

Introduction to R - Part 2

September 11, 2019

Contents

Objectives for this part	1
R packages	2
Data visualization with <i>ggplot2</i>	2
Create a scatter plot	2
Exercise 1	4
Types of graphics: the <i>geom</i> functions	4
Exercise 2	8
Divide a graph in facets	8
Exercise 3	11
Customize the appearance of graphs	11
Summary	16
Data manipulation with <i>dplyr</i>	16
Data organization	16
Select observations with <i>filter</i>	17
Exercise 4	18
Sort observations with <i>arrange</i>	18
Pick variables with <i>select</i>	19
Chain operations with <i>%>%</i>	19
Exercise 5	20
Derive new variables with <i>mutate</i>	20
Compute summary statistics per group: <i>group_by</i> and <i>summarize</i>	20
Exercise 6	21
Join two data frames	21
Summary of <i>dplyr</i> functions	22
References	22
Solutions to the exercises	23
Exercise 1	23
Exercise 2	23
Exercise 3	24
Exercise 4	25
Exercise 5	25
Exercise 6	26

Objectives for this part

- Load *packages* to add functions to R.
- Produce graphics with the *ggplot2* package.
 - Create different types of graphics: point clouds, boxplots, histograms and more.
 - Create composite graphics.

- Customize the appearance of graphical elements.
- Manipulate data frames with the *dplyr* package.
 - Filter and sort observations.
 - Transform the variables.
 - Calculate statistics by groups of observations.
 - Join two tables according to common variables.

R packages

- A package is a collection of functions developed by R users that increases the basic capabilities of the R language in a certain domain. The CRAN (<https://cran.r-project.org/web/packages/>) package repository currently has more than 12,000!
- To **install** a package on your computer, use the `install.packages` function with the package name in quotation marks, e.g. `install.packages("ggplot2")`. In RStudio, you can view the list of packages already installed under the *Packages* tab in the lower right quadrant.
- To **load** a package into your current R session and have access to its functions, use the `library` function, e.g. `library(ggplot2)`. Note that quotation marks are not required this time.

Data visualization with *ggplot2*

There are several ways to produce graphs in R. We will use in this course the *ggplot2* package, which provides a consistent code structure for multiple types of graphs and offers a wide range of options to customize the appearance of the graphics.

```
library(ggplot2)
```

Create a scatter plot

The `msleep` dataset included with *ggplot2* contains data on sleep of 83 mammal species.

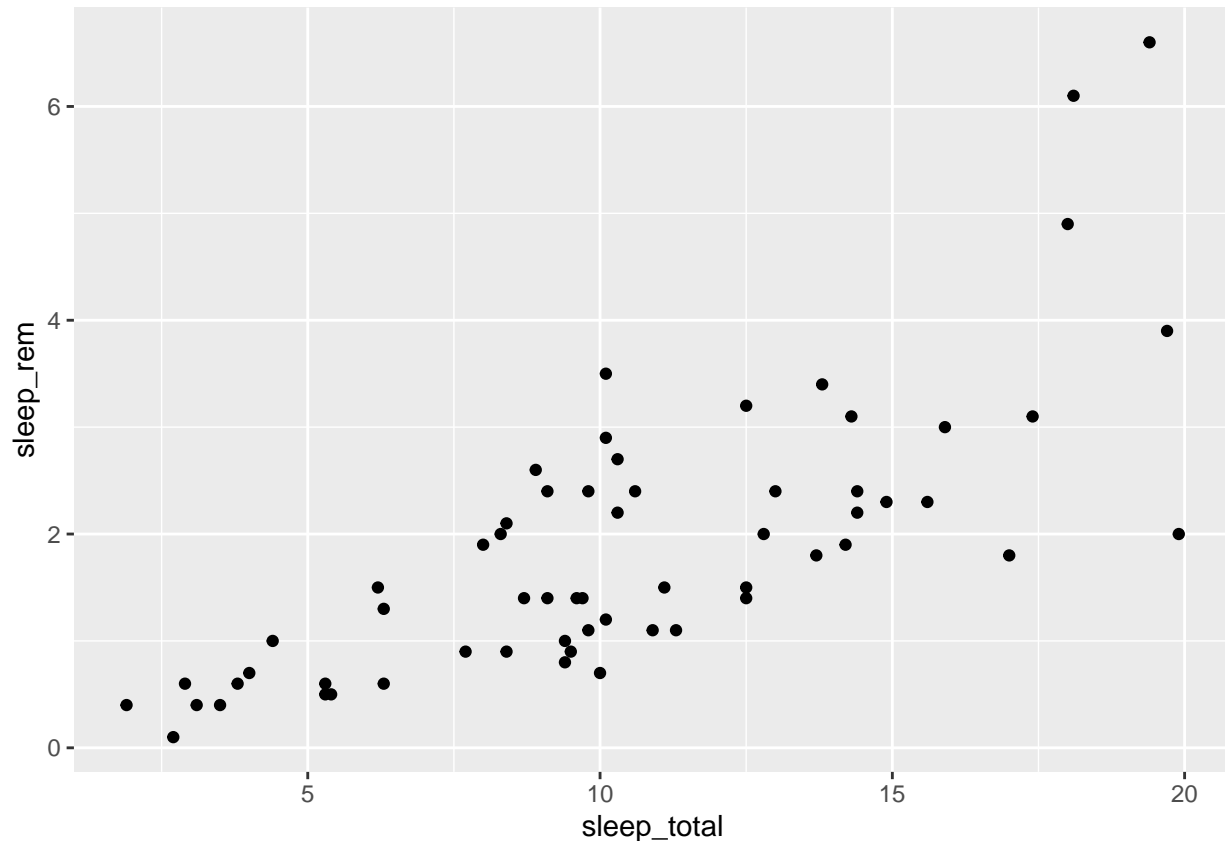
```
msleep
```

```
## # A tibble: 83 x 11
##   name genus vore order conservation sleep_total sleep_rem sleep_cycle
##   <chr> <chr> <chr> <chr> <chr>          <dbl>      <dbl>      <dbl>
## 1 Chee~ Acin~ carni Carn~ lc          12.1        NA        NA
## 2 Owl ~ Aotus omni Prim~ <NA>         17         1.8        NA
## 3 Moun~ Aplo~ herbi Rode~ nt          14.4        2.4        NA
## 4 Grea~ Blar~ omni Sori~ lc          14.9        2.3        0.133
## 5 Cow   Bos   herbi Arti~ domesticated      4         0.7        0.667
## 6 Thre~ Brad~ herbi Pilo~ <NA>         14.4        2.2        0.767
## 7 Nort~ Call~ carni Carn~ vu           8.7        1.4        0.383
## 8 Vesp~ Calo~ <NA> Rode~ <NA>           7         NA        NA
## 9 Dog   Canis carni Carn~ domesticated     10.1        2.9        0.333
## 10 Roe ~ Capr~ herbi Arti~ lc           3         NA        NA
## # ... with 73 more rows, and 3 more variables: awake <dbl>, brainwt <dbl>,
## #   bodywt <dbl>
```

Enter the following code to produce a scatter plot showing the number of hours of the REM phase (*sleep_rem*) based on the total number of hours of sleep.

```
ggplot(data = msleep, mapping = aes(x = sleep_total, y = sleep_rem)) +  
  geom_point()
```

```
## Warning: Removed 22 rows containing missing values (geom_point).
```



The graph appears in the *Plots* tab of the lower right quadrant in RStudio. You can export it in .PNG or .PDF format using the *Export* menu.

This first example illustrates the basic structure for producing a graph with *ggplot2*:

- A call to the `ggplot` function with two arguments:
 - a data frame (`data`) and
 - a mapping specified with the `aes` function. Within this function, we associate elements of the graph with variables in the dataset (eg `sleep_total` on the *x* axis).
- The `+` symbol to indicate that we will add other components to the graph.
- A *geom* function to specify the geometric elements represented (here `geom_point`, for a scatter plot).

Note: After naming the data frame in the `data` argument, the `aes` function only requires the variable names, without quotation marks and without specifying the data frame again (`sleep_rem` instead of `msleep$sleep_rem`). This is a special property of the *ggplot2* package functions, which we will also find with the *dplyr* package below, as well as with the regression functions later in the lesson.

Exercise 1

First, load the Kejimikujik National Park Dataset that we used in the first lab:

```
kejim <- read.csv("cours1_kejimkujik.csv", stringsAsFactors = FALSE)
head(kejim)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece  dhp
## 1  BD         A   31    8  2004         1         1  TSCA 16.3
## 2  BD         A   31    8  2004         2         1  TSCA 24.0
## 3  BD         A   31    8  2004         6         1  TSCA 29.8
## 4  BD         A   31    8  2004         7         1  ACRU 29.0
## 5  BD         A   31    8  2004         8         1  TSCA 15.5
## 6  BD         A   31    8  2004         9         1  TSCA 32.0
```

