

# Basics of R Graphics

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Lab Note: STA321 Topics of Advanced Statistics

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## 1 Introduction

This note introduces the graphical capabilities of base R graphical functions. Base R graphical system contains a set of **high-level plotting functions** such as `plot()`, `hist()`, `barplot()`, etc., and also a set of **low-level functions** that are used jointly with the high-level plotting functions such as `points()`, `lines()`, `text()`, `segments()`, etc. to make a flexible graphical system.

## 2 R Graphic Devices

R is able to output graphics to the screen or save them directly to a file (e.g. postscript, pdf, svg, png, jpeg, etc.). The different functions for producing graphical output are known as **Graphic Devices**. For example, `pdf()` would invoke the `pdf device`, while `png()` would invoke the `png device`. Type `?Devices` into the R console to see a list of graphical devices that are available to R on your system.

By default, graphical output is sent to the screen. As R is cross-platform, the graphics device for producing `**screen**` graphics differs by the system. The available fonts may also differ by the system and graphical device.

## 3 Plotting with Base Graphics: plot()

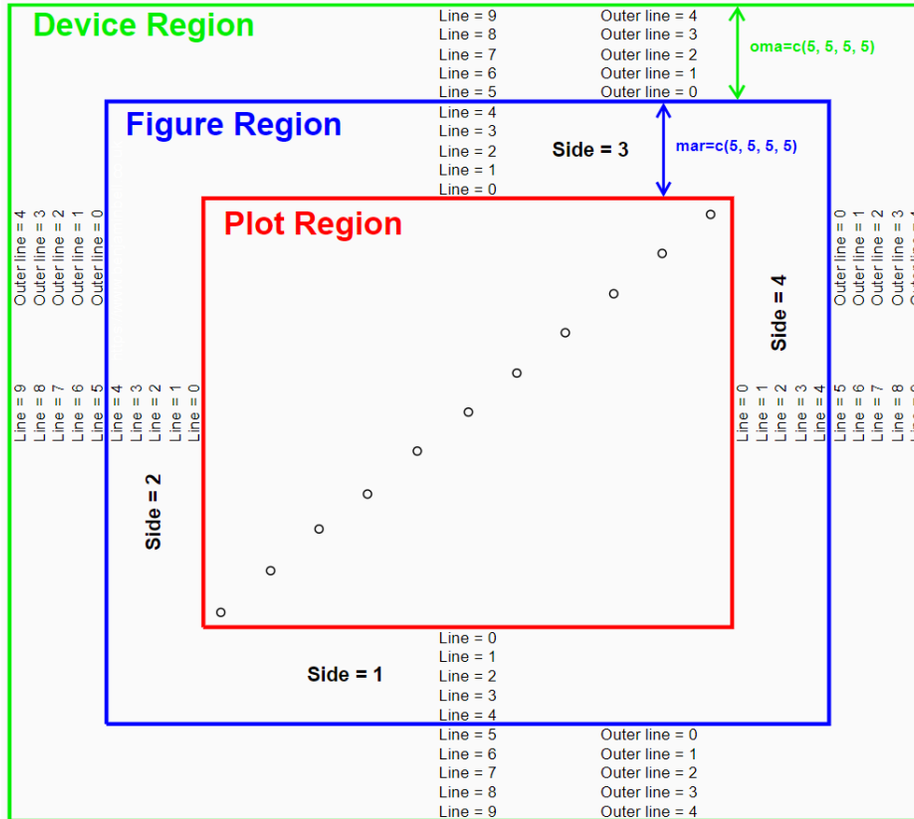
`plot()` is the most important high-level graphic function. When using `plot()`, the output is sent to the **graphics device**. This creates a “plot region” on the graphics device. The visible area of the graphics device is also known as the “device region”. The plot region on the other hand is the area of the plot (usually bounded by a box) and does not include the plot axes, labels, or margins.

The plot region, plus the axes, labels, and margins are known as the “figure region”. Often, the device region and figure region can be the same size - but they are not the same thing.

### 3.1 Outer and Inner Margins

```
par(mar=c(x, x, x, x), oma = c(x, x, x, x))
```

Almost all kinds of plots, charts, and graphs can be produced using base graphics. These plots can be fully customized using `par()` (graphical parameter). The following figure explains the graphical parameters we can use to customize the graphical layout.



Example 1:

```
iris = read.table("https://raw.githubusercontent.com/pengdsci/sta321/main/ww02/w02-iris.txt", header = TRUE)
par(mfrow = c(1,1), oma=c(1, 1, 1, 1), mar=c(2, 2, 2, 2))
plot(iris$SepalLength, iris$PetalLength, pch = 16, cex = 1.5)
title(main="Sepal Length vs Petal Length (Outer)",
      outer=TRUE,
      col.main = "red",
      cex.main=1)
title(main="Sepal Length vs Petal Length (Inner)",
      outer=FALSE,
      col.main = "blue",
      cex.main=0.8)
## Coloring points with the transparency level
## species ID
setosa = which(iris$Classification == "Iris-setosa")
versicolor = which(iris$Classification == "Iris-versicolor")
virginica = which(iris$Classification == "Iris-virginica")
## adding points
points(iris$SepalLength[setosa], iris$PetalLength[setosa],
       pch=16,
```

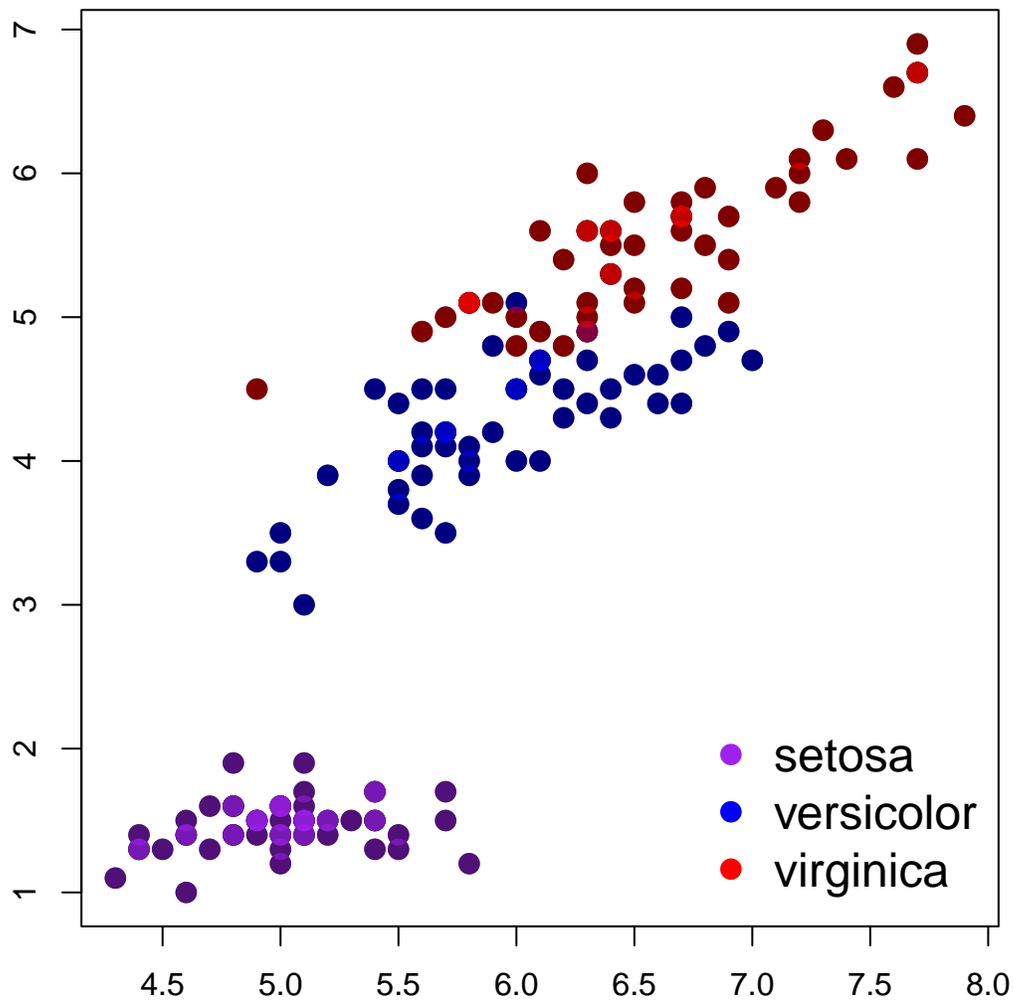
```

col = alpha("purple", 0.5),    # add a transparency level
cex = 1.5)
points(iris$SepalLength[versicolor], iris$PetalLength[versicolor],
pch=16,
col = alpha("blue", 0.5),
cex = 1.5)
points(iris$SepalLength[virginica], iris$PetalLength[virginica],
pch=16,
col = alpha("red", 0.5),
cex = 1.5)
legend("bottomright", c("setosa", "versicolor", "virginica"),
col=c("purple", "blue", "red"),
pch=rep(16,3), cex=rep(1.5,3), bty="n")

```

### Sepal Length vs Petal Length (Outer)

#### Sepal Length vs Petal Length (Inner)



### 3.2 Simple multi-panel Plots

layout() or par(mfrow = c())

## Example 2

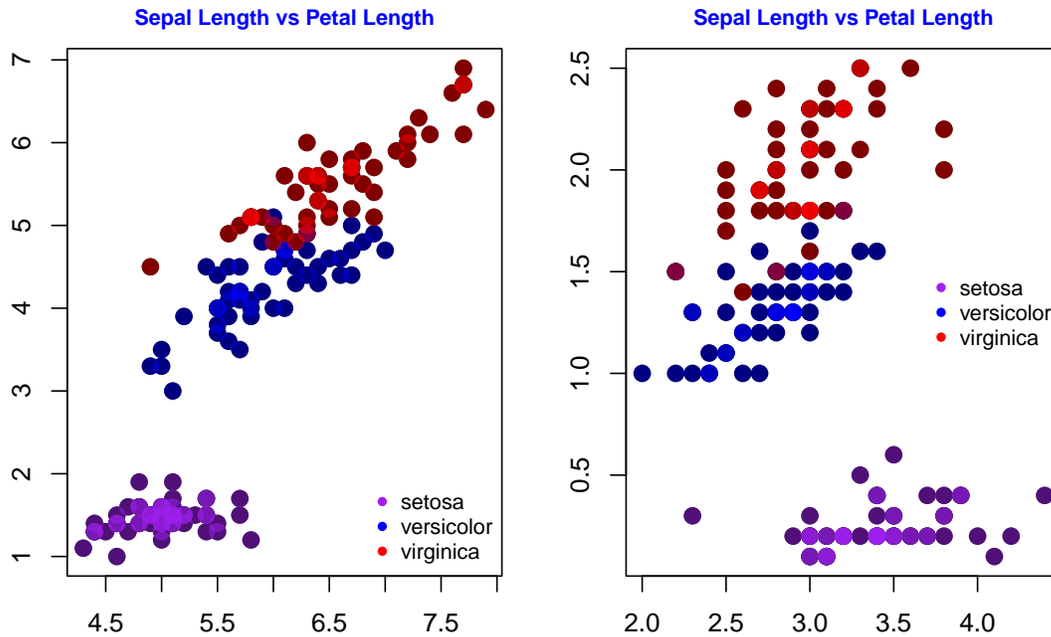
```
iris = read.table("https://raw.githubusercontent.com/pengdsci/sta321/main/ww02/w02-iris.txt", header = TRUE)
## species ID
setosa = which(iris$Classification == "Iris-setosa")
versicolor = which(iris$Classification == "Iris-versicolor")
virginica = which(iris$Classification == "Iris-virginica")
###
par(mfrow = c(1,2), oma=c(2, 2, 2, 2), mar=c(2, 2, 2, 2))
plot(iris$SepalLength, iris$PetalLength, pch = 16, cex = 1.5,
      xlab = "Sepal Width",
      ylab = "Petal Width")
title(main="Sepal Length vs Petal Length",
      outer=FALSE,
      col.main = "blue",
      cex.main=0.8)
## adding points
points(iris$SepalLength[setosa], iris$PetalLength[setosa],
       pch=16,
       col = alpha("purple", 0.5),
       cex = 1.5)
points(iris$SepalLength[versicolor], iris$PetalLength[versicolor],
       pch=16,
       col = alpha("blue", 0.5),
       cex = 1.5)
points(iris$SepalLength[virginica], iris$PetalLength[virginica],
       pch=16,
       col = alpha("red", 0.5),
       cex = 1.5)
legend("bottomright", c("setosa", "versicolor", "virginica"),
      col=c("purple", "blue", "red"),
      pch=rep(16,3), cex=0.8, bty="n")
##
plot(iris$SepalWidth, iris$PetalWidth, pch = 16, cex = 1.5,
      xlab = "Sepal Width",
      ylab = "Petal Width")
title(main="Sepal Length vs Petal Length",
      outer=FALSE,
      col.main = "blue",
      cex.main=0.8)
## adding points
points(iris$SepalWidth[setosa], iris$PetalWidth[setosa],
       pch=16,
       col = alpha("purple", 0.5),
       cex = 1.5)
points(iris$SepalWidth[versicolor], iris$PetalWidth[versicolor],
       pch=16,
       col = alpha("blue", 0.5),
       cex = 1.5)
points(iris$SepalWidth[virginica], iris$PetalWidth[virginica],
       pch=16,
       col = alpha("red", 0.5),
       cex = 1.5)
legend("right", c("setosa", "versicolor", "virginica"),
```

```

col=c("purple", "blue", "red"),
pch=rep(16,3), cex=0.8, bty="n")
## Overall Title
title(main="Comaring Sepal and Petal Sizes",
      outer=TRUE,
      col.main = "darkred",
      cex.main=1.2)

```

## Comaring Sepal and Petal Sizes



## 4 Making Your Own Color

You can make transparent colors using R and the `rgb()` command. These colors can be useful for charts and graphics with overlapping elements.

The `rgb()` command defines a new color using numerical values (0–255) for red, green and blue. In addition, we also set an alpha value (also 0–255), which sets the transparency (0 being fully transparent and 255 being “solid”).

The following example takes the standard blue and makes it transparent (~50%):

```
mycol <- rgb(0, 0, 255, max = 255, alpha = 125, names = "blue50")
```

You can also use `col2rgb()` to check the composition of an existing named R color and modify it to define your own favorite colors.

**Example 3:** We check the RGB code from several named R colors in the following

```
col2rgb("darkred")
```

```
##      [,1]
## red   139
## green   0
```

```
## blue    0
col2rgb("skyblue")
```

```
##      [,1]
## red   135
## green 206
## blue  235
```

```
col2rgb("gold")
```

```
##      [,1]
## red   255
## green 215
## blue   0
```

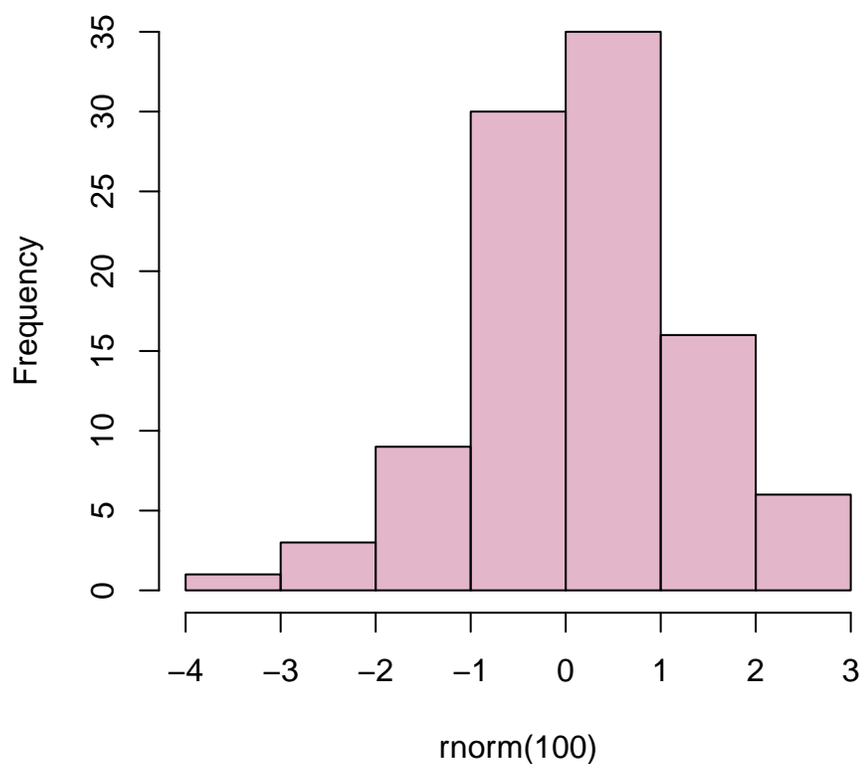
```
col2rgb("purple")
```

```
##      [,1]
## red   160
## green  32
## blue  240
```

**Example 3** We define a color by modifying “gold” and “purple”. I take the average of the RGB codes of “gold” and “purple” to see what the color looks like.

```
ratio = 0.4
avggoldpurple = round(ratio*col2rgb("gold") + (1-ratio)*col2rgb("purple"))
##
mycol.1 = rgb(avggoldpurple[1,1], avggoldpurple[2,1], avggoldpurple[3,1], max = 255, alpha = 125, names = "mycol.1")
###
hist(rnorm(100), col = mycol.1)
```

## Histogram of rnorm(100)



```
ratio = 0.8
avvgoldpurple = round(ratio*col2rgb("gold") + (1-ratio)*col2rgb("purple"))
##
mycol.2 = rgb(avvgoldpurple[1,1], avvgoldpurple[2,1], avvgoldpurple[3,1], max = 255, alpha = 125, names = "mycol.2")
###
hist(rnorm(100,1,1), col = mycol.2)
```

**Histogram of  $\text{rnorm}(100, 1, 1)$**

