Joining Data Tables

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Joining Data Tables

Shortcut: Use crtl + opt + i to create a code block.

```
library(readr)
```

```
library(dplyr)
##
## Attaching package: 'dplyr'
## The following objects are masked from 'package:stats':
##
##
      filter, lag
## The following objects are masked from 'package:base':
##
      intersect, setdiff, setequal, union
##
surveys <- read_csv("../197-raw_storage/surveys.csv")</pre>
## Rows: 35549 Columns: 9
## -- Column specification -----
## Delimiter: ","
## chr (2): species_id, sex
## dbl (7): record_id, month, day, year, plot_id, hindfoot_length, weight
##
## i Use 'spec()' to retrieve the full column specification for this data.
## i Specify the column types or set 'show_col_types = FALSE' to quiet this message.
species <- read.csv( "../197-raw_storage/species.csv")</pre>
plots <- read.csv("../197-raw_storage/plots.csv")</pre>
```

How do we combine tables?

We can use the *shared columns* between different data frames to combine them.

Specifically, we will use _join functions to combine two or mate data tables.

The different functions of _join allow us to combine in different ways: inner_join - connects data sets, and removes all outliers.

left_join - keeps all values in x, and adds matching values for y right_join - keeps all values in y, and adds matching values in x full_join - keeps all values of x AND y

```
surveys %>%
inner_join(species, by = "species_id") -> join_table
head(join_table)
```

```
## # A tibble: 6 x 12
##
     record_id month
                        day year plot_id speci~1 sex
                                                         hindf~2 weight genus species
         <dbl> <dbl> <dbl> <dbl> <dbl>
##
                                                                  <dbl> <chr> <chr>
                                    <dbl> <chr>
                                                   <chr>
                                                           <dbl>
## 1
             1
                   7
                         16 1977
                                        2 NL
                                                              32
                                                                      NA Neot~ albigu~
                                                   M
             2
                   7
                                        3 NL
## 2
                         16 1977
                                                   Μ
                                                              33
                                                                      NA Neot~ albigu~
## 3
             3
                   7
                         16 1977
                                        2 DM
                                                   F
                                                              37
                                                                      NA Dipo~ merria~
## 4
             4
                   7
                         16 1977
                                        7 DM
                                                              36
                                                                      NA Dipo~ merria~
## 5
                   7
                                        3 DM
             5
                         16 1977
                                                              35
                                                                      NA Dipo~ merria~
                                                   M
## 6
             6
                   7
                         16
                            1977
                                        1 PF
                                                   М
                                                              14
                                                                      NA Pero~ flavus
## # ... with 1 more variable: taxa <chr>, and abbreviated variable names
       1: species_id, 2: hindfoot_length
```

str(join_table)

```
## spc_tbl_ [34,786 x 12] (S3: spec_tbl_df/tbl_df/tbl/data.frame)
##
   $ record_id
                    : num [1:34786] 1 2 3 4 5 6 7 8 9 10 ...
## $ month
                     : num [1:34786] 7 7 7 7 7 7 7 7 7 7 ...
## $ day
                     : num [1:34786] 16 16 16 16 16 16 16 16 16 ...
## $ year
                     : num [1:34786] 1977 1977 1977 1977 ...
##
                    : num [1:34786] 2 3 2 7 3 1 2 1 1 6 ...
   $ plot_id
                     : chr [1:34786] "NL" "NL" "DM" "DM" ...
##
  $ species_id
                     : chr [1:34786] "M" "M" "F" "M" ...
## $ sex
##
   $ hindfoot_length: num [1:34786] 32 33 37 36 35 14 NA 37 34 20 ...
                    : num [1:34786] NA ...
##
   $ weight
                     : chr [1:34786] "Neotoma" "Neotoma" "Dipodomys" "Dipodomys" ...
## $ genus
                     : chr [1:34786] "albigula" "albigula" "merriami" "merriami" ...
## $ species
                     : chr [1:34786] "Rodent" "Rodent" "Rodent" ...
##
   $ taxa
##
   - attr(*, "spec")=
##
     .. cols(
##
         record_id = col_double(),
##
         month = col double(),
##
         day = col_double(),
     . .
##
         year = col_double(),
##
         plot_id = col_double(),
##
         species_id = col_character(),
     . .
##
         sex = col_character(),
##
         hindfoot_length = col_double(),
     . .
##
         weight = col_double()
##
     ..)
   - attr(*, "problems")=<externalptr>
```

This is good! Now, let's filter out any plot types that aren't "Control".

```
surveys %>%
  inner_join(plots, by = "plot_id") %>%
  filter(plot_type == 'Control') %>%
  head() %>%
  str()
```

```
## tibble [6 x 10] (S3: tbl_df/tbl/data.frame)
## $ record_id
                   : num [1:6] 1 3 7 14 16 18
## $ month
                   : num [1:6] 7 7 7 7 7
## $ day
                   : num [1:6] 16 16 16 16 16 16
## $ year
                   : num [1:6] 1977 1977 1977 1977 ...
## $ plot_id
                   : num [1:6] 2 2 2 8 4 2
                   : chr [1:6] "NL" "DM" "PE" "DM" ...
## $ species_id
                   : chr [1:6] "M" "F" "F" NA ...
## $ sex
## $ hindfoot_length: num [1:6] 32 37 NA NA 36 22
## $ weight
                   : num [1:6] NA NA NA NA NA NA
## $ plot_type
                   : chr [1:6] "Control" "Control" "Control" ...
```

Finding Relevant Data

What if we don't know what columns both data sets have?

What if sorting through columns is too much of a tedious task.

```
intersect(colnames(surveys), colnames(species))
```

```
## [1] "species_id"
```

Joining Multiple Tables

```
surveys %>%
  inner_join(species, by = "species_id") %>%
  inner_join(plots, by = "plot_id") -> combined
head(combined)
```

```
## # A tibble: 6 x 13
##
     record_id month
                                                        hindf~2 weight genus species
                       day year plot_id speci~1 sex
##
         <dbl> <dbl> <dbl> <dbl>
                                    <dbl> <chr>
                                                  <chr>>
                                                          <dbl>
                                                                 <dbl> <chr> <chr>
## 1
                                                                    NA Neot~ albigu~
             1
                   7
                        16 1977
                                       2 NL
                                                  М
                                                             32
## 2
             2
                   7
                        16 1977
                                        3 NL
                                                             33
                                                  М
                                                                    NA Neot~ albigu~
                   7
## 3
                                        2 DM
                                                  F
                                                             37
             3
                        16 1977
                                                                    NA Dipo~ merria~
## 4
             4
                   7
                        16 1977
                                       7 DM
                                                  М
                                                             36
                                                                    NA Dipo~ merria~
## 5
             5
                   7
                        16 1977
                                       3 DM
                                                             35
                                                  М
                                                                    NA Dipo~ merria~
                   7
                        16 1977
                                       1 PF
                                                             14
             6
                                                  М
                                                                    NA Pero~ flavus
## # ... with 2 more variables: taxa <chr>, plot_type <chr>, and abbreviated
       variable names 1: species_id, 2: hindfoot_length
```

str(combined)

```
## spc_tbl_ [34,786 x 13] (S3: spec_tbl_df/tbl_df/tbl/data.frame)
                   : num [1:34786] 1 2 3 4 5 6 7 8 9 10 ...
## $ record_id
## $ month
                    : num [1:34786] 7 7 7 7 7 7 7 7 7 7 ...
## $ day
                    : num [1:34786] 16 16 16 16 16 16 16 16 16 ...
                    : num [1:34786] 1977 1977 1977 1977 ...
## $ year
## $ plot_id
                    : num [1:34786] 2 3 2 7 3 1 2 1 1 6 ...
## $ species_id
                   : chr [1:34786] "NL" "NL" "DM" "DM" ...
## $ sex
                    : chr [1:34786] "M" "M" "F" "M" ...
## $ hindfoot_length: num [1:34786] 32 33 37 36 35 14 NA 37 34 20 ...
                   : num [1:34786] NA ...
## $ weight
                    : chr [1:34786] "Neotoma" "Neotoma" "Dipodomys" "Dipodomys" ...
## $ genus
## $ species
                   : chr [1:34786] "albigula" "albigula" "merriami" "merriami" ...
                    : chr [1:34786] "Rodent" "Rodent" "Rodent" "Rodent" ...
## $ taxa
##
   $ plot_type
                    : chr [1:34786] "Control" "Long-term Krat Exclosure" "Control" "Rodent Exclosure"
##
  - attr(*, "spec")=
##
     .. cols(
##
         record_id = col_double(),
##
     .. month = col_double(),
##
     .. day = col_double(),
##
     .. year = col_double(),
     .. plot_id = col_double(),
##
##
     .. species_id = col_character(),
##
     .. sex = col_character(),
##
        hindfoot_length = col_double(),
##
         weight = col_double()
    . .
##
     ..)
   - attr(*, "problems")=<externalptr>
surveys %>%
  inner_join(species, by = "species_id") %>%
  inner_join(plots, by = "plot_id") %>%
  filter(plot_type == 'Control' | plot_type == 'Long-term Krat Exclosure') %>%
  filter(taxa == 'Rodent', !is.na(weight)) %>%
  select(year, genus, species, weight, plot_type)
## # A tibble: 19,344 x 5
##
      year genus
                       species weight plot_type
      <dbl> <chr>
                       <chr>>
                                 <dbl> <chr>
## 1 1977 Dipodomys
                       merriami
                                    40 Long-term Krat Exclosure
## 2 1977 Dipodomys
                       merriami
                                    29 Control
## 3 1977 Dipodomys
                       merriami
                                    46 Control
                                   52 Control
## 4 1977 Dipodomys
                       ordii
## 5 1977 Perognathus flavus
                                    8 Control
## 6 1977 Onychomys
                                   22 Long-term Krat Exclosure
                       sp.
## 7 1977 Perognathus flavus
                                   7 Control
## 8 1977 Dipodomys merriami
                                   22 Control
## 9 1977 Perognathus flavus
                                    8 Control
## 10 1977 Dipodomys merriami
                                    41 Control
## # ... with 19,334 more rows
```