Recently I came across a question where someone was looking to take a bunch of CSV files, each of which contained numerical columns, and (a) get them into R, (b) calculate the mean and standard deviation of every column in every CSV file, and (c) calculate some overall summary like the mean of all the means and the mean of all the standard deviations.

I already know how to use $map_dfr()$ to read a lot of CSVs with the same structure into a nice tidy tibble. (I'll show you below in a moment if you don't know how to do this.) It's a nice demonstration of the utility of iterating over a vector of filenames with a tidy result. But in thinking about the question I also wanted to provide a reproducible answer, which meant thinking about how to *create* a bunch of CSV files to read in by way of an example. So, here's one way to do that. Along the way we'll take advantage of a few small features of common tidyverse functions that I've found very helpful but whose existence is sometimes a little hard to discover. Or rather, their utility is sometimes hard to see when reading the help pages.

First let's make a bunch of CSVs in a folder called tmpdat. Each CSV will have five columns containing 100 normally-distributed observations with a mean of zero and a standard deviation of one.

Here's the code all at once:

```
library(tidyverse)
  ## Make a "tmpdat" folder in the working dir if one doesn't exist
  ifelse(!dir.exists(file.path("tmpdat")), dir.create(file.path("tmpdat")), FALSE)
  ## Number of columns in each of our made-up data files.
  nvars <- 5
8
9
  ## Create the CSV files
10
paste0("csv_", 1:100) %>%
   set names() %>%
12
    map(~ replicate(n = nvars, rnorm(100, 0, 1))) %>%
13
    map dfr(as tibble, .id = "id", .name repair = ~ paste0("v", 1:nvars)) %>%
14
    group by(id) %>%
15
    nest() %>%
16
    pwalk(\sim write csv(x = .y, file = paste0("tmpdat/", .x, ".csv")))
17
```

The first few lines load the tidyverse and create the tmpdir folder if it doesn't already exist. We could do this in a full-on tidyverse way with the fs package, but here I just use the Base R equivalent.

Next we create nvars which is the number of columns each of our CSV files will have.

Now the fun starts. The first line creates a vector of 100 identifiers. The call to <code>set_names()</code> gives each element of the vector a name, which by default is the same as the value of that element. We do this so that the elements of the list we're about to create will have recognizable names as well.

```
1 paste0("csv_", 1:100) %>%
2  set_names()
3 #>    csv_1    csv_2    csv_3    csv_4    csv_5    csv_6    csv_7
4 csv_8
5 #>    "csv_1"    "csv_2"    "csv_3"    "csv_4"    "csv_5"    "csv_6"    "csv_7"
6 "csv_8"
7
```

```
## (Cut off to save space)
```

Next, we use map() to feed each of our hundred elements to the replicate() function. We don't pass through any arguments to replicate at all (except for nvars which is the same for all of them). Instead, what happens is that replicate() creates a list of one hundred random matrices, each one of dimensions 100×5. One matrix lives inside each named list item, from csv 10 csv 100.

```
paste0("csv_", 1:100) %>%
    set names() %>%
   map(\sim replicate(n = nvars, rnorm(100, 0, 1)))
  #> $csv 1
5
                                [,2]
                                                [,4]
  #>
                   [,1]
                                          [,3]
 6
     [1,] -0.116601453 -2.339554484 -2.10008625 0.48430248 0.73549398
  #>
     [2,] -1.281593689 0.979867725 0.12217918 0.67191402 -2.10950592
8
     [3,] -0.131462279 2.025939426 1.01061386 -0.97849787 -0.86495635
10
  ## (Cut off to save space)
```

Step three, convert each matrix to a tibble and then bind them all together into one large tibble:

```
paste0("csv_", 1:100) %>%
    set names() %>%
3
    map(\sim replicate(n = nvars, rnorm(100, 0, 1))) %>%
 4
    map dfr(as tibble, .id = "id", .name repair = ~ paste0("v", 1:nvars))
 5
 6
  #> # A tibble: 10,000 x 6
        id
                 v1
                       v2
                                             v5
  #>
                               v3
                                      \nabla 4
8
  #>
9
  #> 1 csv 1 -0.867 0.119 0.171 1.92 1.03
10
  #> 2 csv 1 0.163 0.714 -1.31 -3.06 0.470
11
  #> 3 csv_1 0.604 -0.282 -0.228 0.647 0.302
  #> 4 csv 1 -0.775 -2.31 -0.522 -0.661 -0.160
13
  #> 5 csv 1 -0.862 1.03 -0.781 -0.115 -1.61
14
  #> 6 csv 1 0.943 1.12 0.251 -0.170 -0.356
15
  #> 7 csv 1 -0.277 -1.03 -0.864 -1.62 1.70
16
  #> 8 csv 1 0.613 -0.360 -0.491 1.01 0.436
17
  #> 9 csv 1 -0.520 0.711 -2.77 2.10 0.450
  #> 10 csv 1 1.01 0.657 0.519 -0.630 -0.927
  \#> \# ... with 9,990 more rows
20
21
```

It's starting to look a little tidier now. Like map(), $map_dfr()$ feeds each element of the data—in this case, 100 list items, each of which contains a matrix—to the $as_tibble()$ function. This converts each matrix to a tibble. Unlike map(), which always returns a list, $map_dfr()$ will return a data frame or tibble (bound by row). Along the way we add an id column, to keep track of which imaginary dataset this will be from, and we also name the columns of the made-up data. The $.name_repair$ argument has several useful pre-sets, but you can also send it a function. We do that here, to create the column names v1, v2, etc, all the way to however many columns there are. Tibbles require unique column names.

Onward:

```
paste0("csv_", 1:100) %>%
   set names() %>%
    map(~ replicate(n = nvars, rnorm(100, 0, 1))) %>%
    map_dfr(as_tibble, .id = "id", .name_repair = ~ paste0("v", 1:nvars)) %>%
 5
    group by(id) %>%
 6
    nest()
 7
 8
  #> # A tibble: 100 x 2
  #> # Groups: id [100]
11 #>
       id
            data
  #>
12
#> 1 csv_1
#> 2 csv_2
  #> 3 csv_3
#> 4 csv_4
15
#> 5 csv_5
#> 6 csv_6
10 #> 7 csv_7
#> 8 csv_8
20 #> 9 csv_9
21 #> 10 csv_10
  #> # ... with 90 more rows
23
```

Having created our data, we $group_by()$ the id column and use nest() to create a list column of datasets, which are now all tibbles and all have consistent column names. Finally, we write each one out to a file:

```
1
2 paste0("csv_", 1:100) %>%
2    set_names() %>%
3    map(~ replicate(n = nvars, rnorm(100, 0, 1))) %>%
4    map_dfr(as_tibble, .id = "id", .name_repair = ~ paste0("v", 1:nvars)) %>%
5    group_by(id) %>%
6    nest() %>%
7    nest() %>%
8    pwalk(~ write_csv(x = .y, file = paste0("tmpdat/", .x, ".csv")))
```

We use walk() when we want to iterate over a list, just as with map(). But walk() is for those times when the result of whatever we do is not an object we'll use further in R, but rather a "side effect", like a file or a plot. The pwalk() function is a special case of walk() designed for tibbles and data frames. It iterates over each row of the tibble. Here it takes the second argument of the tibble, the data list-column, and writes out its contents to a file whose name is constructed from the id column.

Now we have conjured up one hundred CSVs of made-up data. Perhaps a career in Social Psychology awaits us.

```
immood ~[D/c/s/d/impdat git:master ) is

csv_1.csv csv_18.csv csv_27.csv csv_36.csv csv_45.csv csv_54.csv csv_63.csv csv_72.csv csv_81.csv csv_98.csv csv_18.csv csv_19.csv csv_28.csv csv_37.csv csv_46.csv csv_55.csv csv_64.csv csv_73.csv csv_82.csv csv_92.csv csv_38.csv csv_47.csv csv_56.csv csv_65.csv csv_74.csv csv_83.csv csv_92.csv csv_11.csv csv_28.csv csv_38.csv csv_48.csv csv_57.csv csv_66.csv csv_75.csv csv_84.csv csv_92.csv csv_12.csv csv_21.csv csv_38.csv csv_48.csv csv_57.csv csv_66.csv csv_75.csv csv_84.csv csv_93.csv csv_12.csv csv_21.csv csv_38.csv csv_48.csv csv_58.csv csv_67.csv csv_76.csv csv_86.csv csv_94.csv csv_13.csv csv_22.csv csv_31.csv csv_48.csv csv_55.csv csv_68.csv csv_77.csv csv_86.csv csv_94.csv csv_14.csv csv_23.csv csv_32.csv csv_56.csv csv_56.csv csv_68.csv csv_77.csv csv_86.csv csv_95.csv csv_15.csv csv_24.csv csv_33.csv csv_42.csv csv_56.csv csv_68.csv csv_77.csv csv_78.csv csv_96.csv csv_15.csv csv_24.csv csv_34.csv csv_56.csv csv_68.csv csv_77.csv csv_88.csv csv_97.csv csv_16.csv csv_28.csv csv_97.csv csv_16.csv csv_
```

All the data

With our data now on disk, we can read it all back in and do our calculations. Once again, map_dfr() is our friend. We feed it a vector, filenames, which is just the full path to each CSV file in tmpdat.

```
1
  filenames %>%
    map dfr(read csv, .id = "id", col types = cols()) %>%
    group_by(id) %>%
    summarize(across(everything(),
5
                    list(mean = mean, sd = sd))) %>%
 6
    pivot longer(-id,
7
                names to = c("col", ".value"), names sep= " ") %>%
8
   group_by(col) %>%
9
    summarize(avg mean = mean(mean),
10
             avg_sd = mean(sd))
  #> # A tibble: 5 x 3
12
  #>
      col avg mean avg sd
13
  #>
14
  #> 1 v1
            0.0119 0.993
15
  #> 2 v2
            0.000967 0.992
  #> 3 v3 -0.000190 0.991
17
  #> 4 v4
          -0.00533 0.994
18
  #> 5 v5 -0.0172 0.986
19
20
```

Three things here. First, sending filenames down the pipe results in each element of it (i.e. each filename) getting fed one at a time to $read_csv()$. Because we're using $map_dfr()$ to do the feeding, we get a tibble back. Second, we know we're pivoting all the columns except id, so instead of naming the column range we're including, we drop the single column that's not part of the pivot, using the shorthand of putting a minus sign in front of its name, like this: -id. Third, given that we know what we're dealing with inside each CSV, we use $col_types = cols()$ to suppress the column specification message that would otherwise be displayed at the console for each file as it's read in.

```
1 filenames %>%
   map_dfr(read_csv, .id = "id", col_types = cols())
3 #> # A tibble: 10,000 x 6
4 #>
       id
               v1
                      v2
                            v3
                                  v 4
                                          v_5
5 #>
           -2.72
 6 #> 1 1
                   1.44 -0.883 0.603 -0.739
7 #> 2 1
           -1.31 -0.516 0.701 0.0594 1.56
            0.834 1.00 -2.13
8 #> 3 1
                                1.70
                                      -0.591
           -0.139 0.601 -0.356 -1.12
9 #> 4 1
                                       0.167
           -1.43 0.194 1.20 -0.284
10 #> 5 1
                                       0.457
11 #> 6 1
           -0.0937 0.116 -0.725 -0.521 -0.677
12 #> 7 1
            0.556 -1.87 1.20 -0.449 1.92
```

Next we group by id (i.e., by CSV file) and use across() to get the means and standard deviations for everything. There's no missing data, so we don't need to add na.rm = TRUE in the summarize statement.

```
filenames %>%
   map dfr(read csv, .id = "id", col types = cols()) %>%
   group by(id) %>%
   summarize(across(everything(),
               list(mean = mean, sd = sd)))
1
2 #> # A tibble: 100 x 11
3 #> id v1 mean v1 sd v2 mean v2 sd v3 mean v3 sd v4 mean v4 sd v5 mean
4 v5_sd
5 #>
6 #> 1 1
          0.0649 1.06 0.0335 0.991 0.0182 1.06 -0.131 1.07 -0.0231
7 1.05
8 #> 2 10 0.00919 0.917 0.0283 1.04 -0.0184 0.970 0.0950 1.04 0.0243
90.973
10 #> 3 100  0.0760  1.08  -0.172  1.13  0.0303  1.05  0.00903  1.01  0.148
11 1.03
12 #> 4 11 0.0506 0.949 0.0667 1.00 0.0904 0.884 -0.0381 1.10 0.0538
13 0.954
14 #> 5 12 0.0938 1.00 0.117 1.05 -0.105 0.994 -0.0474 0.926 0.0981
15 0.992
16 #> 6 13 -0.112 0.933 -0.178 1.04 0.162 0.934 -0.0837 0.990 0.00821
17 1.01
191.09
20 #> 8 15 -0.103 1.13 0.0927 0.994 0.186 1.02 -0.166 1.07 -0.110
21 0.989
0.830
 0.992
  #> # ... with 90 more rows
```

Now, what we'd like next is to end up with a tibble with four columns that looks like this:

```
5 1
        v5 -0.0231 1.05
9
  6 10
              0.00919 0.917
       v1
10
  7 10 v2
              0.0283 1.04
11
  8 10 v3 -0.0184 0.970
12
  9 10 v4
              0.0950 1.04
13
  10 10 v5
              0.0243 0.973
# ... with 490 more rows
16
```

That is, a tidy or long-form summary, where the first column is the CSV id, the second is which variable we're talking about, and the third and fourth columns are the summary statistics for that variable in that CSV.

If you use pivot longer() to do this in the default way, you will not get quite what you want:

```
filenames %>%
   map_dfr(read_csv, .id = "id", col_types = cols()) %>%
   group by(id) %>%
   summarize(across(everything(),
 5
                   list(mean = mean, sd = sd))) %>%
 6
   pivot longer(-id)
 7
  #> # A tibble: 1,000 x 3
 8
      id name value
  #>
 9
  #>
10
  #> 1 1 v1 mean 0.0649
11
  #> 2 1
           v1 sd 1.06
#> 3 1 v2_mean 0.0335
#> 4 1
14 ...
          v2 sd 0.991
15 #> 5 1
           v3_mean 0.0182
#> 6 1
           v3 sd 1.06
  #> 7 1
           v4 mean -0.131
" #> 8 1 v4_sd
17
                   1.07
  #> 9 1
           v5_mean -0.0231
19
#> 10 1 v5_sd 1.05
  #> # ... with 990 more rows
21
22
```

And if you read in the docs for pivot_longer() you might also try giving it a regular expression to remove the variable prefixes and get them in to their own column:

```
1 filenames %>%
   map_dfr(read_csv, .id = "id", col_types = cols()) %>%
   group_by(id) %>%
   summarize (across (everything (),
5
                    list(mean = mean, sd = sd))) %>%
 6 pivot_longer(-id,
7
                 names pattern = "(v\d?) (.*)",
8
                 names_to = c("mean", "sd")) #< this is wrong</pre>
9
10 \#> \# A tibble: 1,000 x 4
11 #>
      id mean sd value
12 #>
```

```
13
  #>
     1 1
             771
                          0.0649
                   mean
  #> 2 1
                  sd
             17.1
                         1.06
15
  #> 3 1
                         0.0335
             v2
                  mean
16
  #> 4 1
            v2
                 sd
                         0.991
17
  #> 5 1
             v3
                         0.0182
                   mean
18
  #> 6 1
             v3
                        1.06
19
20 #> 7 1
                   mean -0.131
             \nabla 4
  #> 8 1
             v4
                         1.07
                   sd
-
#> 9 1
             v5
                   mean -0.0231
  #> 10 1
             v5
                   sd
                         1.05
23
  \#> \# ... with 990 more rows
24
25
```

Whoops, that's not right either. And even if we clean up the column headers we would still be left wanting to pivot wider again to get mean and sd in their own columns. But *that* would create a tibble with alternating NAs on each row for mean and sd. All we want is for the mean and sd parts to become the column names, and get their respective value. We should be able to do this in one step.

We can:

```
filenames %>%
   map_dfr(read_csv, .id = "id", col_types = cols()) %>%
3
    group by(id) %>%
 4
    summarize(across(everything(),
 5
                   list(mean = mean, sd = sd))) %>%
    pivot_longer(-id,
7
               names_sep= " ",
8
                names to = c("col", ".value"))
9
10
  #> # A tibble: 500 x 4
11
       id col mean
  #>
                              sd
12
  #>
13
  #> 1 1
                  0.0649 1.06
             v1
14
  #> 2 1
             v2
                  0.0335 0.991
15
  #> 3 1
           v3
                  0.0182 1.06
16
  #> 4 1
             \nabla 4
                  -0.131 1.07
17
  #> 5 1
            v5 -0.0231 1.05
18
  #> 6 10
             v1
                   0.00919 0.917
19
  #> 7 10
             v2
                  0.0283 1.04
20
  #> 8 10
             v3
                  -0.0184 0.970
21
  #> 9 10
             v4
                  0.0950 1.04
22
  #> 10 10 v5
                   0.0243 0.973
23
  #> # ... with 490 more rows
24
25
```

We create the names of the new columns by splitting the existing names ($v1_mean$, $v1_sd$ etc) on the _character. In the first place the split gives us v1, v2, v3, etc, which we put into a column named col. This leaves us with mean and sd names, each with its own particular value. Now, we don't want to put alternating mean and sd names in a single column named, say, "measure", with their values in a value column, as is the default. We want a single column of mean values and a single column of sd values. The trick is the special .value sentinel in the names_to argument. As noted in the help, the names_to argument is "a string specifying the name of the column to create from the data stored in the column

names of data." This can be a character vector, thus enabling the pivoting of multiple columns. And in addition, the help goes on to note,

. value indicates that component of the name defines the name of the column containing the cell values, overriding values to.

It might not jump out at you how handy this is. What it means is that we just take whatever unique name elements are left over when we split the original column names, and we use those as the names of the new columns. The corresponding values get inserted in a column with that name. This is a *very* useful option, because we find ourselves wanting to pivot out summary statistics quite a lot. The basic action of summarize() in conjunction with group_by() will do a lot for us a lot of the time. But sometimes we want to change the shape of the data we have in just this way. In those cases, being aware of ".value" in pivot_longer() is your friend.

From there we can get the overall statistics we originally wanted, grouping by col to return the mean of all means and the mean of all sds per column:

```
1
 2 filenames %>%
 3 map dfr(read csv, .id = "id", col types = cols()) %>%
 4 group_by(id) %>%
 5 summarize(across(everything(),
           list(mean = mean, sd = sd))) %>%
 7 pivot longer(-id,
 8
               names to = c("col", ".value"), names sep= " ") %>%
9 group_by(col) %>%
10   summarize(avg_mean = mean(mean),
11
       avg sd = mean(sd)
12
13 #> # A tibble: 5 x 3
14 #> col avg_mean avg_sd
15 #>
16 #> 1 v1 0.0119 0.993
17 #> 2 v2 0.000967 0.992
18 #> 3 v3 -0.000190 0.991
19 #> 4 v4 -0.00533 0.994
20 #> 5 v5 -0.0172 0.986...
21
```