Introduction to the tidyverse: dplyr, tidyr and purrr

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Introduction

Thi is a RMarkdown document generated to illustrate the exercices suggested as part of the Workshop "Introduction to the tidyverse", that will be celebrated in Barcelona within the 1st Meeting of the Iberian Ecological Society and the XIV AEET Meeting. All the code and the data needed to produce this document can be found in GitHub (https://github.com/ameztegui/dplyr_workshop). For any doubt about the exercises, you can contact Aitor Ameztegui (ameztegui@gmail.com) or V?ctor Granda (ameztegui@gmail.com).

The packagestidyr, dplyr, and purr are part of a set of packages known as the tidyverse, created by Hadley Wickham, Chief Scientist in RStudio. The tidyverse was created to easen data analysis and data management. It consists on packages to import and read data, to organize and modify them, to analyze and model them and to visualize them. In this seminar we will focus on tidyr, conceived to help organize data, and dplyr, that focuses on data transformation. We must firs install these packages (if we haven't done it yet). To install all the packages from the tidyverse at once we need to write install.packages("tidyverse"). Then we need to load the packages by typing:

```
library(tidyverse)
```

We also need to load the dataset we will use for this workshop, we can download them from GitHub.

```
load("../data/data_workshop.Rdata")

trees <- tbl_df(trees)
plots <- tbl_df(plots)
species <- tbl_df(species)
coordinates <- tbl_df(coordinates)</pre>
```

Through the function tbl_df we will convert normal data frames into tibbles. A tibble is just a data frame with some particularities: for example, they only print the first 10 rows by default (instead of the whole data frame), and printing them provides information on all the variables and their class. Besides that, we can treat tibbles as normal data frames, because they behave like them at all effects.

In these exercises we will use four data frames with information from the 2nd and 3rd Spanish National Forest Inventory (IFN2 e IFN3) in Catalonia. The data frames are:

- plots [11,858 x 15]: all the IFN3 plots in Catalonia, with info about the date and time of measurement, soil texture and soil pH, total canopy cover and tree canopy cover, etc.
- trees [111,756 x 12]: contains all the adult trees (diam > 7.5 cm) measured both in IFN2 and IFN3. Contains info about the plot, the species, diameter class, diameter measured at IFN2 and IFN3...
- species [14,778 x 15]: contains the number of trees/ha per species and diameter class.
- coordinates [11,858 x 6]: contains the X & Y coordinates of each IFN3 plot.

First thing to do is to have a look at the data, to get familiar with the data they contain. We will use the function glimpse for that.

```
glimpse(plots)
glimpse(trees)
glimpse(species)
glimpse(coordinates)
```

dplyr: transforming data frames

dplyr can be used to transform our data frames in the way we need it in each case: we can create bew variables, select those of interest, execute filters, etc. The dplyr package contains 5 main verbs:

- filter selects rows based on a given set of conditions
- select select columns based on their name
- arrange sort the data frame based on one or several variables
- mutate create new variables
- summarise create new variables that summarize values of an existing variable (mean, sum, etc.)

All of them have a similar structure: the first argument in the function is the data frame to which it will be applied, and the rest of arguments specify what to do with this data frame, depending on the verbwe are using.

filter

filter selects those rows of a data frame that accomplish a certain criterion. The first argument is the data frame, and the rest are the criteria, that can be specified in chain, separated by commas.

```
###Exercice 1
```

To practice with filter let's try to find those plots of IFN that:

• 1.1 Are located in Barcelona (08) or Girona (17). We have two options:

```
# Option 1
   filter (plots, Provincia =="08" | Provincia =="17")
# Option 2
   filter (plots, Provincia %in% c("08", "17"))
```

We see that both options produce exactly the same result. But the next option wouldn't be valid, since we need to specify explicitly the variable every time we add a new condition:

```
filter(plots, Provincia =="08" | "17")
```

• 1.2 plots that were measured completely in January 2001

To do this we need to find the plots for which the completion date is later than 31 December 2000 and earlier than 1 February 2001. We can do this in two ways: the first one is to use the & operator to indicate we want to get the rows that meet both criteria. The second options would be simply to concatenate both criteria with a comma, since filter assumes all of them must be met.

```
# Option 1
filter (plots, FechaFin < "2001-02-01" & FechaFin > "2000-12-31")

# Option 2
filter (plots, FechaFin < "2001-02-01", FechaFin > "2000-12-31")
```

• 1.3 Those plots that took more than 2 hours to be measured (7200 seg)

```
filter(plots, (HoraFin - HoraIni) >7200)
```

As we see, we can do operations within filter conditions. In this case, we want that the difference between EndDate and StartDate be < 7200 s (2 hours).

select

select allows to retain only some columns, based on their name. To help us find the columns, there are some specific functions such as starts_with or contains, that only work within select. We can see the list of special functions by typing help("select")

###Exercise 2 To practice with select let's try to find 4 different ways of selecting the variables that specify the starting and ending date of measurement of the plots (FechaIni y FechaFin)

• For example, we could specify the name of the columns we want to keep in an explicit way

```
select(plots, FechaIni, FechaFin)
```

• We can also specify them as a range, so that all columns between the two indicated will be selected

```
select(plots, FechaIni:FechaFin)
```

• Or we could select all the columns that contain the text 'fecha'. In this case, since we are not interested in the pH measurement date, we can decide to delete it from our selection:

```
select(plots, contains ("Fecha"), -FechaPh)
```

• At last, we could also select all the variables that start with 'fecha' (in this case we also need to eliminate FechaPh):

```
select(plots, starts_with("Fecha"), -FechaPh)
```

arrange

arrange sorts the data frame based on the values of ine or more variables (columns). The first argument will be, as usual, the data frame we want to sort, and then we must specify the variables that determine the ordering. If we specify more than 1 variable, the succesive variables will be used to decide order when there are ties (i.e. secondary sorting variables). We can also use 'desc(x)' to sort in decreasing order. Let's try with a few exercises:

###Exercise 3

• Ex.3.1 Sort the plots by measurement date and time

```
arrange(plots, FechaFin, HoraFin)
```

• Ex. 3.2 Which plots were started to be measured later in the day?

```
arrange(plots,desc(HoraIni))
```

• Ex. 3.3 Which took longer to be measured?

```
arrange(plots, desc(HoraFin-HoraIni))
```

We see that, as it happens with filter, we can also sort data frames based on the result of an arithmetic operation.

mutate

mutate allows us to create new variables with a certain value or as combination of existing variables. We just need to specify the data frame, and indicate the new variables name and its value. Let's see some examples:

###Exercise 4 Let's create two new variables:

• Ex.4.1 A variable with individual tree growth (in cm) between IFN2 and IFN3.

```
trees <- mutate (trees, growth= DiamIf3 - DiamIf2)</pre>
```

• Ex.4.2 Create two new variables with the basal area per hectare that each tree represents, both in IFN2 and IFN3. Which species was the fastest growing tree in basal area?

As we see, we can calculate new variables based on the variables we just created. Also, we can combine mutate and arrange to know which is the fastest growing tree.

summarise

summarise allows us to make calculations with the variables in the data frame, but using summary functions, that transform the variability in a given variable into a single value. Functions such as sum, mean, max, IQR, etc. are examples of summary functions. However, this function by itself often lacks any interest, cause it would reduce all the data frame to a single value. It is commonly used together with group_by, that classifies the data frame in groups based on a categorical variable.

To use group_by we just need to indicate the data frame and the variable we want to group it by. To be more efficient, dplyr does not create a copy of the data frame, but it creates a hidden variable that indexes the groups, so that when we ask it to perform operations by group, it know to which group belongs each observation.

In the case of our data frame trees, there are several groups that could be of interest el caso de nuestra base de datos de pies trees, hay varios grupos que pueden tener inter?s:

```
# Por provincia
by_province <- group_by (trees, Provincia)

# Por parcela
by_plot <- group_by (trees, Codi)

# Por especie
by_species <- group_by (trees, Especie)

#Por clase diam?trica
by_CD <- group_by (trees, CD)

#Por parcela y especie
by_plot_species <- group_by (trees, Codi, Especie)</pre>
```

We can see, by typing glimpse(by_plot) that the resulting data frame is not different at all from the original, at least apparently. However, if we type class(by_plot) we see it has now a new class grouped_df.

###Exercise 5 What statistics could be of interest to characterize the diameter values of each plot? We can, for example, calculate te mean, minimum and maximum value, percentile 0.9 and interquartile range for each plot. We can also compute the number of trees measured in each plot and the number of different species, using the functions n() y n_distinct(x). In this case, the resulting data frame will have less rows, one per plot, and will only contain the new variables created.

```
summarise(by_plot,
    media = mean(DiamIf3),
    min = min (DiamIf3),
    max = max(DiamIf3),
    q90 = quantile(DiamIf3, 0.9),
    IQ = IQR(DiamIf3),
```

```
n =n(),
sps = n_distinct(Especie) )
```

```
## # A tibble: 7,713 x 8
    Codi media min max q90 IQ
##
                                    n sps
##
    <fct> <dbl> <dbl> <dbl> <dbl> <dbl> <int> <int>
## 1 080001 26.3 13.4 38
                          34.4 4.15
                                     15
## 2 080002 35.2 24.8 44.4 43.0 10.2
                                     13
## 3 080003 32.0 14.2 51 46.1 12.2
                                     7
                                          2
## 4 080004 24.3 16.8 31.7 30.2 7.42
                                     2
## 5 080005 28.4 16.2 59.8 39.7 15.3
                                     12
                                          3
## 6 080006 35.9 14 55.9 52.3 17.9
                                    23
## 7 080007 30.8 15.2 63.6 49.4 12.3
                                    35
                                          2
## 8 080008 16.0 9
                     21.4 17.8 2.05
                                     11
## 9 080009 16.7 9
                     36.0 24 3.88
                                     16
                                          2
## 10 080010 31.6 9.2 95.5 61.5 18.2
                                     13 3
## # ... with 7,703 more rows
```

Pipelines (%>%)

We will often use several dplyr verbs together, creating nested functions. However, when we need to perform several operations, these nested functions can easily get complex and difficult to understand. For example, by having a look at this code, would you be able to say what it will do?

The code gets those observations, from the data frame trees, that have a value of diameter (!is.na(DiamIf3)), it then groups them by plot and species (group_by(Codi, Especie)), calculates for each combination the mean diameter (diam = mean (DiamIf3)), and the number of trees per plot (n = n()), and finally selects only those cases in which there are at least 5 trees (filter (n>5)).

Although this syntaxis is not operationally complex, it is hard to understand. Often a solution is to save each step as a different data frame, but this is an important source of errors.

We can however simplify this code using the *pipe* operator (%>%) from the magrittr package, which is installed and loaded with tidyr and dplyr. When we use %>%, the result of the left side is processed by the righ side function as first argument. In the case of dplyr and tidyr, since the first argument is always a data frame, %>% makes that a function be applied to the data frame resulting from the previous function. Thus, we can express filter (df, color == "blue") as df %>% filter(color == "blue"). This allows to concatenate several functions in a logical and understandable way, so that the operator %>% could be read as then. Let's see how this would be in the previous function

```
diam_medio_especie <- trees %>%  # take the df 'trees' and THEN
filter(!is.na(DiamIf3)) %>%  # eliminate NA values and THEN
group_by(Codi, Especie) %>%  # group y plot and species and THEN
summarise(diam=mean(DiamIf3), n = n()) %>%  # calculate mean and number of trees and THEN
filter(n > 5)  # filter those with n> 5
```

###Exercise 6 Let's do some exercises. Using the pipe operator, let's create pipelines to solve the next exercises:

• Ex.6.1 Which plots have the greatest average growth between IFN2 and IFN3?

We first define the data frame we will work with. THEN (%>%) we create a new variable with the growth of each tree, THEN we group by plot, THEN we calculate, for each plot, the mean growth, and THEN we arrange the results in decreasing order. The resulting code would be:

```
trees %>%
  mutate(growth=DiamIf3-DiamIf2) %>%
  group_by(Codi) %>%
  summarise(av_growth=mean(growth), n=n()) %>%
  arrange(desc(av_growth))
```

A tibble: 7,713 x 3

```
##
      Codi
              av_growth
                              n
##
      <fct>
                   <dbl> <int>
##
    1 171089
                    23.1
                              3
    2 170819
                    21.6
##
                              1
##
    3 172607
                    17.6
                              6
##
    4 172216
                    17.4
                              6
##
    5 172690
                    16.0
                             17
##
    6 171682
                    15.4
                              6
##
    7 083267
                    15.3
                              1
##
    8 431363
                    15.1
                              4
    9 171664
                    14.8
                              5
## 10 171976
                    14.4
                              1
## # ... with 7,703 more rows
```

• Ex.6.2 Which is the plot with highest species richness?

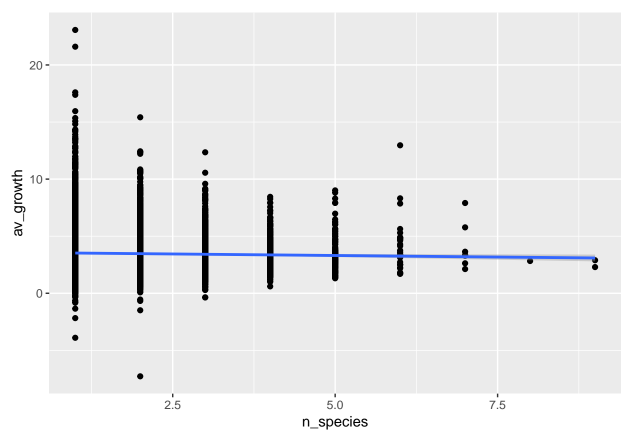
First, we define the data frame (trees), *THEN* we group by Code, *THEN* we determine the number of species per plot and *THEN* we arrange in decreasing order:

```
trees %>%
  group_by(Codi) %>%
  summarise (n_species=n_distinct(Especie)) %>%
  arrange(desc(n_species))
```

```
## # A tibble: 7,713 x 2
##
      Codi
              n_species
##
      <fct>
                  <int>
##
    1 170195
                      9
##
    2 171036
                      9
##
    3 170218
                       8
##
    4 170121
##
    5 170596
##
    6 170635
                       7
                       7
##
    7 170799
    8 171398
                       7
                       7
##
   9 171481
## 10 172650
                       7
## # ... with 7,703 more rows
```

• Ex.6.3 Are both variables (species richness and average growth) related?

First, we would need to indicate the data frame we will work with, *THEN* we will group by plot, *THEN* we will calculate the variables. To see the relationship between both variables, we will create a plot with ggplot2, just to show how all the packages in the tidyverse relate to each other. The aim of this workshop is not to learn ggplot2, so we won't go into further details. To know more about ggplot you can visit this website: http://ggplot2.org/.



Here we see one of the advantages of the tidyverse, the fact that all the packages and functions can communicate one with another. In this way, we just created a plot without the need of creating intermediate objects or data frames, starting directly from the raw data frame, and chianing orders in a logical and intuitive way.

Grouped mutate/grouped filter

Most of the times we use group_by, we will do it with the *summary functions*, that is, functions that take n values as input, and give back 1 value as output. Examples of *summary functions* are mean(), sd(), min(), sum(), etc.

However, some times we will need to do some operation by group, but we will need to produce one output per input, that is n inputs -> n outputs. This can be done using mutate or filter in combination with group_by.

###Exercise 7

Taking this into account, let's try to:

• Ex.7.1 Identify those trees that grow much faster than the average of the plot

In the previous code we see we first calculate the growth of each tree, and after grouping by plot, we calculate

a new variable, where the plot average is substracted from the growth of each tree, and the result is divided by the standard deviation of the plot. We calculate in this way the standardized growth of each tree with respect to the plot, making it easy to identify those trees that grow suspiciously more than the average for their plot.

• Ex. 7.2 Identify those plots where a species grows much more than the average for the species

```
trees %>%
  mutate(growth=DiamIf3-DiamIf2) %>%
  group_by(Especie) %>%
  mutate(growth_sp = mean(growth)) %>%
  group_by(Codi, Especie) %>%
  mutate(growth_sp_plot = mean(growth),
        inc = (growth_sp_plot /growth_sp))%>%
  arrange(desc(inc))
```

As we did before, we first calculate the growth of eaxh tree, we then group by species, so that we can calculate the mean growth for each species (growth_sp). Finally, we group again, now for plot and species, to calculate the mean growth of each species on each plot (growth_sp_plot). Once we have this, we can calculate the ratio between the two variables, identifying those plots where the species is performing better)assuming no mistakes, of course).

Let's see one last example:

*Ex.7.3 Select those species of the IFN3 occupied by "pure" Pinus nigra stands (Especie = 025)

Note: a forest is considered as pure stand if more than 80% of their Basal Area corresponds to a single species. Let's see how we would do that:

```
## # A tibble: 648 x 5
## # Groups:
               Codi [648]
##
      Codi
             Especie BA_sp BA_tot ratio
##
      <fct> <fct>
                     <dbl>
                             <dbl> <dbl>
    1 080132 025
                             49.9 0.830
##
                     41.4
##
    2 080307 025
                     51.2
                             54.8 0.933
    3 080313 025
##
                     28.1
                             35.0 0.804
   4 080318 025
                     23.7
                             26.6 0.894
##
##
    5 080322 025
                     29.7
                             32.3 0.919
##
    6 080323 025
                      1.61
                             1.61 1
##
   7 080324 025
                     18.2
                             19.7 0.924
##
   8 080325 025
                     43.8
                             44.6 0.982
    9 080326 025
                      6.01
                             6.01 1
## 10 080328 025
                             65.7 0.945
                     62.1
## # ... with 638 more rows
```

In this case, we first calculate BA per plot, using summarise. We then calculate the sum of BA per plot, but in this case we use mutate, because we don't want to aggregate tge data by plot, but calculate them separately for each plot but keeping the rest of the data as it was. Once we have both values, we can filter to select those plots with Pinus nigra, in which percentage of basal area for that species be > 80%.

Joins: working with to tables

Very often, the information we will work with more than a table. The *join* functions will allow us to work with several data frames, joining them in different ways. Within dplyr there are two types of joins:

Mutating joins

They add the columns of a data frame to the other, depending on whether they share some observations or not. There are four types.

- left_join(x, y) adds the columns of y to the observations of x that are also in y. Those that are not present in y will receive the value NA. With this function we ensure that we will not lose any observation.
- right_join(x, y) adds the columns of x to those observations in y that are also in x. Those that are not present will receive NA. It is equivalent to left_join, but the columns will be ordered differently.
- full_join(x,y) includes all observations in x and y. If they do not coincide, they assign NA.
- inner_join(x, y) includes only those observations both in x and y (repeats rows if it is necessary).

Filtering joins

The second type of joins are the **filtering joins**, that affect only to the observation, not to the variables. That is, they never add new columns, but they keep or delete the rows of the original frame as a function of their correspondence or not with a second data frame. There are only two types:

- semi_join(x, y) keeps the observations in x that match observations in y.
- anti_join(x, y) deletes the observations in x that match observations in y.

You can find more information about the join functions typing vignette("two-table").

###Exercise 8 To try the join functions, let's add the geographic information (X & Y coordinates), contained in the data frame coordinates to the data frame plots.

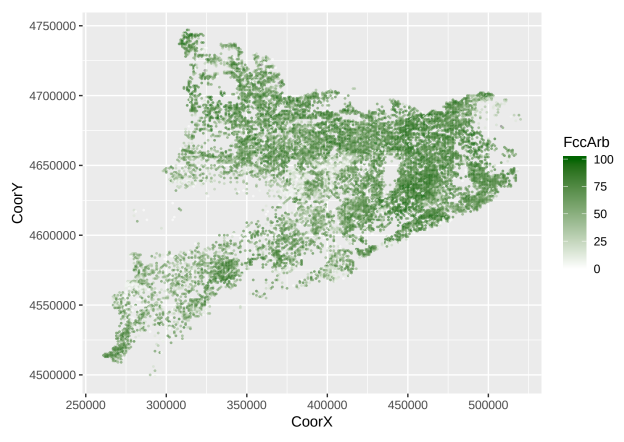
```
left_join(plots,coordinates, "Codi")
```

In this case, since we want to keep all the plots in the original plotsdata frame, we use left_join. In this case, since the number of observations in coordinates and plots is the same, the function inner_join should give us the same results.

Now we added the coordinates, we can represent in a map any variable in the data frame. We could, for example, represent the values of tree canopy cover (FccArb). We need to load the package "maps". (If we don't have it installed, we can install it typing in the console install.packages("maps")).

```
library(maps)
```

```
##
## Attaching package: 'maps'
## The following object is masked from 'package:purrr':
##
## map
left_join(plots,coordinates, "Codi") %>%
ggplot( aes(CoorX, CoorY)) +
    geom_point(aes(color=FccArb), size=0.3, alpha=0.6) +
    scale_color_continuous(low= "white", high="dark green")
```



Once again, we see we don't even need to create a new data frame with the new information, we can chain the functions in dplyr and ggplot2, producing the results in a very easy and fast way.

tidyr: changing the shape of the data frames

One of the main ideas behind the tidyverse is the concept of tidy data, that we have introduced before. According to Hadley Wickham, we can say that our data are tidy when two conditions are met:

- Each column corresponds to a variable
- Each row is a different observation

Of course, the data are not always organized in this way, sometimes other formats are more efficient (for example, for gathering data). For instance, if we have a look at the table **species**we will see that the number of trees for the different size classes are in different columns. This format is more convenient for entering the data or for some kinds of analyses, but in general the *tidy* format easens the processing and analysis, specially in vectorized languages such as R.

View(species)

The tidyr package allows to change the way in which data are organized, so that we can arrange them in the way we need to our analysis. It has four basic verbs:

- gather aggregates variables that are in several columns and converts them into two variables: a factor (key) and a numeric variablea (value).
- spread is the inverse of gather, it takes the levels of a factor and a numeric variable and creates a new variable for each level of the factor.
- separate divides the content of a column into several columns
- unite inverse of separate, concatenates the values of several columns

gather & separate

gather transforms data in *wide* format into *long* format. gather takes a series of columns and transforms them into two variables: a factor (*key*) and a numeric variable (*value*). The first parameter in gather is the data frame, the second and third are the names we will give to *key* and *value*, and the rest are the variables to group.

###Exercise 9 Let's use gather and separate to transform the data frame species into a *tidy* format, where each column is a variable and each row, an observation. First, let's have a look at the data frame we want to transform:

glimpse(species)

To convert it into the 'long' format we specify the data frame, then the new factor to create (key), and the numeric variable (value), and last, the columns to aggregate. For the last part we have three equivalent options:

• (A) Explicitly define the variables we want to gather:

• (B) Define the interval containing the variables we want to gather

```
gather(species, CD, n, CD_10:CD_70)
```

• (C) Define the variables with the helpers functions

```
gather(species, CD, n, starts_with('CD'))
```

• (D) Define the variables we DO NOT want to include (with -). The function will assume we want to gather the rest of variables.

```
gather(species,CD, n,-Codi, -Especie)
```

The three pieces of code above produce the same result. Once we have converted the data frame into the new format, we can divide the new variable "CD" into two new variables, that we will name "Name" and "CD", using separate. If we do not specify where to make the separation, the function takes by defualt the first non alphanumeric character in the string.

```
species_long <- gather(species,CD, n,-Codi, -Especie)
species_long<-separate(species_long, col=CD, into = c("Nombre", "CD"))
species_long</pre>
```

spread & unite

If we have a data frame in *long* format, we can use **spread** and **unite** to transform it back into the *wide* format. This is what we will do in the next exercise, converting back the data frame with species and diameter classes into its original format. As with **gather**, **spread** takes the data frame as the first argument. The second parameter is the factor we will use to create the new columns, and the third parameter is the name of the column that contains the values. We can see this with an example:

###Exercise 10 Use unite and spread to transform back the data to its original format.

First we create a new variable, that will be useful to create the new columns:

```
species_unite <- unite(species_long, CD, Nombre, CD)</pre>
```

Now we will transform the data frame, specifying the variable that will produce the new columns ("CD") and the variable that contains the values (n)

```
spread(species_unite, CD,n)
```

purrr fuctional programming

purr allows for functional programming in R, meaning that functions becomes "first citizens'in the R environment (they can be used as arguments or returned from other functions).

Making loops pipe friendly

You can think of map, the main verb of purrer, as a *pipe-friendly* version of the apply family. It will accept functions as arguments and they will be applied to all elements of the list/vector provided.

Exercise 11

First thing is grouping by the plot code (Codi), but also, and this is important, by Province as we will need it later. Then we summarise the height with the mean, join the leaf dataset, split by province an map the lineal model. Optionally, we can use broom or summary to see the results:

```
trees %>%
  group_by(Codi, Provincia) %>%
  summarise(height = mean(HeiIf3, na.rm = TRUE)) %>%
  left_join(leaf, by = 'Codi') %>%
  split(.$Provincia) %>%
  map(~ lm(height ~ leaf_biomass, data = .)) %>%
  map_dfr(broom::tidy)
```

Other interesting functionalities of the tidyverse

Communication between packages

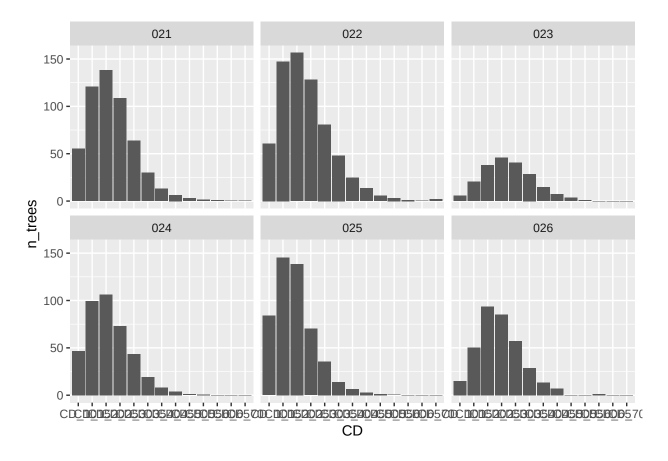
All the packages in the tidyverse are designed to communicate with each other. That means that we can combine dplyr and tidyr functions, connecting them trhough pipes (%>%). For example, if we wanted to know which the mean diameter distribution of all the pine species we could first filter the species we want to study (pines) then transform the data frame, group the data by species and diameter class and finally calculate the mean number of trees:

```
species %>%
  filter(Especie %in% c("021", "022","023", "024", "025", "026")) %>%
  gather(CD, n, CD_10:CD_70) %>%
  group_by(Especie,CD) %>%
  summarise(n_trees=mean(n))
```

```
## # A tibble: 78 x 3
               Especie [?]
## # Groups:
      Especie CD
##
                     n_trees
                       <dbl>
##
      <fct>
              <chr>>
##
   1 021
              CD_10
                       55.4
##
    2 021
              CD 15 121.
    3 021
              CD 20
##
                      138.
   4 021
              CD_25
                     109.
##
##
   5 021
              CD 30
                       63.7
##
    6 021
              CD_35
                       29.8
##
    7 021
              CD_40
                       13.2
##
   8 021
              CD_45
                        6.43
##
   9 021
              CD_50
                        3.00
              CD_55
                        1.39
## 10 021
## # ... with 68 more rows
```

But dplyr and tidyr also can connect to other packages in the tidyverse, such as ggplot2 or broom, so we could expand on the previous code to generate a plot by species.

```
species %>%
    filter(Especie %in% c("021", "022","023", "024", "025", "026")) %>%
    gather(CD, n, CD_10:CD_70) %>%
    group_by(Especie,CD) %>%
    summarise(n_trees=mean(n)) %>%
    ggplot(aes(x=CD, y=n_trees)) +
    geom_col() +
    facet_wrap(~Especie)
```



Functional sequences

Another interesting aspect of dplyris that we can save sequences of orders as an object, so they can later be applied to different data frames, as if it was a function. To do this, we must use the pronoun . as data frame in the sequence of orders to save. Let's see an example:

```
av_growth <- . %%
mutate(growth=DiamIf3-DiamIf2) %>%
group_by(Codi) %>%
summarise(mean=mean(growth), n=n())
```

If we print the object, we will see it has a class functional sequence, and it specifies the orders to execute: av_growth

```
## Functional sequence with the following components:
##
## 1. mutate(., growth = DiamIf3 - DiamIf2)
## 2. group_by(., Codi)
## 3. summarise(., mean = mean(growth), n = n())
##
## Use 'functions' to extract the individual functions.
```

We can then apply this sequence to a data frame...

```
trees %>% av_growth()

## # A tibble: 7,713 x 3

## Codi mean n

## <fct> <dbl> <int>
```

```
##
    1 080001
               3.33
                        15
##
    2 080002
                        13
              3.63
    3 080003
##
               5.93
                        7
                        2
##
    4 080004
               6.55
##
    5 080005
               2.08
                        12
    6 080006
               2.28
                        23
##
    7 080007
##
               2.45
                        35
##
    8 080008
               1.79
                        11
##
    9 080009
               1.86
                        16
## 10 080010 3.33
                        13
## # ... with 7,703 more rows
```

 \dots or combine it with new dplyr or tidyr functions

```
trees %>%
  filter(Provincia=="17") %>%
  av_growth()
```

```
## # A tibble: 2,113 x 3
##
      Codi
                        n
               mean
##
      <fct>
              <dbl> <int>
##
    1 170004
               2.99
                       39
##
    2 170005
               1.86
                        28
    3 170006
##
               1.75
                        31
##
    4 170007
               3.14
                        29
##
    5 170008
               1.68
                        9
    6 170009
               1.48
                        28
##
                        26
##
    7 170010
               2.40
    8 170012
               2.11
                        21
##
    9 170013
               1.55
                        34
## 10 170014
              5.38
                         8
## # ... with 2,103 more rows
```

Databases

In this workshop we've seen how to work with dplyr and tidyr using data stored in our computer, but dplyralso allows working woth remote databases, admiting most formats and standards: PostgreSQL, MySQL, SQLite, MonetDB, BigQuery, Oracle...

When working with databases we will use the same verbs and coding we've seen so far, but <code>dplyrtransforms</code> the R code into <code>SQLsequences</code>, so we don't need to change the language to read and analyze the data. Also, it is much faster than R. The details of working with databases are beyond the scope of this workshop, but you can find more information in the resources listed in the following section.

More info

Both the code and the data needed to generate this document and execute the examples can be found in GitHub (https://github.com/ameztegui/dplyr_workshop). You can also find more information about these packages and their functions in the book R for data science by Hadley Wickham, or in the vignettes for each function.

```
# Sobre dplyr
vignette("introduction")

# Sobre tidyr
vignette("tidy-data")

# Sobre unir dos tablas mediante join
vignette("two-table")

# Sobre trabajo con databases
vignette("databases")
```