## Getting to plot in ${\bf R}$

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## Plotting in **R**

- Introduction

### Why plot in **R**?

- Powerful (large range of plot types)
- Fully customizable (make your own style)
- Practical (integrate your plots and your code together)

# Graphics paradigms in R

They are three dominant graphics paradigms in R:

- traditional graphics (based on graphics)
- lattice (based on grid)
- ggplot2 (based on grid)

# Graphics paradigms in R

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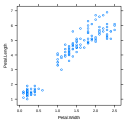
#### Note:

- graphics and grid are part of any basic installation of R
- lattice is part of the so-called list of CRAN recommended packages
- ggplot2 is part of the tidyverse universe (from RStudio)
- we will focus on traditional graphics and ggplot2, but lattice is excellent too!
- some other packages are sometimes useful too (e.g. rgl, plotly)

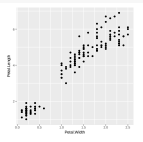
# An example (using default settings)

```
plot(Petal.Length ~ Petal.Width, data = iris)
```

```
library(lattice)
xyplot(Petal.Length ~ Petal.Width,
data = iris)
```



```
library(ggplot2)
ggplot(data = iris,
   aes(x = Petal.Width, y = Petal.Length)) +
geom_point()
```



Petal.Width

2.0

# How to learn on your own?

1. Check the examples readily available in R, e.g.

```
demo(graphics)
demo(image)
demo(persp)
demo(colors)
demo(plotmath)
demo(Hershey)
example(plot)
example(boxplot)
example(hist)
example(bartplot)
browseVignettes(package = "ggplot2")
```

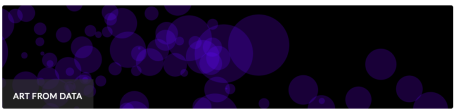
## How to learn on your own?

### 2. Scroll the web:

(e.g. http://www.r-graph-gallery.com/all-graphs/)



HOME GGPLOT2 ALL GRAPHS BLOG ABOUT PYTHON



ALL GRAPHS













If you are looking for something in particular, please use the search tool below. It works even if you are looking for informations concerning an R graph function that is used in the website.

Type an R function, graph type, graph number...

Search ...

# How to learn on your own?

#### 3. Read books:



### Plotting in R

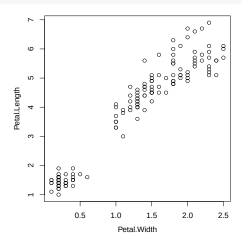
- Plotting with traditional graphics

## Plotting in **R**

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- Plotting with traditional graphics
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  - Aesthetics
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- Conclusion: traditional graphics vs. ggplot

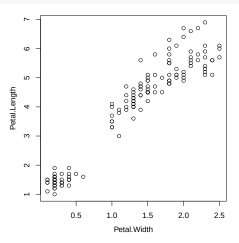
### In traditional graphics, use plot() to draw a scatter plot:

plot(Petal.Length ~ Petal.Width, data = iris)



You can choose what type of scatter plot to display with argument type:

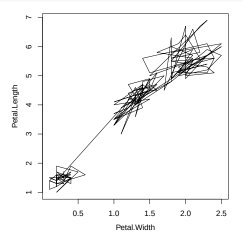
plot(Petal.Length ~ Petal.Width, data = iris, type = "p")



Note: see "?plot.default"

You can choose what type of scatter plot to display with argument type:

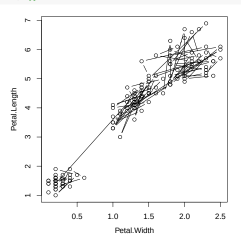
plot(Petal.Length ~ Petal.Width, data = iris, type = "1")



Note: it makes more sense when data are ordered...

You can choose what type of scatter plot to display with argument type:

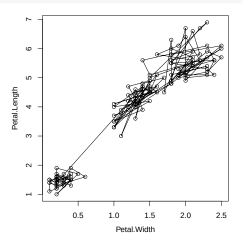
plot(Petal.Length ~ Petal.Width, data = iris, type = "b")



Note: it makes more sense when data are ordered...

You can choose what type of scatter plot to display with argument type:

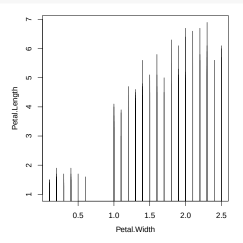
plot(Petal.Length ~ Petal.Width, data = iris, type = "o")



Note: it makes more sense when data are ordered...

You can choose what type of scatter plot to display with argument type:

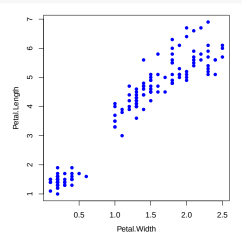
plot(Petal.Length ~ Petal.Width, data = iris, type = "h")



Note: it makes more sense when x-values are unique. . .

You can change point shapes (pch) and colour (col):

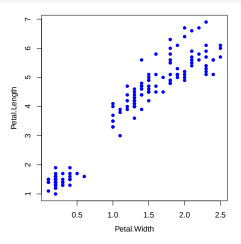
plot(Petal.Length ~ Petal.Width, data = iris, pch = 16, col = "blue")



Note: you can use colour names

You can change point shapes (pch) and colour (col):

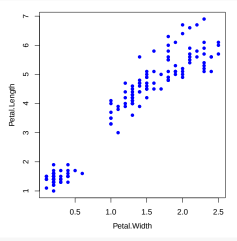
plot(Petal.Length ~ Petal.Width, data = iris, pch = 16, col = 4)



Note: you can use number of basic colours

### You can change point shapes (pch) and colour (col):

```
plot(Petal.Length ~ Petal.Width, data = iris, pch = 16, col = "#0000FFFF")
```

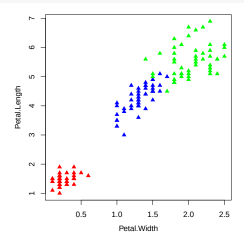


```
rgb(red = 0, green = 0, blue = 255, alpha = 255, maxColorValue = 255)
## [1] "#0000FFFF"
```

Note: you can have full control using hexadecimal!!

You can change point shapes (pch) and colour (col):

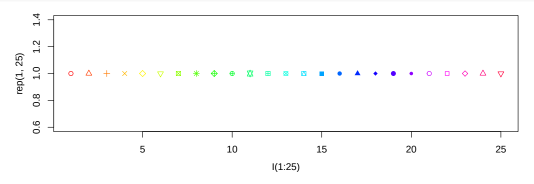
```
palette(c("red", "blue", "green"))
plot(Petal.Length ~ Petal.Width, data = iris, pch = 17, col = iris$Species)
```



Note: you can use a palette to match the levels of a factor

You can change point shapes (pch) and colour (col):

```
plot(rep(1, 25) ~ I(1:25), data = iris, pch = 1:25, col = rainbow(25))
```



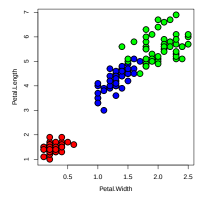
#### Note:

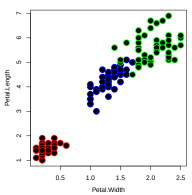
- there are 25 basic symbols (but other ways allow to use many more)
- check "?rainbow" for a list of different color palettes
- the I() allows for the creation of the vector before being interpreted by plot()

For many elements you can set both an outline colour (col) and background colour (bg):

```
palette(c("red", "blue", "green"))
plot(Petal.Length - Petal.Width, data = iris, cex = 2,
    pch = 21, col = "black", bg = iris$Species, lwd = 2)
```

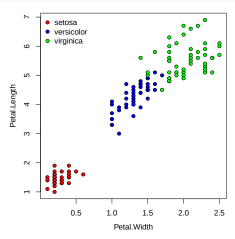
```
palette(c("red", "blue", "green"))
plot(Petal.Length ~ Petal.Width, data = iris, cex = 2,
    pch = 21, col = iris$Species, bg = "black", lwd = 2)
```





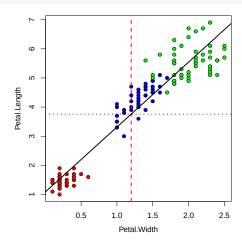
### Add a legend to make colours understandable:

```
palette(c("red", "blue", "green"))
plot(Petal.Length ~ Petal.Width, data = iris, pch = 21, bg = iris$Species)
legend(x = "topleft", legend = c("setosa", "versicolor", "virginica"), pch = 21, pt.bg = c("red", "blue", "green"), bty = "n")
```



### You can add lines to the plot:

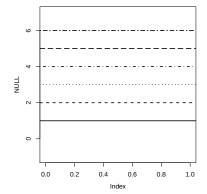
```
palette(c("red", "blue", "green"))
plot(Petal.Length ~ Petal.Width, data = iris, pch = 21, bg = iris$Species)
abline(v = mean(iris$Petal.Width), lty = 2, col = "red", lwd = 2)
abline(h = mean(iris$Petal.Length), lty = 3, col = "black", lwd = 2)
abline(a = 1.084, b = 2.23, lty = 1, lwd = 2)
```

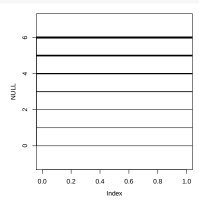


#### You can add lines to the plot:

```
plot(NULL, xlim = c(0, 1), ylim = c(-1, 7))
abline(h = 0, lty = 0, lwd = 2)
abline(h = 1, ltv = 1, lwd = 2)
abline(h = 2, lty = 2, lwd = 2)
abline(h = 3, lty = 3, lwd = 2)
abline(h = 4, ltv = 4, lwd = 2)
abline(h = 5, lty = 5, lwd = 2)
abline(h = 6, lty = 6, lwd = 2)
```

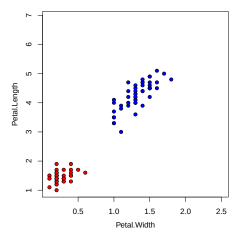
```
plot(NULL, xlim = c(0, 1), ylim = c(-1, 7))
abline(h = 0, lty = 1, lwd = 0.2)
abline(h = 1, ltv = 1, lwd = 0.5)
abline(h = 2, lty = 1, lwd = 1)
abline(h = 3, lty = 1, lwd = 2)
abline(h = 4, ltv = 1, lwd = 3)
abline(h = 5, lty = 1, lwd = 4)
abline(h = 6, lty = 1, lwd = 5)
```





