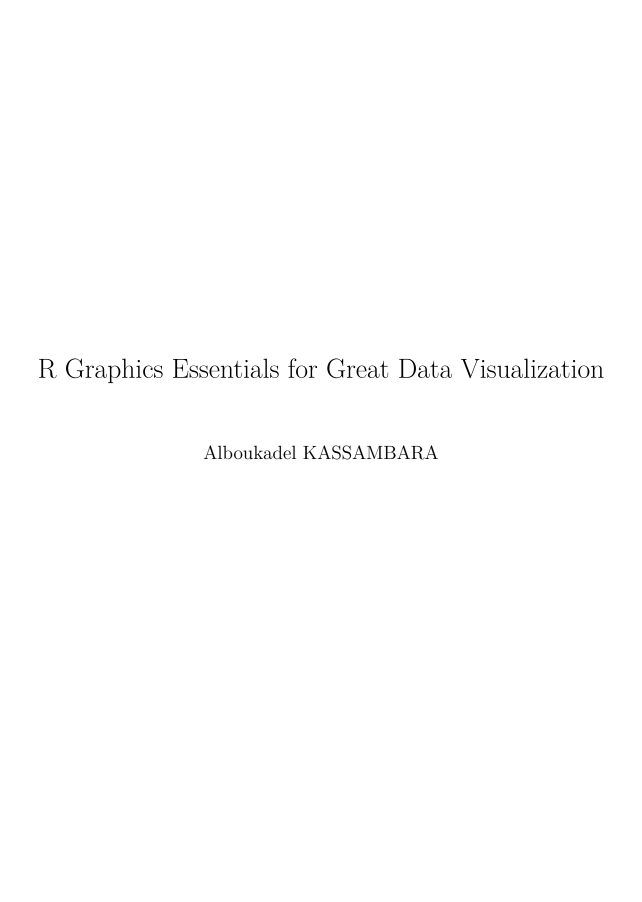
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R Graphics Essentials for Great Data Visualization

+200 Practical Examples You Want to Known

Alboukadel Kassambara



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Preface

0.1 What you will learn

Data visualization is one of the most important part of data science. Many books and courses present a catalogue of graphics but they don't teach you which charts to use according to the type of the data.

In this book, we start by presenting the key graphic systems and packages available in R, including R base graphs, lattice and ggplot2 plotting systems.

Next, we provide practical examples to create great graphics for the right data using either the ggplot2 package and extensions or the traditional R graphics.

With this book, you 'll learn:

- How to quickly create beautiful graphics using ggplot2 packages
- How to properly customize and annotate the plots
- Type of graphics for visualizing categorical and continuous variables
- How to add automatically p-values to box plots, bar plots and alternatives
- How to add marginal density plots and correlation coefficients to scatter plots
- Key methods for analyzing and visualizing multivariate data
- R functions and packages for plotting time series data
- How to combine multiple plots on one page to create production-quality figures.

0.2 Book website

The website for this book is located at: http://www.sthda.com/english/. It contains number of resources.

0.3 Executing the R codes from the PDF

For a single line R code, you can just copy the code from the PDF to the R console.

0.4. COLOPHON vii

For a multiple-line R codes, an error is generated, sometimes, when you copy and paste directly the R code from the PDF to the R console. If this happens, a solution is to:

- Paste firstly the code in your R code editor or in your text editor
- Copy the code from your text/code editor to the R console

0.4 Colophon

This book was built with R 3.3.2 and the following packages :

##		name	version	source
##	1	bookdown	0.5	Github:rstudio/bookdown
##	2	changepoint	2.2.2	CRAN
##	3	cowplot	0.8.0.9000	<pre>Github:wilkelab/cowplot</pre>
##	4	dplyr	0.7.4	cran
##	5	factoextra	1.0.5.999	local:kassambara/factoextra
##	6	${\tt FactoMineR}$	1.38	CRAN
##	7	GGally	1.3.0	CRAN
##	8	ggcorrplot	0.1.1.9000	<pre>Github:kassambara/ggcorrplot</pre>
##	9	ggforce	0.1.1	Github:thomasp85/ggforce
##	10	ggformula	0.6	CRAN
##	11	ggfortify	0.4.1	CRAN
##	12	ggpmisc	0.2.15	CRAN
##	13	ggpubr	0.1.5.999	Github:kassambara/ggpubr
##	14	lattice	0.20-34	CRAN
##	15	readr	1.1.1	cran
##	16	${\tt scatterplot3d}$	0.3-40	cran
##	17	strucchange	1.5-1	CRAN
##	18	tidyr	0.7.2	cran

About the author

Alboukadel Kassambara is a PhD in Bioinformatics and Cancer Biology. He works since many years on genomic data analysis and visualization (read more: http://www.alboukadel.com/).

He has work experiences in statistical and computational methods to identify prognostic and predictive biomarker signatures through integrative analysis of large-scale genomic and clinical data sets.

He created a bioinformatics web-tool named GenomicScape (www.genomicscape.com) which is an easy-to-use web tool for gene expression data analysis and visualization.

He developed also a training website on data science, named STHDA (Statistical Tools for High-throughput Data Analysis, www.sthda.com/english), which contains many tutorials on data analysis and visualization using R software and packages.

He is the author of many popular R packages for:

- multivariate data analysis (factoextra, http://www.sthda.com/english/rpkgs/factoextra),
- survival analysis (survminer, http://www.sthda.com/english/rpkgs/survminer/),
- correlation analysis (ggcorrplot, http://www.sthda.com/english/wiki/ggcorrplot-visualization-of-a-correlation-matrix-using-ggplot2),
- creating publication ready plots in R (ggpubr, http://www.sthda.com/english/rpkgs/ggpubr).

Recently, he published three books on data analysis and visualization:

- 1. Practical Guide to Cluster Analysis in R (https://goo.gl/yhhpXh)
- 2. Practical Guide To Principal Component Methods in R (https://goo.gl/d4Doz9)

Chapter 1

R Basics for Data Visualization

1.1 Introduction

R is a free and powerful statistical software for analyzing and visualizing data.

In this chapter, you'll learn:

- the basics of R programming for importing and manipulating your data:
 - filtering and ordering rows,
 - renaming and adding columns,
 - computing summary statistics
- R graphics systems and packages for data visualization:
 - R traditional base plots
 - Lattice plotting system that aims to improve on R base graphics
 - ggplot2 package, a powerful and a flexible R package, for producing elegant graphics piece by piece.
 - ggpubr package, which facilitates the creation of beautiful ggplot2-based graphs for researcher with non-advanced programming backgrounds.
 - ggformula package, an extension of ggplot2, based on formula interfaces (much like the lattice interface)

1.2 Install R and RStudio

RStudio is an integrated development environment for R that makes using R easier. R and RStudio can be installed on Windows, MAC OSX and Linux platforms.

- 1. R can be downloaded and installed from the Comprehensive R Archive Network (CRAN) webpage (http://cran.r-project.org/)
- 2. After installing R software, install also the RStudio software available at: http://www.rstudio.com/products/RStudio/.
- 3. Launch RStudio and start use R inside R studio.

1.3 Install and load required R packages

An R package is a collection of functionalities that extends the capabilities of base R. To use the R code provide in this book, you should install the following R packages:

- tidyverse packages, which are a collection of R packages that share the same programming philosophy. These packages include:
 - readr: for importing data into R
 - dplyr: for data manipulation
 - ggplot2 and ggpubr for data visualization.
- ggpubr package, which makes it easy, for beginner, to create publication ready plots.
- 1. **Install the tidyverse package**. Installing tidyverse will install automatically readr, dplyr, ggplot2 and more. Type the following code in the R console:

```
install.packages("tidyverse")
```

- 2. Install the ggpubr package.
- We recommend to install the latest developmental version of ggpubr as follow:

```
if(!require(devtools)) install.packages("devtools")
devtools::install_github("kassambara/ggpubr")
```

• If the above R code fails, you can install the latest stable version on CRAN:

```
install.packages("ggpubr")
```

3. Load required packages. After installation, you must first load the package for using the functions in the package. The function library() is used for this task. An alternative function is require(). For example, to load ggplot2 and ggpubr packages, type this:

```
library("ggplot2")
library("ggpubr")
```

Now, we can use R functions, such as ggscatter() [in the ggpubr package] for creating a scatter plot.

If you want to learn more about a given function, say ggscatter(), type this in R console: ?ggscatter.

1.4 Data format

Your data should be in rectangular format, where columns are variables and rows are observations (individuals or samples).

• Column names should be compatible with R naming conventions. Avoid column with blank space and special characters. Good column names: long_jump or long.jump. Bad column name: long jump.

- Avoid beginning column names with a number. Use letter instead. Good column names: sport 100m or x100m. Bad column name: 100m.
- Replace missing values by NA (for not available)

For example, your data should look like this:

```
manufacturer model displ year cyl
                                             trans drv
1
          audi
                         1.8 1999
                                         auto(15)
                   a4
2
          audi
                   a4
                         1.8 1999
                                     4 manual(m5)
                                                     f
3
                   a4
                         2.0 2008
                                     4 manual(m6)
                                                     f
          audi
          audi
                   a4
                         2.0 2008
                                         auto(av)
                                                     f
```

Read more at: Best Practices in Preparing Data Files for Importing into R¹

1.5 Import your data in R

First, save your data into txt or csv file formats and import it as follow (you will be asked to choose the file):

```
library("readr")

# Reads tab delimited files (.txt tab)
my_data <- read_tsv(file.choose())

# Reads comma (,) delimited files (.csv)
my_data <- read_csv(file.choose())

# Reads semicolon(;) separated files(.csv)
my_data <- read_csv2(file.choose())</pre>
```

Read more about how to import data into R at this link: http://www.sthda.com/english/wiki/importing-data-into-r

1.6 Demo data sets

R comes with several demo data sets for playing with R functions. The most used R demo data sets include: **USArrests**, **iris** and **mtcars**. To load a demo data set, use the function **data**() as follow. The function **head**() is used to inspect the data.

```
data("iris")
               # Loading
head(iris, n = 3) # Print the first n = 3 rows
##
     Sepal.Length Sepal.Width Petal.Length Petal.Width Species
## 1
              5.1
                           3.5
                                        1.4
                                                     0.2
                                                          setosa
## 2
              4.9
                           3.0
                                        1.4
                                                     0.2
                                                          setosa
```

 $^{^{1}} http://www.sthda.com/english/wiki/best-practices-in-preparing-data-files-for-importing-into-r\\$

3 4.7 3.2 1.3 0.2 setosa

To learn more about iris data sets, type this:

?iris

After typing the above R code, you will see the description of iris data set: this iris data set gives the measurements in centimeters of the variables sepal length and width and petal length and width, respectively, for 50 flowers from each of 3 species of iris. The species are Iris setosa, versicolor, and virginica.

1.7 Data manipulation

After importing your data in R, you can easily manipulate it using the dplyr package (Wickham et al., 2017), which can be installed using the R code: install.packages("dplyr").

After loading dplyr, you can use the following R functions:

- filter(): Pick rows (observations/samples) based on their values.
- distinct(): Remove duplicate rows.
- arrange(): Reorder the rows.
- select(): Select columns (variables) by their names.
- rename(): Rename columns.
- mutate(): Add/create new variables.
- summarise(): Compute statistical summaries (e.g., computing the mean or the sum)
- group_by(): Operate on subsets of the data set.

Note that, dplyr package allows to use the forward-pipe chaining operator (%>%) for combining multiple operations. For example, x %>% f is equivalent to f(x). Using the pipe (%>%), the output of each operation is passed to the next operation. This makes R programming easy.

We'll show you how these functions work in the different chapters of this book.

1.8 R graphics systems

There are different graphic packages available in R² for visualizing your data: 1) R base graphs, 2) Lattice Graphs (Sarkar, 2016) and 3) ggplot2 (Wickham and Chang, 2017).

In this section, we start by providing a quick overview of R base and lattice plots, and then we move to ggplot2 graphic system. The vast majority of plots generated in this book is based on the modern and flexible **ggplot2** R package.

²http://www.sthda.com/english/wiki/data-visualization

1.8.1 R base graphs

R comes with simple functions to create many types of graphs. For example:

Plot Types	R base function
Scatter plot	plot()
Scatter plot matrix	pairs()
Box plot	boxplot()
Strip chart	stripchart()
Histogram plot	hist()
density plot	density()
Bar plot	barplot()
Line plot	plot() and line()
Pie charts	pie()
Dot charts	dotchart()
Add text to a plot	text()

In the most cases, you can use the following arguments to customize the plot:

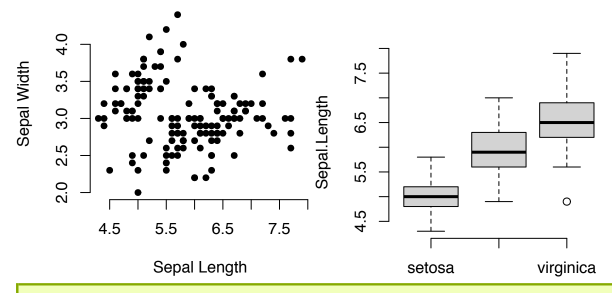
- pch: change point shapes. Allowed values comprise number from 1 to 25.
- cex: change point size. Example: cex = 0.8.
- col: change point color. Example: col = "blue".
- frame: logical value. frame = FALSE removes the plot panel border frame.
- main, xlab, ylab. Specify the main title and the x/y axis labels -, respectively
- las: For a vertical x axis text, use las = 2.

In the following R code, we'll use the iris data set to create a:

- (1) Scatter plot of Sepal.Length (on x-axis) and Sepal.Width (on y-axis).
- (2) Box plot of Sepal.length (y-axis) by Species (x-axis)

```
# (1) Create a scatter lot
plot(
    x = iris$Sepal.Length, y = iris$Sepal.Width,
    pch = 19, cex = 0.8, frame = FALSE,
    xlab = "Sepal Length",ylab = "Sepal Width"
    )

# (2) Create a box plot
boxplot(Sepal.Length ~ Species, data = iris,
         ylab = "Sepal.Length",
         frame = FALSE, col = "lightgray")
```



Read more examples at: R base Graphics on STHDA, http://www.sthda.com/english/wiki/r-base-graphs

1.8.2 Lattice graphics

The lattice R package provides a plotting system that aims to improve on R base graphs. After installing the package, whith the R command install.packages("lattice"), you can test the following functions.

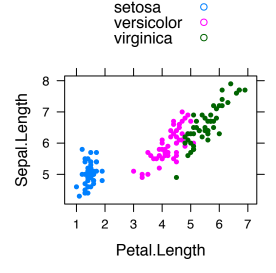
• Main functions in the lattice package:

Plot types	Lattice functions
Scatter plot	xyplot()
Scatter plot matrix	splom()
3D scatter plot	$\operatorname{cloud}()$
Box plot	bwplot()
strip plots (1-D scatter plots)	stripplot()
Dot plot	dotplot()
Bar chart	barchart()
Histogram	histogram()
Density plot	densityplot()
Theoretical quantile plot	qqmath()
Two-sample quantile plot	qq()
3D contour plot of surfaces	contourplot()
False color level plot of surfaces	levelplot()
Parallel coordinates plot	parallel()
3D wireframe graph	wireframe()

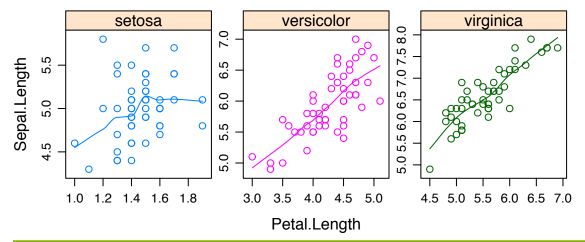
The lattice package uses formula interface. For example, in lattice terminology, the formula $y \sim x \mid$ group, means that we want to plot the y variable according to the x variable, splitting the plot into multiple panels by the variable group.

• Create a basic scatter plot of y by x. Syntax: y ~ x. Change the color by groups and use auto.key = TRUE to show legends:

```
library("lattice")
xyplot(
   Sepal.Length ~ Petal.Length, group = Species,
   data = iris, auto.key = TRUE, pch = 19, cex = 0.5
)
```



• Multiple panel plots by groups. Syntax: y ~ x | group.



Read more examples at: Lattice Graphics on $STHDA^a$

ahttp://www.sthda.com/english/wiki/lattice-graphs

1.8.3 ggplot2 graphics

GGPlot2 is a powerful and a flexible R package, implemented by Hadley Wickham, for producing elegant graphics piece by piece. The $\mathbf{g}\mathbf{g}$ in ggplot2 means *Grammar of Graphics*, a graphic concept which describes plots by using a "grammar". According to the ggplot2 concept, a plot can be divided into different fundamental parts: $\mathbf{Plot} = \mathbf{data} + \mathbf{Aesthetics} + \mathbf{Geometry}$

- data: a data frame
- **aesthetics**: used to indicate the **x** and **y** variables. It can be also used to control the **color**, the **size** and the **shape** of points, etc.....
- **geometry**: corresponds to the type of graphics (histogram, box plot, line plot,)

The ggplot2 syntax might seem opaque for beginners, but once you understand the basics, you can create and customize any kind of plots you want.

Note that, to reduce this opacity, we recently created an R package, named **ggpubr** (ggplot2 Based Publication Ready Plots), for making ggplot simpler for students and researchers with non-advanced programming backgrounds. We'll present ggpubr in the next section.

After installing and loading the ggplot2 package, you can use the following key functions:

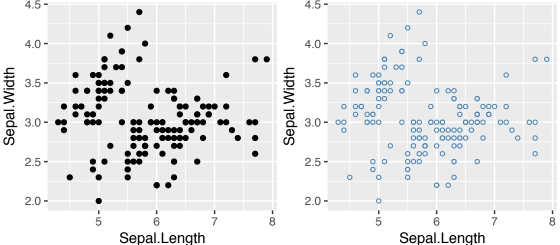
Plot types	GGPlot2 functions							
Initialize a ggplot	ggplot()							
Scatter plot	$geom_point()$							
Box plot	geom_boxplot()							
Violin plot	geom_violin()							
strip chart	geom_jitter()							
Dot plot	geom_dotplot()							
Bar chart	geom_bar()							
Line plot	geom_line()							
Histogram	geom_histogram()							
Density plot	geom_density()							
Error bars	geom_errorbar()							
QQ plot	$\operatorname{stat}_{qq}()$							
ECDF plot	$\operatorname{stat}_{\operatorname{ecdf}}()$							
Title and axis labels	labs()							

The main function in the ggplot2 package is ggplot(), which can be used to initialize the plotting system with data and x/y variables.

For example, the following R code takes the iris data set to initialize the ggplot and then a layer ($geom_point()$) is added onto the ggplot to create a scatter plot of x = Sepal.Length by y = Sepal.Width:

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

```
# Change point size, color and shape
ggplot(iris, aes(x = Sepal.Length, y = Sepal.Width))+
geom_point(size = 1.2, color = "steelblue", shape = 21)
4.5-
```



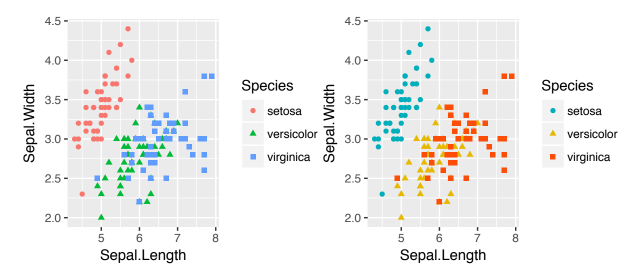
Note that, in the code above, the shape of points is specified as number. To display the different point shape available in R, type this:

```
ggpubr::show_point_shapes()
```

It's also possible to control points shape and color by a grouping variable (here, Species). For example, in the code below, we map points color and shape to the Species grouping variable.

```
# Control points color by groups
ggplot(iris, aes(x = Sepal.Length, y = Sepal.Width))+
  geom_point(aes(color = Species, shape = Species))

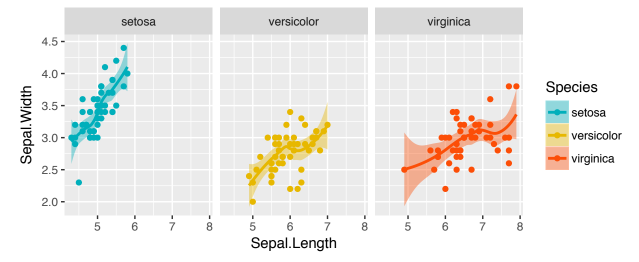
# Change the default color manually.
# Use the scale_color_manual() function
ggplot(iris, aes(x = Sepal.Length, y = Sepal.Width))+
  geom_point(aes(color = Species, shape = Species))+
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
```



You can also split the plot into multiple panels according to a grouping variable. R function: facet_wrap(). Another interesting feature of ggplot2, is the possibility to combine multiple layers on the same plot. For example, with the following R code, we'll:

- Add points with geom point(), colored by groups.
- Add the fitted smoothed regression line using geom_smooth(). By default the function geom_smooth() add the regression line and the confidence area. You can control the line color and confidence area fill color by groups.
- Facet the plot into multiple panels by groups
- Change color and fill manually using the function scale_color_manual() and scale fill manual()

```
ggplot(iris, aes(x = Sepal.Length, y = Sepal.Width))+
  geom_point(aes(color = Species))+
  geom_smooth(aes(color = Species, fill = Species))+
  facet_wrap(~Species, ncol = 3, nrow = 1)+
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))+
  scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
```



Note that, the default theme of ggplots is theme_gray() (or theme_grey()), which is theme with grey background and white grid lines. More themes are available for professional presentations or publications. These include: theme bw(), theme classic() and

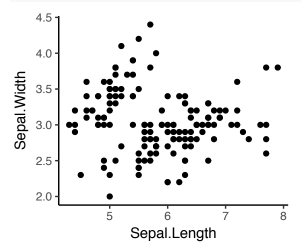
```
theme minimal().
```

To change the theme of a given ggplot (p), use this: p + theme_classic(). To change the default theme to theme_classic() for all the future ggplots during your entire R session, type the following R code:

```
theme_set(
  theme_classic()
)
```

Now you can create ggplots with theme_classic() as default theme:

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

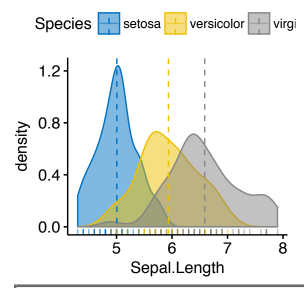


1.8.4 ggpubr for publication ready plots

The **ggpubr** R package facilitates the creation of beautiful ggplot2-based graphs for researcher with non-advanced programming backgrounds (Kassambara, 2017).

For example, to create the density distribution of "Sepal.Length", colored by groups ("Species"), type this:

```
library(ggpubr)
# Density plot with mean lines and marginal rug
ggdensity(iris, x = "Sepal.Length",
   add = "mean", rug = TRUE,  # Add mean line and marginal rugs
   color = "Species", fill = "Species", # Color by groups
   palette = "jco")  # use jco journal color palette
```

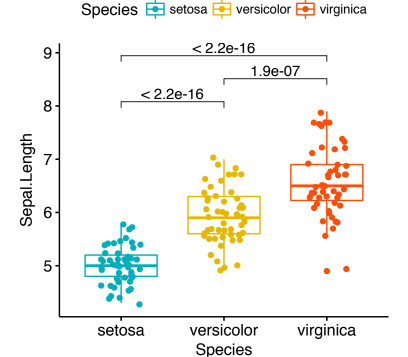


Note that the argument palette can take also a custom color palette. For example palette= c("#00AFBB", "#E7B800", "#FC4E07").

• Create a box plot with p-values comparing groups:

```
# Groups that we want to compare
my_comparisons <- list(
   c("setosa", "versicolor"), c("versicolor", "virginica"),
   c("setosa", "virginica")
)

# Create the box plot. Change colors by groups: Species
# Add jitter points and change the shape by groups
ggboxplot(
   iris, x = "Species", y = "Sepal.Length",
   color = "Species", palette = c("#00AFBB", "#E7B800", "#FC4E07"),
   add = "jitter"
   )+
   stat_compare_means(comparisons = my_comparisons, method = "t.test")</pre>
```



Learn more on STHDA at: ggpubr: Publication Ready Plots^a

ahttp://www.sthda.com/english/articles/24-ggpubr-publication-ready-plots/

1.9 Export R graphics

You can export R graphics to many file formats, including: PDF, PostScript, SVG vector files, Windows MetaFile (WMF), PNG, TIFF, JPEG, etc.

The standard procedure to save any graphics from R is as follow:

- 1. Open a graphic device using one of the following functions:
- pdf("r-graphics.pdf"),
- postscript("r-graphics.ps"),
- svg("r-graphics.svg"),
- png("r-graphics.png"),
- tiff("r-graphics.tiff"),
- jpeg("r-graphics.jpg"),
- win.metafile("r-graphics.wmf"),
- and so on.

Additional arguments indicating the width and the height (in inches) of the graphics region can be also specified in the mentioned function.

- 2. Create a plot
- 3. Close the graphic device using the function dev.off()

For example, you can export R base plots to a pdf file as follow:

```
pdf("r-base-plot.pdf")
# Plot 1 --> in the first page of PDF
plot(x = iris$Sepal.Length, y = iris$Sepal.Width)
# Plot 2 ---> in the second page of the PDF
hist(iris$Sepal.Length)
dev.off()
```

To export ggplot2 graphs, the R code looks like this:

```
# Create some plots
library(ggplot2)
myplot1 <- ggplot(iris, aes(Sepal.Length, Sepal.Width)) +
    geom_point()
myplot2 <- ggplot(iris, aes(Species, Sepal.Length)) +
    geom_boxplot()

# Print plots to a pdf file
pdf("ggplot.pdf")
print(myplot1)  # Plot 1 --> in the first page of PDF
print(myplot2)  # Plot 2 ---> in the second page of the PDF
dev.off()
```

Note that for a ggplot, you can also use the following functions to export the graphic:

- ggsave() [in ggplot2]. Makes it easy to save a ggplot. It guesses the type of graphics device from the file extension.
- ggexport() [in ggpubr]. Makes it easy to arrange and export multiple ggplots at once.

See also the following blog post to save high-resolution ggplots^a

 $^a \verb|http://www.sthda.com/english/wiki/saving-high-resolution-ggplots-how-to-preserve-semi-transparency$

Chapter 2

Plot One Variable

2.1 Introduction

To visualize one variable, the type of graphs to be used depends on the type of the variable:

- For **categorical variable** or grouping variables. You can visualize the count of categories using a bar plot or using a pie chart to show the proportion of each category.
- For **continuous variable**, you can visualize the distribution of the variable using density plots, histograms and alternatives.

In this R graphics tutorial, you'll learn how to:

- Visualize a categorical variable using bar plots, dot charts and pie charts
- Visualize the distribution of a continuous variable using:
 - density and histogram plots,
 - other alternatives, such as frequency polygon, area plots, dot plots, box plots,
 Empirical cumulative distribution function (ECDF) and Quantile-quantile plot (QQ plots).
 - Density ridgeline plots, which are useful for visualizing changes in distributions, of a continuous variable, over time or space.
 - Bar plot and modern alternatives, including lollipop charts and cleveland's dot plots.

2.2 Prerequisites

Load required packages and set the theme function theme_pubr() [in ggpubr] as the default theme:

```
library(ggplot2)
library(ggpubr)
theme_set(theme_pubr())
```

2.3 One categorical variable

2.3.1 Bar plot of counts

- Plot types: Bar plot of the count of group levels
- Key function: geom bar()
- Key arguments: alpha, color, fill, linetype and size

Demo data set: diamonds [in ggplot2]. Contains the prices and other attributes of almost 54000 diamonds. The column cut contains the quality of the diamonds cut (Fair, Good, Very Good, Premium, Ideal).

The R code below creates a bar plot visualizing the number of elements in each category of diamonds cut.

```
ggplot(diamonds, aes(cut)) +
geom_bar(fill = "#0073C2FF") +
theme_pubclean()

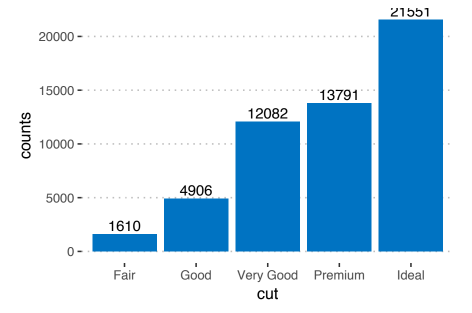
20000 -
15000 -
5000 -
Fair Good Very Good Premium Ideal
cut
```

Compute the frequency of each category and add the labels on the bar plot:

- dplyr package used to summarise the data
- geom_bar() with option stat = "identity" is used to create the bar plot of the summary output as it is.
- geom_text() used to add text labels. Adjust the position of the labels by using hjust (horizontal justification) and vjust (vertical justification). Values should be in [0, 1].

```
## 1 Fair 1610
## 2 Good 4906
## 3 Very Good 12082
## 4 Premium 13791
## 5 Ideal 21551
```

```
# Create the bar plot. Use theme_pubclean() [in ggpubr]
ggplot(df, aes(x = cut, y = counts)) +
  geom_bar(fill = "#0073C2FF", stat = "identity") +
  geom_text(aes(label = counts), vjust = -0.3) +
  theme_pubclean()
```



2.3.2 Pie charts

Pie chart is just a stacked bar chart in polar coordinates.

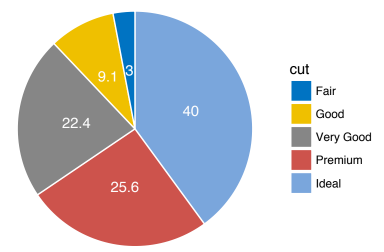
First,

- Arrange the grouping variable (cut) in descending order. This important to compute the y coordinates of labels.
- compute the proportion (counts/total) of each category
- compute the position of the text labels as the cumulative sum of the proportion. To
 put the labels in the center of pies, we'll use cumsum(prop) 0.5*prop as label
 position.

```
##
        <ord> <int> <dbl>
                              <dbl>
## 1
        Ideal 21551 40.0
                               20.0
## 2
      Premium 13791 25.6
                               52.8
## 3 Very Good 12082 22.4
                               76.8
## 4
         Good
                4906
                      9.1
                               92.5
```

• Create the pie charts using ggplot2 verbs. Key function: coord polar().

```
ggplot(df, aes(x = "", y = prop, fill = cut)) +
  geom_bar(width = 1, stat = "identity", color = "white") +
  geom_text(aes(y = lab.ypos, label = prop), color = "white")+
  coord_polar("y", start = 0)+
  ggpubr::fill_palette("jco")+
  theme_void()
```



• Alternative solution to easily create a pie chart: use the function ggpie()[in gg-pubr]:

```
ggpie(
  df, x = "prop", label = "prop",
  lab.pos = "in", lab.font = list(color = "white"),
  fill = "cut", color = "white",
  palette = "jco"
)
```

2.3.3 Dot charts

Dot chart is an alternative to bar plots. Key functions:

- geom_linerange():Creates line segments from x to ymax
- geom_point(): adds dots
- ggpubr::color_palette(): changes color palette.

```
ggplot(df, aes(cut, prop)) +
  geom_linerange(
  aes(x = cut, ymin = 0, ymax = prop),
  color = "lightgray", size = 1.5
```

Easy alternative to create a dot chart. Use ggdotchart() [ggpubr]:

```
ggdotchart(
  df, x = "cut", y = "prop",
  color = "cut", size = 3,  # Points color and size
  add = "segment",  # Add line segments
  add.params = list(size = 2),
  palette = "jco",
  ggtheme = theme_pubclean()
)
```

2.4 One continuous variable

Different types of graphs can be used to visualize the distribution of a continuous variable, including: density and histogram plots.

2.4.1 Data format

Create some data (wdata) containing the weights by sex (M for male; F for female):

```
## sex weight
## 1 F 53.8
## 2 F 55.3
## 3 F 56.1
## 4 F 52.7
```

Compute the mean weight by sex using the dplyr package. First, the data is grouped by sex and then summarized by computing the mean weight by groups. The operator %>% is used to combine multiple operations:

```
library("dplyr")
mu <- wdata %>%
  group_by(sex) %>%
  summarise(grp.mean = mean(weight))
## # A tibble: 2 x 2
##
        sex grp.mean
##
     <fctr>
                <dbl>
## 1
          F
                54.9
## 2
          М
                58.1
```

2.4.2 Basic plots

We start by creating a plot, named a, that we'll finish in the next section by adding a layer.

```
a <- ggplot(wdata, aes(x = weight))
```

Possible layers include: geom_density() (for density plots) and geom_histogram() (for histogram plots).

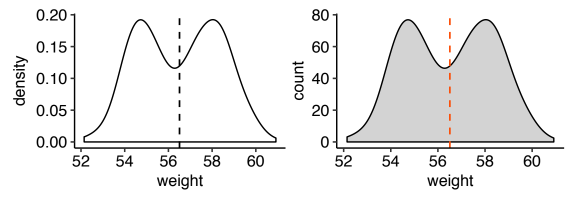
Key arguments to customize the plots:

- color, size, linetype: change the line color, size and type, respectively
- fill: change the areas fill color (for bar plots, histograms and density plots)
- alpha: create a semi-transparent color.

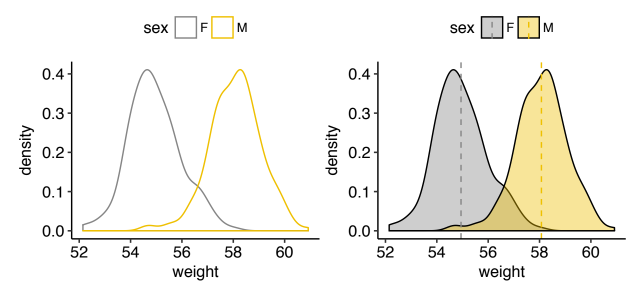
2.4.3 Density plots

Key function: geom density().

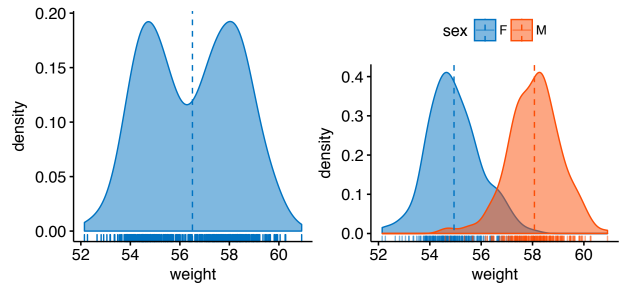
1. Create basic density plots. Add a vertical line corresponding to the mean value of the weight variable (geom vline()):



- 2. Change areas fill and add line color by groups (sex):
- Add vertical mean lines using geom_vline(). Data: mu, which contains the mean values of weights by sex (computed in the previous section).
- Change color manually:
 - use scale_color_manual() or scale_colour_manual() for changing line color
 - use scale_fill_manual() for changing area fill colors.



3. Simple solution to create a ggplot2-based density plots: use ggboxplot() [in ggpubr].

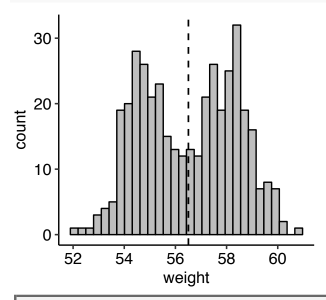


2.4.4 Histogram plots

An alternative to density plots is histograms, which represents the distribution of a continuous variable by dividing into bins and counting the number of observations in each bin.

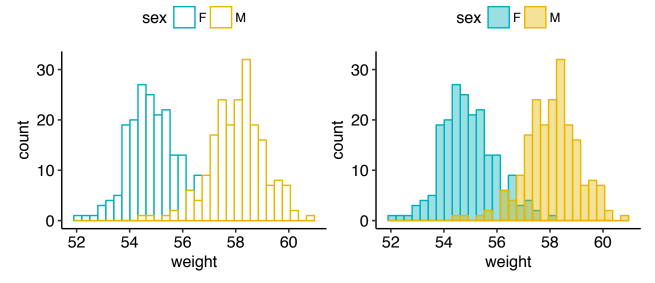
Key function: geom histogram(). The basic usage is quite similar to geom density().

1. **Create a basic plots**. Add a vertical line corresponding to the mean value of the weight variable:



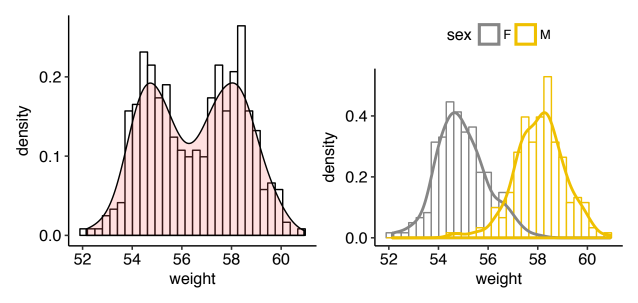
Note that, by default:

- By default, geom_histogram() uses 30 bins this might not be good default. You can change the number of bins (e.g.: bins = 50) or the bin width (e.g.: binwidth = 0.5)
- The y axis corresponds to the count of weight values. If you want to change the plot in order to have the density on y axis, specify the argument y = ..density.. in aes().
- 2. Change areas fill and add line color by groups (sex):
- Add vertical mean lines using geom_vline(). Data: mu, which contains the mean values of weights by sex.
- Change color manually:
 - use scale_color_manual() or scale_colour_manual() for changing line color
 - use scale_fill_manual() for changing area fill colors.
- Adjust the position of histogram bars by using the argument position. Allowed values: "identity", "stack", "dodge". Default value is "stack".

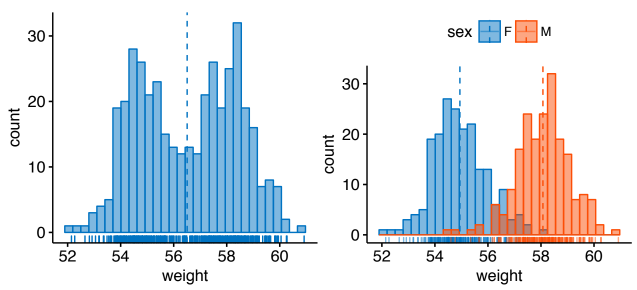


3. Combine histogram and density plots:

- Plot histogram with density values on y-axis (instead of count values).
- Add density plot with transparent density plot



4. Simple solution to create a ggplot2-based histogram plots: use gghistogram() [in ggpubr].



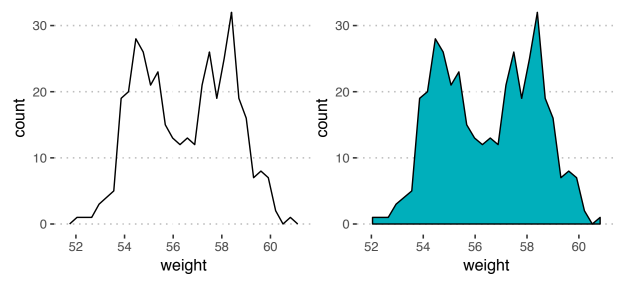
2.4.5 Alternative to density and histogram plots

- 1. **Frequency polygon**. Very close to histogram plots, but it uses lines instead of bars.
 - Key function: geom freqpoly().
 - Key arguments: color, size, linetype: change, respectively, line color, size and type.
- 2. **Area plots**. This is a continuous analog of a stacked bar plot.
 - Key function: geom area().
 - Key arguments:
 - color, size, linetype: change, respectively, line color, size and type.
 - fill: change area fill color.

In this section, we'll use the theme_pubclean() [in ggpubr]. This is a theme without axis lines, to direct more attention to the data. Type this to use the theme:

```
theme_set(theme_pubclean())
```

• Create a basic frequency polygon and basic area plots:



• Change colors by groups (sex):

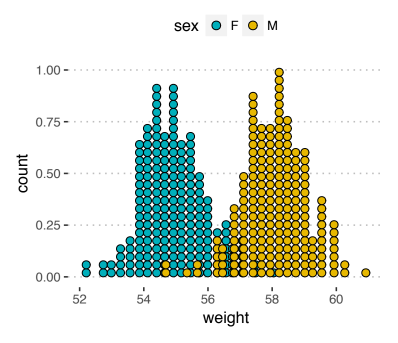
```
# Create a stacked area plots
a + geom_area(aes(fill = sex), color = "white",
               stat ="bin", bins = 30) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800"))
               sex F - M
                                                        sex
  20
                                           20
count
  10 -
                                           10 -
                          58
                                                     54
                                                                          60
              54
                    56
                                 60
                                                            56
                                              52
                                                                   58
       52
                   weight
                                                            weight
```

As in histogram plots, the default y values is count. To have density values on y axis, specify y = ..density.. in aes().

- 3. **Dot plots**. Represents another alternative to histograms and density plots, that can be used to visualize a continuous variable. Dots are stacked with each dot representing one observation. The width of a dot corresponds to the bin width.
- Key function: geom dotplot().
- Key arguments: alpha, color, fill and dotsize.

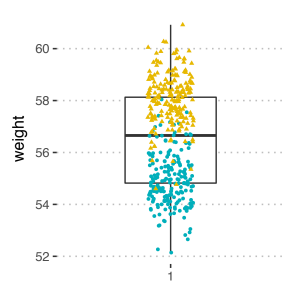
Create a dot plot colored by groups (sex):

```
a + geom_dotplot(aes(fill = sex), binwidth = 1/4) +
scale_fill_manual(values = c("#00AFBB", "#E7B800"))
```



4. Box plot:

- Create a box plot of one continuous variable: geom_boxplot()
- Add jittered points, where each point corresponds to an individual observation: geom_jitter(). Change the color and the shape of points by groups (sex)



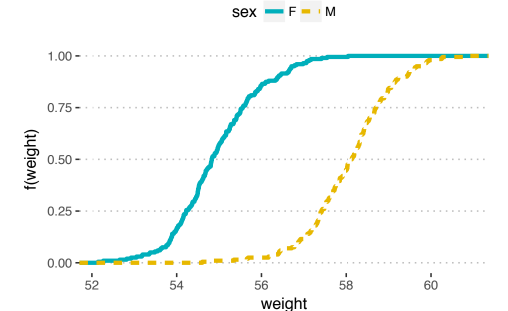
sex • F • M

5. Empirical cumulative distribution function (ECDF). Provides another alternative visualization of distribution. It reports for any given number the percent

of individuals that are below that threshold.

For example, in the following plots, you can see that:

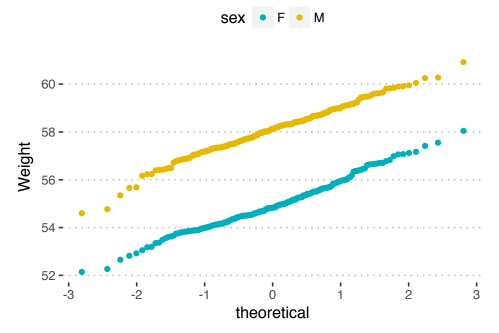
- about 25% of our females are shorter than 50 inches
- about 50% of males are shorter than 58 inches



- 6. Quantile-quantile plot (QQ plots). Used to check whether a given data follows normal distribution.
- Key function: stat qq().
- Key arguments: color, shape and size to change point color, shape and size.

Create a qq-plot of weight. Change color by groups (sex)

```
# Change point shapes by groups
ggplot(wdata, aes(sample = weight)) +
   stat_qq(aes(color = sex)) +
   scale_color_manual(values = c("#00AFBB", "#E7B800"))+
   labs(y = "Weight")
```



Alternative plot using the function ggqqplot() [in ggpubr]. The 95% confidence band is shown by default.

2.4.6 Density ridgeline plots

The density ridgeline plot is an alternative to the standard <code>geom_density()</code> function that can be useful for visualizing changes in distributions, of a continuous variable, over time

or space. Ridgeline plots are partially overlapping line plots that create the impression of a mountain range.

This functionality is provided in the R package ggridges (Wilke, 2017).

1. Installation:

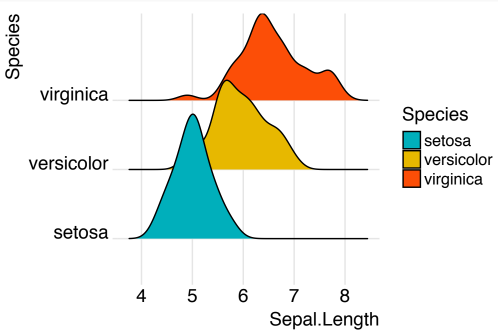
```
install.packages("ggridges")
```

2. Load and set the default theme to theme_ridges() [in ggridges]:

```
library(ggplot2)
library(ggridges)
theme_set(theme_ridges())
```

3. Example 1: Simple distribution plots by groups. Distribution of Sepal.Length by Species using the iris data set. The grouping variable Species will be mapped to the y-axis:

```
ggplot(iris, aes(x = Sepal.Length, y = Species)) +
  geom_density_ridges(aes(fill = Species)) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
```



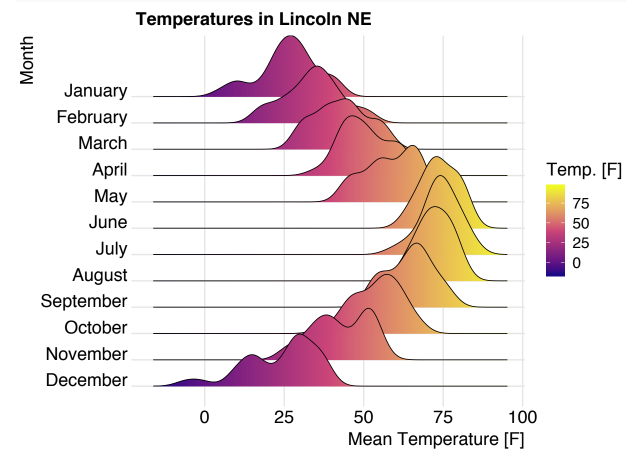
You can control the overlap between the different densities using the scale option. Default value is 1. Smaller values create a separation between the curves, and larger values create more overlap.

```
ggplot(iris, aes(x = Sepal.Length, y = Species)) +
  geom_density_ridges(scale = 0.9)
```

- 4. Example 4: Visualize temperature data.
- Data set: lincoln weather [in ggridges]. Weather in Lincoln, Nebraska in 2016.
- Create the density ridge plots of the Mean Temperature by Month and change the

fill color according to the temperature value (on x axis). A gradient color is created using the function geom density ridges gradient()

```
ggplot(
  lincoln_weather,
  aes(x = `Mean Temperature [F]`, y = `Month`)
) +
geom_density_ridges_gradient(
  aes(fill = ..x..), scale = 3, size = 0.3
) +
scale_fill_gradientn(
  colours = c("#0D0887FF", "#CC4678FF", "#F0F921FF"),
  name = "Temp. [F]"
) +
labs(title = 'Temperatures in Lincoln NE')
```



For more examples, type the following R code:

```
browseVignettes("ggridges")
```

2.4.7 Bar plot and modern alternatives

In this section, we'll describe how to create easily basic and ordered bar plots using ggplot2 based helper functions available in the ggpubr R package. We'll also present some modern alternatives to bar plots, including lollipop charts and cleveland's dot plots.

• Load required packages:

library(ggpubr)

Valiant

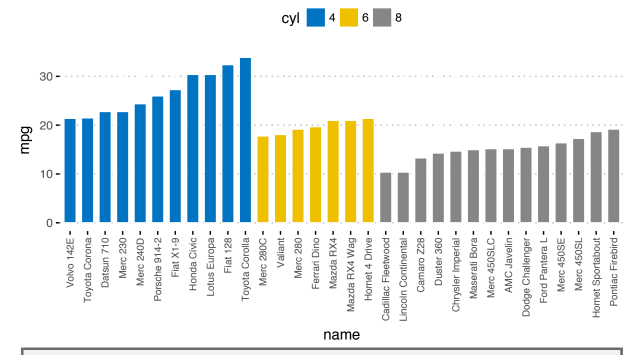
• Load and prepare data:

```
# Load data
dfm <- mtcars
# Convert the cyl variable to a factor
dfm$cyl <- as.factor(dfm$cyl)</pre>
# Add the name colums
dfm$name <- rownames(dfm)</pre>
# Inspect the data
head(dfm[, c("name", "wt", "mpg", "cyl")])
##
                                  name
                                          wt mpg cyl
## Mazda RX4
                             Mazda RX4 2.62 21.0
## Mazda RX4 Wag
                        Mazda RX4 Wag 2.88 21.0
                                                    6
## Datsun 710
                            Datsun 710 2.32 22.8
## Hornet 4 Drive
                        Hornet 4 Drive 3.21 21.4
                                                    6
## Hornet Sportabout Hornet Sportabout 3.44 18.7
                                                    8
```

• Create an ordered bar plot of the mpg variable. Change the fill color by the grouping variable "cyl". Sorting will be done globally, but not by groups.

Valiant 3.46 18.1

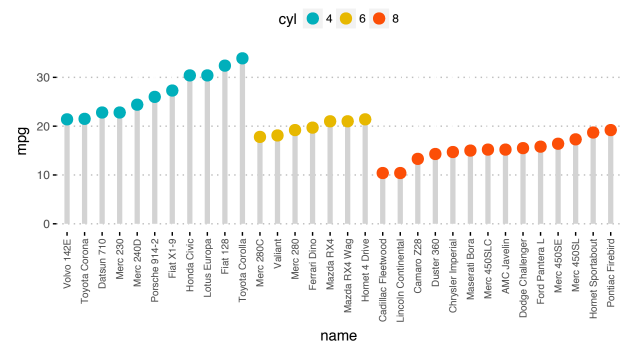
6



To sort bars inside each group, use the argument sort.by.groups = TRUE

- Create a Lollipop chart:
 - Color by groups and set a custom color palette.
 - Sort values in ascending order.
 - Add segments from y = 0 to dots. Change segment color and size.

