Week 2 - Codebook

October 14, 2024

1 Week 2: Importing Data

1.1 Clear the entire workspace

```
[52]: rm(list=ls())
```

1.2 Load required libraries

```
[53]: ReqdLibs = ReddLibs = ReddL
```

1.3 Theme defaults

This is only for the Jupyter Notebook so you the figures axes are larger

```
thm = theme(
    strip.text.x=element_text(size=20,face="bold"),
    strip.text.y=element_text(size=20,face="bold"),
    legend.text=element_text(size=16,face="bold"),
    legend.position = "top",
    legend.title=element_text(size=16,face="bold"),
    title =element_text(size=14, face='bold'),
    text = element_text(colour = "black",size=18),
    plot.title = element_text(colour = "black",size = 22, face = "bold"),
    axis.ticks.length = unit(0.3,"cm"),
    axis.line = element_line(colour = "black",size=0.85),
    axis.ticks = element_line(colour = "black",size=0.85),
    axis.text = element_text(colour = "black",size=24),
    axis.title=element_text(size=25))
```

1.4 Retrieve the folder path to the data files, a.k.a, the Root Folder

Note that this section is slightly different from the R project folder because the Jupyter notebook is outside teh R project folder and is read within the root folder ReproRehab_Bootcamp so we are going to need to direct it to the Materials/Week 2/R project folder.

```
[55]: folder_path = here("Materials/Week 2/R project", "Raw Data")
# output the folder name
print(folder_path)
# make sure it exists
dir.exists(folder_path)
```

[1] "/Users/rinivarghese/Documents/My Documents/JHU/Lab/Professional Development/ReproRehab 2024/ReproRehab_Bootcamp/Materials/Week 2/R project/Raw Data"

TRUE

1.5 What subfolders and files are within the root folder? Let's check.

OK, once we have the 'folder path' to the root folder, i.e., Raw Data, it is important to know exactly what the structure within is. In our case, the structure looks something like this: Raw

```
sub1/
                     sub1 rest.xlsx
                                           sub1 trial1.xlsx
                                                                  sub1 trial2.xlsx
Data/
                     sub2_rest.xlsx
                                          sub2\_trial1.xlsx
                                                                  sub2_trial2.xlsx
   . . .
         sub2/
         sub3/
                     sub3_rest.xlsx
                                           sub3_trial1.xlsx
                                                                   sub3_trial2.xlsx
   . . .
         . . .
```

```
[56]: options(warn=0)
subfolder_path = here(folder_path, 'Sub1')

files.test=list.files(subfolder_path)
files.test

#Let's read in one file to see how ugly the data are
temp0=suppressMessages(read_excel(here(subfolder_path,files.test[1]),))
head(temp0)
```

1. 'Sub1_rest.xlsx' 2. 'Sub1_trial1.xlsx' 3. 'Sub1_trial2.xlsx' 4. 'Sub1_trial3.xlsx' 5. 'Sub1_trial4.xlsx' 6. 'Sub1_trial5.xlsx' 7. 'Sub1_trial6.xlsx'

| | ID code: | 11 | 3 | Test number: | 88 | |
|--------------------------|---|--------------------------------------|----------------|---|---|---|
| A tibble: 6×129 | <chr $>$ | <chr></chr> | <lgl $>$ | <chr $>$ | <chr $>$ | |
| | Last name: | SUBJECT | NA | Test date: | 6/21/2017 | |
| | First name: | NO1 | NA | Test time: | 10:16 |] |
| | Sex: | M | NA | N. of steps: | 53 | |
| | Age: | 26 | NA | Duration (hh:mm:ss): | 00:02:59 |] |
| | Height (in): | 72.834645669291334 | NA | BSA (m^2): | 2.1000404912839743 |] |
| | Weight (lb): | 189.59770013580487 | NA | BMI (Kg/m^2): | 25.127830533235937 |] |
| A tibble: 6×129 | First name: Sex: Age: Height (in): | NO1 M 26 72.834645669291334 | NA NA NA | Test time: N. of steps: Duration (hh:mm:ss): BSA (m^2): | 10:16 53 00:02:59 2.10004049128397 | |