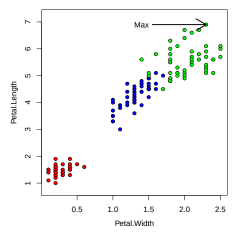
#### You can also add additional points to the plot:

```
versicolor <- subset(iris, Species == "versicolor")</pre>
           <- subset(iris, Species == "setosa")
plot(Petal.Length ~ Petal.Width, data = versicolor, pch = 21, bg = "blue", xlim = range(iris$Petal.Width), ylim = range(iris$Petal.Length))
points (Petal Length ~ Petal Width, data = setosa, pch = 21, bg = "red", xlim = range(iris$Petal Width), ylim = range(iris$Petal Length))
```



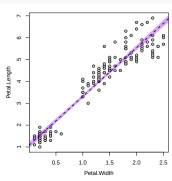
Note: You need to make sure the axis limits are the same!

### You can add text and arrows:



### Including an interval around a prediction line requires you to build a polygon:

```
test_mod <- lm(Petal.Length ~ Petal.Width, data = iris)
         <- data.frame(Petal.Width = seq(0, 3, length.out = 100))
newdat
         <- predict(test_mod, newdata = newdat, interval = "confidence")</pre>
pred
plot(Petal.Length ~ Petal.Width, data = iris, pch = 21, bg = "grey")
polygon(x = c(newdat$Petal.Width, rev(newdat$Petal.Width)),
       v = c(pred[, "lwr"], rev(pred[, "upr"])),
        col = alpha("purple", alpha = 0.4), ## alpha sets transparency!
        border = NA) ## removes black line around the polygon
abline(test_mod, lty = 2, lwd = 2)
```

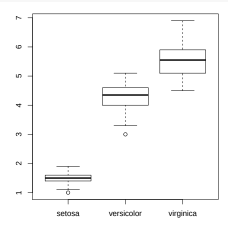


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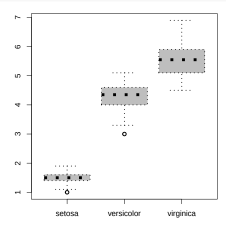
In traditional graphics, use boxplot() to draw a box plot:

boxplot(Petal.Length ~ Species, data = iris)



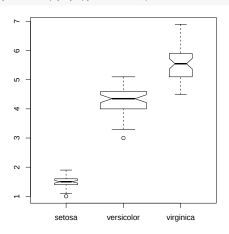
Many of the same changes made to scatter plots can be made here:

boxplot(Petal.Length ~ Species, data = iris, col = "grey", lwd = 2, lty = 3)



### There are also some boxplot specific arguments:

boxplot(Petal.Length ~ Species, data = iris, width = c(1, 2, 1), notch = TRUE)



### You can retrieve information by storing the output in an object:

```
my_boxcox <- boxplot(Petal.Length ~ Species, data = iris, plot = FALSE)
my_boxcox
## $stats
        [,1] [,2] [,3]
## [1,] 1.1 3.30 4.50
## [2,]
        1.4 4.00 5.10
## [3.] 1.5 4.35 5.55
## [4.] 1.6 4.60 5.90
## [5,] 1.9 5.10 6.90
##
## $n
## [1] 50 50 50
## $conf
            [,1]
                   [,2]
                              [,3]
## [1,] 1.455311 4.215933 5.371243
## [2,] 1.544689 4.484067 5.728757
##
## $out
## [1] 1 3
##
## $group
## [1] 1 2
## $names
## [1] "setosa" "versicolor" "virginica"
```

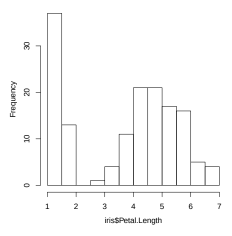
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# Traditional graphics: Histograms

### In traditional graphics, use hist() to draw an histogram:

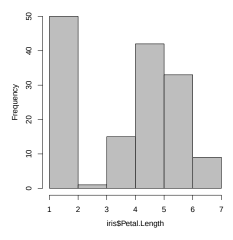
hist(iris\$Petal.Length, main = "") ## main used here to remove the automatic title



# Traditional graphics: Histograms

You can change the number and location of breaks between bins:

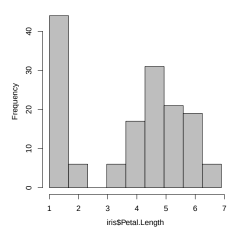
hist(iris\$Petal.Length, main = "", breaks = 5, col = "grey")



# Traditional graphics: Histograms

#### You can change the number and location of breaks between bins:

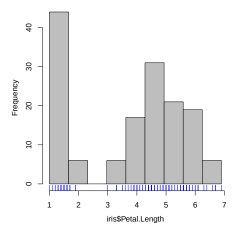
```
hist_breaks <- seq(min(iris$Petal.Length), max(iris$Petal.Length), length.out = 10)
hist(iris$Petal.Length, main = "", breaks = hist_breaks, col = "grey")
```



# Traditional graphics: Histograms

#### You can change the number and location of breaks between bins:

```
hist_breaks <- seq(min(iris$Petal.Length), max(iris$Petal.Length), length.out = 10)
hist(iris$Petal.Length, main = "", breaks = hist_breaks, col = "grey")
rug(x = iris$Petal.Length, col = "blue")</pre>
```



Note: it never hurts to add a rug under an histogram!

