CCESR Intern Hub

CCESR Fellows

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Welcome!

This website / HTML book is intended to collect resources for Cedar Creek summer interns doing independent research projects, and present those resources in an easily accessible way.

For now, the focus is primarily on data analysis and using the R programming language.

Much of the content featured is adapted from the work of past CCESR Fellows, including Mariana Cardenas and Bea Baselga.

This site is structured in different parts, which can be read in any order you choose, depending on your needs / what you already know. Currently, the first part goes over data analysis in general, the second part describes R-related software and workflows, and the third part is intended to give a primer in R coding.

Part I Data Analysis

1 Data Analysis at a Glance

Analyzing your data is usually about transforming long spreadsheets into a form that is relevant to your question/s, and oftentimes including an appropriate statistical approach for inference.

You might use **descriptive statistics**, which is simply *describing* what you observed without presenting every data point, and instead a summary of those data. This can often be helpful in providing a frame of reference to your dataset before looking deeper at trends and comparisons. Alternatively, sometimes descriptive statistics are the main goal - like in surveys of populations and communities (e.g., what is the population size of a certain grass of interest in an old field?). Descriptive statistics include things like the mean and variance, but can also include more niche measures like dispersion.

You could also use **inferential statistics**, which is more about using math or simulation techniques to *infer* some conclusion form the shape of your data. This is directly relevant to when you have an ecological question about cause and effect, associations among variables, comparisons among categories, etc. The results of inferential statistics provide a starting point from which to interpret/discuss an answer to your question. Examples include t-tests and linear regression.

When using both of these types of statistics, you should be mindful of **data types**, which are the form that variables take. For example, the height of a tree is number, but the species of a tree is a category. This contrast is obvious, but there are subtle differences that can be important for how you describe, assess, and plot your data.

2 Data Types

First, let's go over different types of data:

2.1 Numeric Data

Any data that can be described with numbers or have quantifiable relationships between values is numeric. But! There are multiple types of numeric data. The most important distinction is **discrete** vs **continuous**.

Discrete numeric data is data where not every value is possible, but you can still quantify specific differences among the possible values - the major exmaple being integer values (1, 2, 3, the rest). Most programming languages will refer to this type as integer or int. Examples might include number of ants on a log.

Continuous numeric data is data where every valuable is possible! So this is basically all real numbers, including decimals (1.0, 1.1, etc.). Many programming languages will refer to this as simply numeric data, but lower level languages might use "float" or "double". Examples might include th biomass of ants on a log. Note: measures that consist of very large integer values are approximately continuous.

Other things to consider with numeric data is whether the scale of measurement is bound by any values. For example, the number of or biomass of ants on a log cannot be less than zero. In addition, percentages and proportions are bound by 0 and 100 and 0 and 1 respectively. These limitations can lead to special considerations when performing inferential statistics.

2.2 Categorical Data

Any data for which the values have no specifically quantitative difference among them is categorical. Again there is one majorly important distinction: **nominal** vs **ordinal**.

Nominal data is data where categories have no ranking or order, like the species of ants on a log.

Ordinal data is data where categories have some order, like your top 5 favorite breakfast cereals. But wait! You may be thinking - "isn't this quantitative?" Well yes and no. The difference between ordinal data and discrete numeric data is that you can't really quantify the

exact difference between ordinal data values. Say there is a go-kart race between Mario, Luigi, and Peach. The place that each finished would be ordinal, e.g., Peach got 1st and Luigi 2nd, but you wouldn't be able to say how much faster Peach was than Luigi. The time it took for Peach and Luigi each to finish the race would be a numeric variable, and there would be a specific value difference between them.

3 Descriptive Statistics

Now, let's discuss how to describe your data:

3.1 Centrality

You'll often want to describe the central tendency of your data - around where are the values centered?

Mean - the average of the values, or the sum of all values divided by the number of observations

Median - the value at which half of the observations are greater, and the other half are less

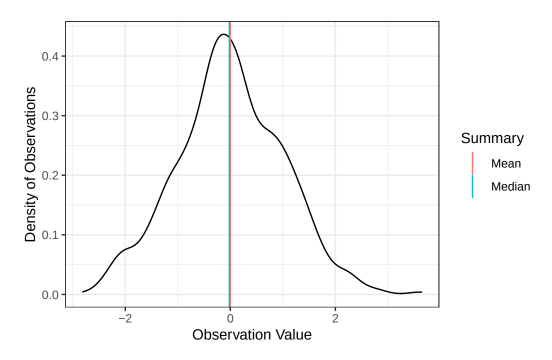
Mode - the most commonly observe value

Usually, the mean is a a perfectly adequate descriptor. You can use it on continuous numeric data, discrete numeric data (though the mean value will often be unrealistic), or even ordinal rankings.

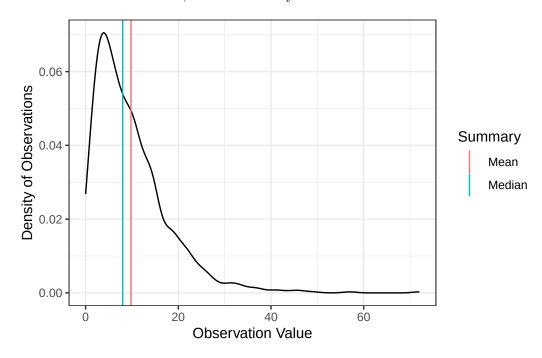
When might you prefer to use the median over the mean?

When the data is skewed such that there are many small values and a few big values, the mean might be inflated by those large values, and thus overestimate the central tendency in some contexts.

When data is roughly normally distributed, the mean and median are roughly the same:



But when data are skewed, the median may be a better estimate of the central tendency:



3.2 Spread

You also might be interested in how varied your data is, how much it deviates from the central tendency. This can be done with the following:

Variance - how variable is the data? Measured as the average squared difference between observations and the mean:

$$Variance = \frac{\sum (Observation_i - Mean)^2}{Number of Observations}$$

 $(\sum means "sum of")$

The differences are squared to get rid of negative differences, because other wise everything would cancel out and our variance would be zero!

Standard Deviation - the square root of the variance. This is useful because it is in the same units as the original measurements!

3.3 Other Descriptors

Another descriptor that may prove useful is the **dispersion**, or the variance divided by the mean. This provides an estimate of how skewed the data is - for example, the first plot above has very low dispersion, while the second plot has high dispersion.

3.4 Ecological Community Descriptors

Many of you are interested in describing the species composition of of community. Here's a few common descriptors:

Species Richness - this is just the number of different species present.

Species Diversity - this is an index that takes into account the richness as well as the relative abundances of each species. E.g. Shannon's Diversity Index, where higher numbers mean more species more evenly distributed.

Species Evenness - this is an index that estimates specifically how evenly distributed species abundances are. E.g., Pielou's Evenness, which ranges from 0 to 1, with 1 meaning that each species has equal numbers.

Note: these measures can apply to any taxonomic distinction, e.g., family richness, order diversity, etc.

4 Inferential Statistics

Now, let's think about how to use your data to answer your questions. There are a couple approaches statisticians use, and we will talk about frequentist statistics, where probabilities are though of like relative frequencies. There is also Bayesian statistics, which is a bit more complex, so we will skip it for now.

Within frequentist statistics, we can run various tests to see how variables are related, which typically make some assumptions about the data. We can also do something called bootstrapping, which makes no assumptions, but can be simplistic from some perspectives.

In any case, we are hoping to estimate two main values:

Effect Size: How related are two variables? How different are two means? How much does one variable affect another?

p-value: What is the chance of observing data like yours (or something more extreme) if there was no relationship among the variables?

The p-value can be a bit tricky, but know that it **isn't** the probability that there is no relationship. P-values are used to draw conclusions from test results, a traditional guideline is that if the p-value is less than 0.05, the results are "significant." Some statisticians bristle at this arbitrary and binary system, so it's often best to report both the effect size and the actual p-value, so readers can interpret for themselves. The smaller the effect size, the weaker the relationship, the smaller the p-value, the stronger the evidence for the relationship.

Obviously, there are entire classes taught on this stuff (which may have taken or will take!), but we are thinking in just the basics for now.

4.1 Classic Frequentist Tests

Now let's go over some statistical tests! For this section, it can be useful to remind ourselves of the variables involved in a research question:

Independent / Explanatory / Predictor Variable: this is either what you are manipulating in an experiment or what your study is designed to capture variation in (e.g., C02 at BioCON, species richness at BigBio).

Dependent / **Response Variable**: these are what you measure or observe throughout your study, generally hypothesizing that they will differ among the levels of your independent variable (e.g., aboveground biomass in BioCON or BigBio).

4.1.1 Assumptions

We should mention what these tests generally assume about your data.

First, they assume that your **data are independent**. This just means that no two observations of your data are more related to eachother in a way that isn't accounted for by a variable. Say you were comparing mean tree height between two forests - individual tree heights in the same forest would be independent, but two measures of the same tree on different days would be non-independent.

Second, they assume that the **errors are normally distributed**. This is a bit more confusing without a statistical background. An example may be illustrative - in the tree height example above, we assume that the individual tree height are normally distributed about the mean. Without getting too much into the weeds, if you collect enough data (i.e., 30+ observations), these errors will likely be approximately normally distributed. However, things get divcey when we deal with data that is not continuous like tree height, for example, discrete count data - more on that below.

Third, they assume **homogeneity of variance**. This is another complicated one, but it mean that the variance of the errors doesn't change with the independent variable. In the tree example, we are assuming that the variance of the differences between observed tree heights and the forest mean does not change between forests.

Data that break the first assumption are difficult to deal with outside of accounting for the non-independence factor (which can severely reduce the size of your sample), but failing to meet the second or third assumptions generally leads to transforming data or using alternative tests.

4.1.2 Categorical Predictor/s, Numeric Response

4.1.2.1 Two Predictor Categories

When you are comparing numeric values from two groups, you can use a **t-test** to compare their means. T-tests can be **paired** when each observation in one group is specifically linked to an observation in the other group (e.g., masses of sibling plants in separate treatments) which can be more powerful. When the variance of values in each group changes, you can do a **t-test with unequal variance**.

The effect size here is the difference between means.

4.1.2.2 More Than Two Predictor Categories

If you have more than two groups/categories, you can use a **Analysis of Variance** or **ANOVA**. This will tell you if the means of each group are equivalent, or if there is at least one inequality. You can test for pairwise comparisons among the groups with **Tukey' test**. If you have multiple categorical predictors, you can do **two-way or three-way ANOVAs**. Tests with more than three categorical predictor variables are uncommon and harder to interpret.

The effect sizes are the pairwise difference in means.

4.1.2.3 Ordinal Predictors

When your predictor variable is ordinal, the quick and easy way to analyze it would be to convert the predictor to a numeric integer data type and proceed from there. However this is imprecise...

This section is under construction

4.1.3 Numeric Predictor/s, Numeric Response

4.1.3.1 Simple Association

When all you are interested in is whether two numeric variables are related to one another, not cause and effect, you can do a **correlation test**. **Pearson's correlation** is generally applicable for continuous data. **Spearman's correlation** is good for when you are dealing with data with non-normal distributions, like count data (it also works for ordinal data!).

The effect size here will be a correlation coefficient ranging from -1 to 1, with -1 means an inverse relationship, 0 means no relationship, and 1 mean a direct positive relationship.

4.1.3.2 Cause and Effect

When you are suppose a causal relationship between numeric variables, you can use a **linear regression**. This will use linear algebra or maximum likelihod estimation (don't worry about it) to find the best fit line that describes the relationship between two variables; where the sum of the squared distances from the observations to the line is minimized. You can also include multiple predictor variables to perform **multiple linear regression** AKA **multivariate linear regression**.

When your response variable is count data, the assumptions of simple linear regression are usually unmet, so you can use generalized forms like a **Poisson regression** or a **Negative Binomial Regression**.

The effect sizes here are the parameter coefficients, i.e., how much does the response change for on unit increase in the predictor? Note: these are not straightforward for Poisson and negative binomial regression, so ask your mentor.

4.1.4 Numeric Predictor/s, Categorical Response

4.1.4.1 Binary Response

When your categorical response is only two categories (e.g., presence or absence), you can use a **binomial regression** AKA **logistic regression**. This works similarly to linear regression, but the effect sizes are measured in log odds, which is difficult to interpret, but can be transformed to estimating how the probability of one category value over the other increases with a variable.

4.1.4.2 Multiple Response Categories

Multinomial regression (under construction)

4.1.5 Categorical Predictor/s, Categorical Response

Chi-square test (under construction)

4.2 Bootstrapping

One alternative to these classic tests has no assumptions: bootstrapping. Essentially, it involves using the sampled data to simulate more samples, and compare your observations to those simulations.

Empirical Bootstrapping is where you take your actual observations and shuffle which value is associated with which observation. For example, you could take measurements of tree heights from two forests, and randomly assign forest ID to each measurement.

Parametric Bootstrapping is where you summarize your observed data and use it to generat simulated data. For example, you could calculate the mean and variance of tree heights in two forests and then generate simulated forests of trees through random pulls from a normal distribution with the appropriate mean and variance.

With both approaches, you simulate a large number of simulated datasets (1000+), and then calculate whatever you are interested in for each of those simulations, and compare the calculation from the observed data to the distribution of simulated values. For example, if you empirically bootstrap the two forests of tree heights 1000 times, and then calculate difference

in means for each you will have 1000 mean difference values. The proportion of those simulated values that are equal to or more extreme than your observed mean difference is your p-value!

Part II R on your Computer

5 R Itself

R is both a programming language and an application that you can install to your computer.

5.1 R, the Language

R is a programming language designed for statistical computing, and is often the language of choice for scientists. R is also used for data science in some business, tech, and health contexts (but many prefer Python in those areas).

As a programming language it is essentially an expandable collection of functions with syntax to perform tasks, and it could be written in any text editor. However, in order for your computer to interpret the language, it needs some software.

5.2 R, the Software

The R application allows you to run R code on your computer, and comes with a basic "console" window where code is run and output is printed, as well as a basic script editor where you can write code to run.

You can download the application from here:

https://cran.r-project.org/

If you are asked to select a mirror, simply select the nearest one (I believe Iowa State should work).

If you have a Windows machine, it should be fairly straightforward to simply download and install the "base" R from the link.

If you have a Mac, you will want to select the .pkg file that matches your processor type: x-86 for Intel processors (mostly Macs pre-2020), arm64 for Macs with the M1 or M2 chip (most Macs post-2020).

If you are using Linux, you know more than me.

5.3 R Packages

As mentioned above, R is *expandable*. You can add more functionality to R by installing packages. Packages contain more options of code to use to process and analyze data, and also do many other things.

Packages can be installed through writing R code, or by clicking some buttons in RStudio. Then they will live in a directory that was built when you installed R for auxiliary packages.

We will discuss more about installing packages in the R coding section.

6 R Studio

While you can use R with just the basic application, it is much easier and beginner-friendly to use RStudio, which is an integrated development environment or IDE. This is just an application that provides a suite of features to make programming easier for users. In fact, I'm typing this in RStudio *right now*!! Note: you must have the R application installed to use RStudio, as it relies on the R application to interpret R code.

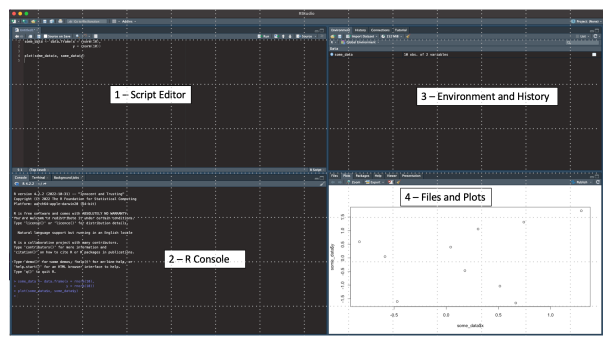
You can download and install RStudio from here:

https://posit.co/download/rstudio-desktop/

Which should be more straightforward than downloading and installing R.

6.1 RStudio at a Glance

If you open up RStudio, you will see something like this:



- 1- Script Editor: Here is where you will write code! You can create an R script (a text document to save code in) with the file tab, and write what you need in the resulting script. It is **highly** recommended to use scripts, because then you can save your code for later, and troubleshoot errors easier. From this window, you can highlight code and run it with the "Run" button on top, or with Ctrl + Enter / Cmd +Enter.
- 2- R Console: Here is where the action happens code will run here, and text output, warnings and messages will be displayed. You can also type code into the console, but that is only recommended for installing packages, entering credentials, rendering documents, and things of that nature. Don't type your data processing or analysis code into the console, use a script instead! There's also a terminal tab if you ever need to perform shell commands.
- **3- Environment and History**: Here you can find a list of the variables and data you have loaded into your "workspace" or "environment" in the Environment tab. These are objects you can do stuff with with code. You can also click the History tab to see the code you have run thus far.
- **4- Files and Plots**: Here is where any figures you draw will pop up (and you can save them from here as well). There is also a Files tab that allows you to navigate through your file directory (helpful with projects, described below). The Packages tab shows which packages you have installed and loaded (you can also click "Install" at top to easily install new ones!). Finally, the help tab is where you can search for the documentation on any R function.

6.2 R Projects

It is highly recommended to use R Projects when working with RStudio. Projects are essentially just subdirectories in your file folders, but they come with a special .Rproj file that RStudio can read and use. This helps you organize your work, and makes your code more easily portable.

You can create a new projects from the File tab at upper left, or in the project dropdown menu at upper right. You can just create one in a new directory. Then you can select a name and where you want to save it.

There are many different types of projects - this book/website is one!

If you want to backup your work with version control or collaborate with others using git and GitHub, you will need to use projects. (Well, technically you don't need to, but you'd be doing many things manually).

7 Optional: Git and Github

If you are interested in:

1. Backing up your code using a version control system that allows you to roll back changes and monitor incremental progress

and/or

2. Sharing your code and collaborating with others

You may like to try using git (a program for your computer) and GitHub (a website that hosts code projects).

We won't go into detail here, but Jenny Bryan's excellent introduction and tutorial on the topic can be found here:

https://happygitwithr.com/

Part III R Programming

8 The Basics

8.1 Intro

I'm sure those of you reading this come from a wide variety of backgrounds regarding computer programming - some of you may be very familiar with it, others total novices. Some of you may love computing, others might hate it. If you're apprehensive about learning R, or if you find yourself struggling with it - don't worry! Scientific computing presents a challenge at some point to everyone who does it. Just remember a few things:

- 1. Everyone makes mistakes.
- 2. Don't be afraid to ask questions!
- 3. Don't compare yourself to others, compare you today to you yesterday.
- 4. Everyone is constantly learning new things, including those who seem like experts.

That said, learning a programming language is a little like learning a human language, except there's a much smaller vocabulary and the grammar is *very* strict. And where human language has parts of speech like nouns and verbs, R has a certain syntax as well. Some of the main components of the R language are **operators**, **functions**, **arguments**, and **data**.

8.2 Operators

Operators are short symbols that tell the computer to do certain simple things. You are already familiar with many operators - the **math operators** like +, -, *, and /. R at its simplest is a calculator:

```
## This is block of R code! Anything that starts with # is a comment, and doesn't run.
## adding
2 + 2
```

[1] 4

```
## subtracting
  5 - 4
[1] 1
  ## multiplying
  3 * 3
[1] 9
  ## dividing
  6 / 2
[1] 3
There are a couple other math operators too:
  ## exponentiate with ^
  3^2
[1] 9
  ## find the remainder with the modulus, %%
  10 %% 3
[1] 1
  ## perform integer division with %/%
  10 %/% 3
[1] 3
```

But math operators aren't the only type! There are also the closely related **comparison operators**, which will return TRUE or FALSE instead of calculated numbers:

```
## equals, ==
2 + 2 == 4
```

[1] TRUE

```
## does not equal, !=
2 + 2 != 4
```

[1] FALSE

```
## greater than, > 5 > 4
```

[1] TRUE

```
## less than, < 5 < 4
```

[1] FALSE

There are also greater than or equal to (>=) and less than or equal to (<=).

You can combine comparisons with **logical operators** - and (&), or (|), and not (!):

```
## and: are both true?
(3 > 2) & (4 > 3)
```

[1] TRUE

```
## or: is at least one true?
(2 == 1) | (4 < 3)
```

[1] FALSE

```
## not: is this false?
!(2 == 1)
```

[1] TRUE

There are few other important operators, but they will make more sense once we talk about the other parts of R.

8.3 Functions

Functions are words (though not necessarily real words) or letters that instruct the computer to perform more complicated tasks. They generally are followed by parentheses ().

```
## here's a function that returns the current date
Sys.Date()
```

[1] "2023-07-28"

```
## and here is a function that returns the date with the time
Sys.time()
```

```
[1] "2023-07-28 15:25:37 CDT"
```

No you may be thinking - "this is pretty basic" and "what are the parentheses for?", which brings use to arguments!

