# Intro to Base Riterations

And Lab 1

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Week 2

# Agenda

- For loops
- Apply family of loops
  - o lapply()
  - o sapply()
  - o vapply()
  - o apply() (briefly)

# Learning objectives

- Understand the basics of what it means to loop through a vector
- Begin to recognize use cases
- Be able to apply basic for loops and write their equivalents with lapply.

# Basic overview: for loops

```
a <- letters[1:26]
a

## [1] "a" "b" "c" "d" "e" "f" "g" "h" "i" "j" "k" "l" "m" "n" "o" "p" "q"
## [22] "v" "w" "x" "y" "z"

For(i in 1:5){
    print(a[i])</pre>
Note these are five different
character appliess (atomic
```

```
## [1] "a'
## [1] "b'
## [1] "c'
## [1] "d'
## [1] "e'
```

Note these are five different character scalars (atomic vectors of length one). It is NOT a single vector.

# Creating indexes

You will commonly see code like

```
for(i in 1:nrow(df))
```

Instead, preference seq\_along() or seq\_len().

#### Examples

```
x <- c(3, 2.6, 8)
seq_along(x)

## [1] 1 2 3

seq_len(length(x))

## [1] 1 2 3</pre>
```

## seq\_\*

- seq\_along() will create an index from 1 to the length of the vector
- seq\_len() takes a single positive integer argument, and creates an index from 1 to that integer

Why are these preferable?

# Avoid the 1 o problem

```
means <- c()
out <- rep(NA, length(means))

for (i in 1:length(means)) {
  out[i] <- rnorm(10, means[i])
}</pre>
```

## Error in rnorm(10, means[i]): invalid arguments

#### seq\_\* version

```
means <- c()
out <- rep(NA, length(means))

for (i in seq_along(means)) {
  out[i] <- rnorm(10, means[i])
}</pre>
```

Basically, the loop is just not executed.

# Another basic example

#### Simulate tossing a coin, record results

For a single toss

```
sample(c("Heads", "Tails"), 1)
```

```
## [1] "Heads"
```

• For multiple tosses, first allocate a vector with **length** equal to the number of iterations

```
result <- rep(NA, 10)
result
```

## [1] NA NA NA NA NA NA NA NA NA

ullet Next, run the trial  $oldsymbol{n}$  times, storing the result in your preallocated vector.

```
for(i in seq_along(result)) {
    result[i] <- sample(c("Heads", "Tails"), 1)
}
result</pre>
```

## [1] "Tails" "Heads" "Tails" "Tails"

# Growing vectors

- Always pre-allocate a vector for storage before running a for loop.
- Contrary to some opinions you may see out there, for loops are not actually slower than lapply, etc., provided the for loop is written well
- This primarily means not growing a vector

# Example

100,000 coin flips by growing a vector

```
library(tictoc)

set.seed(1)
tic()
not_allocated <- sample(c("Heads", "Tails"), 1)
for(i in seq_len(le5 - 1)) {
    not_allocated <- c(
        not_allocated,
        sample(c("Heads", "Tails"), 1)
    )
}
toc()</pre>
```

## 50.637 sec elapsed

#### same exact thing with pre-allocated vector

```
set.seed(1)
tic()
allocated <- rep(NA, 1e5)
for(i in seq_len(1e5)) {
    allocated[i] <- sample(c("Heads", "Tails"), 1)
}
toc()</pre>
```

## 0.935 sec elapsed

## Result

• The result is the same, regardless of the approach (notice I forced the random number generator to start at the same place in both samples)

identical(not\_allocated, allocated)

## [1] TRUE

Speed is obviously not identical

# You try

#### Base R comes with letters and LETTERS

- Make an alphabet of upper/lower case. For example,
   create "Aa" with paste0(LETTERS[1], letters[1])
- Write a for loop for all letters



## Answer

```
alphabet <- rep(NA, length(letters))

for(i in seq_along(alphabet)) {
    alphabet[i] <- paste0(LETTERS[i], letters[i])
}
alphabet</pre>
```

```
## [1] "Aa" "Bb" "Cc" "Dd" "Ee" "Ff" "Gg" "Hh" "Ii" "Jj" "Kk" "Ll" "Mm" "N
## [17] "Qq" "Rr" "Ss" "Tt" "Uu" "Vv" "Ww" "Xx" "Yy" "Zz"
```

# Another example

- Say we wanted to simulate 100 cases from random normal distribution, where we varied the standard deviation in increments of 0.2, ranging from 1 to 5
- Hopefully this is a relatable example, but if not, that's okay – focus on the process.
- First, specify a vector standard deviations

```
increments \leftarrow seq(1, 5, by = 0.2)
```

• Next, allocate a vector. There are many ways I could store this result (data frame, matrix, list). I'll do it in a list.

```
simulated <- vector("list", length(increments))</pre>
```

## Look at our vector

#### str(simulated)

```
List of 21
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
## $ : NULL
## $ : NULL
## $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
##
  $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
##
  $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
##
    $ : NULL
```

# Write for loop

```
for(i in seq_along(simulated)) {
    simulated[[i]] <- rnorm(100, 0, increments[i])
    # note use of `[[` above
}</pre>
```

Now look at our vector

#### str(simulated)

```
## List of 21
   $: num [1:100] -2.387 0.405 -1.599 -0.285 0.288 ...
## $ : num [1:100] 0.298 0.433 -1.021 1.384 -0.323 ...
   $: num [1:100] 0.893 -1.799 -0.819 -1.11 -2.198 ...
##
   $: num [1:100] -0.332 1.067 -0.823 2.899 1.863 ...
##
   $: num [1:100] -2.568 -0.672 -0.244 -1.645 2.221 ...
## $ : num [1:100] 2.4 -1.95 1.13 3.05 3.56 ...
##
  $ : num [1:100] -2.978 0.798 2.212 2.15 -2.197 ...
   $: num [1:100] -0.211 -1.768 3.35 2.06 0.213 ...
##
   $: num [1:100] 0.718 -4.029 -1.093 0.417 -3.952 ...
##
   $: num [1:100] 0.632 3.084 -2.62 -1.282 -2.965 ...
##
  $: num [1:100] 2.1759 -0.4681 1.6349 -0.0809 -0.7611 ...
## $ : num [1:100] 1.236 3.055 -2.575 -0.868 4.369 ...
   $: num [1:100] 0.7795 -1.0125 -6.465 0.0926 1.8629 ...
```

# List/data frame

- Remember, if all the vectors of our list are the same length, it can be transformed into a data frame.
- First, let's provide meaningful names

```
names(simulated) <- paste0("sd_", increments)
str(simulated)</pre>
```

```
## List of 21
   $ sd 1 : num [1:100] -2.387 0.405 -1.599 -0.285 0.288 ...
   $ sd 1.2: num [1:100] 0.298 0.433 -1.021 1.384 -0.323 ...
##
   $ sd 1.4: num [1:100] 0.893 -1.799 -0.819 -1.11 -2.198 ...
   $ sd 1.6: num [1:100] -0.332 1.067 -0.823 2.899 1.863 ...
   $ sd 1.8: num [1:100] -2.568 -0.672 -0.244 -1.645 2.221 ...
##
##
   $ sd 2 : num [1:100] 2.4 -1.95 1.13 3.05 3.56 ...
   $ sd 2.2: num [1:100] -2.978 0.798 2.212 2.15 -2.197 ...
##
   $ sd 2.4: num [1:100] -0.211 -1.768 3.35 2.06 0.213 ...
##
   $ sd 2.6: num [1:100] 0.718 -4.029 -1.093 0.417 -3.952 ...
   $ sd 2.8: num [1:100] 0.632 3.084 -2.62 -1.282 -2.965 ...
##
##
   $ sd 3 : num [1:100] 2.1759 -0.4681 1.6349 -0.0809 -0.7611 ...
   $ sd 3.2: num [1:100] 1.236 3.055 -2.575 -0.868 4.369 ...
   $ sd 3.4: num [1:100] 0.7795 -1.0125 -6.465 0.0926 1.8629 ...
   $ sd 3.6: num [1:100] 3.466 -1.245 0.496 3.67 -2.207 ...
```

## Convert to df

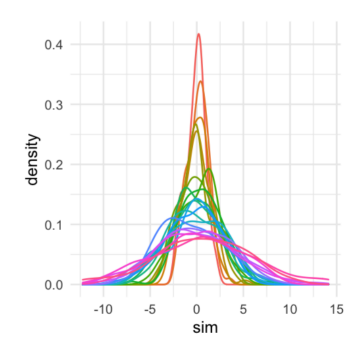
```
sim_d <- data.frame(simulated)
head(sim_d)</pre>
```

```
##
                   sd 1.2
                              sd 1.4
                                         sd 1.6
                                                    sd 1.8
                                                                 sd 2
          sd 1
## 1 -2.3872613 0.2979273 0.8930471 -0.3319310 -2.5676229 2.3954045 -2.9
## 2 0.4051212 0.4329239 -1.7989656 1.0673114 -0.6722755 -1.9542566
## 3 -1.5992856 -1.0209222 -0.8192303 -0.8232251 -0.2435614 1.1309855
                                                                      2.1
## 4 -0.2847246 1.3838266 -1.1097486 2.8991565 -1.6445420 3.0545722
## 5 0.2881735 -0.3233308 -2.1982580 1.8633398 2.2213515 3.5578607
                                                                     -2.1
## 6 0.1175257 -0.4150724 0.6353818 0.7142314 -0.5219616 -0.8029945
                                                                      0.1
##
        sd 2.4
                   sd 2.6 sd 2.8
                                            sd 3
                                                     sd 3.2
                                                                 sd 3.4
## 1 -0.210\overline{5}980 0.717\overline{5}135 0.632\overline{1}448 2.175892\overline{8}2 1.236\overline{2}364 0.7794\overline{7}966
## 2 -1.7675364 -4.0290962 3.0843027 -0.46812673 3.0554725 -1.01245886 -1
## 3 3.3501949 -1.0927053 -2.6196216 1.63492841 -2.5751022 -6.46499466
## 4 2.0601974 0.4174713 -1.2824915 -0.08085208 -0.8678742 0.09259855
## 5 0.2125117 -3.9521276 -2.9646399 -0.76111234 4.3687915 1.86290325 -2
## 6 1.7822910 -0.1081454 4.5420524 3.53122922 -2.4194781 -1.14660593 -1
##
        sd 3.8
                    sd 4 sd 4.2 sd 4.4 sd 4.6
                                                             sd 4.8
                         0.3347596 - 2.122876  2.4479578  0.6723229
## 1 -2.7119876 2.834307
                                                                    -3.46
## 2 -4.2104097 6.083823
                         0.5735697 3.164909 -1.4721622 -4.7240885
                                                                     2.82
## 3 -3.6861183 -3.000288
                         4.1063037 1.104068 4.4111688 3.3779921 -11.48
## 4 -0.7277885 4.293730
                         4.4141833 -4.064636 2.3400425 -1.8110032
                                                                   -1.85
## 5 -0.1416683 4.178344
                         0.8965279 0.578414 -0.3461786 10.3301558
                                                                     7.54
## 6 -2.2110984 4.644032 -5.7370545 -5.072939 4.7260451 0.4489877
                                                                     2.95
```

# tidyverse

 One of the best things about the tidyverse is that it often does the looping for you

```
library(tidyverse)
pd <- sim_d %>%
    pivot_longer(
      everything(),
      names_to = "sd",
      values_to = "sim",
      names_prefix = "sd_",
      names_ptypes = list(
        sd = factor()
ggplot(pd, aes(sim)) +
geom_density(
   aes(color = sd)
 ) +
guides(color = "none")
```



Of course, someone has to write loops. It doesn't have to be you. --- Jenny Bryan

## Base R Method

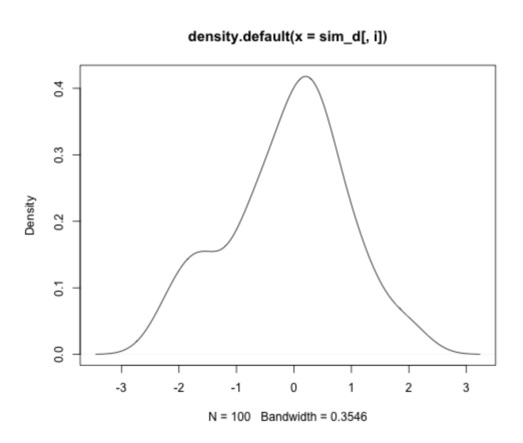
densities <- vector("list", length(sim\_d))</pre>

Calculate all the densities

```
for(i in seq_along(densities)) {
    densities[[i]] <- density(sim_d[ ,i])</pre>
str(densities)
## List of 21
## $ :List of 7
## ..$ x : num [1:512] -3.45 -3.44 -3.42 -3.41 -3.4 ...
## ..$ y : num [1:512] 0.000173 0.000195 0.000219 0.000245 0.00027
## ..$ bw : num 0.355
## ..$ n : int 100
## ..$ call : language density.default(x = sim_d[, i])
## ..$ data.name: chr "sim d[, i]"
## ..$ has.na : logi FALSE
   ..- attr(*, "class") = chr "density"
##
   $:List of 7
##
## ..$ x : num [1:512] -3.39 -3.38 -3.36 -3.35 -3.33 ...
## ..$ y : num [1:512] 0.000261 0.000296 0.000334 0.000377 0.00042
## ..$ bw : num 0.405
## ..$ n : int 100
## ..$ call : language density.default(x = sim_d[, i])
```

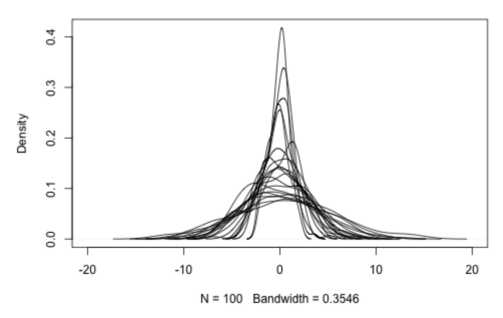
• Next, plot the first density

#### plot(densities[[1]])



• Finally, loop through all the other densities

#### density.default(x = sim\_d[, i])



# Skipping iterations

- On the prior slide, I set the index to skip over the first by using seq(2, length(densities))
- Alternatively, the loop could have been written like this

```
plot(densities[[1]], xlim = c(-20, 20))

for(i in seq_along(densities)) {
    if(i == 1) next
    lines(x = densities[[i]]$x,
        y = densities[[i]]$y)
}
```

# Breaking loops

• Similarly, if a condition is met, you may want to break out of the loop

```
set.seed(1)

rand_unif <- vector("double", 10)

for(i in seq_along(rand_unif)) {
    rand_unif[i] <- runif(1, 0, 10)
    if(any(rand_unif > 5)) {
        break
    }
}

rand_unif
```

# \*apply

# lapply

- One of numerous functionals in R
- A functional "takes a function as an input and returns a vector as output" (adv-r, Chpt 9)
- lapply will always return a list

## Revisiting our simulation with

### n = 10

Our for loop version

```
increments <- seq(1, 5, by = 0.2)
simulated <- vector("list", length(increments))

for(i in seq_along(simulated)) {
    simulated[[i]] <- rnorm(10, 0, increments[i])
}
simulated</pre>
```

```
## [[1]]
## [1] 1.329799263 1.272429321 0.414641434 -1.539950042 -0.928567035 -0
## [7] -0.005767173 2.404653389 0.763593461 -0.799009249
##
## [[2]]
## [[2]]
## [1] -1.3771884 -0.3473539 -0.3590581 -0.4938130 0.3026681 -1.0703054
## [8] -1.4850461 -0.2691215 0.4528748
##
## [[3]]
## [1] 0.18667091 1.12586531 -0.07994948 0.70505116 1.52007711 30.9673
```

# The lapply version

##

## [[6]]

```
increments \leftarrow seq(1, 5, by = 0.2)
sim l <- lapply(increments, function(sd) rnorm(10, 0, sd))
sim l
## [[1]]
## [1] -1.06620017 -0.23845635 1.49522344 1.17215855 -1.45770721 0.0950
## [8] -1.62436453 1.40856336 -0.54176036
##
## [[2]]
## [1] 0.33439767 -0.23276729 1.89138982 -1.77065716 -0.17352985 -1.1438
## [8] 2.67511464 -1.81739641 -0.07404891
##
## [[3]]
## [1] -0.2061791 2.1582303 -1.3745979 0.6952094 2.3757270 -0.3650308 -
## [8] -0.2256499 0.7018506 -1.4189555
##
## [[4]]
## [1] 2.583603577 0.009027176 -4.647838497 -1.771463710 2.476107092 -1
## [7] -0.162405516  0.068240400 -2.554748823  0.785547796
##
## [[5]]
## [1] 0.7588861 3.3730270 1.8621258 0.1472586 -0.1485428 1.0909322 -
## [8] 0.1897585 0.6351741 0.9907080
```

# Some more examples

### Loop through a data frame

• Remember – a data frame is a list. We can loop through it easily

```
library(palmerpenguins)
lapply(penguins, is.double)
```

```
## $species
## [1] FALSE
##
## $island
## [1] FALSE
##
## $bill_length_mm
## [1] TRUE
##
## $bill_depth_mm
## [1] TRUE
##
## $flipper_length_mm
## [1] FALSE
```

#### lapply(mtcars, mean)

```
## $mpg
## [1] 20.09062
##
## $cyl
## [1] 6.1875
##
## $disp
## [1] 230.7219
##
## $hp
## [1] 146.6875
##
## $drat
## [1] 3.596563
##
## $wt
## [1] 3.21725
##
## $qsec
## [1] 17.84875
##
## $vs
## [1] 0.4375
##
## $am
## [1] 0.40625
##
## $gear
```

## Add a condition

```
lapply(penguins, function(x) {
   if(is.numeric(x)) {
      mean(x, na.rm = TRUE)
   }
})
```

```
## $species
## NULL
##
## $island
## NULL
##
## $bill_length_mm
## [1] 43.92193
##
## $bill depth mm
## [1] 17.15117
##
## $flipper_length_mm
## [1] 200.9152
##
## $body_mass_g
## [1] 4201.754
##
## $sex
```

## Add a second condition

```
lapply(penguins, function(x) {
    if(is.numeric(x)) {
       return(mean(x, na.rm = TRUE))
    }
    else if(is.character(x) | is.factor(x)) {
       return(table(x))
    }
})
```

```
## $species
## X
## Adelie Chinstrap Gentoo
##
        152
                   68
                             124
##
## $island
## x
##
                Dream Torgersen
     Biscoe
##
        168
                   124
                              52
##
## $bill length mm
##
  [1] 43.92193
##
## $bill depth mm
   [1] 17.15117
##
```

# Passing arguments

There's missing data, so this won't work

#### head(airquality)

```
## Ozone Solar.R Wind Temp Month Day
## 1 41 190 7.4 67 5 1
## 2 36 118 8.0 72 5 2
## 3 12 149 12.6 74 5 3
## 4 18 313 11.5 62 5 4
## 5 NA NA 14.3 56 5 5
## 6 28 NA 14.9 66 5 6
```

#### lapply(airquality, mean)

```
## $Ozone
## [1] NA
##
## $Solar.R
## [1] NA
##
## $Wind
## [1] 9.957516
##
```

#### But this will.

#### lapply(airquality, mean, na.rm = TRUE)

```
## $Ozone
## [1] 42.12931
##
## $Solar.R
## [1] 185.9315
##
## $Wind
## [1] 9.957516
##
## $Temp
## [1] 77.88235
##
## $Month
## [1] 6.993464
##
## $Day
## [1] 15.80392
```

#### Alternative notation

The prior code could also be written like this

```
lapply(airquality, function(x) mean(x, na.rm = TRUE))
```

```
## $Ozone
## [1] 42.12931
##
## $Solar.R
## [1] 185.9315
##
## $Wind
## [1] 9.957516
##
## $Temp
## [1] 77.88235
##
## $Month
## [1] 6.993464
##
## $Day
## [1] 15.80392
```

# Simulation again

```
lapply(seq(1, 5, 0.2), rnorm, n = 10, mean = 0)
```

```
## [[1]]
## [1] -0.02516264 -0.16367334 0.37005975 -0.38082454 0.65295237 2.0613
## [8] 0.58407712 -0.72275312 -0.62916466
##
## [[2]]
## [1] -2.1794473 -0.3111469 0.4015587 -1.7126011 2.3263539 -0.9114363 -
## [8] -0.1368609 2.8222280 1.9155850
##
## [[31]
## [1] 1.7884237 1.1045592 0.6460515 -0.6132968 -2.1109298 -3.1121246 -
## [8] -2.4958643 -1.3830868 1.0198842
##
## [[4]]
## [1] -1.4154959 -2.4615063 -1.6710007 -2.7490179 1.2860121 -2.4028595 -
## [8] 0.9271338 1.9224409 3.0302573
##
## [[5]]
## [1] -3.1684074 1.6641842 -1.0017759 -0.3250514 2.6053925 -1.0928366
## [8] -0.1684038 -0.8821553 2.5391869
##
## [[6]]
## [1] -0.4491476 -0.4249910 1.3927569 1.8303650 -1.8467486 2.2937465 -
## [8] -1.7728866 -4.6662734 -0.2909816
##
```

# Operations by group

# Mimic dplyr::group\_by

```
by_cyl <- split(mtcars, mtcars$cvl)</pre>
str(by_cyl)
```

```
## List of 3
##
    $ 4:'data.frame': 11 obs. of 11 variables:
##
  ..$ mpg : num [1:11] 22.8 24.4 22.8 32.4 30.4 33.9 21.5 27.3 26 30.4 .
##
   ..$ cyl : num [1:11] 4 4 4 4 4 4 4 4 4 ...
##
   ..$ disp: num [1:11] 108 146.7 140.8 78.7 75.7 ...
##
   ..$ hp : num [1:11] 93 62 95 66 52 65 97 66 91 113 ...
##
   ..$ drat: num [1:11] 3.85 3.69 3.92 4.08 4.93 4.22 3.7 4.08 4.43 3.77
   ..$ wt : num [1:11] 2.32 3.19 3.15 2.2 1.61 ...
##
##
    ..$ qsec: num [1:11] 18.6 20 22.9 19.5 18.5 ...
##
    ..$ vs : num [1:11] 1 1 1 1 1 1 1 0 1 ...
##
    ..$ am : num [1:11] 1 0 0 1 1 1 0 1 1 1 ...
##
   ..$ qear: num [1:11] 4 4 4 4 4 4 3 4 5 5 ...
##
     ..$ carb: num [1:11] 1 2 2 1 2 1 1 1 2 2 ...
##
    $ 6:'data.frame': 7 obs. of 11 variables:
##
    ..$ mpg : num [1:7] 21 21 21.4 18.1 19.2 17.8 19.7
##
    ..$ cyl : num [1:7] 6 6 6 6 6 6 6
##
    ..$ disp: num [1:7] 160 160 258 225 168 ...
##
    ..$ hp : num [1:7] 110 110 110 105 123 123 175
##
    ..$ drat: num [1:7] 3.9 3.9 3.08 2.76 3.92 3.92 3.62
##
    ..$ wt : num [1:7] 2.62 2.88 3.21 3.46 3.44 ...
##
    ..$ qsec: num [1:7] 16.5 17 19.4 20.2 18.3 ...
##
     ..$ vs : num [1:7] 0 0 1 1 1 1 0
```

#### lapply(by\_cyl, function(x) mean(x\$mpg))

```
## $`4`
## [1] 26.66364
##
## $`6`
## [1] 19.74286
##
## $`8`
## [1] 15.1
```

#### Your turn

Try splitting the penguins dataset by species and calculating the average bill\_length\_mm

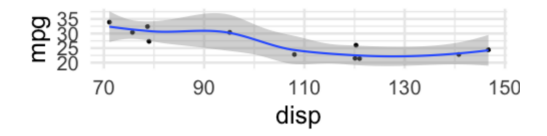


# 

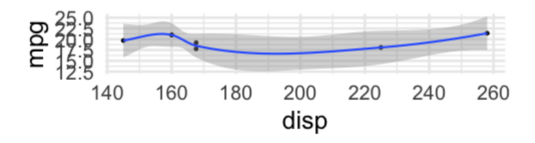
# Produce separate plots

```
lapply(by_cyl, function(x) {
    ggplot(x, aes(disp, mpg)) +
        geom_point() +
        geom_smooth()
})
```

## \$`4`



## ## \$`6`



## ## \$`8`

#### Your turn

Produce separate plots of the relation between bill\_length\_mm and body\_mass\_g



# Saving

- You can extend this example further by saving the plot outputs to an object, then looping through that object to save the plots to disk.
- Using functionals, this would require parallel iterations, which we'll cover later (need to loop through plots and a file name)
- Could extend it fairly easily with a for loop

# Saving w/for loop

Save plots to an object (list)

```
plots <- lapply(by_cyl, function(x) {
    ggplot(x, aes(disp, mpg)) +
        geom_point() +
        geom_smooth()
})</pre>
```

Specify file names/directory

```
#dir.create(here::here("plots"))
filenames <- here::here(
    "plots",
    paste0("cyl", names(by_cyl), ".png")
)
filenames</pre>
```

```
## [1] "/Users/daniel/Teaching/data_sci_specialization/2021-22/c3-fp-2022/g
## [2] "/Users/daniel/Teaching/data_sci_specialization/2021-22/c3-fp-2022/g
## [3] "/Users/daniel/Teaching/data_sci_specialization/2021-22/c3-fp-2022/g
```

# Saving

```
for(i in seq_along(plots)) {
    ggsave(filenames[i], # single bracket
        plots[[i]], # double bracket
        device = "png",
        width = 6.5,
        height = 8)
}
```

# You try!



# More than one groups

Let's say we wanted to create a plot of bill length vs bill depth by for each combination of species/island.

Just split by both! Pass both variables as a list().

```
splt2 <- split(
  penguins,
  list(penguins$species, penguins$island)
)</pre>
```

#### Inspect it

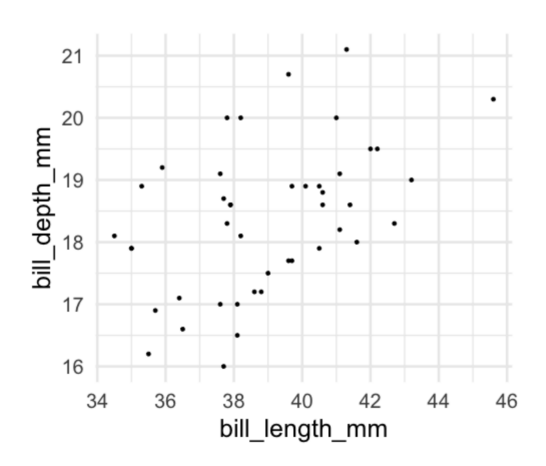
#### splt2

```
## $Adelie.Biscoe
## # A tibble: 44 × 8
##
     species island bill length mm bill depth mm flipper length mm body ma
                             <dbl>
                                           <dbl>
##
   <fct>
            <fct>
                                                             <int>
##
   1 Adelie Biscoe
                              37.8
                                            18.3
                                                               174
## 2 Adelie Biscoe
                              37.7
                                            18.7
                                                              180
                                           19.2
##
   3 Adelie Biscoe
                            35.9
                                                              189
##
   4 Adelie Biscoe
                              38.2
                                           18.1
                                                              185
##
   5 Adelie Biscoe
                              38.8
                                           17.2
                                                              180
##
                              35.3
                                           18.9
   6 Adelie Biscoe
                                                              187
## 7 Adelie Biscoe
                                           18.6
                             40.6
                                                              183
   8 Adelie Biscoe
##
                              40.5
                                           17.9
                                                              187
##
   9 Adelie Biscoe
                              37.9
                                           18.6
                                                              172
## 10 Adelie Biscoe
                              40.5
                                            18.9
                                                              180
## # ... with 34 more rows, and 1 more variable: year <int>
##
## $Chinstrap.Biscoe
## # A tibble: 0 × 8
## # ... with 8 variables: species <fct>, island <fct>, bill length mm <dbl>,
## # bill depth mm <dbl>, flipper length mm <int>, body mass g <int>, sex
####
     year <int>
##
## $Gentoo.Biscoe
## # A tibble: 124 × 8
```

# Create plots

```
plots2 <- lapply(splt2, function(x) {
   ggplot(x, aes(bill_length_mm, bill_depth_mm)) +
     geom_point()
})</pre>
```

#### plots2[[1]]



plots2[[2]]

bill\_depth\_mm

bill\_length\_mm

```
plots2[[3]]
```

## Warning: Removed 1 rows containing missing values (geom\_point).

#### Uh oh

Let's get rid of our empty plots - how? Ideas?

Create a logical vector that checks the number of rows.

We'll do this in a moment.

# Variants of lapply

#### sapply

- Will try to simplify the output, if possible. Otherwise it will return a list.
- Fine for interactive work, but I strongly recommend against it if writing a function (difficult to predict the output)

#### vapply

- Strict you specify the output
- Use if writing functions (or just always stick with lapply), or consider jumping to {purrr} (next week)

### Examples

#### Our simulation

##

```
sim_s \leftarrow sapply(seq(1, 5, by = 0.2), function(x) {
   rnorm(10, 0, x)
 })
 class(sim s)
## [1] "matrix" "array"
dim(sim s)
## [1] 10 21
 sim s
```

```
## [2,] 0.002415809 -1.60656089 -1.4772590 -1.0498872 -4.39616038 -3.0463
## [3,] 0.509665571 0.82578723 -1.0263566 1.5350309 1.04457363 -0.8439
## [4,] -1.084720001 0.08553678 0.2952702 2.4896842 1.17909360 2.7218
## [5,] 0.704832977 2.62770283 -1.3984890 -1.6652743 -0.54811591 603.5075
```

## [1,] -2.939773695 -0.38696441 -2.0067288 -0.6857975 0.26119194 -2.6882

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

#### sapply(penguins, is.double)

```
##
             species
                                  island
                                            bill length mm
                                                                 bill depth mm
##
               FALSE
                                                       TRUE
                                                                          TRUE
                                   FALSE
## flipper length mm
                            body mass g
                                                        sex
                                                                          year
##
               FALSE
                                   FALSE
                                                      FALSE
                                                                         FALSE
```

Now that it's a vector we can easily use it for subsetting

#### head(penguins)

```
## # A tibble: 6 × 8
## species island bill length mm bill depth mm flipper length mm body
                             <dbl>
## <fct> <fct>
                                         <dbl>
                                                          <int>
## 1 Adelie Torgersen
                             39.1
                                          18.7
                                                           181
## 2 Adelie Torgersen
                             39.5
                                          17.4
                                                           186
## 3 Adelie Torgersen
                             40.3
                                          18
                                                           195
## 4 Adelie Torgersen
                         NA
                                          NA
                                                           NA
## 5 Adelie Torgersen
                        36.7
                                          19.3
                                                           193
## 6 Adelie Torgersen
                      39.3
                                          20.6
                                                           190
## # ... with 1 more variable: year <int>
```

#### head( penguins[ ,sapply(penguins, is.double)] )

```
## # A tibble: 6 × 2
     bill length mm bill depth mm
##
                               \langle d\overline{b}1 \rangle
##
               <dbl>
                               18.7
## 1
                39.1
## 2
                39.5
                                17.4
## 3
                40.3
                                18
## 4
                NA
                                NA
## 5
            36.7
                               19.3
## 6
                39.3
                                20.6
```

# Challenge

Can you make return the opposite? In other words – all those that are *not* double?

02:00

#### head( penguins[ ,!sapply(penguins, is.double)] )

```
## # A tibble: 6 × 6
## species island flipper length mm body mass g sex
                                                        year
## <fct>
           <fct>
                                <int>
                                           <int> <fct>
                                                       <int>
## 1 Adelie
                                            3750 male
                                                        2007
           Torgersen
                                  181
## 2 Adelie Torgersen
                                 186
                                            3800 female 2007
## 3 Adelie Torgersen
                                 195
                                           3250 female 2007
## 4 Adelie Torgersen
                                             NA <NA> 2007
                                 NA
## 5 Adelie
                                 193
                                           3450 female 2007
           Torgersen
## 6 Adelie
                                            3650 male
                                                       2007
           Torgersen
                                 190
```

# Clean up our plots

Can you recreate the plots while omitting the empty ones now?

Try!

03:00

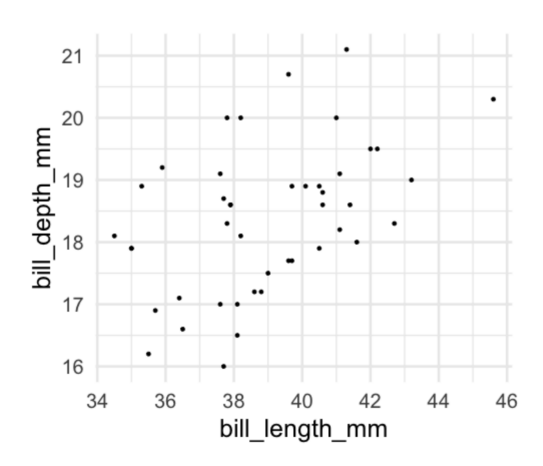
#### Remove zero-row dfs

```
# check if n rows > 0
keep <- sapply(splt2, function(x) nrow(x) > 0)
keep
##
        Adelie.Biscoe Chinstrap.Biscoe
                                                Gentoo.Biscoe
                                                                      Adeli
##
                  TRUE
                                     FALSE
                                                          TRUE
##
      Chinstrap.Dream
                        Gentoo.Dream
                                              Adelie.Torgersen Chinstrap.To
##
                  TRUE
                                                          TRUE
                                     FALSE
##
     Gentoo.Torgersen
##
                 FALSE
# Use this to subset the list
splt3 <- splt2[keep]</pre>
```

## Recreate plots

```
plots3 <- lapply(splt3, function(x) {
   ggplot(x, aes(bill_length_mm, bill_depth_mm)) +
     geom_point()
})</pre>
```

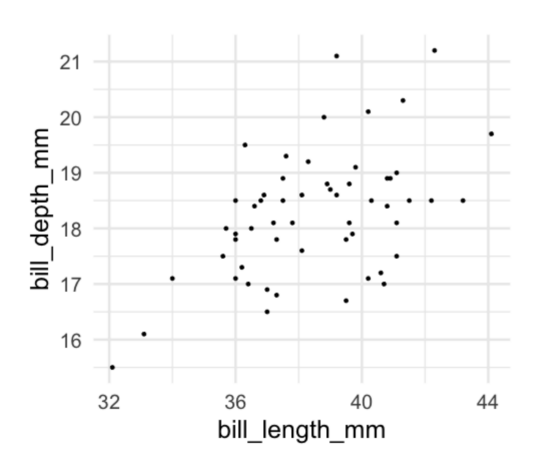
#### plots3[[1]]



```
plots3[[2]]
```

## Warning: Removed 1 rows containing missing values (geom\_point).

#### plots3[[<mark>3</mark>]]



# vapply

 As you can probably see, simplifying can be really helpful for interactive work.



- Not ideal for programmatic work need to be able to reliably predict the output
- vapply solves this issue.

## Error in vapply(penguins, is.double, FUN.VALUE = character(1)): values n
## but FUN(X[[1]]) result is type 'logical'

```
vapply(penguins, is.double, FUN.VALUE = logical(1))
```

```
##
                                         bill length mm
            species
                              island
                                                            bill depth mm
##
              FALSE
                               FALSE
                                                   TRUE
                                                                     TRUE
## flipper length mm
                         body mass g
                                                    sex
                                                                     year
##
                                FALSE
               FALSE
                                                  FALSE
                                                                    FALSE
```

### Coercion with vapply

## flipper\_length\_mm

##

• If it can coerce the vector without loss of information, it will

body\_mass\_g

```
vapply(penguins, is.double, FUN.VALUE = double(1))

## species island bill_length_mm bill_depth_mm
## 0 0 1
```

sex

year

## Count missing data

```
vapply(airquality, function(col) {
   sum(is.na(col))
   },
   FUN.VALUE = double(1)
)

## Ozone Solar.R Wind Temp Month Day
## 37 7 0 0 0 0
```

## sapply alternative

For interactive work, the code on the previous slide is maybe too much. Could be reduced to

```
sapply(airquality, function(col) sum(is.na(col)))
## Ozone Solar.R Wind Temp Month Day
## 37 7 0 0 0 0
```

### Summary

- for loops are incredibly flexible and there's nothing inherently "wrong" about them
  - Do require more text, and often repetitive text, which can lead to errors/bugs
  - The flexibility can actually be more of a curse than a blessing

### Summary

- The lapply family of functions help put the focus on a given function, and what values are being looped through the function
  - lapply will always return a list
  - sapply will try to simplify, which is problematic for programming, but fine for interactive work
  - vapply is strict, and will only return the type specified

# 

Quickly

# apply(x, dimension, function, function\_args)

- x The thing to loop over (usually a matrix or data frame)
- dimension
  - 1 = apply the function to each row
  - 2 = apply the function to each column
  - $\circ$  n = apply to nth dimension of an array (rare for the work I do)

### Rows example

```
people <- data.frame(
  first = c("Frederick", "Anna", "Julia"),
  last = c("Douglass", "Murray", "Griffiths")
)
people</pre>
```

```
## first last
## 1 Frederick Douglass
## 2 Anna Murray
## 3 Julia Griffiths
```

Suppose we wanted a new column that was the first and last name.

### We might try this

```
people %>%
  mutate(full_name = paste(first, last, collapse = " "))
```

... but it pastes together the full columns

#### Instead, do it by row

```
apply(people, 1, paste, collapse = " ")
## [1] "Frederick Douglass" "Anna Murray" "Julia Griffiths"
Or within a mutate() call
people %>%
  mutate(full_name = apply(people, 1, paste, collapse = " "))
## first last full name
## 1 Frederick Douglass Frederick Douglass
## 2 Anna Murray Anna Murray
## 3 Julia Griffiths Julia Griffiths
```

### Column example

Back to the airquality example - standardize all columns.

Notice it returns a matrix though, which is less than ideal.

```
apply(airquality, 2, function(x) {
  as.numeric(scale(x))
})
```

```
Ozone Solar.R Wind
##
                                                   Temp
                                                              Month
## [1,] -0.03423409 0.0451761540 -0.72594816 -1.14971398 -1.40729432 -1.
## [2,] -0.18580489 -0.7543048742 -0.55563883 -0.62146702 -1.40729432 -1.
## [3,] -0.91334473 -0.4100838759 0.75006604 -0.41016823 -1.40729432 -1.
## [4,] -0.73145977 1.4109562438 0.43783226 -1.67796094 -1.40729432 -1.
## [5,]
                 NA
                               NA 1.23260914 -2.31185730 -1.40729432 -1.
## [6,] -0.42831817
                               NA 1.40291847 -1.25536337 -1.40729432 -1.
## [7,] -0.57988897 1.2555015994 -0.38532950 -1.36101276 -1.40729432 -0.
## [8,] -0.70114561 -0.9652790344 1.09068470 -1.99490912 -1.40729432 -0.
## [9,] -1.03460136 -1.8535912880 2.87893266 -1.78361034 -1.40729432 -0.
##
   [10,]
                 NA 0.0895917667 -0.38532950 -0.93841519 -1.40729432 -0.
##
   [11,] -1.06491552
                               NA -0.86787260 -0.41016823 -1.40729432 -0.
## [12,] -0.79208809 0.7780337632 -0.07309573 -0.93841519 -1.40729432 -0.
##
   [13,] -0.94365889 1.1555664709 -0.21502017 -1.25536337 -1.40729432 -0.
##
   [14,] -0.85271641 0.9779040202 0.26752293 -1.04406459 -1.40729432 -0.
```

### Last bit

There are other loops, like tapply(), but I tend to not use them much (instead just use lapply(split(x)).

All of this stuff takes practice, both to understand how it works and to start to see use cases

Careful not to get into too deep of nested loops — if you're nesting beyond two levels (and I honestly never go beyond one anymore) there's probably better ways to approach it.

Lab 1