# Lesson 3 - Intro to Tidy Data: Go Long!

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Objective: At the end of this session you will know the principles of tidy data, a transform your data to perform simple calculations. You will be able to add new r data frame.  Highlighting				

# Packages Used in This Lesson

The following packages are used in this lesson:

tidyverse (tidyr, dplyr, tibble)

Please install and load these packages for the lesson. In this document I will load each package separately, but I will not be reminding you to install the package. Remember: these packages may be from CRAN OR Bioconductor.

# Data Files Used in This Lesson

- $-ENV\_pitlatrine.csv$
- -gapminder\_wide.csv
- $\hbox{-SPE\_pitlatrine.csv}$
- $\hbox{-fellowship.csv}$
- -towers.csv
- -returnking.csv
- -Female.csv
- -Male.csv



Figure 1:

These files can be downloaded at https://github.com/eacton/CAGEF/tree/master/Lesson\_3/data. Right-click on the filename and select 'Save Link As...' to save the file locally. The files should be saved in the same folder you plan on using for your R script for this lesson.

Or click on the blue hyperlink at the start of the README.md at https://github.com/eacton/CAGEF/tree/master/Lesson\_3 to download the entire folder at DownGit.

## Dataset: Sequencing of the V3-V5 hypervariable regions of the 16S rRNA gene

16S rRNA sequencing of 30 latrines from Tanzania and Vietnam at different depths (multiples of 20cm). Microbial abundance is represented in Operational Taxonomic Units (OTUs). Operational Taxonomic Units (OTUs) are groups of organisms defined by a specified level of DNA sequence similarity at a marker gene (e.g. 97% similarity at the V4 hypervariable region of the 16S rRNA gene). Intrinsic environmental factors such as pH, temperature, organic matter composition were also recorded.

We have 2 csv files:

- 1. A metadata file (Naming conventions: [Country\_LatrineNo\_Depth]) with sample names and environmental variables.
- 2. OTU abundance table.

B Torondel, JHJ Ensink, O Gunvirusdu, UZ Ijaz, J Parkhill, F Abdelahi, V-A Nguyen, S Sudgen, W Gibson, AW Walker, and C Quince. Assessment of the influence of intrinsic environmental and geographical factors on the bacterial ecology of pit latrines Microbial Biotechnology, 9(2):209-223, 2016. DOI:10.1111/1751-7915.12334

In this lesson we want to answer 3 simple questions:

- Which latrine depth has the greatest mean number of OTUs? (remember the Country\_LatrineNo\_Depth site encoding)
- Is there more Clostridia in Tanzania or Vietnam?
- Which site had the greatest number of Taxa represented?

Last lesson, we learned how to filter and select data subsets we were interested in. However, we can make data manipulation more efficient by controlling the overall structure or format of our data.



Let's read in our dataset, store it in a variable, and remind ourselves about the original structure.

```
library(readr)

dat <- read_csv("data/SPE_pitlatrine.csv")
head(dat)</pre>
```

```
## # A tibble: 6 x 82
     Taxa T_2_1 T_2_10 T_2_12 T_2_2 T_2_3 T_2_6 T_2_7 T_2_9 T_3_2 T_3_3 T_3_5
                          <dbl> <
##
     <chr> <dbl>
                   <dbl>
## 1 Acid~
               0
                       0
                              0
                                           0
                                                 0
                                                        0
                                                              0
## 2 Acid~
               0
                       0
                              0
                                                 0
                                                              0
                                                                     0
                                     0
                                           0
                                                        0
                                                                           0
                                                                                 0
## 3 Acid~
               0
                       0
                              0
                                     0
                                           0
                                                 0
## 4 Acid~
               0
                       0
                              0
                                                 0
                                                        0
                                                              0
                                                                     0
                                     0
                                           0
                                                                           0
                                                                                 0
## 5 Acid~
                       0
                              0
                                                 0
                                                              0
                                     0
                                           0
                                                                                 0
## 6 Acid~
               0
                       0
                              0
                                     2
                                           2
                                                 5
                                                        0
                                                                                 0
     ... with 70 more variables: T_4_3 < bl>, T_4_4 < bl>, T_4_5 < bl>,
       T_4_6 < db1>, T_4_7 < db1>, T_5_2 < db1>, T_5_3 < db1>, T_5_4 < db1>,
       T_5_5 <dbl>, T_6_2 <dbl>, T_6_5 <dbl>, T_6_7 <dbl>, T_6_8 <dbl>,
## #
       T_9_1 <dbl>, T_9_2 <dbl>, T_9_3 <dbl>, T_9_4 <dbl>, T_9_5 <dbl>,
## #
       V_1_2 <dbl>, V_10_1 <dbl>, V_11_1 <dbl>, V_11_2 <dbl>, V_11_3 <dbl>,
## #
       V_12_1 <dbl>, V_12_2 <dbl>, V_13_1 <dbl>, V_13_2 <dbl>, V_14_1 <dbl>,
## #
       V_14_2 <dbl>, V_14_3 <dbl>, V_15_1 <dbl>, V_15_2 <dbl>, V_15_3 <dbl>,
## #
## #
       V_16_1 <dbl>, V_16_2 <dbl>, V_17_1 <dbl>, V_17_2 <dbl>, V_18_1 <dbl>,
## #
       V_18_2 <dbl>, V_18_3 <dbl>, V_18_4 <dbl>, V_19_1 <dbl>, V_19_2 <dbl>,
       V_{19_3} < bl>, V_{2_1} < bl>, V_{2_2} < bl>, V_{2_3} < bl>, V_{20_1} < bl>,
## #
## #
       V_21_1 <dbl>, V_21_4 <dbl>, V_22_1 <dbl>, V_22_3 <dbl>, V_22_4 <dbl>,
       V 3 1 <dbl>, V 3 2 <dbl>, V 4 1 <dbl>, V 4 2 <dbl>, V 5 1 <dbl>,
       V_5_3 <dbl>, V_6_1 <dbl>, V_6_2 <dbl>, V_6_3 <dbl>, V_7_1 <dbl>,
## #
## #
       V_7_2 <dbl>, V_7_3 <dbl>, V_8_2 <dbl>, V_9_1 <dbl>, V_9_2 <dbl>,
## #
       V_9_3 <dbl>, V_9_4 <dbl>
```

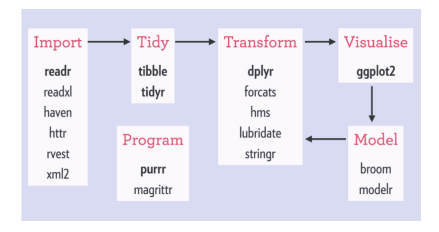


Figure 2:

# Intro to tidy data

Why tidy data?

