

## **Getting started (again)**

Open RStudio
Open your NRI project
Open a **new** script for today:
File > New File > R Script

Make sure to load packages at the top:
library(tidyverse)
library(palmerpenguins)

# Take Heart! I KNOW\_LOTS! HOW MUCH I THINK I KNOW ABOUT R I KNOW ABOUT R Artwork by @allison horst on Twitter -- "Knowing so little never felt so fun. #rstats" 3/50

# **Exploring everything at once**

# Visualize with ggpairs()

• From **GGally** package

```
library(GGally)
penguins_sub <- select(penguins, -sex, -island, -year)
ggpairs(penguins_sub)</pre>
```

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# Side Note: **tidyverse** functions

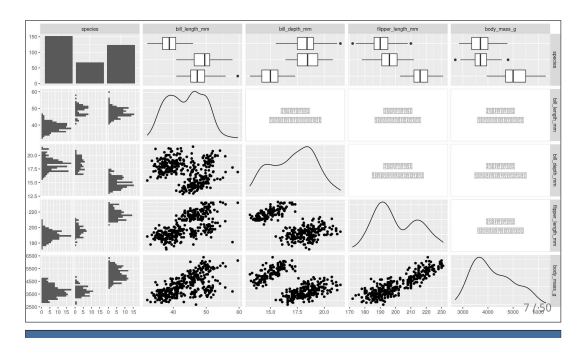
• From **GGally** 

```
library(GGally)

penguins_sub <- select(penguins, -sex, -island, -year)
ggpairs(penguins_sub)</pre>
```

#### select()

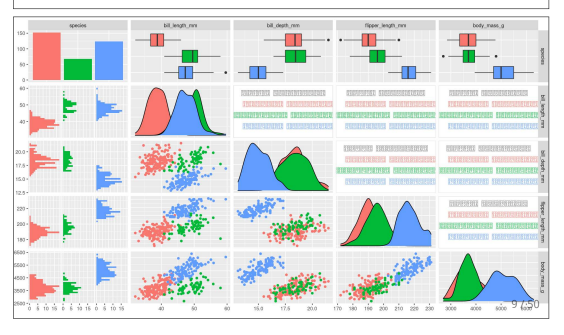
- tidyverse functions always start with the data, followed by other arguments
- you can reference any column from 'data'
- select() chooses columns to keep or to remove (with -)



# Visualize with ggpairs()

library(GGally)
ggpairs(select(penguins, -sex, -island, -year), aes(colour = species))

ggpairs() builds on ggplot() so we can use an aes() specification



#### Summarize with **skim()**

#### skim() from skimr package

```
library(skimr)
 skim(penguins)
## -- Data Summary --
## Name
## Number of rows
                                 344
## Number of columns
##
## Column type frequency:
## factor
## numeric
## Group variables
##
## -- Variable type: factor --
## skim_variable n_missing complete_rate ordered n_unique top_counts
## 1 species 0 1 FALSE 3 Ade: 152, Gen: 124, Chi: 68
## 2 island 0 1 FALSE 3 Bis: 168, Dre: 124, Tor: 52
## 3 sex 11 0.968 FALSE 2 mal: 168, fem: 165
                                                                                                                    10 / 50
##
```

#### Summarize with **skim()**

#### skim() from skimr package

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#### Summarize with **skim()**

#### skim() from skimr package

library(skimr)
skim(penguins)

Your Turn!

#### Summarize with **skim()**

• From **skimr** package

```
penguins_sp <- group_by(penguins, species)
skim(penguins_sp)</pre>
```

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## Side Note: tidyverse functions

• From **skimr** package

```
penguins_sp <- group_by(penguins, species)
skim(penguins_sp)</pre>
```

#### group\_by()

- tidyverse functions always start with the data, followed by other arguments
- you can reference any column from 'data'
- group\_by() assigns grouping to a data frame. Here, we group penguins by species

#### xtra:

In the console look at penguins (type in penguins and hit enter), and then look at penguins\_sp (type in penguins\_sp and it enter). How does the output differ? (Hint very little! But there is one difference...)

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## Summarize with **skim()**

#### skim() from skimr package

```
penguins_sp <- group_by(penguins, species)</pre>
 skim(penguins_sp)
## -- Data Summary -
## Name
                                   skimp
## Number of rows
                                  344
## Number of columns
## Column type frequency:
## factor
## numeric
## Group variables
## -- Variable type: factor -
## skim_variable species n_missing complete_rate ordered n_unique top_counts
## 1 island Adelie 0 1 FALSE 3 Dre: 56, To
## 2 island Chinstrap 0 1 FALSE 1 Dre: 68, Bi
                     Adelie 0 1 FALSE 3 Dre: 56, Tor: 52, Bis: 44
Chinstrap 0 1 FALSE 1 Dre: 68, Bis: 0, Tor: 0
Gentoo 0 1 FALSE 1 Bis: 124, Dre: 0, Tor: 0
## 2 island
                      Gentoo 6
## 3 island
                                                                        2 fem: 73, mal: 73
                                                                                                                             14/50
## 4 sex
                 Adelie
                                                  0.961 FALSE
```

## 

#### Summarize with **skim()**

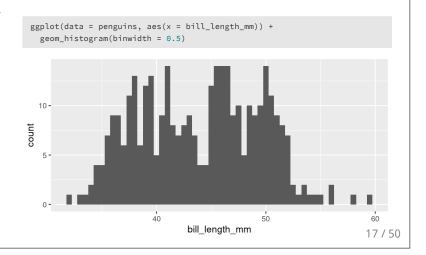
```
## — Variable type: factor ## 1 island Adelie 0 1 FALSE 3 Dre: 56, Tor: 52, Bis: 44
## 2 island Chinstrap 0 1 FALSE 1 Bis: 124, Dre: 0, Tor: 0
## 4 sex Adelie 6 0.961 FALSE 2 fem: 73, mal: 73
## 5 sex Chinstrap 0 1 FALSE 2 fem: 73, mal: 73
## 6 sex Gentoo 5 0.960 FALSE 2 mal: 61, fem: 58
##
## — Variable type: numeric — wissing complete_rate mean sd p0 p25 p50 p75
## 1 bill_length_mm Adelie 1 0.993 38.8 2.66 32.1 36.8 38.8 40.8
## 2 bill_length_mm Gentoo 1 0.992 47.5 3.08 40.9 45.3 47.3 49.6
## 3 bill_length_mm Adelie 1 0.993 18.3 1.22 15.5 17.5 18.4 19
## 5 bill_depth_mm Adelie 1 0.993 18.3 1.22 15.5 17.5 18.4 19
## 5 bill_depth_mm Chinstrap 0 1 18.4 1.14 16.4 17.5 18.4 19.4
## 6 bill_depth_mm Gentoo 1 0.992 15.0 0.981 13.1 14.2 15 15.7
```

## **Exploring variable by variable**

# Visualize with ggplot()

#### From last week...

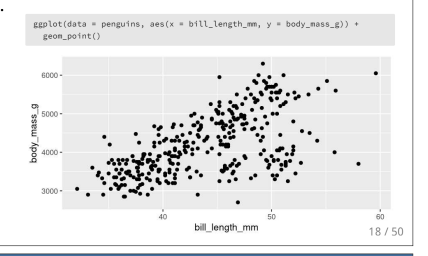
• Histograms



# Visualize with ggplot()

#### From last week...

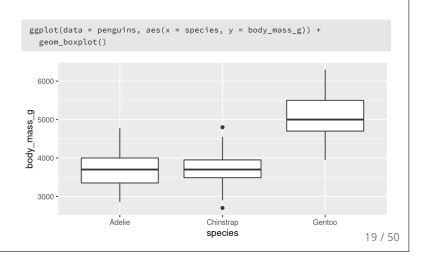
- Histograms
- Scatterplots



# Visualize with ggplot()

#### From last week...

- Histograms
- Scatterplots
- Boxplots



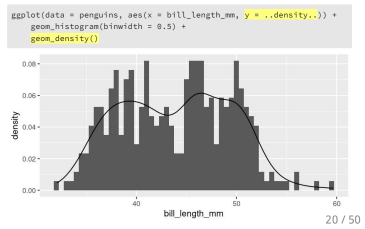
## Visualize with ggplot()

#### **Histogram with Density**

- Default uses counts
- Here use density

y = ..density..

- Same as density curve geom\_density()
- Use to assess shape and distribution of data



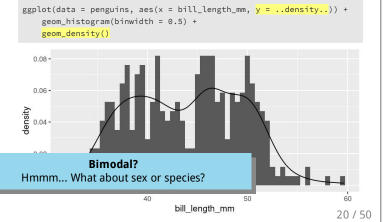
#### Visualize with ggplot()

#### **Histogram with Density**

- Default uses counts
- Here use density

y = ..density..

- Same as density curve geom\_density()
- Use to assess shape and distribution of data



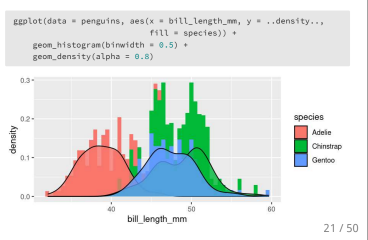
## Visualize with ggplot()

#### **Histogram with Density**

- Default uses counts
- Here use density

y = ..density..

- Same as density curve geom\_density()
- Use to assess shape and distribution of data

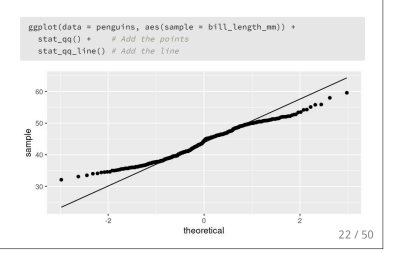


#### Visualize with ggplot() **Histogram with Density** ggplot(data = penguins, aes(x = bill\_length\_mm, y = ..density.., • Default uses counts fill = species)) + • Here use density $geom\_histogram(binwidth = 0.5) +$ y = ..density.. geom\_density(alpha = 0.8) • Same as density curve geom\_density() • Use to assess shape and species distribution of data 0.2 density Adelie Chinstrap Gentoo **Extra** Perhaps sex plays a role in those remaining patterns...? bill\_length\_mm 21 / 50

## Visualize with ggplot()

#### **QQ Norm plots**

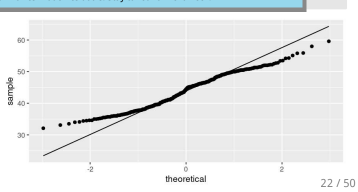
 Assess whether data follows normal distribution



# Visualize with **ggplot()**

#### **QQ Norm plots**

Asses
 Here we are **NOT** assessing assumptions of normality for a model follow
 That involves model residuals: Stay tuned for next week!



## Summarize with summarize() Ha!

• From dplyr package (part of tidyverse)

```
summarize(penguins,
    mean_mass = mean(body_mass_g),
    sd_mass = sd(body_mass_g),
    median_mass = median(body_mass_g))
```

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#### Side Note: tidyverse functions

• From dplyr package (part of tidyverse)

```
summarize(penguins,
    mean_mass = mean(body_mass_g),
    sd_mass = sd(body_mass_g),
    median_mass = median(body_mass_g))
```

#### summarize()

- tidyverse functions always start with the data, followed by other arguments
- you can reference any column from 'data'
- summarize() creates a data frame with new columns (summarizes your data)

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## Summarize with **summarize()**

• From dplyr package (part of tidyverse)

## Summarize with **summarize()**

• From dplyr package (part of tidyverse)

Why all NAs?

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## Summarize with summarize()

mean(), sd(), median()

```
summarize(penguins,
    mean_mass = mean(body_mass_g, na.rm = TRUE),
    sd_mass = sd(body_mass_g, na.rm = TRUE),
    median_mass = median(body_mass_g, na.rm = TRUE))
```

## mean\_mass sd\_mass median\_mass ## <dbl> <dbl> <dbl> ## 1 4202. 802. 4050

## # A tibble: 1 x 3

Need to tell summary statistic functions to remove missing values na.rm = TRUE

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#### Summarize with **summarize()**

mean(), sd(), median(), quantile(), n()\*

\* n() only works inside summarize()/mutate()

## Your Turn: summarize()

#### Calculate summary statistics for Bill Length

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#### Side Note: Removing NAs

• With arguments

```
    na.rm = TRUE (summary stats i.e. mean(), sd())
    na.action = na.exclude (models i.e., lm(), lmer())
```

- You can remove all NAs from your data (drop\_na())
- You can selectively remove NAs from your data (filter())

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## Side Note: Removing NAs

#### **Remove all NAs**

- This removes **every** row that has an **NA** in **any** column
- drop\_na() function from tidyr package (part of tidyverse)

penguins\_no\_na <- drop\_na(penguins)</pre>

• Consider removing columns with lots of NAs first (assuming you don't need them)

```
penguins_no_na <- select(penguins, -sex)
penguins_no_na <- drop_na(penguins_no_na)</pre>
```

## Side Side Note: **tidyverse** functions

• From tidyr package (part of tidyverse)

penguins\_no\_na <- drop\_na(penguins)</pre>

#### drop\_na()

- tidyverse functions always start with the data, followed by other arguments
- here, there are no other arguments

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#### Side Note: Removing NAs

#### Selective remove NAs with filter()

• From dplyr package (part of tidyverse)

filter(penguins, !is.na(body\_mass\_g))

- is.na() checks if there is an NA and returns TRUE if so
- ! turns a TRUE into a FALSE
- filter() only keeps rows that are TRUE
- Thus any row with an NA in body\_mass\_g is removed

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## Side Side Note: **tidyverse** functions

• From dplyr package (part of tidyverse)

filter(penguins, !is.na(body\_mass\_g))

#### filter()

- tidyverse functions always start with the data, followed by other arguments
- you can reference any column from 'data'
- filter() keeps only rows that return TRUE to the logical statements

#### Summarize with **summarize()** (and **group\_by()**)

• Can also use group\_by() to calculate summaries by groups

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#### Summarize with **summarize()** (and **group\_by()**)

• Can also use group\_by() to calculate summaries by groups

```
penguins_sp_sex <- group_by(penguins, species, sex)</pre>
 summarize(penguins sp sex.
           mean_mass = mean(body_mass_g, na.rm = TRUE),
           sd_mass = sd(body_mass_g, na.rm = TRUE),
           {\tt median\_mass = median(body\_mass\_g, na.rm = TRUE))}
## # A tibble: 8 x 5
## # Groups: species [3]
## species sex mean_mass sd_mass median_mass
## <fct> <fct> <dbl> <dbl> <dbl> <dbl> 3369. 269. 3400
## 2 Adelie male 4043. 347. 4000
## 3 Adelie <NA> 3540 477.
## 4 Chinstrap female 3527. 285.
                                              3550
## 5 Chinstrap male 3939. 362. 3950
## 6 Gentoo female 4680. 282.
## 7 Gentoo male 5485. 313.
                                              4700
## 7 Gentoo male 5485. 313.
## 8 Gentoo <NA> 4588. 338.
                                              5500
                                                                                                   35 / 50
                                           4688.
```

#### Summarize with **summarize()** (and **group\_by()**)

• Can also use **group\_by()** to calculate summaries by groups

```
## # A tibble: 8 x 5
                                Where are the decimal points?
## # Groups: species
## species sex mean_mass sd_mass median_mass
## <fct> <fct> <dbl> <dbl> <dbl> ## 1 Adelie female 3369. 269.
                                               3400
## 2 Adelie male 4043. 347.
## 3 Adelie <NA> 3540 477.
## 4 Chinstrap female 3527. 285.
                                   477.
                                               3475
                                                3550
## 5 Chinstrap male 3939. 362.
## 6 Gentoo female 4680. 282.
## 7 Gentoo male 5485. 313.
                                               4700
## 7 Gentoo male 5485. 313.
## 8 Gentoo <NA> 4588. 338.
                                               5500
                                                                                                     35 / 50
                                            4688.
```

## Side Note: Where are the decimal points?

• tibble hides them for easy viewing

```
penguins_sum <- summarize(penguins_sp_sex,</pre>
                            mean_mass = mean(body_mass_g, na.rm = TRUE),
                             sd_mass = sd(body_mass_g, na.rm = TRUE),
                             median_mass = median(body_mass_g, na.rm = TRUE))
 penguins_sum
## # A tibble: 8 x 5
## # Groups: species [3]
## species sex mean_mass sd_mass median_mass
## <fct> <fct> <dbl> <dbl> <dbl>
## 1 Adelie female 3369. 269.
## 2 Adelie male 4043. 347.
## 3 Adelie <NA> 3540 477.
                                                 4000
                                                  3475
## 4 Chinstrap female 3527. 285.
                                               3550
## 5 Chinstrap male 3939. 362.
                                                3950
## 6 Gentoo female 4680. 282.
## 7 Gentoo male 5485. 313.
## 8 Gentoo <NA> 4588. 338.
                                                 5500
                                                                                                         36 / 50
```

#### Side Note: Where are the decimal points?

• tibble hides them for easy viewing

```
penguins_sum <- summarize(penguins_sp_sex,</pre>
                             mean_mass = mean(body_mass_g, na.rm = TRUE),
                             sd_mass = sd(body_mass_g, na.rm = TRUE),
                              median_mass = median(body_mass_g, na.rm = TRUE))
 penguins_sum
## # A tibble: 8 x 5
                                                                                    Note
## # Groups: species [3]
                                                                       If you want to keep the output, you
## species sex mean_mass sd_mass median_mass ## <fct> <fct> <dbl> <dbl> <dbl> <dbl>
                                                                       need to assign (<-) it to an object.
## 1 Adelie female 3369. 269.
                                                                            Here, penguins_sum
## 2 Adelie male 4043. 347.
                                                  4000
                            3540
                                      477.
                                                   3475
## 3 Adelie
                <NA>
## 4 Chinstrap female 3527. 285.
                                                 3550
## 5 Chinstrap male 3939. 362.

## 6 Gentoo female 4680. 282.

## 7 Gentoo male 5485. 313.

## 8 Gentoo <NA> 4588. 338.
                                                 3950
                                                   4700
                                              5500
                                                                                                            36 / 50
```

#### Side Note: Where are the decimal points?

• as.data.frame() to see the raw data

```
## species sex mean_mass sd_mass median_mass
## 1 Adelie female 3368.836 269.3801 3400.0
## 2 Adelie male 4043.493 346.8116 4000.0
## 3 Adelie <NA> 3540.000 477.1661 3475.0
## 4 Chinstrap female 3527.206 285.3339 3550.0
## 5 Chinstrap male 3938.971 362.1376 3950.0
## 6 Gentoo female 4679.741 281.5783 4700.0
## 7 Gentoo male 5484.836 313.1586 5500.0
## 8 Gentoo <NA> 4587.500 338.1937 4687.5
```

• Or click on the name in the Environment pane

## Side Note: Where are all my data?

```
penguins
## # A tibble: 344 x 8
## species island bill_length_mm bill_depth_mm flipper_length_... body_mass_g
     <fct> <fct> <dbl> <dbl> <int>
                          39.1
39.5
                                                        181
186
## 1 Adelie Torge...
                                        18.7
                                        17.4
## 2 Adelie Torge...
                                                                   3800
## 3 Adelie Torge...
                                  NA
19.3
20.6
17.8
                          NA
36.7
                                                        NA
193
## 4 Adelie Torge...
## 5 Adelie Torge...
                                                                    3450
                          39.3
                                                        190
## 6 Adelie Torge...
                                                                   3650
                                                        181
## 7 Adelie Torge...
                          38.9
                                                                   3625
## 10 Adelie Torge... 42
## # ... with 334 more rows 254
   8 Adelie Torge...
                            39.2
                                        19.6
                                       18.1
                                                        193
                                                                   3475
                                       20.2
                                                        190
## # ... with 334 more rows, and 2 more variables: sex <fct>, year <int>
```

... with 334 more rows, and 2 more variables: sex <fct>, year <int>

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## Side Note: Where are all my data?

```
print(penguins, n = Inf)
## # A tibble: 344 x 8
   species island bill_length_mm bill_depth_mm flipper_length_... body_mass_g
##
     <fct> <fct> <fct> <dbl> <int> <int>
                    1 Adelie Torge...
                                                   181
                                                  186
  2 Adelie Torge...
                                                            3800
  3 Adelie Torge...
                                                           3250
                                                 NA
193
   4 Adelie Torge…
##
  5 Adelie Torge...
                                                            3450
  6 Adelie Torge...
                                                190
                                                          3650
                                                           3625
##
  7 Adelie Torge…
                                                 181
    8 Adelie Torge...
                                                  193
## 9 Adelie Torge...
                                                            3475
## 10 Adelie Torge...
                                                  190
                                                            4250
## 11 Adelie Torge...
                         37.8
                                                   186
                                                            3300
                                    17.1
                         37.8
## 12 Adelie Torge...
                                    17.3
                                                   180
                                                            3700
## 13 Adelie Torge...
                         41.1
                                   17.6
## 14 Adelie Torge...
                         38.6
                                                   191
                                                            3800
                                     21.2
                                                                              39 / 50
## 15 Adelie Torge...
                    34.6
                                21.1
                                                    198
                                                            4400
```

#### Side Note: Where are all my data?

```
as.data.frame(penguins)
      species island bill_length_mm bill_depth_mm flipper_length_mm body_mass_g
                                                                      sex year
       Adelie Torgersen 39.1 18.7 181 3750 male 2007
## 1
## 2
       Adelie Torgersen
                            39.5
                                       17.4
                                                               3800 female 2007
                           40.3
NA
                                      18.0
                                                      195
NA
                                                              3250 female 2007
## 3
       Adelie Torgersen
                                        NA
## 4
       Adelie Torgersen
                                                                NA <NA> 2007
       Adelie Torgersen
                         36.7 19.3
                                                      193 3450 female 2007
                                                      190
                                                               3650 male 2007
       Adelie Torgersen
                          39.3
                                       20.6
## 6
## 7
       Adelie Torgersen
                             38.9
                                        17.8
                                                       181
                                                                3625 female 2007
                           39.2
## 8
       Adelie Torgersen
                                      19.6
                                                      195
                                                               4675 male 2007
                           34.1 18.1
                                                              3475 <NA> 2007
                                                      193
## 9
       Adelie Torgersen
       Adelie Torgersen
                                        20.2
                                                       190
                                                                4250
                                                                     <NA> 2007
                           37.8 17.1
                                                      186
                                                              3300 <NA> 2007
       Adelie Torgersen
## 11
## 12
       Adelie Torgersen
                           37.8 17.3
                                                      180
                                                               3700 <NA> 2007
## 13
                             41.1
                                        17.6
                                                       182
                                                                3200 female 2007
       Adelie Torgersen
## 14
       Adelie Torgersen
                            38.6
                                       21.2
                                                       191
                                                                3800 male 2007
## 15
       Adelie Torgersen
                           34.6
                                                              4400 male 2007
                                                       185
                                                               3700 female 2007
## 16
       Adelie Torgersen
                             36.6
                                        17.8
                                                           3450 female 200<del>/</del>40 / 50
## 17
      Adelie Torgersen
                            38.7
                                        19.0
                                                       195
```

#### Summarize with **summarize()**

#### skewness(), kurtosis()

• From moments package

\* Excess kurtosis would be 0 for a normal distribution, but this functions measures kurtosis

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#### Summarize with **summarize()**

#### **Confidence Intervals**

- By hand!
- 95% Confidence interval ranges from [mean (1.96 SE)] to [mean + (1.96 SE)]
- You can also express this interval as: mean +/- (1.96 \* SE)
- Standard Errors (SE) can be calculated by SD / sqrt(n)

```
summarize(penguins,
    mean_mass = mean(body_mass_g, na.rm = TRUE),
    sd_mass = sd(body_mass_g, na.rm = TRUE),
    n = n(),
    se_mass = sd_mass / sqrt(n),  # Calculate Standard Error
    ci_mass = 1.96 * se_mass,  # CI margin of error
    ci_low_mass = mean_mass - ci_mass,  # The lower range
    ci_high_mass = mean_mass + ci_mass) # The upper range
```

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#### Summarize with **summarize()**

#### **Confidence Intervals**

- By hand!
- 95% Confidence interval ranges from [mean (1.96 SE)] to [mean + (1.96 SE)]
- You can also express this interval as: mean +/- (1.96 \* SE)
- Standard Errors (SE) can be calculated by SD / sqrt(n)

#### Put it All Together

```
penguins_sp <- group_by(penguins, species)</pre>
summarize(penguins_sp,
         mean_mass = mean(body_mass_g, na.rm = TRUE),
          sd_mass = sd(body_mass_g, na.rm = TRUE),
         q25_mass = quantile(body_mass_g, probs = 0.25, na.rm = TRUE),
         median_mass = median(body_mass_g, na.rm = TRUE),
         q75_mass = quantile(body_mass_g, probs = 0.25, na.rm = TRUE),
         n = n(),
         n_no_missing = sum(!is.na(body_mass_g)),
         skew_mass = skewness(body_mass_g, na.rm = TRUE),
         kurt_mass = kurtosis(body_mass_g, na.rm = TRUE),
         se_mass = sd_mass / sqrt(n),
         ci_mass = 1.96 * se_mass,
          ci_low_mass = mean_mass - ci_mass,
          ci_high_mass = mean_mass + ci_mass)
```

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## Put it All Together

```
species mean_mass sd_mass q25_mass median_mass q75_mass n n_no_missing skew_mass kurt_mass
       Adelie 3700.662 458.5661 3350.0 3700 3350.0 152 151 0.28249381 2.405611 cinstrap 3733.088 384.3351 3487.5 3700 3487.5 68 68 0.24194125 3.463681
## 2 Chinstrap 3733.088 384.3351 3487.5
                                                                      123 0.06878276 2.257871
## 3 Gentoo 5076.016 504.1162 4700.0
                                               5000 4700.0 124
## se_mass ci_mass ci_low_mass ci_high_mass
## 1 37.19462 72.90146 3627.761 3773.564
## 2 46.60747 91.35065 3641.738 3824.439
## 3 45.27097 88.73111 4987.285 5164.747
```

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# Put it All Together (Advanced!)

## 3 Adelie Torgersen male 2007 flipper\_length\_mm 181

## 7 Adelie Torgersen female 2007 flipper\_length\_mm 186

## 5 Adelie Torgersen female 2007 bill\_length\_mm

## 6 Adelie Torgersen female 2007 bill\_depth\_mm

#### pivot\_longer() transposes data

• from tidyr package (part of tidyverse)

4 Adelie Torgersen male

```
penguins_long <- pivot_longer(penguins,</pre>
                                 cols = c(bill_length_mm, bill_depth_mm, flipper_length_mm,
 body_mass_g),
                                 names_to = "measurement", values_to = "values")
 penguins_long
## # A tibble: 1.376 x 6
## species island sex year measurement
     <fct> <fct> <fct> <fct> <fct> <int> <chr>
                                                            <dbl>
##
## 1 Adelie Torgersen male 2007 bill_length_mm
## 2 Adelie Torgersen male 2007 bill_depth_mm
                                                             39.1
                                                            18.7
```

39.5

17.4

2007 body\_mass\_g 3750

18

#### Put it All Together (Advanced!)

#### pivot\_longer() transposes data

• from tidyr package (part of tidyverse)

```
## # A tibble: 1,376 x 6
##
    species island sex year measurement
                                                    values
                     <fct> <int> <chr>
      <fct> <fct>
                                                      <dbl>
## 1 Adelie Torgersen male 2007 bill_length_mm
## 2 Adelie Torgersen male 2007 bill_depth_mm
                                                       39.1
                                                      18.7
## 3 Adelie Torgersen male 2007 flipper_length_mm 181
   4 Adelie Torgersen male
                               2007 body_mass_g
## 5 Adelie Torgersen female 2007 bill_length_mm
                                                      39.5
## 6 Adelie Torgersen female 2007 bill_depth_mm
                                                      17.4
## 7 Adelie Torgersen female 2007 flipper_length_mm 186
```

#### **Extra**

Compare penguins to penguins\_long.
Can you see what the pivot\_longer()
function is doing?

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#### Put it All Together (Advanced!)

```
penguins_long_sp <- group_by(penguins_long, species, measurement)
summarize(penguins_long_sp,
    mean = mean(values, na.rm = TRUE),
    sd = sd(values, na.rm = TRUE),
    q25 = quantile(values, probs = 0.25, na.rm = TRUE),
    median = median(values, na.rm = TRUE),
    q75 = quantile(values, probs = 0.25, na.rm = TRUE),
    n = n(),
    n_no_missing = sum(!is.na(values)),
    skew = skewness(values, na.rm = TRUE),
    kurt = kurtosis(values, na.rm = TRUE))</pre>
```

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## Put it All Together (Advanced!)

```
## `summarise()` regrouping output by 'species' (override with `.groups` argument)
## # A tibble: 12 x 11
## # Groups: species [3]
     species measurement
                                            q25 median
                                                              n n_no_missing
kurt
##
     <fct>
                              <dbl> <dbl> <dbl> <dbl> <dbl> <int>
                                                                         <int>
                                                                                <dbl>
<dbl>
## 1 Adelie
             bill_depth_mm
                              18.3 1.22
                                            17.5 18.4 17.5 152
                                                                        151 0.318
2.90
             bill_length_mm
## 2 Adelie
                              38.8 2.66
                                           36.8 38.8 36.8 152
                                                                        151 0.160
2.81
## 3 Adelie
             body_mass_g
                             3701. 459.
                                           3350 3700 3350
2.41
## 4 Adelie
              flipper_length_mm 190.
                                      6.54 186
                                                        186
                                                               152
                                                                          151 0.0865
3.28
## 5 Chinstrap bill_depth_mm
                               18.4 1.14 17.5 18.4 17.5
                                                                68
                                                                           68 0.00673
## 6 Chinstrap bill length mm
                               48.8 3.34 46.3 49.6 46.3
                                                                68
                                                                           68 -0.0886
2.95
                                                                                    48 / 50
```

## All Data vs. Variable by Variable

#### Depends on what you need

- ggpairs() and skim()
  - Lots of data quickly summarized and examined
  - · Less easily customized
- ggplot() and summarize()
  - Take a bit longer to write out
  - Very customizable
  - Can easily include stats not available in ggpairs() and skim()

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# Wrapping up: Further reading (all Free!)

- RStudio > Help > Cheatsheets > Data Transformation with dplyr
- R for Data Science
  - Data transformation
  - Exploratory Data Analysis