tutorial017-Pandas-DataWranglingAutomation

November 16, 2022

1 Automating data wrangling

Sometimes we require a "one off" solution to a unique data analysis problem. In this situation, we write code to do a particular analysis on a particular data set. Then, if the analysis is part of a publication, we make the code and data publically available and... we're done.

Often, however, we require a *reusable* solution that operates on data of a given format even though some of the particulars, such as sample size or variable names, might change. In this case, we want our code to be "dynamic" in the sense that it should be able to handle any anticipated changes to the details of the input data.

Here, we'll tackle the same problem as last time – reformatting a data set from a cumbersome format into a more useful and "tidy" format.

1.0.1 Learning goals:

- write reusable code for a data wrangling problem
- create a function to make the code handy to use

1.1 Import pandas and look at the data from last time

```
[57]: import pandas as pd
```

Read in the data from last time.

```
[58]: my_input_data = pd.read_csv('datasets/017DataFile.csv')
```

Take a peek to remind ourselves of the data format.

```
[59]: my_input_data.head()
```

```
[59]:
         Male Mutant
                       Female Mutant
                                       Male Wild Type
                                                        Female
                                                                 Wild Type
      0
            10.485451
                             8.250013
                                             20.127063
                                                                 25.946384
      1
            11.747948
                             8.453839
                                             20.068147
                                                                 23.464870
      2
            13.412580
                             9.706605
                                             21.215148
                                                                 22.989480
      3
            12.910095
                             9.522116
                                             20.706416
                                                                 25.324376
            10.367770
                             8.583212
                                             18.074795
                                                                 22.607487
```

In this data set, there are two "independent variables", sex and genotype of laboratory rats, and one "dependent variable", response time. The data are formatted such that each column contains the

data from a unique combination of the two independent variables, *i.e.* a "cell" of the experimental design. Like this:

	male	female
mutant	mm	fm
${\bf wildtype}$	mw	fw

This format might seem to make sense, but it's actually not very flexible. For analysis purposes, it's generally better to have data in a format that obeys a couple of rules:

- each row should correspond to a single observation (measurement)
- each column should correspond to a single variable

Data in this format are also referred to as "tidy".

So in this case, our goal is to take the above data and put it into a format like this:

response time	sex	genotype
rt value	male or female	wild or not

Once the data are in this format, we can easily use our tools to do things like compare wild to mutant, or compare wild to mutant only in females, etc.

Last time, we stacked the reaction time values into a single column using pandas functions. This relied on us knowing and "hard coding" the column names ("Male Mutant", etc.). If we're going to automate things, we want our code to be agnostic about these. One way would be to somehow read the column names into variables and work with them somehow...

But what about numpy arrays? We already know how to manipulate those and, since they are just numbers, there are no column names or pesky row indexes to worry about. So let's try using numpy!

```
[60]: import numpy as np
```

Pandas dataframes know how to convert themselves to numpy arrays. They have a to_numpy() method that will pull *just the numbers* out of our dataframe, ignoring the column labels and row indexs.

```
[61]: raw_data = my_input_data.to_numpy()
```

Let's take a look!

```
[62]: raw_data
```

```
[11.69842177, 9.83500171, 20.36762403, 23.05218737], [11.58315277, 10.53209602, 20.15252058, 25.3690367], [11.44734892, 9.39416641, 19.39247581, 23.37270897], [10.85227619, 8.73947266, 18.52434071, 25.21564644], [11.28589742, 10.89239399, 20.32502629, 24.99050453]])
```

1.2 Get some useful information from the original data

So far so good! Now we are going to put the data into the format we want. To automate this, we are going to get

- the number of observations in each group (which is the number of rows), and
- the number of groups (which is the number of columns)

and store them in variables.

```
[63]: obs_per_grp, grps = raw_data.shape print("We have ", obs_per_grp, " observations per group and ", grps, " groups.")
```

We have 10 observations per group and 4 groups.

Now we'll calculate the total number of observations, which is also how long we want our new data frame to be.

```
[64]: new_length = obs_per_grp*grps
print("We have ", new_length, " total observations.")
```

We have 40 total observations.

Complete the following exercise.

• Use the cell below and explain in your own words why we used Numpy Arrays in the previous cells. What was our final goal? Why did we dump the data into a Numpy Array?

It allows us to view the data and overall a quicker way to see everything rather than seeing the raw data.

1.3 Build our response time (dependent variable) column

We could now play legos "by hand", stacking the columns of our numpy array on top of each other to make a new array (and we already know how to do that).

Or we could take advantage of the fact that one of the things numpy arrays know how to do – one of the methods they have – is to change their shape. So we'll take our obs by cols array and numpy.reshape() into a new_length by 1 array.

What this command does (effectively) is read out the data values from the original array one-by-one, and places them in the cells of a new array of a shape you specify. The only catch is that the total number of cells in the new array has to be the same as in the old array – in other words, each and every data value has to have one and only one place to go in the new array. Which makes sense.

```
[65]: values_col = np.reshape(raw_data, (new_length, 1))
```

I called it values_col because it will eventually become the values column of our new pandas data frame.

```
Let's see if that worked:
[66]: values_col
[66]: array([[10.48545088],
              [8.2500131],
              [20.12706278],
              [25.94638414],
              [11.74794775],
              [8.45383932],
              [20.06814699],
              [23.46487013],
              [13.41258004],
              [ 9.70660484],
              [21.21514789],
              [22.98948034],
              [12.91009526],
              [ 9.52211638],
              [20.70641578],
              [25.32437595],
              [10.36777045],
              [8.58321246],
              [18.07479515],
              [22.60748688],
              [11.69842177],
              [ 9.83500171],
              [20.36762403],
              [23.05218737],
              [11.58315277],
              [10.53209602],
              [20.15252058],
              [25.3690367],
              [11.44734892],
              [ 9.39416641],
              [19.39247581],
              [23.37270897],
              [10.85227619],
              [8.73947266],
              [18.52434071],
              [25.21564644],
              [11.28589742],
              [10.89239399],
              [20.32502629],
```

```
[24.99050453]])
```

[23.46487013],

Nice! But let's make absolutely sure that worked. What we want is for the columns of the original data to be stacked on top of one another. Is that what we have?

Nope, it's not right. What happened is that the values got read out *left to right, top to bottom* (or row-wise) and placed into the new array one-by-one. But what we want is for the values to be read *top to bottom*, *left to right* (or columnwise). We can make this happen with the order= argument of numpy.reshape().

```
[67]: values_col = np.reshape(raw_data, (new_length, 1), order = 'F')
```

```
Let's make sure that worked:
[68]:
     values_col
[68]: array([[10.48545088],
              [11.74794775],
              [13.41258004],
              [12.91009526],
              [10.36777045],
              [11.69842177],
              [11.58315277],
              [11.44734892],
              [10.85227619],
              [11.28589742],
              [8.2500131],
              [8.45383932],
              [ 9.70660484],
              [ 9.52211638],
              [8.58321246],
              [ 9.83500171],
              [10.53209602],
              [ 9.39416641],
              [8.73947266],
              [10.89239399],
              [20.12706278],
              [20.06814699],
              [21.21514789],
              [20.70641578],
              [18.07479515],
              [20.36762403],
              [20.15252058],
              [19.39247581],
              [18.52434071],
              [20.32502629],
              [25.94638414],
```

```
[22.98948034],
[25.32437595],
[22.60748688],
[23.05218737],
[25.3690367],
[23.37270897],
[25.21564644],
[24.99050453]])
```

Yay! It did!

Useless trivia: Two of Ye Olde Major Programming Languages are **C** (used mainly by programmers) and **Fortran** (used mainly by scientists). C (the language used to write Python) uses row-wise indexing, whereas Fortran uses columnwise indexing. That's why "F" is used to specify columnwise indexing above: the "F" is for "Fortran".

Minor annoying thing: (there is always at least one that pops up in any coding task, amirite?) values_col is a (40x1) 2-dimensional numpy array but, when we go to build our new data frame, we'll need it to be a 40 long (40,) 1-dimensional array.