Data cleaning (or dealing with 'messy' data) accounts for a huge chunk of data scientist's time. Ultimately, we want to get our data into a 'tidy' format where it is easy to manipulate, model and visualize. Having a consistent data structure and tools that work with that data structure can help this process along.

Tidy data has:

- 1. Each variable forms a column.
- 2. Each observation forms a row.
- 3. Each type of observational unit forms a table.

This seems pretty straight forward, and it is. It is the datasets you get that will not be straight forward. Having a map of where to take your data is helpful to unraveling its structure and getting it into a usable format.

The 5 most common problems with messy datasets are:

- common headers are values, not variable names
- multiple variables in one column
- variables stored in both rows and columns
- a single variable stored in multiple tables
- multiple types of observational units stored in the same table

Fortunately, Hadley has also given us the tools to solve these problems.

## Intro to the tidyverse

The tidyverse is the universe of packages created by Hadley Wickham for data analysis. There are packages to help import, tidy, transform, model and visualize data. His packages are pretty popular, so he made a package to load all of his packages at once. This wrapper package is tidyverse. In this lesson series we have used readr and readxl, and we will be using dplyr and tidyr today, and stringr and ggplot2 in future lessons. Install the package now.

install.packages("tidyverse")

Hadley has a large fan-base. Someone even made a plot of Hadley using his own package, ggplot2.

Back to the normal verse...

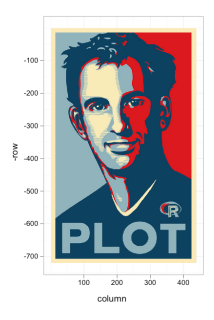


Figure 3:

#### Assessing our data frame

Which tidy data rules might our data frame break?

At first glance we can see that the column names are actually 3 different variables: 'Country', 'LatrineNumber', and 'Depth'. This information will likely be useful in our study, as we expect different bacteria at different depths, sites, and geographical locations. Each of these is a variable and should have its own separate column.

We could keep the column names as the sample names (as they are meaningful to the researcher) and add the extra variable columns, or we could make up sample names (ie. Sample\_1) knowing that the information is not being lost, but rather stored in a more useful format.

Some of the Taxa also appear to have an additional variable of information (ie. \_Gp1), but not all taxa have this information. We could also make a separate column for this information.

Each result is the same observational unit (ie. relative abundances of bacteria), so having one table is fine.

## Intro to tidyr

tidyr is a package with functions that help us turn our 'messy' data into 'tidy' data. It has 2 major workhorse functions and 2 other tidying functions:

- 1. gather() convert a data frame from wide to long format
- 2. spread() convert a data frame from long to wide format
- 3. separate() split a column into 2 or more columns based on a string separator
- 4. unite() merge 2 or more columns into 1 column using a string separator

gather() and spread() rely on key-value pairs to collapse or expand columns.

First let's load our library. Loading tidyverse will include the tidyr package that the gather() function is from.

library(tidyverse)

## -- Attaching packages

Note that 8 different packages are loaded, and that 2 functions from the stats package have been replaced by functions of the same name by dplyr. Note that you can still access the stats version of the function by calling it directly as stats::filter().

We can use the gather() function to collect our columns. This will make our dataset 'long' instead of 'wide'.

We need to provide gather() with information on our new columns. The first argument is our data frame, the second argument is the key, which is the name for the variable we want to gather (a set of column names). In this case our columns represent latrine sites, so we can assign the name 'Sites' to our key. The next argument is the value, which is the name for the measurements (usually numeric or integer) we have. In this case our values are Operational Taxonomic Units, so we can assign the name 'OTUs' to our value. The third argument is all of the columns that we want to gather. You can specify the columns by listing their names or positions.

```
dat %>% gather(key = Site, value = OTUs, T_2_1:V_9_4) %>% head()
```

```
## # A tibble: 6 x 3
##
     Taxa
                                OTUs
                         Site
##
     <chr>>
                         <chr> <dbl>
## 1 Acidobacteria Gp1 T 2 1
                                   0
## 2 Acidobacteria Gp10 T 2 1
## 3 Acidobacteria_Gp14 T_2_1
                                   0
## 4 Acidobacteria Gp16 T 2 1
                                   0
## 5 Acidobacteria_Gp17 T_2_1
                                   0
## 6 Acidobacteria_Gp18 T_2_1
#equivalent to
dat %>% gather(key = Site, value = OTUs, 2:82) %>% head()
#equivalent to
dat %>% gather(Site, OTUs, -1) %>% head()
```

In the above examples "-" means gather every column except the 1st, or gather every column except Taxa. Taxa would still be retained as a column but its elements are not grouped in with 'Sites' (ie. we do not want 'V\_9\_4', 'T\_2\_9', and 'Clostridia' gathered into the same column).

Let's save the last variation into a data frame called gather dat.

dat %>% gather(Site, OTUs, -Taxa) %>% head()

#equivalent to

```
gather_dat <- dat %>% gather(Site, OTUs, -Taxa)
```

Note how the dimensions of your dataframe have changed relative to dat. Instead of 52 rows and 82 columns, we now have a data frame with 4212 rows and 3 columns (which is the 81 columns we gathered x 52 rows). gather\_dat is now in a long format instead of wide.

