Data transformation with dplyr

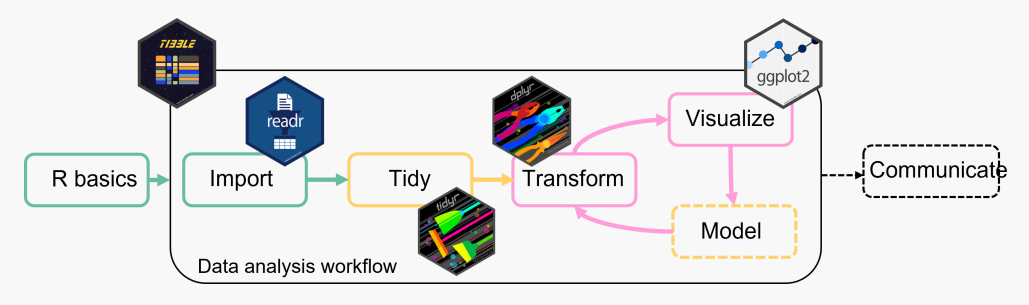
Day 2 - Introduction to Data Analysis with R

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Data transformation

Data transformation is an important step in **understanding** the data and **preparing** it for further analysis.



We can use the tidyverse package dplyr for this.

Data transformation

With dplyr we can (among other things)

- Filter data to analyse only a part of it
- Create new variables
- Summarize data
- Combine multiple tables
- Rename variables
- Reorder observations or variables

To get started load the package dplyr:

```
library(dplyr)
# or
library(tidyverse)
```

Dplyr basic vocabulary

All of the dplyr functions work similarly:

- First argument is the data (a tibble)
- Other arguments specify what to do exactly
- Return a tibble

The data

Data set and_vertebrates with measurements of a trout and 2 salamander species in different forest sections.

- year: observation year
- section: CC (clear cut forest) or OG (old growth forest)
- unittype: channel classification (C = Cascade, P = Pool, ...)
- species: Species measured
- length_1_mm: body length [mm]
- weight_g: body weight [g]



Coastal giant salamander (terrestrial form)
Andrews Forest Program by Lina DiGregorio
via CC-BY from

https://andrewsforest.oregonstate.edu

The data

Data set and_vertebrates with measurements of a trout and 2 salamander species in different forest sections.

```
library(lterdatasampler)
vertebrates <- and vertebrates |>
 select(year, section, unittype, species, length 1 mm, weight g) >
 filter(species != "Cascade torrent salamander")
vertebrates
#> # A tibble: 32,191 × 6
      year section unittype species
                                    length 1 mm weight g
     <dbl> <chr>
                   <chr>>
                                                  <dbl>
                            <chr>>
                                                           <dbl>
   1 1987 CC
                            Cutthroat trout
                                                     58
                                                            1.75
   2 1987 CC
                                                           1.95
                            Cutthroat trout
                                                     61
                            Cutthroat trout
                                                           5.6
   3 1987 CC
   4 1987 CC
                            Cutthroat trout
                                                     58
                                                            2.15
   5 1987 CC
                            Cutthroat trout
                                                           6.9
                                                     93
     1987 CC
                            Cutthroat trout
                                                           5.9
                                                     86
                            Cutthroat trout
   7 1987 CC
                                                          10.5
                                                    107
      1987 CC
                            Cutthroat trout
                                                    131
                                                           20.6
                            Cutthroat trout
      1987 CC
                                                          9.55
                                                    103
#> 10
     1987 CC
                            Cutthroat trout
                                                    117
                                                           13
#> # i 32,181 more rows
```

filter()

picks rows based on their value

filter()

Filter only the trout species:

```
filter(vertebrates, species == "Cutthroat trout")
#> # A tibble: 20,433 × 6
      year section unittype species
                                            length 1 mm weight g
      <dbl> <chr>
                    <chr>>
                             <chr>>
                                                   <dbl>
                                                            <dbl>
#>
   1 1987 CC
                             Cutthroat trout
                                                      58
                                                             1.75
     1987 CC
                             Cutthroat trout
                                                             1.95
                                                             5.6
   3 1987 CC
                             Cutthroat trout
      1987 CC
                            Cutthroat trout
                                                      58
                                                             2.15
      1987 CC
                                                             6.9
                            Cutthroat trout
                            Cutthroat trout
                                                            5.9
      1987 CC
                                                      86
   7 1987 CC
                            Cutthroat trout
                                                            10.5
                                                     107
      1987 CC
                            Cutthroat trout
                                                            20.6
                                                     131
                            Cutthroat trout
      1987 CC
                                                            9.55
                                                     103
      1987 CC
#> 10
                             Cutthroat trout
                                                     117
                                                            13
#> # i 20,423 more rows
```

filter() goes through each row of the data and return only those rows where the value for species is "Cutthroat trout"

filter()