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1.6 So, as you see above, the data are quite messy

we will modify the code so we only import a certain range Let's just do the first 5 rows where the data is in long format

```
[57]: temp=suppressMessages(read_excel(here(subfolder_path,files.test[1]),range = cell_cols("J:0")))
head(temp)
# the first two rows are header-like information so remove it
temp=temp[-c(1,2),-2]
head(temp)
```

| | \mathbf{t} | 2 | Rf | | VT | | VE | | VO2 |
|------------------------|--------------|-----------|------------|-----------|-------------|-----------|------------|-----------|----------|
| | <chr $>$ | <dbl $>$ | <chr $>$ | | <chr $>$ | | <chr $>$ | | <chr $>$ |
| | hh:mm:ss | NA | b/min | | 1 | | l/min | | ml/mii |
| | 0 | NA | NA | | NA | | NA | | NA |
| A tibble: 6×6 | 00:00:02 | 2 | 18.5185185 | 18518519 | 0.754853053 | 861151733 | 13.9787602 | 252065134 | 408.698 |
| | 00:00:05 | 3 | 20.9790209 | 79020977 | 0.553898929 | 987980249 | 11.6202572 | 270205645 | 304.786 |
| | 00:00:08 | 3 | 18.2370820 | 66869302 | 0.705889612 | 229617564 | 12.8733667 | 789595908 | 347.704 |
| | 00:00:10 | 2 | 25.5319148 | 93617021 | 0.782394989 | 35139695 | 19.9760422 | 28131226 | 614.813 |
| | _ | Df | | V | | VE | | VO9 | |
| A tibble: 6×5 | t | Rf | | VT | | VE | | VO2 | |
| | <chr $>$ | <chr $>$ | | <chr $>$ | | <chr $>$ | | <chr $>$ | |
| | 00:00:02 | 18.518518 | 518518519 | 0.7548530 | 05361151733 | 13.978760 | 0252065134 | 408.6987 | 04538219 |
| | 00:00:05 | 20.979020 | 979020977 | 0.5538989 | 92987980249 | 11.62025 | 7270205645 | 304.7867 | 51427508 |
| | 00:00:08 | 18.237082 | 066869302 | 0.7058896 | 61229617564 | 12.873366 | 6789595908 | 347.7043 | 21728538 |
| | 00:00:10 | 25.531914 | 893617021 | 0.7823949 | 98935139695 | 19.976042 | 228131226 | 614.8134 | 54296942 |
| | 00:00:13 | 21.978021 | 978021978 | 0.4957548 | 84331783432 | 10.895710 | 0842150205 | 253.6430 | 90772051 |
| | 00:00:16 | 18.927444 | 794952681 | 0.7874953 | 34782174502 | 14.905274 | 472217814 | 437.0966 | 60485628 |

1.7 Now we do this iteratively.

We go to our root folder Raw Data then we loop through all the subjects' folders within it to repeat the steps described above.

1.7.1 create a list of all the folder names within the root folder

```
[58]: dir.list = dir(folder_path)
    dir.list
```

1. 'Sub1' 2. 'Sub10' 3. 'Sub11' 4. 'Sub12' 5. 'Sub13' 6. 'Sub2' 7. 'Sub3' 8. 'Sub4' 9. 'Sub5' 10. 'Sub6' 11. 'Sub7' 12. 'Sub8' 13. 'Sub9'

1.7.2 create an empty data.frame that can accommodate any variable type

data.frame(list()): Converts the empty list into a data frame. Lists can accommodate any data type, i.e., numeric, string, characters, booleans etc. Since the list is empty, the resulting data frame will have no columns and no rows. In other words, this command initializes an empty data frame, which can later be filled with data, but starts with no content (i.e., no columns and no rows).

```
[59]: data.all = data.frame(list())
```

1.8 Method 1: using a for loop to compile the master dataset

We use a for loop in R to iteratively "stack" individual participant data tables. For each participant, the steps are the samea as we did above.

```
[60]: for(i in 1:length(dir.list)){
        files.import=list.files(here(folder_path,dir.list[i]))
        for(j in 1:length(files.import)){
          #Give me only the rows I need
          temp=suppressMessages(read excel(here(folder path,dir.list[i],files.
       →import[j]),
                                            range = cell_cols("J:0")))
          #Remove the random stuff
          temp=temp[-c(1,2),-2]
          #Convert to numeric
          temp[,c(2:5)]=apply(temp[,c(2:5)],2,as.numeric)
          #Covert to seconds
          temp$t=seconds(times(temp$t))+(minutes(times(temp$t))*60)
          #Assign Sub id
          temp$Sub=dir.list[i]
          #Assign trial id
          if(nchar(files.import[j])<16){</pre>
          temp$trial="rest"
          }else{
            temp$trial=paste("trial",as.numeric(substr(files.import[j],nchar(files.
       →import[j])-5,nchar(files.import[j])-5)))
          # this final step is where the 'stacking' happens
          data.all=rbind(data.all,temp)
        }
      }
```

1.8.1 Last step! Check the compiled dataset dimensions

```
[61]: head(data.all) dim(data.all)
```

```
Rf
                                    VT
                                                VE
                                                           VO2
                t
                                                                     Sub
                                                                              trial
                                                <dbl>
                                                                              <chr>
                < dbl >
                         <dbl>
                                    <dbl>
                                                           <dbl>
                                                                      <chr>
                2
                         18.51852 0.7548531
                                                13.97876
                                                          408.6987
                                                                     Sub1
                                                                              rest
                         20.97902 \quad 0.5538989
                                                11.62026
                                                          304.7868
                                                                     Sub1
                                                                              rest
A tibble: 6 \times 7
                         18.23708 0.7058896
                                                12.87337 \quad 347.7043
                                                                     Sub1
                                                                              rest
                10
                         25.53191 0.7823950
                                                19.97604 614.8135
                                                                     Sub1
                                                                              rest
                13
                         21.97802 \quad 0.4957548
                                                10.89571
                                                          253.6431
                                                                     Sub1
                                                                              rest
                16
                         18.92744 0.7874953
                                                14.90527 \quad 437.0967
                                                                     Sub1
                                                                              rest
```

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1.9 Method 2: using map_df in the purrr package

map_df is used in two layers: * Outer Layer: Iterates over each directory in *dir.list*, listing files within each directory. * Inner Layer: Processes each file in the current directory by: 1) reading the Excel file, 2) cleaning the data, 3) converting time to seconds, and 4) assigning identifiers.

This nesting efficiently combines results from all files into a single data frame, allowing for streamlined data aggregation from multiple directories.

```
[62]: data.all <- map df(dir.list, function(dir name) {
        # List all files in the current directory
        files.import <- list.files(here(folder_path, dir_name))</pre>
        map_df(files.import, function(file_name) {
          # Read the Excel file
          temp <- suppressMessages(read_excel(here(folder_path, dir_name, file_name),_
       ⇔range = cell_cols("J:0")))
          # Clean the data
          temp <- \text{temp}[-c(1, 2), -2]
          temp[, 2:5] <- apply(temp[, 2:5], 2, as.numeric)
          # Convert time to seconds
          temp$t <- seconds(times(temp$t)) + (minutes(times(temp$t)) * 60)</pre>
          # Assign Sub id
          temp$Sub <- dir_name</pre>
          # Assign trial id
          temp$trial <- ifelse(nchar(file_name) < 16, "rest", paste("trial", as.</pre>
       -numeric(substr(file name, nchar(file name) - 5, nchar(file name) - 5))))
          return(temp)
        })
      })
      # Now data.all contains all the processed data
      head(data.all)
      dim(data.all)
```

| | \mathbf{t} | Rf | VT | VE | VO2 | Sub | trial |
|------------------------|--------------|----------|-----------|----------|----------|----------|-----------------------|
| A tibble: 6×7 | <d b l $>$ | <dbl $>$ | <dbl $>$ | <dbl $>$ | <dbl $>$ | <chr $>$ | <chr $>$ |
| | 2 | 18.51852 | 0.7548531 | 13.97876 | 408.6987 | Sub1 | rest |
| | 5 | 20.97902 | 0.5538989 | 11.62026 | 304.7868 | Sub1 | rest |
| | 8 | 18.23708 | 0.7058896 | 12.87337 | 347.7043 | Sub1 | rest |
| | 10 | 25.53191 | 0.7823950 | 19.97604 | 614.8135 | Sub1 | rest |
| | 13 | 21.97802 | 0.4957548 | 10.89571 | 253.6431 | Sub1 | rest |
| | 16 | 18.92744 | 0.7874953 | 14.90527 | 437.0967 | Sub1 | rest |

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1.10 Method 3: combined power of map_df and dplyr

all the same things as aboe, but note how we transform the data in the inner layer with dplyr.

```
[63]: data.all <- map_df(dir.list, function(dir_name) {
        # List all files in the current directory
        files.import <- list.files(here(folder_path, dir_name))</pre>
        map_df(files.import, function(file_name) {
          # Read the Excel file
          temp <- suppressMessages(read_excel(here(folder_path, dir_name, file_name),_
       →range = cell_cols("J:0"))) %>%
            # Clean and transform the data
            slice(-c(1, 2)) %>%
                                                    # Remove the first two rows
            select(-2) %>%
                                                   # Remove the second column
            mutate(across(2:5, as.numeric),  # Convert columns 2 to 5 to numeric
                   t = seconds(times(t)) + (minutes(times(t)) * 60), # Convert time_
       →to seconds
                                                   # Assign Sub id
                   Sub = dir_name,
                   trial = ifelse(nchar(file_name) < 16, "rest",</pre>
                                   # Assign trial id below by concatening diff pieces_
       \hookrightarrow of info
                                    paste("trial", as.numeric(substr(file_name, __
       →nchar(file_name) - 5, nchar(file_name) - 5)))))
          return(temp)
        })
      })
      # Now data.all contains all the processed data
      head(data.all)
      dim(data.all)
```

| | \mathbf{t} | Rf | VT | VE | VO2 | Sub | trial |
|------------------------|--------------|----------|-----------|----------|----------|----------|------------------------|
| A tibble: 6×7 | <dbl $>$ | <dbl $>$ | <dbl $>$ | <dbl $>$ | <dbl $>$ | <chr $>$ | <chr $>$ |
| | 2 | 18.51852 | 0.7548531 | 13.97876 | 408.6987 | Sub1 | rest |
| | 5 | 20.97902 | 0.5538989 | 11.62026 | 304.7868 | Sub1 | rest |
| | 8 | 18.23708 | 0.7058896 | 12.87337 | 347.7043 | Sub1 | rest |
| | 10 | 25.53191 | 0.7823950 | 19.97604 | 614.8135 | Sub1 | rest |
| | 13 | 21.97802 | 0.4957548 | 10.89571 | 253.6431 | Sub1 | rest |
| | 16 | 18.92744 | 0.7874953 | 14.90527 | 437.0967 | Sub1 | rest |

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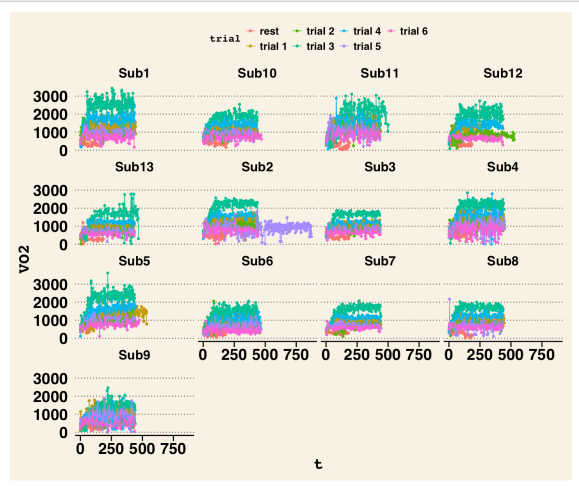
1.11 Visualization

Now that we have our compiled dataset, let's visualize it at different levels

1.11.1 Each timepoint for each trial for per participant

```
[64]: options(repr.plot.width = 12, repr.plot.height = 10)

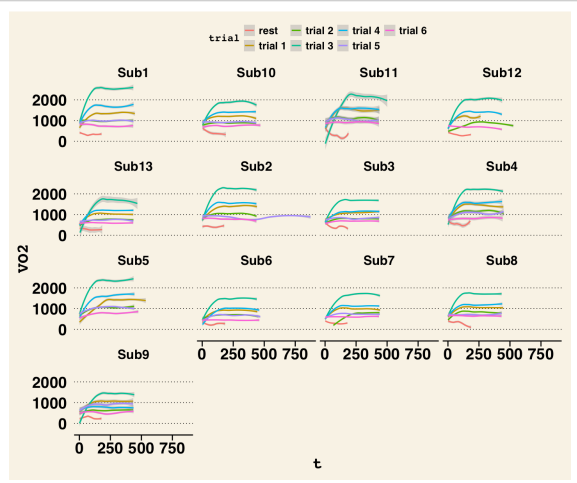
#Visualize raw data by subject
ggplot(data.all,aes(x=t,y=V02,color=trial))+
    geom_point()+
    geom_line()+
    facet_wrap(~Sub) + theme_wsj() + thm
```



1.11.2 Smoothed traces across time for each trial per participant

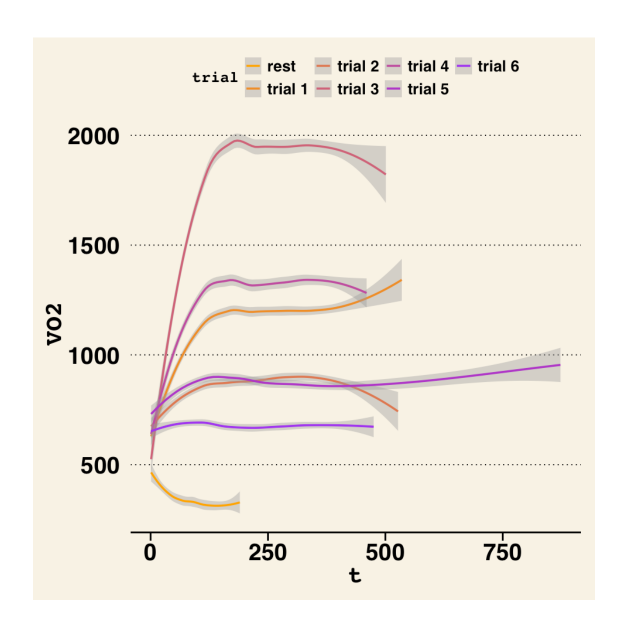
In smoothing, we use a 'loess' procedure. This function is in-built within the geom_smooth function of ggplot. The 'loess' or 'lowess' procedure follows a kind of windowed smoothing procedure. It assigns weights to each point within the window, with closer points given higher weights, to calculate the smoothed value for each data point.

```
[65]: #Visualize data using loess by subject
ggplot(data.all,aes(x=t,y=V02,color=trial))+
    geom_smooth(method = 'loess', formula = 'y~x')+
    facet_wrap(~Sub) + theme_wsj() + thm
```



1.11.3 Each trial averaged across particpants

```
[66]: options(repr.plot.width = 8, repr.plot.height = 8)
#create a color gradient
colfunc <- colorRampPalette(c("orange", "purple"))
#Visualize each trial across all participants
ggplot(data.all,aes(x=t,y=V02,color=trial))+
    geom_smooth(method = 'loess', formula = 'y~x')+
    scale_colour_manual(values = c(colfunc(7))) + theme_wsj() + thm</pre>
```



1.12 The End