Lesson notes | Data dive: Ebola in Sierra Leone

Created by the GRAPH Courses team

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Learning objectives

- 1. You can use RStudio's graphic user interface to import CSV data into R.
- 2. You can explain the concept of reproducibility.
- 3. You can use the nrow(), ncol() and dim() functions to get the dimensions of a dataset, and the summary() function to get a summary of the dataset's variables.
- 4. You can use vis_dat(), inspect_num() and inspect_cat() to obtain visual summaries of a dataset.
- 5. You can inspect a numeric variable:
 - with the summary functions mean(), median(), max(), min(), length() and sum();
 - with esquisse-generated ggplot2 code.
- 6. You can inspect a categorical variable:
 - with the summary functions table() and janitor::tabyl();
 - with the graphical functions barplot() and pie().

Introduction

With your newly-acquired knowledge of functions and objects, you now have the basic building blocks required to do simple data analysis in R. So let's get started. The goal is to start working with data as quickly as possible, even before you feel ready.

Here you will analyze a dataset of confirmed and suspected cases of Ebola hemorrhagic fever in Sierra Leone in May and June of 2014 (Fang et al., 2016). The data is shown below:

You will import and explore this dataset, then use R to answer the following questions about the outbreak:

- When was the first case reported?
- What was the median age of those affected?
- Had there been more cases in men or women?
- What district had had the most reported cases?
- By the end of June 2014, was the outbreak growing or receding?

Script setup

First, open a new script in RStudio with File > New File > R Script. (If you are on RStudio, you can open up any of your previously-created projects.)



Next, save the script with File > Save As or press Command/Control + S to bring up the Save File dialog box. Save the file with the name "ebola_analysis" or something similar



Header

Add a title, name and date to the start of the script, as code comments. This is generally good practice for writing R scripts, as it helps give you and your collaborators context about your script. Your header may look like this:

```
# Ebola Sierra Leone analysis
# John Sample-Name Doe
# 2024-01-01
```

Packages

Next, use the $p_load()$ function from {pacman} to load the packages you will be using. Put this under a section header called "Load packages", with four hyphens, as shown below:

```
# Load packages ----
if(!require(pacman)) install.packages("pacman")
pacman::p_load(
   tidyverse, # meta-package
   inspectdf,
   plotly,
   janitor,
   visdat,
   esquisse
)
```

Remember that the *full signifier* of a function includes both the package name and the function name, package::function(). This full signifier is handy if you want to use a function before you have loaded its source package. This is the case in the code chunk above: we want use $p_load()$ from {pacman} without formally loading the {pacman} package, so we type $pacman::p_load()$

REMINDER



We could also first load {pacman} before using the p_load function:

(Also recall that the benefit of $p_load()$ is that it automatically installs a package if it is not yet installed. Without $p_load()$, you have to first install the package with install.packages() before you can load it with library().)

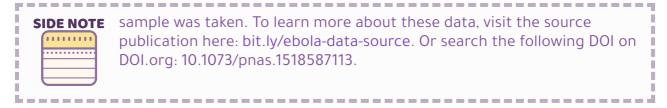
Importing data into R

Now that the needed packages are loaded, you should import the dataset.

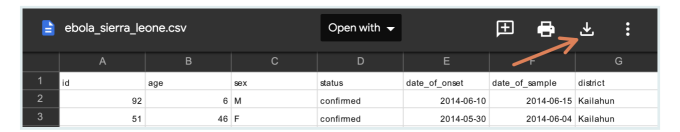
About the Ebola dataset



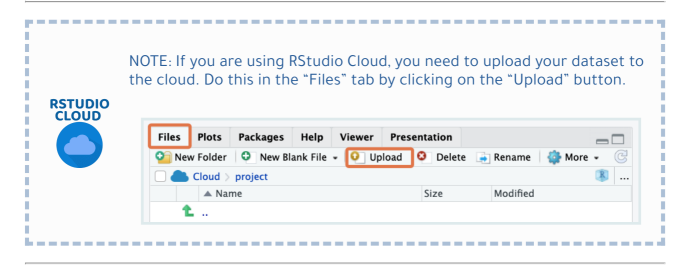
The data you will be working on contains a sample of patient information from the 2014-2016 Ebola outbreak in Sierra Leone. It comes from a research paper which analyzed the transmission dynamics of that outbreak. Key variables include the status of a case, whether the case



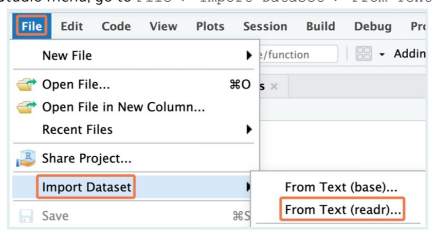
Go to bit.ly/view-ebola-data to view the dataset you will be working on. Then click the download icon at the top to download it to your computer.



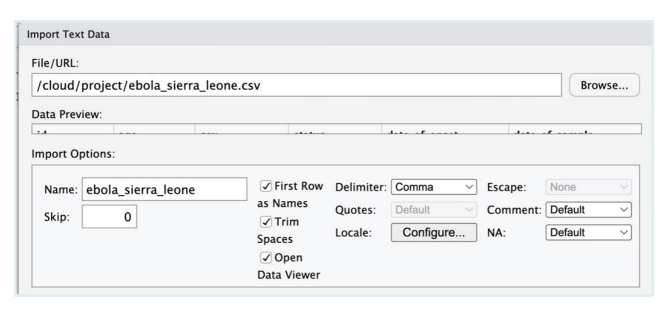
You can leave the dataset in your downloads folder, or move it to somewhere more respectable; the upcoming steps will work independent of where the data is stored. In the next lesson, you will learn how to organize your data analysis projects properly, and we will think about the ideal folder setup for storing data.



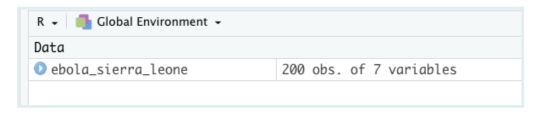
Next, on the RStudio menu, go to File > Import Dataset > From Text (readr).



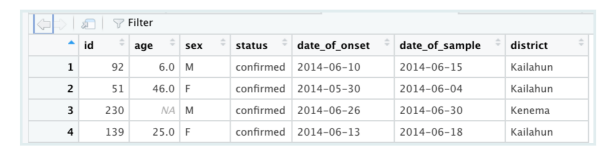
Browse through the computer's files and navigate to the downloaded dataset. Click to open it. You should see an import dialog box like this:



Leave all the import settings at the default values; simply click on "Import" at the bottom; this should load the dataset into R. You can tell this by looking at your environment pane, which should now feature an object called "ebola_sierra_leone" or something similar:



RStudio should also have called the View() function on your dataset, so you should see a familiar spreadsheet view of this data:



Now take a look at your console. Do you observe that your actions in the graphical user interface actually triggered some R code to be run? Copy the line of code that includes the $read_csv()$ function, leaving out the > symbol.

```
> Copy this
> library(readr)
> ebola_sierra_leone <- read_csv("ebola_sierra_leone.csv") (or something similar)
Rows: 200 Columns: 7
— Column specification
```

Paste the copied code into your R script, and label this section "Load data". This may look something like the below (the file path inside quotes will differ from computer to computer.

```
# Load data ----
ebola_sierra_leone <- read_csv("~/Downloads/ebola_sierra_leone.csv")</pre>
```

Nice work so far!

Your R script should look similar to this:



Intro to reproducibility

Now that the code for importing data is in your R script, you can easily rerun this script anytime to reimport the dataset; there will be no need to redo the manual point-and-click procedure for data import.

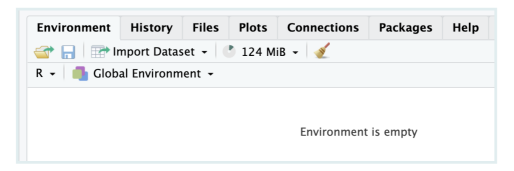
Try restarting R and rerunning the script now. Save your script with Control/Command + s, then restart R with the RStudio Menu, at Session > Restart R. On RStudio Cloud, the menu option looks like this:



If restarting is successful, your console should print this message:



You should also see the phrase "Environment is empty" in the Environment tab, indicating that the dataset you imported is no longer stored by R—you are starting with a fresh workspace.



To re-run your script, use Command/Control + a to highlight all the code, then Command/Control + Enter to run it.

If this worked, congratulations; you have the beginnings of your first "reproducible" analysis script!





When you do things with code rather than by pointing and clicking, it is easy for anyone to re-run, or *reproduce* these steps, by simply re-running your script.

While you can use RStudio's graphical user interface to point-and-click your way through the data import process, you should always copy the relevant code to your script so that your script remains a reproducible record of all your analysis steps.

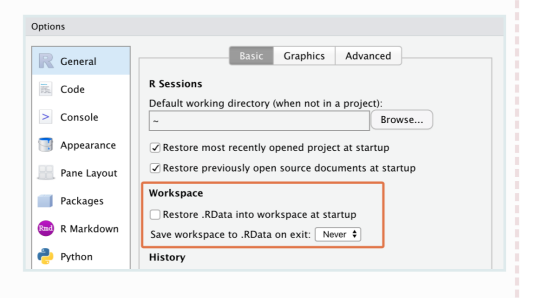


Of course, your script so far is not yet *entirely* reproducible, because the file path for the dataset (the one that looks like this: "...intro-to-data-analysis-with-r/ch01_getting_started/data...") is specific to just your computer. Later on we will see how to use relative file paths, so that the code for importing data can work on anyone's computer.

If your environment was not empty after restarting R, it means you skipped a step in a previous lesson. Do this now:

- In the RStudio Menu, go to Tools > Global Options to bring up RStudio's options dialog box.
- Then go to General > Basic, and uncheck the box that says "Restore .RData into workspace at startup".
- For the option, "save your workspace to .RData on exit", set this to "Never".





Quick data exploration

Now let's walk through some basic steps of data exploration—taking a broad, bird's eye look at the dataset. You should put this section under a heading like "Explore data" in your script.

To view the top and bottom 6 rows of the dataset, you can use the head() and tail() functions:

```
# Explore data ----
head(ebola_sierra_leone)
```

```
## # A tibble: 6 × 7
##
       id
                                date of onset date of sample
            age sex
                      status
##
    <dbl> <dbl> <chr> <chr>
                                <date>
                                              <date>
## 1
                      confirmed 2014-06-10
                                              2014-06-15
       92
             6 M
## 2
             46 F
                      confirmed 2014-05-30
       51
                                              2014-06-04
## 3
      230
                      confirmed 2014-06-26
                                              2014-06-30
             NA M
## 4
      139
             25 F
                      confirmed 2014-06-13
                                              2014-06-18
## 5
       8
              8 F
                      confirmed 2014-05-22
                                              2014-05-27
    215
             49 M
## 6
                      confirmed 2014-06-24
                                              2014-06-29
## # ... with 1 more variable: district <chr>
```

```
tail(ebola sierra leone)
```

```
## # A tibble: 6 × 7
                                date of onset date of sample
##
       id age sex
                      status
    <dbl> <dbl> <chr> <chr>
##
                                <date>
                                              <date>
## 1
      214
            6 F
                      confirmed 2014-06-24
                                              2014-06-30
             45 F
                      confirmed 2014-05-27
## 2
       28
                                              2014-06-01
## 3
       12
             27 F
                      confirmed 2014-05-22
                                              2014-05-27
## 4
      110
              6 M
                      confirmed 2014-06-10
                                              2014-06-15
## 5
      209
             40 F
                      confirmed 2014-06-24
                                              2014-06-27
## 6
      35
             29 M
                      suspected 2014-05-28
                                              2014-06-01
## # ... with 1 more variable: district <chr>
```

To view the whole dataset, use the View() function.

```
View(ebola_sierra_leone)
```

This will again open a familiar spreadsheet view of the data:



You can close this tab and return to your script.

The functions nrow(), ncol() and dim() give you the dimensions of your dataset:

```
nrow(ebola_sierra_leone) # number of rows
```

[1] 200

```
ncol(ebola_sierra_leone) # number of columns
```

[1] 7

```
dim(ebola_sierra_leone) # number of rows and columns
```

[1] 200 7

REMINDER



If you're not sure what a function does, remember that you can get function help with the question mark symbol. For example, to get help on the ncol() function, run:

?ncol

Another often-helpful function is summary():

```
summary(ebola_sierra_leone)
```

```
##
        id
                        age
                                      sex
                                                       status
date of onset
## Min. : 1.00
                   Min. : 1.80
                                  Length:200
                                                   Length:200
Min. :2014-05-18
## 1st Qu.: 62.75
                   1st Qu.:20.00
                                  Class :character
                                                   Class : character
1st Qu.:2014-06-01
## Median :131.50
                   Median :35.00
                                  Mode :character Mode :character
Median :2014-06-13
## Mean :136.72
                   Mean :33.85
Mean :2014-06-12
## 3rd Qu.:208.25
                   3rd Qu.:45.00
3rd Qu.:2014-06-23
## Max. :285.00
                   Max. :80.00
Max. :2014-06-29
                   NA's :4
## date of sample
                        district
## Min. :2014-05-23 Length:200
## 1st Qu.:2014-06-07 Class :character
## Median :2014-06-18 Mode :character
```

```
## Mean :2014-06-17
## 3rd Qu.:2014-06-29
## Max. :2014-07-17
##
```

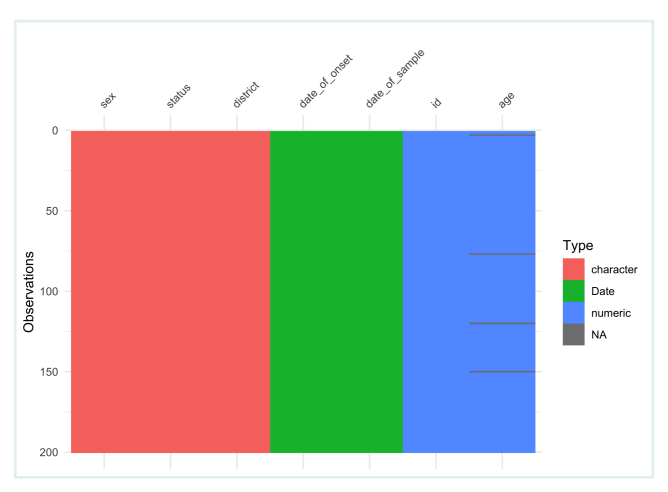
As you can see, for numeric columns in your dataset, summary () gives you the minimum value, the maximum value, the mean, median and the 1st and 3rd quartiles.

For character columns it gives you just the length of the column (the number of rows), the "class" and the "mode". We will discuss what "class" and "mode" mean later.

```
vis dat()
```

The <code>vis_dat()</code> function from the {visdat} package is a wonderful way to quickly visualize the data types and the missing values in a dataset. Try this now:

```
vis_dat(ebola_sierra_leone)
```



From this figure, you can quickly see the character, date and numeric data types, and you can note that age is missing for some cases.

```
inspect cat() and inspect num()
```

Next, inspect_cat() and inspect_num() from the {inspectdf} package give you visual summaries of the distribution of variables in the dataset.

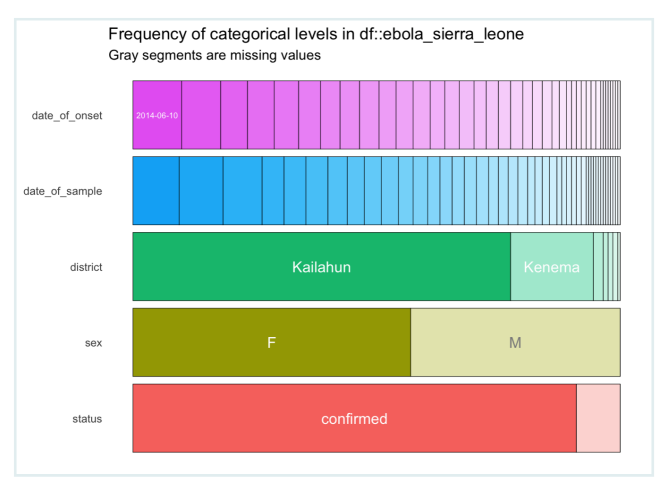
If you run <code>inspect_cat()</code> on the data object, you get a tabular summary of the categorical variables in the dataset, with some information hidden in the <code>levels</code> column (later you will learn how to extract this information).

```
inspect_cat(ebola_sierra_leone)
```

But the magic happens when you run show plot() on the result from inspect cat():

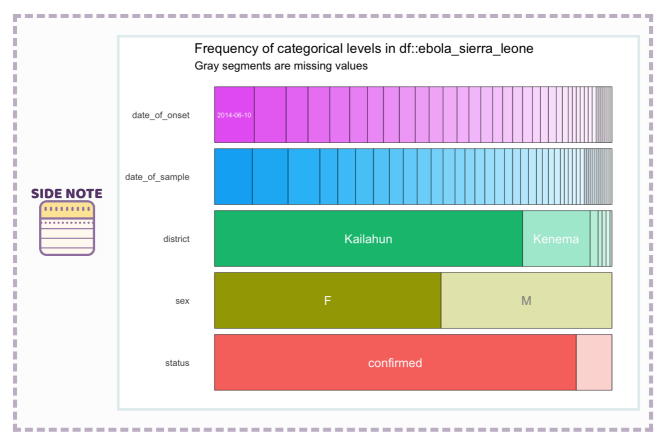
```
# store the output of `inspect_cat()` in `cat_summary`
cat_summary <- inspect_cat(ebola_sierra_leone)

# call the `show_plot()` function on that summmary.
show_plot(cat_summary)</pre>
```



You get a wonderful figure showing the distribution of all categorical and date variables!





From this plot, you can quickly tell that most cases are in Kailahun, and that there are more cases in women than in men ("F" stands for "female").

One problem is that in this plot, the smaller categories are not labelled. So, for example, we are not sure what value is represented by the white section for "status" at the bottom right. To see labels on these smaller categories, you can turn this into an interactive plot with the ggplotly() function from the {plotly} package.

```
cat_summary_plot <- show_plot(cat_summary)
ggplotly(cat_summary_plot)</pre>
```

Wonderful! Now you can hover over each of the bars to see the proportion of each bar section. For example you can now tell that 9% (0.090) of the cases have a suspected status:



REMINDER



The assignment arrow, <-, can be written with the RStudio shortcut alt +- (alt AND minus) on Windows or option +- (option AND minus) on macOS.

You can obtain a similar plot for the numerical (continuous) variables in the dataset with inspect num(). Here, we show all three steps in one go.

```
num_summary <- inspect_num(ebola_sierra_leone)
num_summary_plot <- show_plot(num_summary)
ggplotly(num_summary_plot)</pre>
```

This gives you an overview of the numerical columns, age and id. (Of course, the distribution of the id variable is not meaningful.)

You can tell that individuals aged 35 to 40 (mid-point 37.5) are the largest age group, making up 13.8% (0.1377...) of the cases in the dataset.

Analyzing a single numeric variable

Now that you have a sense of what the entire dataset looks like, you can isolate and analyze single variables at a time—this is called *univariate analysis*.

Go ahead and create a new section in your script for this univariate analysis.

```
# Univariate analysis, numeric variables ----
```

Let's start by analyzing the numeric age variable.

Extract a column vector with \$

To extract a single variable/column from a dataset, use the dollar sign, \$ operator:

```
ebola_sierra_leone$age # extract the age column in the dataset
```

```
## [1] 6.0 46.0 NA 25.0 8.0 49.0 13.0 50.0 35.0 38.0 60.0 18.0 10.0 14.0 50.0 35.0 43.0 17.0 3.0 ## [20] 60.0 38.0 41.0 49.0 12.0 74.0 21.0 27.0 41.0 42.0 60.0 30.0 50.0 50.0 22.0 40.0 35.0 19.0 3.0 ## [39] 34.0 21.0 73.0 65.0 30.0 70.0 12.0 15.0 42.0 60.0 14.0 40.0 33.0 43.0 45.0 14.0 14.0 40.0 35.0 ## [58] 30.0 17.0 39.0 20.0 8.0 40.0 42.0 53.0 18.0 40.0 20.0 45.0 40.0 60.0 44.0 33.0 23.0 45.0 7.0
```

```
## [96] 26.0 37.0 30.0 3.0 56.0 32.0 35.0 54.0 42.0 48.0 11.0 1.8 63.0 55.0 20.0 62.0 62.0 42.0 65.0 ## [115] 29.0 20.0 33.0 30.0 35.0 NA 50.0 16.0 3.0 22.0 7.0 50.0 17.0 40.0 21.0 9.0 27.0 52.0 50.0 ## [134] 25.0 10.0 30.0 32.0 38.0 30.0 50.0 26.0 35.0 3.0 50.0 60.0 40.0 34.0 4.0 42.0 NA 54.0 18.0 ## [153] 45.0 30.0 35.0 35.0 16.0 26.0 23.0 45.0 45.0 45.0 38.0 45.0 35.0 30.0 60.0 5.0 18.0 2.0 70.0 ## [172] 35.0 3.0 30.0 80.0 62.0 20.0 45.0 18.0 28.0 48.0 38.0 39.0 26.0 60.0 35.0 20.0 50.0 11.0 36.0 ## [191] 29.0 57.0 35.0 26.0 6.0 45.0 27.0 6.0 40.0 29.0
```



This list of values is called a *vector* in R. A vector is a kind of data structure that has elements of one *type*. In this case, the type is "numeric". We will formally introduce you to vectors and other data structures in a future chapter. In this lesson, you can take "vector" and "variable" to be synonyms.

Basic operations on a numeric variable

To get the mean of these ages, you could run:

```
mean(ebola_sierra_leone$age)
```

```
## [1] NA
```

But it seems we have a problem. R says the mean is NA, which means "not applicable" or "not available". This is because there are some missing values in the vector of ages. (Did you notice this when you printed the vector?) By default, R cannot find the mean if there are missing values. To ignore these values, use the argument na.rm (which stands for "NA remove") setting it to T, or TRUE:

```
mean(ebola_sierra_leone$age, na.rm = T)
```

```
## [1] 33.84592
```

Great! This need to remove the NAs before computing a statistic applies to many functions. The median() function for example, will also return NA by default if it is called on a vector with any NAs:

```
median(ebola_sierra_leone$age) # does not work
```

```
median(ebola sierra leone$age, na.rm = T) # works
  ## [1] 35
mean and median are just two of many R functions that can be used to inspect a
numerical variable. Let's look at some others.
But first, we can assign the age vector to a new object, so you don't have to keep typing
ebola sierra leone$age each time.
age vec <- ebola sierra leone$age # assign the vector to the object "age vec"
Now run these functions on age vec and observe their outputs:
sd(age vec, na.rm = T) # standard deviation
  ## [1] 17.26864
max(age vec, na.rm = T) # maximum age
  ## [1] 80
min(age_vec, na.rm = T) # minimum age
 ## [1] 1.8
 summary(age vec) # min, max, mean, quartiles and NAs
                                                       NA's
  ##
       Min. 1st Qu. Median
                               Mean 3rd Qu.
                                               Max.
  ##
       1.80 20.00 35.00
                               33.85 45.00
                                               80.00
 length(age vec) # number of elements in the vector
  ## [1] 200
```

sum(age_vec, na.rm = T) # sum of all elements in the vector

20

[1] 6633.8

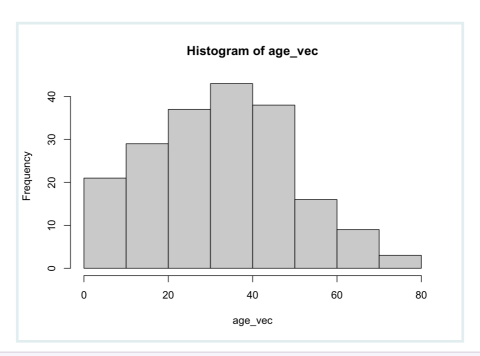
Do not feel intimidated by the long list of functions! You should not have to memorize them; rather you should feel free to Google the function for whatever operation you want to carry out. You might search something like "what is the function for standard deviation in R". One of the first results should lead you to what you need.

Visualizing a numeric variable

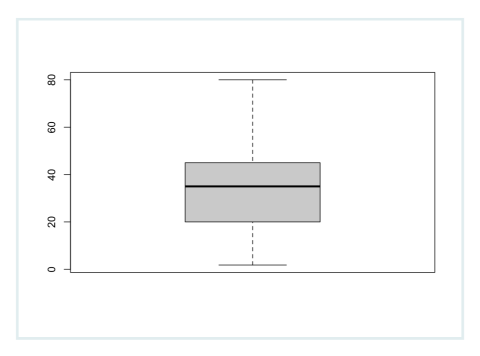
Now let's create a graph to visualize the age variable. The two most common graphics for inspecting the distribution of numerical variables are histograms (like the output of the <code>inspect_num()</code> function you saw earlier) and boxplots.

R has built-in functions for these:

hist(age vec)



boxplot(age vec)



Nice and easy!

Graphical functions like boxplot() and hist() are part of R's base graphics package. These functions are quick and easy to use, but they do not offer a lot of flexibility, and it is difficult to make beautiful plots with them. So most people in the R community use an extension package, {ggplot2}, for their data visualization.

In this course, we'll use ggplot indirectly; by using the {esquisse} package, which provides a user-friendly interface for creating ggplot2 plots.

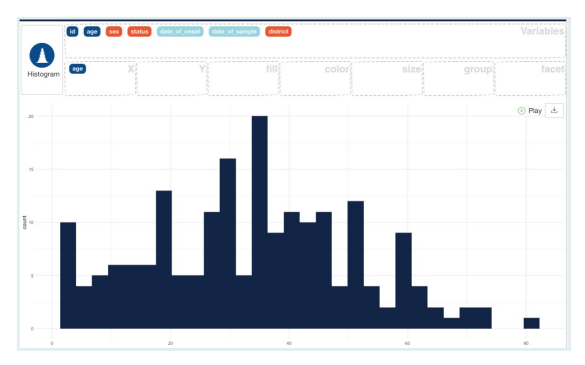
The workhorse function of the {esquisse} package is <code>esquisser()</code>, and this function takes a single argument—the dataset you want to visualize. So we can run:

```
esquisser(ebola_sierra_leone)
```

This should bring a graphic user interface that you can use to plot different variables. To visualize the age variable, simply drag age from the list of variables into the x axis box:



When age is in the x axis box, you should automatically get a histogram of ages:



You can change the plot type by clicking on the "Histogram" button and selecting one of the other valid plot types. Try out the boxplot, violin plot and density plot and observe the outputs.



When you are done creating a plot with {esquisse}, you should copy the code that was created by clicking on the "Code" button at the bottom right then "Copy to clipboard":



Now, paste that code into your script, and make sure you can run it from there. The code should look something like this:

```
ggplot(ebola_sierra_leone) +
  aes(x = age) +
  geom_histogram(bins = 30L, fill = "#112446") +
  theme_minimal()
```

By copying the generated code into your script, you ensure that the data visualization you created is fully reproducible.



{esquisse} can only create fairly simple graphics, so when you want to make highly customized or complex plots, you will need to learn how to write {ggplot} code manually. This will be the focus of a later course.

You should also test out the other tabs on the bottom toolbar to see what they do: Labels & Title, Plot options, Appearance and Data.

Easy bivariate and multivariate plots





In this lesson we are focusing on univariate analysis: exploring and visualizing one variable at a time. But with esquisse; it is *so* easy to make a bivariate or multivariate plot, so you can already get your feet wet with this.

Try the following plots:

CHALLENGE



- Drag age to the X box and sex to the Y box.
- Drag age to the X box, sex to the Y box, and sex to the fill box.
- Drag age to the X box and district to the Y box.

Analyzing a single categorical variable

Next, let's look at a categorical variable, the districts of reported cases:

Univariate analysis, categorical variables ---- ebola sierra leone\$district

	-	-		"Kailahun"	"Kenema"	"Kailahun"	
			"Kailahun				
	## [7] "Kai	lahun"	"Kailahun"	"Kenema"	"Kailahun"	
"Kailahun" "Kailahur							
	## [1	3] " Kai	lahun"	"Kailahun"	"Kailahun"	"Kailahun"	
	"Kaila	.hun"	"Kenema"				
	## [1	9] " Kon	0"	"Kailahun"	"Kailahun"	"Kailahun"	
	"Kenem	ıa"	"Kailahun	1"			
	## [2	5] " Kai	lahun"	"Kailahun"	"Kailahun"	"Kailahun"	
	"Kenem	ıa"	"Kenema"				
	## [3	1] "Ken	ema"	"Kailahun"	"Kailahun"	"Bo"	
	"Kaila	hun"	"Kailahun	1"			
	## [3	7] " Kai	lahun"	"Kenema"	"Kenema"	"Kenema"	
	"Kaila	.hun"	"Kailahun	1"			
	## [4	3] " Kai	lahun"	"Kailahun"	"Kailahun"	"Kailahun"	
	"Western Urban" "Kailahun"						
	## [4	9] "Kai	lahun"	"Kailahun"	"Kailahun"	"Kailahun"	
	"Kaila	.hun"	"Kailahun	1"			
			lahun"		"Kailahun"	"Kailahun"	
"Kailahun" "Kailahu			"Kailahun	1"			
## [61] "Kailahun" "			lahun"	"Kenema"	"Western Urban"	"Kambia"	
			"Kailahun				
	## [6	71 " Kai	lahun"	"Kailahun"	"Kailahun"	"Kailahun"	
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			ema"		"Kailahun"	"Kailahun"	
			"Kailahun				
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			"Kenema"				
			lahun"		"Kailahun"	"Kono"	
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				"Kailahun"	"Kailahun"	"Kailahun"	
	11 L D	, j nal		1.411411	1.411411	-varranun	

```
"Kenema"
            "Kailahun"
## [103] "Kailahun" "Kenema"
                                   "Kailahun"
                                                 "Kailahun"
"Kailahun" "Kailahun"
## [109] "Kailahun" "Kailahun"
                                   "Kenema"
                                                 "Western Urban"
"Kailahun" "Kailahun"
## [115] "Kailahun" "Kailahun"
                                  "Kailahun"
                                                 "Kailahun"
"Kailahun" "Kailahun"
## [121] "Kailahun" "Kailahun"
                                   "Kenema"
                                                 "Kailahun"
"Kailahun" "Kenema"
## [127] "Kailahun" "Port Loko"
                                   "Kailahun"
                                                 "Kailahun"
"Kailahun" "Kailahun"
## [133] "Kailahun" "Kailahun"
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                                                 "Kailahun"
"Kailahun" "Kailahun"
## [139] "Kailahun" "Kailahun"
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"Kailahun" "Kenema"
## [145] "Kenema" "Kailahun"
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                                                 "Kailahun"
"Kailahun" "Kailahun"
## [151] "Kailahun" "Kailahun"
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                                                 "Kailahun"
"Kailahun"
         "Kenema"
                    "Kenema"
## [157] "Kailahun"
                                   "Kailahun"
                                                 "Kailahun"
"Kenema"
        "Kailahun"
## [163] "Kailahun" "Kailahun"
                                   "Kailahun"
                                                 "Bo"
"Kailahun" "Kailahun"
## [169] "Kailahun" "Kailahun"
                                   "Kailahun"
                                                 "Kailahun"
"Kenema"
             "Kailahun"
## [175] "Kailahun" "Kenema"
                                   "Kailahun"
                                                 "Kailahun"
"Kailahun"
             "Kailahun"
## [181] "Kailahun" "Kailahun"
                                   "Kailahun"
                                                 "Western Urban"
"Kailahun" "Kailahun"
## [187] "Kenema" "Kailahun"
                                   "Kailahun"
                                                 "Kailahun"
"Kailahun" "Kailahun"
## [193] "Kailahun" "Kenema"
                                   "Kenema"
                                                 "Kailahun"
"Kailahun" "Kailahun"
## [199] "Kailahun" "Kenema"
```

Sorry for printing that very long vector!

Frequency tables

You can use the table() function to create a frequency table of a categorical variable:

table(ebola sierra leone\$district)

##					
##	Во	Kailahun	Kambia	Kenema	Kono
Port Loko	Western	Urban			
##	2	155	1	34	2
2	4				

You can see that most cases are in Kailahun and Kenema.

table() is auseful "base" function. But there is a better function for creating frequency tables, called tabyl(), from the {janitor} package.

To use it, you supply the name of your data frame as the first argument, then the name of variable to be tabulated:

```
tabyl(ebola_sierra_leone, district)
```

```
## district n percent
## Bo 2 0.010
## Kailahun 155 0.775
## Kambia 1 0.005
## Kenema 34 0.170
## Kono 2 0.010
## Port Loko 2 0.010
## Western Urban 4 0.020
```

As you can see, tabyl() gives you both the counts and the percentage proportions of each value. It also has some other attractive features you will see later.

You can also easily make cross-tabulations with tabyl(). Simply add additional variables separated by a comma. For example, to create a cross-tabulation by district and sex, run:

```
tabyl(ebola_sierra_leone, district, sex)
```



```
## district F M
## Bo 0 2
## Kailahun 91 64
## Kambia 0 1
## Kenema 20 14
## Kono 0 2
## Port Loko 1 1
## Western Urban 2 2
```

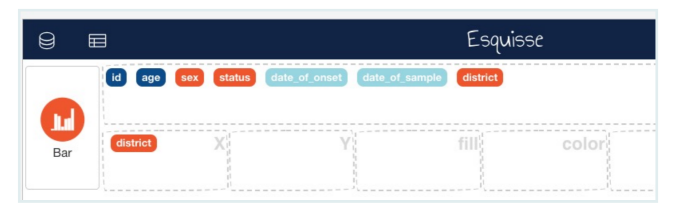
The output shows us that there were 0 women in the Bo district, 2 men in the Bo district, 91 women in the Kailahun district, and so on.

Visualizing a categorical variable

Now, let's try to visualize the district variable. As before, the best way to do this is with the esquisser() function from {esquisse}. Run this code again:

```
esquisser(ebola_sierra_leone)
```

Then drag the district variable to the X axis box:



You should get a bar chart showing the count of individuals across districts. Copy the generated code and paste it into your script.

Answering questions about the outbreak

With the functions you have just learned, you have the tools to answer the questions about the Ebola outbreak that were listed at the top. Give it a go. Attempt these questions on your own, then look at the solutions below.

- When was the first case reported? (Hint: look at the date of sample)
- As at the end of June 2014, which 10-year age group had had the most cases?
- What was the median age of those affected?
- Had there been more cases in men or women?
- What district had had the most reported cases?
- By the end of June 2014, was the outbreak growing or receding?

Solutions

When was the first case reported?

```
min(ebola_sierra_leone$date_of_sample)
```

```
## [1] "2014-05-23"
```

We don't have the date of report, but the first "date_of_sample" (when the Ebola test sample was taken from the patient) is May 23rd. We can use this as a proxy for the date of first report.

What was the median age of cases?

```
median(ebola_sierra_leone$age, na.rm = T)
## [1] 35
```

The median age of cases was 35.

Are there more cases in men or women?

```
tabyl(ebola_sierra_leone$sex)

## ebola_sierra_leone$sex n percent
## F 114 0.57
## M 86 0.43
```

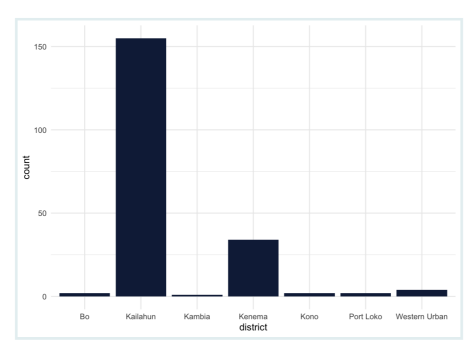
As seen in the table, there were more cases in women. Specifically, 57% of cases are of women.

What district has had the most reported cases?

```
tabyl(ebola_sierra_leone$district)
```

```
## ebola_sierra_leone$district n percent
##
                         Bo 2 0.010
##
                    Kailahun 155 0.775
                      Kambia 1 0.005
##
                      Kenema 34 0.170
##
##
                        Kono 2
                                0.010
##
                   Port Loko 2 0.010
##
               Western Urban 4 0.020
```

```
# We can also plot the following chart (generated with esquisse)
ggplot(ebola_sierra_leone) +
  aes(x = district) +
  geom_bar(fill = "#112446") +
  theme_minimal()
```

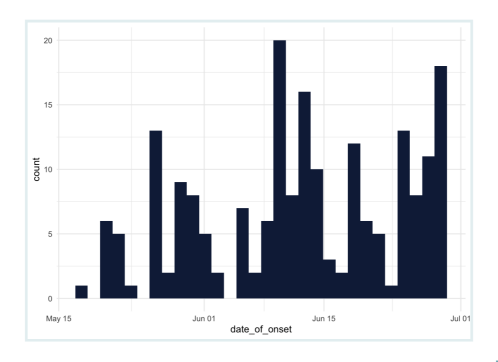


As seen, the Kailahun district had the majority of cases.

• By the end of June 2014, was the outbreak growing or receding?

For this, we can use esquisse to generate a bar chart that shows a count of cases in each day. Simply drag the <code>date_of_onset</code> variable to the x axis. The output code from esquisse should resemble the below:

```
ggplot(ebola_sierra_leone) +
  aes(x = date_of_onset) +
  geom_histogram(bins = 30L, fill = "#112446") +
  theme_minimal()
```



Great! But it is debatable whether the outbreak was growing or receding at the end of June 2014; a precise trend is not really clear!

Haven't had enough?

If you would like to practice some of the methods and functions you learned on a similar dataset, try downloading the data that is stored on this page: https://bit.ly/view-yaounde-covid-data

That dataset is in the form of an Excel spreadsheet, so when you are importing the dataset with RStudio, you should use the "From Excel" option (File > Import Dataset > From Excel).

This dataset contains the results of a COVID-19 serological survey conducted in Yaounde, Cameroon in late 2020. The survey estimated how many people had been infected with COVID-19 in the region, by testing for IgG and IgM antibodies. The full dataset can be obtained from here: go.nature.com/3R866wx

Wrapping up

Congratulations! You have now taken your first baby steps in analyzing data with R: you imported a dataset, explored its structure, performed basic univariate analysis and visualization on its numeric and categorical variables, and you were able to answer important questions about the outbreak based on this.

Of course, this was only a *sneak peek* of the data analysis process—a lot was left out. Hopefully, though, this sneak peek has gotten you a bit excited about what you can do with R. And hopefully, you can already start to apply some of these to your own datasets. The journey is only beginning! See you soon.

Contributors

The following team members contributed to this lesson:



References

Some material in this lesson was adapted from the following sources:

- Barnier, Julien. "Introduction à R Et Au Tidyverse." Partie 13 Diffuser et publier avec rmarkdown, May 24, 2022. https://juba.github.io/tidyverse/13-rmarkdown.html.
- Yihui Xie, J. J. Allaire, and Garrett Grolemund. "R Markdown: The Definitive Guide." Home, April 11, 2022. https://bookdown.org/yihui/rmarkdown/.

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