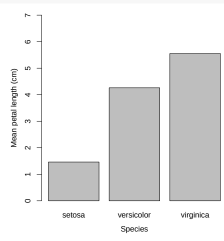
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```
spp_means <- data.frame(Species = c("setosa", "versicolor", "virginica"),</pre>
                        mean = as.numeric(by(iris$Petal.Length, iris$Species, mean)),
                        SE = as.numeric(by(iris$Petal.Length, iris$Species, function(x)sd(x)/sqrt(length(x))))
spp_means
        Species mean
         setosa 1.462 0.02455980
## 2 versicolor 4.260 0.06645545
## 3 virginica 5.552 0.07804970
```

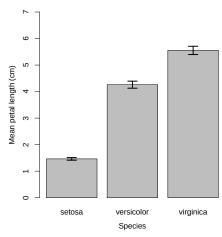
#### In traditional graphics, use barplot() to draw a bar plot:

```
barplot(height = spp_means$mean, names.arg = spp_means$Species, ylim = c(0, 7),
       xlab = "Species", ylab = "Mean petal length (cm)")
```

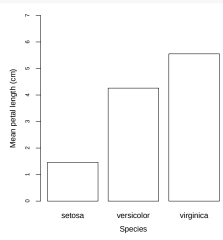


#### Adding error bars can be done with the arrows function:

```
bar_locations <- barplot(height = spp_means mean, names.arg = spp_means Species, plot = FALSE)
barplot(height = spp_means$mean, names.arg = spp_means$Species, ylim = c(0, 7),
        xlab = "Species", ylab = "Mean petal length (cm)")
arrows(x0 = bar_locations[, 1], x1 = bar_locations[, 1],
      y0 = spp_means$mean - 2*spp_means$SE, y1 = spp_means$mean + 2*spp_means$SE,
      angle = 90, lwd = 2, code = 3, length = 0.1)
```



#### As before, there are similar arguments available:



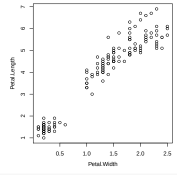
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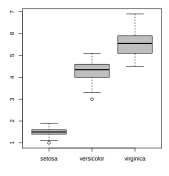
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### Combining plots

You can easily combine multiple traditional graphics using global parameters:

```
oldpar <- par(mfrow = c(1, 2))
plot(Petal.Length ~ Petal.Width, data = iris)
boxplot(Petal.Length ~ Species, data = iris, col = "grey")</pre>
```





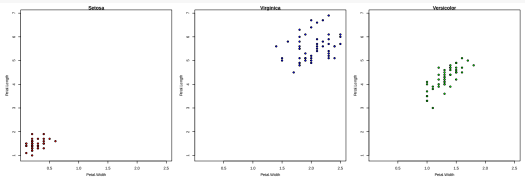
```
par(oldpar) ## or par(mfrow = c(1, 1))
```

Note: if you want to combine graphics of different sizes it is a bit more complex (see ?layout).

### **Facetting**

#### This can be used to created 'facet' plots:

```
oldpar \leftarrow par(mfrow = c(1, 3))
split_data <- split(iris, iris$Species)</pre>
plot(Petal.Length ~ Petal.Width, data = split data$setosa, main = "Setosa", pch = 21,
     bg = "red", col = "black", xlim = range(iris$Petal.Width), ylim = range(iris$Petal.Length))
plot(Petal.Length ~ Petal.Width, data = split_data$virginica, main = "Virginica", pch = 21,
     bg = "blue", col = "black", xlim = range(iris$Petal.Width), ylim = range(iris$Petal.Length))
plot(Petal.Length ~ Petal.Width, data = split_data$versicolor, main = "Versicolor", pch = 21,
     bg = "green", col = "black", xlim = range(iris$Petal, Width), vlim = range(iris$Petal, Length))
```



par(oldpar)

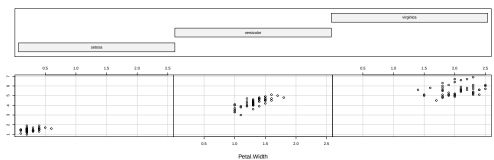
### Facetting

Petal.Length

#### Note: there is also an easier function for that:

coplot(Petal.Length ~ Petal.Width | Species, data = iris, rows = 1)

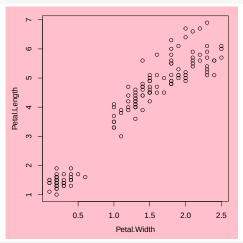
Given : Species



### Background

### Global parameters can also be used to change the background colour:

```
oldpar <- par(bg = "pink")</pre>
plot(Petal.Length ~ Petal.Width, data = iris)
```

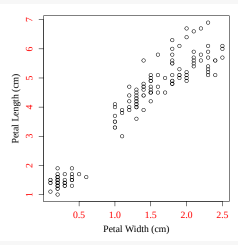


par(oldpar)

Note: you can also plot a background image but you need to use specific packages for that.

# Modifying typefaces

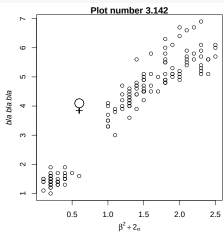
#### Global parameters can also be used to change the typeface:



Aesthetics

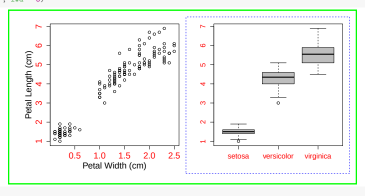
# Special characters

#### You can use weird characters:



## Plot margins

#### Global parameters can also be used to change individual plot margins (blue) and outer margins (green):

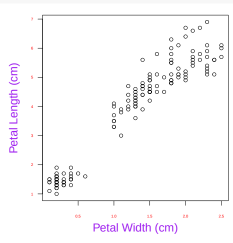


par(oldpar)

### Change axes

You can change the size, colour and orientation of the axis labels and text easily:

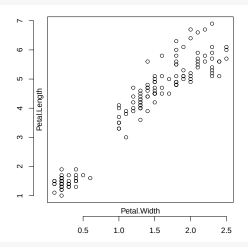
```
plot(Petal.Length ~ Petal.Width, data = iris,
    xlab = "Petal Width (cm)", ylab = "Petal Length (cm)",
    cex.lab = 1.5, cex.axis = 0.5, col.axis = "red", col.lab = "purple", las = 1)
```



## Change axes

#### You can change the position of the axes:

```
oldpar <- par(mgp = c(0.4, 2, 1.5))
plot(Petal.Length ~ Petal.Width, data = iris)
```



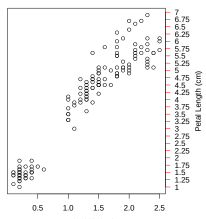
par(oldpar)

Note: check ?par for a huge list of the things you can easily change!

#### Change axes

#### You can change the axes themselves:

```
oldpar <- par(mar = c(4, 4, 1, 4))
plot(Petal.Length ~ Petal.Width, data = iris, axes = FALSE,
     xlab = "Petal Width (cm)", ylab = "")
axis(side = 1)
axis(side = 4, at = seq(1, 7, 0.25), labels = seq(1, 7, 0.25), col.ticks = "red", las = 1)
mtext("Petal Length (cm)", side = 4, line = 3)
box()
```



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## Traditional graphics: Exporting

```
?pdf ?jpeg ?tiff ?bmp ?postscript
```

```
pdf("base_plot.pdf", width = 15, height = 5)
plot(Petal.Length ~ Petal.Width, data = iris)
dev.off()
```

## Plotting in **R**

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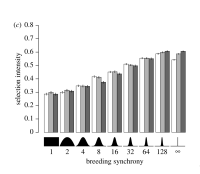
# Challenge

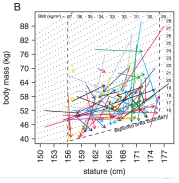
Run the examples of the following traditional plot functions:

- plot.ecdf()
- curve()
- pie()
- cdplot()
- image()
- contour()
- persp()
- o dotchart()
- mosaicplot()
- stars()
- matplot()
- pairs()
- sunflowerplot()

Note: there are a few other ones, but they do not seem very interesting. . .

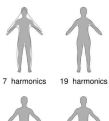
## You can create home-made R graphics too







36 harmonics







316 harmonics

### Plotting in **R**

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### Plotting in **R**

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### ggplot: An introduction

ggplot (defunct) and ggplot2 were both created by Hadley Wickham (now chief Scientist at Rstudio) during his PhD: http://had.co.nz/thesis/practical-tools-hadley-wickham.pdf

The idea was to create a grammar of graphics for  $\mathbf{R}$ .

It is inspired from the seminal Leland Wilkson's book The Grammar of Graphics: "This book [...] presents a unique foundation for producing almost every quantitative graphic found in scientific journals, newspapers, statistical packages, and data visualization systems. This foundation was designed for a distributed computing environment (Internet, Intranet, client-server), with special attention given to conserving computer code and system resources."

In plain english, that means that the graphics are build by considering successive layers.

The originality is that the user directly handles different functions corresponding to each layer.

Other graphics systems do use layers too, but only behind the curtain.

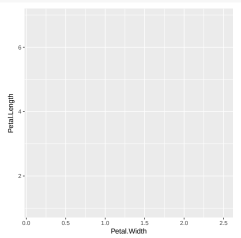
The conceptual unit is thus shifted from the type of plot to the type of layer!

It has pros and cons.

# ggplot: An introduction

Unlike traditional graphics, ggplot works around a single function:

ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length))



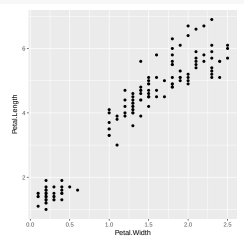
Note: we will then use different functions to add layers on top.

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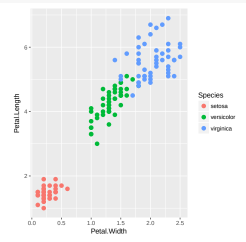
#### We add points on the empty plot:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
 geom_point()
```



You can change point shapes shape (= pch in traditional graphics) and colour (= col in traditional graphics):

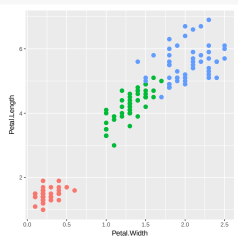
```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
 geom_point(aes(colour = Species), shape = 16, size = 3)
```



Note: everytime we want to create a connection between the data and some elements of the plot, we use the same function: aes()!

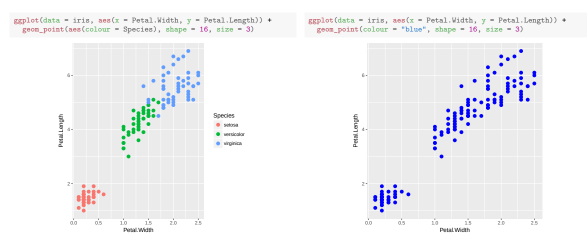
### In ggplot the legend is included by default but you can remove it:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
  geom_point(aes(colour = Species), shape = 16, size = 3) +
 theme(legend.position = "none")
```



The aesthetic function in ggplot is a powerful tool for changing plot aesthetics.

If you specify an aesthetic argument inside aes(), it will give each point a different aesthetic based on its value. If you specify the same aesthetic argument outside aes(), it will give all data points the same aesthetic:



Note: we have used aes() to apply colours but it can be used to make other changes (see later).

Again, for many elements you can set both an outline colour (= col for traditional graphics) and background fill colour (= bg for traditional graphics):

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
                                                                                      ggplot(data = iris, aes(x = Petal.Width, v = Petal.Length)) +
  geom point(aes(colour = Species), fill = "black", shape = 21)
                                                                                         geom point(aes(fill = Species), colour = "black", shape = 21)
                                                           Species
                                                                                                                                                 Species
                                                                                                   Petal.Length
            Petal.Length

    versicolor

    versicolo

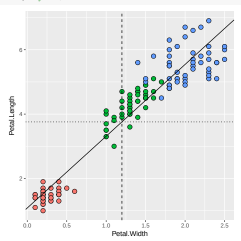
    virginica

    virginica

                                                                                                                      Petal Width
```

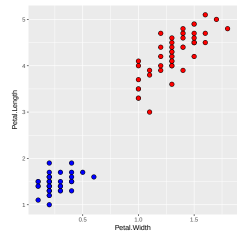
#### You can also add lines to the plot:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
  geom_point(aes(fill = Species), shape = 21, size = 3) +
  theme(legend.position = "none") +
  geom_hline(yintercept = mean(iris$Petal.Length), lty = 3) +
  geom_vline(xintercept = mean(iris$Petal.Width), lty = 2) +
  geom abline(intercept = 1.084, slope = 2.23, lty = 1)
```



#### You can add extra points to the plot by using two geom\_point layers:

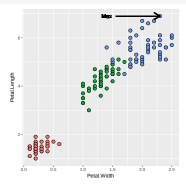
```
versicolor <- subset(iris, Species == "versicolor")</pre>
           <- subset(iris, Species == "setosa")
ggplot() +
  geom_point(data = versicolor, aes(x = Petal.Width, y = Petal.Length), fill = "red", shape = 21, size = 3) +
 geom_point(data = setosa, aes(x = Petal.Width, y = Petal.Length), fill = "blue", shape = 21, size = 3)
```



# Traditional graphics: Scatter plots

#### You can add text and arrows:

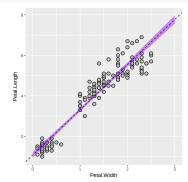
```
max.value <- iris[iris$Petal.Length == max(iris$Petal.Length), ]</pre>
ggplot(data = iris, aes(x = Petal.Width, v = Petal.Length)) +
  geom point(aes(fill = Species), shape = 21, size = 3) +
  theme(legend.position = "none") +
  geom segment(aes(x = max.value$Petal.Width - 0.75,
                   xend = max.value$Petal.Width, y = max.value$Petal.Length,
                   yend = max.value$Petal.Length), size = 1,
               arrow = arrow(length = unit(0.5, "cm"))) +
  geom text(aes(x = max.value$Petal.Width - 0.9, v = max.value$Petal.Length, label = "Max"))
```



# ggplot: Scatter plots

In ggplot, confidence intervals can be added with specialised function geom\_ribbon():

```
test_mod <- lm(Petal.Length ~ Petal.Width, data = iris)
        <- data.frame(Petal.Width = seq(0, 3, length.out = 100))
newdat
        <- predict(test_mod, newdata = newdat, interval = "confidence")</pre>
pred
ggplot() +
                           aes(x = Petal.Width, y = Petal.Length), fill = "grey", shape = 21, size = 3) +
 geom point(data = iris,
 geom_ribbon(data = newdat, aes(x = Petal.Width, ymin = pred[, 2], ymax = pred[, 3]), fill = "purple", alpha = 0.5) +
 geom abline(intercept = coef(test mod)[1], slope = coef(test mod)[2], ltv = 2)
```



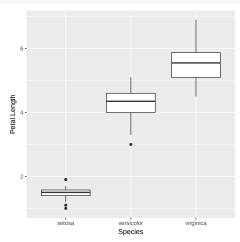
Note: because we are building the plot with two different datasets (i.e. iris & newdat), we specify the data separately in each line of the ggplot code.

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# ggplot: Box plots

## In ggplot, use geom\_boxplot() to draw a box plot:

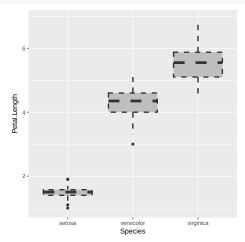
```
ggplot(data = iris, aes(x = Species, y = Petal.Length)) +
 geom_boxplot()
```



# ggplot: Box plots

## Many of the same changes made to scatter plots can be made here:

```
ggplot(data = iris, aes(x = Species, y = Petal.Length)) +
 geom_boxplot(fill = "grey", size = 1, lty = 2)
```

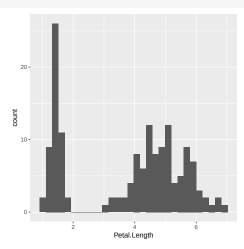


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# ggplot: Histograms

## In ggplot, use geom\_histogram() to draw an histogram:

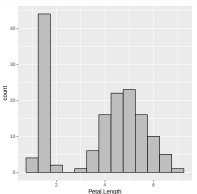
```
ggplot(data = iris, aes(x = Petal.Length)) +
  geom_histogram()
## 'stat_bin()' using 'bins = 30'. Pick better value with
## 'binwidth'.
```



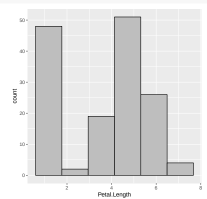
# ggplot: Histograms

You can change the width of bins with binwidth or their number with bins:

```
ggplot(data = iris, aes(x = Petal.Length)) +
  geom_histogram(binwidth = 0.5, colour = "black", fill = "grey")
```



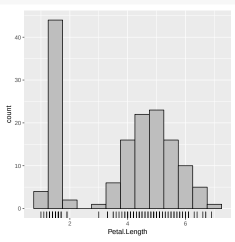




# ggplot: Histograms

## Add a rug below the histogram with geom\_rug:

```
ggplot(data = iris, aes(x = Petal.Length)) +
  geom_histogram(binwidth = 0.5, colour = "black", fill = "grey")+
 geom_rug()
```

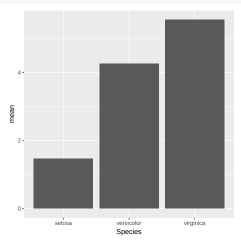


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# ggplot: Bar plots

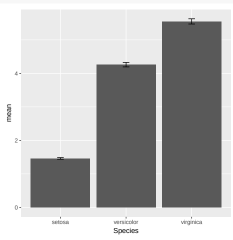
### In ggplot, use geom\_col() to draw a bar plot:

```
ggplot(data = spp_means, aes(x = Species, y = mean)) +
 geom_col()
```



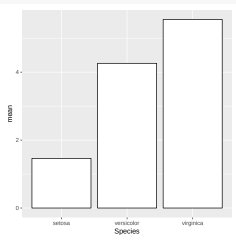
# Adding error bars in ggplot is much easier than in traditional graphics:

```
ggplot(data = spp_means, aes(x = Species, y = mean)) +
  geom_col() +
  geom_errorbar(aes(ymin = mean - SE, ymax = mean + SE), width = 0.1)
```



### As before, modifications are straigthforward:

```
ggplot(data = spp_means, aes(x = Species, y = mean)) +
  geom_col(fill = "white", colour = "black")
```



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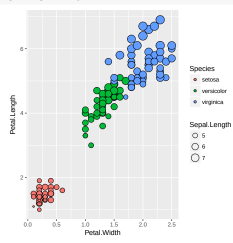
### Introduction to aesthetics

In ggplot, you can change aesthetics in individual segments of the code (aes()) or you can change information for the whole plot using themes (theme()).

# Using the aesthetic function more

We'll start by looking back at aes(). We can use it to change multiple aesthetics of a plot:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
   geom_point(aes(fill = Species, size = Sepal.Length), shape = 21)
```

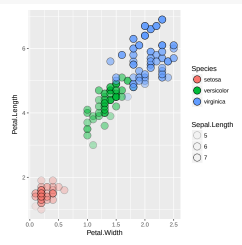


Aesthetics

# Using the aesthetic function more

## Here we change the transparency:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
 geom_point(aes(fill = Species, alpha = Sepal.Length), shape = 21, size = 5)
```

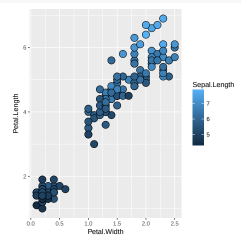


Aesthetics

# Using the aesthetic argument more

## Applying aesthetics to continuous variables will be different to categorical variables:

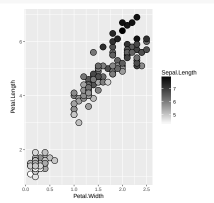
```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
 geom_point(aes(fill = Sepal.Length), shape = 21, size = 5)
```



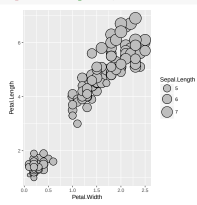
# Using the aesthetic argument more

#### We can adjust the way the aesthetics are applied:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
geom_point(aes(fill = Sepal.Length), shape = 21, size = 5) +
scale_fill_continuous(low = "white", high = "black")
```



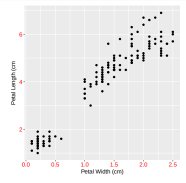
```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
  geom_point(aes(size = Sepal.Length), shape = 21, fill = "grey") +
  scale size continuous(range = c(3, 10))
```



Aesthetics

## You can change size and colour of axis text easily:

```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
 geom_point() +
 xlab("Petal Width (cm)") +
 ylab("Petal Length (cm") +
 theme(axis.text = element_text(size = 12, colour = "red"),
       axis.title = element_text(size = 12))
```

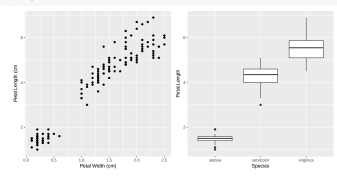


Note: the text size uses different measurement units than traditional graphics.