You can also combine filters using logical operators (&, |, !):

```
filter(vertebrates, species == "Cutthroat trout" & year == 1987)
#> # A tibble: 603 × 6
      year section unittype species
                                           length 1 mm weight g
#>
      <dbl> <chr>
                   <chr>
                                                   <dbl>
                            <chr>
                                                            <dbl>
   1 1987 CC
                             Cutthroat trout
                                                             1.75
                                                      58
   2 1987 CC
                            Cutthroat trout
                                                            1.95
                                                      61
   3 1987 CC
                            Cutthroat trout
                                                             5.6
     1987 CC
                            Cutthroat trout
                                                      58
                                                             2.15
      1987 CC
                            Cutthroat trout
                                                            6.9
                                                      93
      1987 CC
                            Cutthroat trout
                                                            5.9
                                                      86
   7 1987 CC
                            Cutthroat trout
                                                            10.5
                                                     107
      1987 CC
                            Cutthroat trout
                                                            20.6
                                                     131
                            Cutthroat trout
      1987 CC
                                                            9.55
                                                     103
      1987 CC
                            Cutthroat trout
#> 10
                                                     117
                                                            13
#> # i 593 more rows
```

filter() + %in%

Use the %in% operator to filter rows based on multiple values, e.g. unittypes

```
unittype select <- c("R", "C", "S")</pre>
filter(vertebrates, unittype %in% unittype select)
#> # A tibble: 19,619 × 6
      year section unittype species
                                    length 1 mm weight g
     <dbl> <chr>
                   <chr>
                           <chr>>
                                                 <dbl>
                                                          <dbl>
   1 1987 CC
                           Cutthroat trout
                                                          1.75
                                                    58
   2 1987 CC
                           Cutthroat trout
                                                          1.95
                                                    61
   3 1987 CC
                                                          5.6
                       Cutthroat trout
   4 1987 CC
                       Cutthroat trout
                                                          2.15
                                                    58
                        Cutthroat trout
   5 1987 CC
                                                          6.9
                                                    93
     1987 CC
                           Cutthroat trout
                                                         5.9
                                                    86
   7 1987 CC
                           Cutthroat trout
                                                         10.5
                                                   107
                           Cutthroat trout
     1987 CC
                                                         20.6
                                                   131
                           Cutthroat trout
     1987 CC
                                                   103
                                                         9.55
#> 10 1987 CC
                           Cutthroat trout
                                                         13
                                                   117
#> # i 19,609 more rows
```

filter() + is.na()

Filter only rows that don't have a value for the weight

```
filter(vertebrates, is.na(weight g))
#> # A tibble: 13,259 × 6
      year section unittype species
                                       length 1 mm weight g
      <dbl> <chr>
                    <chr>>
                            <chr>>
                                                   <dbl>
                                                            <dbl>
#>
     1993 CC
                                                      93
                             Cutthroat trout
                                                               NA
      1993 CC
                            Cutthroat trout
                                                               NA
                                                     175
     1993 CC
                            Cutthroat trout
                                                     104
                                                               NΑ
      1993 CC
                            Cutthroat trout
                                                      98
                                                               NΑ
      1993 CC
                            Cutthroat trout
                                                     97
                                                               NA
                            Cutthroat trout
      1993 CC
                                                               NA
                                                     123
      1993 CC
                            Cutthroat trout
                                                     149
                                                               NA
      1993 CC
                            Cutthroat trout
                                                               NA
                                                     100
                            Cutthroat trout
      1993 CC
                                                     118
                                                               NA
#> 10
      1993 CC
                             Cutthroat trout
                                                     163
                                                               NA
#> # i 13,249 more rows
```

Or the opposite: filter only the rows that have a value for the weight

```
filter(vertebrates, !is.na(weight_g))
```

filter() + between()

Combine different filters:

Filter rows where the value for year is between 2000 and 2005

```
filter(vertebrates, between(year, 2000, 2005))
#> # A tibble: 6,662 × 6
      year section unittype species length 1 mm weight g
     <dbl> <chr> <chr> <chr>
                                              <dbl>
                                                      <dbl>
   1 2000 CC
                         Cutthroat trout
                                                84
                                                         NΑ
   2 2000 CC
                       Cutthroat trout
                                                132
                                                         NΑ
                      Cutthroat trout
   3 2000 CC
                                               105
                                                         NA
                     Cutthroat trout
     2000 CC
                                                41
                                                         NΑ
                     Cutthroat trout
      2000 CC
                                                42
                                                         NΑ
      2000 CC
                     Cutthroat trout
                                                42
                                                         NΑ
      2000 CC
                     Cutthroat trout
                                                         NA
                                                 41
                     Cutthroat trout
      2000 CC
                                                 51
                                                         NA
                       Cutthroat trout
      2000 CC
                                                 45
                                                         NA
      2000 CC
                        Cutthroat trout
#> 10
                                                44
                                                         NΔ
#> # i 6,652 more rows
```

Or you could also do it like this:

```
filter(vertebrates, year >= 2000 & year <= 2005)
```

Useful filter() helpers

These functions and operators help you filter your observations:

- relational operators <, >, ==, ...
- logical operators &, |, !
- %in% to filter multiple values
- is.na() to filter missing values
- between() to filter values that are between an upper and lower boundary
- near() to compare floating points (use instead of == for doubles)

select()

picks columns based on their names

select()

Select the columns species, length_1_mm, and year

```
select(vertebrates, species, length 1 mm, year)
#> # A tibble: 32,191 × 3
     species
             length 1 mm year
#>
                         <dbl> <dbl>
     <chr>
   1 Cutthroat trout
                            58 1987
   2 Cutthroat trout
                            61 1987
  3 Cutthroat trout
                            89 1987
   4 Cutthroat trout
                            58 1987
  5 Cutthroat trout
                            93 1987
                       86 1987
  6 Cutthroat trout
  7 Cutthroat trout
                       107 1987
   8 Cutthroat trout
                           131 1987
  9 Cutthroat trout
                           103 1987
#> 10 Cutthroat trout
                           117 1987
#> # i 32,181 more rows
```

Remove variables using -

```
select(vertebrates, -species, -length_1_mm, -year)
```

select() + starts_with()

Select all columns that start with "s"

You can use the same structure for ends_with() and contains().

```
# this does not make sense for the example data
# but combinations like this are helpful for research data
select(vertebrates, starts_with("_location1"))
select(vertebrates, contains("_id_"))
```

Useful select() helpers

- starts_with() and ends_with(): variable names that start/end with a specific string
- contains(): variable names that contain a specific string
- matches(): variable names that match a regular expression
- any_of() and all_of(): variables that are contained in a character vector

mutate()

Adds new columns to your data

mutate()

New columns can be added based on values from other columns

```
mutate(vertebrates, weight_kg = weight_g/1000)
#> # A tibble: 32,191 × 7
     year section unittype species
                                           length_1_mm weight_g weight_kg
#>
     <dbl> <chr>
                  <chr>
                           <chr>>
                                                 <dbl>
                                                          <dbl>
                                                                    <dbl>
#> 1 1987 CC
                           Cutthroat trout
                                                           1.75
                                                                  0.00175
                                                    58
#> 2 1987 CC
                           Cutthroat trout
                                                    61
                                                           1.95
                                                                  0.00195
#> 3 1987 CC
                           Cutthroat trout
                                                                  0.0056
                                                    89
                                                           5.6
#> # i 32,188 more rows
```

Add multiple new columns at once:

```
mutate(vertebrates,
    weight_kg = weight_g/1000,
    length_m = length_1_mm/1000)
```

mutate() + case_when()

Use case_when to add column values conditional on other columns.

case_when() can combine many cases into one.

```
mutate(vertebrates,
      type = case when(
        species == "Cutthroat trout" ~ "Fish",
                                               # case 1
        species == "Coastal giant salamander" ~ "Amphibian", # case 2
        .default = NA
                                                          # all other
))
#> # A tibble: 32,191 × 7
      year section unittype species length 1 mm weight g type
#>
     <dbl> <chr>
                                               <dbl>
                                                        <dbl> <chr>>
                  <chr>
                           <chr>>
   1 1987 CC
                                                       1.75 Fish
                          Cutthroat trout
                                                  58
   2 1987 CC
                          Cutthroat trout
                                                     1.95 Fish
                                                  61
   3 1987 CC
                          Cutthroat trout
                                                        5.6 Fish
                                                  89
   4 1987 CC
                          Cutthroat trout
                                                        2.15 Fish
                                                  58
     1987 CC
                                                        6.9 Fish
                           Cutthroat trout
                                                  93
     1987 CC
                                                      5.9 Fish
                           Cutthroat trout
                                                86
   7 1987 CC
                           Cutthroat trout
                                                 107
                                                        10.5 Fish
     1987 CC
                                                        20.6 Fish
                          Cutthroat trout
                                                 131
     1987 CC
                       Cutthroat trout
                                                     9.55 Fish
                                                 103
#> 10 1987 CC
                        Cutthroat trout
                                                 117
                                                        13
                                                             Fish
#> # i 32,181 more rows
```

summarize()

summarizes data

summarize()

summarize will collapse the data to a single row

summarize() by group

summarize is much more useful in combination with the grouping argument .by

summary will be calculated separately for each group

```
# summarize the grouped data
summarize(vertebrates,
   mean length = mean(length 1 mm, na.rm = TRUE),
   mean weight = mean(weight g, na.rm = TRUE),
    .by = species
\# # A tibble: 2 × 3
    species
                       mean length mean weight
    <chr>>
                                   <dbl>
                                               <dbl>
#> 1 Cutthroat trout
                                    83.5
                                               8.84
#> 2 Coastal giant salamander
                                    57.0
                                                9.03
```

count()

Counts observations by group

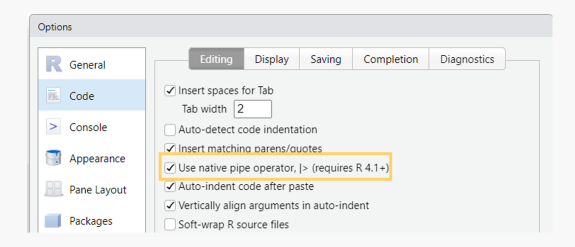
```
# count rows grouped by year
count(vertebrates, year)
#> # A tibble: 33 × 2
#>
      year
     <dbl> <int>
   1 1987
             603
   2 1988
             302
   3 1989
             308
     1990
             513
   5 1991
             626
      1992
             616
   7 1993
             870
      1994
             948
   9 1995
             583
#> 10 1996
             928
#> # i 23 more rows
```

Combine multiple data operations into one command

Data transformation often requires multiple operations in sequence.

The pipe operator |> helps to keep these operations clear and readable.

- You may also see %>% from the magrittr package
- Turn on the native R pipe |> in Tools -> Global Options -> Code



Let's look at an example without pipe:

```
# 1: filter rows that have don't have NA in the unittype column
vertebrates_new <- filter(vertebrates, !is.na(unittype))
# 2: summarize mean values by year
vertebrates_new <- count(vertebrates_new, year, species, section)</pre>
```

How could we make this more efficient?

Use one **nested function** without intermediate results:

```
vertebrates_new <- count(
  filter(vertebrates, !is.na(unittype)),
  year, species, section
)</pre>
```

But this gets complicated and error prone very quickly

The pipe operator makes it very easy to combine multiple operations:

```
vertebrates_new <- vertebrates |>
  filter(!is.na(unittype)) |>
  count(year, species, section)

vertebrates_new
```

You can read from top to bottom and interpret the |> as an "and then do".

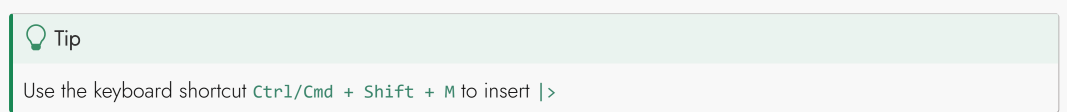
But what is happening?

The pipe is "pushing" the result of one line into the first argument of the function from the next line.

```
vertebrates |>
  count(year)

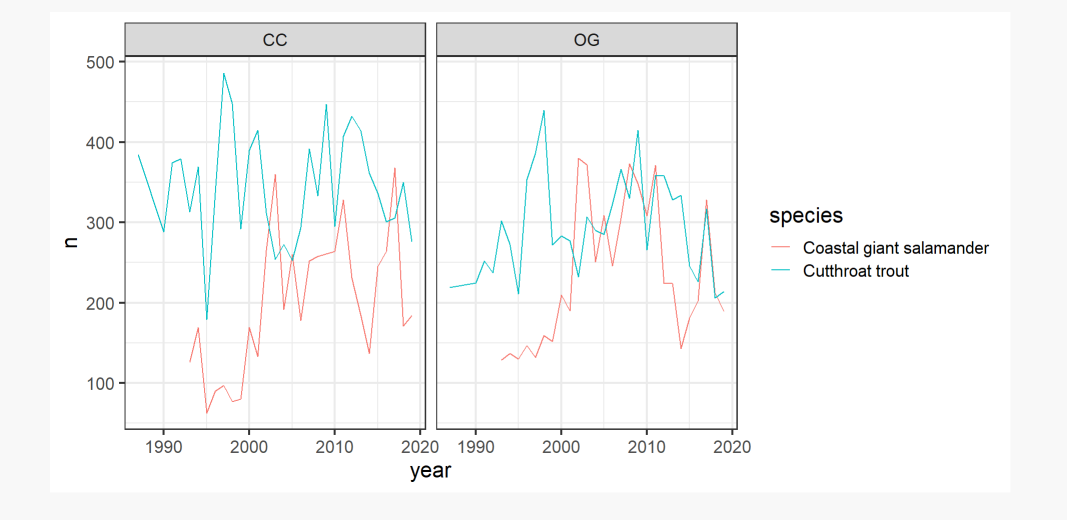
# instead of
count(vertebrates, year)
```

Piping works perfectly with the **tidyverse** functions because they are designed to return a tibble **and** take a tibble as first argument.



Piping also works well together with ggplot

```
vertebrates |>
  filter(!is.na(unittype)) |>
  count(year, species, section) |>
  ggplot(aes(x = year, y = n, color = species)) +
  geom_line() +
  facet_wrap(~section)
```



Combining mulitiple tables

Combine two tibbles by row bind_rows

Situation: Two (or more) tibbles with the same variables (column names)

```
tbl_a <- vertebrates[1:2, ] # first two rows
tbl_b <- vertebrates[2:nrow(vertebrates), ] # the rest</pre>
```

```
tbl a
\# # A tibble: 2 x 6
     year section unittype species
                                         length 1 mm weight g
    <dbl> <chr> <chr>
                                               <dbl>
                          <chr>>
                                                        <dbl>
#> 1 1987 CC
                          Cutthroat trout
                                                  58
                                                        1.75
#> 2 1987 CC
                          Cutthroat trout
                                                        1.95
                                                  61
```

```
tbl b
\# # A tibble: 32,190 × 6
   year section unittype species length 1 mm weight g
    <dbl> <chr>
                 <chr>
                         <chr>>
                                             <dbl>
                                                     <dbl>
                                                   1.95
#> 1 1987 CC
                         Cutthroat trout
                                               61
#> 2 1987 CC
                         Cutthroat trout
                                               89 5.6
#> # i 32,188 more rows
```

Combine two tibbles by row bind_rows

Bind the rows together with bind_rows():

```
bind rows(tbl a, tbl b)
#> # A tibble: 32,192 × 6
     year section unittype species
                                    length 1 mm weight g
     <dbl> <chr>
                  <chr>
                           <chr>>
                                                <dbl>
                                                         <dbl>
#> 1 1987 CC
                           Cutthroat trout
                                                          1.75
                                                   58
#> 2 1987 CC
                           Cutthroat trout
                                                          1.95
                                                   61
#> # i 32,190 more rows
```

You can also add an ID-column to indicate which line belonged to which table:

```
bind rows(a = tbl a, b = tbl b, .id = "id")
#> # A tibble: 32,192 × 7
           year section unittype species length 1 mm weight g
    <chr> <dbl> <chr>
                       <chr>
                                                     <dbl>
                                                              <dbl>
                                <chr>>
           1987 CC
                                Cutthroat trout
                                                             1.75
#> 1 a
                                                        58
       1987 CC
#> 2 a
                                Cutthroat trout
                                                           1.95
       1987 CC
                                Cutthroat trout
                                                              1.95
#> 3 b
                                                        61
#> # i 32,189 more rows
```

You can use bind_rows() to bind as many tables as you want:

```
bind_rows(a = tbl_a, b= tbl_b, c = tbl_c, ..., .id = "id")
```

Join tibbles with left_join()

vertebrates

Situation: Two tables that share some but not all columns.

#> # A tibble: 32,191 × 6 year section unittype species length 1 mm weight g <dbl> <chr> <chr> <chr> <dbl> <dbl> #> 1 1987 CC Cutthroat trout 58 1.75 #> 2 1987 CC Cutthroat trout 1.95 61 #> # i 32,189 more rows

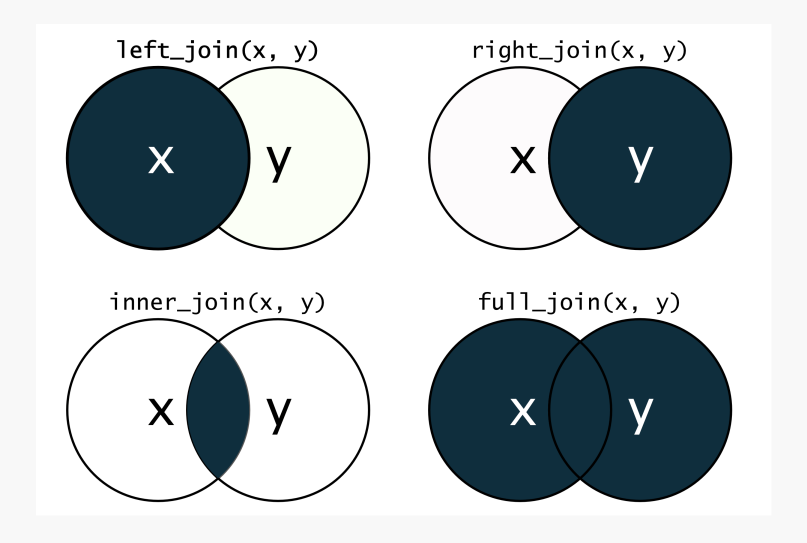
Join tibbles with left_join()

Join the two tables by the common column species

```
left join(vertebrates, species, by = "species")
#> # A tibble: 32,191 × 7
      year section unittype species
                                    length 1 mm weight g type
     <dbl> <chr>
                   <chr>>
                            <chr>
                                                 <dbl>
                                                          <dbl> <chr>
   1 1987 CC
                            Cutthroat trout
                                                    58
                                                           1.75 Fish
   2 1987 CC
                            Cutthroat trout
                                                           1.95 Fish
   3 1987 CC
                            Cutthroat trout
                                                           5.6 Fish
   4 1987 CC
                            Cutthroat trout
                                                    58
                                                           2.15 Fish
     1987 CC
                            Cutthroat trout
                                                           6.9 Fish
                                                    93
                                                          5.9 Fish
      1987 CC
                            Cutthroat trout
                                                    86
   7 1987 CC
                           Cutthroat trout
                                                          10.5 Fish
                                                   107
      1987 CC
                            Cutthroat trout
                                                          20.6 Fish
                                                   131
                            Cutthroat trout
      1987 CC
                                                          9.55 Fish
                                                   103
#> 10
      1987 CC
                            Cutthroat trout
                                                   117
                                                          13
                                                                Fish
#> # i 32,181 more rows
```

left_join() means that the resulting tibble will contain all rows of vertebrates, but not necessarily all rows of species (in this case it does though).

Different *_join() functions



Summary

Data transformation with dplyr

Summary I

All dplyr functions take a tibble as first argument and return a tibble.

```
filter()
```

- pick rows with helpers
 - relational and logical operators
 - %in%
 - is.na()
 - between()
 - near()

Summary II

All dplyr functions take a tibble as first argument and return a tibble.

```
select()
```

- pick columns with helpers
 - starts_with(), ends_with()
 - contains()
 - matches()
 - any_of(), all_of()

Summary III

arrange()

- change order of rows (adscending)
 - or descending with desc()

mutate()

- add columns but keep all columns
 - case_when() for conditional values

Summary IV

summarize()

- collapse rows into one row by some summary
 - use .by argument to summarize by group

count

• count rows based on a group

Summary V

```
bind_rows()
```

- combine rows of multiple tibbles into one
 - the tibbles need to have the same columns
 - add an id column with the argument .id = "id"
 - function bind_cols() works similarly just for columns

left_join()

• combine tables based on common columns

Now you

Task (45 min)

Transform the penguin data set

Find the task description here