Next, we can use the separate() function to get the Country, Latrine\_Number, and Depth information from our Site column. separate() takes in your dataframe, the name of the column to be split, the names of your new columns, and the character that you want to split the columns by (in this case an underscore). Note that the default is to remove your original column - if you want to keep it, you can add the additional argument remove = FALSE, keeping in mind that you now have redundant data.

```
split_dat <- gather_dat %>%
   separate(Site, c("Country", "Latrine_Number", "Depth"), sep = "_")
## #equivalent to
## split_dat <- gather_dat %>%
## separate(Site, c("Country", "Latrine_Number", "Depth"), sep = "_", remove = TRUE)
```

We may also want to do this for the 'Group' of Acidobacteria. Try the code, but do not save the answer in a variable.

```
split_dat %>% separate(Taxa, c("Taxa", "Group"), sep = "_Gp") %>% head(20)
## Warning: Expected 2 pieces. Missing pieces filled with `NA` in 3078 rows
## [15, 16, 17, 18, 19, 20, 21, 22, 23, 24, 25, 26, 27, 28, 29, 30, 31, 32,
## 33, 34, ...].
## # A tibble: 20 x 6
##
      Taxa
                            Group Country Latrine_Number Depth OTUs
##
      <chr>
                            <chr> <chr>
                                           <chr>
                                                           <chr> <dbl>
##
    1 Acidobacteria
                            1
                                  Τ
                                           2
                                                                      0
                                  Т
                                           2
                                                                      0
##
                            10
    2 Acidobacteria
                                                           1
    3 Acidobacteria
                            14
                                  Τ
                                           2
                                                           1
                                                                      0
##
   4 Acidobacteria
                            16
                                  T
                                           2
                                                           1
                                                                      0
                                  Τ
                                           2
    5 Acidobacteria
                            17
                                                           1
                                                                      0
                                  Т
                                           2
    6 Acidobacteria
                            18
                                                                      0
##
                                                           1
                            21
                                  Т
                                           2
##
    7 Acidobacteria
                                                           1
                                                                      0
                            22
                                  Т
                                           2
                                                                      0
##
  8 Acidobacteria
                                                           1
  9 Acidobacteria
                            3
                                  Т
                                           2
                                                           1
                                                                      0
                            4
                                  Τ
                                           2
                                                                      0
## 10 Acidobacteria
                                                           1
## 11 Acidobacteria
                            5
                                  Т
                                           2
                                                           1
                                                                      0
                            6
                                  Τ
                                           2
## 12 Acidobacteria
                                                           1
                                                                      0
## 13 Acidobacteria
                            7
                                  Т
                                           2
                                                                      0
                                                           1
## 14 Acidobacteria
                            9
                                  Τ
                                           2
                                                                      0
## 15 Actinobacteria
                            <NA>
                                  Τ
                                           2
                                                           1
                                                                    110
                                           2
## 16 Alphaproteobacteria <NA>
                                  Τ
                                                           1
                                                                     11
## 17 Anaerolineae
                                           2
                                                                      2
                            <NA>
                                  Τ
                                                           1
                                           2
## 18 Bacilli
                            <NA>
                                  Τ
                                                           1
                                                                     19
                                           2
## 19 Bacteroidia
                            <NA>
                                  Τ
                                                           1
                                                                   1547
## 20 Betaproteobacteria
                            <NA>
                                           2
```

We get a warning from R that it has filled in 'NA' for the bacteria that did not have groups. Note that I chose to split Taxa using '\_Gp' since I did not need 'Gp'.

#### Challenge

Use the glimpse() function to look at the type of each variable in our new data frame, split\_dat. Are those the types you expected? Why or why not? How is glimpse() different from the str() function?

There is a useful function <code>group\_by()</code> that you can use to group variables or sets of variables together. This is useful for calculations and plotting on subsets of your data without having to turn your variables into factors. Say I wanted to look at a combination of Country and Well Depth. While visually, you wouldn't notice any changes to your data frame, if you look at the structure it will now be a 'grouped\_df'. There are 15 groupings resulting from Country and Depth. After I have performed my desired operation, I can return my data frame to its original structure by calling <code>ungroup()</code>. First we will examine the structure of grouped and ungrouped output without any additional operations.

```
str(split_dat)
## Classes 'tbl_df', 'tbl' and 'data.frame':
                                                4212 obs. of 5 variables:
                           "Acidobacteria_Gp1" "Acidobacteria_Gp10" "Acidobacteria_Gp14" "Acidobacteria
                    : chr
                           "T" "T" "T" "T" ...
   $ Country
                    : chr
                           "2" "2" "2" "2" ...
   $ Latrine_Number: chr
##
                           "1" "1" "1" "1" ...
   $ Depth
                    : chr
   $ OTUs
                    : num
                          0 0 0 0 0 0 0 0 0 0
str(split_dat %>% group_by(Country, Depth))
## Classes 'grouped_df', 'tbl_df', 'tbl' and 'data.frame': 4212 obs. of 5 variables:
##
   $ Taxa
                    : chr
                           "Acidobacteria_Gp1" "Acidobacteria_Gp10" "Acidobacteria_Gp14" "Acidobacteria
                           "T" "T" "T" "T" ...
##
   $ Country
                    : chr
   $ Latrine_Number: chr
                           "2" "2" "2" "2" ...
                           "1" "1" "1" "1" ...
##
   $ Depth
                    : chr
                    : num 0000000000...
##
   $ OTUs
                           "Country" "Depth"
   - attr(*, "vars")= chr
   - attr(*, "drop")= logi TRUE
   - attr(*, "indices")=List of 15
##
##
     ..$: int 0123456789 ...
##
     ..$ : int 52 53 54 55 56 57 58 59 60 61 ...
##
     ..$ : int 104 105 106 107 108 109 110 111 112 113 ...
     ..$ : int 156 157 158 159 160 161 162 163 164 165 ...
##
     ..$ : int 208 209 210 211 212 213 214 215 216 217 ...
##
     ..$ : int 624 625 626 627 628 629 630 631 632 633 ...
##
     ..$ : int 520 521 522 523 524 525 526 527 528 529 ...
     ..$ : int 260 261 262 263 264 265 266 267 268 269 ...
##
##
     ..$ : int 312 313 314 315 316 317 318 319 320 321 ...
     ..$: int 1196 1197 1198 1199 1200 1201 1202 1203 1204 1205 ...
##
     ..$ : int 364 365 366 367 368 369 370 371 372 373 ...
##
##
     ..$: int 1560 1561 1562 1563 1564 1565 1566 1567 1568 1569 ...
##
     ..$: int 1508 1509 1510 1511 1512 1513 1514 1515 1516 1517 ...
##
     ..$ : int 1716 1717 1718 1719 1720 1721 1722 1723 1724 1725 ...
     ..$: int 2652 2653 2654 2655 2656 2657 2658 2659 2660 2661 ...
##
##
   - attr(*, "group_sizes")= int 104 52 52 260 260 156 260 104 156 52 ...
   - attr(*, "biggest_group_size")= int 1040
                                       15 obs. of 2 variables:
   - attr(*, "labels")='data.frame':
     ..$ Country: chr "T" "T" "T" "T" ...
##
     ..$ Depth : chr "1" "10" "12" "2" ...
##
     ..- attr(*, "vars")= chr "Country" "Depth"
     ..- attr(*, "drop")= logi TRUE
str(split_dat %>% group_by(Country, Depth) %>% ungroup())
## Classes 'tbl_df', 'tbl' and 'data.frame':
                                                4212 obs. of 5 variables:
##
   $ Taxa
                           "Acidobacteria_Gp1" "Acidobacteria_Gp10" "Acidobacteria_Gp14" "Acidobacteria
                    : chr
                           "T" "T" "T" "T" ...
   $ Country
                    : chr
                           "2" "2" "2" "2"
   $ Latrine_Number: chr
                           "1" "1" "1" "1" ...
##
   $ Depth
                    : chr
   $ OTUs
                    : num
                          0 0 0 0 0 0 0 0 0 0 ...
```

Now we can see an example of how grouped\_by in action with summarize(), can easily calculate summary statistics for groups of data. Whereas in our messy data frame it was difficult to do calculations based on Country, Well Number or Latrine Depth, this is now an easy task. Let's get the mean, median, standard deviation and maximum for the number of OTUs collected in Tanzania vs Vietnam.

```
split_dat %>%
  group_by(Country) %>%
  summarize(mean = mean(OTUs), median = median(OTUs), sd = sd(OTUs), maximum = max(OTUs))
## # A tibble: 2 x 5
##
     Country mean median
                             sd maximum
##
     <chr>>
             <dbl> <dbl> <dbl>
                                   <dbl>
## 1 T
              122.
                        0 896.
                                   17471
## 2 V
              186.
                           863.
                                   17572
                        0
```

In dealing with grouped data, we no longer have to grab a Country by subsetting or using helper functions to grab letters from their names. Group by recognizes that we have 2 countries and will perform calculations for both of them.

Now that we have tidy data, let's answer our questions:

- Which latrine depth has the greatest mean number of OTUs? (remember the Country\_LatrineNo\_Depth site encoding)
- Is there more Clostridia in Tanzania or Vietnam?
- Which site had the greatest number of Taxa represented?

## Okay, Go!

Which latrine depth has the greatest mean number of OTUs?

```
split_dat %>%
  group_by(Latrine_Number) %>%
  summarize(mean = mean(OTUs)) %>%
  arrange(desc(mean))
```

```
## # A tibble: 22 x 2
##
      Latrine_Number mean
##
      <chr>
                     <dbl>
##
   1 17
                      415.
##
  2 16
                      403.
##
  3 11
                      338.
## 4 20
                      335.
## 5 10
                      328.
##
  6 8
                      236.
##
  7 15
                      207.
## 8 3
                      206.
## 9 18
                      201.
## 10 12
## # ... with 12 more rows
```

Without data being in 'tidy' format - with all variables that were mashed together into a Site (Country\_LatrineNo\_Depth) having their own columns - this is a difficult question to answer. Once Latrine\_Number is a variable, we can simply group our data to perform our mean calculation and get an answer.

Is there more Clostridia in Tanzania or Vietnam?

```
split_dat %>%
  filter(Taxa == "Clostridia") %>%
  group_by(Country) %>%
  summarise(sum = sum(OTUs), mean = mean(OTUs)) %>%
  arrange(desc(sum))
```



Figure 4:

```
## # A tibble: 2 x 3
## Country sum mean
## <chr> <dbl> <dbl> <dbl> ## 1 V 176079 3386.
## 2 T 101217 3490.
```

Again, being able to filter by Taxa and group by Country (as an isolated variable) helps a lot. With dplyr syntax we can perform all data manipulations and calculations in a code block that is readable.

Which site had the greatest number of Taxa represented?

```
split_dat %>%
  group_by(Country, Latrine_Number, Depth) %>%
  filter(OTUs != 0) %>%
  summarize(count = n()) %>%
  arrange(desc(count))
```

```
## # A tibble: 81 x 4
## # Groups:
                Country, Latrine_Number [28]
##
      Country Latrine_Number Depth count
##
      <chr>
               <chr>>
                                 <chr> <int>
    1 V
##
               7
                                 3
                                           36
##
    2 V
               22
                                 1
                                           31
    3 V
               7
                                 2
                                           29
##
               2
                                 6
                                           27
##
    4 T
##
    5 V
                                 1
                                           27
               11
    6 V
                                3
                                           27
##
               11
                                9
    7 T
               2
                                           26
##
                                 2
                                           26
##
    8 V
               15
##
    9 V
               15
                                 3
                                           26
## 10 V
               18
                                 4
                                           26
## # ... with 71 more rows
```

Since we can group by the 3 variables that were in the Site name, there is no disadvantage to having our data in tidy format compared to our original wide data frame. However now we are able to filter for non-zero OTUs, which was impossible in the wide format. Since we know from earlier in the lesson that each Taxa is only represented once for each site, we only have to count and order the number of observations to get our answer.

To get data back into its original format, there are reciprocal functions in the tidyr package, making it possible to switch between wide and long formats.

Fair question: But you've just been telling me how great the 'long' format is?!?! Why would I want the wide format again???

**Honest answer:** Note that our original data frame was 52 rows and expanded to 4212 rows in the long format. When you have, say, a genomics dataset you might end up with 6,000 rows expanding to 600,000 rows. You probably want to do your calculations and switch back to the more 'human readable' format. Sure, I can save a data frame with 600,000 rows, but I can't really send it to anyone because LibreOffice or Excel might crash opening it.

# Challenge

Collapse Country, Latrine\_Number and Depth back into one variable, 'Site', using the unite() function. Store the output in a data frame called unite\_dat.

# Challenge

Use the **spread()** function to turn unite\_dat into the wide shape of our original dataset. Save the output into a data frame called 'spread\_dat'.

## Challenge

Read in the gapminder\_wide.csv. What rules of tidy data does it break? Transform the dataset to the format below. How many rows do you have?

country	continent	year	lifeExp	pop	gdpPercap
Afghanistan	Asia	1952	28.801	8425333	779.4453
Afghanistan	Asia	1957	30.332	9240934	820.8530
Afghanistan	Asia	1962	31.997	10267083	853.1007
Afghanistan	Asia	1967	34.020	11537966	836.1971
Afghanistan	Asia	1972	36.088	13079460	739.9811
Afghanistan	Asia	1977	38.438	14880372	786.1134

# Adding rows and columns to data frames

We are going to use a subset of the collected environmental data from the study to practice adding rows and columns to a data frame. We can use our pipe while reading in our data to do this subsetting.

```
jdat <- read_csv("data/ENV_pitlatrine.csv") %>% select(1:4)
```

```
## Parsed with column specification:
## cols(
## Samples = col_character(),
## pH = col_double(),
## Temp = col_double(),
## VS = col_double(),
## VFA = col_double(),
```

```
##
     CODt = col double(),
     CODs = col_double(),
##
##
     perCODsbyt = col double(),
     NH4 = col_double(),
##
##
     Prot = col_double(),
     Carbo = col double()
##
## )
glimpse(jdat)
## Observations: 81
## Variables: 4
## $ Samples <chr> "T 2 1", "T 2 10", "T 2 12", "T 2 2", "T 2 3", "T 2 6"...
             <dbl> 7.82, 9.08, 8.84, 6.49, 6.46, 7.69, 7.48, 7.60, 7.55, ...
## $ pH
             <dbl> 25.1, 24.2, 25.1, 29.6, 27.9, 28.7, 29.8, 25.0, 28.8, ...
## $ Temp
## $ TS
             <dbl> 14.53, 37.76, 71.11, 13.91, 29.45, 65.52, 36.03, 46.87...
```

There is a function from dplyr, bind\_rows(), which tries to save you from human error when joining a row of observations to a data frame by ensuring that your column names match (so data is not entered in the wrong column). If you do not give column names, an error will be thrown.

What data structure should you be using to add a row to a data frame?

If you give bind\_rows() the correct column information in vector format, it will only add a vector as a row IF THE CHARACTER TYPE MATCHES ALL DATA FRAME COLUMN TYPES (ie. if our data frame was all character data). Otherwise, it actually gives you the useful error that your character types are not matching.

```
bind_rows(jdat, c(Samples = "V_2_10", pH = 6, Temp = 5, TS = 5)) %>% tail()
```

## Error in bind\_rows\_(x, .id): Column `pH` can't be converted from numeric to character

A list allows us to add the data we can't add in vector format, by retaining the data types that match the data frame columns.

```
bind_rows(jdat, list(Samples = "V_2_10", pH = 6, Temp = 5, TS = 5)) %>% tail()
```

```
## # A tibble: 6 x 4
##
     Samples
                pH Temp
     <chr>
             <dbl> <dbl> <dbl>
##
## 1 V 8 2
                    19.5 36.8
              9.3
## 2 V 9 1
              7.3
                    18.8 31.3
## 3 V 9 2
              4.04
                    20.5
## 4 V_9_3
              4.36
                    19.9
                          92.6
## 5 V_9_4
              4.3
                    19.8
                          36.5
## 6 V_2_10
                           5
              6
                     5
```

If you give bind\_rows() the correct column information in list format, it will add your row.

A sanity check is making sure things are turning out as you expect them to - it is a way of checking your (or others') assumptions. It is particularly useful in the data exploration stage when you are getting familiar with your data set before you try to run any type of model. However, it is also good to make sure a function is behaving as you expect it to, or trouble-shooting odd behaviours

Here is a sanity check on switching the order of the columns.

```
bind_rows(jdat, list(Samples = "V_2_10", Temp = 5, pH = 6, TS = 5)) %>% tail()
## # A tibble: 6 x 4
## Samples pH Temp TS
```

```
##
     <chr>>
              <dbl> <dbl> <dbl>
## 1 V_8_2
               9.3
                     19.5
                            36.8
                            31.3
## 2 V 9 1
               7.3
                     18.8
                     20.5
## 3 V_9_2
               4.04
                            32
## 4 V_9_3
               4.36
                     19.9
                            92.6
## 5 V 9 4
                            36.5
               4.3
                     19.8
## 6 V 2 10
               6
                       5
                             5
```

Since we have named our values, bind\_rows() will add the value to the appropriate column.

What happens if column name do not match? bind\_rows() will match your columns as much as possible, and then create new columns for the data that does not fit at the end of your data frame. Note that 'NA's will be created for all missing data.

```
bind_rows(jdat, list(Turtles = "V_2_10", pH = 5, Volatility = 5, TS = 5)) %>% tail()
```

```
## # A tibble: 6 x 6
##
     Samples
                 рΗ
                    Temp
                              TS Turtles Volatility
##
     <chr>>
              <dbl> <dbl> <dbl> <chr>
                                               <dbl>
## 1 V_8_2
                            36.8 <NA>
                                                   NA
               9.3
                     19.5
## 2 V_9_1
               7.3
                     18.8
                            31.3 <NA>
                                                   NA
## 3 V_9_2
               4.04
                     20.5
                            32
                                 <NA>
                                                  NA
## 4 V_9_3
               4.36
                     19.9
                            92.6 <NA>
                                                  NA
## 5 V_9_4
               4.3
                     19.8
                            36.5 <NA>
                                                   NA
## 6 <NA>
               5
                     NA
                             5
                                 V_2_10
                                                    5
```

You can also bind data frames together, which is another common use of this function.

```
bind_rows(jdat, data.frame(Samples = "V_2_10", pH = 5, Temp = 5, TS = 5, stringsAsFactors = F)) %>%
tail()
```

```
## # A tibble: 6 x 4
##
     Samples
                 pH Temp
                              TS
     <chr>>
             <dbl> <dbl>
##
                          <dbl>
## 1 V 8 2
               9.3
                     19.5
                           36.8
## 2 V_9_1
               7.3
                     18.8
                           31.3
## 3 V 9 2
                     20.5
               4.04
## 4 V_9_3
               4.36
                     19.9
                           92.6
## 5 V 9 4
                     19.8
                           36.5
               4.3
                             5
## 6 V_2_10
                      5
               5
```

Why doesn't jdat change after all these operations?

Whenever you are adding rows or columns, I strongly advise checking to see that the dimensions of the resulting data frame are what you expect.

```
jdat <- bind_rows(jdat, list(Samples = "V_2_10", pH = 6, Temp = 5, TS = 5))
dim(jdat)
## [1] 82 4</pre>
```

#### Challenge

Make a column and add it to jdat using the dplyr function bind\_cols(). Do you forsee any problems in using this function?

#### Challenge

Given 5 data frames with data on the number of words spoken in the Lord of the Rings trilogy (datacarpentry archive, tidy-data), answer the following questions:

- 1. How many characters are present in all 3 films?
- 2. Which race speaks the most? Is this true across all 3 films?
- 3. Which character is chattiest?
- 4. Which gender speaks the most for Elvish characters? Does this hold true across all 3 films?
- 5. Which Characters are these elves likely to be?

# All of the joins

Often we have more than one data table that shares a common attribute. For example, with our current dataset, we have other variables (such as pH) for a sample, as well as our OTU table, both of which have site IDs (ie. T\_2\_9). We want to merge these into one table.

Joins can be tricky, so we are going to use a subset of our tidy data such that we can easily observe the output of our join.

Joins use a 'key' by which we can match up our data frames. When we look at our data frames we can see that they have matching information in 'Samples'. We are going to reduce our dataset by only keeping non-zero values so we can see how the join functions work a bit more easily.

#### Challenge

Filter unite\_dat to remove all non-zero values, and store it in an object ndat. How many rows in unite\_dat had the value of zero? Sort ndat to keep the top 20 rows with the highest OTUs.

Answer:

```
ndat <- unite_dat %>% filter(OTUs!=0) %>% arrange(desc(OTUs)) %>% .[1:20, ]
```

We will remove some observations from jdat, just so our key columns don't match perfectly.

```
jdat <- jdat[-c(4,7,9,12,20,22) , 1:4]
```

As a starting point for the next exercise ndat has 20 rows and 3 columns, and jdat has 76 rows and 4 columns. There are 2 types of joins:

- 1. *mutating joins* uses a key to match observations and combines variables from 2 tables (adding columns and potentially rows)
- 2. filtering joins uses a key to match observations, to subset observations (rows)

In these examples we are joining 2 dataframes: x and y.

**Inner join** is a mutating join. Inner join returns all rows from x where there are matching values in y, and all columns from x and y. If there are multiple matches between x and y, all combination of the matches are returned.

A set of graphics from 'R for Data Science' makes the description clearer:

```
inner_join(ndat, jdat, by = c("Site" = "Samples"))
```

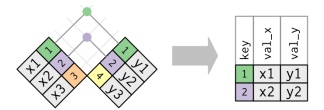
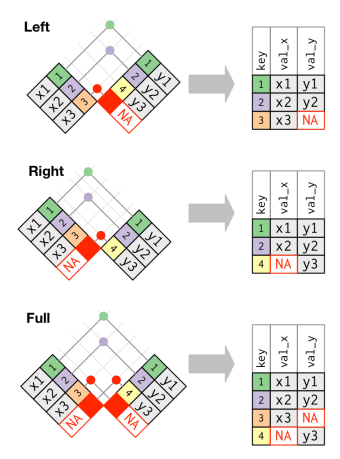


Figure 5: img borrowed from r4ds.had.co.nz

```
## # A tibble: 18 x 6
##
      Taxa
                            Site
                                    0TUs
                                             рН
                                                 Temp
                                                          TS
##
      <chr>
                                   <dbl>
                            <chr>
                                          <dbl>
                                                <dbl>
                                                       <dbl>
##
    1 Clostridia
                            V_16_2 17572
                                           6.26
                                                 16.3
                                                        24
                                           7.6
                                                 25
##
    2 Clostridia
                            T_2_9 17471
                                                        46.9
##
    3 Clostridia
                            V_17_2 13875
                                           8.14
                                                 15.4
                                                        25.6
    4 Clostridia
                                           7.69
                                                 28.7
##
                            T_2_6 12169
                                                        65.5
##
    5 Bacteroidia
                            V_17_1 11392
                                           7.39
                                                 15.7
                                                        25.3
##
    6 Clostridia
                            V_10_1 11180
                                          9.12
                                                 18.6
                                                        46.3
                                           6.46
                                                 27.9
##
    7 Clostridia
                            T_2_3
                                   10944
                                                        29.4
##
    8 Unknown
                            T_9_2
                                   10624
                                           7.86
                                                 28.5
                                                        13.6
##
    9 Clostridia
                            T 4 4
                                   10545
                                           7.84
                                                 26.3
                                                        28.8
## 10 Clostridia
                            V_16_1 10043
                                          7.83
                                                 17.7
                                                        25.1
   11 Clostridia
                            T_5_3
                                    8981
                                           7.53
                                                 27.5
                                                        12.6
## 12 Alphaproteobacteria V_3_2
                                                 20.7
                                                        52
                                    8660
                                           8.08
                            V_15_1
## 13 Clostridia
                                    8244
                                           7.44
                                                 18.9
                                                        25.4
## 14 Clostridia
                                          7.39
                                                 15.7
                                                        25.3
                            V_17_1
                                    7722
  15 Clostridia
                            V_18_1
                                    7481
                                           7.09
                                                 22.5
                                                        35.1
  16 Alphaproteobacteria
                                                 22.4
                           V_3_1
                                    7322
                                           8.9
                                                        56.2
  17 Clostridia
                                                        29.9
                            V_18_4
                                    6246
                                           8.25
                                                 19.4
## 18 Clostridia
                            T_2_1
                                    6213
                                          7.82
                                                 25.1
                                                        14.5
```

We have 6 columns in the resulting data frame - all columns minus the 'Samples' column which was a duplicated id column (equal to ndat's 'Site' column). We can see that there are 18 rows in the resulting data frame. Rows from ndat that did not have a matching site in jdat were removed.

Outer joins are a set of mutating joins. There are 3 outer joins: left, right, and full.



**Left join** returns all rows from x, and all columns from x and y. Rows in x with no match in y will have NA values in the new columns. If there are multiple matches between x and y, all combinations of the matches are returned.

```
left_join(ndat, jdat, by = c("Site" = "Samples"))
```

```
## # A tibble: 20 x 6
##
      Taxa
                           Site
                                    OTUs
                                             ηН
                                                 Temp
                                                         TS
##
      <chr>
                           <chr>
                                   <dbl> <dbl> <dbl> <dbl> <
##
    1 Clostridia
                           V_16_2 17572
                                          6.26
                                                 16.3
                                                       24
                                                 25
##
    2 Clostridia
                           T_2_9 17471
                                          7.6
                                                       46.9
##
    3 Clostridia
                           V 17 2 13875
                                          8.14
                                                 15.4
                                                       25.6
    4 Clostridia
                           T_2_6 12169
                                          7.69
                                                 28.7
                                                       65.5
##
##
    5 Bacteroidia
                           V 17 1 11392
                                          7.39
                                                 15.7
                                                       25.3
                           V_10_1 11180
##
    6 Clostridia
                                          9.12
                                                 18.6
                                                       46.3
    7 Clostridia
                           T 2 3
                                  10944
                                          6.46
                                                 27.9
                                                       29.4
##
                           T_9_2
                                                 28.5
##
    8 Unknown
                                  10624
                                          7.86
                                                       13.6
   9 Clostridia
                                   10545
                                          7.84
                                                 26.3
                                                       28.8
##
                           T 4 4
                                          7.83
## 10 Clostridia
                           V_16_1 10043
                                                 17.7
                                                       25.1
  11 Clostridia
                           T_2_2
                                    8999 NA
                                                 NA
                                                       NA
## 12 Clostridia
                           T_5_3
                                                 27.5
                                    8981
                                          7.53
                                                       12.6
## 13 Alphaproteobacteria V_3_2
                                    8660
                                          8.08
                                                 20.7
                                                       52
## 14 Clostridia
                           V_{15_{1}
                                    8244
                                          7.44
                                                 18.9
                                                       25.4
## 15 Clostridia
                           V_17_1
                                    7722
                                          7.39
                                                 15.7
                                                       25.3
## 16 Betaproteobacteria
                           T_4_3
                                    7710 NA
                                                 NA
                                                       NA
## 17 Clostridia
                           V_18_1
                                    7481
                                          7.09
                                                 22.5
                                                       35.1
## 18 Alphaproteobacteria V_3_1
                                    7322
                                          8.9
                                                 22.4
                                                       56.2
```

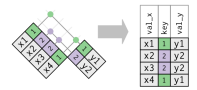


Figure 6: img borrowed from r4ds.had.co.nz

```
## 19 Clostridia V_18_4 6246 8.25 19.4 29.9
## 20 Clostridia T_2_1 6213 7.82 25.1 14.5
```

That means that we will have our 20 rows from ndat and 6 columns; any Sites that weren't found in jdat (in this case T 2 2 and T 4 3) will be filled with NA.

**Right join** returns all rows from y, and all columns from x and y. Rows in y with no match in x will have NA values in the new columns. If there are multiple matches between x and y, all combinations of the matches are returned.

```
right_join(ndat, jdat, by = c("Site" = "Samples"))
```

```
# A tibble: 77 x 6
##
##
      Taxa
                  Site
                           0TUs
                                        Temp
                                                 TS
                                    рΗ
##
       <chr>
                   <chr>
                          <dbl> <dbl>
                                       <dbl>
                                              <dbl>
##
    1 Clostridia T_2_1
                           6213
                                  7.82
                                         25.1
                                               14.5
##
    2 <NA>
                  T 2 10
                             NA
                                  9.08
                                         24.2
                                               37.8
##
    3 <NA>
                  T 2 12
                             NA
                                  8.84
                                         25.1
                                               71.1
    4 Clostridia T_2_3
##
                          10944
                                  6.46
                                         27.9
                                               29.4
    5 Clostridia T_2_6
                          12169
                                  7.69
                                         28.7
                                               65.5
##
##
    6 Clostridia T 2 9
                          17471
                                  7.6
                                               46.9
##
    7 <NA>
                  T_3_3
                             NA
                                  7.68
                                         28.9
                                               14.6
##
    8 <NA>
                  T_3_5
                             NA
                                  7.69
                                         28.7
                                               14.9
##
    9 Clostridia T_4_4
                          10545
                                  7.84
                                         26.3
                                               28.8
## 10 <NA>
                  T_{4}_{5}
                             NA
                                  7.95
                                         27.9
                                               46.8
## # ... with 67 more rows
```

That means that we will have our 76 rows from jdat, and any items that weren't found in ndat will be filled with NA. Since there are 77 rows in the final data frame, this means there must have been multiple rows matching from ndat. In other words, we had a duplicate key (Site) in ndat. Again, we have 6 columns.

Let's try to find which Site was duplicated by using n().

```
right_join(ndat, jdat, by = c("Site" = "Samples")) %>%
  group_by(Site) %>%
  filter(n()>1)
## # A tibble: 2 x 6
## # Groups:
               Site [1]
##
     Taxa
                 Site
                          OTUs
                                               TS
                                  рН
                                      Temp
     <chr>
                         <dbl>
##
                  <chr>
                               <dbl>
                                     <dbl>
                                           <dbl>
## 1 Bacteroidia V_17_1 11392
                                             25.3
                                7.39
                                      15.7
## 2 Clostridia V_17_1 7722
                               7.39
                                      15.7
                                            25.3
```

Note that Bacteriodia and Clostridia have different OTUs (from ndat), but the same pH, Temp and TS (from jdat).

**Full join** returns all rows and all columns from both x and y. Where there are not matching values, returns NA for the one missing.

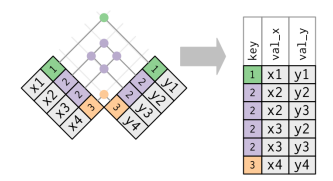


Figure 7: img borrowed from r4ds.had.co.nz

```
full_join(ndat, jdat, by = c("Site" = "Samples"))
## # A tibble: 79 x 6
##
      Taxa
                           0TUs
                                    ηЧ
                                                TS
                                        Temp
##
      <chr>
                   <chr>
                          <dbl> <dbl>
                                       <dbl> <dbl>
                                 6.26
                                        16.3
                                              24
##
    1 Clostridia
                  V_16_2 17572
##
    2 Clostridia
                  T_2_9
                          17471
                                 7.6
                                        25
                                              46.9
##
    3 Clostridia
                  V 17 2 13875
                                 8.14
                                        15.4
                                              25.6
    4 Clostridia T_2_6 12169
                                 7.69
                                        28.7
##
                                              65.5
    5 Bacteroidia V 17 1 11392
                                 7.39
                                        15.7
    6 Clostridia
                  V_10_1 11180
                                        18.6
##
                                 9.12
                                              46.3
##
    7 Clostridia
                  T_2_3
                          10944
                                 6.46
                                        27.9
                                              29.4
    8 Unknown
                   T 9 2
                          10624
                                 7.86
                                        28.5
                                              13.6
##
    9 Clostridia
                          10545
                                 7.84
                                        26.3
                                              28.8
                  T 4 4
                  V_16_1 10043
                                              25.1
## 10 Clostridia
                                 7.83
                                        17.7
## # ... with 69 more rows
```

The full join returns all of the rows from both ndat and jdat as well as the 6 columns - we have the 75 rows from jdat, plus the second V\_17\_1 as seen in the right\_join and T\_2\_2 and T\_4\_3 that we present in ndat but NA in jdat for a total of 79 rows.

Lastly, we have the 2 filtering joins. These will not add columns, but rather filter the rows based on what is present in the second data frame.

**Semi join** returns all rows from x where there are matching values in y, keeping just columns from x. A semi join differs from an inner join because an inner join will return one row of x for each matching row of y, where a semi join will never duplicate rows of x.

```
semi_join(ndat, jdat, by = c("Site" = "Samples"))
```

```
## # A tibble: 18 x 3
##
      Taxa
                           Site
                                    0TUs
##
      <chr>
                           <chr>
                                  <dbl>
##
    1 Clostridia
                           V_16_2 17572
    2 Clostridia
                           T 2 9 17471
##
                           V 17 2 13875
##
    3 Clostridia
    4 Clostridia
                           T_2_6 12169
##
##
    5 Bacteroidia
                           V_17_1 11392
    6 Clostridia
##
                           V_10_1 11180
##
    7 Clostridia
                           T_2_3 10944
##
    8 Unknown
                           T_9_2 10624
```

```
## 9 Clostridia
                          T 4 4 10545
## 10 Clostridia
                          V 16 1 10043
## 11 Clostridia
                          T 5 3
                                  8981
## 12 Alphaproteobacteria V_3_2
                                  8660
## 13 Clostridia
                          V_15_1
                                  8244
## 14 Clostridia
                          V 17 1
                                  7722
## 15 Clostridia
                          V 18 1
                                  7481
## 16 Alphaproteobacteria V 3 1
                                  7322
## 17 Clostridia
                          V_18_4
                                  6246
## 18 Clostridia
                          T_2_1
                                  6213
```

semi\_join() returns the 18 rows of ndat that have a Site match in jdat. Note that we have 3 columns; the columns from jdat have not been added.

It is easier to see the effect of semi-join by switching the order of our data frame input (ie. semi\_join(jdat, ndat, by = c("Samples" = "Site"))). How many rows and columns do you expect in this output?

Anti join returns all rows from x where there are not matching values in y, keeping just columns from x.

This returns our 2 rows in ndat that did not have a match in jdat. Again, we have 3 columns; the columns from jdat have not been added.

# Challenge

Given the definitions for the inner\_, left\_, right\_, full\_, semi\_ and anti\_joins, what would you expect the resulting data frame to be if jdat and ndat were reversed? Write down the number of rows and columns you would expect the data frame to have, then run the code. Did you find any surprises?

## Resources

https://github.com/wmhall/tidyr\_lesson/blob/master/tidyr\_lesson.md
http://vita.had.co.nz/papers/tidy-data.pdf
https://thinkr.fr/tidyverse-hadleyverse/
http://stat545.com/bit001\_dplyr-cheatsheet.html
http://dplyr.tidyverse.org/articles/two-table.html
https://github.com/datacarpentry/archive-datacarpentry/tree/master/data/tidy-data - lotr dataset

Thanks for coming!!!

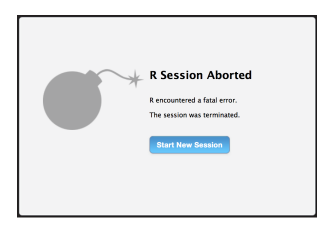


Figure 8: