Data transformation with dplyr

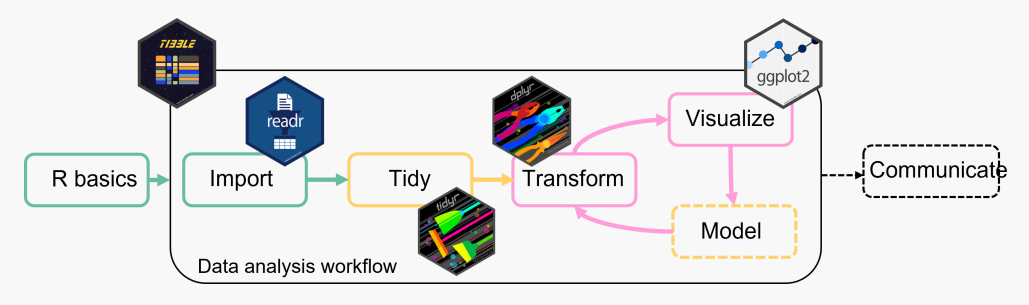
Day 2 - Introduction to Data Analysis with R

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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)
and vertebrates
#> # A tibble: 32,191 × 6
   year section unittype species length 1 mm weight q
#>
   <dbl> <chr> <chr>
                                              <dbl>
                                                      <db1>
#>
                         <chr>
#>
   1 1987 CC
                         Cutthroat trout
                                                 58
                                                   1.75
             R
   2 1987 CC
                          Cutthroat trout
#>
                                                61 1.95
#>
   3 1987 CC
                                                89 5.6
                          Cutthroat trout
                                                       2.15
#>
  4 1987 CC
                          Cutthroat trout
                                                58
#>
   5 1987 CC
                          Cutthroat trout
                                                93
                                                       6.9
#>
  6 1987 CC
                          Cutthroat trout
                                                86 5.9
#>
  7 1987 CC
                          Cutthroat trout
                                               107
                                                      10.5
#>
  8 1987 CC
                          Cutthroat trout
                                               131
                                                      20.6
                                               103 9.55
   9 1987 CC
                          Cutthroat trout
#> 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(and vertebrates, species == "Cutthroat trout")
\#>\# A tibble: 20,433 × 6
#>
     year section unittype species length 1 mm weight q
#>
  <dbl> <chr> <chr>
                       <chr>
                                         <dbl>
                                                <db1>
#>
  1 1987 CC R
                       Cutthroat trout
                                            58 1.75
  2 1987 CC R
#>
                       Cutthroat trout
                                    61 1.95
#>
  3 1987 CC R
                                           89 5.6
                       Cutthroat trout
  4 1987 CC R
#>
                       Cutthroat trout
                                           58 2.15
#>
  5 1987 CC
                                    93 6.9
                       Cutthroat trout
                                       86 5.9
#>
  6 1987 CC R
                       Cutthroat trout
#>
  7 1987 CC
                       Cutthroat trout
                                          107
                                                10.5
                       Cutthroat trout
#>
  8 1987 CC
                                          131 20.6
   9 1987 CC
                       Cutthroat trout
                                          103 9.55
#> 10 1987 CC
                       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 (and vertebrates, species == "Cutthroat trout" & year == 1987)
#> # A tibble: 603 × 6
#>
  year section unittype species length 1 mm weight g
  <dbl> <chr> <chr> <chr>
#>
                                  <dbl> <dbl>
  1 1987 CC R Cutthroat trout
                                       58 1.75
#> 2 1987 CC R Cutthroat trout
                                61 1.95
  3 1987 CC R
#>
                                89 5.6
                    Cutthroat trout
#>
  4 1987 CC R
                    Cutthroat trout 58 2.15
  5 1987 CC R
#>
                    Cutthroat trout
                                93 6.9
  6 1987 CC R
                    Cutthroat trout
                                86 5.9
#> 7 1987 CC R
                    Cutthroat trout
                                      107 10.5
#> 8 1987 CC R
                                      131 20.6
                    Cutthroat trout
  9 1987 CC R
                    Cutthroat trout
                                      103 9.55
#> 10 1987 CC
                     Cutthroat trout
                                      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")
filter (and 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 R Cutthroat trout
                                        58
                                             1.75
#> 2 1987 CC R
                                 61 1.95
                     Cutthroat trout
#> 3 1987 CC R
                     Cutthroat trout 89 5.6
                                 58 2.15
#> 4 1987 CC R
                     Cutthroat trout
                                 93 6.9
#>
  5 1987 CC R
                     Cutthroat trout
#> 6 1987 CC R
                     Cutthroat trout
                                 86 5.9
#> 7 1987 CC R
                                       107 10.5
                     Cutthroat trout
#> 8 1987 CC R
                     Cutthroat trout
                                       131 20.6
  9 1987 CC R
                     Cutthroat trout
                                       103 9.55
#> 10 1987 CC
                     Cutthroat trout
                                       117
                                            13
#> # i 19,609 more rows
```

filter() + is.na()

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

```
filter(and vertebrates, is.na(weight g))
\#>\# A tibble: 13,259 × 6
#>
   year section unittype species length 1 mm weight q
  <dbl> <chr> <chr> <chr>
#>
                                           <dbl>
                                                 <dbl>
  1 1993 CC P
                        Cutthroat trout
                                              93
                                                      NA
  2 1993 CC P
#>
                        Cutthroat trout
                                             175
                                                     NA
  3 1993 CC P
#>
                        Cutthroat trout
                                            104
                                                     NA
#> 4 1993 CC P
                        Cutthroat trout
                                      98
                                                     NA
#> 5 1993 CC P
                        Cutthroat trout
                                      97
                                                     NA
#>
  6 1993 CC
                        Cutthroat trout
                                             123
                                                     NΑ
#> 7 1993 CC
                        Cutthroat trout
                                             149
                                                     NA
#> 8 1993 CC
                        Cutthroat trout
                                            100
                                                     NA
  9 1993 CC
                        Cutthroat trout
                                             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(and_vertebrates, !is.na(weight_g))
```

filter() + between()

Combine different filters:

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

```
filter (and 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 C
#>
                       Cutthroat trout
                                            84
                                                    NA
  2 2000 CC C
#>
                       Cutthroat trout
                                           132
                                                    NA
  3 2000 CC C
#>
                       Cutthroat trout
                                           105
                                                    NA
  4 2000 CC
#>
                                     4.1
                       Cutthroat trout
                                                    NA
#>
  5 2000 CC
                                            42
                       Cutthroat trout
                                                    NA
#> 6 2000 CC C
                                            42
                       Cutthroat trout
                                                    NA
#> 7 2000 CC C
                       Cutthroat trout
                                     41
                                                    NA
  8 2000 CC C
                       Cutthroat trout
                                                    NA
   9 2000 CC C
                       Cutthroat trout
                                     45
                                                    NA
#> 10 2000 CC
                       Cutthroat trout
                                        44
                                                    NA
\# > \# i 6,652 \text{ more rows}
```

Or you could also do it like this:

```
filter(and_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 (and vertebrates, species, length 1 mm, year)
#> # A tibble: 32,191 × 3
  species length 1 mm year
#>
  #>
  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
#> 6 Cutthroat trout 86 1987
#> 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(and_vertebrates, -species, -length_1_mm, -year)
```

select() + ends_with()

Select all columns that start with "s"

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

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

select() + from:to

Multiple consecutive columns can be selected using the from:to structure with either column id or name:

```
select(and_vertebrates, 1:3)
select(and_vertebrates, year:unittype)

#> # A tibble: 32,191 × 3
#> year section unittype
#> <dbl> <chr> <chr> #> 1 1987 CC R
#> 2 1987 CC R
#> 3 1987 CC R
#> 3 1987 CC R
#> #> # i 32,188 more rows
```

Be a bit careful with these commands: They are not robust if you e.g. change the order of your columns at some point.

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 (and 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>
                                          <dbl>
                                                 <dbl>
                                                          <dbl>
                       \langle chr \rangle
#> 1 1987 CC R Cutthroat trout
                                                 1.75 0.00175
                                            58
#> 2 1987 CC R Cutthroat trout
                                          61 1.95 0.00195
#> 3 1987 CC R Cutthroat trout
                                                  5.6 0.0056
                                            89
#> # i 32,188 more rows
```

Add multiple new columns at once:

```
mutate(and_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 (and 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> <chr> <chr>
                                    <dbl> <dbl> <chr>
                                          58 1.75 Fish
#>
  1 1987 CC R Cutthroat trout
  2 1987 CC R
#>
                      Cutthroat trout
                                   61 1.95 Fish
#>
  3 1987 CC R
                       Cutthroat trout 89 5.6 Fish
#>
  4 1987 CC R
                      Cutthroat trout
                                   58 2.15 Fish
#>
  5 1987 CC
                       Cutthroat trout
                                   93 6.9 Fish
           R
#>
  6 1987 CC
                       Cutthroat trout
                                   86 5.9 Fish
                                         107 10.5 Fish
#> 7 1987 CC
                      Cutthroat trout
#> 8 1987 CC
                       Cutthroat trout
                                         131
                                               20.6 Fish
                                         103 9.55 Fish
                       Cutthroat trout
  9 1987 CC
#> 10 1987 CC
                                         117
                                               13 Fish
                       Cutthroat trout
```

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

count()

Counts observations by group

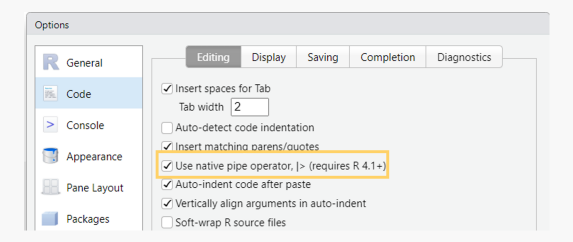
```
# count rows grouped by year
count(and vertebrates, year)
#> # A tibble: 33 × 2
    year
   <dbl> <int>
  1 1987
            603
  2 1988
          302
#>
   3 1989 308
#>
  4 1990
          513
  5 1991
          626
#>
          616
  6 1992
#>
  7 1993
          870
#> 8 1994
          948
   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
and_vertebrates_new <- filter(and_vertebrates, !is.na(unittype))
# 2: summarize mean values by year
and_vertebrates_new <- count(and_vertebrates_new, year, species, section)</pre>
```

How could we make this more efficient?

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

```
and_vertebrates_new <- count(
  filter(and_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:

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

and_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.

```
and_vertebrates |>
  count(year)

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

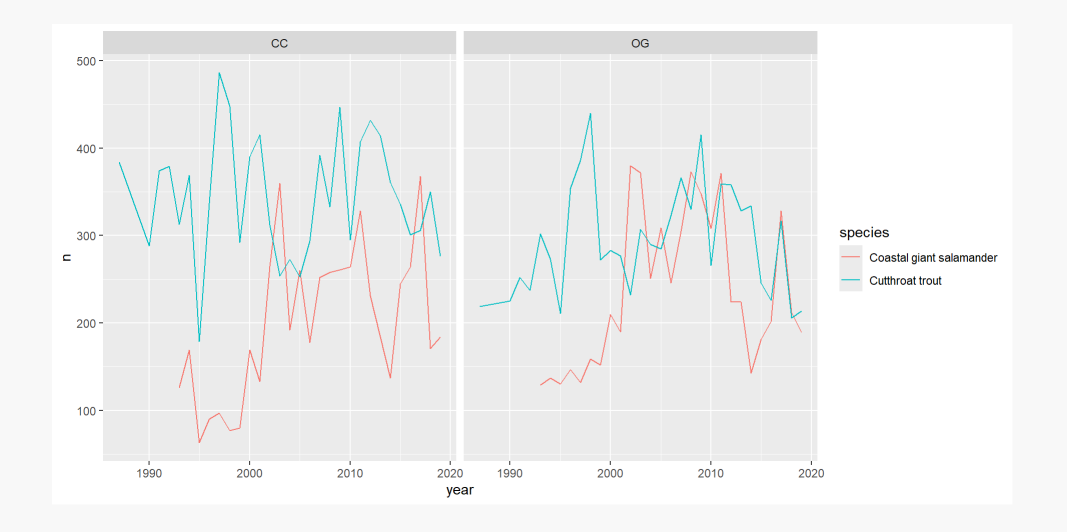
Piping works perfectly with the **tidyverse** functions because they are designed to return a tibble **and** take a tibble as first argument.

```
    □ Tip

Use the keyboard shortcut Ctrl/Cmd + Shift + M to insert |>
```

Piping also works well together with ggplot

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

```
#> # A tibble: 2 × 6
#> year section unittype species length_1_mm weight_g
#> <dbl> <chr> <chr> <chr> < dbl> <chr> < Cutthroat trout 58 1.75
#> 2 1987 CC R Cutthroat trout 61 1.95
```

```
#> # A tibble: 32,190 × 6
#> year section unittype species length_1_mm weight_g
#> <dbl> <chr> <chr> <chr> <chr> <chr> < dbl> <fl> 1 1987 CC R Cutthroat trout 61 1.95
#> 2 1987 CC R 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> <chr> <chr> < dbl> <dbl> <dbl> <dbl> </dbl>
#> 1 1987 CC R Cutthroat trout 58 1.75

#> 2 1987 CC R Cutthroat trout 61 1.95

#> # i 32,190 more rows
```

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

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()

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> <chr> < Color <
```

```
# table with more information on the species
species
#> # A tibble: 2 × 2
#> species type
#> <chr>
#> 1 Cutthroat trout Fish
#> 2 Coastal giant salamander Amphibian
```

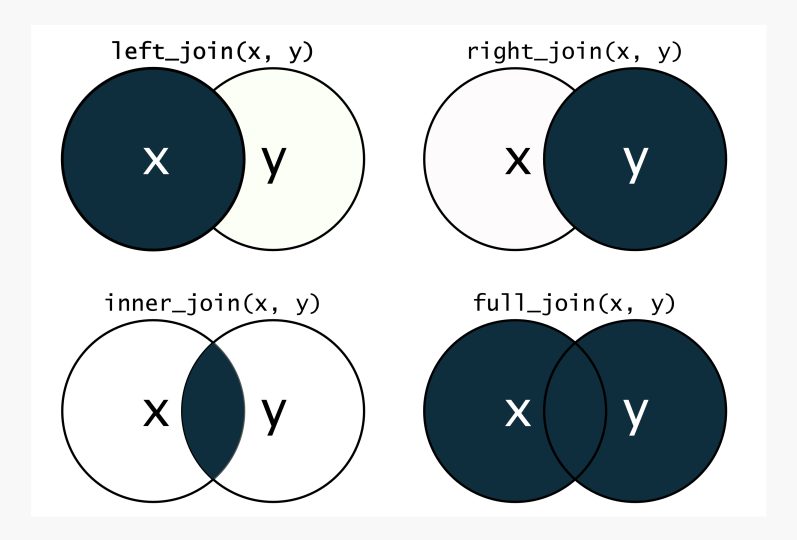
Join tibbles with left_join()

Join the two tables by the common column species

```
left join(and vertebrates, species, by = "species")
#> # A tibble: 32,191 × 7
    year section unittype species length 1 mm weight g type
#>
   <dbl> <chr> <chr>
                                           \langle dbl \rangle \langle dbl \rangle \langle chr \rangle
#>
                        <chr>
  1 1987 CC R
                                              58 1.75 Fish
#>
                        Cutthroat trout
#> 2 1987 CC R
                                      61 1.95 Fish
                        Cutthroat trout
                                              89 5.6 Fish
#>
  3 1987 CC R
                        Cutthroat trout
#>
  4 1987 CC R
                                              58 2.15 Fish
                        Cutthroat trout
#> 5 1987 CC
                        Cutthroat trout
                                      93 6.9 Fish
                                      86 5.9 Fish
#>
  6 1987 CC
                        Cutthroat trout
            R
#> 7 1987 CC
                                                   10.5 Fish
                        Cutthroat trout
                                             107
#> 8 1987 CC
                                                   20.6 Fish
                        Cutthroat trout
                                             131
   9 1987 CC
                        Cutthroat trout
                                             103 9.55 Fish
#> 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
and_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