# Introduction to Programming with R - June 15

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# Taking a random sample from a vector

The sample function is a very useful function for drawing random samples from vectors either with or without replacement.

If it is called with just a vector, it returns a permuted form of that vector:

```
sample(1:10) Size not specified, defaults to range from 1st argument
```

```
[1] 8 6 9 7 2 5 1 3 10 4
```

A smaller vector can be created by specifying the size argument:

```
sample(1:10, size = 5)
```

```
[1] 10 9 5 4 8
```

In this case, 5 unique values are returned. If you want to sample with replacement, specify replace = TRUE:

```
sample(1:10, size = 5, replace = TRUE)
```

```
[1] 8 7 7 8 10
```

If a larger vector is to be sampled, then replace has to be TRUE:

```
sample(1:10, size = 100, replace = TRUE)
```

```
2
                  7
                    1 6 8 7
                               3 3
                                    4
                                       4 2
                                            8
                                              9 10 10
                    2
                       6
                         5 1
                              4 10
                                      4 1
                                            9 5
          4 1 10
                  5
                                    6
                         4 6
                    5
                       6
                                 7
                                    9
                                       9
                                         8
                         5 10
                                       2
[70]
          9 10
               1
                  7
                    4
                       1
                               2
                                 9
                                    9
       5 7 9 2 7 10
```

By default, all elements in the vector have the same likelihood of being sampled. Weights can be applied by specifying them in the prob argument:

```
sample(letters[1:5], size = 20, replace = TRUE, prob = c(10, 10, 1, 1, 0))
```

#### Discrete values

The function unique() will list the unique values in a vector in the order it finds them:

```
x <- sample(letters, 10, replace = TRUE)
x</pre>
```

```
[1] "t" "d" "x" "f" "e" "s" "c" "e" "i" "e"
unique(x)
```

```
[1] "t" "d" "x" "f" "e" "s" "c" "i"
```

The function duplicated() will identify those elements in a vector that occur at an earlier position:

```
duplicated(x)
 [1] FALSE FALSE FALSE FALSE FALSE FALSE TRUE FALSE
# the negation of duplicated is the same as unique
x[!duplicated(x)]
[1] "t" "d" "x" "f" "e" "s" "c" "i"
unique(x)
[1] "t" "d" "x" "f" "e" "s" "c" "i"
To calculate the frequency of values in a vector (the number of occurences), use table():
x <- sample(letters, 20, replace = TRUE)
table(x)
x
bcdeflnopqrwx
1 1 3 2 1 1 2 2 1 1 2 2 1
table can be used for cross-tabulation as well - counting frequency of occurrence of a combination of categories
months <- sample(month.abb, 20, replace = TRUE)</pre>
sex <- sample(c("m", "f"), 20, replace = TRUE)</pre>
freq <- table(sex, months)</pre>
freq
   months
sex Apr Aug Feb Jan Jun Mar May Nov Oct Sep
  f
      0
          3
                   1
                       1
                            3
                                0
                                    3
               0
                   0
                       0
                            0
                                1
                                    1
The values in a table can be accessed like a vector or matrix
freq["m", ]
Apr Aug Feb Jan Jun Mar May Nov Oct Sep
              0
                       0
freq["f", c("Jan", "Feb", "Mar")]
Jan Feb Mar
  1
      1
```

# Data selection and manipulation

Х

To identify values of one vector that are within another one, use %in%:

```
letters %in% c("a", "f", "G", "b")

[1] TRUE TRUE FALSE FALSE FALSE TRUE FALSE FALSE FALSE FALSE FALSE
[12] FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE FALSE
[23] FALSE FALSE FALSE
To identify values of a logical vector that are TRUE, use which:

x <- sample(1:100, 20)
```

```
64 90 100 66 35 36 22 52 78 63 83 58 25 93 20 17 68
[18]
     79 67 41
which(x < 50)
[1] 5 6 7 13 15 16 20
To identify the minimum and maximum values, use which.min and which.max:
which.min(x)
[1] 16
which.max(x)
[1] 3
To see if any values meet a condition, use any:
any(x < 50)
[1] TRUE
any(x > 200)
[1] FALSE
To see if all values meet a condition, use all:
all(x < 50)
[1] FALSE
all(x < 200)
[1] TRUE
Vectors can be reversed with rev:
x \leftarrow sample(1:5, 10, replace = T)
 [1] 3 4 2 5 4 3 2 1 5 5
rev(x)
 [1] 5 5 1 2 3 4 5 2 4 3
and sorted with sort:
sort(x)
 [1] 1 2 2 3 3 4 4 5 5 5
# in decreasing order
sort(x, decreasing = TRUE)
 [1] 5 5 5 4 4 3 3 2 2 1
However, sort can't be applied to a matrix or data.frame to sort the rows. For that, you need order. order
returns a vector of indices in the order they should be as if they were sorted:
```

x <- data.frame(
v1 = sample(letters, 20, replace = TRUE),
v2 = sample(letters, 20, replace = TRUE),</pre>

v3 = sample(letters, 20, replace = TRUE)

```
)
х
  v1 v2 v3
1 kgi
2 s b v
3 kak
4
  s r y
5 g v e
6 h b x
7 q o b
8 b f y
9 q z s
10 g u r
11 h x w
12 t u u
13 s h h
14 u p n
15 n w q
16 r b x
17 d n z
18 r r x
19 a d b
20 f k f
x.ord <- order(x$v1)</pre>
x[x.ord, ]
  v1 v2 v3
19 a d b
8 b f y
17 d n z
20 f k f
5 g v e
10 g u r
6 h b x
11 h x w
1 k g i
3 kak
15 n w q
7 q o b
9 q z s
16 r b x
18 r r x
2 s b v
4 s r y
13 s h h
12 t u u
14 u p n
# also in decreasing order
x[order(x$v1, decreasing = TRUE), ]
  v1 v2 v3
14 u p n
```

```
12 t u u
2
   s
     b v
   s r y
13 s h h
16 r b x
18 r r x
   q o b
7
9
   q
     z
        s
15 n
     W
        q
1
   k
     g
3 \quad k \quad a \quad k
6
  h b x
11 h x w
5
     v
   g
        е
10 g u r
20 f
     k f
17 d n z
   b f y
8
19 a d b
```

order can take several vectors to do hierarchical sorting.

```
i <- order(x$v2, x$v1, x$v3)
i</pre>
```

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

# x[i,]

```
v1 v2 v3
  k a k
6
  h b x
16 r b x
2
   s b v
19 a
     d b
   b f y
8
   k g
1
13 s h h
20 f k f
17 d n z
   q o
       b
14 u
     p n
18
  r
     r
       Х
4
   s r y
10
   g
     u r
12 t
     u
       u
5
   g
     V
       е
15 n w
       q
11 h x w
   q z s
```

# Character and string manipulation

#### nchar

A character vector is a vector where every element is a character string of any length. The length() of a character vector is the number of elements in it:

```
x <- c("This is a sentence", "Hello World!", "This is the third element")
length(x)</pre>
```

[1] 3

To get the number of characters in each element, use nchar():

```
\mathtt{nchar}(\mathtt{x})
```

[1] 18 12 25

#### substr

Strings can be extracted from elements using substr(). You specify the first and last characters to be extracted from each string:

```
# get the first three characters from every string
substr(x, 1, 3)
```

```
[1] "Thi" "Hel" "Thi"
```

```
# get the 3rd character from every string
substr(x, 3, 3)
```

```
[1] "i" "l" "i"
```

substr can also be used to replace values within strings by assigning:

```
substr(x, 1, 4) <- "That"
x</pre>
```

- [1] "That is a sentence" "Thato World!"
- [3] "That is the third element"

# strsplit

Strings can be split based on some common delimiter using strsplit():

```
# split based on spaces
x.split <- strsplit(x, " ")
x.split

[[1]]
[1] "That" "is" "a" "sentence"</pre>
```

```
[[2]]
```

[1] "Thato" "World!"

[[3]]
[1] "That" "is" "the" "third" "element"

```
str(x.split)
```

```
List of 3
$ : chr [1:4] "That" "is" "a" "sentence"
$ : chr [1:2] "Thato" "World!"
$ : chr [1:5] "That" "is" "the" "third" ...
```

Note that the return value from strsplit is a list. Each element in the list corresponds to a vector resulting from splitting every element in the original vector

```
x.split[[1]]
```

```
[1] "That" "is" "a" "sentence"
```

#### paste

To create strings from combinations of strings (or numbers) we use paste(). This function takes a set of vectors, and pastes the elements together using recycling:

```
# vectors are equal length
paste(letters[1:6], 1:6)
```

```
[1] "a 1" "b 2" "c 3" "d 4" "e 5" "f 6"

# one vector is a multiple of the other
paste(letters[1:6], 1:2)
```

```
[1] "a 1" "b 2" "c 1" "d 2" "e 1" "f 2"

# one vector is not a multiple of the other

paste(letters[1:6], 1:4)
```

```
[1] "a 1" "b 2" "c 3" "d 4" "e 1" "f 2"
```

The argument sep determines what character is used as a separator between the characters:

```
paste(letters[1:6], 1:2, sep = "-")
```

```
[1] "a-1" "b-2" "c-1" "d-2" "e-1" "f-2"
```

If you do not want a separator character, either set sep = "" or use pasteO():

```
paste0(letters[1:6], 1:2)
```

```
[1] "a1" "b2" "c1" "d2" "e1" "f2"
```

If you want to paste all of the arguments to create a single element vector, set the collapse argument:

```
paste(letters[1:6], 1:2, sep = "-", collapse = "#")
```

```
[1] "a-1#b-2#c-1#d-2#e-1#f-2"
```

### tolower, toupper

Character case can be changed with tolower and toupper:

```
tolower(x)
```

```
[1] "that is a sentence" "thato world!"
```

<sup>[3] &</sup>quot;that is the third element"

# 

# Regular Expressions

[3] "THAT IS THE THIRD ELEMENT"

For finer control on searching and replacing text within strings, you will have to turn to "regular expressions", which is a kind of syntax of its own and is common across several platforms. The help page for regular expressions in R is ?regex. The functions that are most commonly used with regular expressions are given in grep. The most commonly used on this page are:

grep and grepl: Identify elements that have the sought after pattern sub and gsub: Replace a desired pattern with other text

# apply Functions

Many times, we want to execute the same function on sequential elements of some object. This could be things like the elements of a vector or list, the rows of a matrix, or the columns of a data frame. For these, R provides a family of functions that usually end in -apply or are based on them.

### lapply

The most basic of these functions is lapply. The "l" refers to the fact that lapply will always return a list. There are two main arguments to lapply: the first is the object to be iterated over, and the second is a function that takes sequential elements of that object. As an example, let's use the sample function. Recall that if you execute sample with a single integer(n), it will return a permutation of the vector 1:n:

```
sample(5)
[1] 3 5 2 4 1
sample(10)
[1] 6 2 7 4 5 1 9 10 3 8
Here is a list resulting from calls to sample with the elements of the vector 1:5:
x <- lapply(1:5, sample)
str(x)</pre>
```

```
List of 5
 $ : int 1
$ : int [1:2] 1 2
$ : int [1:3] 3 2 1
$ : int [1:4] 3 4 2 1
$ : int [1:5] 1 4 3 5 2
[[1]]
[1] 1
[[2]]
[1] 1 2
[[3]]
[1] 3 2 1
[[4]]
[1] 3 4 2 1
[[5]]
[1] 1 4 3 5 2
Note that the result is a list, the elements of which are the result of calls to sample(1), sample(2), sample(3),
etc. The elements of the return value are in the same order as the original object being iterated over:
lapply(c(5, 3, 1, 8), sample)
[[1]]
[1] 1 4 5 2 3
[[2]]
[1] 1 3 2
[[3]]
[1] 1
[[4]]
[1] 4 2 7 6 3 5 8 1
The first argument can be a list too:
lapply(x, sum)
[[1]]
[1] 1
[[2]]
[1] 3
[[3]]
[1] 6
[[4]]
[1] 10
```

```
[[5]]
[1] 15
```

### sapply

If the return value from every iteration was the same length, you may want to simplify the result. This is what sapply is for. If every call to the function returns a scalar, then sapply will return a vector. If every call to the function returns a vector of equal length, then sapply will return a matrix. If every call to the function returns a value of different lengths, then sapply defaults to returning a list:

```
# every return value from sum is a scalar - sapply returns a vector
sapply(x, sum)
[1] 1 3 6 10 15
# every return value from sample is a 5 element vector - sapply returns a matrix
sapply(rep(5, 8), sample)
     [,1] [,2] [,3] [,4] [,5] [,6] [,7] [,8]
[1,]
             3
                   2
                        1
                              1
                                   4
                                         3
                                              2
[2,]
        4
             2
                   1
                        3
                              5
                                   2
                                         5
                                              1
[3,]
        2
             4
                   4
                        2
                              2
                                   5
                                         4
                                              3
[4,]
                        4
                              3
        1
             5
                   5
                                   3
                                              4
                                         1
                   3
                        5
[5,]
        5
                              4
                                   1
                                         2
                                              5
              1
# this is the same as our lapply example - sapply returns a list
sapply(c(5, 3, 1, 8), sample)
[[1]]
[1] 4 1 5 2 3
[[2]]
[1] 1 2 3
[[3]]
[1] 1
[[4]]
[1] 8 3 2 7 6 4 5 1
Arguments to the function can be specified in the lapply or sapply call:
```

```
sapply(c(5, 3, 1, 8), sample, size = 5, replace = TRUE)
```

```
[,1] [,2] [,3] [,4]
[1,]
               2
                     1
                           6
         1
[2,]
                           7
         4
                     1
               1
[3,]
         1
               3
                     1
                           4
[4,]
                     1
                           8
         4
               1
[5,]
         4
               2
                     1
                           4
```

# apply

If you are dealing with a multi-dimensional object (matrix, array, or data frame) and you want to apply a function to a given dimension (i.e, each row or each column), use apply. You have to specify the dimension that you will be iterating over as the second argument (1 = rows, 2 = columns, etc). apply will try to simplify the results like sapply:

```
x <- matrix(sample(1:100, 24, replace = TRUE), nrow = 4)
     [,1] [,2] [,3] [,4] [,5] [,6]
             81
                             90
[1,]
       27
                  83
                        65
                                   28
[2,]
        8
             51
                  21
                         1
                             20
                                    8
                             72
[3,]
        1
             43
                  67
                        85
                                   78
[4,]
     100
             15
                  83
                             96
                                   17
# median of each row
apply(x, 1, median)
[1] 73.0 14.0 69.5 50.0
# difference of each column
apply(x, 2, diff)
     [,1] [,2] [,3] [,4] [,5]
[1,]
      -19
            -30
                 -62
                       -64
                            -70
                                  -20
[2,]
       -7
             -8
                  46
                        84
                             52
                                   70
[3,]
       99
            -28
                  16
                       -71
                             24
                                  -61
```

### tapply

If you want to execute a function on groups of values, tapply can often be a good choice. The arguments are a vector that will be summarized, another vector or set of vectors that represent identify elements to groups, and the function that will get the sequential subsets of the original vector. As a simple example, we calculate the means of subsets of a random vector of numbers

```
x <- sample(1:100, 100, replace = TRUE)
grp <- sample(letters[1:5], 100, replace = TRUE)
tapply(x, grp, mean)</pre>
```

```
a b c d e
49.94444 55.76190 61.09091 48.27273 50.29412
```

As a more practical example, we can calculate the average temperature at each station in our ctd dataset:

```
ctd <- read.csv("ctd.csv")
tapply(ctd$temp, ctd$station, mean)</pre>
```

```
Station.1 Station.10 Station.11 Station.12 Station.13 Station.14
  13.56772
             14.57675
                        14.93466
                                    14.40093
                                               13.84903
                                                          14.42424
Station.15 Station.16 Station.17 Station.18 Station.19
                                                         Station.2
             14.42280
                                               16.32791
  14.10233
                        14.27389
                                   14.57620
                                                          14.24618
Station.20 Station.21 Station.22 Station.23 Station.24 Station.25
  13.34375
             13.77648
                        14.18930
                                    14.60143
                                               16.35819
                                                          15.82101
Station.26 Station.27 Station.28 Station.29
                                              Station.3 Station.30
  15.84702
             14.10141
                        13.21872
                                               14.23830
                                   13.76510
                                                          14.19600
Station.31 Station.32 Station.33 Station.34 Station.35 Station.36
  14.85625
             16.73097
                        14.07543
                                   14.62647
                                               14.74482
                                                          15.53890
Station.37 Station.38 Station.39
                                  Station.4 Station.40
                                                         Station.5
  15.12451
             15.57174
                        15.14841
                                   14.51093
                                               16.37120
                                                          14.65009
Station.6 Station.7 Station.8 Station.9
  14.23170
            13.63123
                        13.94914
                                   14.18727
```

We can use two grouping variables to return a matrix. However, when we do this, the second argument must be specified as a list.

```
# What is the average temperature at each station and depth?
mean.temp <- tapply(ctd$temp, list(station = ctd$station, depth = ctd$depth), mean)
head(mean.temp[, 1:5])
            depth
station
                             2
                                      3
                    1
  Station.1 17.22627 17.18102 17.07373 16.92864 16.75797
  Station.10 16.67695 16.55271 16.24712 15.95271 15.66983
  Station.11 16.39310 16.17458 15.89458 15.59407 15.34814
  Station.12 16.86448 16.74119 16.52642 16.35448 16.15030
  Station.13 17.05638 16.86203 16.68034 16.49610 16.24271
  Station.14 16.98061 16.81493 16.59866 16.41940 16.16791
aggregate
If we want to apply the same grouped summary to every column in a data frame, we can use aggregate:
# what is the median of each measurement at each station?
st.medians <- aggregate(ctd[, 3:8], list(station = ctd$station), median, na.rm = TRUE)
head(st.medians)
     station
               temp salinity dox
                                    ph pct light density
1 Station.1 13.070 33.4570 7.05 8.05
                                          88.330 25.1380
2 Station.10 14.445 33.4695 7.90 8.15
                                          81.530 24.8515
3 Station.11 14.940 33.4625 7.88 8.15
                                          76.270 24.7710
4 Station.12 14.095
                     33.4530 7.66 8.12
                                          85.035 24.8905
5 Station.13 13.500 33.4635 7.42 8.10
                                          86.640 25.0325
6 Station.14 14.170 33.4640 7.67 8.13
                                          84.700 24.8860
Be careful if the function returns more than one thing though.
st.range <- aggregate(ctd[, 3:8], list(station = ctd$station), range, na.rm = TRUE)
head(st.range)
     station temp.1 temp.2 salinity.1 salinity.2 dox.1 dox.2 ph.1 ph.2
1 Station.1
               9.92 22.74
                               33.130
                                          34.033 2.06 10.61 7.66 8.62
2 Station.10 10.36
                               33.162
                                          33.864 2.14 13.03 7.66 8.55
                     22.65
3 Station.11 10.58 23.06
                               33.209
                                          33.817 2.52 11.77 7.69 8.50
4 Station.12 10.24 23.00
                               32.561
                                          34.311 2.28 11.38 7.67 8.63
5 Station.13 10.00
                     22.99
                               33.090
                                          33.879 2.51 10.88 7.69 8.59
6 Station.14 10.20 22.74
                               33.069
                                          33.891 2.25 11.35 7.65 8.61
  pct_light.1 pct_light.2 density.1 density.2
                             22.923
                                       26.196
        69.45
                    92.25
1
                                       25.995
2
        30.53
                    89.64
                             22.945
3
         5.34
                    89.20
                             22.822
                                       25.895
4
        47.59
                    90.87
                             22.841
                                       26.041
5
        55.29
                    91.79
                             22.841
                                       26.076
        41.77
                    90.71
                             22.909
                                       26.052
str(st.range)
                40 obs. of 7 variables:
'data.frame':
$ station : Factor w/ 40 levels "Station.1", "Station.10",...: 1 2 3 4 5 6 7 8 9 10 ...
            : num [1:40, 1:2] 9.92 10.36 10.58 10.24 10 ...
 $ temp
```

```
$ salinity : num [1:40, 1:2] 33.1 33.2 33.2 32.6 33.1 ...
           : num [1:40, 1:2] 2.06 2.14 2.52 2.28 2.51 2.25 2.26 2.24 2.33 2.33 ...
           : num [1:40, 1:2] 7.66 7.66 7.69 7.67 7.69 7.65 7.67 7.66 7.65 7.67 ...
 $ pct_light: num [1:40, 1:2] 69.45 30.53 5.34 47.59 55.29 ...
 $ density : num [1:40, 1:2] 22.9 22.9 22.8 22.8 22.8 ...
Note that the column names seem to have .1 and .2 when you print the object, but they aren't in the
structure. In this case, every measurement column is itself a two column matrix:
dim(st.range$temp)
[1] 40 2
head(st.range$temp)
      [,1] [,2]
[1,] 9.92 22.74
[2,] 10.36 22.65
[3,] 10.58 23.06
[4,] 10.24 23.00
[5,] 10.00 22.99
[6,] 10.20 22.74
by
To apply a function to an entire data frame, use by(), which is works much like tapply:
# How many records per station?
st.rows <- by(ctd, ctd$station, nrow)</pre>
head(st.rows)
ctd$station
Station.1 Station.10 Station.11 Station.12 Station.13 Station.14
      3535
                 1120
                              762
                                        1876
                                                    2229
                                                               1865
str(st.rows)
 'by' int [1:40(1d)] 3535 1120 762 1876 2229 1865 1826 1865 1474 1120 ...
- attr(*, "dimnames")=List of 1
 ...$ ctd$station: chr [1:40] "Station.1" "Station.10" "Station.11" "Station.12" ...
 - attr(*, "call") = language by.data.frame(data = ctd, INDICES = ctd$station, FUN = nrow)
st.rows["Station.5"]
Station.5
      809
You can also summarize with multiple groups, which have to be included as a list:
# How many records per station?
st.depth.rows <- by(ctd, list(station = ctd$station, depth = ctd$depth), nrow)
str(st.depth.rows)
 'by' int [1:40, 1:60] 59 59 58 67 58 66 59 67 59 59 ...
- attr(*, "dimnames")=List of 2
 ..$ station: chr [1:40] "Station.1" "Station.10" "Station.11" "Station.12" ...
  ..$ depth : chr [1:60] "1" "2" "3" "4" ...
 - attr(*, "call") = language by.data.frame(data = ctd, INDICES = list(station = ctd$station, depth = ct
```

```
# The object can be indexed like a matrix
st.depth.rows["Station.1", "12"]
```

[1] 59

### mapply

To apply a function to sequential elements of multiple vectors, use mapply. The first argument is a function, and every argument afterwards is an argument to that function composed of vectors being iterated over. For example, the following creates a list of random numbers of alternating length with increasing range:

```
mapply(sample, x = 5:10, size = c(20, 4), replace = TRUE)

[[1]]
  [1] 3 1 3 3 1 2 1 1 3 3 2 3 4 1 4 5 3 4 2 2

[[2]]
  [1] 3 1 3 4

[[3]]
  [1] 4 7 6 3 2 3 4 1 1 7 3 1 3 5 5 5 5 1 5 4

[[4]]
  [1] 8 6 3 5

[[5]]
  [1] 6 6 6 7 9 9 9 9 3 1 9 6 8 1 5 3 5 8 1 8

[[6]]
  [1] 2 10 7 9
```

# split

A handy function for creating lists based on a grouping variable is **split**. It will split a vector, matrix, or data frame. For instance, here is a list where every element is a data frame containing only one station's data:

```
st.list <- split(ctd, ctd$station)
head(st.list[[1]])</pre>
```

```
station sample_date temp salinity dox
                                             ph pct_light density depth
1 Station.1 2012-11-08 16.81
                               33.420 8.07 8.20
                                                    90.32 24.346
2 Station.1 2012-04-19 10.52
                               33.805 3.16 7.73
                                                    88.14 25.930
                                                                     18
3 Station.1 2010-01-06 15.11
                                                    88.97 24.725
                                                                     32
                               33.415 7.22 8.13
4 Station.1 2014-02-06 14.00
                               33.430 7.31
                                                    88.01 24.974
                                                                     41
                                             NA
5 Station.1 2011-01-05 14.20
                               33.286 7.91 8.16
                                                           24.822
                                                    86.17
                                                                      3
6 Station.1 2015-02-03 13.92
                                                    87.68 24.953
                               33.382 6.45 8.05
                                                                     51
head(st.list[[2]])
```

```
ph pct_light density
       station sample_date temp salinity dox
3536 Station.10 2010-05-10 14.99
                                   33.479 9.62 8.35
                                                        70.32 24.799
3537 Station.10 2011-02-02 13.10
                                   33.337 7.24 8.06
                                                        65.39 25.085
                                   33.406 8.62 8.17
3538 Station.10 2010-03-17 13.45
                                                        73.64
                                                               25.069
3539 Station.10 2016-08-02 19.91
                                   33.465 8.98 8.28
                                                        82.14
                                                               23.616
3540 Station.10 2016-11-02 14.00
                                   33.279 6.68 8.02
                                                        79.46 24.858
```

```
3541 Station.10 2010-03-17 13.53
                                     33.404 8.62 8.20
                                                            72.08 25.050
     depth
3536
         4
3537
         6
3538
         6
        12
3539
3540
        19
3541
         4
Here's the same creating an elment for each cast (station x date):
st.dt.list <- split(ctd, list(station = ctd$station, date = ctd$sample_date))
st.dt.list[[1]]
[1] station
                sample_date temp
                                          salinity
                                                      dox
                                                                   ph
[7] pct_light
                density
                             depth
<0 rows> (or 0-length row.names)
Because it does all combinations of the grouping factors, a lot will be empty. Let's find them:
num.rows <- sapply(st.dt.list, nrow)</pre>
zero.rows <- which(num.rows == 0)
st.dt.list <- st.dt.list[-zero.rows]</pre>
st.dt.list[[1]]
        station sample_date temp salinity dox
                                                    ph pct_light density
                 2010-01-05 14.72
                                     33.374 7.61 8.18
                                                            78.87
5929 Station.12
                                                                   24.779
6294 Station.12
                 2010-01-05 14.72
                                      33.374 7.61 8.18
                                                            79.14
                                                                   24.778
                                                            79.32
6295 Station.12
                 2010-01-05 14.72
                                     33.373 7.59 8.18
                                                                   24.778
                 2010-01-05 14.72
                                      33.373 7.59 8.18
6750 Station.12
                                                            79.11
                                                                   24.778
6775 Station.12
                 2010-01-05 14.72
                                     33.374 7.60 8.18
                                                            78.97
                                                                   24.779
                                      33.375 7.50 8.18
6778 Station.12
                 2010-01-05 14.64
                                                            77.96
                                                                   24.796
6794 Station.12
                 2010-01-05 14.72
                                      33.373 7.59 8.18
                                                            79.13
                                                                   24.778
6856 Station.12
                 2010-01-05 14.71
                                      33.374 7.56 8.18
                                                            79.24
                                                                   24.781
6957 Station.12
                 2010-01-05 14.59
                                     33.373 7.43 8.16
                                                            68.05
                                                                   24.807
6973 Station.12
                 2010-01-05 14.59
                                      33.373 7.44 8.16
                                                            69.51
                                                                   24.806
6992 Station.12
                 2010-01-05 14.72
                                      33.365 7.59 8.18
                                                            78.82
                                                                   24.773
7061 Station.12
                 2010-01-05 14.71
                                     33.363 7.62 8.19
                                                            78.36
                                                                   24.772
7067 Station.12
                                     33.245 7.52 8.17
                                                            74.31
                 2010-01-05 14.75
                                                                   24.674
7087 Station.12
                 2010-01-05 14.72
                                     33.368 7.58 8.18
                                                            79.09
                                                                   24.775
7094 Station.12
                 2010-01-05 14.59
                                     33.373 7.45 8.17
                                                            70.50
                                                                   24.805
7103 Station.12
                 2010-01-05 14.60
                                      33.374 7.45 8.17
                                                            72.98
                                                                   24.804
7108 Station.12
                 2010-01-05 14.75
                                      33.243 7.52 8.17
                                                            74.02
                                                                   24.671
                 2010-01-05 14.73
                                                            76.89
7131 Station.12
                                      33.328 7.61 8.18
                                                                   24.742
7161 Station.12
                 2010-01-05 14.71
                                      33.364 7.61 8.18
                                                            79.00
                                                                   24.773
7172 Station.12
                 2010-01-05 14.71
                                     33.360 7.64 8.18
                                                            78.11
                                                                   24.770
7199 Station.12
                 2010-01-05 14.72
                                      33.366 7.58 8.18
                                                            78.80
                                                                   24.773
7218 Station.12
                 2010-01-05 14.72
                                      33.347 7.63 8.18
                                                            77.19
                                                                   24.759
7260 Station.12
                 2010-01-05 14.73
                                      33.310 7.58 8.18
                                                            74.98
                                                                   24.727
7266 Station.12
                 2010-01-05 14.72
                                      33.372 7.59 8.18
                                                            78.90
                                                                   24.777
7271 Station.12
                 2010-01-05 14.72
                                     33.368 7.60 8.18
                                                            78.87
                                                                   24.775
7275 Station.12
                 2010-01-05 14.74
                                     33.268 7.55 8.17
                                                            73.94
                                                                   24.693
7284 Station.12
                 2010-01-05 14.73
                                     33.324 7.60 8.18
                                                            76.35
                                                                   24.739
7285 Station.12
                 2010-01-05 14.72
                                     33.371 7.58 8.18
                                                            79.13
                                                                   24.777
     depth
5929
        21
6294
        20
```

```
6295
         19
         18
6750
6775
         22
         24
6778
6794
         17
         23
6856
6957
         28
6973
         27
6992
         11
7061
          9
7067
          2
7087
         14
7094
         26
7103
         25
7108
          1
7131
7161
         10
7172
          8
7199
         12
7218
          7
7260
          4
7266
         16
7271
         13
7275
          3
7284
          5
7285
         15
```

# **Function Definition**

The basic format for declaring a function actually uses a function called (wait for it...) function. I like to think of every function as having four components:

- Name The name a user will use to call the function.
- **Arguments** The input values a function needs to operate on.
- Body The code that processes the arguments.
- Return Value The output from the result of the processing in Body.

However, in any given function, depending on its purpose, one or more of the above items may be missing. Here is a simple function that has all four components designed to determine if x is between a and b:

```
isBetween <- function(x, a, b) {
   gt.a <- x > a
   lt.b <- x < b
   btwn <- gt.a & lt.b
   return(btwn)
}</pre>
```

In this function, the name is is Between, and if we call it with the arguments x = 6, a = 2, and b = 10, it will return the value TRUE:

```
isBetween(x = 6, a = 2, b = 10)
```

[1] TRUE

# Name

A function's name should be short(ish), but also meaningful and easy to understand. This is an art and you should take the time to play with names until they fit. Pretend like you are a naive user who has no idea what the function does. You should be able to get most of that information from the name. Also pay attention to how other functions that your function will be working with are named. For example, if you have a function that reads a particular data file and formats it, but is only one of several data files that will be read, it would be bad form to call that function readData. It might be better to call it something like, readSalinityData. Even better might be, readSalinityData. However, it would be unnecessarily long and mean to users to call it something like, readSalinityDataFromCSVFileAndRemoveMissingDataPoints.

# Arguments

These are the input values that the function needs to operate on. It is good programming practice to make them both as short and as long as necessary to be descriptive. In general, names for arguments should also be short and descriptive. However, some argument names are frequently used, such as x for the first argument, and y for the second argument, especially in mathematically-based functions or for data for axes in plotting functions. It is good practice to not refer to anything in the function body that is either not in the arguments, or is not created in the function body.

### Body

This is the code that is the heart of the function. It operates on the arguments to convert them to a value to be returned or perform an action. Curly braces ({ and }) are used to denote the code that composes the body and belongs to the function. If the function only has one line for a body then the curly braces can be omitted. For example:

```
isBetween.2 <- function(x, a, b) return(x > a & x < b)
isBetween.2(x = 6, a = 2, b = 10)
```

[1] TRUE

#### Return Value

A function can only return one object. The function called return is often used to denote what this object is as in the above examples. However, if there is no call to return, then the result of the last line in a function is its return value. For example, our addAndRaise function could also be written as:

```
isBetween.3 <- function(x, a, b) {
  gt.a <- x > a
  lt.b <- x < b
  gt.a & lt.b
}
isBetween.3(x = 6, a = 2, b = 10)</pre>
```

#### [1] TRUE

Sometimes you want a function to do an action, but only return a value if it is assigned to something. In this case, use the invisible function. In this example, our isBetween function will not print the result when called by itself:

```
isBetween <- function(x, a, b) {
  gt.a <- x > a
  lt.b <- x < b</pre>
```

```
invisible(gt.a & lt.b)
}
# nothing printed
isBetween(x = 6, a = 2, b = 10)

# assign to object
result <- isBetween(x = 6, a = 2, b = 10)
result</pre>
```

[1] TRUE

# Arguments

To better understand how arguments are handled, let's first create a function that abbreviates vectors of scientific names to shorter versions. For instance, we want a function that takes "Homo" and "sapiens" and creates "H sap":

```
abbrev <- function(genus, species) {
    # get the first character from genus names
    g <- substr(genus, 1, 1)
    # get the first three characters from the species names
    spp <- substr(species, 1, 3)
    # paste the two together and return the result
    paste(g, spp)
}</pre>
```

Let's also load some data to use with it:

```
spp.codes <- read.csv("tblCodeSpecies.csv", stringsAsFactors = FALSE)
head(spp.codes)</pre>
```

```
SPCODE
           ORDER
                   SUBORDER
                                  FAMILY FAMILY.NAMES
                                                             GENUS
     001 CETACEA ODONTOCETI
1
                              ZIPHIIDAE BEAKED WHALES Mesoplodon
                                              DOLPHINS
     002 CETACEA ODONTOCETI DELPHINIDAE
                                                          Stenella
2
3
     003 CETACEA ODONTOCETI DELPHINIDAE
                                              DOLPHINS
                                                          Stenella
    004 CETACEA ODONTOCETI DELPHINIDAE
4
                                                          Stenella
                                              DOLPHINS
5
     005 CETACEA ODONTOCETI DELPHINIDAE
                                              DOLPHINS Delphinus
6
    006 CETACEA ODONTOCETI DELPHINIDAE
                                                          Stenella
                                              DOLPHINS
              SPECIES
                                        COMMON.NAME
                                 Pygmy beaked whale
1
           peruvianus
            attenuata Pantropical spotted dolphin
2
3 longirostris subsp. unidentified spinner dolphin
4
              clymene
                                    Clymene dolphin
5
                       Unidentified common dolphin
                  sp.
6 attenuata graffmani
                            Coastal spotted dolphin
gns <- spp.codes$GENUS</pre>
spp <- spp.codes$SPECIES</pre>
... and test it out:
gspp <- abbrev(genus = gns, species = spp)</pre>
head(gspp)
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

#### **Arguments - Matching**

Argument are matched according to two rules. First, arguments that are specifically named, like **genus = gns** are matched. Then any remaining unnamed arguments are matched based on the order in which they are found. This is simple to understand in our two argument function, which we can call as we have before, or like this:

```
# 'species' is not named
gspp <- abbrev(genus = gns, spp)
head(gspp)

[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"

# 'genus' is not named
gspp <- abbrev(species = spp, gns)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

If you know the name and order of the arguments, none of them have to be named as long as they are always supplied in the correct order. In many commonly used functions, this is normal for the first few arguments. So we would normally call this function like this:

```
gspp <- abbrev(gns, spp)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

Let's add a third argument to specify the number of characters in the species name to demonstrate the name and order matching further:

```
abbrev <- function(genus, species, num.spp) {
    # get the first character from genus names
    g <- substr(genus, 1, 1)
    # get the 'num.spp' characters from the species names
    spp <- substr(species, 1, num.spp)
    # paste the two together and return the result
    paste(g, spp)
}
gspp <- abbrev(gns, spp, 3)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
# we can also call it like this:
gspp <- abbrev(num.spp = 3, gns, spp)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
# this is fine too:
gspp <- abbrev(gns, num.spp = 3, spp)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
# but this will not produce the desired output:
gspp <- abbrev(num.spp = 3, spp, gns)
head(gspp)</pre>
```

```
[1] "p Mes" "a Ste" "l Ste" "c Ste" "s Del" "a Ste"
```

Argument names can also be abbreviated as long as the abbreviations are unique. Let's add a fourth argument specifying the number of characters in the genus name to return:

```
abbrev <- function(genus, species, num.g, num.spp) {
    # get the first 'num.g' characters from genus names
    g <- substr(genus, 1, num.g)
    # get the first 'num.spp' characters from the species names
    spp <- substr(species, 1, num.spp)
    # paste the two together and return the result
    paste(g, spp)
}
gspp <- abbrev(gns, spp, num.g = 1, num.spp = 3)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

We can abbreviate gns as just g because no other arguments start with "g":

```
gspp <- abbrev(s = spp, g = gns, num.g = 1, num.spp = 3)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

However if we want to abbreviate num.spp, the shortest we can make it is num.s because any shorter than that and you couldn't differentiate it from num.g:

```
gspp <- abbrev(s = spp, g = gns, num.g = 1, num.s = 3)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

This will produce an error:

```
gspp <- abbrev(s = spp, g = gns, num = 1, 3)</pre>
```

Error in abbrev(s = spp, g = gns, num = 1, 3): argument 3 matches multiple formal arguments

# **Arguments - Defaults**

Sometimes it is useful to specify default values for arguments. This means that users do not have to enter the default values every time an argument is called, but they can be modified if need be. Default values are specified by setting them directly in the argument list:

```
abbrev <- function(genus, species, num.g = 1, num.spp = 3) {
    # get the first 'num.g' characters from genus names
    g <- substr(genus, 1, num.g)
    # get the first 'num.spp' characters from the species names
    spp <- substr(species, 1, num.spp)
    # paste the two together and return the result
    paste(g, spp)
}
gspp <- abbrev(gns, spp)
head(gspp)</pre>
```

```
[1] "M per" "S att" "S lon" "S cly" "D sp." "S att"
```

But they can be changed by specifying them by name or position in the function call:

```
# set num.g to 3
gspp <- abbrev(gns, spp, 3)
head(gspp)

[1] "Mes per" "Ste att" "Ste lon" "Ste cly" "Del sp." "Ste att"
# set num.spp to 1
gspp <- abbrev(gns, spp, num.spp = 1)
head(gspp)

[1] "M p" "S a" "S l" "S c" "D s" "S a"</pre>
```

### **Arguments - Ellipses**

There are times when you want to be able to pass arguments on to functions within your function, but you don't want to have to specify all possible arguments for that function in your argument list. To solve this, you can use the ellipses or dot-dot-dot notation, .... Here we use them to pass on formatting arguments like sep and collapse to the paste function:

```
abbrev <- function(genus, species, num.g = 1, num.spp = 3, ...) {
  # get the first 'num.g' characters from genus names
  g <- substr(genus, 1, num.g)
  # get the first 'num.spp' characters from the species names
  spp <- substr(species, 1, num.spp)
  # paste the two together and return the result
  paste(g, spp, ...)
}
gspp <- abbrev(gns, spp, sep = ".")
head(gspp)</pre>
```

```
[1] "M.per" "S.att" "S.lon" "S.cly" "D.sp." "S.att"
```

# **Disposable Functions**

There are times when an argument to a function is another function. In these cases, the function being passed can be predefined or, if it is being defined only for this purpose, it can just be defined directly in the argument list. This latter method creates what is called an "anonymous" or "disposable" function. One place this is frequently done is with the 'apply family of functions where one function is applied to every element in an object and the results are collected and returned in a convenient form. For example, let's say that species code data.frame is actually a list of data.frames with one element per Family:

```
families <- split(spp.codes, spp.codes$FAMILY)
names(families)</pre>
```

```
[1] ""
                        "BALAENIDAE"
                                           "BALAENOPTERIDAE"
[4] "CHELONIDAE"
                        "DELPHINIDAE"
                                           "DERMOCHELYIDAE"
[7] "ESCHRICTIDAE"
                        "INIIDAE"
                                           "KOGIIDAE"
[10] "MONODONTIDAE"
                        "MUSTELIDAE"
                                           "NEOBALAENIDAE"
[13] "ODOBENIDAE"
                        "OTARIIDAE"
                                           "PHOCIDAE"
[16] "PHOCOENIDAE"
                        "PHYSETERIDAE"
                                           "PLATANISTIDAE"
[19] "PONTOPORIIDAE"
                        "RHINCODONTIDAE"
                                           "SIRENIA"
[22] "UNIDENTIFIED"
                        "URSIDAE"
                                           "ZIPHIIDAE"
```

Now we want to know how many genera are in each Family. Let's first make a function that takes a data.frame and returns the number of unique genera in that data.frame:

```
num.genera <- function(df) length(unique(df$GENUS))
# Number of genera in entire data.frame
num.genera(spp.codes)</pre>
```

#### [1] 82

```
# Number of genera in Ziphiidae
num.genera(families$ZIPHIIDAE)
```

#### [1] 6

To get the number of genera in each family, we want to use sapply on the families list, where the second argument is the function we want to apply to every element in families:

```
sapply(families, num.genera)
```

	BALAENIDAE	BALAENOPTERIDAE	CHELONIDAE
2	2	2	6
DELPHINIDAE	DERMOCHELYIDAE	ESCHRICTIDAE	INIIDAE
20	2	1	1
KOGIIDAE	MONODONTIDAE	MUSTELIDAE	NEOBALAENIDAE
1	2	1	1
ODOBENIDAE	OTARIIDAE	PHOCIDAE	PHOCOENIDAE
1	8	8	5
PHYSETERIDAE	PLATANISTIDAE	PONTOPORIIDAE	RHINCODONTIDAE
1	1	2	1
SIRENIA	UNIDENTIFIED	URSIDAE	ZIPHIIDAE
2	7	1	6

However, if we only need the num.genera function for the sapply then we don't have to predefine it. We can create the disposable function directly in the sapply argument list for the FUN argument, which is second (see ?sapply):

```
sapply(families, function(df) length(unique(df$GENUS)))
```

	BALAENIDAE	BALAENOPTERIDAE	CHELONIDAE
2	2	2	6
DELPHINIDAE	DERMOCHELYIDAE	ESCHRICTIDAE	INIIDAE
20	2	1	1
KOGIIDAE	MONODONTIDAE	MUSTELIDAE	NEOBALAENIDAE
1	2	1	1
ODOBENIDAE	OTARIIDAE	PHOCIDAE	PHOCOENIDAE
1	8	8	5
PHYSETERIDAE	PLATANISTIDAE	PONTOPORIIDAE	RHINCODONTIDAE
1	1	2	1
SIRENIA	UNIDENTIFIED	URSIDAE	ZIPHIIDAE
2	7	1	6

The disposable function may be more than one line in which case we just need to use curly braces to collect the statements. In this example the function returns the number of genera and the number of species. Make sure to include the closing curly brace for the function and the closing parentheses for the sapply:

```
sapply(families, function(df) {
  num.gen <- length(unique(df$GENUS))
  num.spp <- length(unique(df$SPECIES))
  c(num.gen = num.gen, num.spp = num.spp)
})</pre>
```

	BALAENIDAE	BALAENO	PTERIDAE	CHELON	IDAE I	DELPHINIDAE	DERN	OCHELYID!	ΑE
num.gen	2 2		2		6	20			2
num.spp	1 3		9		7	45			2
	ESCHRICTIDAE	INIIDAE	KOGIIDA	E MONOD	ONTIDA	AE MUSTELIDA	4E		
num.gen	1	1		1		2	1		
num.spp	1	1	;	3		2	1		
	NEOBALAENIDA	E ODOBEN	IDAE OTAI	RIIDAE	PHOCII	DAE PHOCOENI	[DAE		
${\tt num.gen}$	:	1	1	8		8	5		
num.spp	:	1	1	10		14	7		
	PHYSETERIDAE	PLATANIS	STIDAE PO	ONTOPOR	IIDAE	RHINCODONT	[DAE	SIRENIA	
${\tt num.gen}$	1		1		2		1	2	
num.spp	1		2		2		1	3	
	UNIDENTIFIED	URSIDAE	ZIPHIID	ΑE					
${\tt num.gen}$	7	1		6					
${\tt num.spp}$	1	1	:	24					

### Flow Control

Normally when code is executed, the "flow" proceeds in a linear fashion. The first line is executed, then the second, and so forth until the last line of code is reached. There can be situations where you want to direct this flow either in branching form: some piece of code is executed based on one condition, while another piece is executed based on another condition, or in a looping manner: the same code is executed repeatedly until some stopping criterion is reached (either number of iterations, or a condition is met). There are several functions that allow you to manage this flow control, the help for which can be found with ?Control.

### Branching

There are three standard branching functions:

- if (cond) cons.expr else alt.expr: executes a set of code (cons.expr) if cond evaluates to TRUE or (optionally) alternative code (alt.expr) if it is FALSE.
- ifelse(test, yes, no): returns elements from yes matching to elements in test that are TRUE and elements in no for elements in test that are FALSE.
- switch(EXPR, ...): executes individual code for named or numbered values in EXPR.

The thing to remember is that if is used for a single branching event (when else is not used) or a bifurcating branch (when else is used) that is based on a single condition (one T or F). ifelse is used to return a vector that is the same length as the logical vector with one set of values of for TRUE elements and another set for FALSE elements. switch is used in places where you want different pieces of code run for different discrete values. This is usually preferred if there are more than two possible conditions.

#### if

As an example of if, lets construct some checks of argument ranges in our species abbreviation code:

```
abbrev <- function(genus, species, num.g = 1, num.spp = 3, ...) {
    # 'num.g' must be 1 or greater
    if(num.g < 1) num.g <- 1
    # 'num.g' shouldn't be too big
    if(num.g > 3) num.g <- 3
    # get the first 'num.g' characters from genus names
    g <- substr(genus, 1, num.g)
    # get the first 'num.spp' characters from the species names
    spp <- substr(species, 1, num.spp)</pre>
```

```
# paste the two together and return the result
paste(g, spp, ...)
}
gspp <- abbrev(gns, spp, num.g = 0, sep = ".")
head(gspp)

[1] "M.per" "S.att" "S.lon" "S.cly" "D.sp." "S.att"
gspp <- abbrev(gns, spp, num.g = 10, sep = ".")</pre>
```

```
[1] "Mes.per" "Ste.att" "Ste.lon" "Ste.cly" "Del.sp." "Ste.att"
```

head(gspp)

Let's say that there are just two abbreviation formats, a short one like: "H.sap" and just the abbreviated genus: "H. sapiens". We control this with a simple argument, called **short**:

```
abbrev <- function(genus, species, short = T) {
  if(short) {
    g <- substr(genus, 1, 1)
    spp <- substr(species, 1, 3)
    g.spp <- paste(g, spp, sep = ".")
    return(g.spp)
} else {
    g <- substr(genus, 1, 1)
    g.spp <- paste(g, species, sep = ".")
    return(g.spp)
}

# The short form
head(abbrev(gns, spp))</pre>
```

```
[1] "M.per" "S.att" "S.lon" "S.cly" "D.sp." "S.att"

# The longer form
head(abbrev(gns, spp, F))
```

We used return(g.spp) in order to make sure the function returns the result from the execution branch for each condition. We can simplify this code in several convenient ways. The first is based on the fact that the last line in the expression for each condition is the return value of the if function. So we can assign g.spp based on each branch then return it once at the end:

```
abbrev <- function(genus, species, short = T) {
   g.spp <- if(short) {
      g <- substr(genus, 1, 1)
      spp <- substr(species, 1, 3)
      paste(g, spp, sep = ".")
   } else {
      g <- substr(genus, 1, 1)
      paste(g, species, sep = ".")
   }
   return(g.spp)
}
# The short form</pre>
```

```
head(abbrev(gns, spp))

[1] "M.per" "S.att" "S.lon" "S.cly" "D.sp." "S.att"

# The longer form
head(abbrev(gns, spp, F))

[1] "M. peruvianus" "S. attenuata"
[3] "S. longirostris subsp." "S. clymene"
[5] "D. sp." "S. attenuata graffmani"
```

Also notice that we create g <- substr(genus, 1, 1) in each expression, so we can move that to the outside. Also, because the result of the last line in a function is its return value we can remove the return(g.spp) line:

```
abbrev <- function(genus, species, short = T) {
   g <- substr(genus, 1, 1)
   if(short) {
      spp <- substr(species, 1, 3)
      paste(g, spp, sep = ".")
   } else {
      paste(g, species, sep = ".")
   }
}
# The short form
head(abbrev(gns, spp))</pre>
```

```
[1] "M.per" "S.att" "S.lon" "S.cly" "D.sp." "S.att"
# The longer form
head(abbrev(gns, spp, F))
```

#### ifelse

The ifelse function returns a vector that is as long as its first argument and chooses from the corresponding yes and no vectors to fill the elements. As an example, here's a function that will create the abbreviation "H. sapiens", but if the species name is longer than 8 characters, it will abbreviate that too:

```
abbrev <- function(genus, species) {
  g <- substr(genus, 1, 1)
  spp <- ifelse(nchar(species) > 8, substr(species, 1, 8), species)
  paste(g, spp, sep = ". ")
}
head(abbrev(gns, spp))
```

```
[1] "M. peruvian" "S. attenuat" "S. longiros" "S. clymene" "D. sp." [6] "S. attenuat"
```

The expressions in **ifelse** can be multiple lines too, but must be wrapped by curly braces. This modification adds a "." to the end of the abbreviated species name:

```
abbrev <- function(genus, species) {
  g <- substr(genus, 1, 1)
  spp <- ifelse(</pre>
```

```
nchar(species) > 8,
      spp.sub <- substr(species, 1, 8)</pre>
      paste0(spp.sub, ".")
    },
    species
  paste(g, spp, sep = ". ")
head(abbrev(gns, spp))
[1] "M. peruvian." "S. attenuat." "S. longiros." "S. clymene"
                    "S. attenuat."
[5] "D. sp."
switch
The final branching function is switch which allows us to choose one of a series of expressions to execute
based on a numeric or character value. For example, our abbreviation code will have an argument, type that
will allow for three formats: short = "H.sap", medium = "H. sapiens", and long = "Homo sapiens":
abbrev <- function(genus, species, type) {</pre>
  g <- substr(genus, 1, 1)
  # we only need an `if` statement to format the species
  spp <- if(type == "short") substr(species, 1, 3) else species</pre>
  # choose the pasting format based on `type`
  switch(type,
    short = paste0(g, ".", spp),
    medium = paste0(g, ". ", spp),
    long = paste(genus, species)
  )
}
# The short form
head(abbrev(gns, spp, "short"))
```

```
[1] "M.per" "S.att" "S.lon" "S.cly" "D.sp." "S.att"

# The medium form
head(abbrev(gns, spp, "medium"))
```

```
[1] "Mesoplodon peruvianus" "Stenella attenuata"
[3] "Stenella longirostris subsp." "Stenella clymene"
[5] "Delphinus sp." "Stenella attenuata graffmani"
```

# Looping

There are three functions to control looping:

• for(var in seq): Executes a set of code for a number of iterations equal to the length of seq. In each iteration var gets sequential values of seq.

- while(cond) expr : Executes a set of code as long as cond is TRUE.
- repeat expr : Repeats code. To stop looping, execute break.

for

With for we execute a set of code for each element in a vector and in each execution, a variable takes sequential values of that vector. In the below example, we calculate the first n values of the fibonacci series.

```
fib <- function(n) {
    x <- 0
    for(i in 2:n) {
        if(i == 2) {
            x[i] <- 1
        } else {
            x[i] <- x[i - 1] + x[i - 2]
        }
    }
    x
}
fib(10)</pre>
```

### [1] 0 1 1 2 3 5 8 13 21 34

A for loop can be stopped with the break function. For example, we will put in a function that stops the loop the first time a number greater than 50 is reached:

```
fib <- function(n) {
    x <- 0
    for(i in 2:n) {
        if(i == 2) {
            x[i] <- 1
        } else {
            x[i] <- x[i - 1] + x[i - 2]
        }
        if(x[i] > 50) break
    }
    x
}
fib(13)
```

### [1] 0 1 1 2 3 5 8 13 21 34 55

We can also force the for loop to iterate again before reaching the natural end of code in an iteration with the next command. In this example, we only print values greater than the number specified in print.num:

```
fib <- function(n, print.num = 10) {
    x <- 0
    for(i in 2:n) {
        if(i == 2) {
            x[i] <- 1
        } else {
            x[i] <- x[i - 1] + x[i - 2]
        }
        if(x[i] < print.num) next
        cat(i, ": ", x[i], "\n")
}</pre>
```

```
X
}
fib(15)
     13
9
     21
10 : 34
11
      55
12
     89
13
   : 144
14 : 233
15 : 377
 [1]
                   3 5 8 13 21 34 55 89 144 233 377
             1
                2
```

### while

The while function is designed to repeat some code until a condition is met. If the condition is never met, the loop will continue indefinitely. The break and next commands will perform the same with function and repeat. In this example, we print the fibonacci series until the specified number is exceeded:

```
fib <- function(n) {
  first <- 0
  second <- 1
  cat(first, " ")
  while(second <= n) {
    cat(second, " ")
    new.val <- first + second
    first <- second
    second <- new.val
  }
}
fib(20)</pre>
```

0 1 1 2 3 5 8 13

### repeat

The repeat function will continuously execute code until it is stopped by a break. Here, we do the same loop as above, but replace while with a repeat and break:

```
fib <- function(n) {
  first <- 0
  second <- 1
  cat(first, " ")
  repeat {
    cat(second, " ")
    new.val <- first + second
    if(new.val > n) break
    first <- second
    second <- new.val
  }
}
fib(20)</pre>
```

# Argument error checking

Despite our best efforts, functions are susceptible to users entering improper arguments or errors showing up that keeps a function from completing. We have some tools at our disposal to mitigate this. First, we can check that arguments what we expect and require using if statements. If they're aren't we can do something like return NA or NULL:

```
addTwo <- function(a, b) {
    # confirm that a and b are numbers
    if(!(is.numeric(a) | is.numeric(b))) return(NULL)
    a + b
}
addTwo(1, "x")</pre>
```

Error in a + b: non-numeric argument to binary operator

```
addTwo(1, 2)
```

[1] 3

We can also issue warnings when something unexpected happens:

```
addTwo <- function(a, b) {
    # confirm that a and b are numbers
    if(!(is.numeric(a) | is.numeric(b))) {
        warning("'a' and 'b' must be numbers. NULL returned.")
        return(NULL)
    }
    a + b
}
addTwo(1, "x")</pre>
```

Error in a + b: non-numeric argument to binary operator

If execution cannot or should not continue, then an error can be thrown with the stop function:

```
divideTwo <- function(a, b) {
    # confirm that a and b are numbers
    if(!(is.numeric(a) | is.numeric(b))) {
        stop("'a' and 'b' must be numbers. NULL returned.")
    }
    if(b == 0) {
        stop("'b' cannot be 0")
    }
    a / b
}
divideTwo(1, "x")</pre>
```

Error in a/b: non-numeric argument to binary operator

```
divideTwo(5, 0)
```

Error in divideTwo(5, 0): 'b' cannot be 0

# Scope

It is important to note that objects declared in a function only exist within that function. On the flipside, objects declared outside of a function are accessible by that function. However, it is very bad form to refer to an object in a function that has not been passed as an argument or declared in the function itself. For example:

```
a <- 2
my.func <- function(x, y) (x + y) * a

# this works
my.func(2, 3)

[1] 10
# remove a from the workspace
rm(a)

# a can't be found, so this produces an error
my.func(2, 3)

Error in my.func(2, 3): object 'a' not found</pre>
```

# Homework

Answer all questions in a script (.R) file. Use comments (# or #') to explain steps in code.

- 1. In the "tblCodeSpecies.csv" data set, how many named species of the family "Balaenopteridae" are there?
- 2. In the "tblCodeSpecies.csv" data set, what is the mean number of species per family of Cetaceans (order Cetacea) and Pinnipeds (suborder Pinnipedia)?
- 3. In the "ctd.csv" data set, what is the mean difference in temperature between 10 meters and the surface?
- 4. In the "ctd.csv" data set, which stations have the lowest and highest mean surface temperature?
- 5. Write a function that returns the following summary statistics for a numerical vector: mean, standard deviation, number of values, number missing, minimum, maximum, and median.
- 6. Add to the function in 5 some code to make sure that the input is valid numerical vector.
- 7. Write a function that uses the function in 6 to summarize the measurements from a data.frame like " ${\it ctd.csv}$ "
- 8. Write a function that uses the function in 6 to summarize the measurements from a CTD data.frame by station and depth.
- 9. Write a function that uses the function in 8 to identify outliers (> 3 standard deviations from mean). The function should take as input a data.frame of CTD casts like "ctd.csv", a depth value of interest, and a measurement of interest. The output should be a three column data.frame that contains the station, sample date, and value of that measurement for every outlier identified.
- 10. Write a function that uses the function in 9 to identify outliers for every depth and measurement.