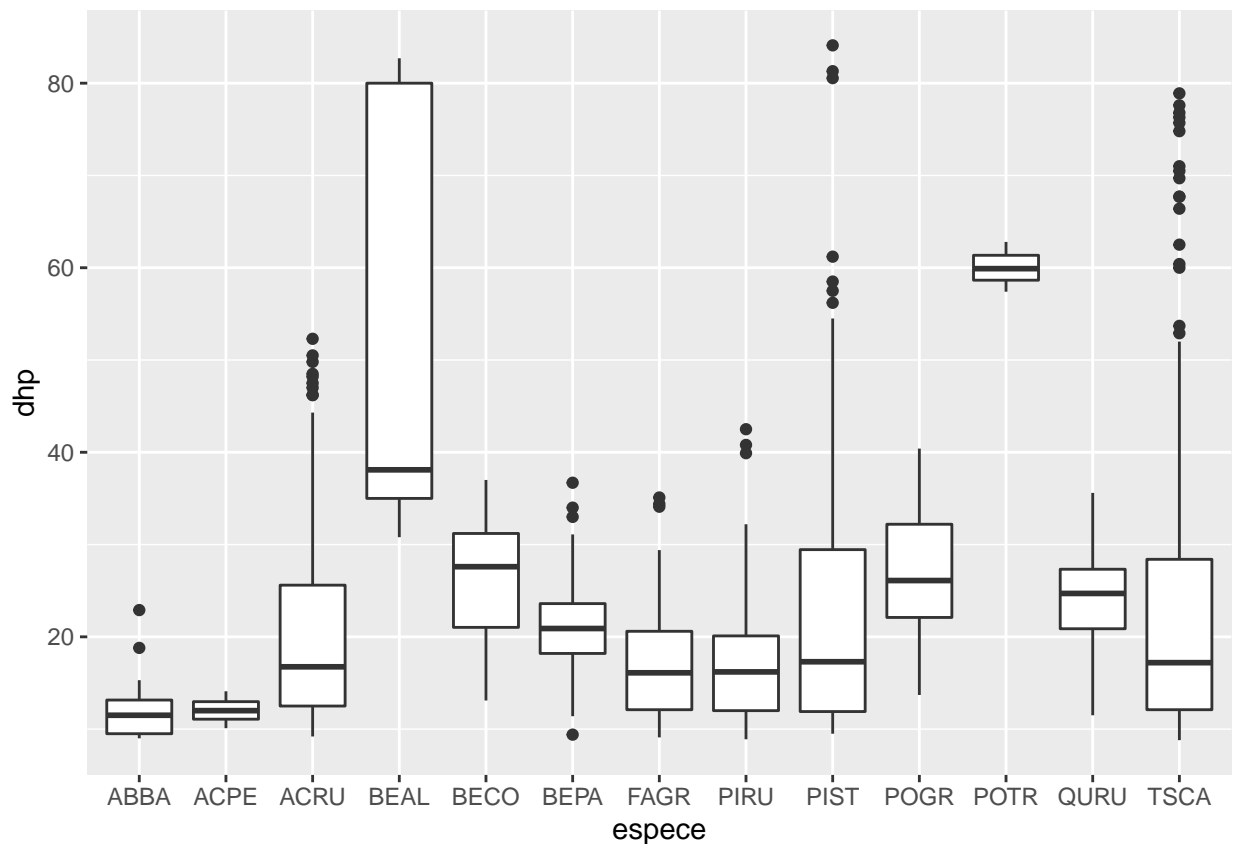
From this data frame, produce a scatter plot of the DBH (`dhp`, on the y axis) according to the species (`espece`, on the x axis).

Solution

Types of graphics: the *geom* functions

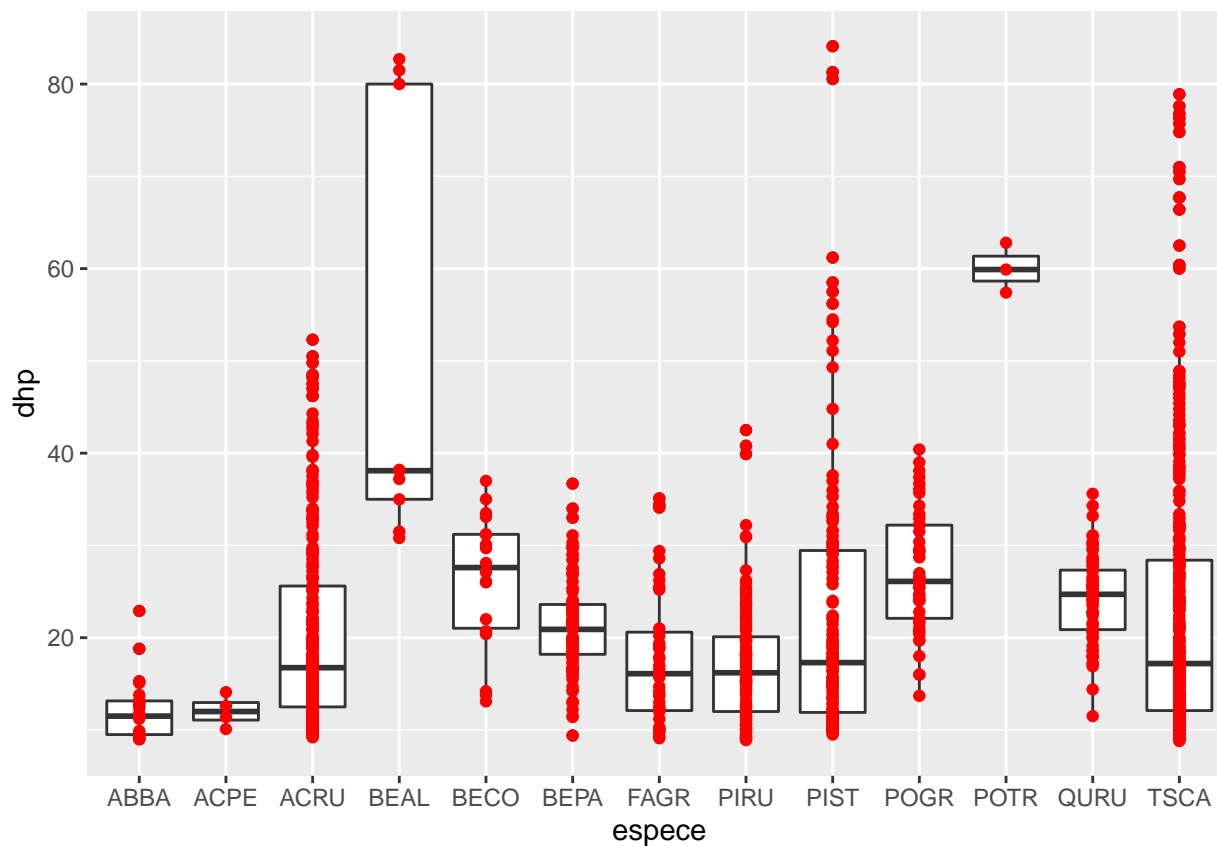
Here, there are too many trees to see the distribution of DBH by species with a scatter plot. To view the same data with boxplots, just replace `geom_point` with `geom_boxplot` in our previous code.

```
ggplot(kejim, aes(x = espece, y = dhp)) +
  geom_boxplot()
```



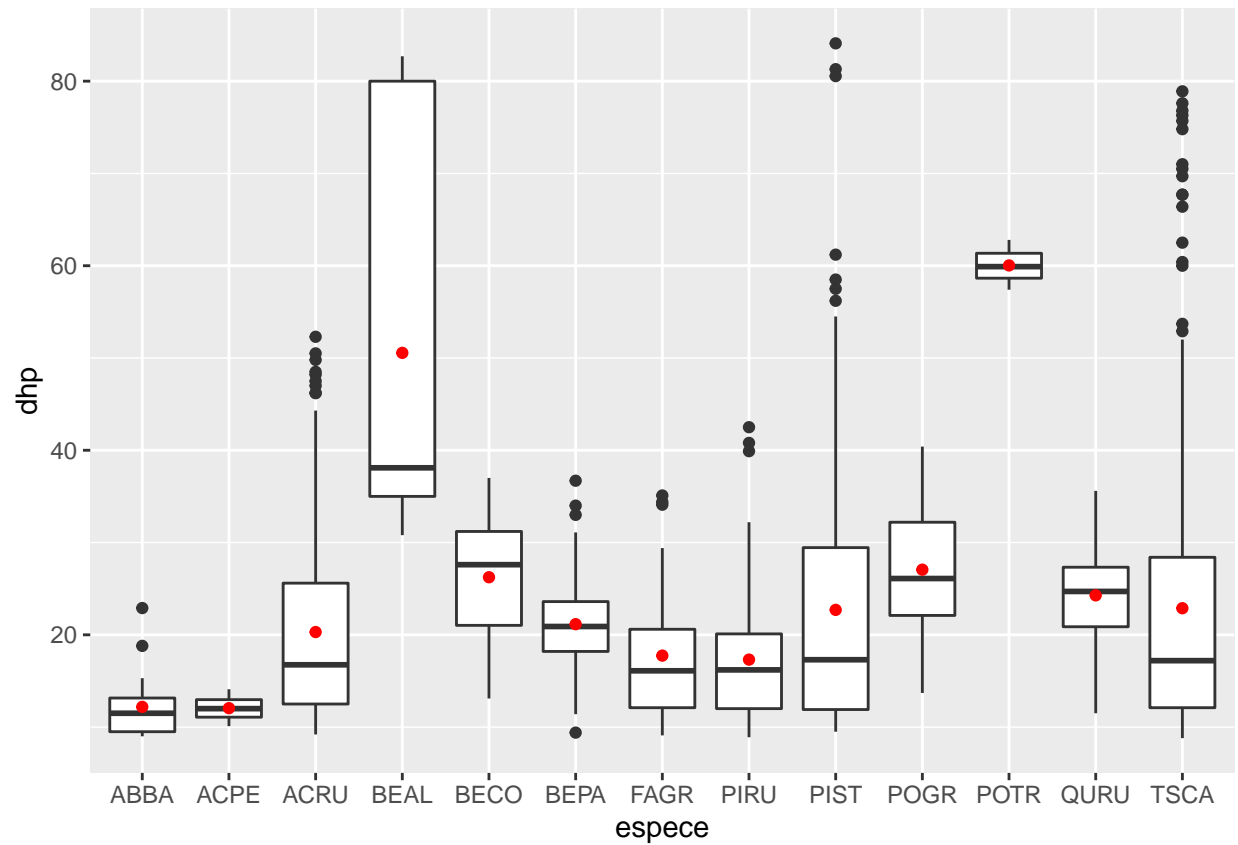
Multiple `geom_...` functions can be used to overlay elements on the same graph. So we add the dots to the boxplot, specifying a different color for `geom_point`.

```
ggplot(kejim, aes(x = espece, y = dhp)) +  
  geom_boxplot() +  
  geom_point(color = "red")
```



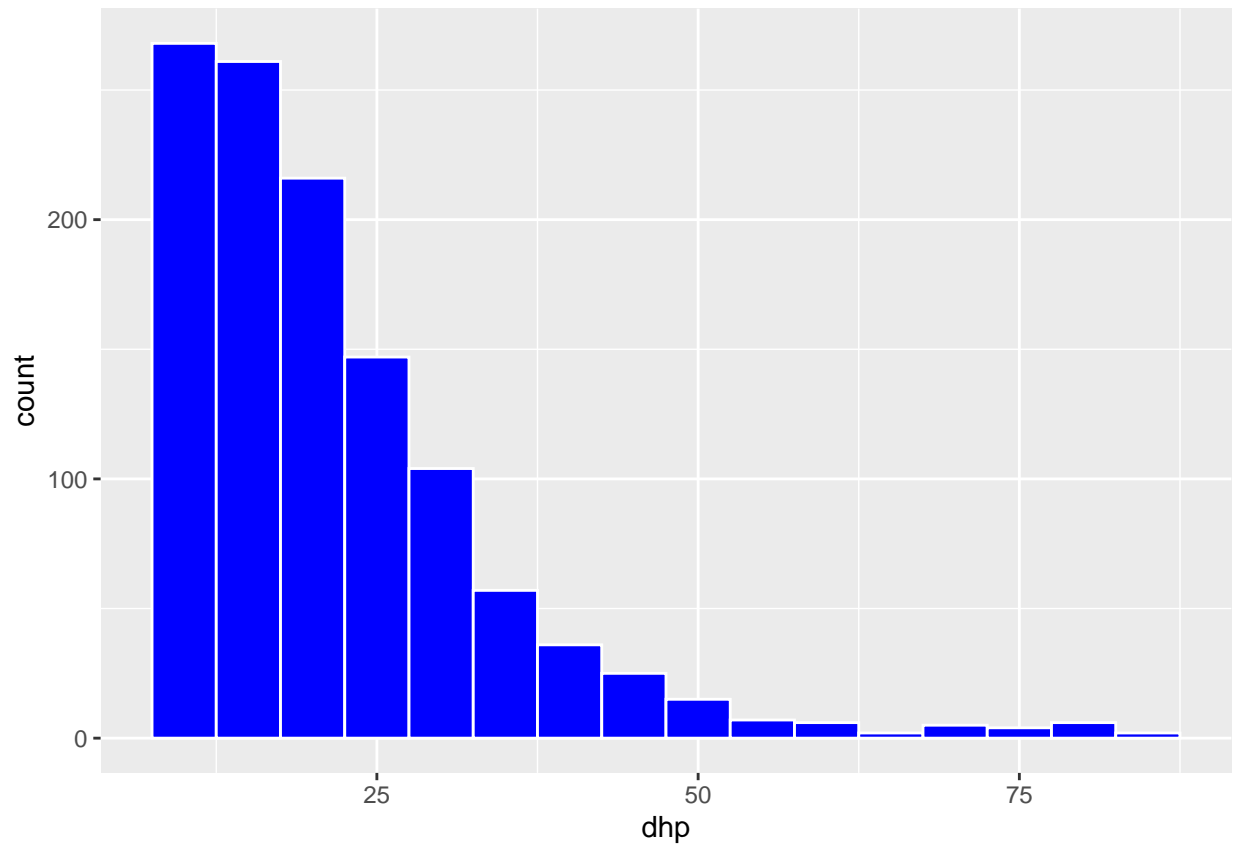
We can also represent summary statistics of a set of points, such as their mean.

```
ggplot(kejim, aes(x = espece, y = dhp)) +  
  geom_boxplot() +  
  geom_point(color = "red", stat = "summary", fun.y = "mean")
```



Now, we produce a histogram of the DBH for all trees with `geom_histogram`. The histogram represents a single variable, so we do not need to specify `y` in `aes`.

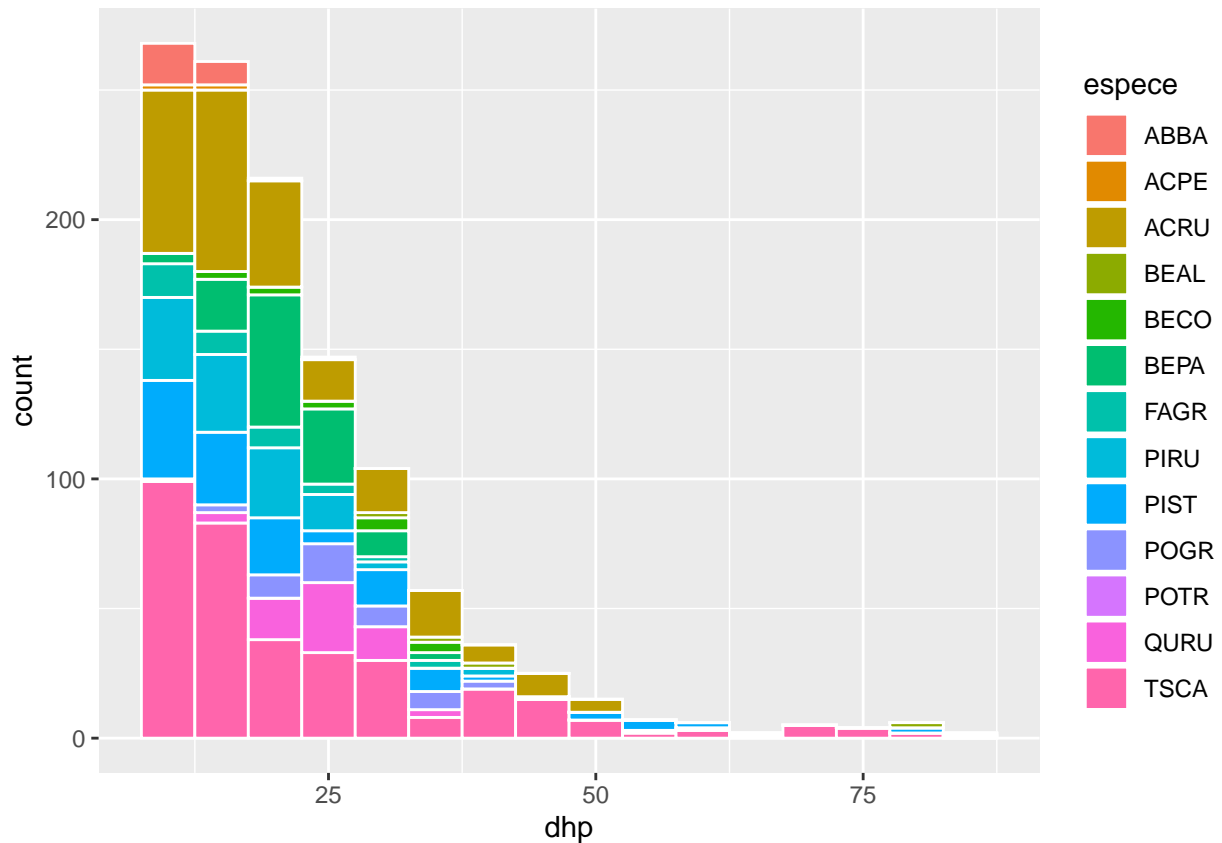
```
ggplot(kejim, aes(x = dhp)) +  
  geom_histogram(binwidth = 5, color = "white", fill = "blue")
```



In the `geom_histogram` function, we specified several arguments in order to choose the size of the bins (`binwidth`), the color of the border (`color`) and the fill color of the bars (`fill`).

Rather than using one fill color for all bars, we could represent different species by bars of different colors on the histogram. In this case, we must associate `fill` to the variable `espece` inside the `aes` function.

```
ggplot(kejim, aes(x = dhp, fill = espece)) +  
  geom_histogram(binwidth = 5, color = "white")
```



Exercise 2

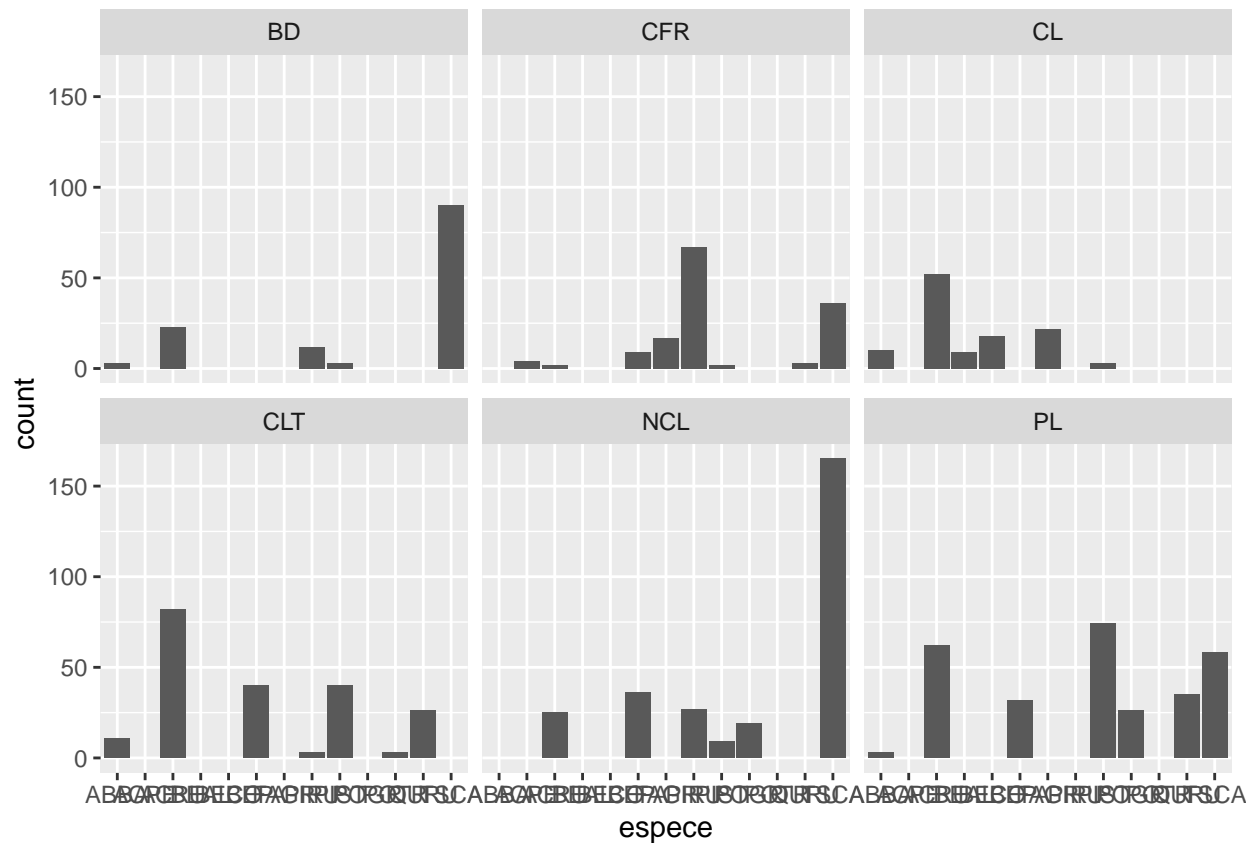
With `geom_bar`, create a bar graph of the number of individuals per species (different species on the x-axis). Use a color code to identify individuals from different sites.

Solution

Divide a graph in facets

In the exercise, we produced a graph of the number of individuals per species. If we want to visualize separately the distribution of species on different sites, we could divide the data by site and produce several graphs. Fortunately, *ggplot2* simplifies this task with the facet concept.

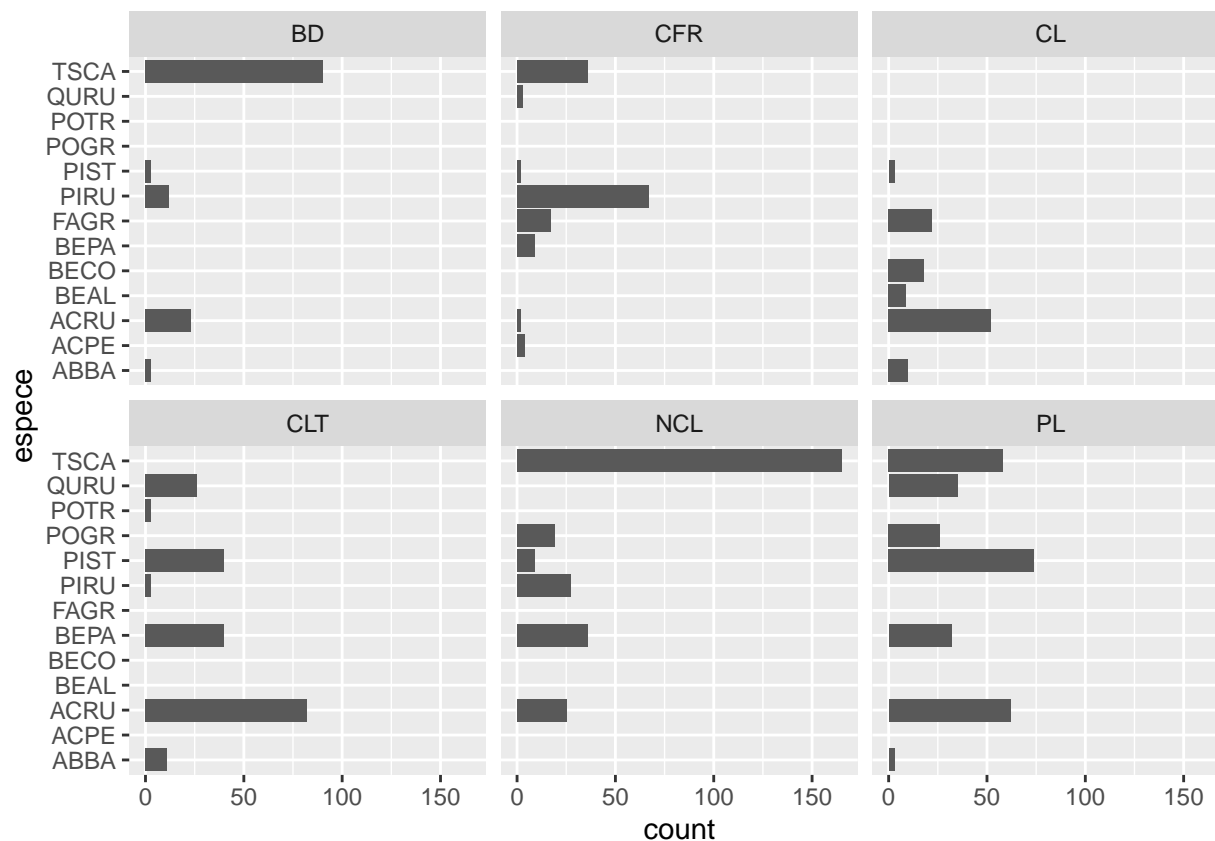
```
ggplot(kejim, aes(x = espece)) +
  geom_bar() +
  facet_wrap(~ site)
```

Note that you must put a *tilde* (~) before the name of the variable in `facet_wrap`.

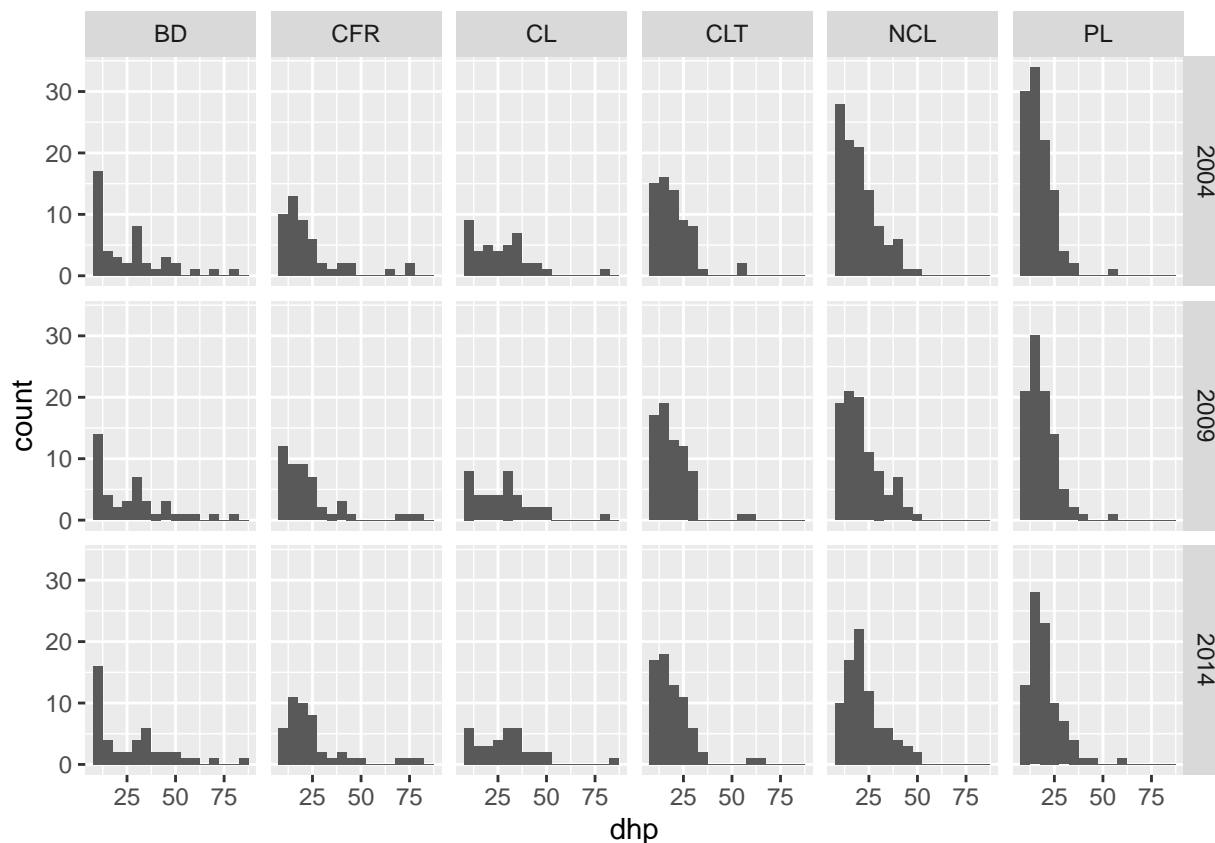
In the previous graph, species codes are not visible on the x axis. So we add the `coord_flip` function to invert the axes and produce horizontal bars.

```
ggplot(kejim, aes(x = espece)) +  
  geom_bar() +  
  facet_wrap(~ site) +  
  coord_flip()
```



We can also create a grid of facets (`facet_grid`) with two variables. For example, here are histograms of the DBH by site and by year.

```
ggplot(kejim, aes(x = dhp)) +
  geom_histogram(binwidth = 5) +
  facet_grid(annee ~ site)
```



Exercise 3

From the `msleep` dataset, create a scatter plot of total sleep (`sleep_total`) and REM sleep (`sleep_rem`) with one facet for each type of diet (`vore`).

Solution

Customize the appearance of graphs

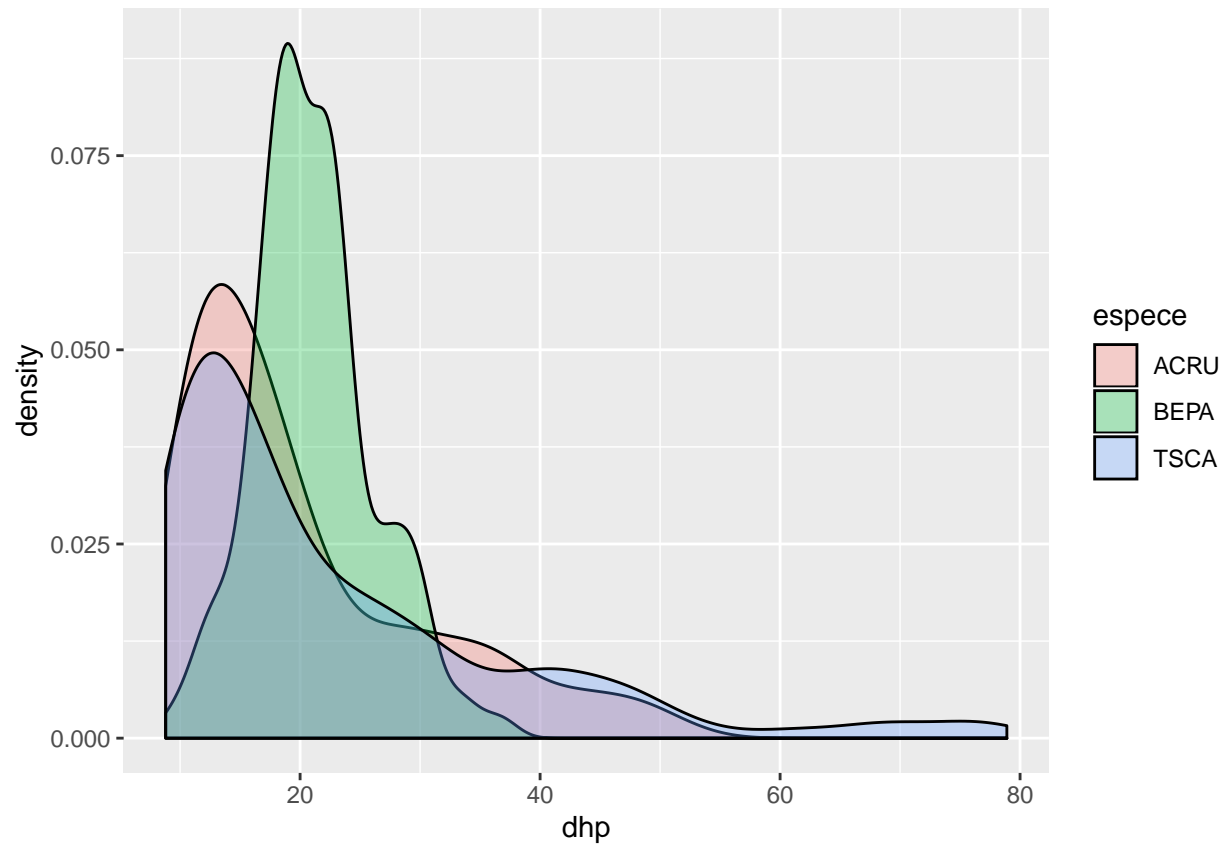
All visual aspects of *ggplot2* graphics can be customized: title and scale of axes, size and font, colors, margins, etc. If the default values of these parameters are sufficient for a quick exploration of the data, the production of figures for publications or presentations requires further adjustments.

Let's start with a density chart for the DBH of three species (TSCA, ACRU and BEPA). We save this graph in an object (`dens`) so that we can modify it without rewriting the code of the basic graph.

```
trois_esp <- kejim[kejim$espece %in% c("TSCA", "ACRU", "BEPA"), ]
```

```
dens <- ggplot(trois_esp, aes(x = dhp, fill = espece)) +  
  geom_density(alpha = 0.3)
```

```
dens
```

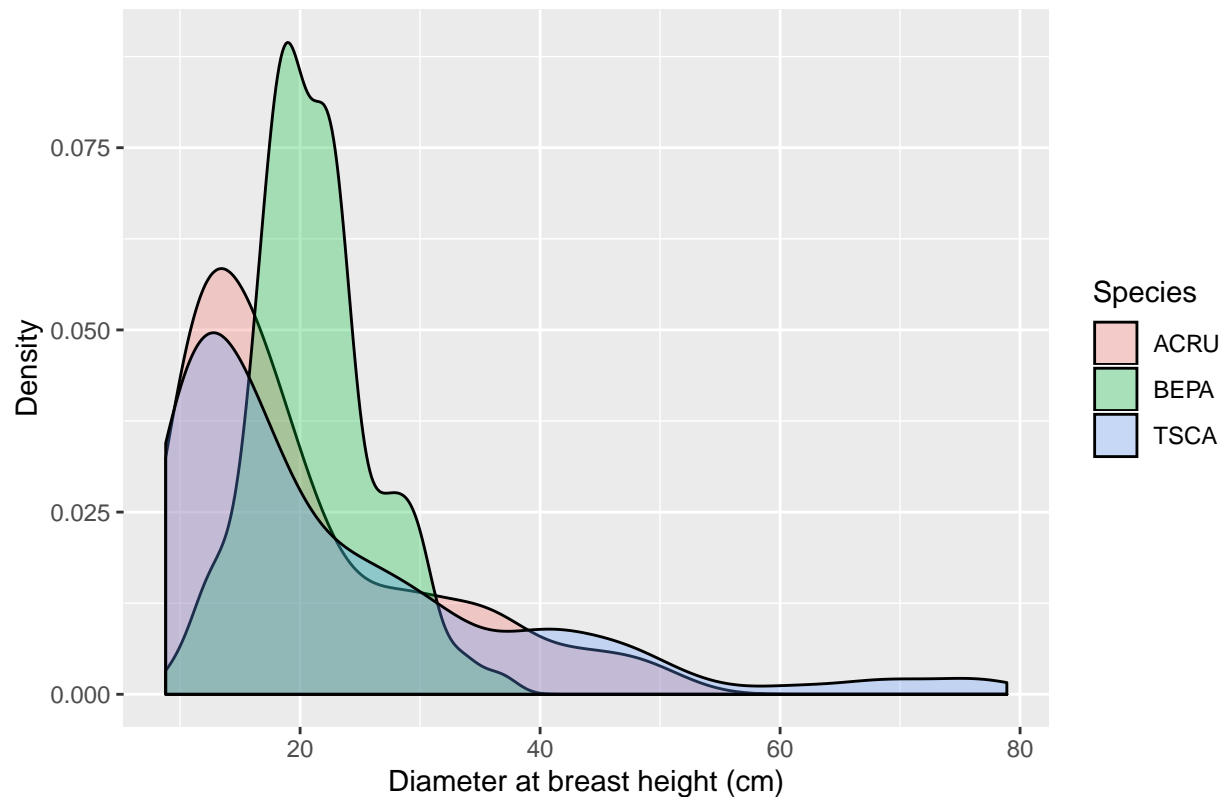


What does the `alpha` parameter means? Try changing its value.

The title of the chart, axes, and legend are specified with the `labs` function (for *labels*).

```
dens <- dens +
  labs(title = "Distribution of the diameter of three species at Kejimikujik National Park",
        x = "Diameter at breast height (cm)", y = "Density", fill = "Species")
dens
```

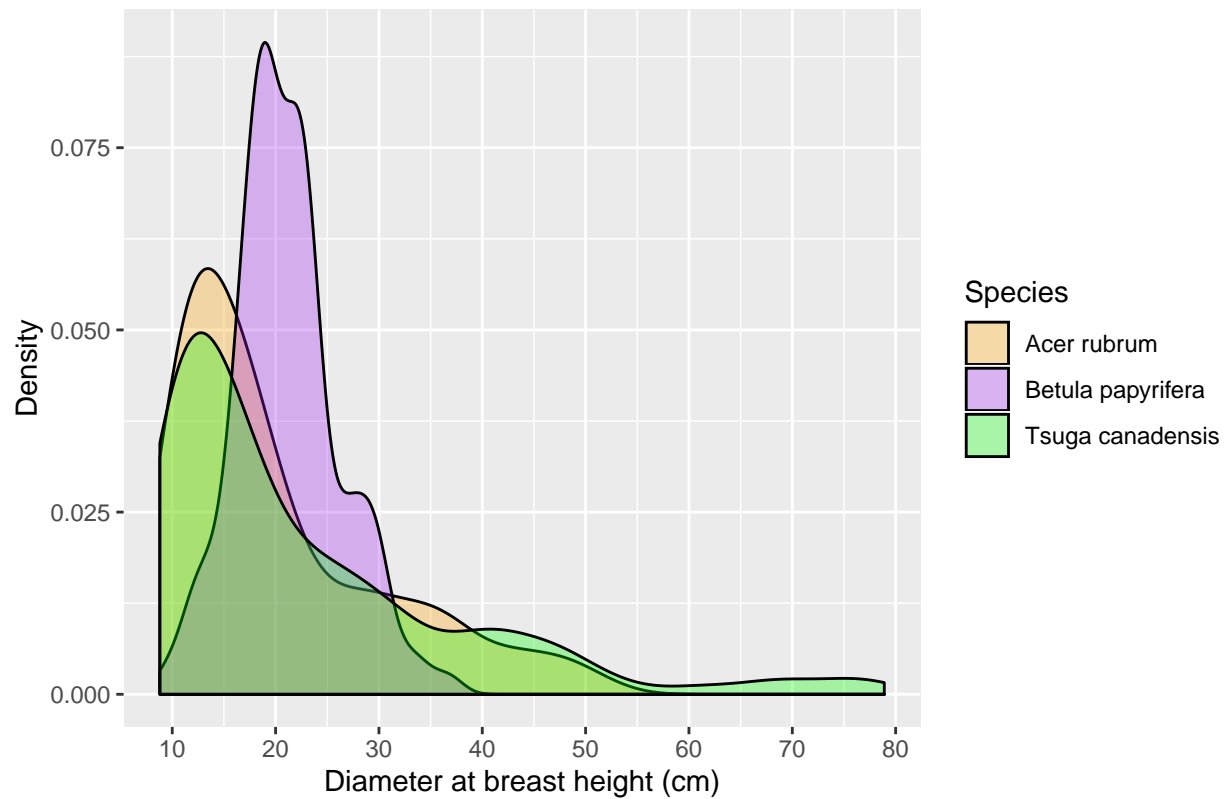
Distribution of the diameter of three species at Kejimikujik National Park



Functions starting with `scale_...` let us adjust different elements of the scales represented on the graph. In the following example, we change the values shown on the x axis with the `breaks` argument of `scale_x_continuous`. With the `scale_fill_manual` function, we specify the fill colors for the density curves (`values`), and then rename the legend elements (`labels`).

```
dens <- dens +
  scale_x_continuous(breaks = seq(10, 80, 10)) +
  scale_fill_manual(values = c("orange", "purple", "green"),
    labels = c("Acer rubrum", "Betula papyrifera", "Tsuga canadensis"))
dens
```

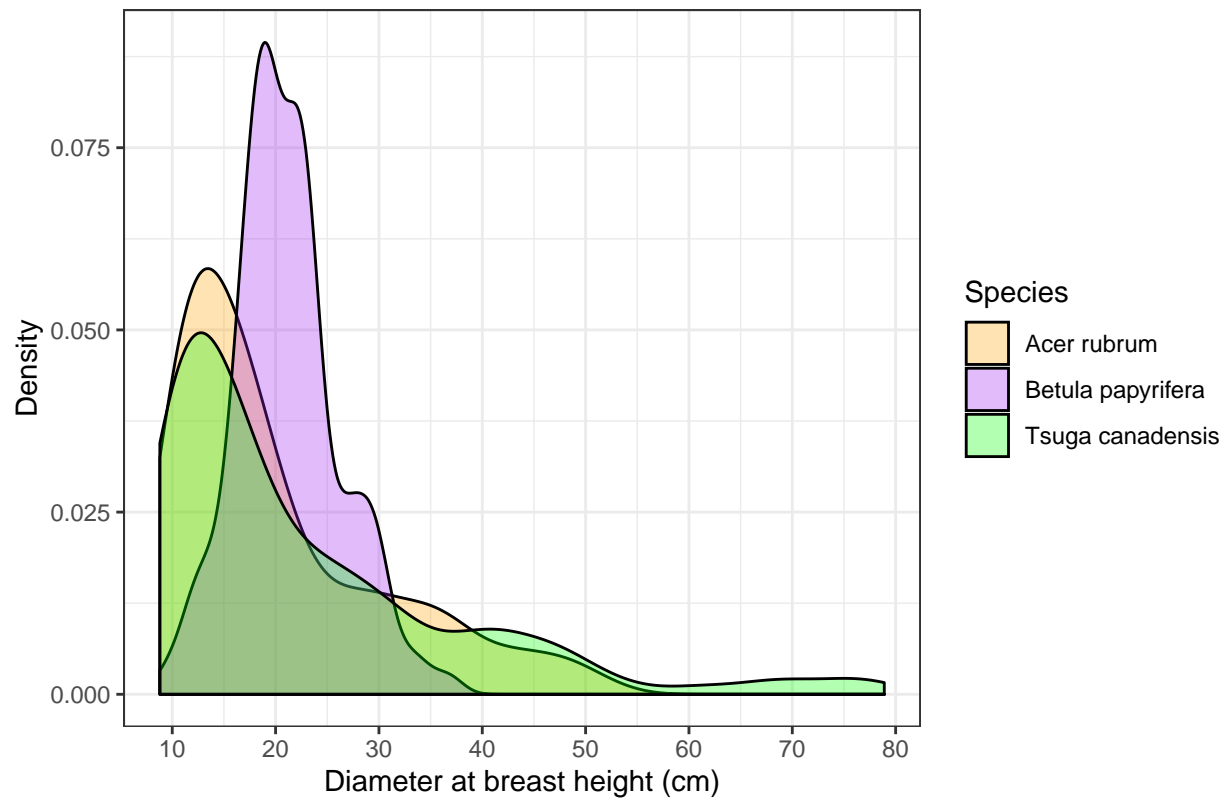
Distribution of the diameter of three species at Kejimikujik National Park



Finally, *ggplot2* has themes that change the overall appearance of the chart. Here we replace the default theme (with the gray background) with `theme_bw`.

```
dens +  
  theme_bw()
```

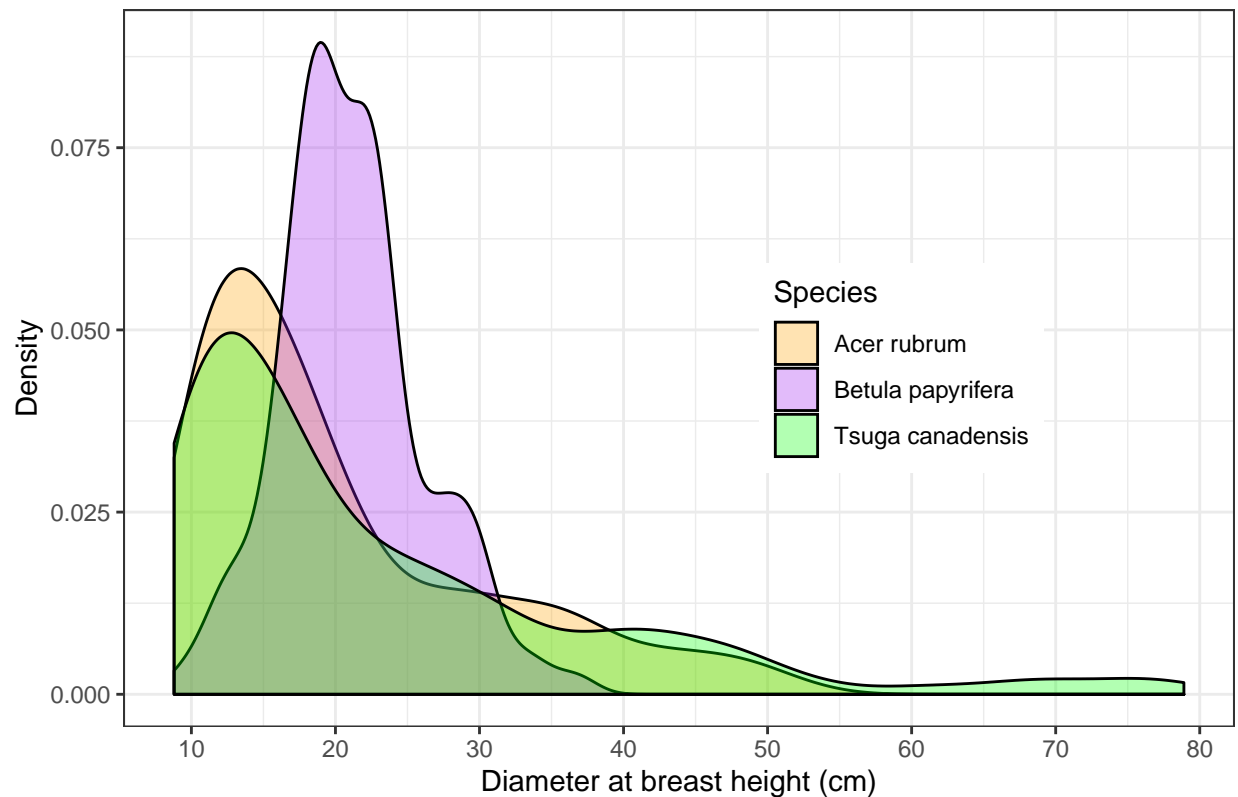
Distribution of the diameter of three species at Kejimikujik National Park



In addition to changing the entire theme, you can edit specific theme parameters with the `theme` function. For example, `legend.position` indicates where to put the legend relative to *x* and *y* (0.7 and 0.5 mean 70% of the way along *x* and 50% along *y*).

```
dens +  
  theme_bw() +  
  theme(legend.position = c(0.7, 0.5))
```

Distribution of the diameter of three species at Kejimikujik National Park



The purpose of this part was to give an overview of the possibilities of *ggplot2*. More complete references on this package are indicated at the bottom of the page.

Summary

- The code of a graph is composed of several functions separated by `+`.
- We start with the `ggplot` function, which requires two arguments: the data frame, and the association between variables and elements of the graph (with the `aes` function).
- We then add one or more geometric layers (`geom_...`) indicating the type of graph (points, lines, bars, histogram, etc.).
- If necessary, we can then modify the appearance of the graph with functions like `scale_...`, `facet_...`, `labs` and `theme`.

Data manipulation with *dplyr*

Data organization

The *dplyr* package contains functions that simplify several common operations on data frames in R. These functions are best suited for data frames in a normal form:

- each row corresponds to an observation and
- each column corresponds to a variable.

These criteria are close to the principles of data organization in a relational database (e.g. Access). In addition, the *dplyr* functions correspond fairly well to basic operations in the SQL database language.

Question: What are the variables in the following table, which indicates the number of individuals per site and species? Does it follow the normal form described above?

```
##   site fir pine birch
## 1   A  35  10  26
## 2   B  24  12  45
## 3   C  51  19   8
```

Answer: The last three columns represent the same variable (number of individuals) for different values of the species. A normalized table would then have three columns: site, species, and number.

```
##   site species number
## 1   A     fir     35
## 2   B     fir     24
## 3   C     fir     51
## 4   A    pine     10
## 5   B    pine     12
## 6   C    pine     19
## 7   A   birch     26
## 8   B   birch     45
## 9   C   birch      8
```

By formatting your data in this way, it will also be easier to visualize (as we will see later) and to model in R.

The *tidyr* package contains functions to automatically convert an array of data between the two formats above. We do not have time to cover this package today, but you can find more information in the references at the bottom of the page.

For this lab, we will use the Kejimikujik National Park dataset, which is already in normal form.

Select observations with *filter*

At the last class, we saw how to choose rows from a data frame according to a given condition. For example, the following code retrieves all rows corresponding to the species *ACRU* (red maple).

```
acru <- kejim[kejim$espece == "ACRU", ]
head(acru)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece dhp
## 4   BD         A  31   8  2004         7         1  ACRU 29.0
## 12  BD         A  31   8  2004        15         1  ACRU 42.9
## 13  BD         A  31   8  2004        16         1  ACRU 18.0
## 23  BD         B  26   8  2004         1         1  ACRU 32.5
## 25  BD         B  26   8  2004         4         1  ACRU 29.8
## 34  BD         B  26   8  2004        14         1  ACRU 33.0
```

Here is the same operation with the function *filter*.

```
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
acru <- filter(kejim, espece == "ACRU")
head(acru)

##   site parcelle jour mois annee num_arbre nb_tiges espece  dhp
## 1   BD         A   31    8  2004        7        1   ACRU 29.0
## 2   BD         A   31    8  2004       15        1   ACRU 42.9
## 3   BD         A   31    8  2004       16        1   ACRU 18.0
## 4   BD         B   26    8  2004        1        1   ACRU 32.5
## 5   BD         B   26    8  2004        4        1   ACRU 29.8
## 6   BD         B   26    8  2004       14        1   ACRU 33.0
```

All *dplyr* functions have the same structure:

- the name of the function indicates the operation to be performed;
- the first argument is the input data frame;
- the other arguments specify the operation (here, the conditions of the filter);
- the function outputs a transformed data frame.

The benefits of using *dplyr* will become more clear after seeing many of the functions.

As in *ggplot2*, after specifying the data frame as the first argument, the following arguments can simply use the variable names, without quotation marks.

Multiple filters can be applied at a time by separating the conditions with commas. For example, `acru30` contains red maple trees with a DBH (*dhp*) greater than 30 cm.

```
acru30 <- filter(kejim, espece == "ACRU", dhp > 30)
head(acru30)

##   site parcelle jour mois annee num_arbre nb_tiges espece  dhp
## 1   BD         B   26    8  2004        1        1   ACRU 32.5
## 2   BD         B   26    8  2004       14        1   ACRU 33.0
## 3   BD         B   26    8  2004       25        1   ACRU 33.8
## 4   CL         A   18    8  2004        7        1   ACRU 39.8
## 5   CL         A   18    8  2004       14        1   ACRU 46.2
```

Exercise 4

Produce a table of observations for the year 2014, excluding the individuals of the species *TSCA* (eastern hemlock).

Solution

Sort observations with *arrange*

The `arrange` function sorts the rows of a data frame according to the value of one or more variables.

```
acru_tri_dhp <- arrange(acru, dhp)
head(acru_tri_dhp)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece dhp
## 1 CLT         A   13   8  2004         37         1  ACRU 9.20
## 2 CLT         A   13   8  2004         26         1  ACRU 9.25
## 3 CLT         A   13   8  2004         31         1  ACRU 9.40
## 4 CLT         A   20  10  2014         37         1  ACRU 9.50
## 5 CLT         A   14  10  2009         46         1  ACRU 9.70
## 6 CLT         A   14  10  2009         37         1  ACRU 9.80
```

To sort in descending order, use the `desc()` function. The following code sorts the data in ascending order of year, then in descending order of DBH.

```
acru_tri_an_dhp <- arrange(acru, annee, desc(dhp))
head(acru_tri_an_dhp)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece dhp
## 1 CL         B   18   8  2004         7         1  ACRU 48.5
## 2 CL         B   18   8  2004        14         1  ACRU 47.5
## 3 CL         A   18   8  2004        14         1  ACRU 46.2
## 4 BD         A   31   8  2004        15         1  ACRU 42.9
## 5 CL         B   18   8  2004        13         1  ACRU 42.1
## 6 CL         A   18   8  2004         7         1  ACRU 39.8
```

Pick variables with *select*

The `select` function selects columns from a data frame based on a comma-separated list of variable names.

```
acru_select <- select(acru_tri_dhp, site, annee, dhp)
head(acru_select)
```

```
##   site annee dhp
## 1 CLT  2004 9.20
## 2 CLT  2004 9.25
## 3 CLT  2004 9.40
## 4 CLT  2014 9.50
## 5 CLT  2009 9.70
## 6 CLT  2009 9.80
```

Chain operations with `%>%`

We often want to apply a series of operations to a data frame, without keeping the intermediate results. *dplyr* provides a useful shortcut for this purpose with the `%>%` (called *pipe*) operator.

The `%>%` operator has the effect of passing the result of a function as the first argument of the following function. For example, three of the operations we did earlier (select ACRU trees, sort by DBH and extract site, year, and DBH) can be combined in the following sequence.

```
acru <- kejin %>%
  filter(espece == "ACRU") %>%
  arrange(dhp) %>%
  select(site, annee, dhp)

head(acru)
```

```
##   site annee dhp
## 1 CLT  2004 9.20
## 2 CLT  2004 9.25
```

```
## 3 CLT 2004 9.40
## 4 CLT 2014 9.50
## 5 CLT 2009 9.70
## 6 CLT 2009 9.80
```

The first `%>%` passes the initial data frame `kejim` to the `filter` function, then the chain continues up to the last output, which is assigned to `acru`.

Exercise 5

Using `%>%`, produce a table of balsam fir (ABBA) observations with DBH > 15 cm, sorted in chronological order.

Solution

Derive new variables with *mutate*

The `mutate` function creates variables derived from existing columns in the data frame. In this example, we calculate the DBH in millimeters.

```
kejim_dhp_mm <- mutate(kejim, dhp_mm = dhp * 10)
head(kejim_dhp_mm)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece  dhp dhp_mm
## 1  BD         A   31    8  2004         1         1  TSCA  16.3   163
## 2  BD         A   31    8  2004         2         1  TSCA  24.0   240
## 3  BD         A   31    8  2004         6         1  TSCA  29.8   298
## 4  BD         A   31    8  2004         7         1  ACRU  29.0   290
## 5  BD         A   31    8  2004         8         1  TSCA  15.5   155
## 6  BD         A   31    8  2004         9         1  TSCA  32.0   320
```

Compte summary statistics per group: *group_by* and *summarize*

These two functions are often used together: `group_by` divides a data frame into groups according to the value of one or more variables and `summarize` calculates one or more summary variables for each group. The following example calculates the mean and standard deviation of the DBH by species.

```
group_by(kejim, espece) %>%
  summarize(moyDHP = mean(dhp), etDHP = sd(dhp))
```

```
## # A tibble: 13 x 3
##   espece moyDHP etDHP
##   <chr>   <dbl> <dbl>
## 1 ABBA    12.2  3.26
## 2 ACPE    12.0  1.71
## 3 ACRU    20.3 10.3
## 4 BEAL    50.6 23.3
## 5 BECO    26.2  7.39
## 6 BEPA    21.2  4.98
## 7 FAGR    17.7  7.51
## 8 PIRU    17.3  6.76
## 9 PIST    22.7 15.3
##10 POGR    27.1  6.59
##11 POTR    60.0  2.70
```

```
## 12 QURU      24.3  4.80
## 13 TSCA      22.9 15.0
```

Notes:

- Among the columns in the input data frame, only those defining the groups are retained by `summarize`.
- The `summarize` function is similar to `mutate`: both create new columns. However, `mutate` outputs a new value for each row of the input data frame, while `summarize` outputs a single value per group.

In addition to `mean`, `summarize` accepts any function that calculates a value from a group of values, e.g. `sum`, `min`, `max`, `sd`, and many others. A special function, `n()`, returns the number of observations in each group. The following example shows how to count the number of trees by site and year.

```
compte_site_annee <- kejim %>%
  group_by(site, annee) %>%
  summarize(compte = n())
head(compte_site_annee)
```

```
## # A tibble: 6 x 3
## # Groups:   site [2]
##   site annee compte
##   <chr> <int> <int>
## 1 BD    2004     45
## 2 BD    2009     42
## 3 BD    2014     44
## 4 CFR    2004     48
## 5 CFR    2009     47
## 6 CFR    2014     45
```

Exercise 6

Sort the species in `kejim` in descending order of maximum diameter.

Solution

Join two data frames

The file `codes_especes.csv` contains a data frame matching species codes from `kejim` to the species common names in French and English.

```
codes_esp <- read.csv("codes_especes.csv", stringsAsFactors = FALSE,
                     encoding = "UTF-8")
head(codes_esp)
```

```
##   espece      nom_espece species_name
## 1  ABBA      Sapin baumier    Balsam Fir
## 2  ACPE Érable de Pennsylvanie Striped Maple
## 3  ACRU      Érable rouge      Red Maple
## 4  ACSA      Érable à sucre    Sugar Maple
## 5  ACSP      Érable à épis Mountain Maple
## 6  AMLA      Amélanchier      Indian Pear
```

Note: The `encoding = "UTF-8"` argument is required to properly read diacritic marks in French.

To add these species names to the `kejim` data frame, we must join the two data frames with `inner_join`.

```
kejim_esp <- inner_join(kejim, codes_esp)
```

```
## Joining, by = "espece"
```

```
head(kejim_esp)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece dhp
## 1  BD         A   31    8  2004         1         1  TSCA 16.3
## 2  BD         A   31    8  2004         2         1  TSCA 24.0
## 3  BD         A   31    8  2004         6         1  TSCA 29.8
## 4  BD         A   31    8  2004         7         1  ACRU 29.0
## 5  BD         A   31    8  2004         8         1  TSCA 15.5
## 6  BD         A   31    8  2004         9         1  TSCA 32.0
##           nom_espece species_name
## 1 Pruche du Canada Eastern Hemlock
## 2 Pruche du Canada Eastern Hemlock
## 3 Pruche du Canada Eastern Hemlock
## 4   Érable rouge      Red Maple
## 5 Pruche du Canada Eastern Hemlock
## 6 Pruche du Canada Eastern Hemlock
```

The concept of a join originates in relational databases. As we can see here, the `inner_join` function has attached to each row of the first data frame (`kejim`) the data from a row in the second data frame (`codes_esp`) that has a matching value in `espece`. By default, *dplyr* assumes that the match must be made on columns of the same name, but it is possible to specify otherwise.

Note that the `kejim_esp` data frame has 1070 rows, 91 fewer than `kejim`. This is because species codes in `kejim` are missing from `codes_esp`. To keep the rows from the first data frame with no match in the second (and add missing values to the species name columns), you must use a different join type, `left_join`. We do not have time to discuss the different join types, but you can read the *dplyr* cheatsheet in the footnotes for more information.

Summary of *dplyr* functions

Fonction	Description
<code>filter</code>	select rows matching certain conditions
<code>arrange</code>	sort rows based on the values of specific variables
<code>select</code>	select columns by name
<code>mutate</code>	create new variables derived from existing columns
<code>group_by</code>	divide observations into groups based on grouping variables
<code>summarize</code>	calculate summaries of multiple observations (often by group)
<code>inner_join</code>	join two data frames based on common variables

References

- *R for Data Science* (<http://r4ds.had.co.nz>), specifically Chapters 3 (*Data Visualisation*) and 5 (*Data transformation*).
- Cheat sheet for *dplyr* and *tidyr*: <https://www.rstudio.com/wp-content/uploads/2015/02/data-wrangling-cheatsheet.pdf>
- Cheat sheet for *ggplot2*: <https://www.rstudio.com/wp-content/uploads/2016/11/ggplot2-cheatsheet-2.1.pdf>.

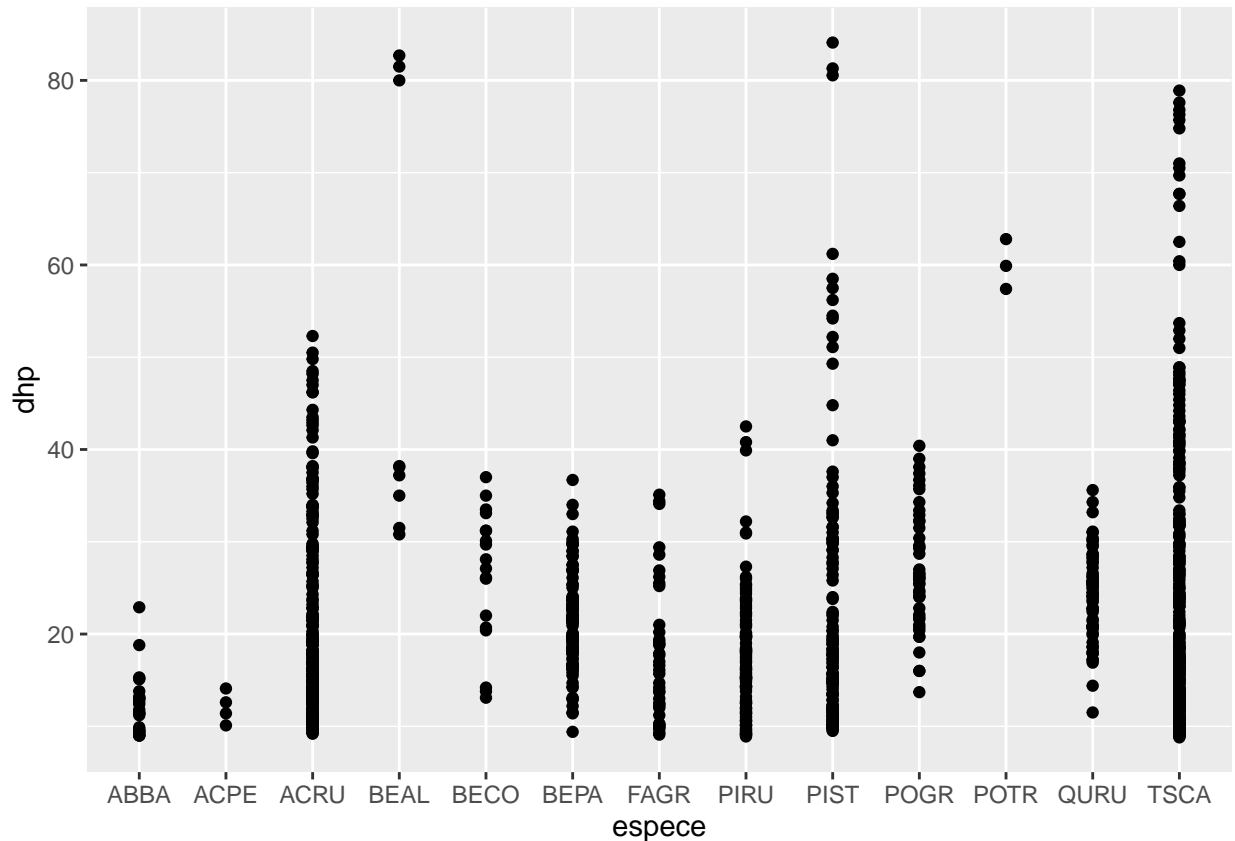
- Reference website for all *ggplot2* functions: <https://ggplot2.tidyverse.org/reference/index.html>.

Solutions to the exercises

Exercise 1

From the `kejim` data frame, produce a scatter plot of the DBH (`dhp`, on the y axis) according to the species (`espece`, on the x axis).

```
ggplot(data = kejim, mapping = aes(x = espece, y = dhp)) +  
  geom_point()
```

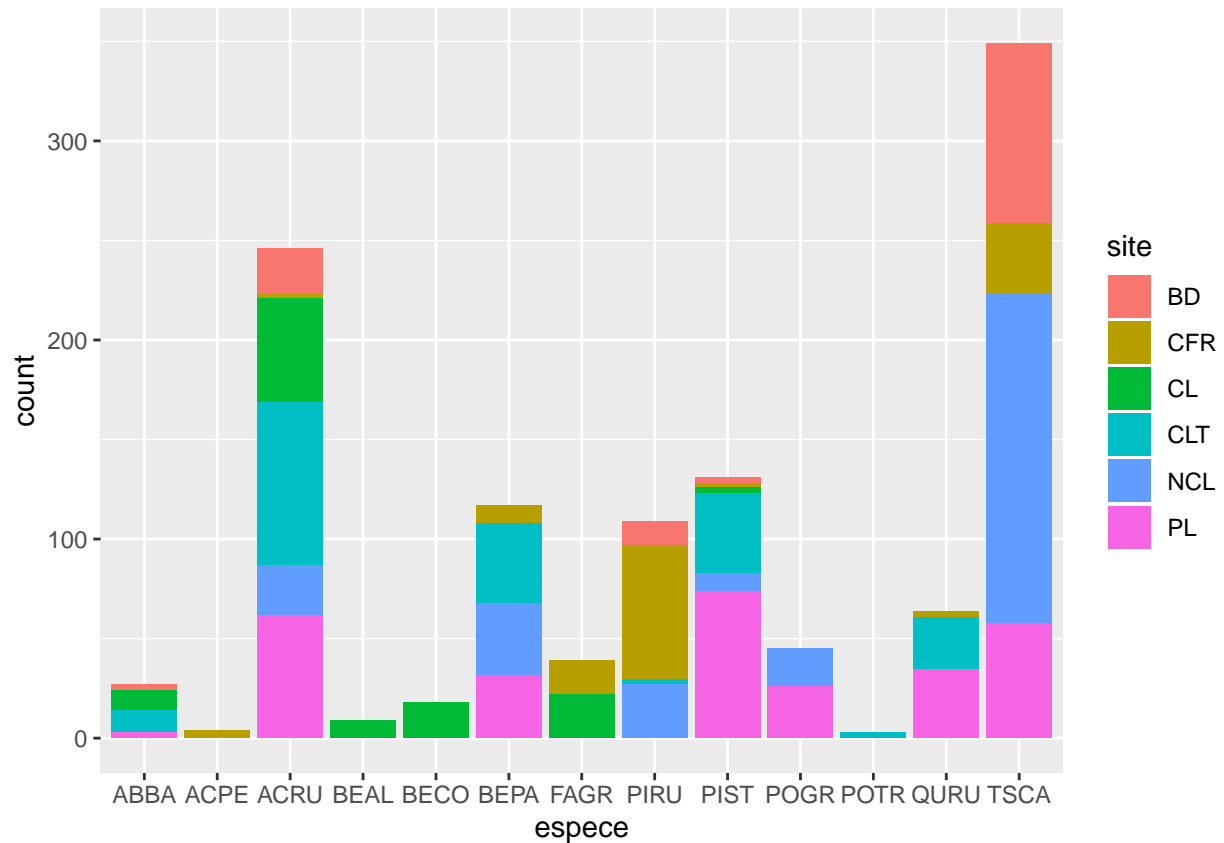


Return

Exercise 2

With `geom_bar`, create a bar graph of the number of individuals per species (different species on the x-axis). Use a color code to identify individuals from different sites.

```
ggplot(kejim, aes(x = espece, fill = site)) +  
  geom_bar()
```



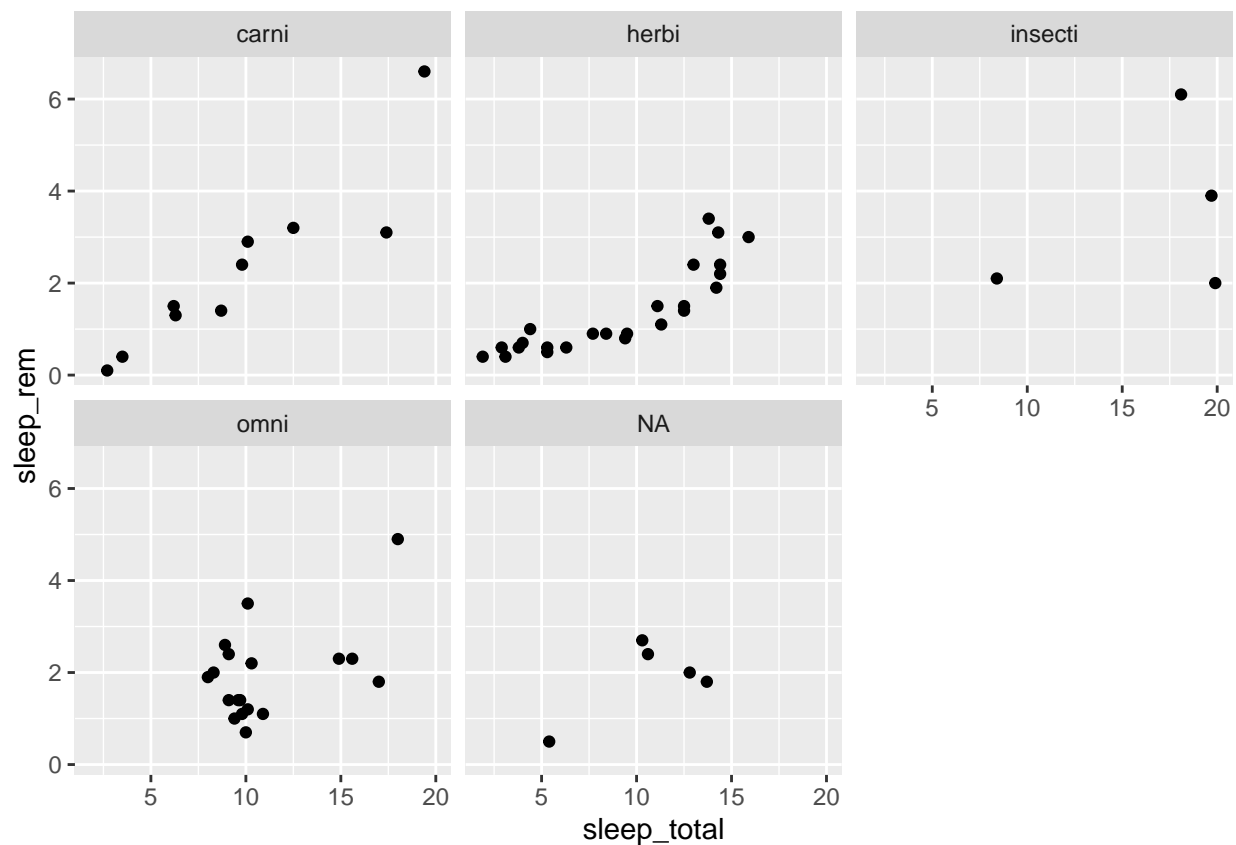
Return

Exercise 3

From the `msleep` dataset, create a scatter plot of total sleep (`sleep_total`) and REM sleep (`sleep_rem`) with one facet for each type of diet (`vore`)

```
ggplot(msleep, aes(x = sleep_total, y = sleep_rem)) +
  geom_point() +
  facet_wrap(~ vore)
```

Warning: Removed 22 rows containing missing values (geom_point).



Return

Exercise 4

Produce a table of observations for the year 2014, excluding the individuals of the species *TSCA* (eastern hemlock).

```
kejim2014 <- filter(kejim, annee == 2014, espece != "TSCA")
head(kejim2014)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece dhp
## 1  BD         A   11    8  2014         4         1  ACRU 38.2
## 2  BD         A   11    8  2014         7         1  ACRU 32.9
## 3  BD         A   11    8  2014        15         1  ACRU 44.3
## 4  BD         A   11    8  2014        16         1  ACRU 18.2
## 5  BD         B   11    8  2014         1         1  ACRU 33.9
## 6  BD         B   11    8  2014         3         1  PIRU  9.8
```

Return

Exercise 5

Using `%>%`, produce a table of balsam fir (ABBA) observations with DBH > 15 cm, sorted in chronological order.

```
kejim %>%
  filter(espece == "ABBA", dhp > 15) %>%
  arrange(annee, mois, jour)
```

```
##   site parcelle jour mois annee num_arbre nb_tiges espece  dhp
## 1 CLT         A   13    8  2004        35         1  ABBA 18.8
## 2 CL         A   18    8  2004         8         1  ABBA 15.1
## 3 CLT         A   14   10  2009        35         1  ABBA 22.9
## 4 CL         A   21   10  2009         8         1  ABBA 15.3
## 5 CLT         B   20   10  2014        10         1  ABBA 15.2
```

Return

Exercise 6

Sort the species in kejim in descending order of maximum diameter.

```
group_by(kejim, espece) %>%
  summarize(maxDHP = max(dhp)) %>%
  arrange(desc(maxDHP))
```

```
## # A tibble: 13 x 2
##   espece maxDHP
##   <chr>   <dbl>
## 1 PIST     84.1
## 2 BEAL     82.7
## 3 TSCA     78.9
## 4 POTR     62.8
## 5 ACRU     52.3
## 6 PIRU     42.5
## 7 POGR     40.4
## 8 BECO      37
## 9 BEPA     36.7
## 10 QURU     35.6
## 11 FAGR     35.1
## 12 ABBA     22.9
## 13 ACPE     14.1
```

Return