2.5. CONCLUSION 35



Read more: Bar Plots and Modern Alternatives¹

2.5 Conclusion

• Create a bar plot of a grouping variable:

```
ggplot(diamonds, aes(cut)) +
geom_bar(fill = "#0073C2FF") +
theme_minimal()
```

• Visualize a continuous variable:

Start by creating a plot, named a, that we'll be finished by adding a layer.

```
a <- ggplot(wdata, aes(x = weight))
```

Possible layers include:

```
geom_density(): density plot
geom_histogram(): histogram plot
geom_freqpoly(): frequency polygon
geom_area(): area plot
geom_dotplot(): dot plot
stat_ecdf(): empirical cumulative density function
stat_qq(): quantile - quantile plot
```

Key arguments to customize the plots:

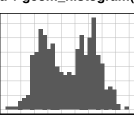
- color, size, linetype: change the line color, size and type, respectively
- fill: change the areas fill color (for bar plots, histograms and density plots)

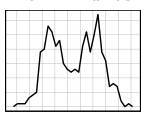
¹https://goo.gl/eSggcW

• alpha: create a semi-transparent color.

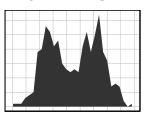
a + geom_density()

a + geom_histogram() a + geom_freqpoly()

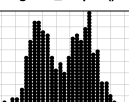




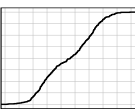
a + geom_area()



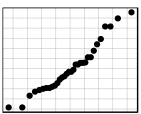
a + geom_dotplot()



a + stat_ecdf()



 $x + stat_qq()$



Chapter 3

Plot Grouped Data

3.1 Introduction

In this chapter, we start by describing how to plot grouped or stacked frequencies of two categorical variables. Tis can be done using bar plots and dot charts. You'll also learn how to add labels to dodged and stacked bar plots.

Next we'll show how to display a continuous variable with multiple groups. In this situation, the grouping variable is used as the x-axis and the continuous variable as the y-axis. You'll learn, how to:

- Visualize a grouped continuous variable using **box plot**, **violin plots**, **stripcharts** and alternatives.
- Add automatically t-test / wilcoxon test p-values comparing groups.
- Create mean and median plots of groups with error bars

3.2 Prerequisites

Load required packages and set the theme function theme_pubclean() [in ggpubr] as the default theme:

```
library(dplyr)
library(ggplot2)
library(ggpubr)
theme_set(theme_pubclean())
```

3.3 Grouped categorical variables

- Plot types: grouped bar plots of the frequencies of the categories. Key function: geom_bar().
- Demo dataset: diamonds [in ggplot2]. The categorical variables to be used in the demo example are:

cut: quality of the diamonds cut (Fair, Good, Very Good, Premium, Ideal)
color: diamond colour, from J (worst) to D (best).

In our demo example, we'll plot only a subset of the data (color J and D). The different steps are as follow:

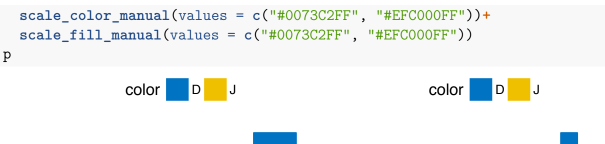
- Filter the data to keep only diamonds which colors are in ("J", "D").
- Group the data by the quality of the cut and the diamond color
- Count the number of records by groups
- Create the bar plot
- 1. Filter and count the number of records by groups:

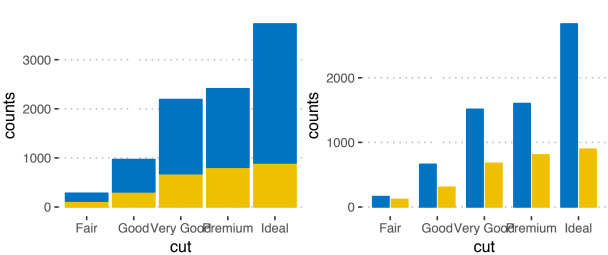
```
df <- diamonds %>%
  filter(color %in% c("J", "D")) %>%
  group_by(cut, color) %>%
  summarise(counts = n())
head(df, 4)
## # A tibble: 4 x 3
## # Groups:
               cut [2]
##
       cut color counts
##
     <ord> <ord>
                 <int>
## 1 Fair
               D
                    163
## 2 Fair
               J
                    119
## 3 Good
               D
                    662
## 4 Good
               J
                    307
```

- 2. Creare the grouped bar plots:
 - Key function: geom_bar(). Key argument: stat = "identity" to plot the data as it is.
 - Use the functions scale_color_manual() and scale_fill_manual() to set manually the bars border line colors and area fill colors.

```
# Stacked bar plots of y = counts by x = cut,
# colored by the variable color
ggplot(df, aes(x = cut, y = counts)) +
geom_bar(
    aes(color = color, fill = color),
    stat = "identity", position = position_stack()
    ) +
scale_color_manual(values = c("#0073C2FF", "#EFC000FF"))+
scale_fill_manual(values = c("#0073C2FF", "#EFC000FF"))

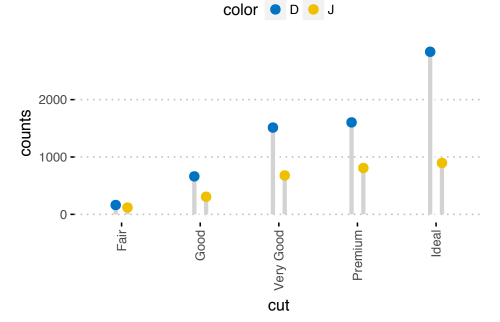
# Use position = position_dodge()
p <- ggplot(df, aes(x = cut, y = counts)) +
geom_bar(
    aes(color = color, fill = color),
    stat = "identity", position = position_dodge(0.8),
    width = 0.7
    ) +</pre>
```





Note that, position_stack() automatically stack values in reverse order of the group aesthetic. This default ensures that bar colors align with the default legend. You can change this behavior by using position = position_stack(reverse = TRUE).

Alternatively, you can easily create a dot chart with the ggpubr package:

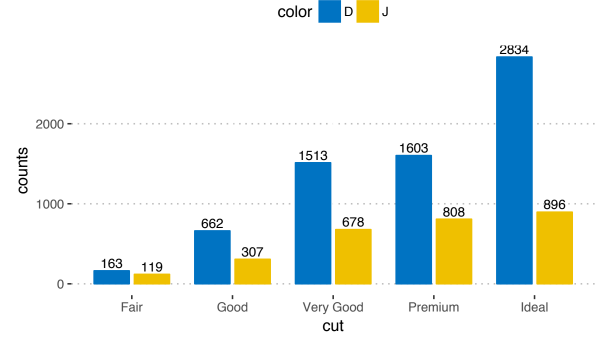


Or, if you prefer the ggplot2 verbs, type this:

```
ggplot(df, aes(cut, counts)) +
  geom_linerange(
    aes(x = cut, ymin = 0, ymax = counts, group = color),
    color = "lightgray", size = 1.5,
    position = position_dodge(0.3)
    )+
  geom_point(
    aes(color = color),
    position = position_dodge(0.3), size = 3
    )+
  scale_color_manual(values = c("#0073C2FF", "#EFC000FF"))+
  theme_pubclean()
```

3. Add labels to the dodged bar plots:

```
p + geom_text(
  aes(label = counts, group = color),
  position = position_dodge(0.8),
  vjust = -0.3, size = 3.5
)
```

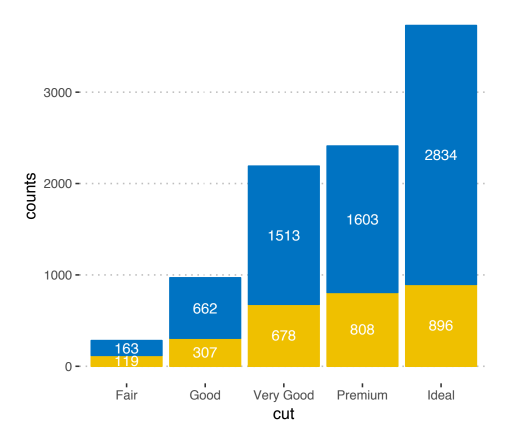


- 4. Add labels to a stacked bar plots. 3 steps required to compute the position of text labels:
 - Sort the data by cut and color columns. As position_stack() reverse the group order, color column should be sorted in descending order.
 - Calculate the cumulative sum of counts for each cut category. Used as the y coordinates of labels. To put the label in the middle of the bars, we'll use cumsum(counts) 0.5 * counts.
 - Create the bar graph and add labels

```
# Arrange/sort and compute cumulative summs
 df <- df %>%
  arrange(cut, desc(color)) %>%
  mutate(lab_ypos = cumsum(counts) - 0.5 * counts)
head(df, 4)
## # A tibble: 4 x 4
## # Groups:
               cut [2]
##
       cut color counts lab ypos
##
     <ord> <ord>
                  <int>
                            <dbl>
## 1 Fair
               J
                    119
                             59.5
## 2
     Fair
               D
                    163
                            200.5
## 3
      Good
               J
                    307
                            153.5
## 4 Good
               D
                    662
                           638.0
# Create stacked bar graphs with labels
ggplot(df, aes(x = cut, y = counts)) +
  geom_bar(aes(color = color, fill = color), stat = "identity") +
  geom_text(
    aes(y = lab_ypos, label = counts, group = color),
    color = "white"
```

```
scale_color_manual(values = c("#0073C2FF", "#EFC000FF"))+
scale_fill_manual(values = c("#0073C2FF", "#EFC000FF"))

color
D
J
```



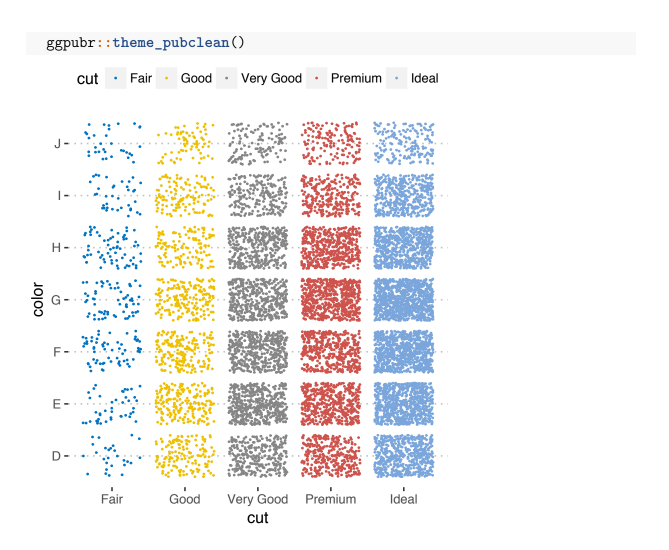
Alternatively, you can easily create the above plot using the function ggbarplot() [in ggpubr]:

6. Alternative to bar plots. Instead of the creating a bar plot of the counts, you can plot two discrete variables with discrete x-axis and discrete y-axis. Each individual points are shown by groups. For a given group, the number of points corresponds to the number of records in that group.

Key function: geom_jitter(). Arguments: alpha, color, fill, shape and size.

In the example below, we'll plot a small fraction (1/5) of the diamonds dataset.

```
diamonds.frac <- dplyr::sample_frac(diamonds, 1/5)
ggplot(diamonds.frac, aes(cut, color)) +
  geom_jitter(aes(color = cut), size = 0.3)+
  ggpubr::color_palette("jco")+</pre>
```



3.4 Grouped continuous variables

In this section, we'll show to plot a grouped continuous variable using box plot, violin plot, strip chart and alternatives.

We'll also describe how to add automatically p-values comparing groups.

In this section, we'll set the theme theme_bw() as the default ggplot theme:

```
theme_set(
  theme_bw()
)
```

3.4.1 Data format

- Demo dataset: ToothGrowth
 - Continuous variable: len (tooth length). Used on y-axis
 - Grouping variable: dose (dose levels of vitamin C: 0.5, 1, and 2 mg/day). Used on x-axis.

First, convert the variable dose from a numeric to a discrete factor variable:

```
data("ToothGrowth")
ToothGrowth$dose <- as.factor(ToothGrowth$dose)
head(ToothGrowth)</pre>
```

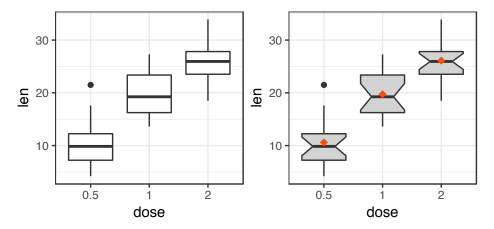
```
## 1 4.2 VC 0.5
## 2 11.5 VC 0.5
## 3 7.3 VC 0.5
## 4 5.8 VC 0.5
## 5 6.4 VC 0.5
## 6 10.0 VC 0.5
```

3.4.2 Box plots

- Key function: geom boxplot()
- Key arguments to customize the plot:
 - width: the width of the box plot
 - notch: logical. If TRUE, creates a notched box plot. The notch displays a confidence interval around the median which is normally based on the median +/- 1.58*IQR/sqrt(n). Notches are used to compare groups; if the notches of two boxes do not overlap, this is a strong evidence that the medians differ.
 - color, size, linetype: Border line color, size and type
 - fill: box plot areas fill color
 - outlier.colour, outlier.shape, outlier.size: The color, the shape and the size for outlying points.

1. Create basic box plots:

• Standard and notched box plots:



• Change box plot colors by groups:

```
# Color by group (dose)
e + geom_boxplot(aes(color = dose))+
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
# Change fill color by group (dose)
e + geom_boxplot(aes(fill = dose)) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
  30
                                          30
                                  dose
                                                                          dose
                                                                            0.5
                                        <u>e</u>u
                                          20
<u>e</u>u
  20
  10
                                           10
                         2
         0.5
                 i
                                                 0.5
                                                                  ż
                dose
                                                        dose
```

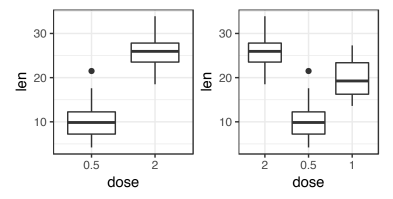
Note that, it's possible to use the function scale x discrete() for:

- choosing which items to display: for example c("0.5", "2"),
- changing the order of items: for example from c("0.5", "1", "2") to c("2", "0.5", "1")

For example, type this:

```
# Choose which items to display: group "0.5" and "2"
e + geom_boxplot() +
    scale_x_discrete(limits=c("0.5", "2"))

# Change the default order of items
e + geom_boxplot() +
    scale_x_discrete(limits=c("2", "0.5", "1"))
```

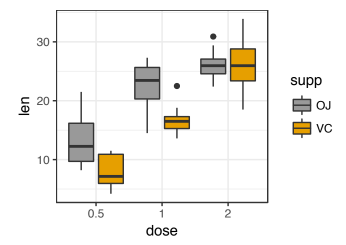


2. Create a box plot with multiple groups:

Two different grouping variables are used: dose on x-axis and supp as fill color (legend variable).

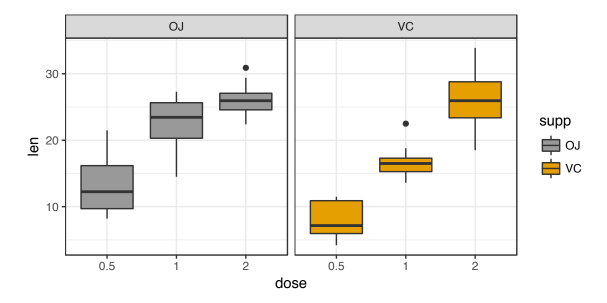
The space between the grouped box plots is adjusted using the function position_dodge().

```
e2 <- e + geom_boxplot(
  aes(fill = supp),
  position = position_dodge(0.9)
  ) +
  scale_fill_manual(values = c("#9999999", "#E69F00"))
e2</pre>
```



Split the plot into multiple panel. Use the function facet_wrap():

```
e2 + facet_wrap(~supp)
```



3.4.3 Violin plots

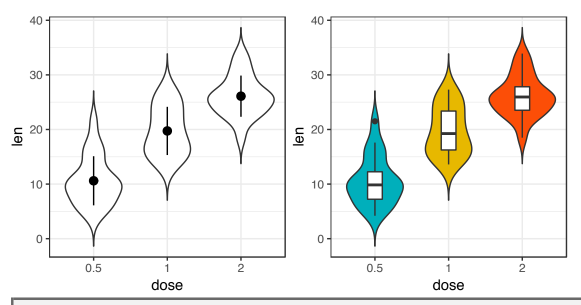
Violin plots are similar to box plots, except that they also show the kernel probability density of the data at different values. Typically, violin plots will include a marker for the median of the data and a box indicating the interquartile range, as in standard box plots.

Key function:

- geom violin(): Creates violin plots. Key arguments:
 - color, size, linetype: Border line color, size and type
 - fill: Areas fill color
 - trim: logical value. If TRUE (default), trim the tails of the violins to the range of the data. If FALSE, don't trim the tails.
- stat summary(): Adds summary statistics (mean, median, ...) on the violin plots.
- 1. Create basic violin plots with summary statistics:

```
# Add mean points +/- SD
# Use geom = "pointrange" or geom = "crossbar"
e + geom_violin(trim = FALSE) +
    stat_summary(
    fun.data = "mean_sdl", fun.args = list(mult = 1),
        geom = "pointrange", color = "black"
    )

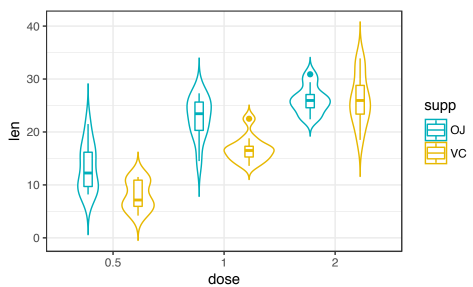
# Combine with box plot to add median and quartiles
# Change color by groups
e + geom_violin(aes(fill = dose), trim = FALSE) +
    geom_boxplot(width = 0.2)+
    scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))+
    theme(legend.position = "none")
```



The function $mean_sdl$ is used for adding mean and standard deviation. It computes the mean plus or minus a constant times the standard deviation. In the R code above, the constant is specified using the argument mult (mult = 1). By default mult = 2. The mean +/- SD can be added as a crossbar or a pointrange.

2. Create violin plots with multiple groups:

```
e + geom_violin(
  aes(color = supp), trim = FALSE,
  position = position_dodge(0.9)
  ) +
  geom_boxplot(
   aes(color = supp), width = 0.15,
   position = position_dodge(0.9)
   ) +
  scale_color_manual(values = c("#00AFBB", "#E7B800"))
```



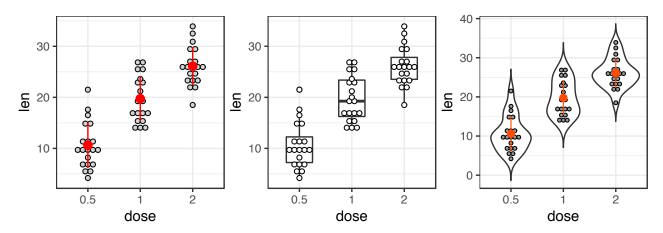
3.4.4 Dot plots

- Key function: geom_dotplot(). Creates stacked dots, with each dot representing one observation.
- Key arguments:
 - stackdir: which direction to stack the dots. "up" (default), "down", "center", "centerwhole" (centered, but with dots aligned).
 - stackratio: how close to stack the dots. Default is 1, where dots just just touch. Use smaller values for closer, overlapping dots.
 - color, fill: Dot border color and area fill
 - dotsize: The diameter of the dots relative to binwidth, default 1.

As for violin plots, summary statistics are usually added to dot plots.

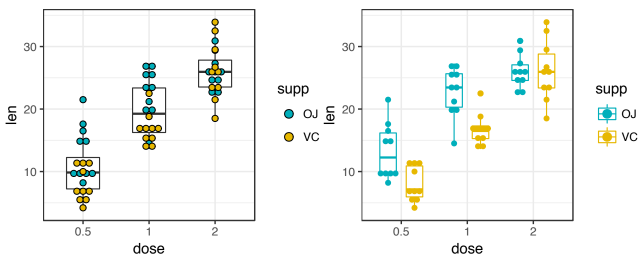
1. Create basic dot plots:

```
# Violin plots with mean points +/- SD
e + geom_dotplot(
 binaxis = "y", stackdir = "center",
 fill = "lightgray"
 ) +
 stat_summary(
    fun.data = "mean_sdl", fun.args = list(mult=1),
    geom = "pointrange", color = "red"
# Combine with box plots
e + geom_boxplot(width = 0.5) +
 geom_dotplot(
   binaxis = "y", stackdir = "center",
   fill = "white"
# Dot plot + violin plot + stat summary
e + geom_violin(trim = FALSE) +
 geom_dotplot(
   binaxis='y', stackdir='center',
    color = "black", fill = "#999999"
    ) +
 stat_summary(
    fun.data="mean_sdl", fun.args = list(mult=1),
    geom = "pointrange", color = "#FC4E07", size = 0.4
```



2. Create dot plots with multiple groups:

```
# Color dots by groups
e + geom_boxplot(width = 0.5, size = 0.4) +
 geom_dotplot(
    aes(fill = supp), trim = FALSE,
    binaxis='y', stackdir='center'
 )+
 scale_fill_manual(values = c("#00AFBB", "#E7B800"))
# Change the position : interval between dot plot of the same group
e + geom_boxplot(
 aes(color = supp), width = 0.5, size = 0.4,
 position = position_dodge(0.8)
  ) +
 geom_dotplot(
    aes(fill = supp, color = supp), trim = FALSE,
   binaxis='y', stackdir='center', dotsize = 0.8,
   position = position_dodge(0.8)
 )+
 scale_fill_manual(values = c("#00AFBB", "#E7B800"))+
 scale_color_manual(values = c("#00AFBB", "#E7B800"))
```

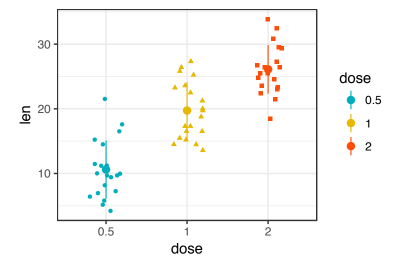


3.4.5 Stripcharts

Stripcharts are also known as one dimensional scatter plots. These plots are suitable compared to box plots when sample sizes are small.

- Key function: geom_jitter()
- key arguments: color, fill, size, shape. Changes points color, fill, size and shape
- 1. Create a basic stripchart:
- Change points shape and color by groups
- Adjust the degree of jittering: position_jitter(0.2)
- Add summary statistics:

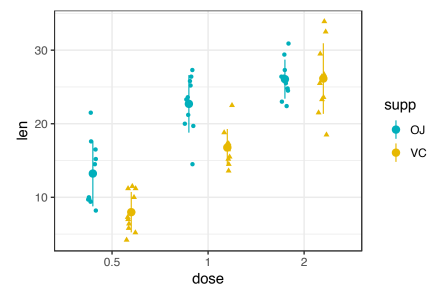
```
e + geom_jitter(
  aes(shape = dose, color = dose),
  position = position_jitter(0.2),
  size = 1.2
) +
  stat_summary(
   aes(color = dose),
   fun.data="mean_sdl", fun.args = list(mult=1),
   geom = "pointrange", size = 0.4
  )+
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
```



2. Create stripcharts for multiple groups. The R code is similar to what we have seen in dot plots section. However, to create dodged jitter points, you should use the function position_jitterdodge() instead of position_dodge().

```
e + geom_jitter(
  aes(shape = supp, color = supp),
  position = position_jitterdodge(jitter.width = 0.2, dodge.width = 0.8),
  size = 1.2
  ) +
  stat_summary(
```

```
aes(color = supp),
fun.data="mean_sdl", fun.args = list(mult=1),
geom = "pointrange", size = 0.4,
position = position_dodge(0.8)
)+
scale_color_manual(values = c("#00AFBB", "#E7B800"))
```



3.4.6 Sinaplot

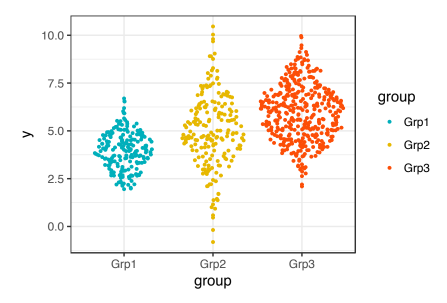
sinaplot is inspired by the strip chart and the violin plot. By letting the normalized density of points restrict the jitter along the x-axis, the plot displays the same contour as a violin plot, but resemble a simple strip chart for small number of data points (Sidiropoulos et al., 2015).

In this way the plot conveys information of both the number of data points, the density distribution, outliers and spread in a very simple, comprehensible and condensed format.

Key function: geom_sina() [ggforce]:

```
library(ggforce)
# Create some data
d1 <- data.frame(
    y = c(rnorm(200, 4, 1), rnorm(200, 5, 2), rnorm(400, 6, 1.5)),
    group = rep(c("Grp1", "Grp2", "Grp3"), c(200, 200, 400))
)

# Sinaplot
ggplot(d1, aes(group, y)) +
    geom_sina(aes(color = group), size = 0.7)+
    scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))</pre>
```



3.4.7 Mean and median plots with error bars

In this section, we'll show how to plot summary statistics of a continuous variable organized into groups by one or multiple grouping variables.

Note that, an easy way, with less typing, to create mean/median plots, is provided in the ggpubr package. See the associated article at: ggpubr-Plot Means/Medians and Error Bars¹

Set the default theme to theme_pubr() [in ggpubr]:

```
theme_set(ggpubr::theme_pubr())
```

- 1. Basic mean/median plots. Case of one continuous variable and one grouping variable:
- Prepare the data: ToothGrowth data set.

```
df <- ToothGrowth
df$dose <- as.factor(df$dose)
head(df, 3)</pre>
```

```
## 1 4.2 VC 0.5
## 2 11.5 VC 0.5
## 3 7.3 VC 0.5
```

• Compute summary statistics for the variable len organized into groups by the variable dose:

```
library(dplyr)
df.summary <- df %>%
  group_by(dose) %>%
  summarise(
```

¹http://www.sthda.com/english/articles/24-ggpubr-publication-ready-plots/

```
sd = sd(len, na.rm = TRUE),
len = mean(len)
)
df.summary
```

```
## # A tibble: 3 x 3
## dose sd len
## <fctr> <dbl> <dbl> ## 1 0.5 4.50 10.6
## 2 1 4.42 19.7
## 3 2 3.77 26.1
```

- Create error plots using the summary statistics data. Key functions:
 - geom_crossbar() for hollow bar with middle indicated by horizontal line
 - geom_errorbar() for error bars
 - geom_errorbarh() for horizontal error bars
 - geom_linerange() for drawing an interval represented by a vertical line
 - geom_pointrange() for creating an interval represented by a vertical line,
 with a point in the middle.

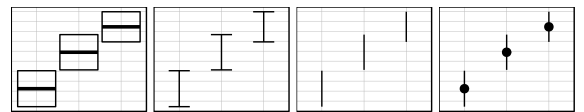
Start by initializing ggplot with the summary statistics data:

- Specify x and y as usually - Specify ymin = len-sd and ymax = len+sd to add lower and upper error bars. If you want only to add upper error bars but not the lower ones, use ymin = len (instead of len-sd) and ymax = len+sd.

```
# Initialize ggplot with data
f <- ggplot(
  df.summary,
  aes(x = dose, y = len, ymin = len-sd, ymax = len+sd)
)</pre>
```

Possible error plots:

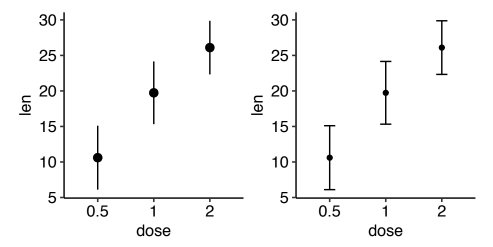
f + geom_crossbar() f + geom_errorbar() f + geom_linerange() f + geom_pointrange()



Create simple error plots:

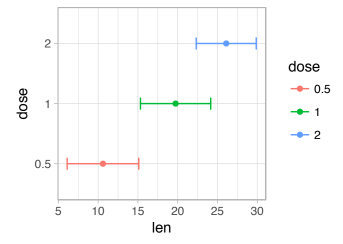
```
# Vertical line with point in the middle
f + geom_pointrange()

# Standard error bars
f + geom_errorbar(width = 0.2) +
geom_point(size = 1.5)
```



Create horizontal error bars. Put dose on y axis and len on x-axis. Specify xmin and xmax.

```
# Horizontal error bars with mean points
# Change the color by groups
ggplot(
    df.summary,
    aes(x = len, y = dose, xmin = len-sd, xmax = len+sd)
    ) +
    geom_point(aes(color = dose)) +
    geom_errorbarh(aes(color = dose), height=.2)+
    theme_light()
```

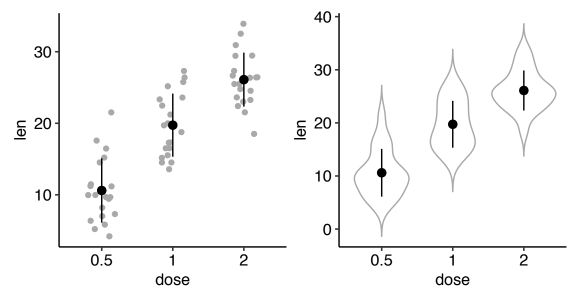


• Add jitter points (representing individual points), dot plots and violin plots. For this, you should initialize ggplot with original data (df) and specify the df.summary data in the error plot function, here geom_pointrange().

```
# Combine with jitter points
ggplot(df, aes(dose, len)) +
geom_jitter(
   position = position_jitter(0.2), color = "darkgray"
   ) +
geom_pointrange(
   aes(ymin = len-sd, ymax = len+sd),
```

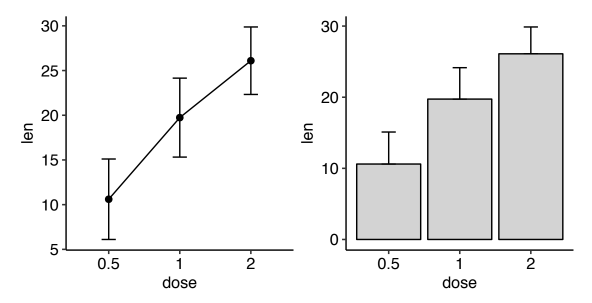
```
data = df.summary
)

# Combine with violin plots
ggplot(df, aes(dose, len)) +
  geom_violin(color = "darkgray", trim = FALSE) +
  geom_pointrange(
   aes(ymin = len-sd, ymax = len+sd),
   data = df.summary
)
```



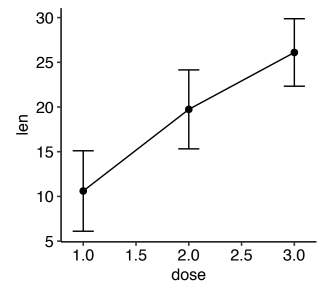
- Create basic bar/line plots of mean +/- error. So we need only the df.summary data.
 - (1) Add lower and upper error bars for the line plot: ymin = len-sd and ymax = len+sd.
 - (2) Add only upper error bars for the bar plot: ymin = len (instead of len-sd) and ymax = len+sd.

Note that, for line plot, you should always specify group = 1 in the aes(), when you have one group of line.



For line plot, you might want to treat x-axis as numeric:

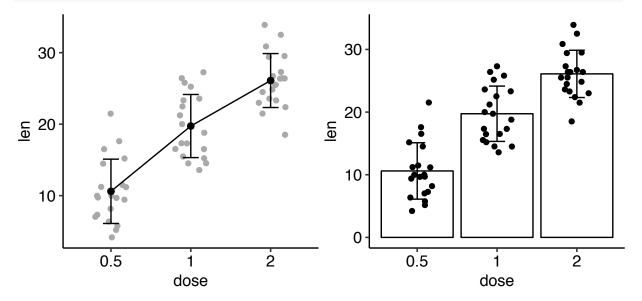
```
df.sum2 <- df.summary
df.sum2$dose <- as.numeric(df.sum2$dose)
ggplot(df.sum2, aes(dose, len)) +
   geom_line() +
   geom_errorbar( aes(ymin = len-sd, ymax = len+sd), width = 0.2) +
   geom_point(size = 2)</pre>
```



- Bar and line plots + jitter points. We need the original df data for the jitter points and the df.summary data for the other geom layers.
 - (1) For the line plot: First, add jitter points, then add lines + error bars + mean points on top of the jitter points.
 - (2) For the bar plot: First, add the bar plot, then add jitter points + error bars on top of the bars.

```
# (1) Create a line plot of means +
# individual jitter points + error bars
ggplot(df, aes(dose, len)) +
```

```
geom_jitter( position = position_jitter(0.2),
               color = "darkgray") +
 geom_line(aes(group = 1), data = df.summary) +
  geom errorbar(
    aes(ymin = len-sd, ymax = len+sd),
    data = df.summary, width = 0.2) +
 geom_point(data = df.summary, size = 2)
# (2) Bar plots of means + individual jitter points + errors
ggplot(df, aes(dose, len)) +
  geom_bar(stat = "identity", data = df.summary,
           fill = NA, color = "black") +
 geom_jitter( position = position_jitter(0.2),
               color = "black") +
 geom_errorbar(
    aes(ymin = len-sd, ymax = len+sd),
    data = df.summary, width = 0.2)
```



- 2. Mean/median plots for multiple groups. Case of one continuous variable (len) and two grouping variables (dose, supp).
- Compute the summary statistics of len grouped by dose and supp:

```
library(dplyr)
df.summary2 <- df %>%
  group_by(dose, supp) %>%
  summarise(
   sd = sd(len),
   len = mean(len)
  )
df.summary2
```

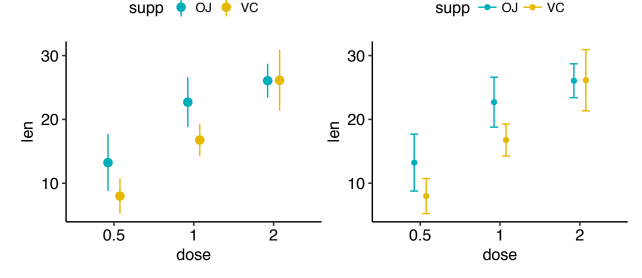
```
## # A tibble: 6 x 4
## # Groups: dose [?]
```

```
##
       dose
               supp
                        sd
                              len
##
     <fctr> <fctr> <dbl> <dbl>
                      4.46 13.23
## 1
        0.5
                 OJ
## 2
        0.5
                 VC
                      2.75 7.98
## 3
                      3.91 22.70
           1
                 OJ
## 4
           1
                 VC
                      2.52 16.77
## 5
           2
                      2.66 26.06
                 OJ
           2
## 6
                 VC
                      4.80 26.14
```

- Create error plots for multiple groups:
 - (1) pointrange colored by groups (supp)
 - (2) standard error bars + mean points colored by groups (supp)

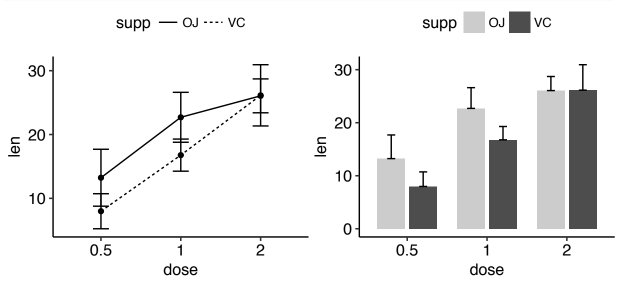
```
# (1) Pointrange: Vertical line with point in the middle
ggplot(df.summary2, aes(dose, len)) +
  geom_pointrange(
    aes(ymin = len-sd, ymax = len+sd, color = supp),
    position = position_dodge(0.3)
    )+
  scale_color_manual(values = c("#00AFBB", "#E7B800"))

# (2) Standard error bars
ggplot(df.summary2, aes(dose, len)) +
  geom_errorbar(
    aes(ymin = len-sd, ymax = len+sd, color = supp),
    position = position_dodge(0.3), width = 0.2
    )+
  geom_point(aes(color = supp), position = position_dodge(0.3)) +
  scale_color_manual(values = c("#00AFBB", "#E7B800"))
```

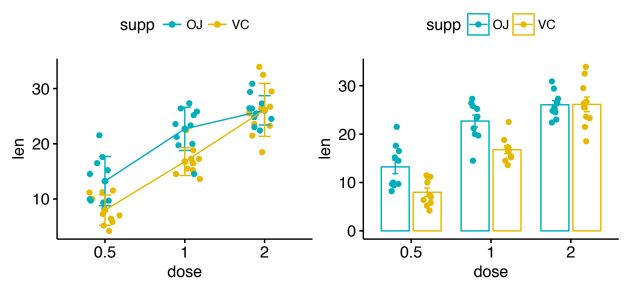


- Create simple line/bar plots for multiple groups.
 - (1) Line plots: change linetype by groups (supp)
 - (2) Bar plots: change fill color by groups (supp)

```
# (1) Line plot + error bars
ggplot(df.summary2, aes(dose, len)) +
 geom_line(aes(linetype = supp, group = supp))+
 geom_point()+
 geom_errorbar(
    aes(ymin = len-sd, ymax = len+sd, group = supp),
    width = 0.2
    )
# (2) Bar plots + upper error bars.
ggplot(df.summary2, aes(dose, len)) +
 geom_bar(aes(fill = supp), stat = "identity",
           position = position_dodge(0.8), width = 0.7)+
 geom_errorbar(
    aes(ymin = len, ymax = len+sd, group = supp),
    width = 0.2, position = position_dodge(0.8)
    )+
  scale_fill_manual(values = c("grey80", "grey30"))
```



• Create easily plots of mean +/- sd for multiple groups. Use the ggpubr package, which will automatically calculate the summary statistics and create the graphs.



• Use the standard ggplot2 verbs, to reproduce the line plots above:

```
# Create line plots
ggplot(df, aes(dose, len)) +
geom_jitter(
   aes(color = supp),
   position = position_jitter(0.2)
   ) +
geom_line(
   aes(group = supp, color = supp),
   data = df.summary2
   ) +
geom_errorbar(
   aes(ymin = len-sd, ymax = len+sd, color = supp),
   data = df.summary2, width = 0.2
   )+
scale_color_manual(values = c("#00AFBB", "#E7B800"))
```

3.4.8 Add p-values and significance levels

In this section, we'll describe how to easily i) compare means of two or multiple groups; ii) and to automatically add p-values and significance levels to a ggplot (such as box plots, dot plots, bar plots and line plots, ...).

Key functions:

- compare_means() [ggpubr package]: easy to use solution to performs one and multiple mean comparisons.
- stat_compare_means() [ggpubr package]: easy to use solution to automatically add p-values and significance levels to a ggplot.

The most common methods for comparing means² include:

²http://www.sthda.com/english/wiki/comparing-means-in-r

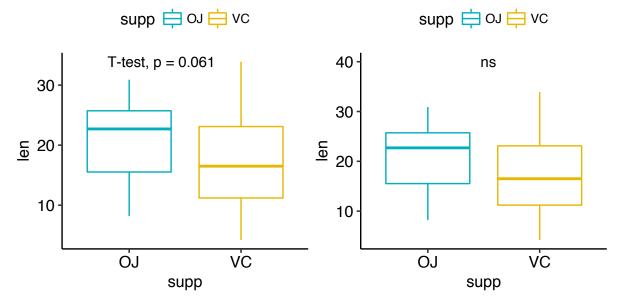
Methods	R function	Description
T-test	t.test()	Compare two groups (parametric)
Wilcoxon test	wilcox.test()	Compare two groups (non-parametric)
ANOVA	aov() or anova()	Compare multiple groups (parametric)
Kruskal-Wallis	kruskal.test()	Compare multiple groups (non-parametric)

- 1. Compare two independent groups:
- Compute t-test:

• Create a box plot with p-values. Use the option method = "t.test" or method = "wilcox.test". Default is wilcoxon test.

```
# Create a simple box plot and add p-values
p <- ggplot(ToothGrowth, aes(supp, len)) +
    geom_boxplot(aes(color = supp)) +
    scale_color_manual(values = c("#00AFBB", "#E7B800"))
p + stat_compare_means(method = "t.test")

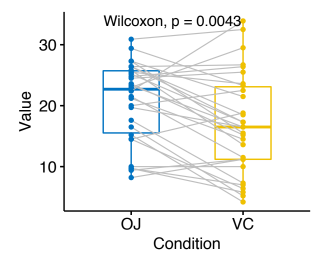
# Display the significance level instead of the p-value
# Adjust label position
p + stat_compare_means(
    aes(label = ..p.signif..), label.x = 1.5, label.y = 40
    )</pre>
```



2. Compare two paired samples. Use ggpaired() [ggpubr] to create the paired

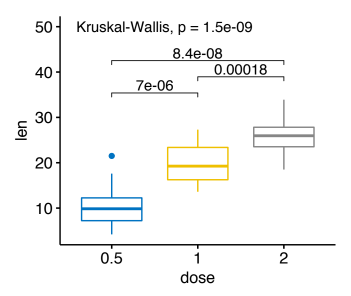
box plot.

```
supp 🖨 OJ 嶭 VC
```



3. Compare more than two groups. If the grouping variable contains more than two levels, then pairwise tests will be performed automatically. The default method is "wilcox.test". You can change this to "t.test".

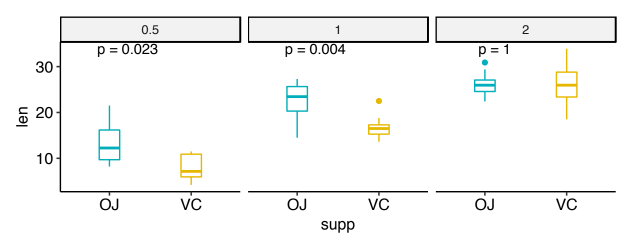
```
# Perorm pairwise comparisons
compare_means(len ~ dose, data = ToothGrowth)
## # A tibble: 3 x 8
       .y. group1 group2
##
                                      p.adj p.format p.signif
                                                                 method
                                 p
##
           <chr> <chr>
     <chr>
                             <dbl>
                                      <dbl>
                                               <chr>
                                                                  <chr>>
## 1
       len
              0.5
                       1 7.02e-06 1.40e-05
                                             7.0e-06
                                                          **** Wilcoxon
## 2
       len
              0.5
                       2 8.41e-08 2.52e-07 8.4e-08
                                                          **** Wilcoxon
## 3
                       2 1.77e-04 1.77e-04 0.00018
       len
                1
                                                           *** Wilcoxon
# Visualize: Specify the comparisons you want
my_{comparisons} \leftarrow list(c("0.5", "1"), c("1", "2"), c("0.5", "2"))
ggboxplot(ToothGrowth, x = "dose", y = "len",
          color = "dose", palette = "jco")+
  stat_compare_means(comparisons = my comparisons)+
  stat_compare_means(label.y = 50)
```



- 4. Multiple grouping variables:
- (1/2). Create a multi-panel box plots facetted by group (here, "dose"):

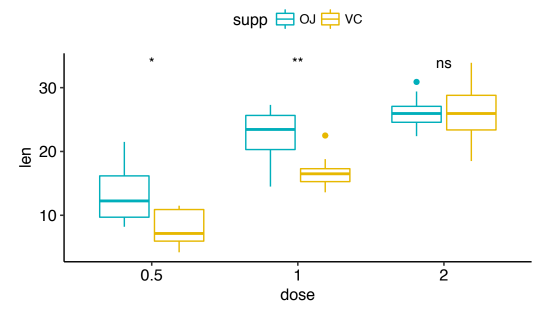
```
# Use only p.format as label. Remove method name.
ggplot(ToothGrowth, aes(supp, len)) +
  geom_boxplot(aes(color = supp))+
  facet_wrap(~dose) +
  scale_color_manual(values = c("#00AFBB", "#E7B800")) +
  stat_compare_means(label = "p.format")
```





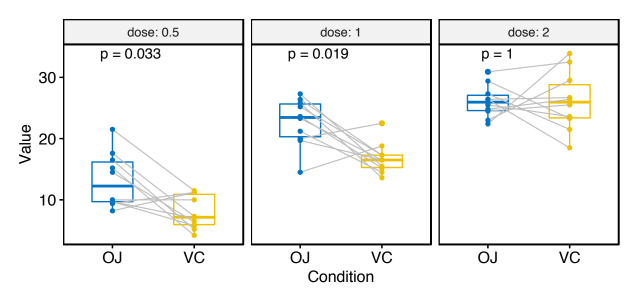
• (2/2). Create one single panel with all box plots. Plot y = "len" by x = "dose" and color by "supp". Specify the option group in stat_compare_means():

```
ggplot(ToothGrowth, aes(dose, len)) +
  geom_boxplot(aes(color = supp))+
  scale_color_manual(values = c("#00AFBB", "#E7B800")) +
  stat_compare_means(aes(group = supp), label = "p.signif")
```



• Paired comparisons for multiple groups:





Read more at: Add P-values and Significance Levels to ggplots³

 $^{^3} http://www.sthda.com/english/articles/24-ggpubr-publication-ready-plots/76-add-p-values-and-significance-levels-to-ggplots/$

3.5 Conclusion

1. Visualize the distribution of a grouped continuous variable: the grouping variable on x-axis and the continuous variable on y axis.

The possible ggplot2 layers include:

- geom boxplot() for box plot
- geom_violin() for violin plot
- geom_dotplot() for dot plot
- geom_jitter() for stripchart
- geom line() for line plot
- geom_bar() for bar plot

Examples of R code: start by creating a plot, named e, and then finish it by adding a layer:

- 2. Create mean and median plots with error bars: the grouping variable on x-axis and the summarized continuous variable (mean/median) on y-axis.
- Compute summary statistics and initialize ggplot with summary data:

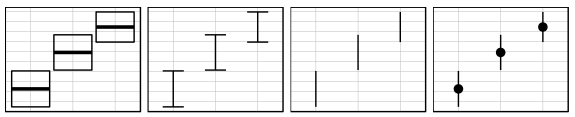
```
# Summary statistics
library(dplyr)
df.summary <- ToothGrowth %>%
  group_by(dose) %>%
  summarise(
    sd = sd(len, na.rm = TRUE),
    len = mean(len)
  )
# Initialize ggplot with data
f <- ggplot(</pre>
```

3.6. SEE ALSO 67

```
df.summary,
aes(x = dose, y = len, ymin = len-sd, ymax = len+sd)
)
```

• Possible error plots:

f + geom_crossbar() f + geom_errorbar() f + geom_linerange() f + geom_pointrange()



3. Combine error bars with violin plots, dot plots, line and bar plots:

```
# Combine with violin plots
ggplot(ToothGrowth, aes(dose, len))+
  geom_violin(trim = FALSE) +
 geom_pointrange(aes(ymin = len-sd, ymax = len + sd),
                  data = df.summary)
# Combine with dot plots
ggplot(ToothGrowth, aes(dose, len))+
 geom_dotplot(stackdir = "center", binaxis = "y",
               fill = "lightgray", dotsize = 1) +
 geom_pointrange(aes(ymin = len-sd, ymax = len + sd),
                  data = df.summary)
# Combine with line plot
ggplot(df.summary, aes(dose, len))+
 geom_line(aes(group = 1)) +
 geom_pointrange(aes(ymin = len-sd, ymax = len + sd))
# Combine with bar plots
ggplot(df.summary, aes(dose, len))+
 geom_bar(stat = "identity", fill = "lightgray") +
 geom_pointrange(aes(ymin = len-sd, ymax = len + sd))
```

3.6 See also

- ggpubr: Publication Ready Plots. https://goo.gl/7uySha
- Facilitating Exploratory Data Visualization: Application to TCGA Genomic Data. https://goo.gl/9LNsRi
- Add P-values and Significance Levels to ggplots. https://goo.gl/VH7Yq7
- Plot Means/Medians and Error Bars. https://goo.gl/zRwAeV

Chapter 4

Plot Two Continuous Variables

4.1 Introduction

Scatter plots are used to display the relationship between two continuous variables x and y. In this article, we'll start by showing how to create beautiful scatter plots in R.

We'll use helper functions in the ggpubr R package¹ to display automatically the **correlation coefficient** and the **significance level** on the plot.

We'll also describe how to color points by groups and to add concentration ellipses around each group. Additionally, we'll show how to create **bubble charts**, as well as, how to add **marginal plots** (histogram, density or box plot) to a scatter plot.

We continue by showing show some alternatives to the standard scatter plots, including rectangular binning, hexagonal binning and 2d density estimation. These plot types are useful in a situation where you have a large data set containing thousands of records.

R codes for zooming in scatter plot are also provided. Finally, you'll learn how to add fitted regression trend lines and equations to a scatter plot.

4.2 Prerequisites

1. **Install cowplot package**. Used to arrange multiple plots. Will be used here to create a scatter plot with marginal density plots. Install the latest developmental version as follow:

```
devtools::install_github("wilkelab/cowplot")
```

2. **Install ggpmisc** for adding the equation of a fitted regression line on a scatter plot:

```
install.packages("ggpmisc")
```

3. Load required packages and set ggplot themes:

¹http://www.sthda.com/english/articles/24-ggpubr-publication-ready-plots/

- Load ggplot2 and ggpubr R packages
- Set the default theme to theme minimal() [in ggplot2]

```
library(ggplot2)
library(ggpubr)
theme_set(
  theme_minimal() +
    theme(legend.position = "top")
)
```

4. Prepare demo data sets:

Dataset: mtcars². The variable cyl is used as grouping variable.

```
# Load data
data("mtcars")
df <- mtcars
# Convert cyl as a grouping variable
df$cyl <- as.factor(df$cyl)</pre>
# Inspect the data
head(df[, c("wt", "mpg", "cyl", "qsec")], 4)
##
                   wt mpg cyl qsec
## Mazda RX4
                2.62 21.0
                             6 16.5
## Mazda RX4 Wag 2.88 21.0
                             6 17.0
## Datsun 710
                2.32 22.8 4 18.6
## Hornet 4 Drive 3.21 21.4 6 19.4
```

4.3 Basic scatter plots

Key functions:

- geom_point(): Create scatter plots. Key arguments: color, size and shape to change point color, size and shape.
- geom_smooth(): Add smoothed conditional means / regression line. Key arguments:
 - color, size and linetype: Change the line color, size and type.
 - fill: Change the fill color of the confidence region.

```
b <- ggplot(df, aes(x = wt, y = mpg))
# Scatter plot with regression line
b + geom_point()+
geom_smooth(method = "lm")</pre>
```

²http://www.sthda.com/english/wiki/r-built-in-data-sets#mtcars-motor-trend-car-road-tests



To remove the confidence region around the regression line, specify the argument se = FALSE in the function geom_smooth().

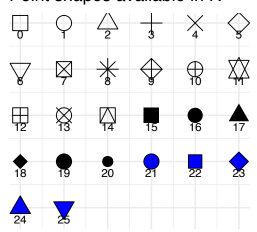
Change the point shape, by specifying the argument shape, for example:

```
b + geom_point(shape = 18)
```

To see the different point shapes commonly used in R, type this:

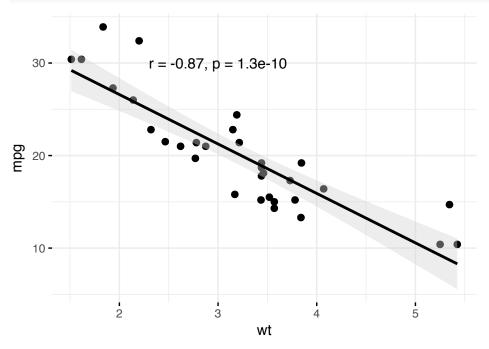
```
ggpubr::show_point_shapes()
```

Point shapes available in R



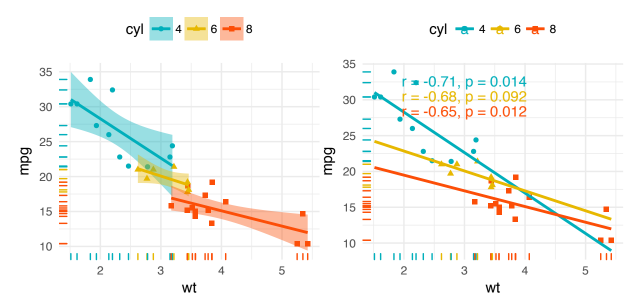
Create easily a scatter plot using ggscatter() [in ggpubr]. Use stat_cor() [ggpubr] to add the correlation coefficient and the significance level.

```
)+
stat_cor(method = "pearson",
    label.x = 3, label.y = 30)
```

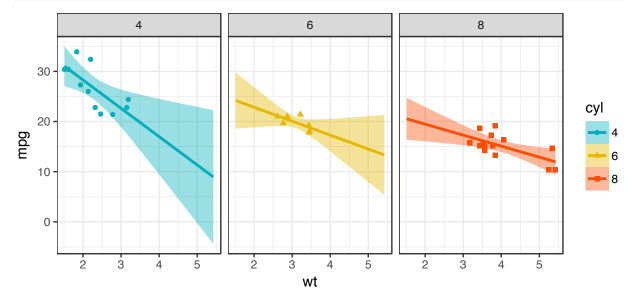


4.4 Multiple groups

- Change point colors and shapes by groups.
- Add marginal rug: geom rug().

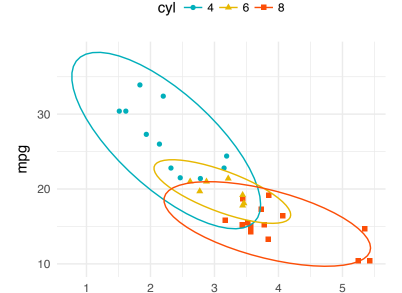


• Split the plot into multiple panels. Use the function facet_wrap():



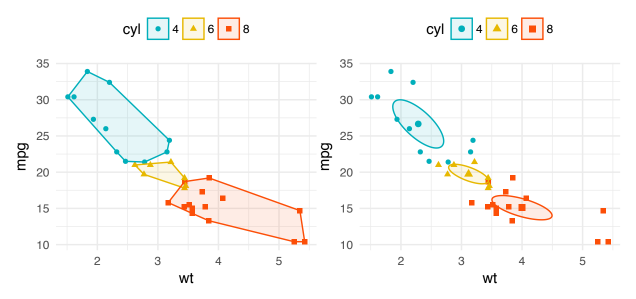
- Add concentration ellipse around groups. R function stat_ellipse(). Key arguments:
 - type: The type of ellipse. The default "t" assumes a multivariate t-distribution, and "norm" assumes a multivariate normal distribution. "euclid" draws a circle with the radius equal to level, representing the euclidean distance from the center.
 - level: The confidence level at which to draw an ellipse (default is 0.95), or, if type="euclid", the radius of the circle to be drawn.

```
b + geom_point(aes(color = cyl, shape = cyl))+
stat_ellipse(aes(color = cyl), type = "t")+
scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
```



wt

Instead of drawing the concentration ellipse, you can: i) plot a convex hull of a set of points; ii) add the mean points and the confidence ellipse of each group. Key R functions: stat_chull(), stat_conf_ellipse() and stat_mean() [in ggpubr]:



• Easy alternative using ggpubr. See this article: Perfect Scatter Plots with Correlation and Marginal Histograms³

```
# Add group mean points and stars
ggscatter(df, x = "wt", y = "mpg",
           color = "cyl", palette = "npg",
           shape = "cyl", ellipse = TRUE,
           mean.point = TRUE, star.plot = TRUE,
           ggtheme = theme_minimal())
# Change the ellipse type to 'convex'
ggscatter(df, x = "wt", y = "mpg",
           color = "cyl", palette = "npg",
           shape = "cyl",
           ellipse = TRUE, ellipse.type = "convex",
           ggtheme = theme_minimal())
                                         35 -
  40 -
                                         30
  30 -
                                 cyl
                                                                       cyl
                                         25 -
mpg
                                         20 -
  20 -
                                         15 -
   10 -
                                         10 -
                                                                5
                                                     3
                                               2
```

 $^{^3}$ http://www.sthda.com/english/articles/24-ggpubr-publication-ready-plots/79-plot-meansmedians-and-error-bars/

4.5 Add point text labels

Key functions:

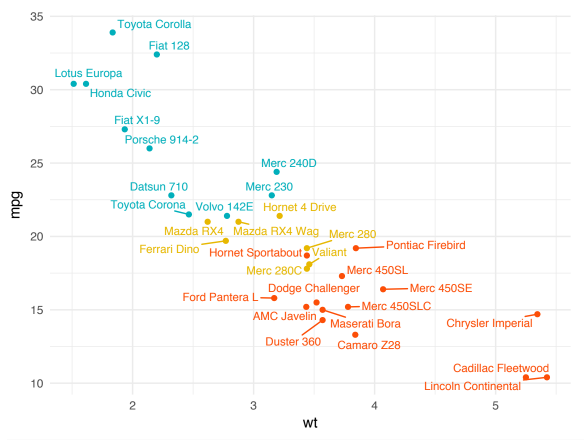
- geom_text() and geom_label(): ggplot2 standard functions to add text to a plot.
- geom_text_repel() and geom_label_repel() [in ggrepel package]. Repulsive textual annotations. Avoid text overlapping.

First install ggrepel (install.packages("ggrepel")), then type this:

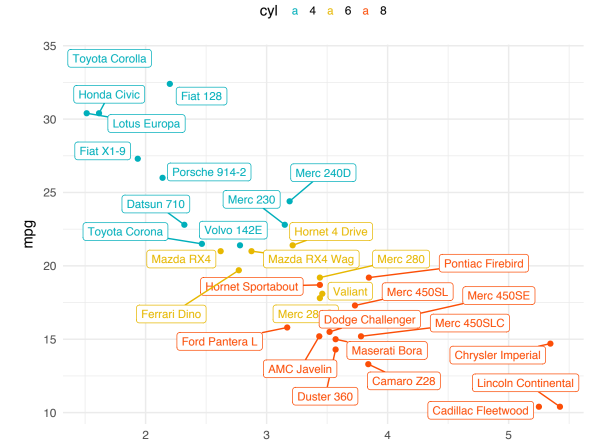
```
library(ggrepel)

# Add text to the plot
.labs <- rownames(df)
b + geom_point(aes(color = cyl)) +
  geom_text_repel(aes(label = .labs, color = cyl), size = 3)+
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))</pre>
```





```
# Draw a rectangle underneath the text, making it easier to read.
b + geom_point(aes(color = cyl)) +
geom_label_repel(aes(label = .labs, color = cyl), size = 3)+
scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))
```

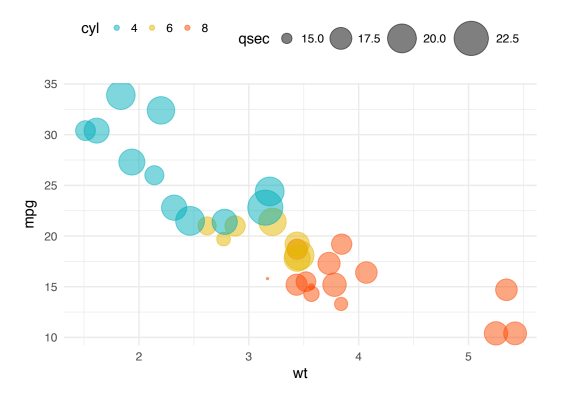


4.6 Bubble chart

In a bubble chart, points size is controlled by a continuous variable, here qsec. In the R code below, the argument alpha is used to control color transparency. alpha should be between 0 and 1.

wt

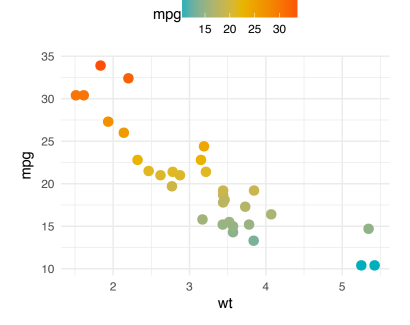
```
b + geom_point(aes(color = cyl, size = qsec), alpha = 0.5) +
scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07")) +
scale_size(range = c(0.5, 12)) # Adjust the range of points size
```



4.7 Color by a continuous variable

- Color points according to the values of the continuous variable: "mpg".
- Change the default blue gradient color using the function scale_color_gradientn() [in ggplot2], by specifying two or more colors.

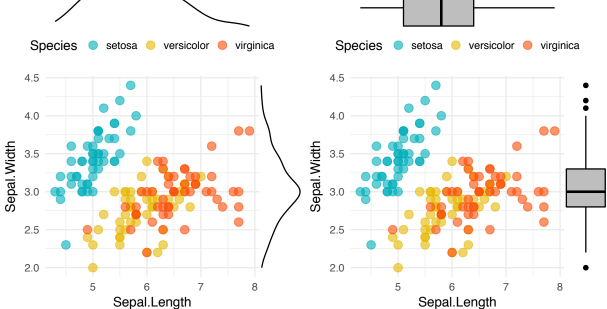
```
b + geom_point(aes(color = mpg), size = 3) +
scale_color_gradientn(colors = c("#00AFBB", "#E7B800", "#FC4E07"))
```



4.8 Add marginal density plots

The function ggMarginal() [in ggExtra package] (Attali, 2017), can be used to easily add a marginal histogram, density or box plot to a scatter plot.

First, install the ggExtra package as follow: install.packages("ggExtra"); then type the following R code:

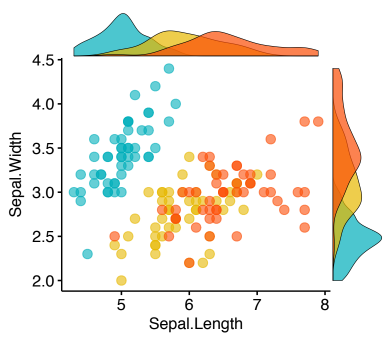


One limitation of ggExtra is that it can't cope with multiple groups in the scatter plot and the marginal plots.

A solution is provided in the function ggscatterhist() [ggpubr]:

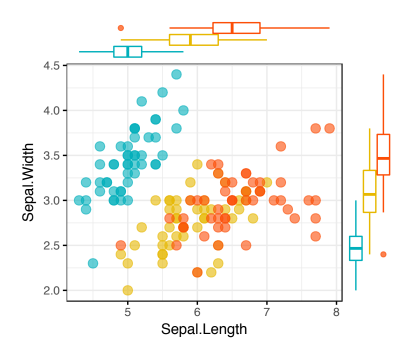
```
library(ggpubr)
# Grouped Scatter plot with marginal density plots
ggscatterhist(
  iris, x = "Sepal.Length", y = "Sepal.Width",
  color = "Species", size = 3, alpha = 0.6,
  palette = c("#00AFBB", "#E7B800", "#FC4E07"),
  margin.params = list(fill = "Species", color = "black", size = 0.2)
  )
```





```
# Use box plot as marginal plots
ggscatterhist(
  iris, x = "Sepal.Length", y = "Sepal.Width",
  color = "Species", size = 3, alpha = 0.6,
  palette = c("#00AFBB", "#E7B800", "#FC4E07"),
  margin.plot = "boxplot",
  ggtheme = theme_bw()
)
```





4.9 Continuous bivariate distribution

In this section, we'll present some alternatives to the standard scatter plots. These include:

- Rectangular binning. Rectangular heatmap of 2d bin counts
- Hexagonal binning: Hexagonal heatmap of 2d bin counts.
- 2d density estimation

1. Rectangular binning:

Rectangular binning is a very useful alternative to the standard scatter plot in a situation where you have a large data set containing thousands of records.

Rectangular binning helps to handle overplotting. Rather than plotting each point, which would appear highly dense, it divides the plane into rectangles, counts the number of cases in each rectangle, and then plots a heatmap of 2d bin counts. In this plot, many small hexagon are drawn with a color intensity corresponding to the number of cases in that bin.

Key function: geom_bin2d(): Creates a heatmap of 2d bin counts. Key arguments: bins, numeric vector giving number of bins in both vertical and horizontal directions. Set to 30 by default.

2. **Hexagonal binning**: Similar to rectangular binning, but divides the plane into regular hexagons. Hexagon bins avoid the visual artefacts sometimes generated by the very regular alignment of 'geom_bin2d().

Key function: geom_hex()

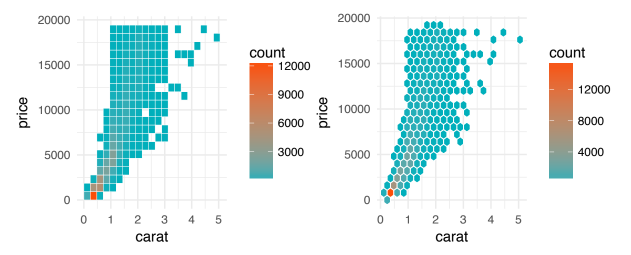
3. Contours of a 2d density estimate. Perform a 2D kernel density estimation and display results as contours overlaid on the scatter plot. This can be also useful for dealing with overplotting.

Key function: geom density 2d()

• Create a scatter plot with rectangular and hexagonal binning:

```
# Rectangular binning
ggplot(diamonds, aes(carat, price)) +
    geom_bin2d(bins = 20, color ="white")+
    scale_fill_gradient(low = "#00AFBB", high = "#FC4E07")+
    theme_minimal()

# Hexagonal binning
ggplot(diamonds, aes(carat, price)) +
    geom_hex(bins = 20, color = "white")+
    scale_fill_gradient(low = "#00AFBB", high = "#FC4E07")+
    theme_minimal()
```



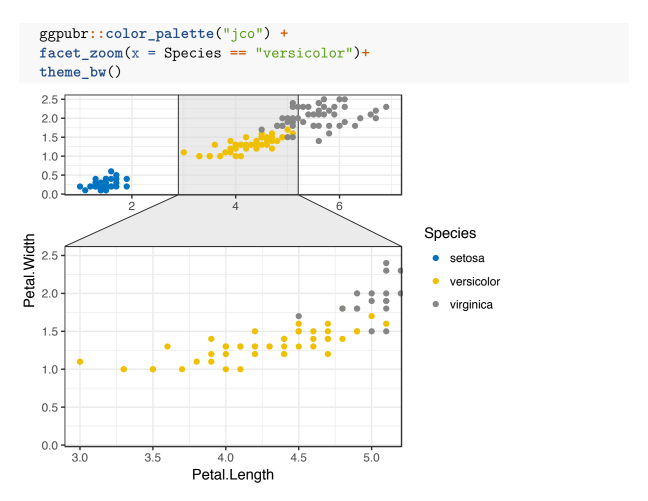
• Create a scatter plot with 2d density estimation:

```
# Add 2d density estimation
sp <- ggplot(iris, aes(Sepal.Length, Sepal.Width)) +</pre>
  geom_point(color = "lightgray")
sp + geom_density_2d()
# Use different geometry and change the gradient color
sp + stat_density_2d(aes(fill = ..level..), geom = "polygon") +
  scale_fill_gradientn(colors = c("#FFEDAO", "#FEB24C", "#F03B20"))
   4.5
                                                         level
                                                                  0.2 0.3 0.4
                                                              0.1
   4.0
                                              4.5
Sepal.Width
   3.5
                                              4.0
                                           Sepal.Width
                                              3.5
                                              3.0
   2.5
                                              2.5
   2.0
                                              2.0
              5
                                         8
                                                         5
                                                                                    8
                  Sepal.Length
                                                             Sepal.Length
```

4.10 Zoom in a scatter plot

- Key function: facet zomm() [in ggforce] (Pedersen, 2016).
- Demo data set: iris. The R code below zoom the points where Species == "versicolor".

```
library(ggforce)
ggplot(iris, aes(Petal.Length, Petal.Width, colour = Species)) +
   geom_point() +
```



To zoom the points, where Petal.Length < 2.5, type this:

```
ggplot(iris, aes(Petal.Length, Petal.Width, colour = Species)) +
  geom_point() +
  ggpubr::color_palette("jco") +
  facet_zoom(x = Petal.Length < 2.5)+
  theme_bw()</pre>
```

4.11 Add trend lines and equations

In this section, we'll describe how to add trend lines to a scatter plot and labels (equation, R2, BIC, AIC) for a fitted lineal model.

1. Load packages and create a basic scatter plot facetted by groups:

```
# Load packages and set theme
library(ggpubr)
library(ggpmisc)

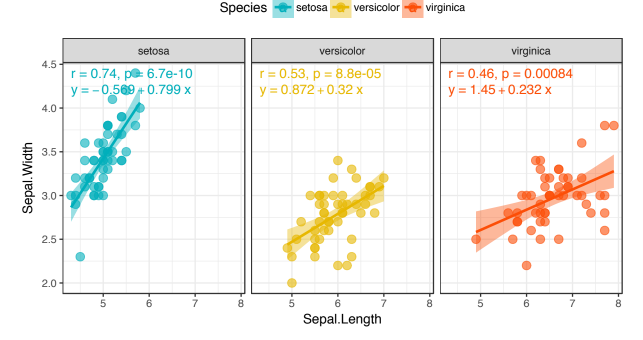
theme_set(
   theme_bw() +
      theme(legend.position = "top")
)
```

```
# Scatter plot
p <- ggplot(iris, aes(Sepal.Length, Sepal.Width)) +
  geom_point(aes(color = Species), size = 3, alpha = 0.6) +
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07")) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))+
  facet_wrap(~Species)</pre>
```

- 2. Add regression line, correlation coefficient and equantions of the fitted line. Key functions:
 - stat_smooth() [ggplot2]
 - stat_cor() [ggpubr]
 - stat poly eq()[ggpmisc]

```
formula <- y ~ x

p +
    stat_smooth( aes(color = Species, fill = Species), method = "lm") +
    stat_cor(aes(color = Species), label.y = 4.4)+
    stat_poly_eq(
        aes(color = Species, label = ..eq.label..),
        formula = formula, label.y = 4.2, parse = TRUE)</pre>
```

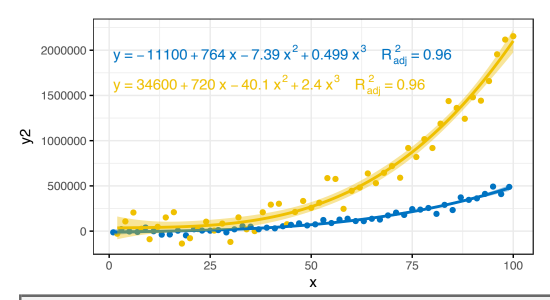


- 3. Fit polynomial equation:
- Create some data:

• Fit polynomial regression line and add labels:

```
# Polynomial regression. Sow equation and adjusted R2
formula <- y ~ poly(x, 3, raw = TRUE)
p <- ggplot(my.data, aes(x, y2, color = group)) +
    geom_point() +
    geom_smooth(aes(fill = group), method = "lm", formula = formula) +
    stat_poly_eq(
    aes(label = paste(..eq.label.., ..adj.rr.label.., sep = "~~~~")),
    formula = formula, parse = TRUE
    )
ggpar(p, palette = "jco")</pre>
```





Note that, you can also display the AIC and the BIC values using ..AIC.label.. and ..BIC.label.. in the above equation.

Other arguments (label.x, label.y) are available in the function stat_poly_eq() to adjust label positions.

For more examples, type this R code: browseVignettes("ggpmisc").

4.12 Conclusion

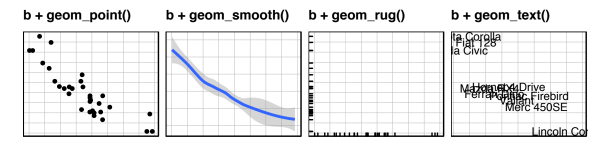
1. Create a basic scatter plot:

```
b <- ggplot(mtcars, aes(x = wt, y = mpg))</pre>
```

Possible layers, include:

- geom_point() for scatter plot
- geom_smooth() for adding smoothed line such as regression line
- geom rug() for adding a marginal rug
- geom text() for adding textual annotations

4.13. SEE ALSO 85

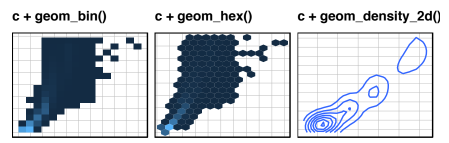


2. Continuous bivariate distribution:

c <- ggplot(diamonds, aes(carat, price))</pre>

Possible layers include:

- geom_bin2d(): Rectangular binning.
- geom_hex(): Hexagonal binning.
- geom_density_2d(): Contours from a 2d density estimate



4.13 See also

- ggpubr: Publication Ready Plots. https://goo.gl/7uySha
- Perfect Scatter Plots with Correlation and Marginal Histograms. https://goo.gl/3o4ddg

Chapter 5

Plot Multivariate Continuous Data

5.1 Introduction

Here, multivariate data are data that contains 3 or more variables.

When you have only three continuous variables in your data set, you can create a **3d** scatter plot.

For a small data set with more than three variables, it's possible to visualize the relationship between each pairs of variables by creating a **scatter plot matrix**. You can also perform a correlation analysis between each pairs of variables.

For a large multivariate data set, it is more difficult to visualize their relationships. Discovering knowledge from these data requires specific techniques. **Multivariate analysis** (MVA) refers to a set of techniques used for analyzing a data set containing multiple variables.

Among these techniques, there are:

- Cluster analysis¹ for identifying groups of observations with similar profile according to a specific criteria.
- Principal component methods², which consist of summarizing and visualizing the most important information contained in a multivariate data set.

In this chapter we provide an overview of methods for visualizing multivariate data sets.

5.2 Demo data set and R package

```
library("magrittr") # for piping %>%
head(iris, 3)
```

Sepal.Length Sepal.Width Petal.Length Petal.Width Species

http://www.sthda.com/english/articles/25-cluster-analysis-in-r-practical-guide/
http://www.sthda.com/english/articles/31-principal-component-methods-in-r-

practical-guide/

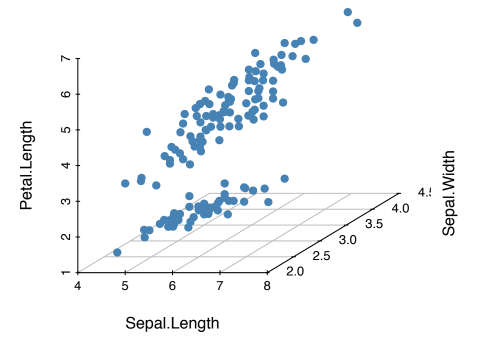
##	1	5.1	3.5	1.4	0.2	setosa
##	2	4.9	3.0	1.4	0.2	setosa
##	3	4.7	3.2	1.3	0.2	setosa

5.3 Create a 3d scatter plot

You can create a 3d scatter plot using the R package **scatterplot3d** (Ligges et al., 2017), which contains a function of the same name.

- Install: install.packages("scatterplot3d")
- Create a basic 3d scatter plot:

```
library(scatterplot3d)
scatterplot3d(
  iris[,1:3], pch = 19, color = "steelblue",
    grid = TRUE, box = FALSE,
    mar = c(3, 3, 0.5, 3)
)
```



• See more examples at: http://www.sthda.com/english/wiki/3d-graphics

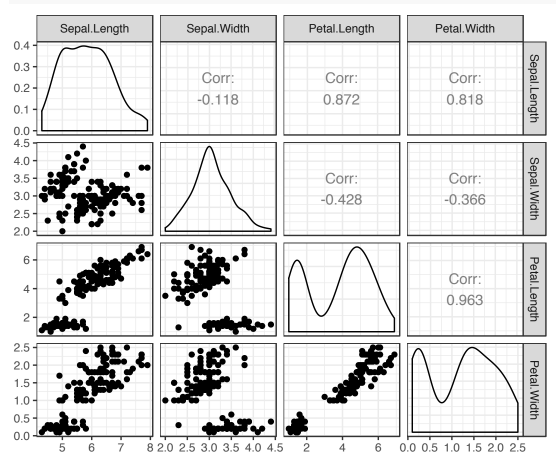
5.4 Create a scatter plot matrix

To create a scatter plot of each possible pairs of variables, you can use the function **gg-pairs**() [in **GGally** package, an extension of ggplot2](Schloerke et al., 2016) . It produces a pairwise comparison of multivariate data.

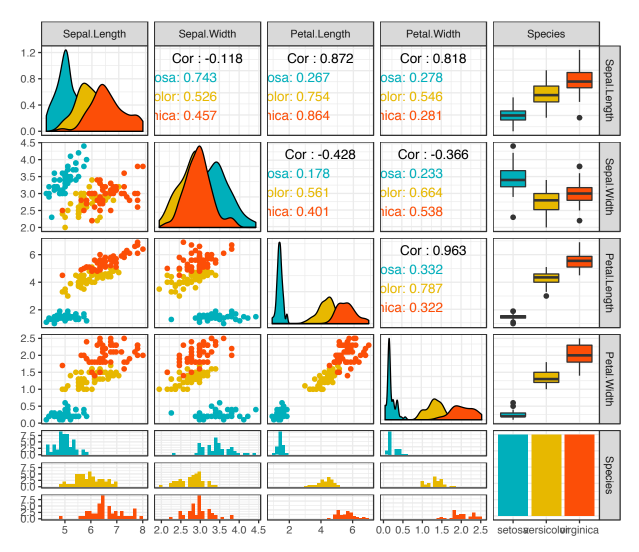
• Install: install.packages("GGally")

- Create a simple scatter plot matrix. The plot contains the:
 - Scatter plot and the correlation coefficient between each pair of variables
 - Density distribution of each variable

```
library(GGally)
library(ggplot2)
ggpairs(iris[,-5])+ theme_bw()
```



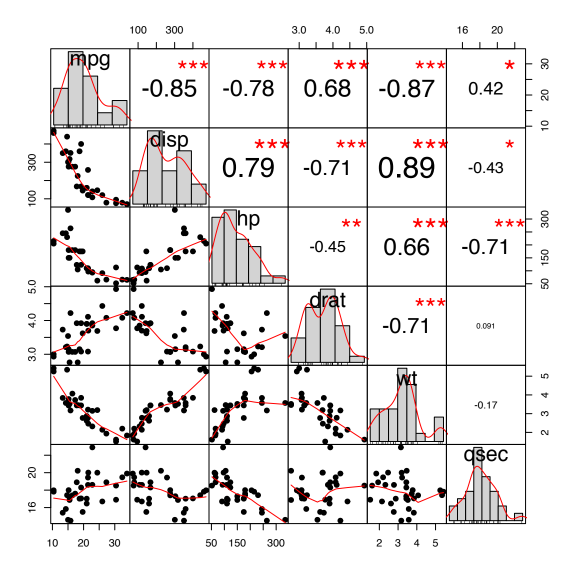
- Create a scatter plot matrix by groups. The plot contains the :
 - Scatter plot and the correlation coefficient, between each pair of variables, colored by groups
 - Density distribution and the box plot, of each continuous variable, colored by groups



An alternative to the function ggpairs() is provided by the R base plot function chart.Correlation() [in PerformanceAnalytics packages]. It displays the correlation coefficient and the significance levels as stars.

For example, type the following R code, after installing the ${\tt PerformanceAnalytics}$ package:

```
# install.packages("PerformanceAnalytics")
library("PerformanceAnalytics")
my_data <- mtcars[, c(1,3,4,5,6,7)]
chart.Correlation(my_data, histogram=TRUE, pch=19)</pre>
```



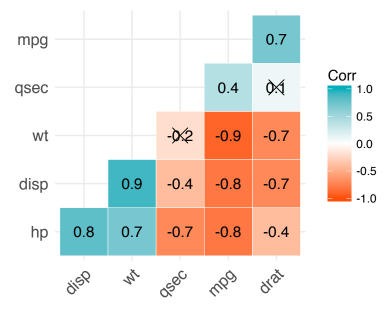
5.5 Correlation analysis

Recall that, correlation analysis is used to investigate the association between two or more variables. Read more at: Correlation analyses in \mathbb{R}^3 .

- 1. Compute correlation matrix between pairs of variables using the R base function cor()
- 2. Visualize the output. Two possibilities:
 - Use the function ggcorrplot() [in ggcorplot package]. Extension to the ggplot2 system. See more examples at: http://www.sthda.com/english/wiki/ggcorrplot-visualization-of-a-correlation-matrix-using-ggplot2.
 - Use the function corrplot() [in corrplot package]. R base plotting system. See examples at: http://www.sthda.com/english/wiki/visualize-correlation-matrix-using-correlogram.

Here, we'll present only the ggcorrplot package (Kassambara, 2016), which can be installed as follow: install.packages("ggcorrplot").

³http://www.sthda.com/english/wiki/correlation-analyses-in-r



In the plot above:

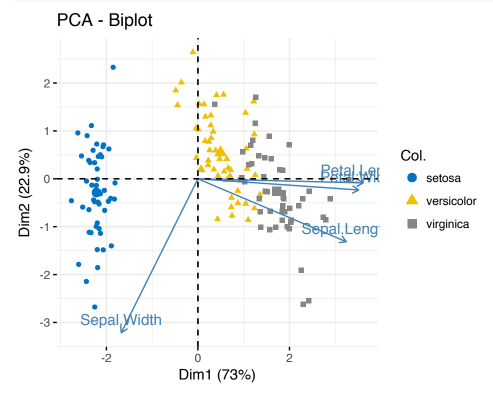
- Positive correlations are shown in blue and negative correlation in red
- Variables that are associated are grouped together.
- Non-significant correlation are marked by a cross (X)

5.6 Principal component analysis

Principal component analysis (PCA) is a multivariate data analysis approach that allows us to summarize and visualize the most important information contained in a multivariate data set.

PCA reduces the data into few new dimensions (or axes), which are a linear combination of the original variables. You can visualize a multivariate data by drawing a scatter plot of the first two dimensions, which contain the most important information in the data. Read more at: https://goo.gl/kabVHq

- Demo data set: iris
- Compute PCA using the R base function prcomp()
- Visualize the output using the factoextra R package (an extension to ggplot2) (Kassambara and Mundt, 2017)



In the plot above:

- Dimension (Dim.) 1 and 2 retained about 96% (73% + 22.9%) of the total information contained in the data set.
- Individuals with a similar profile are grouped together
- Variables that are positively correlated are on the same side of the plots. Variables that are negatively correlated are on the opposite side of the plots.

5.7 Cluster analysis

Cluster analysis is one of the important data mining methods for discovering knowledge in multidimensional data. The goal of clustering is to identify pattern or groups of similar objects within a data set of interest. Read more at: http://www.sthda.com/english/articles/25-cluster-analysis-in-r-practical-guide/.

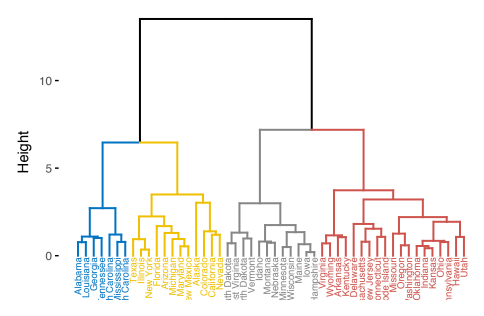
This section describes how to compute and visualize hierarchical clustering, which output is a tree called dendrogram showing groups of similar individuals.

- Computation. R function: hclust(). It takes a dissimilarity matrix as an input, which is calculated using the function dist().
- Visualization: fviz_dend() [in factoextra]
- Demo data sets: USArrests

5.8. CONCLUSION 93

Before cluster analysis, it's recommended to scale (or normalize) the data, to make the variables comparable. R function: scale(), applies scaling on the column of the data (variables).

Cluster Dendrogram



A heatmap is another way to visualize hierarchical clustering. It's also called a false colored image, where data values are transformed to color scale. Heat maps allow us to simultaneously visualize groups of samples and features. You can easily create a pretty heatmap using the R package pheatmap.

In heatmap, generally, columns are samples and rows are variables. Therefore we start by scaling and then transpose the data before creating the heatmap.

5.8 Conclusion

For a multivariate continuous data, you can perform the following analysis or visualization depending on the complexity of your data:

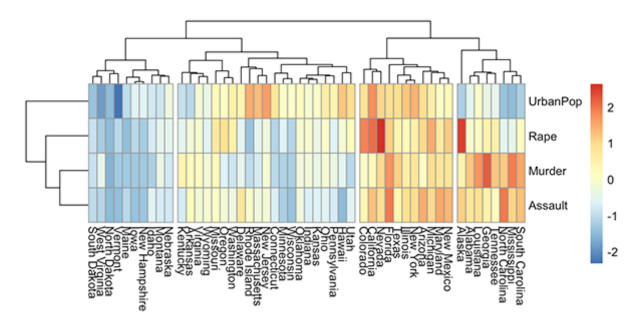


Figure 5.1: Multivariate data Heatmap

- 3D scatter plot : scatterplot3d() [scatterplot3d]
- Create a scatter plot matrix: ggpairs [GGally]
- Correlation matrix analysis and visualization: cor()[stats] and ggcorrplot() [ggcorrplot] for the visualization.
- Principal component analysis: prcomp() [stats] and fviz_pca() [factoextra]
- Cluster analysis: hclust() [stats] and fviz_dend() [factoextra]

Chapter 6

Visualizing Multivariate Categorical Data

6.1 Introduction

To visualize a small data set containing multiple categorical variables, you can create either a bar plot, a balloon plot or a mosaic plot.

For a large multivariate categorical data, you need specialized techniques dedicated to categorical data analysis, such as simple and multiple correspondence analysis¹. These methods make it possible to visualize the association between a large number of categorical variables.

Here, we'll describe simple examples of graphs for visualizing the frequency distribution of categorical variables contained in small contingency tables.

6.2 Prerequisites

Load required R packages and set the default theme:

```
library(ggplot2)
library(ggpubr)
theme_set(theme_pubr())
```

6.3 Bar plots of contingency tables

Demo data set: HairEyeColor (distribution of hair and eye color and sex in 592 statistics students)

• Prepare and inspect the data:

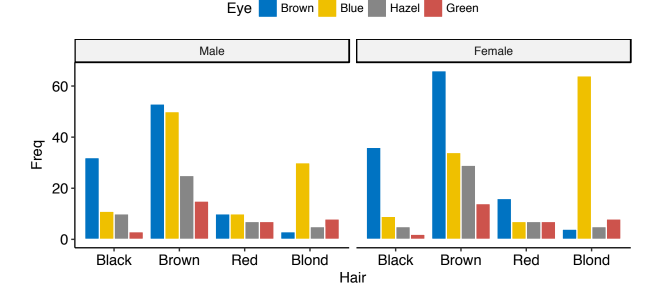
 $^{^{1}}$ http://www.sthda.com/english/articles/31-principal-component-methods-in-r-practical-guide/

```
data("HairEyeColor")
df <- as.data.frame(HairEyeColor)
head(df)</pre>
```

```
##
      Hair
             Eye Sex Freq
## 1 Black Brown Male
## 2 Brown Brown Male
                         53
## 3
       Red Brown Male
                         10
## 4 Blond Brown Male
                          3
## 5 Black Blue Male
                         11
## 6 Brown
            Blue Male
                         50
```

- Create the bar graph:
 - Hair color on x-axis
 - Change bar fill by Eye color
 - Split the graph into multiple panel by Sex

```
ggplot(df, aes(x = Hair, y = Freq))+
  geom_bar(
    aes(fill = Eye), stat = "identity", color = "white",
    position = position_dodge(0.9)
    )+
  facet_wrap(~Sex) +
  fill_palette("jco")
```



6.4 Balloon plot

Balloon plot is an alternative to bar plot for visualizing a large categorical data. We'll use the function ggballoonplot() [in ggpubr], which draws a graphical matrix of a contingency table, where each cell contains a dot whose size reflects the relative magnitude of the corresponding component.

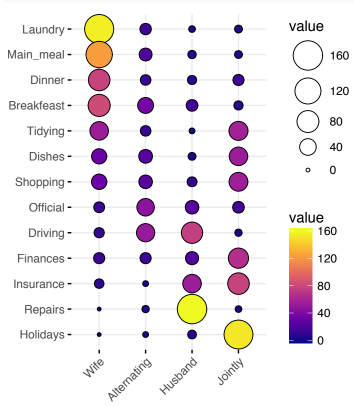
Demo data sets: Housetasks (a contingency table containing the frequency of execution of 13 house tasks in the couple.)

```
housetasks <- read.delim(
   system.file("demo-data/housetasks.txt", package = "ggpubr"),
   row.names = 1
   )
head(housetasks, 4)</pre>
```

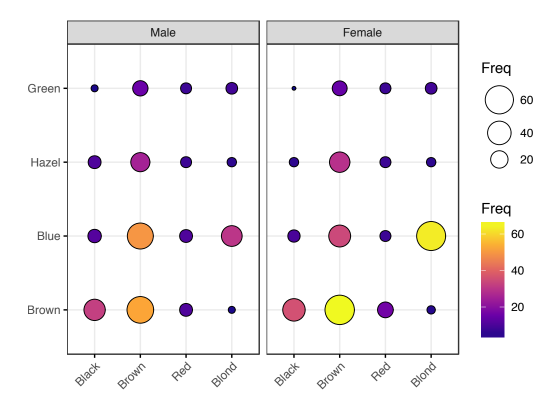
```
Wife Alternating Husband Jointly
##
## Laundry
                156
                               14
                                         2
## Main_meal
                               20
                                         5
                                                  4
                124
## Dinner
                 77
                                         7
                                                 13
                               11
## Breakfeast
                 82
                               36
                                        15
                                                 7
```

• Create a simple balloon plot of a contingency table. Change the fill color by the values in the cells.

```
ggballoonplot(housetasks, fill = "value")+
scale_fill_viridis_c(option = "C")
```



• Visualize a grouped frequency table. Demo data set: HairEyeColor. Create a multi-panel plot by Sex



6.5 Mosaic plot

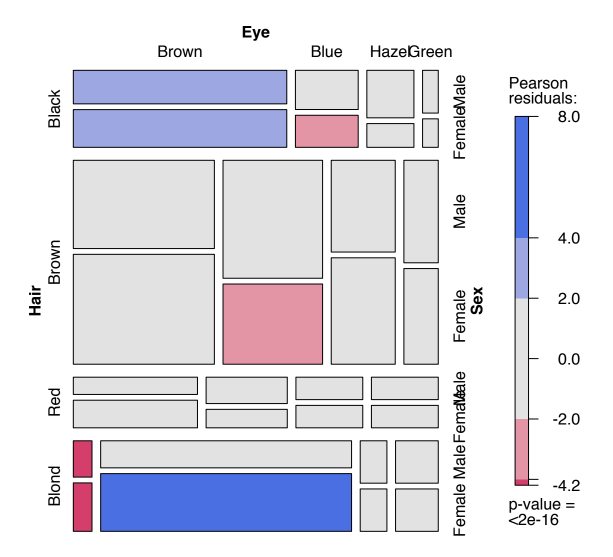
A mosaic plot is basically an area-proportional visualization of observed frequencies, composed of tiles (corresponding to the cells) created by recursive vertical and horizontal splits of a rectangle. The area of each tile is proportional to the corresponding cell entry, given the dimensions of previous splits.

Mosaic graph can be created using either the function mosaicplot() [in graphics] or the function mosaic() [in vcd package]. Read more at: Visualizing Multi-way Contingency Tables with vcd².

Example of mosaic plot:

```
library(vcd)
mosaic(HairEyeColor, shade = TRUE, legend = TRUE)
```

²https://cran.r-project.org/web/packages/vcd/vignettes/strucplot.pdf

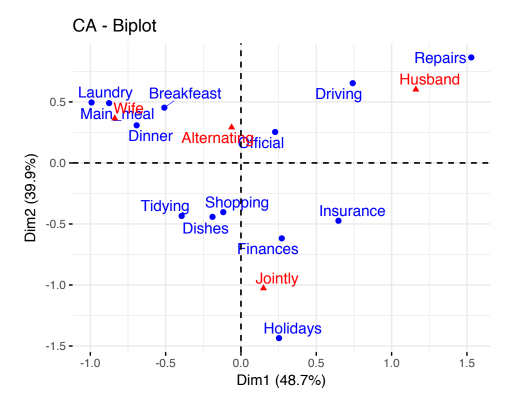


6.6 Correspondence analysis

Correspondence analysis can be used to summarize and visualize the information contained in a large contingency table formed by two categorical variables.

Required package: FactoMineR for the analysis and factoextra for the visualization

```
library(FactoMineR)
library(factoextra)
res.ca <- CA(housetasks, graph = FALSE)
fviz_ca_biplot(res.ca, repel = TRUE)</pre>
```



From the graphic above, it's clear that:

- Housetasks such as dinner, breakfeast, laundry are done more often by the wife
- Driving and repairs are done more frequently by the husband

Read more at: Correspondence analysis in \mathbb{R}^3

³https://goo.gl/7CnpXq

Chapter 7

Plot Time Series Data

7.1 Introduction

In this chapter, we start by describing how to plot simple and multiple time series data using the function <code>geom_line()</code>. Next, we show how to set date axis limits and add trend smoothed line to a time series graphs. Finally, we introduce some extensions to the <code>ggplot2</code> package for easily handling time series objects.

7.2 Basic ggplot of time series

- Plot types: line plot with dates on x-axis
- Demo data set: economics [ggplot2] time series data sets are used.

In this section we'll plot the variables psavert (personal savings rate) and uempmed (number of unemployed in thousands) by date (x-axis).

• Load required packages and set the default theme:

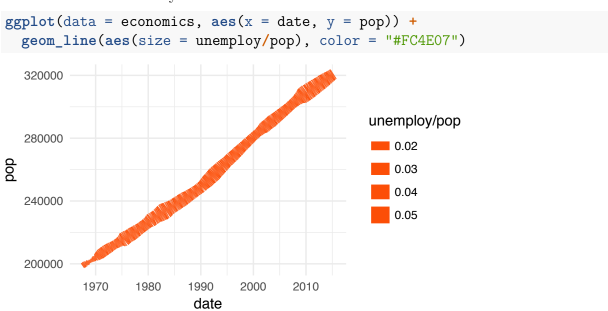
```
library(ggplot2)
theme_set(theme_minimal())
# Demo dataset
head(economics)
```

```
## # A tibble: 6 x 6
##
           date
                          pop psavert uempmed unemploy
                   рсе
##
         <date> <dbl>
                        <int>
                                 <dbl>
                                          <dbl>
                                                   <int>
## 1 1967-07-01
                   507 198712
                                  12.5
                                            4.5
                                                    2944
## 2 1967-08-01
                   510 198911
                                  12.5
                                            4.7
                                                    2945
## 3 1967-09-01
                   516 199113
                                  11.7
                                                    2958
                                            4.6
## 4 1967-10-01
                   513 199311
                                  12.5
                                            4.9
                                                    3143
## 5 1967-11-01
                   518 199498
                                  12.5
                                            4.7
                                                    3066
## 6 1967-12-01
                   526 199657
                                  12.1
                                            4.8
                                                    3018
```

• Create basic line plots

```
# Basic line plot
ggplot(data = economics, aes(x = date, y = pop))+
  geom_line(color = "#00AFBB", size = 2)
# Plot a subset of the data
ss <- subset(economics, date > as.Date("2006-1-1"))
ggplot(data = ss, aes(x = date, y = pop)) +
  geom_line(color = "#FC4E07", size = 2)
  320000
                                          320000
                                          315000
  280000
                                          310000
  240000
                                          305000
                                          300000
  200000
               1980
                     1990
                           2000
                                2010
                                                     2008
                                                            2010
                                                                 2012
                                                                        2014
                                                2006
                      date
                                                             date
```

• Control line size by the value of a continuous variable:



7.3 Plot multiple time series data

Here, we'll plot the variables psavert and uempmed by dates. You should first reshape the data using the tidyr package: - Collapse psavert and uempmed values in the same column (new column). R function: gather()[tidyr] - Create a grouping variable that with levels = psavert and uempmed

```
library(tidyr)
library(dplyr)
```

```
df <- economics %>%
  select(date, psavert, uempmed) %>%
  gather(key = "variable", value = "value", -date)
head(df, 3)
## # A tibble: 3 x 3
           date variable value
##
         <date> <chr> <dbl>
## 1 1967-07-01 psavert 12.5
## 2 1967-08-01 psavert
                           12.5
## 3 1967-09-01 psavert
                           11.7
# Multiple line plot
ggplot(df, aes(x = date, y = value)) +
  geom_line(aes(color = variable), size = 1) +
  scale_color_manual(values = c("#00AFBB", "#E7B800")) +
  theme_minimal()
  25
  20
                                                              variable
value
  15
                                                                psavert
                                                                uempmed
   5
        1970
                  1980
                             1990
                                       2000
                                                 2010
                              date
# Area plot
ggplot(df, aes(x = date, y = value)) +
  geom_area(aes(color = variable, fill = variable),
            alpha = 0.5, position = position_dodge(0.8)) +
  scale_color_manual(values = c("#00AFBB", "#E7B800")) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800"))
  25
  20
                                                              variable
value
  15
                                                                 psavert
  10
                                                                 uempmed
   5
   0
                             1990
                                                 2010
        1970
                  1980
                                       2000
                              date
```

2005

2010

date

2015

7.4 Set date axis limits

Key R function: scale_x_date()

7.5 Format date axis labels

1990

date

2000

2010

Key function: scale x date().

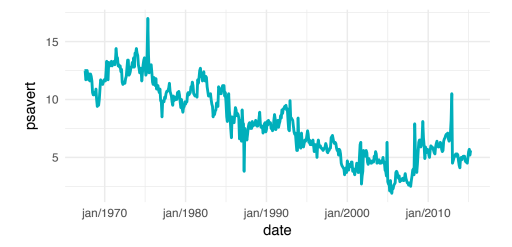
1980

1970

To format date axis labels, you can use different combinations of days, weeks, months and years:

- Weekday name: use %a and %A for abbreviated and full weekday name, respectively
- Month name: use %b and %B for abbreviated and full month name, respectively
- %d: day of the month as decimal number
- %Y: Year with century.
- See more options in the documentation of the function ?strptime

```
# Format : month/year
p + scale_x_date(date_labels = "%b/%Y")
```



7.6 Add trend smoothed line

Key function: stat_smooth()

```
p + stat_smooth(
  color = "#FC4E07", fill = "#FC4E07",
  method = "loess"
)
15
15
1970 1980 1990 2000 2010
date
```

7.7 ggplot2 extensions for ts objects

The ggfortify package is an extension to ggplot2 that makes it easy to plot time series objects (Horikoshi and Tang, 2017). It can handle the output of many time series packages, including: zoo::zooreg(), xts::xts(), timeSeries::timSeries(), tseries::irts(), forecast::forecast(), vars:vars().

Another interesting package is the ggpmisc package (Aphalo, 2017), which provides two useful methods for time series object:

- stat peaks() finds at which x positions local y maxima are located, and
- stat valleys() finds at which x positions local y minima are located.

Here, we'll show how to easily:

- Visualize a time series object, using the data set AirPassengers (monthly airline passenger numbers 1949-1960).
- Identify shifts in mean and/or variance in a time series using the **changepoint** package.
- Detect jumps in a data using the strucchange package and the data set Nile (Measurements of the annual flow of the river Nile at Aswan).
- Detect peaks and valleys using the ggpmisc package and the data set lynx (Annual Canadian Lynx trappings 1821–1934).

First, install required R packages:

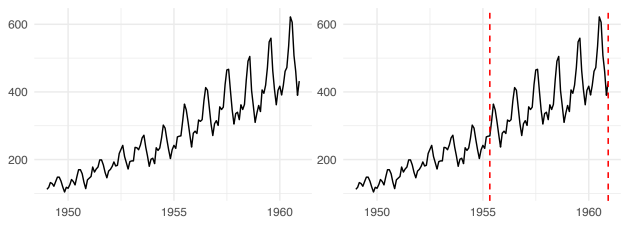
Then use the autoplot.ts() function to visualize time series objects, as follow:

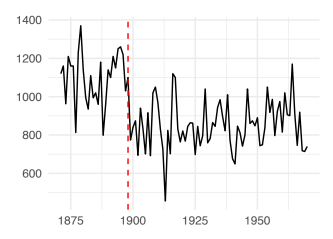
```
library(ggfortify)
library(magrittr) # for piping %>%

# Plot ts objects
autoplot(AirPassengers)

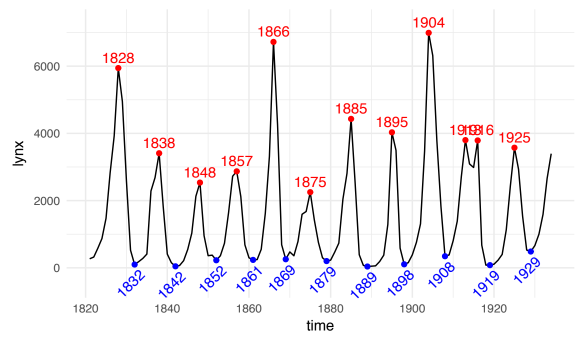
# Identify change points in mean and variance
AirPassengers %>%
    changepoint:: cpt.meanvar() %>% # Identify change points
autoplot()

# Detect jump in a data
strucchange::breakpoints(Nile ~ 1) %>%
autoplot()
```





Detect peaks and valleys:



Chapter 8

Facets: Multi-Panels GGPlot

8.1 Introduction

This chapter describes how to create a multi-panel ggplots or facets. Facets divide a ggplot into subplots based on the values of one or more categorical variables. There are two main functions for faceting: facet_grid() and facet_wrap()

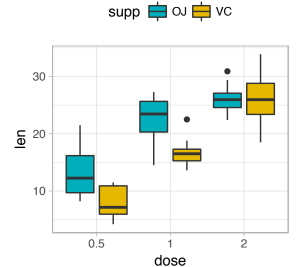
8.2 Prerequisites

Load required packages and set the theme function theme_light() [ggplot2] as the default theme:

```
library(ggplot2)
theme_set(
   theme_light() + theme(legend.position = "top")
)
```

Create a box plot filled by groups:

```
# Load data and convert dose to a factor variable
data("ToothGrowth")
ToothGrowth$dose <- as.factor(ToothGrowth$dose)
# Box plot
p <- ggplot(ToothGrowth, aes(x = dose, y = len)) +
  geom_boxplot(aes(fill = supp), position = position_dodge(0.9)) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800"))
p</pre>
```



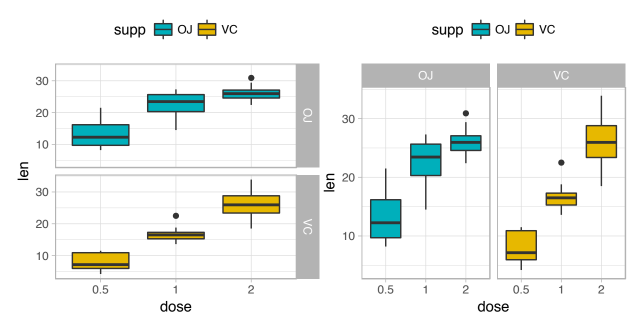
8.3 Split the plot into a matrix of panels

The following functions can be used for facets:

- p + **facet_grid**(supp \sim .): Facet in vertical direction based on the levels of supp variable.
- p + facet_grid(. \sim supp): Facet in horizontal direction based on the levels of supp variable.
- p + facet_grid(dose \sim supp): Facet in horizontal and vertical directions based on two variables: dose and supp.
- $p + facet_wrap(\sim fl)$: Place facet side by side into a rectangular layout
- 1. Facet with one discrete variable: Split by the levels of the group "supp"

```
# Split in vertical direction
p + facet_grid(supp ~ .)

# Split in horizontal direction
p + facet_grid(. ~ supp)
```



2. Facet with two discrete variables: Split by the levels of the groups "dose" and "supp"

```
# Facet by two variables: dose and supp.
# Rows are dose and columns are supp
p + facet_grid(dose ~ supp)
# Facet by two variables: reverse the order of the 2 variables
# Rows are supp and columns are dose
p + facet_grid(supp ~ dose)
                                                        supp 📛 OJ 📛 VC
             supp 🖨 OJ 🖨 VC
  30
                                             30
  20
                                             20
  10
                                             10
  30
                                          <u>e</u>n
<u>e</u>n
  20
  10
                                             30
  30
                                             20
  20
                                             10
  10
                  2
                        0.5
                                   2
                                                            0.5
                                                                1
                                                                   2
                                                                       0.5
       0.5
             1
                                                0.5
                                                    1
                                                        2
                    dose
                                                              dose
```

Note that, you can use the argument margins to add additional facets which contain all the data for each of the possible values of the faceting variables

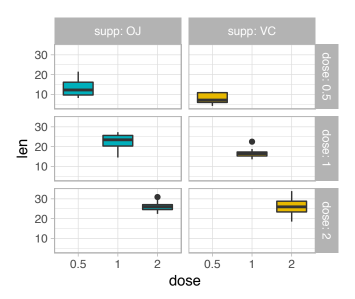
```
p + facet_grid(dose ~ supp, margins=TRUE)
```

3. Facet scales

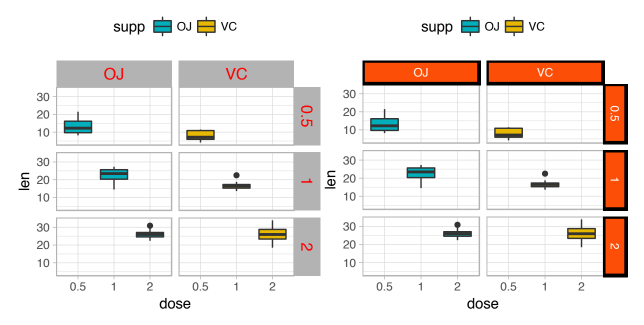
By default, all the panels have the same scales (scales="fixed"). They can be made independent, by setting scales to free, free_x, or free_y.

```
p + facet_grid(dose ~ supp, scales='free')
```

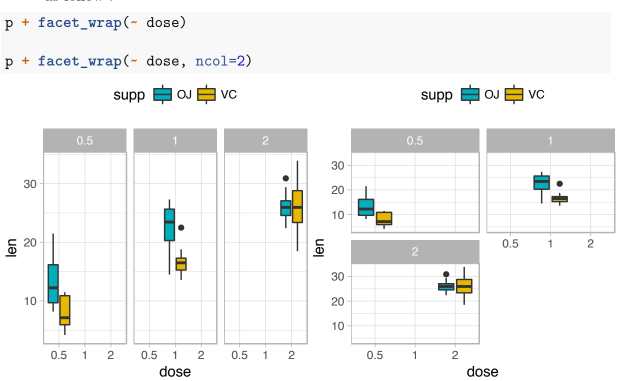
4. **Facet labels**: The argument labeller can be used to control the labels of the panels.



The appearance of facet labels can be modified as follow:



5. **facet_wrap**: Facets can be placed side by side using the function **facet_wrap()** as follow:



8.4 See also

• Create and Customize Multi-panel ggplots: Easy Guide to Facet. https://goo.gl/eRKHV7

Chapter 9

Arrange Multiple GGPlot on One Page

9.1 Introduction

This chapter describes, step by step, how to combine **multiple ggplot** on one page, as well as, over multiple pages, using helper functions available in the *ggpubr* R package. We'll also describe how to export the arranged plots to a file.

9.2 Prerequisites

Load required packages and set the theme function theme_pubr() [in ggpubr] as the default theme:

```
library(ggplot2)
library(ggpubr)
theme_set(theme_pubr())
```

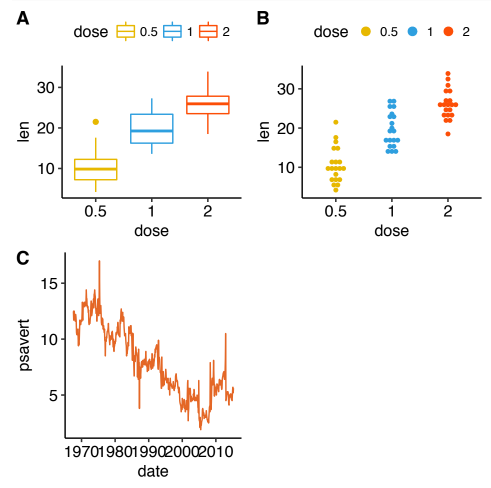
9.3 Arrange on one page

• Create some basic plots as follow:

```
# 0. Define custom color palette and prepare the data
my3cols <- c("#E7B800", "#2E9FDF", "#FC4E07")
ToothGrowth$dose <- as.factor(ToothGrowth$dose)

# 1. Create a box plot (bp)
p <- ggplot(ToothGrowth, aes(x = dose, y = len))
bxp <- p + geom_boxplot(aes(color = dose)) +
    scale_color_manual(values = my3cols)</pre>
```

• Combine multiple ggplot on one page. Use the function ggarrange()[ggpubr package], a wrapper around the function plot_grid() [cowplot package]. Compared to plot_grid(), ggarange() can arrange multiple ggplots over multiple pages.



9.4 Annotate the arranged figure

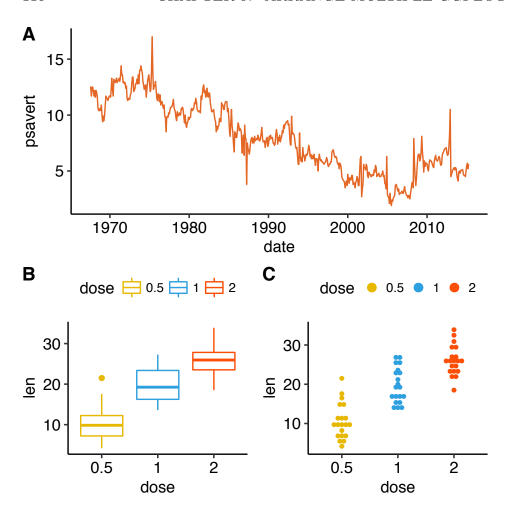
Key R function: annotate_figure() [in ggpubr].

9.5 Change column and row span of a plot

We'll use nested ggarrange() functions to change column/row span of plots. For example, using the R code below:

- the line plot (lp) will live in the first row and spans over two columns
- the box plot (bxp) and the dot plot (dp) will be first arranged and will live in the second row with two different columns

```
ggarrange(
  lp,  # First row with line plot
  # Second row with box and dot plots
  ggarrange(bxp, dp, ncol = 2, labels = c("B", "C")),
  nrow = 2,
  labels = "A"  # Label of the line plot
)
```

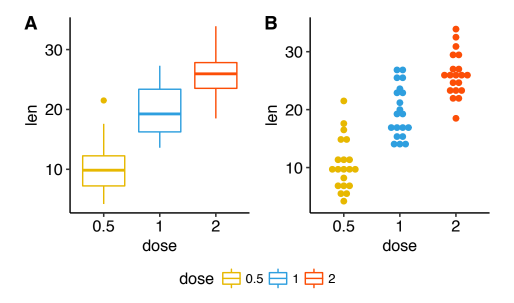


9.6 Use shared legend for combined ggplots

To place a common unique legend in the margin of the arranged plots, the function ggarrange() [in ggpubr] can be used with the following arguments:

- common.legend = TRUE: place a common legend in a margin
- legend: specify the legend position. Allowed values include one of c("top", "bottom", "left", "right")

```
ggarrange(
  bxp, dp, labels = c("A", "B"),
  common.legend = TRUE, legend = "bottom"
)
```



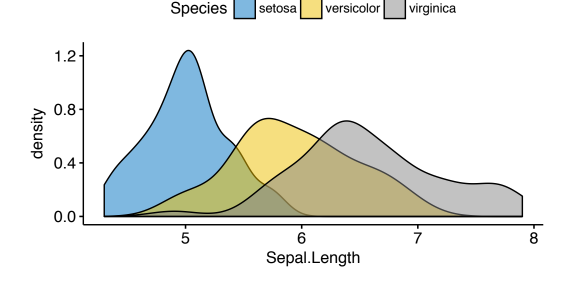
9.7 Mix table, text and ggplot2 graphs

In this section, we'll show how to plot a table and text alongside a chart. The iris data set will be used.

We start by creating the following plots:

- 1. a **density plot** of the variable "Sepal.Length". R function: **ggdensity**() [in ggpubr]
- 2. a plot of the **summary table** containing the descriptive statistics (mean, sd, ...) of Sepal.Length.
 - R function for computing descriptive statistics: **desc_statby**() [in ggpubr].
 - R function to draw a textual table: **ggtexttable**() [in ggpubr].
- 3. a plot of a text **paragraph**. R function: **ggparagraph**() [in ggpubr].

We finish by arranging/combining the three plots using the function **ggarrange**() [in ggpubr]



Species	length	mean	sd
setosa	50	5.01	0.352
versicolor	50	5.94	0.516
virginica	50	6.59	0.636

iris data set gives the measurements in cm of the variables sepal length and width and petal length and width, respectively, for 50 flowers from each of 3 species of iris. The species are Iris setosa, versicolor, and virginica.

9.8 Arrange over multiple pages

If you have a long list of ggplots, say n=20 plots, you may want to arrange the plots and to place them on multiple pages. With 4 plots per page, you need 5 pages to hold the 20 plots.

The function ggarrange() [ggpubr] provides a convenient solution to arrange multiple

ggplots over multiple pages. After specifying the arguments nrow and ncol, ggarrange()' computes automatically the number of pages required to hold the list of the plots. It returns a list of arranged ggplots.

For example the following R code,

returns a list of two pages with two plots per page. You can visualize each page as follow:

```
multi.page[[1]] # Visualize page 1
multi.page[[2]] # Visualize page 2
```

You can also export the arranged plots to a pdf file using the function ggexport() [ggpubr]:

```
ggexport(multi.page, filename = "multi.page.ggplot2.pdf")
```

See the PDF file: Multi.page.ggplot2¹

9.9 Export the arranged plots

R function: ggexport() [in ggpubr].

• Export the arranged figure to a pdf, eps or png file (one figure per page).

```
ggexport(figure, filename = "figure1.pdf")
```

• It's also possible to arrange the plots (2 plot per page) when exporting them.

Export individual plots to a pdf file (one plot per page):

```
ggexport(bxp, dp, lp, bxp, filename = "test.pdf")
```

Arrange and export. Specify nrow and nool to display multiple plots on the same page:

9.10 See also

ggplot2 - Easy Way to Mix Multiple Graphs on The Same Page. https://goo.gl/WrieY4

^{1//}www.slideshare.net/kassambara/multipageggplot2

Chapter 10

Customize GGPlot

In this chapter, we'll show how to change the global appearance of a ggplot.

10.1 Prerequisites

1. Load packages and set the default theme:

```
library(ggplot2)
library(ggpubr)
theme_set(
  theme_pubr() +
    theme(legend.position = "right")
)
```

- 2. Create a box plot (bxp) and a scatter plot (sp) that we'll customize in the next section:
- Box plot using the ToothGrowth dataset:

```
# Convert the variable dose from numeric to factor variable
ToothGrowth$dose <- as.factor(ToothGrowth$dose)
bxp <- ggplot(ToothGrowth, aes(x=dose, y=len)) +
  geom_boxplot(aes(color = dose)) +
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))</pre>
```

• Scatter plot using the cars dataset

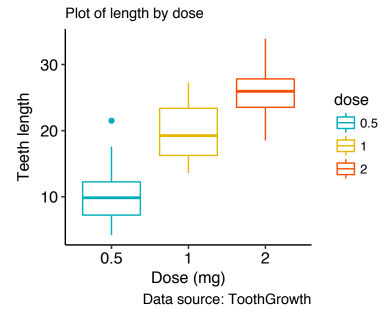
```
sp <- ggplot(cars, aes(x = speed, y = dist)) +
  geom_point()</pre>
```

10.2 Titles and axis labels

Key function: labs(). Used to change the main title, the subtitle, the axis labels and captions.

1. Add a title, subtitle, caption and change axis labels

Effect of Vitamin C on Tooth Growth



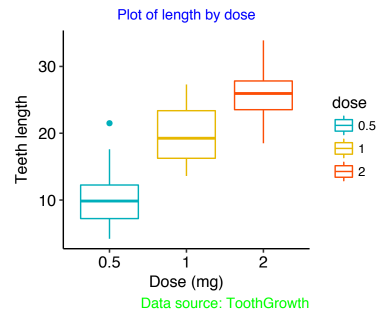
2. Change the appearance of titles

• Key functions: theme() and element text():

```
theme(
  plot.title = element_text(),
  plot.subtitle.title = element_text(),
  plot.caption = element_text()
)
```

- Arguments of the function element text() include:
 - color, size, face, family: to change the text font color, size, face ("plain", "italic", "bold", "bold.italic") and family.
 - lineheight: change space between two lines of text elements. Number between 0 and 1. Useful for multi-line plot titles.
 - hjust and vjust: number in [0, 1], for horizontal and vertical adjustement of titles, respectively.
 - * hjust = 0.5: Center the plot titles.
 - * hjust = 1: Place the plot title on the right
 - * hjust = 0: Place the plot title on the left
- Examples of R code:
 - Center main title and subtitle (hjust = 0.5)
 - Change color, size and face

Effect of Vitamin C on Tooth Growth



3. Case of long titles. If the title is too long, you can split it into multiple lines using \n. In this case you can adjust the space between text lines by specifying the argument lineheight in the theme function element_text():

```
bxp + labs(title = "Effect of Vitamin C on Tooth Growth \n in Guinea Pigs")+
theme(plot.title = element_text(lineheight = 0.9))
```

10.3 Axes: Limits, Ticks and Log

10.3.1 Axis limits and scales

3 Key functions to set the axis limits and scales:

1. Without clipping (preferred). Cartesian coordinates. The Cartesian coordinate system is the most common type of coordinate system. It will zoom the plot, without clipping the data.

```
sp + coord_cartesian(xlim = c(5, 20), ylim = (0, 50))
```

2. With clipping the data (removes unseen data points). Observations not in this range will be dropped completely and not passed to any other layers.

```
# Use this
sp + scale_x_continuous(limits = c(5, 20)) +
```

```
scale_y_continuous(limits = c(0, 50))

# Or this shothand functions
sp + xlim(5, 20) + ylim(0, 50)
```

Note that, scale_x_continuous() and scale_y_continuous() remove all data points outside the given range and, the coord_cartesian() function only adjusts the visible area.

In most cases you would not see the difference, but if you fit anything to the data the functions scale_x_continuous() / scale_y_continuous() would probably change the fitted values.

3. Expand the plot limits to ensure that a given value is included in all panels or all plots.

```
# set the intercept of x and y axes at (0,0)
sp + expand_limits(x = 0, y = 0)

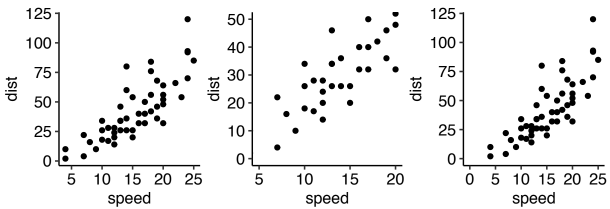
# Expand plot limits
sp + expand_limits(x = c(5, 50), y = c(0, 150))
```

Examples of R code:

```
# Default plot
print(sp)

# Change axis limits using coord_cartesian()
sp + coord_cartesian(xlim =c(5, 20), ylim = c(0, 50))

# set the intercept of x and y axis at (0,0)
sp + expand_limits(x = 0, y = 0)
```



10.3.2 Log scale

Key functions to set a logarithmic axis scale:

1. Scale functions. Allowed value for the argument trans: log2 and log10.

```
sp + scale_x_continuous(trans = "log2")
sp + scale_y_continuous(trans = "log2")
```

2. Transformed cartesian coordinate system. Possible values for x and y are " $\log 2$ ", " $\log 10$ ", "sqrt", ...

```
sp + coord_trans(x = "log2", y = "log2")
```

3. Display log scale ticks. Make sens only for log10 scale:

```
sp + scale_y_log10() + annotation_logticks()
```

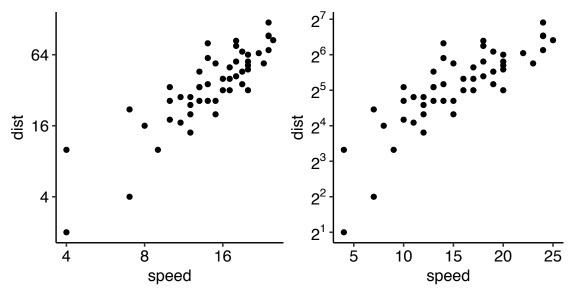
Note that, the scale functions transform the data. If you fit anything to the data it would probably change the fitted values.

An alternative is to use the function $coord_trans()$, which occurs after statistical transformation and will affect only the visual appearance of geoms.

Example of R code

```
# Set axis into log2 scale
# Possible values for trans : 'log2', 'log10','sqrt'
sp + scale_x_continuous(trans = 'log2') +
    scale_y_continuous(trans = 'log2')

# Format y axis tick mark labels to show exponents
require(scales)
sp + scale_y_continuous(
    trans = log2_trans(),
    breaks = trans_breaks("log2", function(x) 2^x),
    labels = trans_format("log2", math_format(2^x))
)
```

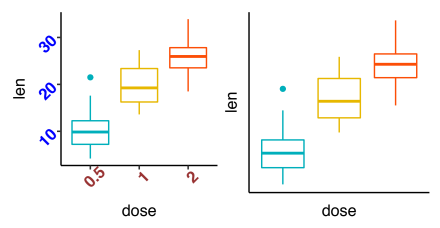


10.3.3 Axis Ticks: Set and Rotate Text Labels

Start by creating a box plot:

```
bxp <- ggplot(ToothGrowth, aes(x=dose, y=len)) +
  geom_boxplot(aes(color = dose)) +
  scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))+
  theme(legend.position = "none")</pre>
```

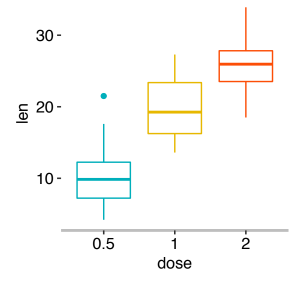
1. Change the style and the orientation angle of axis tick labels. For a vertical rotation of x axis labels use angle = 90.



To adjust the postion of the axis text, you can specify the argument hjust and vjust, which values should be comprised between 0 and 1.

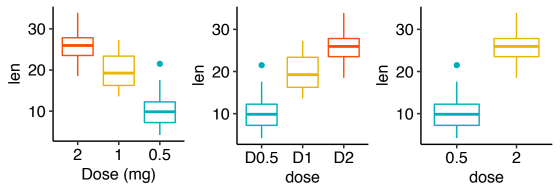
- 2. Change axis lines:
 - Remove the y-axis line
 - Change the color, the size and the line type of the x-axis line:

```
bxp + theme(
  axis.line.y = element_blank(),
  axis.line = element_line(
    color = "gray", size = 1, linetype = "solid"
    )
)
```



3. Customize discrete axis. Use the function scale_x_discrete() or scale_y_discrete() depending on the axis you want to change.

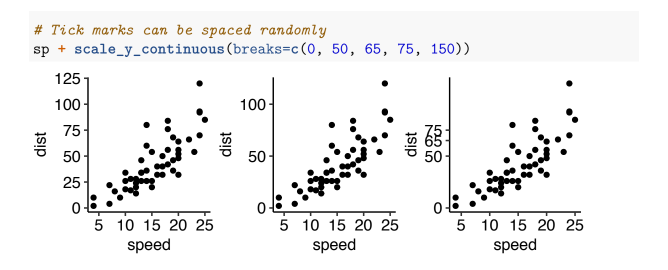
Here, we'll customize the x-axis of the box plot:



4. Customize continuous axis. Change axis ticks interval.

```
# Default scatter plot
sp <- ggplot(cars, aes(x = speed, y = dist)) +
    geom_point()
sp

# Break y axis by a specified value
# a tick mark is shown on every 50
sp + scale_y_continuous(breaks=seq(0, 150, 50))</pre>
```



10.4 Legends: Title, Position and Appearance

Start by creating a box plot using the ToothGrowth data set. Change the box plot fill color according to the grouping variable dose.

```
library(ggplot2)
ToothGrowth$dose <- as.factor(ToothGrowth$dose)
bxp <- ggplot(ToothGrowth, aes(x = dose, y = len))+
  geom_boxplot(aes(fill = dose)) +
  scale_fill_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))</pre>
```

10.4.1 Change legend title and position

- 1. **Legend title**. Use labs() to changes the legend title for a given aesthetics (fill, color, size, shape, . . .). For example:
- Use p + labs(fill = "dose") for geom_boxplot(aes(fill = dose))
- Use p + labs(color = "dose") for geom_boxplot(aes(color = dose))
- and so on for linetype, shape, etc
- 2. **Legend position**. The default legend position is "right". Use the function theme() with the argument legend.position to specify the legend position.

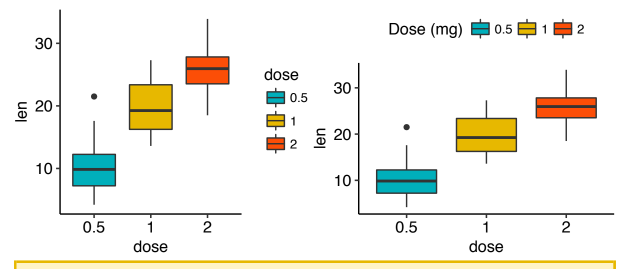
Allowed values for the legend position include: "left", "top", "right", "bottom", "none".

Legend loction can be also a numeric vector c(x,y), where x and y are the coordinates of the legend box. Their values should be between 0 and 1. c(0,0) corresponds to the "bottom left" and c(1,1) corresponds to the "top right" position. This makes it possible to place the legend inside the plot.

Examples:

```
# Default plot
bxp
```

```
# Change legend title and position
bxp +
  labs(fill = "Dose (mg)") +
  theme(legend.position = "top")
```



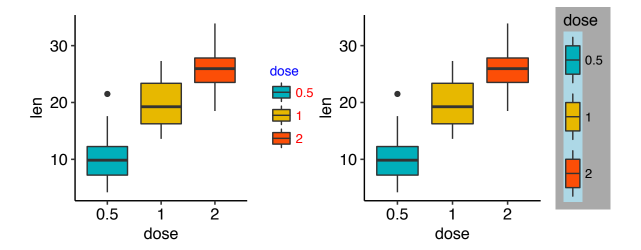
To remove legend, use p + theme(legend.position = "none").

10.4.2 Change the appearance of legends

- Change legend text color and size
- Change the legend box background color

```
# Change the appearance of legend title and text labels
bxp + theme(
  legend.title = element_text(color = "blue", size = 10),
  legend.text = element_text(color = "red")
)

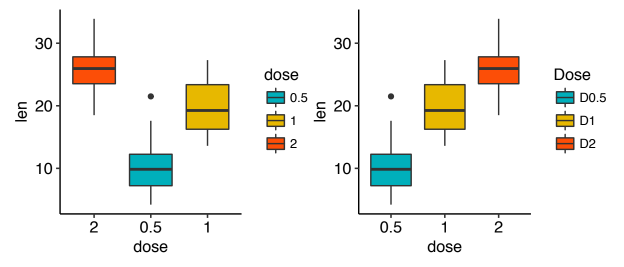
# Change legend background color, key size and width
bxp + theme(
  # Change legend background color
  legend.background = element_rect(fill = "darkgray"),
  legend.key = element_rect(fill = "lightblue", color = NA),
  # Change legend key size and key width
  legend.key.size = unit(1.5, "cm"),
  legend.key.width = unit(0.5, "cm")
)
```



10.4.3 Rename legend labels and change the order of items

```
# Change the order of legend items
bxp + scale_x_discrete(limits=c("2", "0.5", "1"))

# Edit legend title and labels for the fill aesthetics
bxp + scale_fill_manual(
   values = c("#00AFBB", "#E7B800", "#FC4E07"),
   name = "Dose",
   breaks = c("0.5", "1", "2"),
   labels = c("D0.5", "D1", "D2")
   )
```



Other manual scales to set legends for a given aesthetic:

```
# Color of lines and points
scale_color_manual(name, labels, limits, breaks)
# For linetypes
scale_linetype_manual(name, labels, limits, breaks)
# For point shapes
scale_shape_manual(name, labels, limits, breaks)
```

```
# For point size
scale_size_manual(name, labels, limits, breaks)
# Opacity/transparency
scale_alpha_manual(name, labels, limits, breaks)
```

10.5 Themes gallery

Start by creating a simple box plot:

```
bxp <- ggplot(ToothGrowth, aes(x = factor(dose), y = len)) +
  geom_boxplot()</pre>
```

10.5.1 Use themes in ggplot2 package

Several simple functions are available in ggplot2 package to set easily a ggplot theme. These include:

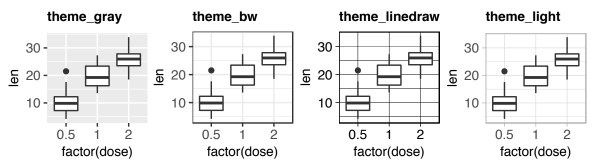
- theme_gray(): Gray background color and white grid lines. Put the data forward to make comparisons easy.
- theme_bw(): White background and gray grid lines. May work better for presentations displayed with a projector.
- theme_linedraw(): A theme with black lines of various widths on white backgrounds, reminiscent of a line drawings.
- theme_light(): A theme similar to theme_linedraw() but with light grey lines and axes, to direct more attention towards the data.

```
bxp + theme_gray(base_size = 14)

bxp + theme_bw()

bxp + theme_linedraw()

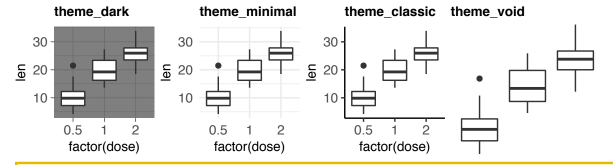
bxp + theme_light()
```



- theme_dark(): Same as theme_light but with a dark background. Useful to make thin coloured lines pop out.
- theme minimal(): A minimal theme with no background annotations
- theme classic(): A classic theme, with x and y axis lines and no gridlines.

• theme_void(): a completely empty theme, useful for plots with non-standard coordinates or for drawings.

```
bxp + theme_dark()
bxp + theme_minimal()
bxp + theme_classic()
bxp + theme_void()
```



Note that, additional themes are availbale in the ggthemes R package^a.

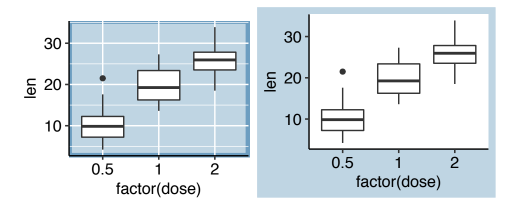
^ahttps://cran.r-project.org/web/packages/ggthemes/vignettes/ggthemes.html

10.6 Background color and grid lines

• Create a simple box plot:

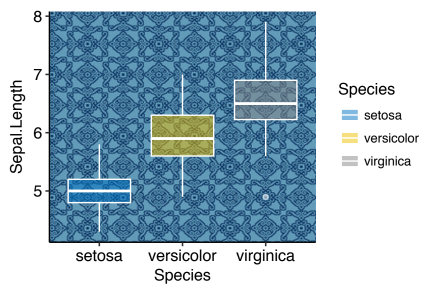
```
p <- ggplot(ToothGrowth, aes(factor(dose), len)) +
  geom_boxplot()</pre>
```

• Change the panel background (1) and the plot background (2) colors:



10.7 Add background image to ggplot2 graphs

- 1. **Import the background image**. Use either the function readJPEG() [in *jpeg* package] or the function 'readPNG()[in *png* package] depending on the format of the background image.
- 2. Combine a ggplot with the background image. R function: background_image() [in ggpubr].



10.8. COLORS 133

10.8 Colors

A color can be specified either by name (e.g.: "red") or by hexadecimal code (e.g.: "#FF1234"). In this section, you will learn how to change ggplot colors by groups and how to set gradient colors.

0. Set ggplot theme to theme minimal():

```
theme_set(
  theme_minimal() +
    theme(legend.position = "top")
)
```

1. **Initialize ggplots** using the iris data set:

```
# Box plot
bp <- ggplot(iris, aes(Species, Sepal.Length))

# Scatter plot
sp <- ggplot(iris, aes(Sepal.Length, Sepal.Width))</pre>
```

2. **Specify a single color**. Change the fill color (in box plots) and points color (in scatter plots).

```
# Box plot
bp + geom_boxplot(fill = "#FFDB6D", color = "#C4961A")
# Scatter plot
sp + geom_point(color = "#00AFBB")
                                                    4.5
   8
                                                    4.0
Sepal.Length
                                                Sepal.Width
                                                    3.5
                                                    3.0
                                                    2.5
   5
                                                    2.0
                       versicolor
                                    virginica
                                                                5
           setosa
                                                                                               8
                       Species
                                                                     Sepal.Length
```

3. Change colors by groups.

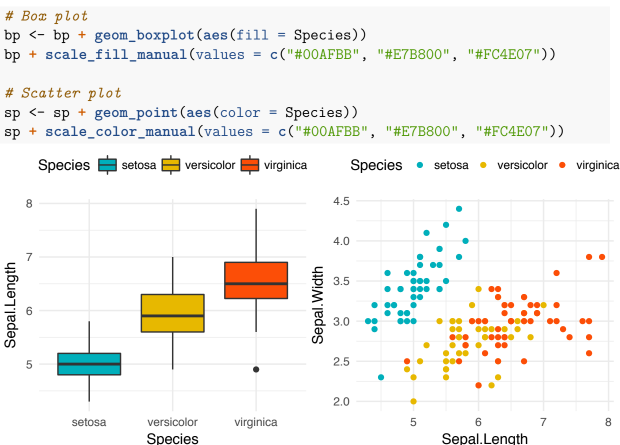
You can change colors according to a grouping variable by:

• Mapping the argument color to the variable of interest. This will be applied to points, lines and texts

• Mapping the argument fill to the variable of interest. This will change the fill color of areas, such as in box plot, bar plot, histogram, density plots, etc.

It's possible to specify manually the color palettes by using the functions:

- scale fill manual() for box plot, bar plot, violin plot, dot plot, etc
- scale_color_manual() or scale_colour_manual() for lines and points



Find below, two color-blind-friendly palettes, one with gray, and one with black (source: http://jfly.iam.u-tokyo.ac.jp/color/).

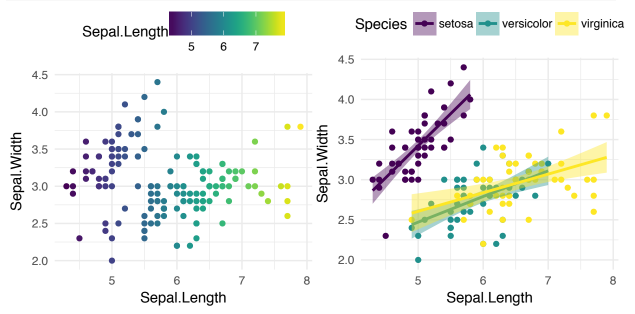
4. Use viridis color palettes. The viridis R package provides color palettes to make beautiful plots that are: printer-friendly, perceptually uniform and easy to read by those with colorblindness. Key functions scale color viridis() and

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```
scale_fill_viridis()
```

```
library(viridis)
# Gradient color
ggplot(iris, aes(Sepal.Length, Sepal.Width))+
    geom_point(aes(color = Sepal.Length)) +
    scale_color_viridis(option = "D")

# Discrete color. use the argument discrete = TRUE
ggplot(iris, aes(Sepal.Length, Sepal.Width))+
    geom_point(aes(color = Species)) +
    geom_smooth(aes(color = Species, fill = Species), method = "lm") +
    scale_color_viridis(discrete = TRUE, option = "D")+
    scale_fill_viridis(discrete = TRUE)
```

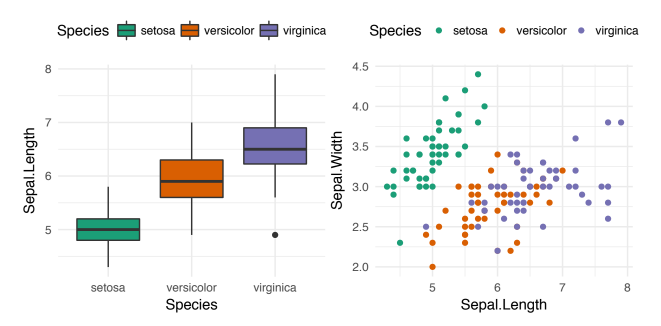


- 5. Use RColorBrewer palettes. Two color scale functions are available in ggplot2 for using the colorbrewer palettes:
- scale fill brewer() for box plot, bar plot, violin plot, dot plot, etc
- scale color brewer() for lines and points

For example:

```
# Box plot
bp + scale_fill_brewer(palette = "Dark2")

# Scatter plot
sp + scale_color_brewer(palette = "Dark2")
```



To display colorblind-friendly brewer palettes, use this R code:





- 6. Other discrete color palettes:
 - Scientific journal color palettes in the ggsci R package. Contains a collection of high-quality color palettes inspired by colors used in scientific journals, data visualization libraries, and more. For example:

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- scale color npg() and scale fill npg(): Nature Publishing Group
- scale_color_aaas() and scale_fill_aaas(): American Association for the Advancement of Science
- scale_color_lancet() and scale_fill_lancet(): Lancet journal
- scale_color_jco() and scale_fill_jco(): Journal of Clinical Oncology
- Wes Anderson color palettes in the wesanderson R package. Contains 16 color palettes from Wes Anderson movies.

For example:

```
# jco color palette from the ggsci package
bp + ggsci::scale_fill_jco()
# Discrete color from wesanderson package
library(wesanderson)
bp + scale_fill_manual(
  values = wes_palette("GrandBudapest1", n = 3)
                                                 Species etosa versicolor virginica
     Species = setosa = versicolor i virginica
   8
                                                8
Sepal.Length
                                             Sepal.Length
   5
                                                5
          setosa
                     versicolor
                                 virginica
                                                       setosa
                                                                 versicolor
                                                                              virginica
```

You can find more examples at ggsci package vignettes¹ and at wesanderson github page²

7. **Set gradient colors**. For gradient colors, you should map the map the argument color and/or fill to a continuous variable. In the following example, we color points according to the variable: **Sepal.Length**.

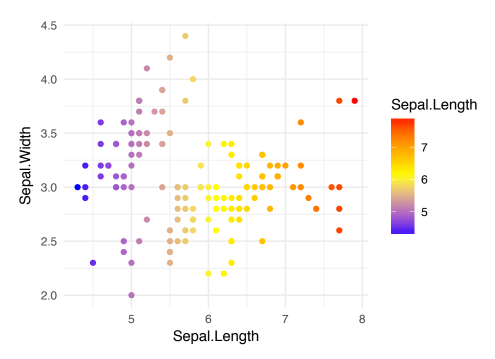
Species

```
ggplot(iris, aes(Sepal.Length, Sepal.Width))+
  geom_point(aes(color = Sepal.Length)) +
  scale_color_gradientn(colours = c("blue", "yellow", "red"))+
  theme(legend.position = "right")
```

Species

https://cran.r-project.org/web/packages/ggsci/vignettes/ggsci.html

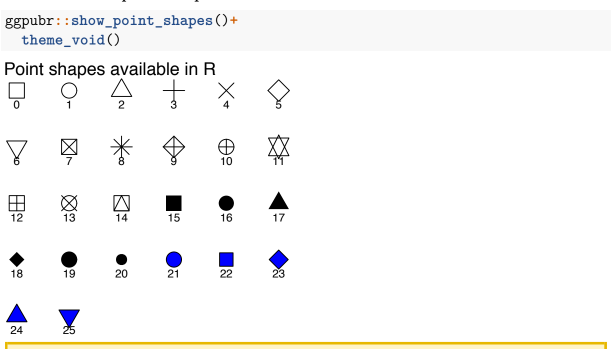
²https://github.com/karthik/wesanderson



8. Design and use the power of color palette at https://goo.gl/F5g3Lb

10.9 Points shape, color and size

1. Common point shapes available in R:



Note that, the point shape options from pch 21 to 25 are open symbols that can be filled by a color. Therefore, you can use the fill argument in geom_point() for these symbols.

2. Change ggplot point shapes. The argument shape is used, in the function geom_point() [ggplot2], for specifying point shapes.

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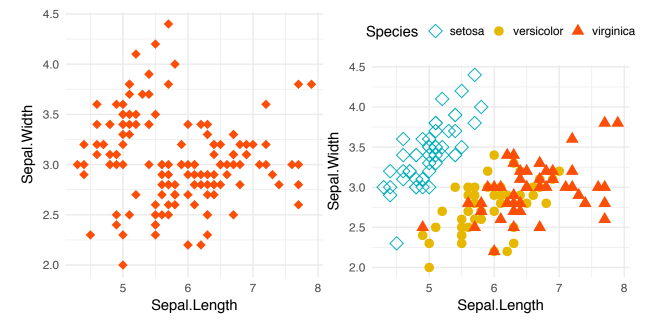
It's also possible to change point shapes and colors by groups. In this case, ggplot2 will use automatically a default color palette and point shapes. You can change manually the appearance of points using the following functions:

- scale_shape_manual(): to change manually point shapes
- scale_color_manual(): to change manually point colors
- scale size manual(): to change manually the size of points

Create a scatter plot and change points shape, color and size:

```
# Create a simple scatter plot
ggplot(iris, aes(Sepal.Length, Sepal.Width)) +
    geom_point(shape = 18, color = "#FC4E07", size = 3)+
    theme_minimal()

# Change point shapes and colors by groups
ggplot(iris, aes(Sepal.Length, Sepal.Width)) +
    geom_point(aes(shape = Species, color = Species), size = 3) +
    scale_shape_manual(values = c(5, 16, 17)) +
    scale_color_manual(values = c("#00AFBB", "#E7B800", "#FC4E07"))+
    theme_minimal() +
    theme(legend.position = "top")
```



10.10 Line types

1. Common line types available in R:

```
ggpubr::show_line_types()+
theme_gray()
```

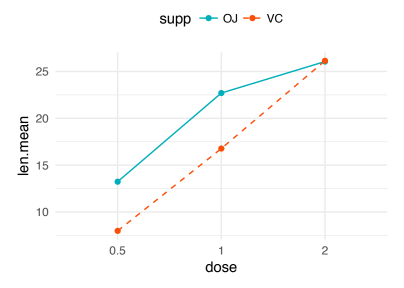
Line types available in R

2. Change line types. To change a single line, use for example linetype = "dashed".

In the following R code, we'll change line types and colors by groups. To modify the default colors and line types, the function scale_color_manual() and scale_linetype_manual() can be used.

```
# Create some data.
# # Compute the mean of `len` grouped by dose and supp
library(dplyr)
df2 <- ToothGrowth %>%
  group_by(dose, supp) %>%
  summarise(len.mean = mean(len))
df2
## # A tibble: 6 x 3
## # Groups:
               dose [?]
##
       dose
              supp len.mean
##
     <fctr> <fctr>
                      <dbl>
## 1
        0.5
                OJ
                      13.23
## 2
                       7.98
        0.5
                VC
## 3
          1
                OJ
                      22.70
          1
                VC
## 4
                      16.77
          2
## 5
                OJ
                      26.06
          2
## 6
                VC
                      26.14
# Change manually line type and color manually
```

```
# Change manually line type and color manually
ggplot(df2, aes(x = dose, y = len.mean, group = supp)) +
  geom_line(aes(linetype = supp, color = supp))+
  geom_point(aes(color = supp))+
  scale_linetype_manual(values=c("solid", "dashed"))+
  scale_color_manual(values=c("#00AFBB","#FC4E07"))
```



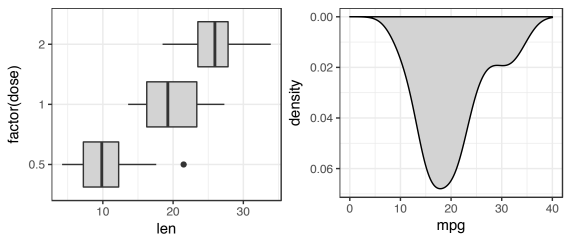
10.11 Rotate a ggplot

Key functions:

- coord_flip(): creates horizontal plots
- scale_x_reverse() and scale_y_reverse(): reverse the axis

```
# Horizontal box plot
ggplot(ToothGrowth, aes(factor(dose), len)) +
    geom_boxplot(fill = "lightgray") +
    theme_bw() +
    coord_flip()

# Reverse y axis
ggplot(mtcars, aes(mpg))+
    geom_density(fill = "lightgray") +
    xlim(0, 40) +
    theme_bw()+
    scale_y_reverse()
```



10.12 Plot annotation

10.12.1 Add straight lines

Key R functions:

- **geom_hline**(yintercept, linetype, color, size): add horizontal lines
- **geom_vline**(xintercept, linetype, color, size): add vertical lines
- **geom_abline**(intercept, slope, linetype, color, size): add regression lines
- **geom_segment()**: add segments

Create a simple scatter plot:

Creating a simple scatter plot

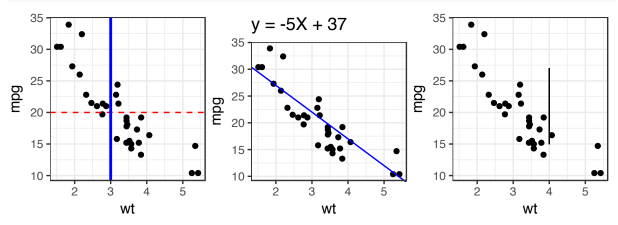
```
sp <- ggplot(data = mtcars, aes(x = wt, y = mpg)) +
  geom_point()+theme_bw()</pre>
```

• Add straight lines and segments

```
# Add horizontal line at y = 20; and vertical line at x = 3
sp + geom_hline(yintercept = 20, linetype = "dashed", color = "red") +
    geom_vline(xintercept = 3, color = "blue", size = 1)

# Add regression line
sp + geom_abline(intercept = 37, slope = -5, color="blue")+
    labs(title = "y = -5X + 37")

# Add a vertical line segment from
# point A(4, 15) to point B(4, 27)
sp + geom_segment(x = 4, y = 15, xend = 4, yend = 27)
```



Add arrows, curves and rectangles:

wt

```
sp + geom_curve(aes(x = 2, y = 15, xend = 3, yend = 15))
# Add rectangles
ggplot(data = mtcars, aes(x = wt, y = mpg)) +
  geom_rect(xmin = 3, ymin = -Inf, xmax = 4, ymax = Inf,
             fill = "lightgray") +
  geom_point() + theme_bw()
  30
                             30
                                                       30
  25
                             25
                                                       25
                             20
                                                       20
  15
                             15
                                                        15
  10
                             10
                                                        10
       2
            3
                      5
                                  2
                                                 5
                                                             2
                                       3
                                                                  3
```

10.12.2 Text annotation

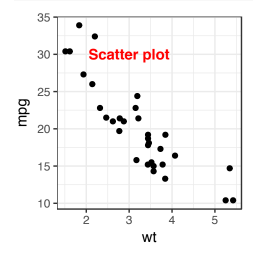
wt

Key ggplot2 function:

- **geom_text**(): adds text directly to the plot
- **geom_label**(): draws a rectangle underneath the text, making it easier to read.
- annotate(): useful for adding small text annotations at a particular location on the plot

wt

• annotation custom(): Adds static annotations that are the same in every panel



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