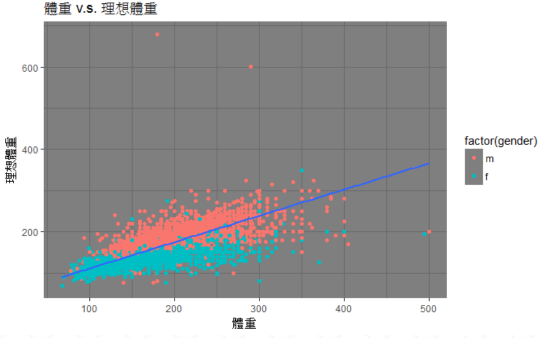
### 修改圖標

```
w + xlab('體重') + ylab('理想體重') + ggtitle('體重 v.s. 理想體重') + scale_color_manual(name = '性別',labels = c("MALE", "FEMALE"), values = c("blue", "red")) + theme(legend.text = element_text(size = 10), legend.title = element_text(size = 10), legend.position = c(1,1), legend.justification = c(1,1))
```



### 修改圖背景樣式

```
w <- ggplot(data=cdc, aes(x=weight, y = wtdesire)) + geom_point(aes(color=factor(gender))) + geom_smooth(method = 'lm') w1 <- w + xlab('體重') + ylab('理想體重') + ggtitle('體重 v.s. 理想體重') w1 + theme_dark()
```



### 匯出圖片

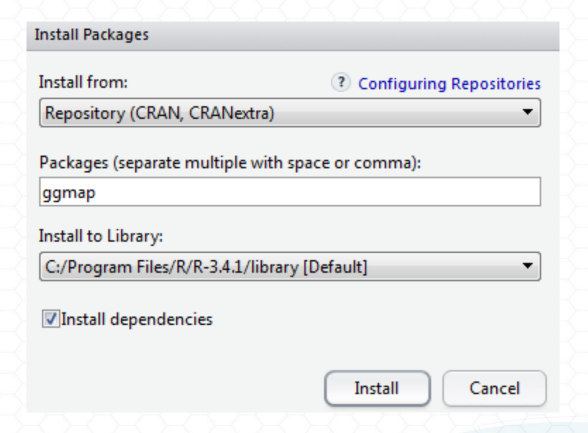
### 使用ggsave 存儲圖片

```
w <- ggplot(data=cdc, aes(x=weight, y = wtdesire)) + geom_point(aes(color=factor(gender))) + geom_smooth(method = 'lm')
w1 <- w + xlab('體重') + ylab('理想體重') + ggtitle('體重 v.s. 理想體重')
w1 + theme_dark()
ggsave("plot1.png")
ggsave(w1, file="plot2.png")
ggsave(w1, file="plot3.png", width=6, height=4)
```

## ggmap

### 安裝ggmap

- ■安裝ggmap
  - □install.packages("ggmap")
- ■使用ggmap
  - □ library(ggmap)
- ■觀看說明頁
  - □help(package='ggmap')



### 顯示台灣地圖

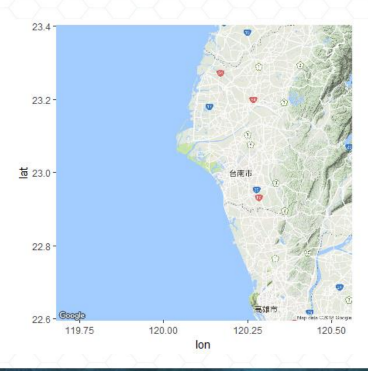
library(ggmap)
map <- get\_map(location = 'Taiwan', zoom = 7)
ggmap(map)</pre>



### 顯示特定經緯度地圖

map <-  $get_map(location = c(lon = 120.12, lat = 23.00),$ zoom = 10, language = "zh-TW")

ggmap(map)



### 蒐集資料

■ 從疫情中心的開放資料網站蒐集到登革熱1998年起每日確定病例統計數字

library(readr)

Dengue <- read\_csv("Dengue\_Daily")
head(Dengue)</pre>

### 將資料點繪製於地圖

```
map <- get_map(location = c(lon = 120.246100, lat = 23.121198), zoom = 10, language = "zh-TW") ggmap(map, darken = c(0.5, "white")) + geom_point(aes(x = 最小統計區中心點X, y = 最小統計區中心點Y), color = "red", data = Dengue_Daily)
```

## 其他繪圖套件

### 高級繪圖套件介紹

- plotly · googleVis
- ■其他:
  - □Ggplot2
  - **□**Ggvis
  - □地圖: RgoogleMaps
  - □社會網路: igraph
  - □熱繪圖:heatmap
  - □類別資料視覺化:vcd
  - □多變量條件式繪圖: lattice
  - □股票: quantmod::chartSeries

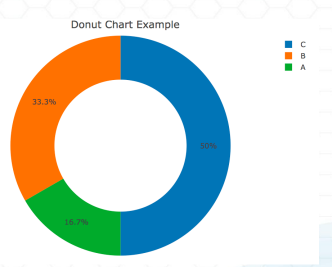
### plotly

■ 可使用Python,R,Matlab,Javascript撰寫的開源繪圖套件,快速、方便

■產生HTML檔

### **Donut Chart**

```
Is a single of the state of th
```



### **Area Chart**

layout(yaxis = list(title = 'rainfall') )

```
Iibrary(plotly)
month<- c(1,2,3,4,5)
taipei <- c(92.5,132.6,168.8,159.1,218.7)
tainan <- c(21.2, 30.6, 37.3, 84.6, 184.3)

plot_ly(x = month, y = taipei, fill = "tozeroy", name="taipei",type='scatter', mode= 'markers') %>%
```

add\_trace(x = month, y = tainan, fill = "tozeroy",name="tainan") %>%

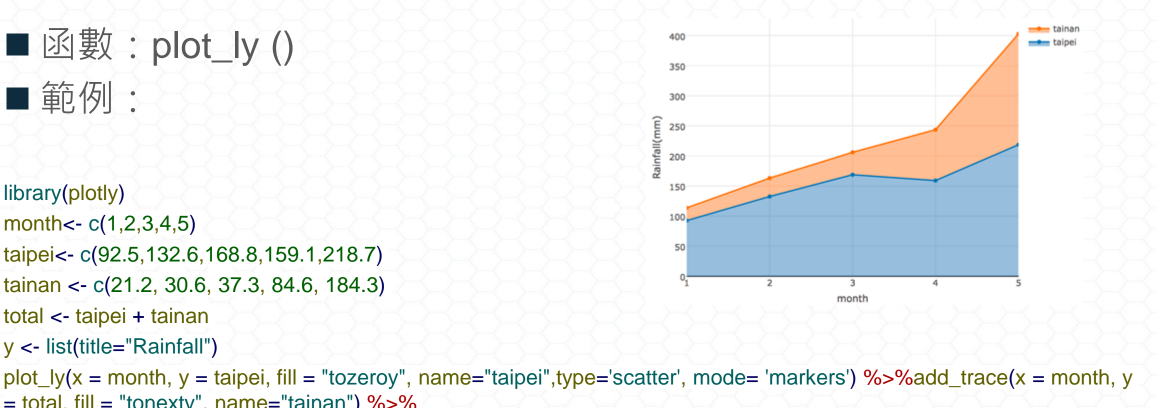
### **Area Chart (Stacked)**

```
■ 函數:plot_ly ()
```

■範例:

layout(yaxis = y)

```
library(plotly)
month<- c(1,2,3,4,5)
taipei<- c(92.5,132.6,168.8,159.1,218.7)
tainan <- c(21.2, 30.6, 37.3, 84.6, 184.3)
total <- taipei + tainan
y <- list(title="Rainfall")</pre>
= total, fill = "tonexty", name="tainan") %>%
```



### **Bubble Chart**

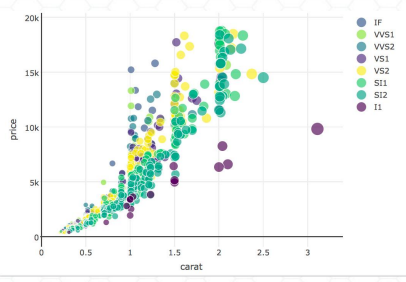
■ 函數: plot\_ly (mode="markers")

data("diamonds")

diamonds

d <- diamonds[sample(nrow(diamonds), 1000), ]

plot\_ly(d, x = d\$carat, y= d\$price, color = d\$clarity, type='scatter', mode= 'markers', size = d\$carat, text= paste("Clarity", d\$clarity))



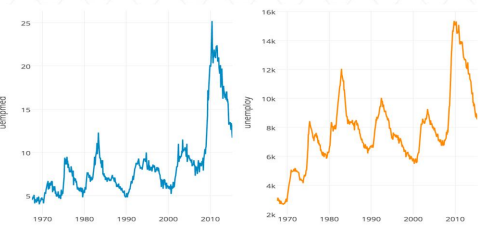
### **Multiple Plots**

p %>% layout(showlegend=FALSE)

■ 函數: subplot(margin,nrows,...)

data("economics")

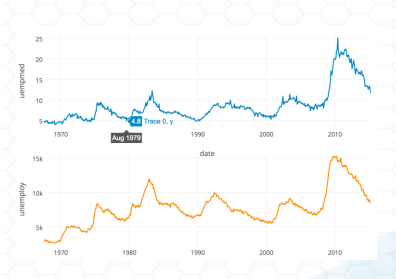
p <- subplot(
plot\_ly(economics, x = economics\$date, y = economics\$unemploy, type='scatter', mode = 'lines'),
plot\_ly(economics, x = economics\$date, y = economics\$uempmed, type='scatter', mode = 'lines'),
margin=0.05
)



### **Multiple Plots**

■ 函數: subplot(margin,nrows,...)

```
p <- subplot(
plot_ly(economics, x = economics$date, y = economics$unemploy, type='scatter', mode = 'lines'),
plot_ly(economics, x = economics$date, y = economics$uempmed, type='scatter', mode = 'lines'),
margin=0.05,
nrows = 2
)
p %>% layout(showlegend=FALSE)
```



# THANK YOU