8.4 Arguments

Arguments are values or objects that go inside the parentheses of functions to specify what you want the function to do. This is what gives functions their power. Arguments are separated inside a function by commas.

```
## the sum function can sum many numbers sum(1,2,3,4,5)
```

[1] 15

In the function above, each number is acting as an argument. In this case, the arguments don't have names. Oftentimes a function's arguments will be explicitly named, and to specify what you want those arguments to be, you use the = operator.

```
## this function pulls values randomly from a normal distribution specified in the argument ## n specifies how many numbers to return, and mean and sd specify shape of the distribution sd rnorm(sd = 10, mean = 5, sd = 1)
```

- [1] 3.377963 6.032801 5.011091 5.136840 4.011441 7.430791 4.382403 5.241645
- [9] 3.945753 4.513725

Operators are actually a special type of function that can be used with syntax that is more intuitive for them. You can also use them in the same way as most functions by surrounding them with back ticks, '.

```
## here we use the + operator in a much more confusing context \dot{}+\dot{}(2, 2)
```

[1] 4

```
## it is equivalent to
2 + 2
```

[1] 4

8.5 Data

We are using the word data here to broadly encompass values (like the numbers we were using above, both with operators and as arguments), variables (stored values), and data structures (organized collections of values).

8.5.1 Values

Values are much like the data types we discuss in the data analysis section. In fact, the different types of values R can deal with are called data types as well!

In R, values can be numeric, character, or logical (among other, more specific types).

```
## numeric values are numbers!
  2
[1] 2
  2.5
[1] 2.5
  ## character values are letters, words, phrases (often referred to as "strings)
  "a"
[1] "a"
  "apple"
[1] "apple"
  "there is a worm in my apple"
[1] "there is a worm in my apple"
  ## note: character values or strings must be surrounded by "" or '' for R to interpret the
  ## Logical values are TRUE or FALSE (you've seen these above)
  TRUE
[1] TRUE
  FALSE
[1] FALSE
```

There are other types of values too: missing values (NA and NaN), infinite values (Inf and -Inf), and something that indicates empty (NULL).

8.5.2 Variables

Variables are named values that are stored in the "environment", or the workspace that R can access to perform its tasks. In order to store a value as a variable, you need to use a special kind of operator called an **assignment operator** (<- or =). As I mentioned variables have names, which are unquoted text.

```
## store 2 as a variable called x
x <- 2

## R returns no output here because you're just storing a value
## but you can return the value by calling the variable
x</pre>
```

[1] 2

```
## store 3 as a variable called y
y <- 3
## you use variables with operators
x + y</pre>
```

[1] 5

```
## store a character value
string <- "hello"

## math doesn't work on strings</pre>
```

Technically, you can use = in place of <-. This is why the equals operator is ==. I generally use <- to prevent any confusion between assignment and comparison.

8.5.2.1 Naming Rules

Variables have rules about how they can be named:

- 1. No special symbols other than _ and .
- 2. You can't start with a number or __.
- 3. They can't be special words that R interprets differently. You can enter ?Reserved in your console to see a list.

8.5.3 Data Structures

Data structures are collections of values with some sort of organization, and also saved in the environment. Plot twist: the variables above are a the simplest data structure, the **scalar**, which is just a single value.

The next data structure is the **vector**, which is a collection of values of the *same data type*. We can store them much like variables.

```
## we use another operator, :, to create a sequence of integers from 1 to 10
my_vector <- 1:5

my_vector

[1] 1 2 3 4 5

## you can also create vectors with the combine function, c()
my_other_vector <- c("a", "b", "c")

my_other_vector

[1] "a" "b" "c"</pre>
```

The next data structure is called a **list**. A list is a collection of values like a vector, but they can be of any data type, or data structure. You can have a list of numeric values and character values, a list of vectors, or even a lists of lists! Every other complex data structure is technically a list with special attributes and/or rules.

```
## you can create lists with the list function
my_list <- list("a", 1, 2:4)

my_list

[[1]]
[1] "a"

[[2]]
[1] 1</pre>
```

```
## can also use the combine function, but it will default to a vector when data types are
my_other_list <- c("b", 2)</pre>
```

Finally, the most common special type of list you will use is the **data frame**. A data frame is a list of vectors that are arranged in a table, much like an excel spreadsheet. Each of the vectors will be named as a column, and all must be the same length. The position of a value in a vector is its row in the data frame.

Next, we will extend these concepts a bit further!

9 Next Steps

Now that we have the basic "parts of speech" of R down, we can move toward what we can do with them.

What follows are a few unconnected topics that will prove as useful background to working with data in the later sections.

9.1 Packages

Packages are collections of R functions that people write to make tasks easier. One of the strengths of R is that countless programmers have taken the time to assemble functions of use in their respective fields, and shared them with the world. For example the "vegan" package contains a number of functions geared towards community ecology, like calculating diversity indices. You could calculate a diversity index with just the base R, but it would be more difficult and take longer.

You can install packages in at least two ways:

1. You can use the following code, with the package names in quotes (this is one of the few times where using the console is recommended, because you only need to install a package once):

```
install.packages("PACKAGE NAME HERE")
```

2. Or you can use the packages tab in RStudio. In the lower right panel, there should be a packages tab in between "Plots" and "Help". Once there, there is an "Install" button. When clicked a window will appear allowing you to search for packages to install.

But Installing packages does not make them automatically accessible to you. When R boots up, it only loads its base functionality by default, so you have to load any packages that you want to use for a given R session. You can do this with the following code (with the package name not in quotes):

```
library(PACKAGE NAME)
```

The code for loading packages should be saved in your r script, because it will need to be done every time you open R.

There is a family of packages that is very popular called the "tidyverse." The aim of the tidyverse is to make data manipulation and visualization streamlined and efficient. Some people are very opinionated about whether you should use the tidyverse or base R, but in my opinion, it's mostly silliness. If you only want to dip into R and don't plan to use it much in the future, you may as well just pick up the specific functions you need to use and not worry about much else. If you'd like to continually use R for data analysis, but don't plan on getting deep into it, getting a handle on the tidyverse may be a good idea. If you want to really get into R, I would recommend learning how to do things in base R (as well as tidyverse functions).

You can install the tidyverse suite with:

```
install.packages("tidyverse")
```

9.2 Subsetting

In the last section we introduced data structures. Now let's talk about what you can do with them.

9.2.1 Vectors

The individual elements of a vector can be accessed with bracket operators - [and]. You can refer to an element by its index, or its numeric place in the sequence of elements (e.g., the 1st, the 10th, etc.). It's important to not here that R starts counting at 1, while many other programming language start counting at 0 (e.g., Python). This is another thing that people are opinionated about, and if you put your mind to it, you can be too! Anyway, here are some examples:

```
## let's create a vector of the first five letters of the alphabet
my_vector <- c("a","b","c","d","e")
my_vector

[1] "a" "b" "c" "d" "e"

## now let's return the 5th element
my_vector[5]</pre>
```

```
[1] "e"

## we can return multiple elements with c()
my_vector[c(2,4)]

[1] "b" "d"

## or as a series with :
my_vector[2:4]

[1] "b" "c" "d"
```

You can also use negative numbers to exclude values from what's returned:

```
## lose the last element
my_vector[-5]

[1] "a" "b" "c" "d"

## everything but the last element
my_vector[-1:-4]

[1] "e"
```

9.2.2 Lists

Subsetting vectors is fairly straightforward, but subsetting lists can be tricky. Since lists have multiple levels of organization, they use both the [] operators and the [[]] operators. Single brackets give you the list element, and double brackets give you what the list element contains. Let's demonstrate:

```
## create a list
my_list <- list(c("a","b","c"), "d", "e")
my_list</pre>
```

```
[[1]]
[1] "a" "b" "c"
[[2]]
[1] "d"
[[3]]
[1] "e"
  ## grab the first list element
  my_list[1]
[[1]]
[1] "a" "b" "c"
  ## grab what's conatined in the first list element (in this case a vector)
  my_list[[1]]
[1] "a" "b" "c"
  \#\# another example with a scalar
  my_list[2]
[[1]]
[1] "d"
  my_list[[2]]
[1] "d"
  ## you can also subset what you have subsetted:
  my_list[[1]][1]
[1] "a"
```

```
## but if you try subsetting a list element, it won't work the same way
my_list[1][1]

[[1]]
[1] "a" "b" "c"

## this is because [] returns the list element as a list of length 1, therefore [1] gives
```

This distinction can be difficult to understand, but don't worry! It takes time. The best analogy I've seen is from Hadley Wickham here:

https://adv-r.hadley.nz/subsetting.html#subset-single

You can think of a list as a train, every list element is a train car, and each has its own contents. Single brackets give you the train car/s, and double brackets gives you what's inside a single train car. And even a single train car is a train (or a list). Also note:

```
## you can grab multiple list elements with []; this give a list with two elements
  my_list[1:2]
[[1]]
[1] "a" "b" "c"
[[2]]
[1] "d"
  ## list elements can be named and indexed by their name as well
  named_list <- list(first = 1:3, second = 10)</pre>
  named_list
$first
[1] 1 2 3
$second
[1] 10
  named_list["first"]
$first
[1] 1 2 3
```

9.2.3 Data Frames

Subsetting data frames is a little easier to get a handle on, you just need to think in two dimensions. When using single brackets to subset data frames, you need to specify the index of the row and the column separately and in that order. You separate each index number by a comma inside the brackets. Check it out:

```
## create data frame
  my_data <- data.frame(letter = c("a", "b", "c"), # each column has a name
                         number = c(1, 2, 3),
                         vowel = c(TRUE, FALSE, FALSE))
  my_data
 letter number vowel
              1 TRUE
1
       a
2
              2 FALSE
       b
3
       С
              3 FALSE
  ## grab the element in the 2nd row, 1st column
  my_data[2,1]
[1] "b"
  ## you can also grab a whole row or column by leaving onse side of the comma blank
  my_data[2,]
  letter number vowel
2
       b
              2 FALSE
  my_data[,1]
[1] "a" "b" "c"
  ## (subsetting a row gives you a data frame, subsetting a column gives you a vector)
```

But data frames also have named columns! Let's use that to our advantage. You can specify a column's name instead of its index in brackets, like for a list, or you can use the \$ operator.

```
## subsetting by name in brackets
my_data[,"vowel"]
```

[1] TRUE FALSE FALSE

```
## subsetting by name with $ (notice no quotes)
my_data$vowel
```

[1] TRUE FALSE FALSE

```
## the downside of $ is that you can't grab more than one column like with brackets
my_data[,c("letter", "vowel")]
```

```
letter vowel
a TRUE
```

2 b FALSE

3 c FALSE

```
## subsetting multiple columns gives you a data.frame
## you can use $ with named lists too
named_list$first
```

[1] 1 2 3

```
## you can mix subsetting operators if you ever need to my_{ata}vowel[1]
```

[1] TRUE

```
my_data[1,]$vowel
```

[1] TRUE

You can also use brackets to select rows by value, not index. You just need to use some comparison operator in a statement that resolves as TRUE or FALSE.

```
## grab the consonant rows
  my_data[my_data$vowel == FALSE,]
 letter number vowel
       b
              2 FALSE
              3 FALSE
3
  ## grab the rows before the third
  my_data[my_data$number < 3,]</pre>
 letter number vowel
              1 TRUE
1
2
       b
              2 FALSE
  ## you can combine criteria
  my_data[my_data$number < 3 & my_data$vowel == FALSE,]</pre>
 letter number vowel
2
              2 FALSE
       b
```

Now the reason that we talked about packages in between data structures and subsetting is because the tidyverse (specifically, the dplyr package) has more functions for subsetting: filter and select. Filter works much like grabbing rows by value, and select works like grabbing columns by name. Let's look at some examples:

```
## load the tidyverse
  library(tidyverse)
-- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
v dplyr
          1.1.1
                    v readr
                                 2.1.4
v forcats
          1.0.0
                     v stringr
                                 1.5.0
v ggplot2 3.4.2
                     v tibble
                                 3.2.1
v lubridate 1.9.2
                     v tidyr
                                 1.3.0
v purrr
           1.0.1
-- Conflicts ----- tidyverse_conflicts() --
x dplyr::filter() masks stats::filter()
                 masks stats::lag()
x dplyr::lag()
i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become
```

```
## filter for consonants
  filter(.data = my_data, vowel == FALSE)
 letter number vowel
      b
              2 FALSE
              3 FALSE
2
       С
  ## select letter related columns
  select(.data = my_data, letter, vowel)
 letter vowel
      a TRUE
1
      b FALSE
2
3
       c FALSE
  ## you can also exclude columns
  select(.data = my_data, !number)
 letter vowel
      a TRUE
1
2
      b FALSE
3
       c FALSE
  ## note: selecting a single column will give a data frame, not a vector
  select(.data = my_data, number)
 number
       1
1
       2
2
       3
  ## another tidyverse/dplyr function, pull, will give just a vector
  pull(.data = my_data, number)
```

[1] 1 2 3

As you can see, filter, select and pull are versatile, consistent and powerful. However, they lack one key ability: assignment. You can use brackets and \$s to assign things:

```
## assign a new value to a data element (NA means missing value)
  my_data[3,2] <- NA</pre>
  my_data
  letter number vowel
              1 TRUE
       a
2
              2 FALSE
       b
3
       С
             NA FALSE
  \#\# create a whole new column with \$ (vector must be of same length as the number of rows)
  my_data$new_column <- c("some", "new", "data")</pre>
  my_data
  letter number vowel new_column
              1 TRUE
       a
       b
              2 FALSE
                             new
       С
             NA FALSE
                            data
```

- 9.3 Optional: Flow Control
- 9.4 Optional: Writing Functions

10 Importing Data

The previous chapters focused on R's functionality, but you might not be feeling any closer to working with *your* data in R. We'll transition to that now! The first step is getting your data into your R environment, so that you can use it with R's functions.

10.1 Reading Data

Importing data into R is often referred to as reading data, as that is what the computer is doing, it's reading the contents of a file (usually a text file). Most ecologists and data scientists work with a text file called a Comma Separated Value file, or csv. This is a small file that's easy for computers to read where each column is separated by a column, and each row by a new line. You can save excel files as csv from the "Save As..." menu, and you can specify csv as the type when downloading a Google sheet.

10.1.1 From Your Computer

So when you have files you want to read locally on your computer, the first thing you need to think about is what's called the "working directory". The working directory is the folder on your computer where R will look for files when prompted, and also where it will save output.

You can check your current working directory:

```
## return current working directory
getwd()
```

[1] "/Users/kit/Documents/UMN/Research/cedar_creek_projects/ccesr_intern_hub"

You can also set your working directory manually

```
## change working directory
setwd("some/different/folder")
```

Or, in RStudio, you can click the Session dropdown menu at the top of the window, then "Set Working Directory", then "Choose Directory."

If you use an R Project (**highly recommended**), you don't have to worry as much about this. If you have a project open, the working directory will be automatically set to the folder that contains the .Rproj file that is created when you create a project. See Section 6.2 for more info!

When you're in an R Project, or have a csv you want in your working directory, you can read it into your environment like so:

```
## read data
my_data <- read.csv("the name of your file in quotes", header = TRUE)</pre>
```

The read.csv function creates a data frame from the csv you specify, and then the <- assigns it to "my_data." The "header = TRUE" argument tells R to interpret the first line of the csv as the column names.

10.1.2 tidyverse Function

The readr package in the tidyverse family also has its own data reading functions.

```
## load tidyverse
library(tidyverse)

## read data (assumes header by default)
my_data <- read_csv("name of your data in quotes")</pre>
```

These functions are pretty similar, with one exception: read.csv gives you a data frame, but read_csv gives you a "tibble." What is a tibble? It's another special type of list, much like a data frame, but with a few differences. IT was designed to work more consistently with tidyverse functions. One important difference between data frames and tibbles that when you subset an individual column with the brackets ([]), data frames will give you vectors, and tibble will give single column tibbles. This has caused me confusion when writing functions, but you may not run into it.

10.1.3 From The Web

You can also read files directly from the web. If you have your data in Google sheets, you can create a URL for R to import it directly. Simply go to the File menu, click "Share" and then "Publish to web". In the box that pops up, you will need to select the file type as ".csv", not web page. Then save the URL that it gives you!

For demonstration, I've created a few data sheets that you too can import into R by copying the following code:

```
## put the url of the data in quotes
fake_mammals <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vQ9mfx88nM33PC6W
fake_insects <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vT0snHMdsxzzzkxt</pre>
```

These two files will be used throughout the next chapters. The first is some made-up data of some mammal captures at 6 sites across forest and savanna habitats (with mass and parasite info), and the second is made up sweep-netting data from the same sites.

10.1.4 Other File Types

Under construction....

10.2 Checking Data

Now that you have data, you will want to look at it!

10.2.1 The Whole Table

You can look at a whole data frame by clicking on its name in the "Environment" pane in RStudio (upper right), or with the View() function:

```
View(fake_mammals)
```

You can also just look at parts:

```
## check top 6 rows
head(fake_mammals)
```

	site	site_type	sp	ecies	mass_g	${\tt tick_count}$	${\tt helminth_mass_mg}$
1	a	forest	White-footed	mouse	20	0	512
2	a	forest	White-footed	mouse	24	10	365
3	a	forest	White-footed	mouse	23	2	0
4	a	forest	White-footed	mouse	19	0	608
5	a	forest	White-footed	mouse	25	12	109
6	a	forest	Deer	mouse	22	3	456

```
## check bottom 6 rows
tail(fake_mammals)
```

	site	site_type	sp	pecies	mass_g	${\tt tick_count}$	${\tt helminth_mass_mg}$
43	f	savanna	${\tt White-footed}$	mouse	21	0	408
44	f	savanna	White-footed	mouse	25	1	197
45	f	savanna	White-footed	mouse	24	0	152
46	f	savanna	Deer	mouse	20	0	508
47	f	savanna	Deer	mouse	22	2	496
48	f	savanna	Meadov	vole	23	NA	56

You can also take a look at the structure of the data with str(), which will tell you how many rows (observations) and how many columns (variables), as well as the type of each column.

```
## check structure
str(fake_mammals)
```

10.2.2 Individual Columns

You can also take a look at individual columns with the \$ operator, and get quick summaries with summary():

```
## summarize mammal masses
summary(fake_mammals$mass_g)

Min. 1st Qu. Median Mean 3rd Qu. Max.
17.00 20.75 23.00 22.60 25.00 28.00

## summarize helminth masses
summary(fake_mammals$helminth_mass_mg)
```

```
Min. 1st Qu. Median Mean 3rd Qu. Max. 0.0 119.5 341.5 306.5 501.2 713.0
```

Note that for the second summary (helminth mass), it tells you how many NA's, or missing values, there are.

10.2.3 Factors: The Pseudo Data Type

Unless you specify, csv reading functions will assume the data type of each column in a data sheet (numeric, character, etc.). Any categorical variable will be considered a character type generally. But the way character types are stored in computer memory does not lend itself well to statistical analysis. To remedy this, R has a special data type called the factor for categorical data. A factor is made up of two parts - the levels, which are stored to the computer as integers, and the labels, which are character strings that we can read as the category names.

In our mammal data, we probably want mammal species, site type, and site all to be factors. We can convert them with the as.factor function!

There are also similar functions for converting data types to numeric (as.numeric()) and character (as.character()).

11 Wrangling Data

Now that we have our data in, let's play with it!

Setup:

```
## read data
fake_mammals <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vQ9mfx88nM33PC6W
fake_insects <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vTOsnHMdsxzzzkxt

## convert species to factor
fake_mammals$species <- as.factor(fake_mammals$species)

## convert site type to factor
fake_mammals$site_type <- as.factor(fake_mammals$site_type)

## convert site to a factor
fake_mammals$site <- as.factor(fake_mammals$site)

## also do for insects
fake_insects$site <- as.factor(fake_insects$site)
fake_insects$site_type <- as.factor(fake_insects$site_type)</pre>
```

11.1 Adding Columns

One simple thing you may want to do is add columns to your data, which may be calculations from existing columns.

11.1.1 Base

In base R, we have already kinda done this. You can assign something to a new column with the \$ and <- operators.

For the insect data, let's say we wanted to calculate average temperature at a given site based on the recorded high (temp_hi) and low (temp_low):

```
## calculate mean temp
fake_insects$temp_mean <- (fake_insects$temp_hi + fake_insects$temp_lo)/2
## check it out
fake_insects$temp_mean</pre>
```

[1] 21.25 23.25 19.25 26.75 24.00 28.00

11.1.2 tidyverse

In the tidyverse, adding new columns is done with the mutate function:

```
## load tidyverse
  library(tidyverse)
-- Attaching core tidyverse packages ----- tidyverse 2.0.0 --
         1.1.1 v readr
v dplyr
                                   2.1.4
v forcats 1.0.0 v stringr 1.5.0
v ggplot2 3.4.2 v tibble 3.2.1
v lubridate 1.9.2 v tidyr 1.3.0
           1.0.1
v purrr
-- Conflicts ----- tidyverse_conflicts() --
x dplyr::filter() masks stats::filter()
x dplyr::lag()
                masks stats::lag()
i Use the conflicted package (<a href="http://conflicted.r-lib.org/">http://conflicted.r-lib.org/</a>) to force all conflicts to become
  ## mutate a new column
  fake_insects <- mutate(.data = fake_insects, ## specify data</pre>
                          temp_mean_mutated = (temp_hi + temp_lo)/2) ## calculate new column
  ## this column should be the same for all six rows (a TRUE should be returned for each)
  fake_insects$temp_mean == fake_insects$temp_mean_mutated
```

[1] TRUE TRUE TRUE TRUE TRUE TRUE

As you can see, using mutate mean you have to write the name of data frame fewer times.

11.2 Pivoting / Reshaping

You also may need to transform your data between the wide and long formats (recall the data management slides, which will be added to this book eventually). I find that the pivot functions from tidyr in the tidyverse are easier to use, so we will go over those. But by all means, if you prefer base, go for it! I'm just less familiar with using base R for this.

11.2.1 Wide to Long

The insect data is partly in wide format: we have a column for each order of insect, where the count is implicitly the values in the cells. IT will be easier to work with if we make one "count" column and one "order" column. We can do this with the pivot_longer function:

A tibble: 6 x 8

	site	site_type	temp_hi	${\tt temp_lo}$	${\tt temp_mean}$	${\tt temp_mean_mutated}$	order	count
	<fct></fct>	<fct></fct>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>	<chr></chr>	<int></int>
1	a	forest	23.5	19	21.2	21.2	${\tt hymenoptera}$	9
2	a	forest	23.5	19	21.2	21.2	lepidoptera	3
3	a	forest	23.5	19	21.2	21.2	coleoptera	16
4	a	forest	23.5	19	21.2	21.2	diptera	29
5	a	forest	23.5	19	21.2	21.2	odonata	4
6	a	forest	23.5	19	21.2	21.2	hemiptera	10

Now we have multiple rows for each site, one for each order! You may not believe me, but this will make things easier down the line.

For reference, the cols argument specifies which columns you want to pivot. The names_to argument names the column that will contain the pivoted column names, and the values_to argument names the column that the cell values will be placed.

11.2.2 Long to Wide

You'll occasionally want to turn long to wide as well. pivot_wider works for this

This is the inverse of what we just did - we made a column for each value in the column given to the names_from argument, the values of which are pulled from the column given to the values_from argument.

11.2.3 Base

You can pivot data in base R with reshape, but this section is under construction...

11.3 String Manipulation

Under construction....

11.3.1 Base

11.3.2 tidyverse

12 Summarizing Data

Now let's describe our data!

Setup:

```
## load tidyverse
library(tidyverse)
## read data
fake_mammals <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vQ9mfx88nM33PC6W
fake_insects <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vTOsnHMdsxzzzkxt
## convert species to factor
fake_mammals$species <- as.factor(fake_mammals$species)</pre>
## convert site type to factor
fake_mammals$site_type <- as.factor(fake_mammals$site_type)</pre>
## convert site to a factor
fake_mammals$site <- as.factor(fake_mammals$site)</pre>
## also do for insects
fake_insects$site <- as.factor(fake_insects$site)</pre>
fake_insects$site_type <- as.factor(fake_insects$site_type)</pre>
## lengthen the order count data
long_insects <- pivot_longer(data = fake_insects,</pre>
                              cols = c(hymenoptera, lepidoptera, coleoptera, diptera,
                                        odonata, hemiptera, orthoptera, ephemeroptera,
                                        tricoptera, plecoptera),
                              names_to = "order",
                              values_to = "count"
```

12.1 Describing With Summaries

You'll often want to give simple, illustrative information about the data you collected. The tidyverse is great for this!

12.1.1 tidyverse

The package dplyr in the tidyverse has two wonderful functions: group_by() and summarize(). You can also use the British spelling, "summarise()", but I use a z because it's what the founders would have wanted.

Before we use these however, we need to introduce a very useful operator, the pipe: %>%. This operator directs data into the first argument of a function, which allows you to chain functions together efficiently. Let's try an example with the filter and select subsetting functions (see Section 9.2.3):

```
## grab only the forest sites from the insect data
forest_sites <- fake_insects %>% ## take fake_insects and pipe it into filter...
  filter(site_type == "forest") %>% ## filter only forest rows, pipe into select
  select(site) ## select only the site column
  ## the whole pipe chain is assigned to "forest_sites"

forest_sites

site
1    a
2    b
3    c

## this is the same as
forest_rows <- filter(fake_insects, site_type == "forest")
forest_sites <- select(forest_rows, site)</pre>
```

Note: base R also has a pipe, |>. It's newer and mostly the same as %>%, so I just haven't transitioned.

Now, let's try with group_by() and summarize()! Let's say you wanted the total number of insects caught at each site (be sure to have pivoted your insect data in the last chapter!):

```
## summarize total insect catch
insect_counts <- long_insects %>%
```

```
group_by(site) %>% ## group observations
    summarize(total_insects = sum(count)) ## sum all insects
  insect_counts
# A tibble: 6 x 2
 site total_insects
  <fct>
                <int>
                   84
1 a
2 b
                   87
3 c
                  136
4 d
                   60
5 e
                   61
6 f
                   51
```

As you can see, the summarize function works a bit like the mutate function: you

Note that group_by doesn't visibly change your data, but it changes some attributes that the computer can see when it runs the summarize function. If you forgot which sites are in which type of habitat, you could also include that variable in the group_by arguments (since it doesn't subdivide the sites, it won't change the calculation).

```
insect counts <- long insects %>%
 group_by(site, site_type) %>%
  summarize(total insects = sum(count))
```

1 coleoptera

You can also calculate means and variances! You can use the mean, var, and sd functions. Let's try for each order across all sites:

```
order_summary <- long_insects %>%
    group_by(order) %>%
    summarize(count mean = mean(count), ## you can do multiple sumaries at once
             count_var = var(count),
             count_sd = sd(count))
  head(order_summary)
# A tibble: 6 x 4
 order count_mean count_var count_sd
 <chr>
                   <dbl>
                            <dbl>
                                     <dbl>
                15.8
                            43.4
```

6.59

2 diptera	17	163.	12.8
3 ephemeroptera	5.33	171.	13.1
4 hemiptera	6.17	42.6	6.52
5 hymenoptera	8.5	9.1	3.02
6 lepidoptera	5.67	5.87	2.42

You could also do this separately by site type:

`summarise()` has grouped output by 'site_type'. You can override using the `.groups` argument.

```
head(orders_by_habitat)
```

```
# A tibble: 6 x 5
# Groups: site_type [1]
```

	-				
	site_type	order	${\tt count_mean}$	${\tt count_var}$	$\verb"count_sd"$
	<fct></fct>	<chr></chr>	<dbl></dbl>	<dbl></dbl>	<dbl></dbl>
1	forest	coleoptera	20.7	17.3	4.16
2	forest	diptera	28.3	9.33	3.06
3	forest	ephemeroptera	10.7	341.	18.5
4	forest	hemiptera	11.7	14.3	3.79
5	forest	hymenoptera	10.3	2.33	1.53
6	forest	lepidoptera	4.33	2.33	1.53

Finally, you can save summarized output with write.csv() or write_csv():

```
## save summary to your working/project directory
## first argument is data, second argument is filename
write.csv(orders_by_habitat, "summary of orders")
```

12.1.2 Base

aggregate function (under construction...)

12.1.3 Making Nice Tables

the gt package is good for this (under construction....)

12.2 Community Ecology

Averages and variances are all well and good but what about ecological measures?

12.2.1 Richness

You may be interested in how many insect orders are represented in each site.

Let's do it in a pipe chain!

```
## order presence
  order_richness_site <- long_insects %>%
    mutate(presence = as.numeric(count > 0)) %>% ## create binary presence column
    group_by(site, site_type) %>%
    summarize(order_richness = sum(presence))
`summarise()` has grouped output by 'site'. You can override using the
`.groups` argument.
  order_richness_site
# A tibble: 6 x 3
# Groups: site [6]
 site site_type order_richness
 <fct> <fct>
                           <dbl>
1 a
        forest
                               8
2 b
        forest
                               7
3 c
        forest
                               9
4 d
                               6
       savanna
5 e
        savanna
                               7
6 f
                               6
        savanna
```

I calculated the presence column by checking if each value is positive (> 0), which returns a logical TRUE or FALSE, and then if you convert a logical variable to a numeric variable, TRUEs become 1s and FALSEs become 0s. Nifty!

12.2.2 Diversity

vegan package (under construction...)

12.3 Related Topic: Joining Data

Sometimes with summaries, you will want to connect them to other pieces of data. Here we have some insect counts by site, and some mammal data by site. Let's connect them! We can use the "merge" function from base R or the "join" functions from the tidyverse.

With merge:

```
## grab only the site and total columns from insect_counts
  ## this prevent doubling the site_type column
 merged_data <- merge(insect_counts[,c("site", "total_insects")], fake_mammals, by = "site"</pre>
  ## look at the new column in your data
  str(merged_data)
'data.frame': 48 obs. of 7 variables:
$ site
                  : Factor w/ 6 levels "a", "b", "c", "d", ...: 1 1 1 1 1 1 1 1 1 1 ...
$ total_insects : int 84 84 84 84 84 84 84 84 84 84 ...
$ site_type
                : Factor w/ 2 levels "forest", "savanna": 1 1 1 1 1 1 1 1 1 ...
$ species
                 : Factor w/ 3 levels "Deer mouse", "Meadow vole", ...: 3 3 3 3 3 1 1 1 2 2 .
$ mass_g
                  : int 20 24 23 19 25 22 22 21 23 20 ...
$ tick_count
                 : int 0 10 2 0 12 3 2 0 NA NA ...
$ helminth_mass_mg: int 512 365 0 608 109 456 521 432 20 129 ...
```

With join:

```
## there are different join functions for different contexts
## left_join keeps every row from the first data frame and adds any matching rows from the
## second. it works in most cases
## inner_join and full_join can also be useful
joined_data <- insect_counts %>%
    select(total_insects, site) %>%
    left_join(fake_mammals, by = "site")

## look at it
str(joined_data)
```

```
tibble [48 x 7] (S3: tbl_df/tbl/data.frame)
```

\$ total_insects : int [1:48] 84 84 84 84 84 84 84 84 84 84 ...

\$ site : Factor w/ 6 levels "a","b","c","d",..: 1 1 1 1 1 1 1 1 1 1 1 ...
\$ site_type : Factor w/ 2 levels "forest","savanna": 1 1 1 1 1 1 1 1 1 1 ...

\$ species : Factor w/ 3 levels "Deer mouse", "Meadow vole",..: 3 3 3 3 1 1 1 2 2 .

\$ mass_g : int [1:48] 20 24 23 19 25 22 22 21 23 20 ...

\$ tick_count : int [1:48] 0 10 2 0 12 3 2 0 NA NA ...

\$ helminth_mass_mg: int [1:48] 512 365 0 608 109 456 521 432 20 129 ...

13 Analyzing Data

Now let's draw some conclusions about our data, and maybe answer questions! Setup:

```
## load tidyverse
library(tidyverse)
## read data
fake_mammals <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vQ9mfx88nM33PC6W
fake_insects <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vTOsnHMdsxzzzkxt
## convert species to factor
fake_mammals$species <- as.factor(fake_mammals$species)</pre>
## convert site type to factor
fake_mammals$site_type <- as.factor(fake_mammals$site_type)</pre>
## convert site to a factor
fake_mammals$site <- as.factor(fake_mammals$site)</pre>
## also do for insects
fake_insects$site <- as.factor(fake_insects$site)</pre>
fake_insects$site_type <- as.factor(fake_insects$site_type)</pre>
## lengthen the order count data
long_insects <- pivot_longer(data = fake_insects,</pre>
                              cols = c(hymenoptera, lepidoptera, coleoptera, diptera,
                                        odonata, hemiptera, orthoptera, ephemeroptera,
                                        tricoptera, plecoptera),
                              names_to = "order",
                              values_to = "count"
```

13.1 Making Comparisons

First off, let's just do some simple comparisons.

13.1.1 t-tests

Let's say we want to compare two groups, like the number of insects caught in forests and savannas. We already created a summary of this in the last chapter:

```
## sum insects by site
  insect_counts <- long_insects %>%
    group_by(site, site_type) %>%
    summarize(total_insects = sum(count))
`summarise()` has grouped output by 'site'. You can override using the
`.groups` argument.
  insect_counts
# A tibble: 6 x 3
# Groups:
            site [6]
 site site_type total_insects
  <fct> <fct>
                          <int>
1 a
        forest
                             84
2 b
                             87
        forest
3 c
        forest
                             136
4 d
        savanna
                             60
5 e
        savanna
                             61
6 f
        savanna
                              51
```

It seems that there may be a difference! So let's run a t-test to test for a difference in means between two groups (see Section 4.1.2.1). Most stats functions in R can use the formula operator, ~. This allows us to connect our dependent variable (insect count in this case) as a function of our independent variable (site habitat type): total_insects ~ site_type.

```
## run the t test
habitat_comparison <- t.test(formula = insect_counts$total_insects ~ insect_counts$site_ty
## check the output</pre>
```

```
data: insect_counts$total_insects by insect_counts$site_type t = 2.6235, df = 2.1422, p-value = 0.1116 alternative hypothesis: true difference in means between group
```

Welch Two Sample t-test

alternative hypothesis: true difference in means between group forest and group savanna is no 95 percent confidence interval:

```
-24.30218 114.30218
sample estimates:
mean in group forest mean in group savanna
102.33333 57.33333
```

If we look at the ouput, it look like the forest mean was 102.333, and the savanna mean was 57.333, for a mean difference or effect size of 45. The p-value, or how strong the evidence for a relationship is, is 0.1116. This is higher than the traditional threshold for significance, likely because we have a very small sample size (6 total).

Note: if you check the help for the t.test function (run?t.test), you can find arguments for paired t-tests (paired) and unequal variances among groups (var.equal).

13.1.2 ANOVA

What if we have more than two categories, and we want to see if any two categories have different means? Let us return to the mammal data and compare the mass of helminths (parasitic worms) in different mammal species. We can run an analysis of variance (see Section 4.1.2.2).

```
## run the anova
helminth_comparison <- aov(fake_mammals$helminth_mass_mg ~ fake_mammals$species)

## check the output (now with the summary function)
summary(helminth_comparison)

Df Sum Sq Mean Sq F value Pr(>F)
fake_mammals$species 2 719721 359861 10.59 0.00017 ***
Residuals 45 1529035 33979
---
Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

Hey, that's a small p-value (0.00017)! That means we have strong evidence that there is at least one difference among the pairs of species, either between white-footed mice and deer mice, white-footed mice, and meadow voles, or deer mice and meadow voles. We can use a Tukey test to find out more:

```
## run tukey on the anova output
  helminth_tukey <- TukeyHSD(helminth_comparison)</pre>
  ## check it out
  helminth tukey
  Tukey multiple comparisons of means
    95% family-wise confidence level
Fit: aov(formula = fake_mammals$helminth_mass_mg ~ fake_mammals$species)
$`fake_mammals$species`
                                     diff
                                                 lwr
                                                             upr
                                                                     p adj
Meadow vole-Deer mouse
                                -356.2222 -544.58911 -167.85534 0.0001061
White-footed mouse-Deer mouse
                               -111.3333 -258.37719
                                                       35.71052 0.1698916
                                244.8889
White-footed mouse-Meadow vole
                                            70.26811 419.50966 0.0039996
```

At the bottom here we can see the pairwise comparisons. The two mouse species differ in helminth mass by $\sim 111 \, \mathrm{mg}$, but the difference is not significant. Meadow voles have a significantly different mean helminth mass from both mouse species. I guess the in the data I made up, voles have less helminth mass than mice.

13.2 Assessing Relationships

But what if you're not dealing with categorical comparisons? Then we can check for numerical associations.

13.2.1 Correlation

We can look for simple associations without cause and effect with correlations (see Section 4.1.3.1). Mice seem to have high helminth loads, so let's check for a correlation between their body mass and helminth mass:

```
## create a subset of only mouse data
## I use the %in% operator to specify that species should be found in a specified vector
## AKA, it could be white-footed mouse OR deer mouse
mouse_data <- filter(fake_mammals, species %in% c("White-footed mouse", "Deer mouse"))

## run correlation with two variables (no formula here)
cor.test(mouse_data$helminth_mass_mg, mouse_data$mass_g)</pre>
```

Pearson's product-moment correlation

Here we get an effect size of -0.874 (correlation coefficient), and a p-value of 3.741e-13, which means 3.741×10^{-13} , or «<0.001. This means there is a strong negative relationship observed between mouse mass and helminth mass, and we have very strong evidence for it.

13.2.2 Linear Regression

If we want to infer cause and effect we can use linear regression (see Section 4.1.3.2). Let's say we want to know if the number of insects at a site is predictive of mammal mass at a site. First let's join the two data frames like wed did in the last chapter:

```
## join our data
mammals_insects <- insect_counts %>%
    select(total_insects, site) %>%
    left_join(fake_mammals, by = "site")

## regress mammal mass on total insects with lm function
## this time I'm specifying the data frame with the data argument
## then I don't have to write it twice
mass_model <- lm(mass_g ~ total_insects, data = mammals_insects)</pre>
```

```
## look at the ouput with summary again
  summary(mass_model)
Call:
lm(formula = mass_g ~ total_insects, data = mammals_insects)
Residuals:
    Min
             1Q Median
                             3Q
                                    Max
-5.6018 -1.5936 0.0221 2.0487
                                 4.3982
Coefficients:
              Estimate Std. Error t value Pr(>|t|)
              20.15860
                          1.09484 18.412
                                            <2e-16 ***
(Intercept)
                          0.01184
total_insects 0.02808
                                    2.372
                                            0.0219 *
Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
Residual standard error: 2.549 on 46 degrees of freedom
```

Multiple R-squared: 0.109, Adjusted R-squared: 0.08959 F-statistic: 5.625 on 1 and 46 DF, p-value: 0.02195

If we look at the coefficient table, we can see that the total insect term has an estimate of 0.028, which is our effect size. For every added insect to a plot, the expected average mass of the mammal community goes up by 0.028g. Connected to that effect size is a p-value of 0.0219, which means we have strong evidence for the relationship.

13.2.3 Binomial Regression

If your response variable is binary (presence absence), you can use a binomial regression with the glm() function. Let's test if mammal mass effects the probability of having ticks attached (tick_count).

```
## first make a presence absence variable for ticks
fake_mammals$tick_presence <- as.numeric(fake_mammals$tick_count > 0)

## now do the regression, with the binomial "family"

tick_pres_model <- glm(tick_presence ~ mass_g, data = fake_mammals, family = "binomial")

## check it</pre>
```

```
summary(tick_pres_model)
```

```
Call:
glm(formula = tick_presence ~ mass_g, family = "binomial", data = fake_mammals)
Deviance Residuals:
    Min
              1Q
                 Median
                                3Q
                                        Max
-1.7550 -0.4805 0.1027
                            0.4404
                                     1.4731
Coefficients:
            Estimate Std. Error z value Pr(>|z|)
(Intercept) -22.3584
                         6.7603 -3.307 0.000942 ***
                                  3.341 0.000835 ***
mass_g
              0.9857
                         0.2950
---
Signif. codes:
                0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
(Dispersion parameter for binomial family taken to be 1)
    Null deviance: 54.040 on 38
                                  degrees of freedom
Residual deviance: 28.181 on 37
                                  degrees of freedom
  (9 observations deleted due to missingness)
AIC: 32.181
Number of Fisher Scoring iterations: 5
```

If we look at this like we looked at the linear regression, the mass_g term has a very small p-value meaning strong evidence for a relationship. It also has an effect size of 0.9857, meaning that the chance of having a tick increases with body mass. However, the units are in log odds, which are hard to interpret. The reason for this is some stats theory that is beyond the scope of this book.

13.2.4 Poisson / Negative Binomial Regression

under construction....

13.3 Multivariate Analysis

You can of course use multiple explanatory variables in your analyses. For example, when we regressed mammal mass on insect count, we ignored mammal species. We could include it like

so:

```
## multiple regression
multi_mod <- lm(mass_g ~ total_insects + species, data = mammals_insects)
summary(multi_mod)</pre>
```

Call:

lm(formula = mass_g ~ total_insects + species, data = mammals_insects)

Residuals:

```
Min 1Q Median 3Q Max -5.1418 -1.1856 0.1335 1.8338 5.1527
```

Coefficients:

	Estimate	Std. Error	t value	Pr(> t)	
(Intercept)	19.10561	1.36855	13.960	< 2e-16 *	**
total_insects	0.03490	0.01272	2.744	0.00875 *	*
speciesMeadow vole	-0.29446	1.08017	-0.273	0.78644	
${\tt speciesWhite-footed\ mouse}$	1.02943	0.86932	1.184	0.24270	

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Residual standard error: 2.545 on 44 degrees of freedom Multiple R-squared: 0.1505, Adjusted R-squared: 0.09254 F-statistic: 2.598 on 3 and 44 DF, p-value: 0.06421

Now we have multiple terms, and since species is categorical, the effect sizes and p-values are based on comparisons to a reference level (deer mouse in this case because it is first alphabetically).

We can look at the overall significance of species by running an ANOVA with aov, and summarizing the outtut:

we can use the model object in our aov function to save time, it will take the formula summary(aov(multi_mod))

```
Df Sum Sq Mean Sq F value Pr(>F)
total_insects 1 36.55 36.55 5.643 0.0219 *
species 2 13.93 6.96 1.075 0.3501
```

```
Residuals 44 285.00 6.48
```

Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

Looks like there is not much evidence of an effect of species.

You may be confused a bit by this code, but essentially t.tests, ANOVAs, and regressions are all "linear models", and we are specifying them and looking at them differently with different R functions. T learn more, I recommend taking stats classes!

Finally, you may be wondering which variable to include in your analyses. Model selection is another thing you would learn in stats, but the tl;dr could be: What is your question? Use those variables.

14 Visualizing Data

Now for what most consider the fun part, visualizing patterns in your data! Setup:

```
## load tidyverse
library(tidyverse)
## read data
fake_mammals <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vQ9mfx88nM33PC6W</pre>
fake_insects <- read.csv("https://docs.google.com/spreadsheets/d/e/2PACX-1vT0snHMdsxzzzkxt</pre>
## convert species to factor
fake_mammals$species <- as.factor(fake_mammals$species)</pre>
## convert site type to factor
fake_mammals$site_type <- as.factor(fake_mammals$site_type)</pre>
## convert site to a factor
fake_mammals$site <- as.factor(fake_mammals$site)</pre>
## also do for insects
fake_insects$site <- as.factor(fake_insects$site)</pre>
fake_insects$site_type <- as.factor(fake_insects$site_type)</pre>
## lengthen the order count data
long_insects <- pivot_longer(data = fake_insects,</pre>
                               cols = c(hymenoptera, lepidoptera, coleoptera, diptera,
                                        odonata, hemiptera, orthoptera, ephemeroptera,
                                        tricoptera, plecoptera),
                               names_to = "order",
                               values_to = "count"
## sum insects by site
insect_counts <- long_insects %>%
  group_by(site, site_type) %>%
```

```
summarize(total_insects = sum(count))
## join data
mammals_insects <- insect_counts %>%
select(total_insects, site) %>%
left_join(fake_mammals, by = "site")
```

14.1 General Notes on Data Visualization

There are a few things to keep in mind in general when creating figures, even outside of R:

Usually, figures should stand alone. This means that your figure can speak for itself, even without a caption. This means that axes and legends are clearly labelled, and trends are emphasized. It can also be helpful to annotate statistical output onto plots themselves.

When you can, **show your actual data**, instead of summary stats. Generally, when it's not too messy, seeing all the data points is more informative to the audience. For example, you could plot a comparison of means with a point for each mean, but you could show more if you plot every point behind those means.

It is advisable to shoot for a **low ink to info ratio**. Basically, you want to make your figures as uncluttered as they can be with extra print. Gray backgrounds, irrelevant fill colors, etc. should be avoided.

Finally, **remember accessibility**. Make color schemes appropriate for color-blindness, and make text large.

14.2 The tidyverse's ggplot2

When it comes to visualizing things in R, there are many methods. You can use the base R functions for plotting (plot, hist, lines, etc.), but I'm not super adept with them. Instead I'll be walking you through using ggplot2, a package in the tidyverse family that is incredibly popular for data visualization. There is a special syntax that may take some getting used to though.

Essentially, you create a ggplot "object" (which is another special type of list with unique attributes), and then you pipe it through a series of ggplot functions to add components, themes, labels, etc. However, ggplot2 is older than the %>% pipe we have used, so it uses an old and deprecated pipe operator: +. R automatically knows to interpret + differently with ggplot objects and functions.

Here is an example of code creating a ggplot figure:

```
## first create the ggplot object
## you need to specify your data in the data argument
## then there is a special set of arguments called aesthetic arguments
## (bound by the aes() sub-function)
## these specify what variables will inform aesthetics of your figure
## (e.g., axes, color, fills, sizes, etc.)
ggplot(data = your_data, aes(x = variable1, y = variable2, color = variable3)) +
geom_point(size = 2) + ## then you add geometry, this "geom" is for a scatterplot
labs(x = "Variable 1") + ## then you can add other things like labels
scale_color_manual(values = c("red", "blue")) + ## or specify scales
theme(axis.text = element_text(size = 12)) ## finally you can modify parts of the theme,
```

It may seem complicated at first, but if you start small and work yourself up, you'll be chaining together code to draw beautiful figures in no time!

14.3 Figure Types

Now we'll go over how to make some common figure types, based on your analyses.

14.3.1 One Variable: Continuous

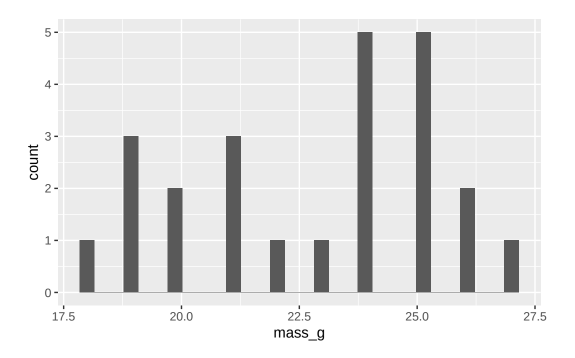
If you want to show the distribution of a single variable, you could use a histogram or a density plot.

For demonstration, let's make a plots of white-footed mouse masses.

```
## create a data frame of only white-footed mice
wf_mice <- filter(fake_mammals, species == "White-footed mouse")

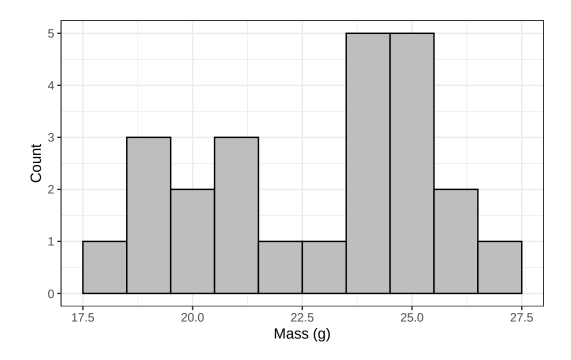
## make a ggplot, use wf_mice data, and specify mass as the x variable
ggplot(data = wf_mice, aes(x = mass_g)) +
    geom_histogram()</pre>
```

`stat_bin()` using `bins = 30`. Pick better value with `binwidth`.



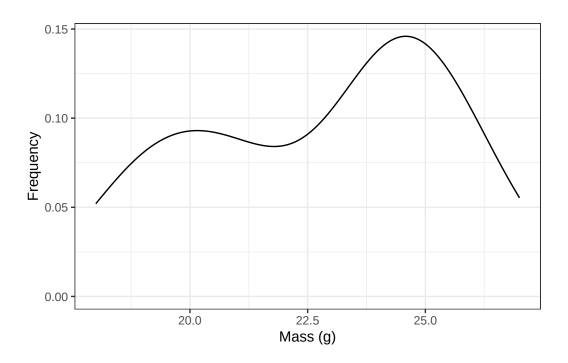
There, a simple histogram. Now let's play with how it looks:

```
## make a ggplot, use wf_mice data, and specify mass as the x variable
ggplot(data = wf_mice, aes(x = mass_g)) +
    ## give a wider binwidth to the histogram, and make it grey bars with black outlines
geom_histogram(binwidth = 1, fill = "grey", color = "black") +
    labs(x = "Mass (g)", y = "Count") + ## nicer labels
    theme_bw() ## my favorite simple theme
```



Cool! We could also look at this as a density plot! This will give more of a smooth line

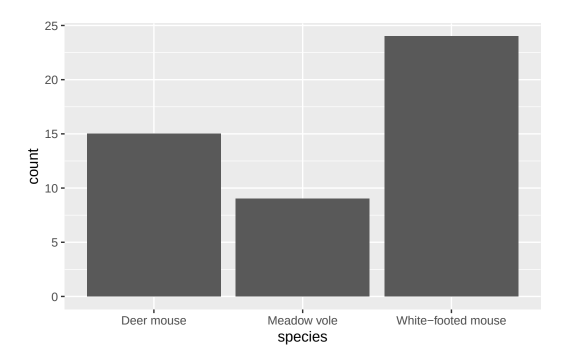
```
## make a ggplot, use wf_mice data, and specify mass as the x variable
ggplot(data = wf_mice, aes(x = mass_g)) +
geom_density() + ## create density plot
labs(x = "Mass (g)", y = "Frequency") + ## nicer labels
theme_bw() ## my favorite simple theme
```



14.3.2 One Variable: Categorical

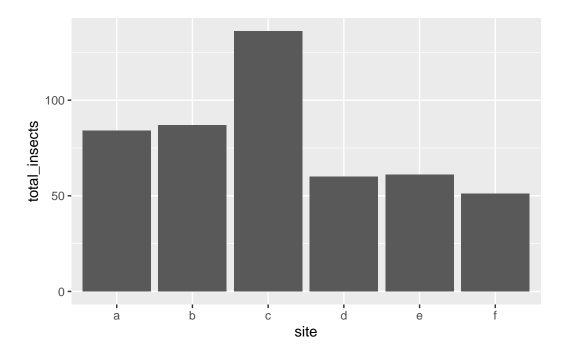
If you want to show how many observations are in each category, you can use a bar plot. In this demo, let's make a bar plot of how many of each mammal species were caught.

```
## specify x as species
ggplot(data = fake_mammals, aes(x = species)) +
  geom_bar() ## make bar plot
```



The geom_bar function will count up all the observations of each species level to inform its bars. Thus, it is assuming you are giving it long data. Another closely related function is geom_col, which just makes a bar as tall as a number value in the data. For example, let's make a bar plot of how many insect were caught at each site.

```
## need to specify two variables this time, one for the category, one for the count value
ggplot(data = insect_counts, aes(x = site, y = total_insects)) +
  geom_col()
```



As you can see, your data format will determine whether you should use geom_col or geom_bar. Note: bar plots are generally only best-suited for counts among categories, when you're dealing with measured variables, there are better options below.

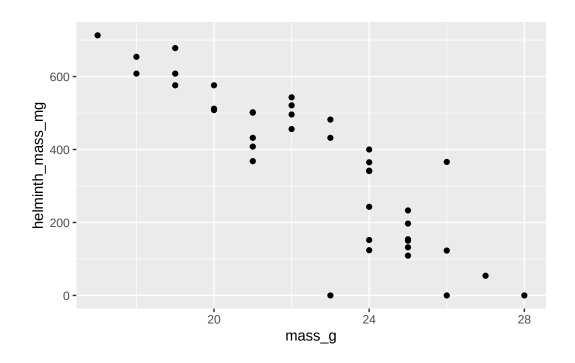
14.3.3 Two Variables: Both Continuous

If you are showing the relationship between two continuous variables, scatterplots with or without lines are usually the best way to go.

Let's try it out with the mammal data on body mass and helminth mass in mice:

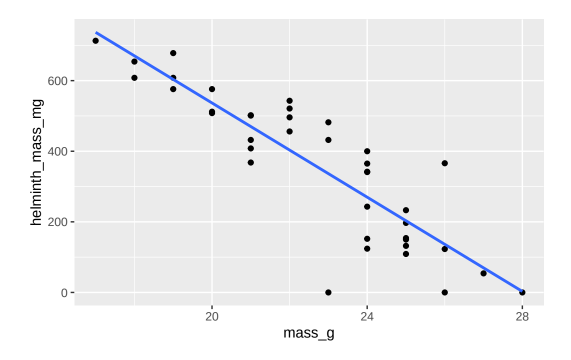
```
## filter for mouse data
mouse_data <- filter(fake_mammals, species %in% c("White-footed mouse", "Deer mouse"))

## create ggplot with your two continuous variables as x and y
ggplot(data = mouse_data, aes(x = mass_g, y = helminth_mass_mg)) +
    geom_point() ## create scatterplot</pre>
```



```
## with trendline
ggplot(data = mouse_data, aes(x = mass_g, y = helminth_mass_mg)) +
    geom_point() + ## create scatterplot
    ## create a trendline; method = "lm" makes it a straight line, se specifys whether there
    geom_smooth(method = "lm", se = FALSE)
```

`geom_smooth()` using formula = 'y ~ x'

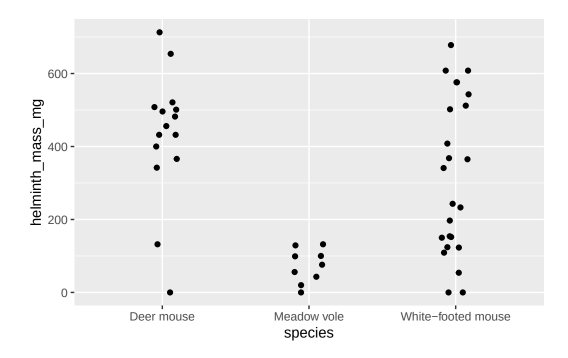


14.3.4 Two Variables: One Continuous, One Categorical

Believe it or not, when one of your variables is categorical, a scatterplot is still appropriate. Why not a bar plot? Because scatterplots show all of your data!

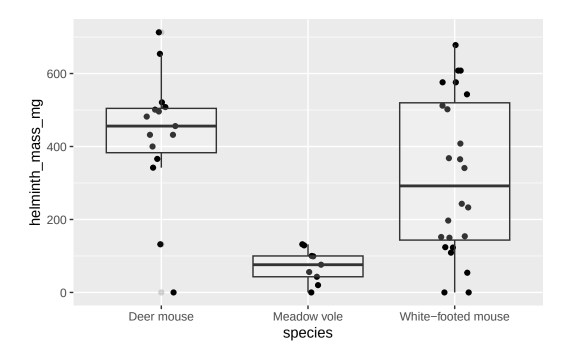
Let's demonstrate with the mammal data by comparing helminth mass among species.

```
## create ggplot with your two variables as x and y
ggplot(data = fake_mammals, aes(x = species, y = helminth_mass_mg)) +
geom_jitter(width = 0.1, height = 0) ## create points that are "jittered" a bit along the
```



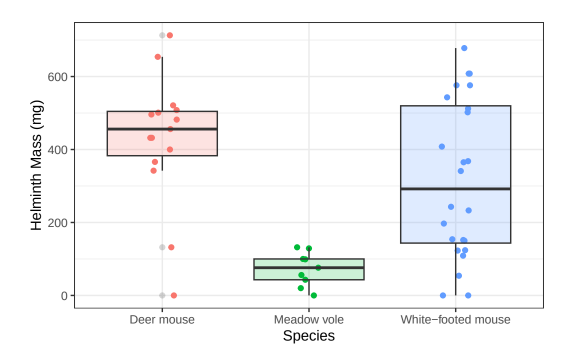
In this plot, we use geom_jitter to make the point spread a bit around each categorical X value so that you can see them better (but we specify height = 0 so as not to mess with the mass information). Instead of a mean helminth mass given by a bar plot, we can see the spread of each set of datapoints, including outliers or lack thereof. Still it's often nice to add some structure to these plots, which can be geom_boxplot or geom_violin (among others). Here is an example:

```
## create ggplot with your two variables as x and y
ggplot(data = fake_mammals, aes(x = species, y = helminth_mass_mg)) +
geom_jitter(width = 0.1, height = 0) + ## create points that are "jittered" a bit along
geom_boxplot(alpha = 0.2) ## create boxplot at 20% transparency with alpha
```



We could also make this plot even clearer by adding color:

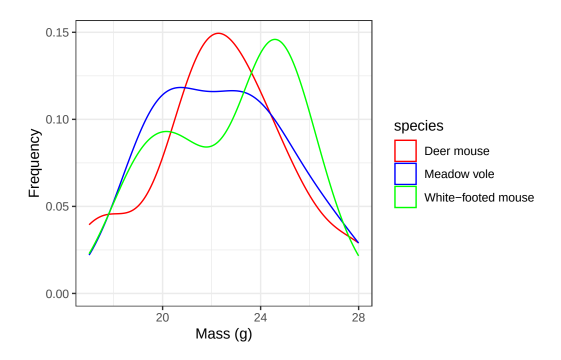
```
## create ggplot with your two variables as x and y
ggplot(data = fake_mammals, aes(x = species, y = helminth_mass_mg)) +
geom_jitter(aes(color = species), width = 0.1, height = 0) + ## you can put aes() inside
geom_boxplot(aes(fill = species), alpha = 0.2) +
labs(x = "Species", y = "Helminth Mass (mg)") +
theme_bw() +
theme(legend.position = "none") ## legend is redundant here, so we can hide it
```



14.3.5 Non-Axis Variables

You can also use other aesthetics to represent variables in your data. For example, you could use color to show the density plots of mammal masses among species. And you can modify the colors with scale functions:

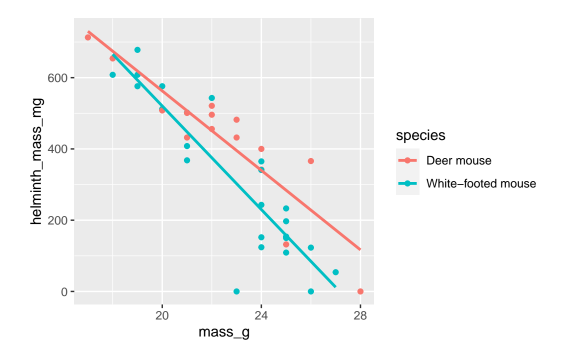
```
## make a ggplot, use wf_mice data, specify mass as the x variable and species as color
ggplot(data = fake_mammals, aes(x = mass_g, color = species)) +
   geom_density() + ## create density plot
   scale_color_manual(values = c("red", "blue", "green")) + ## set my own colors
   labs(x = "Mass (g)", y = "Frequency") + ## nicer labels
   theme_bw() ## my favorite simple theme
```



Similarly, you can add a third variable to a two variable figure. Take the helminth mass by mammal body mass figure from above:

```
ggplot(data = mouse_data, aes(x = mass_g, y = helminth_mass_mg, color = species)) +
   geom_point() + ## create scatterplot
   geom_smooth(method = "lm", se = FALSE)
```

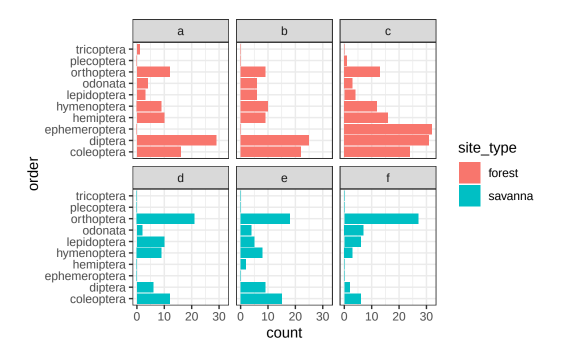
[`]geom_smooth()` using formula = 'y ~ x'



Color isn't the only way to show variables outside of axes, you can also use point shape, size, linetype, etc. In addition, you can split data among plot panels or "facets", with facet_wrap() or facet_grid().

Let's demonstrate with the long insect data, showing the insect communities for each site:

```
## specify order as y variable to show labels better
ggplot(data = long_insects, aes(y = order, x = count, fill = site_type)) +
    geom_col() +
    facet_wrap(vars(site), nrow = 2) + ## specify site variable, two rows to separate habita
    theme_bw()
```

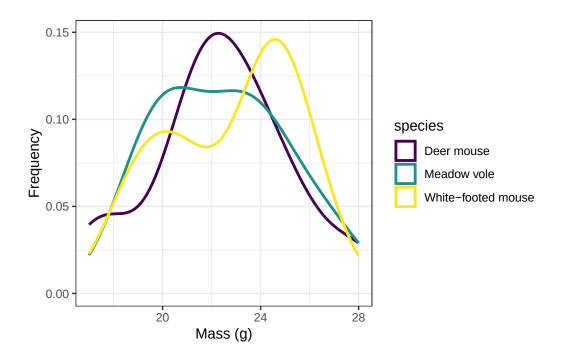


14.3.6 Colorblind Safe Colors

 ${\it ggplot 2}$ has color blind-safe color schemes available from the tidy verse-related package viridis.

Fore example:

```
## make a ggplot, use wf_mice data, specify mass as the x variable and species as color
ggplot(data = fake_mammals, aes(x = mass_g, color = species)) +
   geom_density(linewidth = 1) + ## create density plot, wider lines
   scale_color_viridis_d() + ## set viridis discrete colors
   labs(x = "Mass (g)", y = "Frequency") + ## nicer labels
   theme_bw() ## my favorite simple theme
```



See the following link for more info:

https://cran.r-project.org/web/packages/viridis/vignettes/intro-to-viridis.html

14.4 Further Reading

We have only scratched the surface of what ggplot2 can do! We barely discussed how to edit theme elements, nor did we spend much time on customizing scales.

ggplot2 has an excellent reference website which you can find here:

https://ggplot2.tidyverse.org/reference/index.html

With it you can learn all the ins and outs!

Part IV Appendices

15 Appendix A: More R Resources

Here is a collection of links to other useful resources for learning R!

Ecology-themed tutorial:

https://datacarpentry.org/R-ecology-lesson/index.html

Basic / Base R Materials

Official R manuals:

https://cran.r-project.org/manuals.html

Cookbook for R (features lots of "recipes" for common tasks)

http://www.cookbook-r.com/

Primers and Cheatsheets ("tidyverse"-based)

RStudio primers

https://rstudio.cloud/learn/primers

Tidyverse cheatsheets

https://www.rstudio.com/resources/cheatsheets/

Full Books and Courses

Hadley Wickham's "R for Data Science"

https://r4ds.had.co.nz/index.html

Hadley Wickham's "Advanced R"

https://adv-r.hadley.nz/

Book for "ggplot2" package

https://ggplot2-book.org/

Jenny Bryan's STAT 545 course

http://stat545.com/

Package Function Reference Sites

ggplot2 (data visualization)

https://ggplot2.tidyverse.org/index.html

sf (spatial analysis)

https://r-spatial.github.io/sf/index.html

Miscellaneous Resources

On Style:

http://adv-r.had.co.nz/Style.html

On Reproducibility:

https://reproducible-analysis-workshop.readthedocs.io/en/latest/

https://swcarpentry.github.io/r-novice-gapminder/

References