

# Workshop 5: Programming in R

**QCBS R Workshop Series** 

**Québec Centre for Biodiversity Science** 



```
## Warning in library(package, lib.loc = lib.loc, character.only = TRUE,
## logical.return = TRUE, : there is no package called 'DiagrammeR'
## Installing package into '/home/travis/R/Library'
## (as 'lib' is unspecified)
## also installing the dependencies 'colorspace', 'gtable', 'isoband', 'hms', 'cpp11'
class: inverse, center, middle
```

# **About this workshop**

## Loading required package: DiagrammeR



### **Learning Objectives**

- 1. Recognizing control flow
- 2. Developing your first functions in R
- 3. Accelerating your code
- 4. Demonstrating useful R packages for biologists

# Review

# Objects

### **Review: Vectors**

#### Recall Workshop #1?

#### **Numeric vectors**

```
num.vector <- c(1, 4, 3,
9, 32, -4)
num.vector
# [1] 1 4 3 9 32 -4
```

#### Character vector

### Logical vector

```
bool_vector <- c(TRUE, TRUE, FALSE) # or c(T, T, F)
bool_vector
# [1] TRUE TRUE FALSE</pre>
```

### **Review: Data frames**

We can begin by creating multiple vectors (remember Workshop #1):

```
siteID <- c("A1.01", "A1.02", "B1.01", "B1.02")
soil_pH <- c(5.6, 7.3, 4.1, 6.0)
num.sp <- c(17, 23, 15, 7)
treatment <- c("Fert", "Fert", "No_fert", "No_fert")</pre>
```

We then combine them using the function <code>data.frame()</code>.

```
my.first.df
# siteID soil_pH num.sp treatment
# 1 A1.01    5.6    17    Fert
# 2 A1.02    7.3    23    Fert
# 3 B1.01    4.1    15    No_fert
# 4 B1.02    6.0    7    No_fert
```

my.first.df <- data.frame(siteID, soil pH, num.sp, treatment)</pre>

### Lists

We can also create lists by combining the vectors we created before.

```
my.first.list
# [[1]]
# [1] "A1.01" "A1.02" "B1.01" "B1.02"
#
# [[2]]
# [1] 5.6 7.3 4.1 6.0
#
# [[3]]
# [1] 17 23 15 7
#
# [[4]]
# [1] "Fert" "Fert" "No_fert" "No_fert"
```

my.first.list <- list(siteID, soil\_pH, num.sp, treatment)</pre>

# **Control flow**

### **Control flow**

Program flow control can be simply defined as the order in which a program is executed.

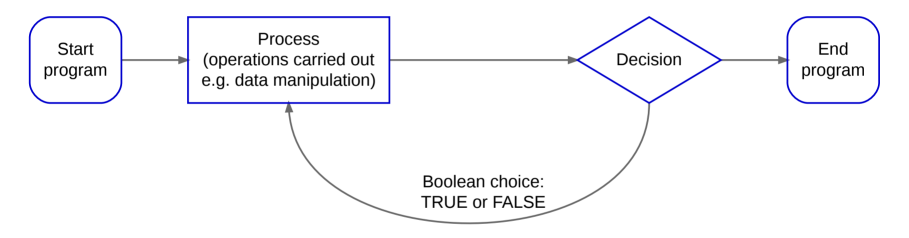
### Why is it advantageous to have structured programs?

- It decreases the complexity and time of the task at hand;
- This logical structure also means that the code has increased clarity;
- It also means that many programmers can work on one program.

This means increased productivity.

### **Control flow**

Flowcharts can be used to plan programs and represent their structure.



### Representing structure

The two basic building blocks of codes are the following:

#### Selection

Program's execution determined by statements

```
if() {}
if() {} else {}
```

### Iteration

Repetition, where the statement will **loop** until a criteria is met

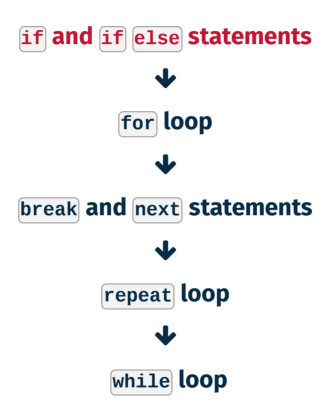
```
for() {}
while() {}
repeat {}
```

Selection and iterative statements can also be controlled by termination and jump statements:

### Termination and Jump

break next

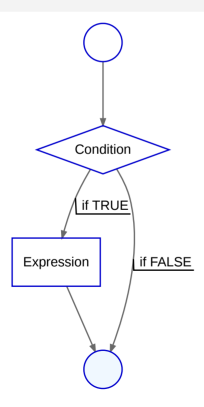
### **Control flow roadmap**



### **Decision making**

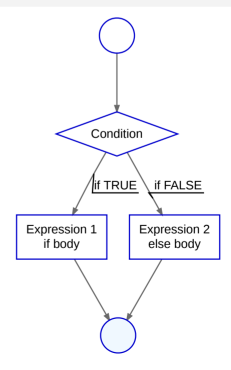
#### if() statement

```
if(condition) {
  expression
}
```



#### if() else statement

```
if(condition) {
   expression 1
} else {
   expression 2
}
```



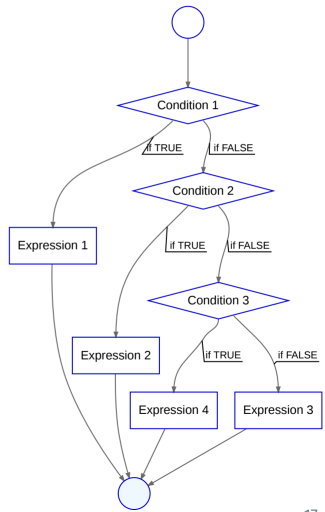
### What if you want to test more than one condition?

- if() and if() else test a single condition.
- You can also use the ifelse() function to:
  - test a vector of conditions;
  - apply a function only under certain conditions.

## Nested if() else statement

While the if() and if() else statements leave you with exactly two options, nested if() else statement allows you consider more alternatives.

```
if(test_expression1) {
  statement1
} else if(test_expression2) {
  statement2
} else if(test_expression3) {
  statement3
} else {
  statement4
}
```



## Beware of R's expression parsing!

What do you think will happen if we try the code below?

This does not work because R evaluates the first line and does not know that you are going to use an else statement

Use curly brackets (1) so that **R** knows to expect more input. Try:

```
if(2+2 == 4) {
  print("Arithmetic works.")
} else {
  print("Houston, we have a problem.")
}
# [1] "Arithmetic works."
```

### **Challenge 1**



#### Consider the following objects:

```
Paws <- "cat"
Scruffy <- "dog"
Sassy <- "cat"
animals <- c(Paws, Scruffy, Sassy)
```

- 1. Use an if() statement to print "meow" if Paws is a "cat".
- 2. Use an if() else statement to print "woof" if you supply an object that is a "dog" and "meow" if it is not. Try it out with Paws and Scruffy.
- 3. Use the <code>ifelse()</code> function to display <code>"woof"</code> for <code>animals</code> that are dogs and <code>"meow"</code> for <code>animals</code> that are cats.

### Challenge 1 - Solution



1. Use an [if()] statement to print "meow" if Paws is a "cat".

```
if(Paws == 'cat') {
   print("meow")
}
# [1] "meow"
```

2. Use an if() else statement to print "woof" if you supply an object that is a "dog" and "meow" if it is not. Try it out with Paws and Scruffy.

```
x = Paws
# x = Scruffy
if(x == 'cat') {
   print("meow")
} else {
   print("woof")
}
# [1] "meow"
```

### Challenge 1 - Solution



3. Use the <code>ifelse()</code> function to display <code>"woof"</code> for <code>animals</code> that are dogs and <code>"meow"</code> for <code>animals</code> that are cats.

```
animals <- c(Paws, Scruffy, Sassy)

ifelse(animals == 'dog', "woof", "meow")
# [1] "meow" "woof" "meow"</pre>
```

Or

```
for(val in 1:3) {
   if(animals[val] == 'cat') {
      print("meow")
   } else if(animals[val] == 'dog') {
      print("woof")
   } else print("what?")
}
# [1] "meow"
# [1] "woof"
# [1] "meow"
```

# Remember the logical operators

Command	Meaning					
==	equal to					
!=	not equal to					
<	less than					
<=	less than or equal to					
>	greater than					
>=	greater than or equal to					
x&y	x AND y					
x y	x OR y					
isTRUE(x)	test if <b>x</b> is true					

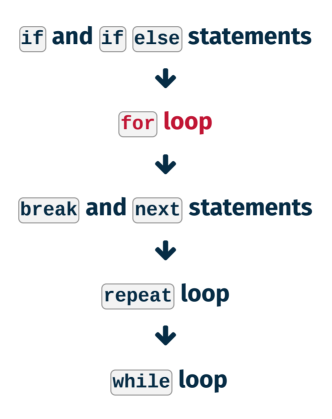
### **Iteration**

Every time some operations have to be repeated, a loop may come in handy.

#### Loops are good for:

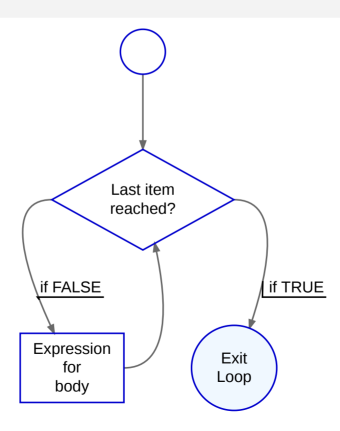
- Doing something for every element of an object;
- Doing something until the processed data runs out;
- Doing something for every file in a folder;
- Doing something that can fail, until it succeeds;
- Iterating a calculation until it converges.

### **Control flow roadmap**



A for() loop works in the following way:

```
for(i in sequence) {
  statement
  }
```



## for loop

The letter **i** can be replaced with any variable name and the sequence can be almost anything, even a list of vectors.

```
# Try the commands below and see what happens:
for(a in c("Hello", "R", "Programmers")) {
  print(a)
for(z in 1:30) {
  a < -rnorm(n = 1, mean = 5, sd = 2)
  print(a)
elements <- list(1:3, 4:10)
for(element in elements) {
  print(element)
```

In the generic example below, Rewould execute the expression 5 times, each one of them by sequentially replacing i by numbers from 1 to 5:

```
for(i in 1:5) {
  expression
}
```

**expression** can be anything:

```
print(i + 1)

vector.a[i] <- 1 + i

matrix.b[i, 1] <- matrix.a[i, 1] * 2</pre>
```

In the following example, every instance of m is being replaced by each number between 1 and 7, until it reaches the last element of the sequence:

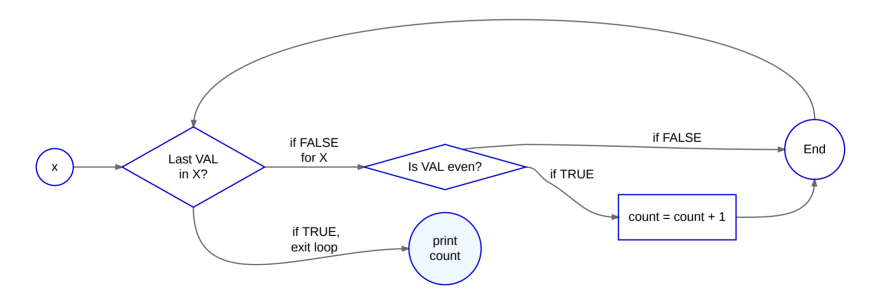
```
for(m in 1:7) {
   print(m*2)
}

# [1] 2
# [1] 4
# [1] 6
# [1] 8
# [1] 10
# [1] 12
# [1] 14
```

Let us perform operations for even elements within x using the modulo operator (%%):

```
x <- c(2, 5, 3, 9, 6)
count <- 0
```

```
for(val in x) {
   if(val %% 2 == 0) {
     count <- count + 1
   }
}
print(count)</pre>
```



**for()** loops are often used to loop over a dataset. We will use loops to perform functions on the **co2** dataset which is built in **R**.

```
data(CO2) # This loads the built in dataset

for(i in 1:length(CO2[,1])) { # for each row in the CO2 dataset
   print(CO2$conc[i]) # print the CO2 concentration
}
```

#### First 40 outputs:

# [:	1] 95	#	[1]	350	#	[1]	1000	#	[1]	250
# [:	1] 175	#	[1]	500	#	[1]	95	#	[1]	350
# [:	1] 250	#	[1]	675	#	[1]	175	#	[1]	500
# [:	1] 350	#	[1]	1000	#	[1]	250	#	[1]	675
# [:	1] 500	#	[1]	95	#	[1]	350	#	[1]	1000
# [:	1] 675	#	[1]	175	#	[1]	500	#	[1]	95
# [:	1] 1000	#	[1]	250	#	[1]	675	#	[1]	175
# [:	1] 95	#	[1]	350	#	[1]	1000	#	[1]	250
# [:	1] 175	#	[1]	500	#	[1]	95	#	[1]	350
# [:	1] 250	#	[1]	675	#	[1]	175	#	[1]	500

#### Another example:

```
for(i in 1:length(CO2[,1])) { # for each row in the CO2 dataset
  if(CO2$Type[i] == "Quebec") { # if the type is "Quebec"
    print(CO2$conc[i]) # print the CO2 concentration
    }
}
```

#### Outputs:

```
# [1] 95
                      # [1] 500
                                                # [1] 175
                                                                       # [1] 675
# [1] 175
                      # [1] 675
                                                  [1] 250
                                                                        [1] 1000
  [1] 250
                      # [1] 1000
                                                  [1] 350
                                                                        [1]
                                                                             95
  [1] 350
                      # [1] 95
                                                  [1] 500
                                                                      # [1] 175
  [1] 500
                      # [1] 175
                                                  [1] 675
                                                                       # [1] 250
  [1] 675
                                                                        [1] 350
                      # [1] 250
                                                  [1] 1000
                      # [1] 350
# [1] 1000
                                                  [1] 95
                                                                       # [1] 500
# [1] 95
                      # [1] 500
                                                  [1] 175
                                                                       # [1] 675
                      # [1] 675
                                                  [1] 250
# [1] 175
                                                                       # [1] 1000
# [1] 250
                      # [1] 1000
                                                # [1] 350
# [1] 350
                      # [1] 95
                                                # [1] 500
```

**Tip 1**. To loop over the number of rows of a data frame, we can use the function (nrow()).

```
for(i in 1:nrow(CO2)) {
    # for each row in
    # the CO2 dataset
  print(CO2$conc[i])
    # print the CO2
    # concentration
}
```

```
[1] 95
                          [1] 1000
                          [1]
                              95
  [1] 175
  [1] 250
                              175
                          [1]
  [1] 350
                          [1] 250
  [1] 500
                          [1] 350
  [1] 675
                          [1]
                              500
  [1] 1000
                          [1] 675
  [1] 95
                         [1] 1000
  [1] 175
                          [1]
                              95
  [1] 250
                          [1] 175
  [1] 350
                         [1] 250
  [1] 500
                          [1] 350
  [1] 675
                          [1] 500
  [1] 1000
                         [1] 675
  [1] 95
                          [1] 1000
  [1] 175
                          [1]
                              95
  [1] 250
                         [1]
                              175
  [1] 350
                         [1] 250
  [1] 500
                          [1] 350
# [1] 675
                       # [1] 500
```

**Tip 2**. To perform operations on the elements of one column, we can directly iterate over it.

```
for(p in CO2$conc) {
    # for each element of
    # the column "conc" of
    # the CO2 df
  print(p)
    # print the p-th element
}
```

```
[1] 95
                        [1] 1000
                        [1]
                             95
  [1] 175
  [1] 250
                         [1] 175
  [1] 350
                         [1] 250
  [1] 500
                       # [1] 350
  [1] 675
                        [1]
                             500
                         [1] 675
  [1] 1000
  [1] 95
                       # [1] 1000
  [1] 175
                        [1]
                             95
  [1] 250
                         [1] 175
  [1] 350
                        [1] 250
  [1] 500
                        [1] 350
  [1] 675
                         [1] 500
  [1] 1000
                       # [1] 675
  [1] 95
                         [1] 1000
  [1] 175
                        [1] 95
  [1] 250
                        [1]
                             175
  [1] 350
                        [1] 250
  [1] 500
                        [1] 350
# [1] 675
                       # [1] 500
```

The expression within the loop can be almost anything and is usually a compound statement containing many commands.

```
for(i in 4:5) { # for i in 4 to 5
  print(colnames(CO2)[i])
  print(mean(CO2[,i])) # print the mean of that column from the CO2 dataset
}
```

#### Output:

```
# [1] "conc"
# [1] 435
# [1] "uptake"
# [1] 27.2131
```

## for() loops within for() loops

In some cases, you may want to use nested loops to accomplish a task. When using nested loops, it is important to use different variables as counters for each of your loops. Here we used **i** and **n**:

```
for(i in 1:3) {
   for(n in 1:3) {
     print(i*n)
   }
}
```

```
# Output
# [1] 1
# [1] 2
# [1] 3
# [1] 4
# [1] 6
# [1] 3
# [1] 9
```

### Getting good: Using the apply() family

R disposes of the apply() function family, which consists of iterative functions that aim at minimizing your need to explicitly create loops.

apply() can be used to apply functions to a matrix.

### lapply()

lapply() applies a function to every element of a list.

It may be used for other objects like dataframes, lists or vectors.

The output returned is a **list** (explaining the "**l**" in **lapply**) and has the same number of elements as the object passed to it.

```
SimulatedData <- list(
   SimpleSequence = 1:4,
   Norm10 = rnorm(10),
   Norm20 = rnorm(20, 1),
   Norm100 = rnorm(100, 5)
   )

# Apply mean to each element
## of the list
lapply(SimulatedData, mean)</pre>
```

```
# $SimpleSequence
# [1] 2.5
#
# $Norm10
# [1] -0.1173473
#
# $Norm20
# [1] 0.6979582
#
# $Norm100
# [1] 4.823129
```

### sapply()

sapply() is a 'wrapper' function for lapply(), but returns a simplified output as
a vector, instead of a list.

#### mapply()

mapply() works as a multivariate version of sapply().

It will apply a given function to the first element of each argument first, followed by the second element, and so on. For example:

```
lilySeeds <- c(80, 65, 89, 23, 21)
poppySeeds <- c(20, 35, 11, 77, 79)

# Output
mapply(sum, lilySeeds, poppySeeds)
# [1] 100 100 100 100 100
```

#### tapply()

tapply() is used to apply a function over subsets of a vector.

It is primarily used when the dataset contains different groups (*i.e.* levels or factors), and we want to apply a function to each of these groups.

```
mtcars[1:10, c("hp", "cyl")]
#
                hp cyl
# Mazda RX4 110 6
# Mazda RX4 Wag 110 6
# Datsun 710
          93 4
# Hornet 4 Drive 110 6
# Hornet Sportabout 175 8
# Valiant
                105 6
# Duster 360 245 8
               62
# Merc 240D
                    4
# Merc 230
            95
                     4
# Merc 280
                123
                     6
```

#### Challenge 2



After coming back from the field, you have realized that your tool for measuring  $\mathbf{CO_2}$  uptake was not calibrated properly at Quebec sites and all measurements are 2 units higher than they should be.

Now, you must do the following:

- 1. Use a loop to correct these measurements for all Quebec sites;
- 2. Use an  $\overline{\text{apply()}}$  family function to calculate the average  $CO_2$  uptake in both Québec and Mississipi sampled sites.

For this, you must load the  $CO_2$  dataset using  $(CO_2)$ , and then use the object  $(CO_2)$ .

Get your hands dirty!

#### Challenge 2: Solution



1. Using for() and if() to correct the measurements:

```
for(i in 1:dim(CO2)[1]) {
   if(CO2$Type[i] == "Quebec") {
      CO2$uptake[i] <- CO2$uptake[i] - 2
   }
}</pre>
```

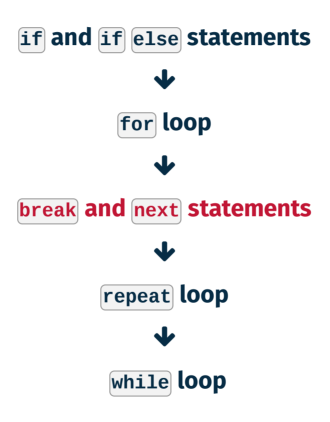
2.Using **tapply()** to calculate the mean for each group:

```
tapply(CO2$uptake, CO2$Type, mean)
# Quebec Mississippi
# 31.54286 20.88333
```

#### **Modifying iterations**

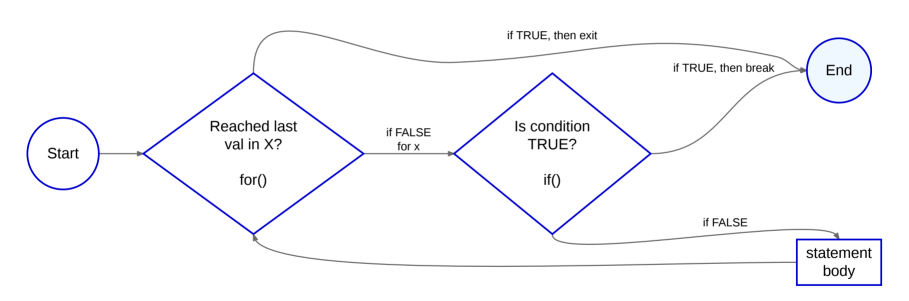
- Normally, loops iterate over and over until they finish.
- Sometimes you may be interested in breaking this behaviour.
- For example, you may want to tell **R** to stop executing the iteration when it reaches a given element or condition.
- You may also want **R** to jump certain elements when certain conditions are met.
- For this, we will introduce break, next and while.

#### **Control flow roadmap**



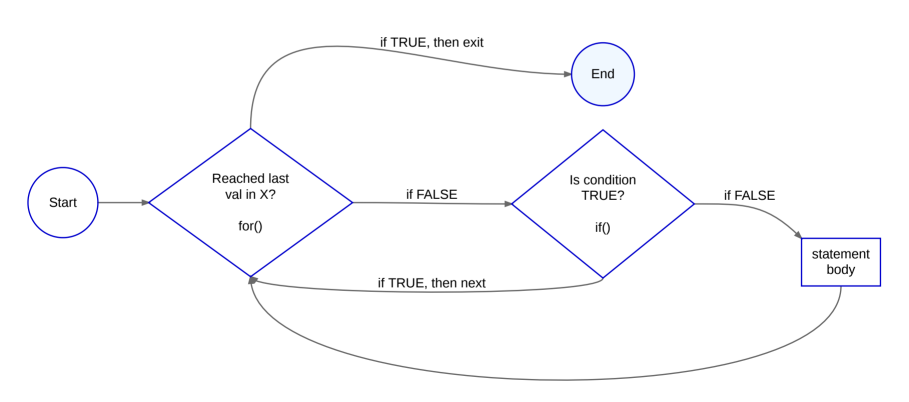
# Modifying iterations: break statement

```
for(val in x) {
  if(condition) { break }
  statement
}
```



# Modifying iterations: next statement

```
for(val in x) {
  if(condition) { next }
  statement
}
```



## Modifying iterations: next statement

Print the  $CO_2$  concentrations for "chilled" treatments and keep count of how many replications were done.

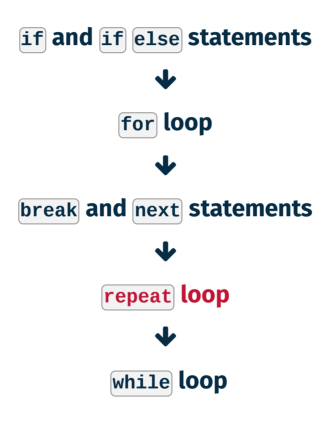
```
count <- 0

for(i in 1:nrow(CO2)) {
    if(CO2$Treatment[i] == "nonchilled") next
    # Skip to next iteration if treatment is nonchilled
    count <- count + 1
# print(CO2$conc[i]) # You can turn this on if you want to
}
print(count) # The count and print command were performed 42 times.

# [1] 42

sum(CO2$Treatment == "nonchilled")
# [1] 42</pre>
```

#### **Control flow roadmap**

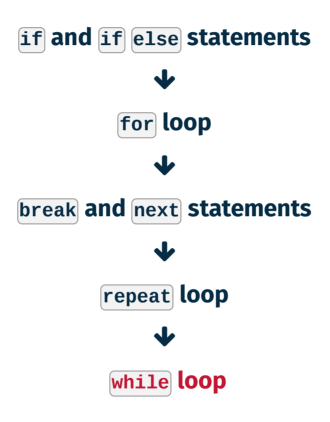


#### Modifying iterations: repeat loop

This could be equivalently written using a repeat loop and break:

```
count <- 0
i <- 0
repeat {
    i <- i + 1
        if(C02$Treatment[i] == "nonchilled") next # skip this loop
        count <- count + 1
        print(C02$conc[i])
        if(i == nrow(C02)) break # stop looping
    }
print(count)</pre>
```

#### **Control flow roadmap**



#### Modifying iterations: while loop

This could also be written using a while loop:

```
i <- 0
count <- 0
while(i < nrow(CO2))
{
   i <- i + 1
   if(CO2$Treatment[i] == "nonchilled") next # skip this loop
   count <- count + 1
   print(CO2$conc[i])
}
print(count)</pre>
```

#### **Challenge 3**



You have realized that another of your tools was not working properly!

At **Mississippi** sites, **concentrations** less than 300 were measured correctly, but concentrations equal or higher than 300 were overestimated by 20 units!

Your *mission* is to use a loop to correct these measurements for all Mississippi sites.

**Tip**. Make sure you reload the  $CO_2$  data so that we are working with the raw data for the rest of the exercise:

data(CO2)

#### Challenge 3: Solution



```
for(i in 1:nrow(CO2)) {
   if(CO2$Type[i] == "Mississippi") {
      if(CO2$conc[i] < 300) next
      CO2$conc[i] <- CO2$conc[i] - 20
   }
}</pre>
```

Note: We could also have written it in this way, which is more concise and clearer:

```
for(i in 1:nrow(CO2)) {
   if(CO2$Type[i] == "Mississippi" && CO2$conc[i] >= 300) {
      CO2$conc[i] <- CO2$conc[i] - 20
   }
}</pre>
```

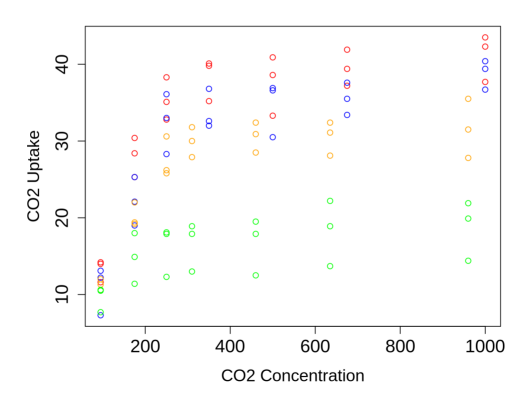
## Edit a plot using for() and if()

Let us plot **uptake** *versus* **concentration** with points of different colors according to their type (*Québec* or *Mississippi*) and **treatment** (*chilled* or *nonchilled*):

```
plot(x = CO2\$conc,
     y = C02$uptake,
     type = "n", # points
     cex.lab = 1.4, cex.axis = 1.5,
     cex.main = 1.5, cex.sub = 1.5,
     xlab = "CO2 concentration",
     ylab = "CO2 uptake")
for(i in 1:length(CO2[, 1])) {
  if(CO2$Type[i] == "Quebec" & CO2$Treatment[i] == "nonchilled") {
    points(CO2$conc[i], CO2$uptake[i], col = "red")
  if(CO2$Type[i] == "Quebec" & CO2$Treatment[i] == "chilled") {
    points(CO2$conc[i], CO2$uptake[i], col = "blue")
  }
  if(CO2$Type[i] == "Mississippi" & CO2$Treatment[i] == "nonchilled") {
    points(CO2$conc[i], CO2$uptake[i], col = "orange")
  }
  if(CO2$Type[i] == "Mississippi" & CO2$Treatment[i] == "chilled") {
    points(CO2$conc[i], CO2$uptake[i], col = "green")
```

## Edit a plot using for() and if()

Plotting **uptake** *versus* **concentration** with points of different colors according to their type using **for()** loop and **if()**.



#### Challenge 4



Generate a plot showing **concentration** *versus* **uptake** where each plant is shown using a point with a different colour.

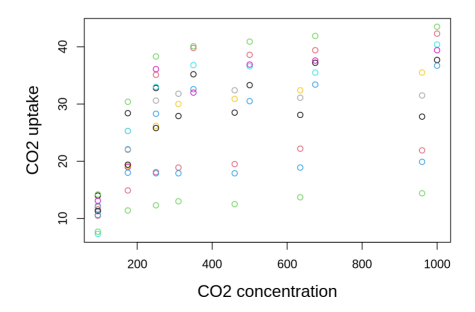
**Bonus points** for doing it with nested loops!

#### Steps:

- 1. Create an empty plot;
- 2. Create a list of plants (hint: ?unique);
- 3. Fill the plot using [for()] and [if()] statements.

#### **Challenge 4: Solution**





# **Writing functions**

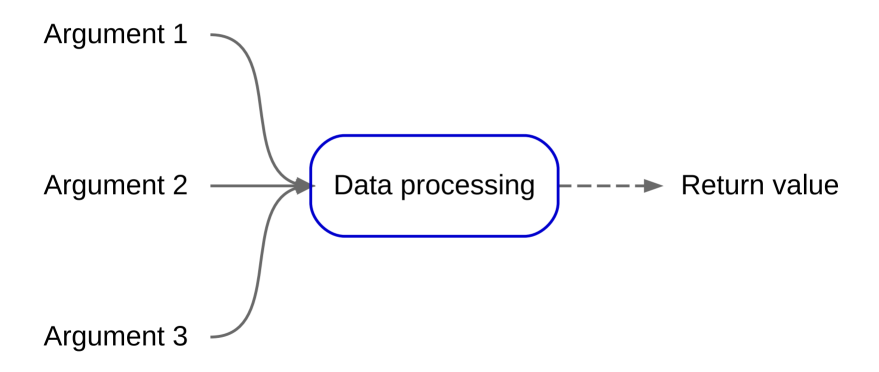
#### Why write functions?

Much of the heavy lifting in R is done by functions.

#### They are useful for:

- 1. Performing a task repeatedly, but configurably;
- 2. Making code more readable;
- 3. Making code easier to modify and maintain;
- 4. Sharing code between different analyses;
- 5. Sharing code with other people;
- 6. Modifying R's built-in functionality.

#### What is a function?



#### **Syntax of a function**

```
function_name <- function(argument1, argument2, ...) {
  expression... # What we want the function to do
  return(value) # Optional
}</pre>
```

#### **Arguments of a function**

```
function_name <- function(argument1, argument2, ...) {
  expression...
  return(value)
}</pre>
```

Arguments are the entry values of your function and will have the information your function needs to be able to perform correctly.

A function can have between 0 and an infinity of arguments. See the following example:

```
operations <- function(number1, number2, number3) {
  result <- (number1 + number2) * number3
  print(result)
}
operations(1, 2, 3)</pre>
```

# [1] 9

#### **Challenge 5**



Using what you learned previously on flow control, create a function print\_animal() that takes an animal as argument and gives the following results:

```
Scruffy <- "dog"
Paws <- "cat"

print_animal(Scruffy)
# [1] "woof"

print_animal(Paws)
# [1] "meow"</pre>
```

#### **Challenge 5: Solution**



Using what you learned previously on flow control, create a function <a href="print\_animal">print\_animal</a> that takes an <a href="mailto:animal">animal</a> as argument and gives the following results:

```
Scruffy <- "dog"
Paws <- "cat"
print_animal(Scruffy)
# [1] "woof"
print_animal(Paws)
# [1] "meow"
print_animal <- function(animal) {</pre>
  if(animal == "dog") {
    print("woof")
  } else if(animal == "cat") {
    print("meow")
```

#### Default values in a function

Arguments can also be optional and be provided with a **default value**.

This is useful when using a function with the same settings, but still provides the flexibility to change its values, if needed.

```
operations <- function(number1, number2, number3 = 3) {
  result <- (number1 + number2) * number3
  print(result)
}

operations(1, 2, 3) # is equivalent to
# [1] 9
operations(1, 2)
# [1] 9
operations(1, 2, 2) # we can still change the value of number3 if needed
# [1] 6</pre>
```

### **Argument** ....

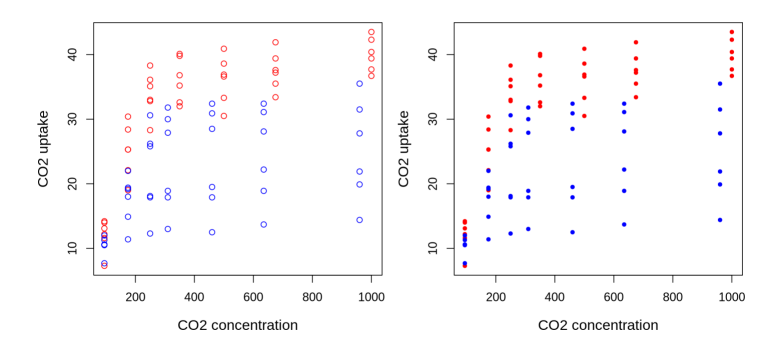
The special argument ... allows you to pass on arguments to another function used inside your function. Here we use ... to pass on arguments to plot() and points().

```
plot.C02 <- function(C02, ...) {
  plot(x=C02\$conc, y=C02\$uptake, type="n", ...)
 for(i in 1:length(CO2[,1])){
     if(CO2$Type[i] == "Ouebec") {
       points(CO2$conc[i], CO2$uptake[i],
              col = "red", type = "p", ...)
     } else if(CO2$Type[i] == "Mississippi") {
       points(CO2$conc[i], CO2$uptake[i],
              col = "blue", type = "p", ...)
plot.C02(C02,
         cex.lab=1.2,
         xlab="CO2 concentration",
         vlab="CO2 uptake")
```

plot.C02(C02,

### **Argument** ....

The special argument ... allows you to pass on arguments to another function used inside your function. Here we use ... to pass on arguments to plot() and points().



## **Argument** ....

The special argument ... allows you to borrow arguments from other functions, *i.e.* allowing for an indefinite number of arguments to be inputed.

```
sum2 <- function(...) {</pre>
  args <- list(...)</pre>
  result <- 0
  for(i in args) {
    result <- result + i
  return(result)
sum2(2, 3)
# [1] 5
sum2(2, 4, 5, 7688, 1)
# [1] 7700
```

#### **Return values**

The last expression evaluated in a **function** becomes the return value.

```
myfun <- function(x) {
   if(x < 10) {
      0
    } else {
      10
   }
}
myfun(5)
# [1] 0
myfun(15)
# [1] 10</pre>
```

function() itself returns the last evaluated value even without including return() function.

#### **Return values**

It can be useful to explicitly **return()** if the routine should end early, jump out of the function and return a value.

```
simplefun1 <- function(x) {
  if(x<0)
  return(x)
}</pre>
```

Functions can return only a single object (and text). But this is not a limitation because you can return a **list** containing any number of objects.

```
# $result
# [1] 3

simplefun2(1, 2)

# $x

# [1] 1

# $y

# [1] 2
```

#### Challenge 6



Using what you have just learned on functions and control flow, create a function named **bigsum** that takes two arguments **a** and **b** and:

- 1. Returns 0 if the sum of **a** and **b** is strictly less than 50;
- 2. Else, returns the sum of **a** and **b**.

#### **Challenge 6: Solution**



Using what you have just learned on functions and control flow, create a function named **bigsum** that takes two arguments **a** and **b** and:

- 1. Returns **o** if the sum of **a** and **b** is strictly less than **50**;
- 2. Else, returns the sum of **a** and **b**.

#### **Answer 1**

```
bigsum <- function(a, b) {
   result <- a + b
   if(result < 50) {
      return(0)
   } else {
      return(result)
   }
}</pre>
```

#### **Answer 2**

```
bigsum <- function(a, b) {
  result <- a + b
  if(result < 50) {
    0
  } else {
    result
  }
}</pre>
```

## Accessibility of variables

It is essential to always keep in mind where your variables are, and whether they are defined and accessible:

- → Variables defined **inside** a function are not accessible outside of it!
- → Variables defined **outside** a function are accessible inside. But it is NEVER a good idea, as your function will not function if the outside variable is erased.

#### **Accessibility of variables**

```
var1 <- 3  # var1 is defined outside our function</pre>
vartest <- function() {</pre>
  a <- 4 # 'a' is defined inside
  print(a) # print 'a'
  print(var1) # print var1
            # we cannot print 'a' as it exists only inside the function
a
  [1] -4 -3 -2 -1 0 1 2 3
# [9] 4 5
vartest() # calling vartest() will print a and var1
# [1] 4
# [1] 3
rm(var1) # remove var1
vartest() # calling the function again does not work anymore
# [1] 4
# Error in print(var1): object 'var1' not found
```

## Accessibility of variables

#### Mandatory tip. Use arguments then!

Also, inside a function, arguments names will take over other variable names.

```
var1 <- 3  # var1 is defined outside our function

vartest <- function(var1) {
   print(var1) # print var1
}

vartest(8)  # Inside our function var1 is now our argument
# [1] 8  # and takes its value

var1  # var1 still has the same value
# [1] 3</pre>
```

#### **Accessibility of variables**

**Tip.** Be very careful when creating variables inside a conditional statement as the variable may never have been created and cause (sometimes imperceptible) errors.

**Tip.** It is good practice to define variables outside the conditions and then modify their values to avoid any problem

```
a <- 3
if(a > 5) {
   b <- 2
}
a + b</pre>
```

```
# Error: object 'b' not found
```

If you had **b** already assigned in your environment, with a different value, you could have had a **bigger** problem!

No error would have been shown and **a** + **b** would have meant another thing!

# Good programming practices

## Why should I care about programming practices?

- To make your life easier;
- To achieve greater readability and makes sharing and reusing your code a lot less painful;
- To reduce the time you will spend to understand your code.

Pay attention to the next tips!

Proper indentation and spacing is the first step to get an easy to read code:

- Use **spaces** between and after your operators;
- Use consistently the same assignation operator.
  - <- is often preferred. = is sometimes OK, but do not switch all the time between the two;
- Use brackets when using flow control statements:
  - Inside brackets, indent by at least two spaces;
  - Put closing brackets on a separate line, except when preceding an else statement.
- Define each variable on its own line.
- Use **Cmd** + **I** or **Ctrl** + **I** in RStudio to indent the highlighted code automatically.

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- Use brackets when using flow control statements:
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- Use **Cmd** + **I** or **Ctrl** + **I** in RStudio to indent the highlighted code automatically.

#### Nay!

# if((a[x,y]>1.0)&(a[x,y]<2.0)){ print("Between 1 and 2") }</pre>

#### Yay!

```
if((a[x, y] > 1.0) & (a[x, y] < 2.0))
  print("Between 1 and 2")
}</pre>
```

On the left, code is not spaced. All brackets are in the same line, and it looks "messy".

```
a<-4; b=3
if(a<b){
if(a==0)print("a zero")}else{
if(b==0){print("b zero")}else print(b)]</pre>
```

On the left, code is not spaced. All brackets are in the same line, and it looks "messy". On the right, it looks more organized, no?

```
a<-4;b=3
if(a<b){
if(a==0)print("a zero")}else{
if(b==0){print("b zero")}else print(b)]</pre>
```

```
a <- 4
b <- 3
if(a < b) {
   if(a == 0) {
      print("a zero")
   }
} else {
   if(b == 0) {
      print("b zero")
   } else {
      print(b)
   }
}</pre>
```

## Use functions to simplify your code

#### Write your own function:

- 1. When portion of the code is repeated more than twice in your script;
- 2. If only a part of the code changes and includes options for different arguments.

This would also reduce the number of potential errors done by copy-pasting, and the time needed to correct them.

## Use functions to simplify your code

Let's modify the example from **Challenge #3** and suppose that all  $CO_2$  uptake from Mississipi plants was overestimated by 20 and Quebec underestimated by 50.

We could write this:

```
for(i in 1:length(CO2[,1])) {
   if(CO2$Type[i] == "Mississippi") {
      CO2$conc[i] <- CO2$conc[i] - 20
   }
}
for(i in 1:length(CO2[,1])) {
   if(CO2$Type[i] == "Quebec") {
      CO2$conc[i] <- CO2$conc[i] + 50
   }
}</pre>
```

#### Or this:

```
recalibrate <- function(CO2, type, bias
for(i in 1:nrow(CO2)) {
   if(CO2$Type[i] == type) {
      CO2$conc[i] <- CO2$conc[i] + bias
   }
   }
   return(CO2)
}

newCO2 <- recalibrate(CO2 = CO2, type = newCO2 <- recalibrate(newCO2, "Quebec"</pre>
```

#### Use meaningful names for functions

Same function as before, but with vague names:

```
rc <- function(c, t, b) {
   for(i in 1:nrow(c)) {
     if(c$Type[i] == t) {
      c$uptake[i] <- c$uptake[i] + b
     }
   }
  return(c)
}</pre>
```

What is c and rc?

Whenever possible, avoid using names of existing  $\mathbb{R}$  functions and variables to avoid confusion and conflits.

#### **Use comments**

**Final tip**. Add comment to describe what your code does, how to use its arguments or a detailed step-by-step description of the function.

```
# Recalibrates the CO2 dataset by modifying the CO2 uptake concentration
# by a fixed amount depending on the region of sampling.
# Arguments
# CO2: the CO2 dataset
# type: the type ("Mississippi" or "Quebec") that need to be recalibrated
# bias: the amount to add or remove to the concentration uptake
recalibrate <- function(CO2, type, bias) {</pre>
  for(i in 1:nrow(CO2)) {
    if(CO2$Type[i] == type) {
      CO2$uptake[i] <- CO2$uptake[i] + bias</pre>
  return(CO2)
```

## **Group** exercise

Using what you learned, write an if() statement that tests whether a numeric variable x is zero. If not, it assigns cos(x)/x to z, otherwise it assigns 1 to z.

Create a function called  $my\_function()$  that takes the variable x as argument and returns z.

If we assign 45, 20, and 0 to  $\boxed{x}$  respectively, which of the following options would represent the results?

- **1.** 0.54, 0.12, and 0;
- **2.** 0.20, 0.54, and 1;
- **3.** 0.12, 0.20, and 1.

## **Group exercise: Solution**



Correct answer is option **3** (\$0.12\$, **0.20**, and **1**).

```
my_function <- function(x) {</pre>
  if(x != 0) {
  z < - \cos(x)/x
  } else { z <- 1 }</pre>
  return(z)
my_function(45)
# [1] 0.01167382
my_function(20)
# [1] 0.0204041
my_function(0)
# [1] 1
```

## Thank you for attending this workshop!