This actually comes up so often that numpy has a squeeze() function to squeeze the dimension of length one into nothingness. It turns (n, 1) things into (n, 1) things.

Let's check the shape of our new array:

```
[69]: values_col.shape
```

[69]: (40, 1)

Now let's squeeze the (uneeded and unwanted) column dimension into oblivion:

```
[70]: values_col = np.squeeze(values_col)
```

And check the shape again:

```
[71]: values_col.shape
```

[71]: (40,)

Okay, that worked, now onto...

Complete the following exercise.

- Use the next cell to explain what happened to the numpy array after the squeeze operation.

 It deleted the 1 column.
 - Type below code demonstrating how you could explore the help for the method .shape() to explore what it does:

```
[72]: values_col.shape?
np.reshape?
```

• Use the cell below to explain the use of the method .resape():

It changes the shape of the array (reshapes it), like changing the order of the data.

1.4 Building the independent variable columns

What we require is that the levels our two independent variables repeat themselves in the right order down their respective columns. We could certainly type this in by hand, but that would be really annoying to change if we required new labels later on or something.

We could also use for() loops; they are designed for exactly such repetitive tasks after all. That might look something like this:

```
[73]: gen_var = list()  # create a python list

for i in range(new_length):  # loop through all observations

if i < new_length/2:  # for the first half, ...

gen_var.append("wildtype")  # set to male

else:  # otherwise...

gen_var.append("mutant")  # set to female
```

```
[74]: print(gen_var)
```

```
['wildtype', 'wildtype', 'wildtype', 'wildtype', 'wildtype',
'wildtype', 'wildtype', 'wildtype', 'wildtype', 'wildtype',
'wildtype', 'wildtype', 'wildtype', 'wildtype', 'wildtype',
'wildtype', 'wildtype', 'mutant', 'mutant', 'mutant', 'mutant', 'mutant',
'mutant', 'mutant', 'mutant', 'mutant', 'mutant', 'mutant', 'mutant',
'mutant', 'mutant', 'mutant', 'mutant', 'mutant', 'mutant']
```

We'd have to get a little bit more fancy with our if... to create the sex variable, that'd be the idea

But pandas provides easy ways to repeat and stack things (numpy does too), so let's try those. The two will use are

- pandas.Series.repeat()
- pandas.concat()

Note: When you see pandas.Series.somefunction() or pandas.DataFrame.somefunction() in the documentation, that means that all Series or DataFrames know how to do somefunction(). So if you had a Series named Phred, you would say Phred.somefunction() to use somefunction().

Complete the following exercise.

• Use the cell below to explain what the variable new_length contain:

It contains the all the observations there are.

• Use the cell below to explain the reason why we use new_length/2 in combination with the if, else:

We use it to assign the strain label. If the data point is in the first half of the list then it is assigned 'wildtype', otherwise it is assigned 'mutant'.

1.4.1 Make the genetic strain variable

In the way we have formatted the data, genetic strain is the "outer" variable, in that it only changes once as we go down the data set: all the wildtypes are on top, and all mutants are on the bottom. The sex variable is the "inner" variable, because it changes once within each value of strain, so it needs to three times as we go down the data set.

This is arbitrary and has nothing to do with the experimental design; we could have formatted the data such that the roles were reversed.

What we will do is

- make a short series containing the two levels of our variable
- repeat each value to make the long series
- deal with annoying index values (there's always something...)

```
[75]: strain = pd.Series(['wildtype', 'mutant'])  # make the short series
strain = strain.repeat(2*obs_per_grp)  # repeat each over two cell's worth_

of data
strain = strain.reset_index(drop=True)  # reset the series's index value
```

Complete the following exercise.

• Use the cell below to explain what is and what it is contained by the variable strain:

The variable strain contains the strain type label for the data point.

