# CB2-101: R introduction

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## 1 History of R

R is the *lingua franca* of statistical software. It is a dialect of S, a computer language developed by John Chambers in 1976 at Bell Labs. In 1988, the system was rewritten in C. The software changed hand frequently, until in 1993, R appeared in the scene. It was a *free* implementation of S, written by Ross Ihaka and Robert Gentleman at the University of Auckland, New Zealand.

## 2 Starting R

If you're in command-line environment you can start R by typing the command R. In our example will will use a far nicer environment called RStudio. It is available from: https://www.rstudio.com/.

#### 3 R environment

- 1. getwd() Get the current directory
- 2. setwd() Set the current directory
- 3. ls() Lists the objects created.
- 4. rm() Will remove objects.
- 5. list.files() or dir() List the files in the current directory
- 6. source() Reads a R file

#### 3.1 Exercise

Create a file called source\_test.R to have the following code.

```
myfunction <- function () {
    x <- rnorm(100)
    mean(x)
}
myfunction()</pre>
```

```
## [1] -0.1980887
```

Type source("source\_test.R") in the R console. Type ls(), you'll see the object myfunction has been created in the workspace. You can also run the script from the command line.

Rscript source\_test.R

```
## WARNING: ignoring environment value of R_{HOME} ## [1] 0.06347507
```

## 4 Getting help

- 1. CRAN The Comprehensive R Archive Network (http://cran.r-project.org/). About ~4000 packages and data available for use.
- 2. R website (http://www.r-project.org/) and FAQs (http://cran.r-project.org/doc/FAQ/R-FAQ.html)
- 3. R related projects are difficult to search using Google. Use R-specific search engine RSEEK (http://www.rseek.org/).
- 4. A very good guide to R is "R Inferno" (http://www.burns-stat.com/pages/Tutor/R inferno.pdf).
- 5. Hadley Wickham's "Advanced R" (http://adv-r.had.co.nz/).

#### 4.1 Some useful online documentation

- 1. An introduction to R (http://cran.r-project.org/doc/manuals/r-release/R-intro.pdf)
- 2. R Data Import/Export (http://cran.r-project.org/doc/manuals/r-release/R-data.pdf)

A full list of such manuals can be found at http://cran.r-project.org/manuals.html.

#### 4.2 Finding help inside R

- 1. help(object) will show the help page of any object including function in R.
- 2. ?object is same thing as above.
- 3. help.search("string") will search for a string pattern in all help files.
- 4. ??string is the same thing as above
- 5. help(package="packagename") will show the start help page of a package.

#### 4.3 R for bioinformatics

- 1. **Bioconductor project http://www.bioconductor.org/** A set of R modules specifically meant for biological data analysis. Distributed separately from CRAN.
- 2. Some biological data related module in CRAN, such as ape, a phylogenetic analysis package. Or, seqinr, a basic sequence analysis package. A brief introduction to using R for bioinformatics can be found here.

### 5 Installing packages in R

install.packages('Packagename') will install any package in R. To use the package you need to use library('Packagename').

## 6 Assignment

Assignment operator in R is <-.

```
x<- 1:20
x
## [1] 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20
```

### 7 Basic data types in R

- 1. character
- 2. numeric All numbers in R are double precision real numbers. If you want an integer, you need to specify L suffix. 1 is numeric, but 1L is integer.
- 3. integer
- 4. complex
- 5. logical (True/False)

Undefined data in R is represented as either 'NaN' or 'NA' types. Infinity is represented as Inf.

#### 8 R data structures

#### 8.1 Vectors

- 1. Basic data structures in R.
- 2. All elements are of same type.
- 3. typeof() returns data-type.
- 4. length() returns the number of elements.

#### 8.1.1 Creating vectors

```
x <- vector()  # Empty vector
x <- c(1,2,3,4)  # Using c()
x <- 1:10  # Using ":" operator
x <- seq(1,10,by=1) # Using seq() funtion</pre>
```

#### 8.1.2 Accessing elements of a vector

Elements of vector in R starts with 1, rather than 0 like in other languages.

```
## [1] 1 2 3 4 5 6 7 8 9 10
x[3] # Access the 3rd element
## [1] 3
x[-1] # Access everything except the first element
## [1] 2 3 4 5 6 7 8 9 10
8.1.3 Modify vector
## [1] 1 2 3 4 5 6 7 8 9 10
x[2] \leftarrow 0; x # Modify 2nd element
## [1] 1 0 3 4 5 6 7 8 9 10
x[x > 5] \leftarrow 1; x # Modify elements greater than 5
## [1] 1 0 3 4 5 1 1 1 1 1
x \leftarrow c(1,2); y \leftarrow c(2,4)
z \leftarrow c(x,y) # Joining two vectors
## [1] 1 2 2 4
8.1.4 Converting vector of one type to another
Use as.x() funtions.
x <- 0:6
as.numeric(x)
## [1] 0 1 2 3 4 5 6
as.logical(x)
## [1] FALSE TRUE TRUE TRUE TRUE TRUE
as.character(x)
## [1] "0" "1" "2" "3" "4" "5" "6"
as.complex(x)
## [1] 0+0i 1+0i 2+0i 3+0i 4+0i 5+0i 6+0i
8.2 Lists
  1. Contains mixed data type.
```

- 2. typeof() returns "list".
- 3. length() returns length.

#### 8.2.1 Creating list

```
x <- list() # Empty list
x \leftarrow list ("a" = 2.5, "b" = TRUE, "c" = 1:3) # Mixed type list
## $a
## [1] 2.5
##
## $b
## [1] TRUE
## $c
## [1] 1 2 3
typeof(x)
## [1] "list"
length(x)
## [1] 3
8.2.2 Access elements of a list
x$a # Access using name
## [1] 2.5
x[c(1:2)] # Using index
## $a
## [1] 2.5
##
## $b
## [1] TRUE
x[["b"]] # Using name
## [1] TRUE
8.2.3 Modify lists
x[["d"]] \leftarrow 25
8.3 Matrix
Special type of vector with "dimension" attributes.
m<-matrix (nrow=2, ncol=3)</pre>
dim(m)
## [1] 2 3
attributes(m)
## $dim
## [1] 2 3
```

Matrix is constructed column-wise.

```
m<-matrix(1:6, nrow=2, ncol=3)</pre>
        [,1] [,2] [,3]
##
## [1,]
           1
              3
## [2,]
            2
                 4
m <- 1:10
\dim(m) < -c(2,5)
        [,1] [,2] [,3] [,4] [,5]
##
## [1,]
           1
                 3
                      5
                            7
## [2,]
            2
                 4
                      6
                            8
                                10
You can create matrix out of vectors using column-binding or row-binding:
y <- 10:12
cbind(x,y)
##
        х у
## [1,] 1 10
## [2,] 2 11
## [3,] 3 12
rbind(x,y)
##
     [,1] [,2] [,3]
## x
             2
       1
                   3
## y 10
            11
                  12
8.4 Factor
Categorical data. Can be ordered or unordered.
x<- factor (c("yes","yes", "no"))</pre>
Factors have levels:
table(x)
## x
## no yes
    1
Factors are basically numerical data under the hood. Each string is given a number. We can view the
underlying assignment using this:
unclass(x)
```

```
## [1] 2 2 1
## attr(,"levels")
## [1] "no" "yes"
```

Explicit ordering can be specified using levels arguments to factor. If we want to make no before yes:

```
x<- factor (c("yes", "no"), levels=c("Yes", "no"))</pre>
```

#### 8.5 Missing values

Undefined values in R are represented by "NaN" and "NA". NA values can have class of numeric, integer, etc. NaN is also NA but the converse is not true. There are two functions to find NaN and NA is R:

```
is.na()
is.nan()

x<- c(1,2, NA, 10, 3)
is.na(x)

## [1] FALSE FALSE TRUE FALSE FALSE
is.nan(x)

## [1] FALSE FALSE FALSE FALSE FALSE

x<- c(1,2,NaN,NA,4)
is.na(x)

## [1] FALSE FALSE TRUE TRUE FALSE
is.nan(x)</pre>
```

#### 8.6 Data frames

Key data type in R. It stores tabular data. Column of a data frame can have different types of data, unlike matrix (same type). Special attributes row.names. To read use read.table() or read.csv(). Data frame can be converted to matrix data.matrix().

```
x <-data.frame (foo=1:4, bar=c(T,T,F,F))
х
##
     foo
           bar
## 1
         TRUE
       1
## 2
       2 TRUE
## 3
       3 FALSE
## 4
       4 FALSE
nrow(x)
## [1] 4
ncol(x)
## [1] 2
```

#### 8.6.1 Reading and writing data-frame

```
data(iris) # Read iris data sets
head(iris) # View the first few line of the iris data
## Sepal.Length Sepal.Width Petal.Length Petal.Width Species
```

```
## 1
                           3.5
              5.1
                                        1.4
                                                     0.2 setosa
## 2
              4.9
                           3.0
                                        1.4
                                                     0.2 setosa
## 3
              4.7
                           3.2
                                        1.3
                                                     0.2 setosa
## 4
              4.6
                           3.1
                                        1.5
                                                     0.2 setosa
## 5
              5.0
                           3.6
                                                     0.2 setosa
                                        1.4
              5.4
                                        1.7
## 6
                           3.9
                                                     0.4 setosa
```

```
write.table(iris,"iris.txt",quote=F) # Write a sample file out
dir() # Check that we have written a file
## [1] "airquality.pdf" "Intro_to_R.pdf" "Intro_to_R.Rmd" "iris.txt"
## [5] "practicals.Rmd" "source_test.R"
test_data<-read.table("iris.txt", header=T) # read the data back into test_data
head(test_data) # Check the first few lines of the data
##
     Sepal.Length Sepal.Width Petal.Length Petal.Width Species
## 1
              5.1
                           3.5
                                        1.4
                                                     0.2 setosa
## 2
              4.9
                           3.0
                                                     0.2 setosa
                                        1.4
## 3
              4.7
                           3.2
                                        1.3
                                                     0.2
                                                          setosa
## 4
              4.6
                           3.1
                                        1.5
                                                     0.2 setosa
## 5
              5.0
                           3.6
                                        1.4
                                                     0.2 setosa
              5.4
                           3.9
## 6
                                        1.7
                                                     0.4 setosa
To read a data frame from a gzipped file:
d<-read.table(gzfile("myzipped.gz"))</pre>
```

### 9 Names

Each type of of data in R can have names. But they are most important for Matrix and Data-frames.

```
m<- matrix(1:4, nrow=2, ncol=2)</pre>
dimnames(m) \leftarrow list (c("a","b"), c("c","d"))
##
     c d
## a 1 3
## b 2 4
d < -data.frame(c(1,2,3),c(4,5,6))
##
     c.1..2..3. c.4..5..6.
## 1
               1
## 2
                2
                            5
## 3
               3
                            6
names(d)<- c("A","B")
##
     A B
## 1 1 4
## 2 2 5
## 3 3 6
```

## 10 Subsetting

Subsetting is a way to select a subset of data. There are 3 basic ways to subset data in R:

- 1. []
- 2. [[]]
- 3. \$

#### 10.1 Subsetting vector

```
x<-c ("a","b", "c")
x[1]
## [1] "a"
x[2]
## [1] "b"
x[1:3]
## [1] "a" "b" "c"
x[-2]
## [1] "a" "c"
x[x > "a"]
## [1] "b" "c"
u <- x > "a"
## [1] FALSE TRUE TRUE
x[u]
## [1] "b" "c"
10.2 Subsetting list
x <- list (foo=1:4, bar=0.6)</pre>
x[1]
## $foo
## [1] 1 2 3 4
x[[1]]
## [1] 1 2 3 4
x$bar
## [1] 0.6
x[["bar"]]
## [1] 0.6
x["bar"]
## $bar
## [1] 0.6
10.3 Removing missing values
x < c(1,2, NA, 4, NA, 5)
bad <- is.na(x)</pre>
x[!bad]
```

```
## [1] 1 2 4 5
```

#### 10.4 Matrix subsetting

```
x<- matrix (1:6, 2, 3) # Create a sample 2x3 matrix
x # print the matrix out
##
        [,1] [,2] [,3]
## [1,]
           1
                3
## [2,]
           2
x[1,2] # what is the value in row 1, column 2
## [1] 3
x[2,1] # what is the value in row 2, column 1
## [1] 2
We can select the whole row or columns:
x[1,] # select whole row 1
## [1] 1 3 5
x[,2] # select whole column 2
## [1] 3 4
```

#### 10.5 Subsetting data-frame

All the operations of matrix also work with data-frame. However, you will more often use \$ to extract individual columns of a data-frame.

```
x<-data.frame(c(1:3),c(4:6)) # create a small data frame
names(x)<-c("A","B") # Give each column a nice name
x

## A B
## 1 1 4
## 2 2 5
## 3 3 6

x$A # Extract the column A
## [1] 1 2 3</pre>
```

## 11 Vectorized operations

Instead of using loops, you should all the time try to use "vectorized" operations. That means if you are interested in operations performed on each element of a "collection", you should use the collection as a whole, not individual elements. This gives the computer to parallelize the operation and usually results in faster runtime.

```
x<-1:4; y<-6:9
x+y
```

```
## [1] 7 9 11 13
```

```
x-y

## [1] -5 -5 -5 -5

x*y

## [1] 6 14 24 36

x/y
```

#### ## [1] 0.1666667 0.2857143 0.3750000 0.4444444

When you multiply two matrices using the "\*", it multiplies element by element multiplications. This is different that standard matrix multiplication. If you are interested standard matrix multiplication, use "%\*%" operator.

```
m1<-matrix(1:4,2,2)
m1
##
        [,1] [,2]
## [1,]
           1
## [2,]
m1*m1
        [,1] [,2]
## [1,]
           1
## [2,]
           4
               16
m1 %*% m1
        [,1] [,2]
##
## [1,]
           7
               15
## [2,]
          10
               22
```

## 12 Some programming concepts in R

#### 12.1 Functions

```
myfunction<-function() {
    cat("Hello world\n");
}</pre>
```

#### 12.2 For loop and if condition

```
for (i in 1:3) {
    if (i > 1) {
        cat("I said hello", i, "times\n")
    }else {
        cat("I said hello", i, "time\n")
    }
}

## I said hello 1 time
## I said hello 2 times
## I said hello 3 times
```

### 12.3 Reading a file line by line in R

```
fh <- file( inputfile, open="r" )
while (length( line <- readLines( fh, n=1, warn= FALSE ) ) > 0) {
    # Do something with the line
}
close(fh)
```

#### 12.4 Regular expression in R

Some useful functions that can be used with regular expression:

- 1. grep() finds a pattern in a vector or character.
- 2. sub() substitute a text for the first occurance.
- 3. gsub() substitute every occurance.
- 4. regexpr() find and extracts values using a pattern.
- 5. regexec() find and extracts values. Usefule for the function regmatches()

```
text <- "gi|123456|ref|ABCDEFG" # Text to search
p <- "gi\\|(\\d+)\\|ref\\|(\\S+)" # Extract gi and acc
m <- regexec(p, text, perl = TRUE) # Find match and extract
s <- regmatches(text, m) # Extract the substring
s[[1]][2] # Print GI

## [1] "123456"
s[[1]][3] # Print accession
## [1] "ABCDEFG"</pre>
```

## 13 Summary statistics

```
x<-rnorm(50)
mean(x)

## [1] -0.0469437

sd(x)

## [1] 1.037544

var(x)

## [1] 1.076498

median(x)

## [1] -0.004295971

sum(x)

## [1] -2.347185

You need to skip NA for doing summary statistics:
data(airquality)
mean(airquality$0zone)</pre>
```

## [1] NA

```
mean(airquality$0zone, na.rm=T)
```

#### ## [1] 42.12931

summary() will give you a nice summary of data:

```
summary(airquality)
```

```
##
        Ozone
                          Solar.R
                                             Wind
                                                               Temp
##
    Min.
           : 1.00
                             : 7.0
                                               : 1.700
                                                                  :56.00
                      Min.
                                       Min.
                                                          Min.
    1st Qu.: 18.00
                      1st Qu.:115.8
                                       1st Qu.: 7.400
                                                          1st Qu.:72.00
    Median : 31.50
                      Median :205.0
                                       Median : 9.700
                                                          Median :79.00
##
           : 42.13
##
    Mean
                      Mean
                              :185.9
                                       Mean
                                               : 9.958
                                                          Mean
                                                                 :77.88
##
    3rd Qu.: 63.25
                      3rd Qu.:258.8
                                       3rd Qu.:11.500
                                                          3rd Qu.:85.00
##
    Max.
            :168.00
                      Max.
                              :334.0
                                       Max.
                                               :20.700
                                                          Max.
                                                                 :97.00
    NA's
                      NA's
##
           :37
                              :7
##
        Month
                          Day
##
            :5.000
   \mathtt{Min}.
                     Min.
                             : 1.0
    1st Qu.:6.000
                     1st Qu.: 8.0
   Median :7.000
                     Median:16.0
##
##
    Mean
            :6.993
                     Mean
                             :15.8
##
    3rd Qu.:8.000
                     3rd Qu.:23.0
##
            :9.000
                             :31.0
    Max.
                     Max.
##
```

#### 14 Random numbers

We can generate random numbers from the normal distribution using rnorm() function.

```
x<- rnorm(10)
x
## [1] -0.9256206 1.5633823 0.7495711 0.1904426 -0.5005065 -0.5616861
## [7] 1.0890480 -0.1941938 0.2313796 2.0793260
```

If we are interested in generating random numbers using a particular mean and standard deviation, we should you this:

```
x<-rnorm(10, mean=5, sd=2)
x
```

```
## [1] 7.276876 2.247866 3.093904 3.212064 4.216682 2.393955 6.212271 6.600028
## [9] 4.031995 3.216107
```

In R, you can sample other distributions, I will leave it up to you to explore.

## 15 Plotting in R

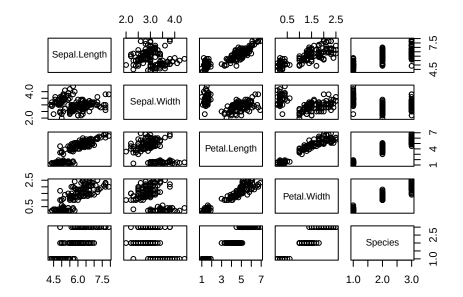
There are 3 graphical libraries in R:

- 1. Base graphics
- 2. Lattice
- 3. gglplot2()

In this tutorial we will discuss only base graphics.

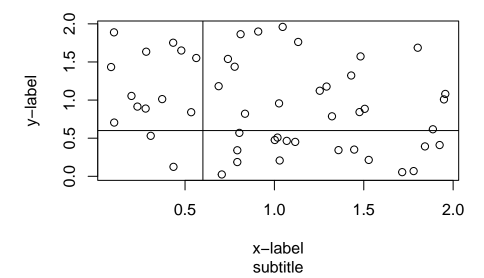
#### 15.1 Plot

plot function is pretty intelligent. If you just give a dataframe, it will plot a graph with all vs all variables.
data(iris)
plot(iris)



```
x<-runif(50,0,2) # create a vector from uniform distribution
y<-runif (50,0,2) # create another one
plot(x,y, main="Main title",sub="subtitle",xlab="x-label", ylab="y-label")
abline(h=0.6,v=0.6)</pre>
```

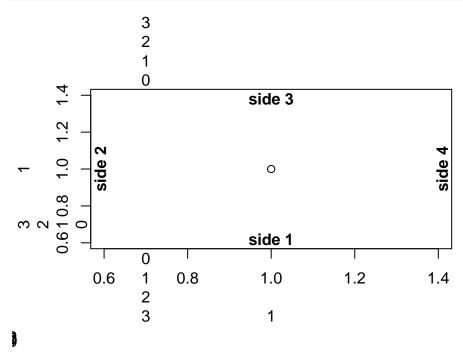
## Main title



Each side of the plot

have specific margins.

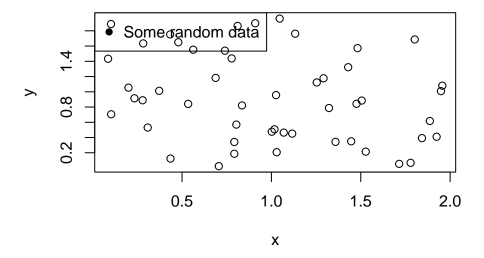
```
plot(1,1)
for (side in 0:3) mtext (0:3,side=side,at=0.7,line=0:3)
mtext(paste("side",1:4),side=1:4,line = -1, font=2)
```



#### 15.1.1 Building a plot from pieces

```
plot (x, y, type="n", xlab="",ylab="", axes=F) # Draw an empty plot area
points(x,y) # Draw point
axis(1) # Draw the first axis
axis(2, at=seq(0.2,1.8,0.2)) # Draw second axis with tick at specific scale
box() # Draw the surrounding box
title(main="Main title",xlab="x",ylab="y") # Write the main title and the labels
legend("topleft",legend="Some random data",pch=16)
```

#### Main title



#### 15.1.2 Plotting characters

pchShow() will show you all the plotting characters.

#### 15.2 Drawing devices in R

Normally when you create a plot, it is drawn on the default computer screen. But you can create all sorts of images, such PDF (good for high resolution publication quality graphs), or bitmap images such as PNG. In all cases, the steps are identical. You open a device, draw your plot and close the device. For e.g., you can create a PDF drawing like this.

```
data(airquality)
pdf("airquality.pdf",width=8.5,height=11)
plot(airquality)
dev.off()
## pdf
## 2
```

### 15.3 par

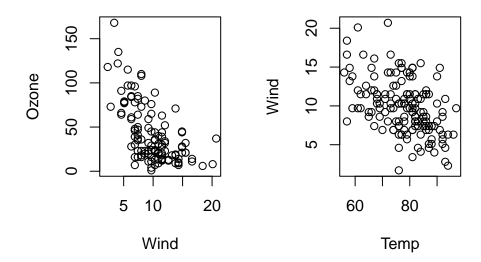
par function controls every parameter of a plot. The function has several parameters. But the most common ones are as follows:

- 1. pch- point type used for plotting
- 2. lty Line type
- 3. lwd line width
- 4. cex character magnification. It has several variations for different regions of the plot.
- 5. las rotation of the axis labels
- 6. bg background color of the plot
- 7. mar margin of the plot
- 8. oma outer margin of the plot
- 9. mfrow draws multiple plots

### 15.4 Drawing multiplots

You can modify mfrow in par(). For e.g, to draw a 1 row two column plot,

```
data(airquality)
par(mfrow=c(1,2))
with(airquality, {plot(Wind,Ozone);plot(Temp,Wind)})
```



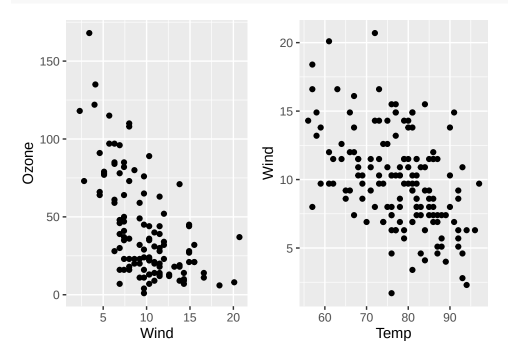
### 15.5 Multiplots in ggplot

There are four ways to draw the multiplots in ggplot2: - grid.arrange() [gridExtra package] - plot\_grid() [cowplot package] - plot\_layout() [patchwork package] - ggarrange() [ggpubr package]

```
library(ggplot2)
```

## Warning: Removed 37 rows containing missing values (geom\_point).

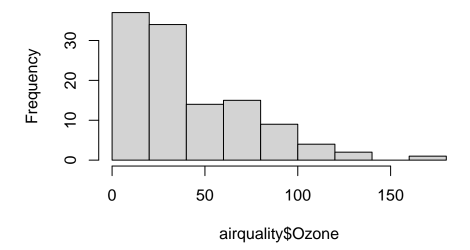
### print(combined\_plot)



## 15.6 Histograms

data("airquality")
hist(airquality\$0zone)

# Histogram of airquality\$Ozone

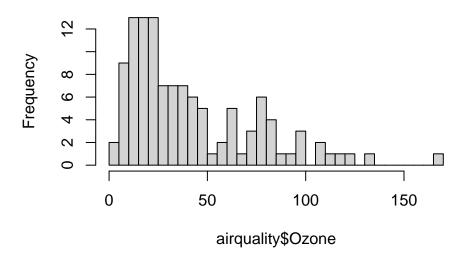


No of breaks in the his-

togram can be controlled:

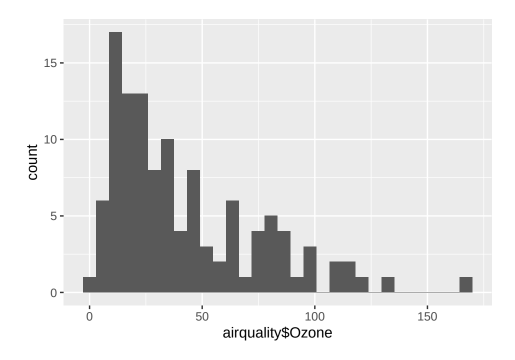
hist(airquality\$0zone, breaks=50)

# Histogram of airquality\$Ozone



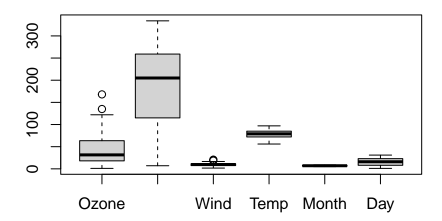
## 15.7 Histogram in ggplot

```
library(ggplot2)
g <- ggplot(data = airquality, aes(x=airquality$0zone)) +
    geom_histogram()
g
## Warning: Use of `airquality$0zone` is discouraged. Use `0zone` instead.
## `stat_bin()` using `bins = 30`. Pick better value with `binwidth`.
## Warning: Removed 37 rows containing non-finite values (stat_bin).</pre>
```



## 15.8 Boxplots

Boxplot is an summary plot showing the median and the distributions of data. You can give it a data frame. boxplot(airquality)

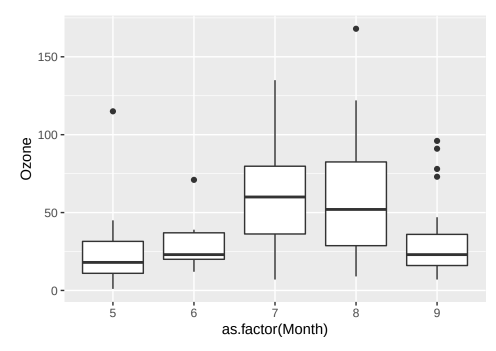


## 15.9 Boxplots in ggplot

```
g <- ggplot(airquality, aes(x=as.factor(Month), y=Ozone)) +
    geom_boxplot()</pre>
```

g

## Warning: Removed 37 rows containing non-finite values (stat\_boxplot).

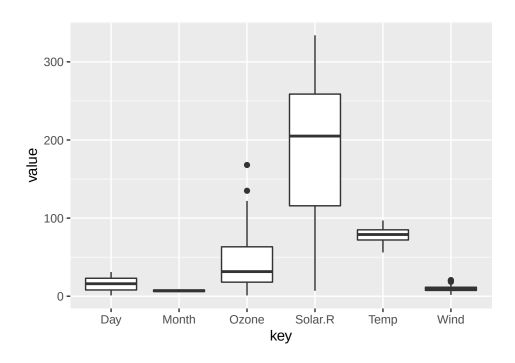


To plot the previous

plot, we need combine the entire dataframe into two variables.

```
library(tidyr)
gathered_data <- gather(airquality)
g <- ggplot(gathered_data, aes(x=key,y=value)) +
    geom_boxplot()
g</pre>
```

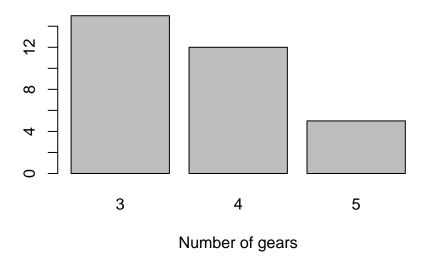
## Warning: Removed 44 rows containing non-finite values (stat\_boxplot).



## 15.10 Barplots

```
data(mtcars)
head(mtcars)
##
                      mpg cyl disp hp drat
                                               wt qsec vs am gear carb
## Mazda RX4
                     21.0
                            6 160 110 3.90 2.620 16.46
                            6 160 110 3.90 2.875 17.02
## Mazda RX4 Wag
                     21.0
                                                                      4
## Datsun 710
                     22.8
                            4 108
                                   93 3.85 2.320 18.61
## Hornet 4 Drive
                     21.4
                            6 258 110 3.08 3.215 19.44
                                                                      1
                                                                      2
## Hornet Sportabout 18.7
                               360 175 3.15 3.440 17.02
## Valiant
                     18.1
                            6 225 105 2.76 3.460 20.22
counts<-table(mtcars$gear)</pre>
barplot(counts, main="Car number of gear distribution", xlab="Number of gears")
```

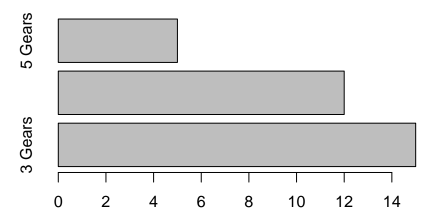
## Car number of gear distribution



You can make a horizontal bar plot.

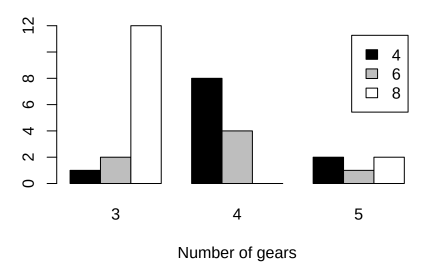
```
barplot(counts, main="Car distribution", horiz=T, names.arg=c("3 Gears", "4 Gears", "5 Gears"))
```

## **Car distribution**

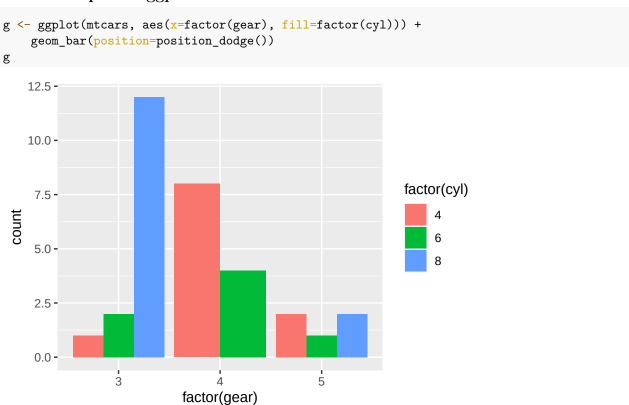


You can draw a grouped bar plot like this.

# Gear and cylinder



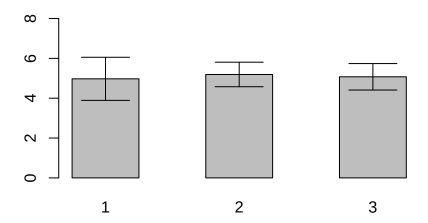
## 15.11 Barplot in ggplot



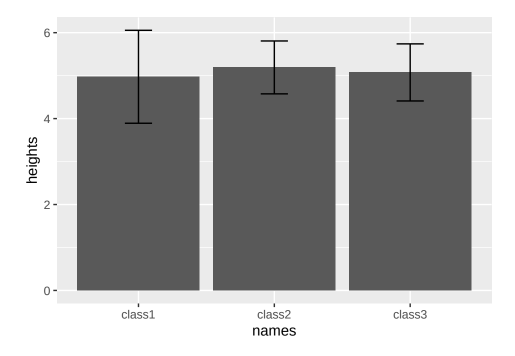
## 15.12 Barplots with error bars

There is no inbuilt way to generate error bars in R. But we have to hack a solution. We will use arrows to generate error bars. The trick is arrows take an argument of angle. If you give angle=90 it will generate

an error bar.



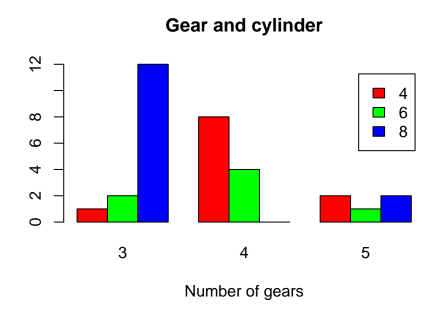
#### 15.13 Errorbars in ggplot



## 16 Colors in R

If you have drawn figures in Excel, you might have used colors like this

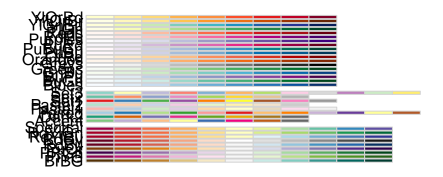
barplot(counts, main="Gear and cylinder", xlab="Number of gears", col=c("red", "green", "blue"), legend=rounder", xlab="Number of gears", col=c("red", "green", "blue"), legend=rounder of gears of gears



This is what I call

"Christmas" plot. You should never use color like this. In R you can use sophisticated color palette.

library(RColorBrewer)
display.brewer.all()

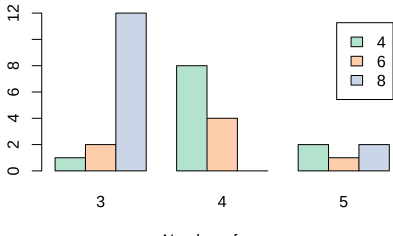


There are 3 palettes: se-

quential, quantitative, and contrasting. Now choose a nice color for our plot.

```
nice<-brewer.pal(3,"Pastel2")
barplot(counts, main="Gear and cylinder", xlab="Number of gears", col=nice,legend=rownames(counts),besi</pre>
```

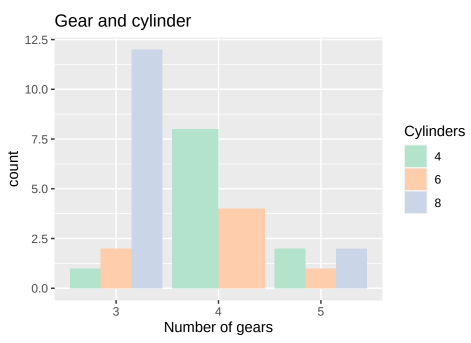
## Gear and cylinder



Number of gears

## 16.1 Colors in ggplot





## 17 Linear regression and correlation

#### 17.1 Linear regression

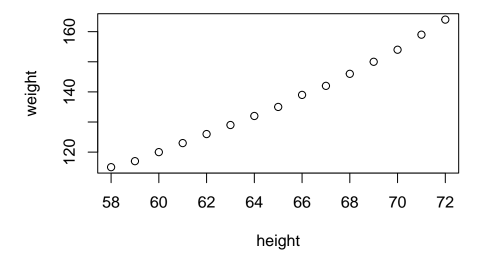
We will use the women datasets in R for this example. These is a simple datasets with "weight" and "height" of women in the US.

```
data(women)
head(women)
```

##		height	weight
##	1	58	115
##	2	59	117
##	3	60	120
##	4	61	123
##	5	62	126
##	6	63	129

If we plot a simple scatterplot of this data, we will see there is an obvious correlation between this two variables.

plot(women)



Let's do a linear regression between this two variables.

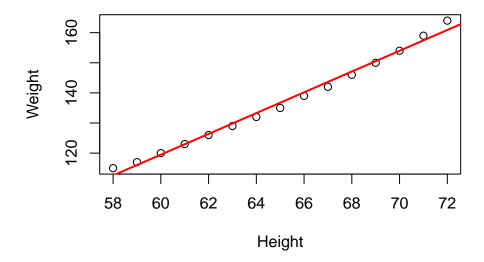
```
1 <- lm(women$weight~women$height)
summary(1)</pre>
```

```
##
## Call:
## lm(formula = women$weight ~ women$height)
##
## Residuals:
##
      Min
                1Q Median
                               3Q
                                      Max
  -1.7333 -1.1333 -0.3833
                           0.7417
##
##
  Coefficients:
##
                Estimate Std. Error t value Pr(>|t|)
## (Intercept) -87.51667
                             5.93694
                                     -14.74 1.71e-09 ***
                                      37.85 1.09e-14 ***
## women$height
                 3.45000
                             0.09114
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 1.525 on 13 degrees of freedom
## Multiple R-squared: 0.991, Adjusted R-squared: 0.9903
## F-statistic: 1433 on 1 and 13 DF, p-value: 1.091e-14
```

You can safely ignore the (Intercept) line. But the adjusted r<sup>2</sup> is 0.99 with P-value 1.1e-14, which is highly significant. Let's put the regression line on the plot.

```
plot(women$height,women$weight,main="Women height vs weight",xlab="Height",ylab="Weight")
abline(1,lwd=2,col="red")
```

# Women height vs weight



You can extract the P-value and the  ${\bf r}^2$  values for printing

```
s<-summary(1)
s$adj.r.squared
## [1] 0.9903183</pre>
```

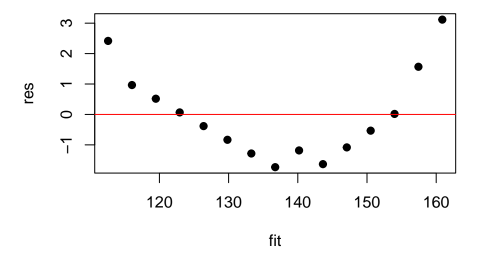
s\$coefficients[2,4] ## [1] 1.090973e-14

#### 17.1.1 Residuals

One way to find out whether a linear model is a correct fit or not is to actually plot the residuals.

```
res <- resid(1)
fit <- fitted(1)

plot(fit, res, pch=19)
abline (0,0, col="red")</pre>
```

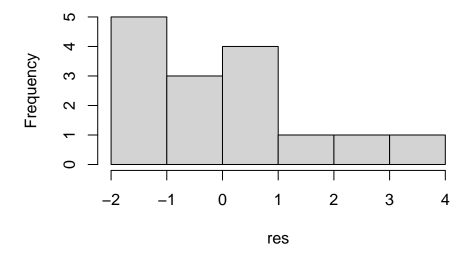


**Homoscedasticiy** is when residuals are equally distributed above and below the horizontal line. **Heteroscedasticity** is there is a bias. For this time looks like scedasticity is pretty good.

The other thing you should check is the whether the residuals are distributed normally.

hist (res)

## Histogram of res



In this case it fit does not look very good. However, our dataset is pretty small. The normality checking should be reliable only for larger dataset.

#### 17.2 Generalized linear model

The test for comparing means between two groups is t.test. The test for comparing means among two or more groups is anova. And we have already seen linear regression. All these are just subclasses of *Generalized Linear Model*. We will not discuss this in details in this course. However, just to show that are all the same see the following code.

First we will do the t.test using iris dataset.

```
data("iris")
iris$Species <- as.character(iris$Species)</pre>
two_species_iris <- iris[iris$Species == "setosa" | iris$Species=="virginica",]
t.test(Petal.Length~Species, data=two_species_iris)
##
   Welch Two Sample t-test
##
## data: Petal.Length by Species
## t = -49.986, df = 58.609, p-value < 2.2e-16
## alternative hypothesis: true difference in means between group setosa and group virginica is not equ
## 95 percent confidence interval:
## -4.253749 -3.926251
## sample estimates:
##
      mean in group setosa mean in group virginica
##
                     1.462
                                             5.552
Note the p-vale above.
summary(aov(Petal.Length~Species, data=two_species_iris))
               Df Sum Sq Mean Sq F value Pr(>F)
                1 418.2
                           418.2
                                    2499 <2e-16 ***
## Species
## Residuals
                    16.4
                             0.2
               98
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
P-value is exactly the same.
summary(lm(Petal.Length~Species,data=two_species_iris))
##
## Call:
## lm(formula = Petal.Length ~ Species, data = two species iris)
##
## Residuals:
##
      Min
                1Q Median
                                3Q
                                       Max
## -1.0520 -0.1620 0.0380 0.1405 1.3480
##
## Coefficients:
##
                    Estimate Std. Error t value Pr(>|t|)
## (Intercept)
                     1.46200
                                0.05786
                                          25.27
                                                  <2e-16 ***
## Speciesvirginica 4.09000
                                0.08182
                                          49.99
                                                  <2e-16 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.4091 on 98 degrees of freedom
## Multiple R-squared: 0.9623, Adjusted R-squared: 0.9619
```

```
## F-statistic: 2499 on 1 and 98 DF, p-value: < 2.2e-16
```

A through introduction to GLM is beyond the scope of of this course. But to show that regression is just and variation of GLM we can use glm function instead of lm:

```
summary(glm(women$weight~women$height, family = gaussian))
```

```
## Call:
## glm(formula = women$weight ~ women$height, family = gaussian)
## Deviance Residuals:
##
      Min
                 1Q
                     Median
                                  3Q
                                          Max
## -1.7333 -1.1333 -0.3833
                              0.7417
                                       3.1167
##
## Coefficients:
                Estimate Std. Error t value Pr(>|t|)
##
## (Intercept) -87.51667
                            5.93694 -14.74 1.71e-09 ***
## women$height
                 3.45000
                            0.09114
                                      37.85 1.09e-14 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for gaussian family taken to be 2.325641)
##
##
      Null deviance: 3362.933 on 14 degrees of freedom
## Residual deviance:
                       30.233 on 13 degrees of freedom
## AIC: 59.082
##
## Number of Fisher Scoring iterations: 2
```

It is identical to the lm result found before.

#### 17.2.1 Predict using linear model

```
petal_model <- lm(Petal.Length~Petal.Width, data = iris)</pre>
summary(petal_model)
##
## Call:
## lm(formula = Petal.Length ~ Petal.Width, data = iris)
## Residuals:
##
       Min
                 1Q
                     Median
                                    30
## -1.33542 -0.30347 -0.02955 0.25776 1.39453
## Coefficients:
##
              Estimate Std. Error t value Pr(>|t|)
## (Intercept) 1.08356
                          0.07297
                                     14.85
                                            <2e-16 ***
## Petal.Width 2.22994
                          0.05140
                                     43.39
                                            <2e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## Residual standard error: 0.4782 on 148 degrees of freedom
## Multiple R-squared: 0.9271, Adjusted R-squared: 0.9266
## F-statistic: 1882 on 1 and 148 DF, p-value: < 2.2e-16
```

```
predict(petal_model, data.frame(Petal.Width=c(0.7)))
## 2.644516
17.3 Correlations
We can also do a correlation between this two variables
s<-cor.test(women$weight,women$height,alternatives="tow.sided")</pre>
##
## Pearson's product-moment correlation
##
## data: women$weight and women$height
## t = 37.855, df = 13, p-value = 1.091e-14
\#\# alternative hypothesis: true correlation is not equal to 0
## 95 percent confidence interval:
## 0.9860970 0.9985447
## sample estimates:
##
         cor
## 0.9954948
s$p.value
## [1] 1.090973e-14
s$estimate
##
## 0.9954948
```