Aesthetics

## Combining plots

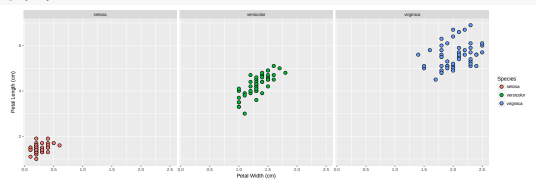
Combining plots is less straightforward in ggplot... You need to use an additional package:



## Faceting

Although combining multiple plots is cumbersome, there is an inbuilt function to create facets:

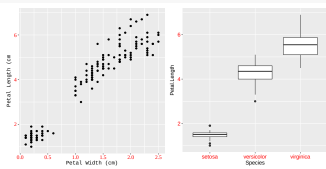
```
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
  geom_point(aes(fill = Species), shape = 21, colour = "black", size = 3) +
 xlab("Petal Width (cm)") +
 vlab("Petal Length (cm)") +
 facet_wrap(~ Species)
```



Aesthetics

# Modifying typefaces

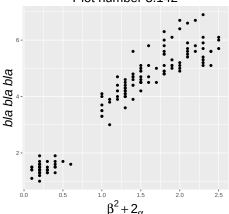
Unlike traditional graphics, in ggplot you can easily change the typeface of individual elements in theme:



## Special characters

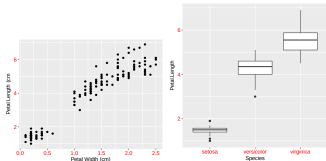
#### You can use weird characters:





## Plot margins

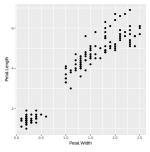
## Plot margins are also controlled in theme of each plot individually:

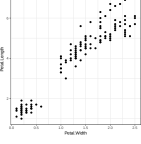


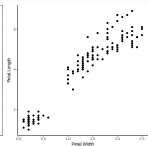
### Preset themes

## ggplot also has a number of preset themes that you can use:

```
grey <- ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +</pre>
  geom_point() +
  theme grey()
bw <- ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +</pre>
  geom point() +
  theme bw()
classic <- ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +</pre>
  geom point() +
  theme_classic()
grid.arrange(grey, bw, classic, nrow = 1)
```





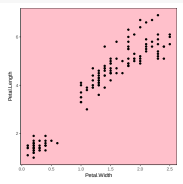


Aesthetics

### Preset themes

### You can create your own theme once for all!

```
theme_pink <- theme_classic() %+replace% theme(panel.background = element_rect(fill = "pink"))</pre>
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
  geom_point()+
 theme_pink
```

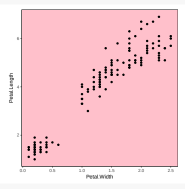


Note: check ?theme for the list of options that can be changed.

### Preset themes

### You can set a theme for all plots with theme\_set

```
theme_old <- theme_set(theme_pink)
ggplot(data = iris, aes(x = Petal.Width, y = Petal.Length)) +
    geom_point()</pre>
```



theme\_set(theme\_old) ## restore original theme

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  - Bar plots
  - Aesthetics
  - Saving your plot
  - Other plots
  - Conclusion: traditional graphics vs. ggplot

# ggplot: Exporting

```
?ggsave
```

```
classic <- ggplot(iris, aes(x = Petal.Width, y = Petal.Length))+
   geom_point()+
   them_classic()

ggsave("ggplot.pdf", plot = classic, width = 15, height = 5)</pre>
```

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# There are other ggplots out there!

There are a few other plot types in ggplot that are worth looking at:

- geom\_violin()
- geom\_area()
- geom\_smooth()
- geom\_density()

But the true force of having developed something as modular as ggplot is that there are many more packages each week providing ggplot extensions!

A few random examples of ggplot extensions:

- ggExtra (for marginal distribution)
- ggthemes (for extra themes, including theme\_excel()!! :-/)
- ggmap (for maps)
- ggrepel (for adding labels to plots)
- ggalt (for creating cluster plots)
- cowplot (for creating nested figures)

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# Which plotting tool should you use?

### Traditional graphics:

- no new packages required
- arguments often differ between functions
- help files useful
- easier for doing something simple
- more difficult for doing something complex

### ggplot:

- requires multiple packages for best results
- uniform grammar
- help files often useless (due to modularity)
- more difficult for doing something simple
- easier for doing something complex
- faster for large datasets
- more accessible than lattice (or grid itself)

ightarrow It may be useful to know both!!