Let's see if that worked:

```
[76]: print(strain)
     0
            wildtype
            wildtype
     1
      2
            wildtype
     3
            wildtype
     4
            wildtype
     5
            wildtype
     6
            wildtype
     7
            wildtype
            wildtype
     8
     9
            wildtype
     10
            wildtype
     11
            wildtype
     12
            wildtype
     13
            wildtype
      14
            wildtype
            wildtype
     15
      16
            wildtype
     17
            wildtype
     18
            wildtype
     19
            wildtype
     20
              mutant
```

```
21
         mutant
22
         mutant
23
        mutant
24
         mutant
25
        mutant
26
         mutant
27
         mutant
28
         mutant
29
        mutant
30
        mutant
31
         mutant
32
         mutant
33
         mutant
34
         mutant
35
         mutant
36
        mutant
37
         mutant
38
        mutant
39
        mutant
dtype: object
```

Complete the following exercise.

• Use the cell below to explain why mutants appear at the bottom of the previous Pandas Series, who decided that order?

We did.

1.4.2 Make the sex variable

As the sex variable is the inner variable, we need it have ['male'..., 'female'...] within each outer block of genotype. So what we'll do is make one block of ['male'..., 'female'...] and then just stack two copies of that to make our variable. So the steps are

- make a short series containing the two levels of our variable (just like above)
- repeat it (just like above)
- stack two copies on top of each other (dropping the annoying indexes in the process)

```
[77]: sexes = pd.Series(['male', 'female'])  # make the short series
sexes = sexes.repeat(obs_per_grp)  # repeat each over one cell's_\text{\text{\text{or "concatonate" two_\text{\text{U}}}}}
sexes = pd.concat([sexes]*2, ignore_index=True)  # stack or "concatonate" two_\text{\text{U}}
copies
```

```
[78]: print(sexes)
```

```
0 male
1 male
```

1 mare

2 male

3 male

```
4
         male
5
         male
6
         male
7
         male
8
         male
9
         male
10
       female
11
       female
12
       female
13
       female
14
       female
15
       female
16
       female
17
       female
18
       female
19
       female
20
         male
21
         male
22
         male
23
         male
24
         male
25
         male
26
         male
27
         male
28
         male
29
         male
30
       female
31
       female
32
       female
33
       female
34
       female
35
       female
36
       female
37
       female
38
       female
39
       female
dtype: object
```

Complete the following exercise.

• Use the cell below to explain in your own words what happened in the previous cell:

We created a series with sex labels, repeated it and stacked the two copies to create the dataset.

• Use the cell below to show your code to create a pandas series called unicorns comprising of 20 mistical equines half of which are white and half pearl-white in color (well ... what what do you want, they are unicorns):

```
[79]: unicorns = pd.Series(['white', 'pearl-white'])
      unicorns = unicorns.repeat(obs_per_grp)
      unicorns = pd.concat([unicorns], ignore_index=True)
      print(unicorns)
     0
                  white
     1
                  white
     2
                  white
     3
                  white
     4
                  white
     5
                  white
     6
                  white
     7
                  white
     8
                  white
     9
                  white
     10
           pearl-white
     11
           pearl-white
     12
           pearl-white
     13
           pearl-white
     14
           pearl-white
     15
           pearl-white
     16
           pearl-white
     17
           pearl-white
           pearl-white
     18
           pearl-white
     dtype: object
```

• Use the cell below to show your code to create a pandas series called Three trees comprising of 30 trees 1/3 of which are Live Oaks, 1/3 White Oaks and 1/3 Red Oaks:

```
[80]: three_trees = pd.Series(['live oaks', 'white oaks', 'red oaks'])
    three_trees = three_trees.repeat(obs_per_grp)
    three_trees = pd.concat([three_trees], ignore_index = True)

print(three_trees)
```

```
0
       live oaks
       live oaks
1
2
       live oaks
3
       live oaks
4
       live oaks
5
       live oaks
6
       live oaks
7
       live oaks
8
       live oaks
9
       live oaks
      white oaks
10
```

```
white oaks
11
12
      white oaks
13
      white oaks
14
      white oaks
15
      white oaks
      white oaks
16
17
      white oaks
18
      white oaks
      white oaks
19
20
        red oaks
21
        red oaks
22
        red oaks
23
        red oaks
24
        red oaks
25
        red oaks
26
        red oaks
27
        red oaks
28
        red oaks
29
        red oaks
dtype: object
```

1.4.3 Build our new data frame!

Data frames are created in pandas by handing it data it can make sense of. There are various ways to accomplish this, and one handy one is to hand it data in a "column label 1: data 1, column label 2: data 2, ..." format.

We can accomplish this with a python "dictionary" (remember those?). A python dict associates a label (the "word") with a value or set of values or whatever (the "definition"). They are very useful, so let's take a look at a simple example before we use one to build out data frame. You create a dictionary using curly braces, and then use colons to bind each word or key with its definition or value. Commas separate each key-value pair.

Complete the following exercise.

• Use the cell below to build a dict() describing a student, with a name, with a student ID, a GPA and a major, make up all the values but use the lables as described here:

So a dictionary associates a label with data values. Perfect!

Time to build our data frame!

Note that the formatting above is just to make the columns we're creating more obvious and human-readable. This will work too:

It's just not as pretty.

Let's look at our creation!

```
[87]: my_tidy_data
```

```
[87]:
                RTs
                        sex
                               strain
      0
          10.485451
                       male
                             wildtype
      1
          11.747948
                       male
                             wildtype
      2
          13.412580
                            wildtype
                       male
      3
          12.910095
                       male wildtype
      4
          10.367770
                       male wildtype
      5
          11.698422
                       male wildtype
          11.583153
      6
                       male wildtype
      7
          11.447349
                       male wildtype
                       male wildtype
      8
          10.852276
      9
          11.285897
                       male wildtype
      10
          8.250013 female wildtype
      11
           8.453839
                     female wildtype
      12
           9.706605
                     female wildtype
      13
           9.522116
                     female wildtype
      14
           8.583212
                     female wildtype
          9.835002
                     female wildtype
      15
      16
          10.532096
                     female wildtype
      17
                     female wildtype
          9.394166
      18
           8.739473
                     female wildtype
      19
          10.892394
                    female wildtype
      20
          20.127063
                       male
                               mutant
```

```
21
    20.068147
                  male
                           mutant
22
    21.215148
                  male
                           mutant
23
    20.706416
                  male
                           mutant
24
    18.074795
                  male
                          mutant
    20.367624
25
                  male
                          mutant
26
    20.152521
                  male
                          mutant
27
    19.392476
                  male
                          mutant
28
    18.524341
                  male
                          mutant
29
    20.325026
                  male
                          mutant
30
    25.946384
               female
                          mutant
31
    23.464870
               female
                          mutant
32
   22.989480
                female
                          mutant
33
    25.324376
                female
                          mutant
34
    22.607487
                female
                          mutant
    23.052187
35
                female
                          mutant
36
    25.369037
                female
                          mutant
37
    23.372709
                female
                          mutant
    25.215646
38
                female
                          mutant
39
    24.990505
                female
                           mutant
```

Yay! We win!

Important point: Crucially, the above code doesn't rely on us knowing much about the input data ahead of time. As long as it's a pandas data frame that contains numerical values, the code will run. It's automatic.

1.5 Look at new data with more observations with same code

We'll make this code self-contained, so it can be run without running anything above. We'll also add comments, so that future-us can read the code more easily without having to wade through the notebook text above.

```
[88]: my_input_data = pd.read_csv('datasets/018DataFile.csv') # read the data

raw_data = my_input_data.to_numpy() # convert to numpy_

obs, grps = raw_data.shape # get the number of_

rows and columns

# get the number of_

# get the number
```

Check the size of the new data real quick:

```
[89]: print("We have ", obs, " observations per group and ", grps, " groups.")
```

We have 20 observations per group and 4 groups.

And now run the "meat" of the code:

```
[90]: new_length = obs*grps
                                                                   # compute total number_
       ⇔of observations
      values_col = np.reshape(raw_data, (new_length, 1),
                               order = 'F')
                                                                  # reshape the array
      values_col = np.squeeze(values_col)
                                                                  # squeeze to make 1D
      # construct the inner grouping variable
      sexes = pd.Series(['male', 'female'])
                                                                  # define the levels
      sexes = sexes.repeat(obs)
                                                                  # make one cycle of \square
       ⇔the levels
      sexes = pd.concat([sexes]*2, ignore_index=True)
                                                                  # and repeat the
      ⇔cycle, ditching the indexes
      # construct the outer grouping variable
      strain = pd.Series(['wildtype', 'mutant'])
                                                                  # define the levels
      strain = strain.repeat(2*obs)
                                                                  # make the one cycle
      strain = strain.reset_index(drop=True)
                                                                  # drop the pesky index
      # construct the data frame
      my_new_tidy_data = pd.DataFrame(
          {
               "RTs": values col,
                                                                  # make a column named
       \hookrightarrow RTs and put the values in
              "sex": sexes,
                                                                  # ditto for sex
               "strain": strain
                                                                  # and for genetic_{\sqcup}
       \hookrightarrow strain
          }
```

[91]: my_new_tidy_data

```
[91]:
             RTs
                          strain
                   sex
     0 12.333785 male wildtype
     1 11.675152 male wildtype
       12.029059 male wildtype
     3 12.126430 male wildtype
     4
        10.307197 male wildtype
     75 24.886821 female
                          mutant
     76 24.475663 female
                          mutant
     77 21.935896 female mutant
     78 23.852748 female mutant
     79 25.515138 female mutant
     [80 rows x 3 columns]
```

Success!

1.6 Making the code even more functional

Now we have a chunk of code that seems handy and re-usable. How could we make it ever more handy?

If we make it into a *function*, then we can run the whole entire thing just by typing one command – no copying, no pasting, fewer ways to make mistakes.

1.6.1 Defining a function

Since we already have all the code, we can literally just indent it and throw a def... in front of it!

```
[92]: def tidyMyData() :
          import pandas as pd
          import numpy as np
          my_input_data = pd.read_csv('datasets/018DataFile.csv') # read the data
          raw_data = my_input_data.to_numpy()
                                                                      # convert to numpy_
       \hookrightarrow array
          obs, grps = raw_data.shape
                                                                      # get the number_
       ⇔of rows and columns
          new_length = obs*grps
                                                                      # compute total
       →number of observations
          values_col = np.reshape(raw_data, (new_length, 1),
                                   order = 'F')
                                                                     # reshape the array
          values_col = np.squeeze(values_col)
                                                                      # squeeze to make_
       \hookrightarrow 1D
          # construct the inner grouping variable
          sexes = pd.Series(['male', 'female'])
                                                                      # define the levels
          sexes = sexes.repeat(obs)
                                                                      # make one cycle_
       ⇔of the levels
          sexes = pd.concat([sexes]*2, ignore_index=True)
                                                               # and repeat the cycle,\Box
       ⇔ditching the indexes
          # construct the outer grouping variable
          strain = pd.Series(['wildtype', 'mutant'])
                                                                      # define the levels
          strain = strain.repeat(2*obs)
                                                                      # make the one
       ⇔cycle
          strain = strain.reset_index(drop=True)
                                                                      # drop the pesky_
       \hookrightarrow index
```

```
datFromFun = tidyMyData()
[93]:
[94]: datFromFun
[94]:
                RTs
                        sex
                               strain
          12.333785
      0
                       male wildtype
      1
          11.675152
                      male wildtype
      2
          12.029059
                      male wildtype
      3
          12.126430
                      male wildtype
      4
          10.307197
                      male wildtype
      . .
      75
         24.886821 female
                               mutant
      76
         24.475663 female
                               mutant
      77
         21.935896 female
                               mutant
      78
         23.852748 female
                               mutant
         25.515138 female
      79
                               mutant
      [80 rows x 3 columns]
```

1.6.2 Defining a function with an argument

A common (very common) scenario in data analysis is wanting to run the same code – like the code we just wrote – on different files. So one really nice addition to this function would be to add the ability for the user to specify a filename to tell the function which data file to read.

This is actually fairly straightforward. All we have to do as add an **argument** to our function, and then replace the hardcoded filename in the function with the **variable** created by the function argument.

```
[95]: def tidyMyData(filename) :
    import pandas as pd
    import numpy as np

my_input_data = pd.read_csv(filename) # read the data
```

```
raw_data = my_input_data.to_numpy()
                                                              # convert to numpy_
\hookrightarrow array
   obs, grps = raw_data.shape
                                                              # get the number
⇔of rows and columns
  new_length = obs*grps
                                                              # compute total
⇔number of observations
   values_col = np.reshape(raw_data, (new_length, 1),
                            order = 'F')
                                                              # reshape the array
  values_col = np.squeeze(values_col)
                                                              # squeeze to make_
→1D
   # construct the inner grouping variable
   sexes = pd.Series(['male', 'female'])
                                                              # define the levels
   sexes = sexes.repeat(obs)
                                                              # make one cycle_
⇔of the levels
   sexes = pd.concat([sexes]*2, ignore_index=True) # and repeat the cycle, __
⇔ditching the indexes
   # construct the outer grouping variable
   strain = pd.Series(['wildtype', 'mutant'])
                                                            # define the levels
   strain = strain.repeat(2*obs)
                                                              # make the one
⇔cycle
   strain = strain.reset_index(drop=True)
                                                              # drop the pesky_
\hookrightarrow index
   # construct the data frame
   my_new_tidy_data = pd.DataFrame(
       {
           "RTs": values col,
                                                              # make a column
→named RTs and put the values in
           "sex": sexes,
                                                              # ditto for sex
           "strain": strain
                                                              # and for genetic_
\hookrightarrowstrain
       }
   )
   return my_new_tidy_data
```

Now we can call the function and specify whatever data files exist. Let's try it with "datasets/018DataFile2.csv"!

```
[96]: newDataFromFun = tidyMyData("datasets/018DataFile2.csv")
```

[97]: newDataFromFun [97]: RTs strain sex 0 12.577226 malewildtype 1 12.778183 malewildtype 2 13.389130 male wildtype 3 12.747877 malewildtype 4 13.615121 male wildtype 163 24.539374 femalemutant 164 23.877924 female mutant 165 23.161896 female mutant 166 24.426455 female mutant 167 21.990136 female mutant [168 rows x 3 columns]

1.6.3 Adding help

It's always a good idea to heavily comment your code!

When writing fuctions, it's also a good idea to add a documentation string, called a docstring, to your function. This way people can get help on your function with the help() function. Like help(tidyMyData).

```
[98]: def tidyMyData(filename):
          tidyMyData() Takes one-column-per-cell rat reaction time data as input.
          Returns tidy one-column-per-variable data.
          User specifies a filename string.
          import pandas as pd
          import numpy as np
          my_input_data = pd.read_csv(filename) # read the data
          raw_data = my_input_data.to_numpy()
                                                                      # convert to numpy_
       \hookrightarrow array
          obs, grps = raw_data.shape
                                                                      # get the number
       →of rows and columns
                                                                      # compute total
          new_length = obs*grps
       →number of observations
          values_col = np.reshape(raw_data, (new_length, 1),
                                   order = 'F')
                                                                      # reshape the array
```

```
values_col = np.squeeze(values_col)
                                                                # squeeze to make_
\hookrightarrow 1D
  # construct the inner grouping variable
  sexes = pd.Series(['male', 'female'])
                                                               # define the levels
  sexes = sexes.repeat(obs)
                                                                # make one cycle
⇔of the levels
  sexes = pd.concat([sexes]*2, ignore_index=True)
                                                         # and repeat the cycle,
⇔ditching the indexes
  # construct the outer grouping variable
  strain = pd.Series(['wildtype', 'mutant'])
                                                               # define the levels
  strain = strain.repeat(2*obs)
                                                               # make the one
⇔cycle
  strain = strain.reset_index(drop=True)
                                                               # drop the pesky_
\hookrightarrow index
  # construct the data frame
  my_new_tidy_data = pd.DataFrame(
       {
           "RTs": values_col,
                                                               # make a column_
⇔named RTs and put the values in
           "sex": sexes,
                                                               # ditto for sex
           "strain": strain
                                                                # and for genetic
\hookrightarrowstrain
  )
  return my_new_tidy_data
```

[99]: help(tidyMyData)

```
Help on function tidyMyData in module __main__:

tidyMyData(filename)
    tidyMyData() Takes one-column-per-cell rat reaction time data as input.
    Returns tidy one-column-per-variable data.
    User specifies a filename string.
```

Complete the following exercise.

• Use the cell below to show how you would modify the previous function so as to make it even more flexible. Let the user specify the output column headers to be whatever they want.

More specifically how would you allos passing in the three labels, sex, RTs and strain, instead of having them 'hard coded' inside the code. This means that instead of using labels such as sex, RTs and strain, we will want to pass parameters for each one of the labels and use the parameters

in the function. For example, instead of sex, RTs and strain we will want to pass others say, s, ReactionTime or type or any three combinations of lables, always three but that can change everytime we call the function.

You would do this with arguments (obviously). But you could do it with multiple arguments, so users would call it like:

```
tidyMyData("datasets/018DataFile2.csv", "Times", "Gender", "Genotype")
or you could do it with one additional arguments, so the user would call it by either:
tidyMyData("datasets/018DataFile2.csv", ["Times", "Gender", "Genotype"])
or
colNames = ["Times", "Gender", "Genotype"]
```

Pro tip: The function would probably be most handy if there were *default* values for the column names, so that user could just type something like

```
myTidyData = tidyMyData("datasets/018DataFile2.csv")
```

tidyMyData("datasets/018DataFile2.csv", colNames)

if they didn't want to specify custom column headers.

```
[53]:
           ReactionTime Gender
                                 Genotype
                           male wildtype
      0
              12.577226
      1
              12.778183
                           male wildtype
      2
              13.389130
                           male wildtype
      3
                                 wildtype
              12.747877
                           male
      4
              13.615121
                                 wildtype
                           \mathtt{male}
      163
              24.539374 female
                                   mutant
      164
              23.877924 female
                                   mutant
      165
              23.161896 female
                                   mutant
      166
              24.426455 female
                                   mutant
      167
              21.990136 female
                                   mutant
```

[168 rows x 3 columns]

```
[100]: def tidyMyData(df1) :
    import pandas as pd
    import numpy as np

my_input_data = pd.read_csv(df1) # read the data
```

```
raw_data = my_input_data.to_numpy()
                                                              # convert to numpy_
\hookrightarrow array
  obs, grps = raw_data.shape
                                                              # get the number
⇔of rows and columns
  new_length = obs*grps
                                                              # compute total
⇔number of observations
  values_col = np.reshape(raw_data, (new_length, 1),
                           order = 'F')
                                                             # reshape the array
  values_col = np.squeeze(values_col)
                                                              # squeeze to make_
→1D
  # construct the inner grouping variable
  sexes = pd.Series(['male', 'female'])
                                                             # define the levels
  sexes = sexes.repeat(obs)
                                                              # make one cycle_
⇔of the levels
  sexes = pd.concat([sexes]*2, ignore_index=True) # and repeat the cycle, __
⇔ditching the indexes
  # construct the outer grouping variable
  strain = pd.Series(['wildtype', 'mutant'])
                                                            # define the levels
  strain = strain.repeat(2*obs)
                                                             # make the one
⇔cycle
  strain = strain.reset_index(drop=True)
                                                             # drop the pesky_
\hookrightarrow index
  # construct the data frame
  my_new_tidy_data = pd.DataFrame(
       {
           "RTs": values col,
                                                              # make a column
→named RTs and put the values in
           "sex": sexes,
                                                             # ditto for sex
           "strain": strain
                                                              # and for genetic_
\hookrightarrowstrain
       }
  return my_new_tidy_data
```

```
[102]: tidyMyData("datasets/018DataFile.csv")
```

```
[102]: RTs sex strain
0 12.333785 male wildtype
1 11.675152 male wildtype
```

```
male wildtype
2
   12.029059
               male wildtype
3
   12.126430
                male wildtype
4
   10.307197
. .
         •••
75 24.886821 female
                       mutant
76 24.475663 female
                       mutant
77 21.935896 female
                       mutant
78 23.852748 female
                       mutant
79 25.515138 female
                       mutant
[80 rows x 3 columns]
```

[]: