Biostat 203B Homework 1

Due Jan 24, 2025 @ 11:59PM

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Display machine information for reproducibility:

```
sessionInfo()
```

```
R version 4.4.1 (2024-06-14)
Platform: x86_64-apple-darwin20
Running under: macOS 15.1
Matrix products: default
        /Library/Frameworks/R.framework/Versions/4.4-x86_64/Resources/lib/libRblas.0.dylib
LAPACK: /Library/Frameworks/R.framework/Versions/4.4-x86_64/Resources/lib/libRlapack.dylib;
locale:
[1] en_US.UTF-8/en_US.UTF-8/en_US.UTF-8/C/en_US.UTF-8/en_US.UTF-8
time zone: America/Los_Angeles
tzcode source: internal
attached base packages:
[1] stats
              graphics grDevices utils
                                             datasets methods
                                                                 base
loaded via a namespace (and not attached):
 [1] compiler_4.4.1
                       fastmap_1.2.0
                                         cli_3.6.3
                                                            tools_4.4.1
                                                            rmarkdown_2.29
 [5] htmltools_0.5.8.1 rstudioapi_0.17.0 yaml_2.3.10
 [9] knitr_1.48
                       jsonlite_1.8.9
                                                            digest_0.6.37
                                         xfun_0.48
[13] rlang_1.1.4
                       evaluate_1.0.1
```

Q1. Git/GitHub

No handwritten homework reports are accepted for this course. We work with Git and GitHub. Efficient and abundant use of Git, e.g., frequent and well-documented commits,

is an important criterion for grading your homework.

- 1. Apply for the Student Developer Pack at GitHub using your UCLA email. You'll get GitHub Pro account for free (unlimited public and private repositories).
- 2. Create a **private** repository biostat-203b-2025-winter and add Hua-Zhou and TA team (Tomoki-Okuno for Lec 1; parsajamshidian and BowenZhang2001 for Lec 82) as your collaborators with write permission.
- 3. Top directories of the repository should be hw1, hw2, ... Maintain two branches main and develop. The develop branch will be your main playground, the place where you develop solution (code) to homework problems and write up report. The main branch will be your presentation area. Submit your homework files (Quarto file qmd, html file converted by Quarto, all code and extra data sets to reproduce results) in the main branch.
- 4. After each homework due date, course reader and instructor will check out your main branch for grading. Tag each of your homework submissions with tag names hw1, hw2, ... Tagging time will be used as your submission time. That means if you tag your hw1 submission after deadline, penalty points will be deducted for late submission.
- 5. After this course, you can make this repository public and use it to demonstrate your skill sets on job market.

Solution Done. I've already created a repository named biostat-203b-2025-winter and here is the repository link.

Q2. Data ethics training

This exercise (and later in this course) uses the MIMIC-IV data v3.1, a freely accessible critical care database developed by the MIT Lab for Computational Physiology. Follow the instructions at https://mimic.mit.edu/docs/gettingstarted/ to (1) complete the CITI Data or Specimens Only Research course and (2) obtain the PhysioNet credential for using the MIMIC-IV data. Display the verification links to your completion report and completion certificate here. You must complete Q2 before working on the remaining questions. (Hint: The CITI training takes a few hours and the PhysioNet credentialing takes a couple days; do not leave it to the last minute.)

Solution Done. Here is the screenshot of mimic data access approval.



Q3. Linux Shell Commands

1. Make the MIMIC-IV v3.1 data available at location ~/mimic. The output of the ls -1 ~/mimic command should be similar to the below (from my laptop).

```
# content of mimic folder
ls -l ~/mimic/
```

```
total 48
                              15199 Oct 10 16:29 CHANGELOG.txt
-rw-r--r-0 1 vickey
                       staff
-rw-r--r-0 1 vickey
                       staff
                               2518 Oct 10 17:30 LICENSE.txt
                               2884 Oct 11 17:55 SHA256SUMS.txt
-rw-r--r-0 1 vickey
                       staff
drwxr-xr-x@ 23 vickey
                       staff
                                736 Jan 24 03:06 hosp
drwxr-xr-x0 11 vickey
                                352 Jan 22 16:08 icu
                       staff
```

Refer to the documentation https://physionet.org/content/mimiciv/3.1/ for details of data files. Do **not** put these data files into Git; they are big. Do **not** copy them into your directory. Do **not** decompress the gz data files. These create unnecessary big files and are not big-data-friendly practices. Read from the data folder ~/mimic directly in following exercises.

Use Bash commands to answer following questions.

2. Display the contents in the folders hosp and icu using Bash command 1s -1. Why are these data files distributed as .csv.gz files instead of .csv (comma separated values) files? Read the page https://mimic.mit.edu/docs/iv/ to understand what's in each folder.

Solution: Display the contents

```
ls -l ~/mimic/hosp
ls -l ~/mimic/icu
```

```
total 10845000
-rw-r--r--@ 1 vickey
                      staff
                               19928140 Jun 24
                                                2024 admissions.csv.gz
-rw-r--r--@ 1 vickey
                      staff
                                 427554 Apr 12
                                                2024 d_hcpcs.csv.gz
-rw-r--r-0 1 vickey
                                 876360 Apr 12
                                                2024 d_icd_diagnoses.csv.gz
                      staff
-rw-r--r-0 1 vickey
                      staff
                                 589186 Apr 12
                                                2024 d_icd_procedures.csv.gz
-rw-r--r--@ 1 vickey
                                            3 09:07 d_labitems.csv.gz
                      staff
                                  13169 Oct
-rw-r--r--@ 1 vickey
                      staff
                               33564802 Oct
                                             3 09:07 diagnoses_icd.csv.gz
                                9743908 Oct 3 09:07 drgcodes.csv.gz
-rw-r--r--@ 1 vickey
                      staff
                                                2024 emar.csv.gz
-rw-r--r--@ 1 vickey
                              811305629 Apr 12
                      staff
-rw-r--r-0 1 vickey
                                2162335 Apr 12 2024 hcpcsevents.csv.gz
                      staff
```

```
2592909134 Oct 3 09:08 labevents.csv.gz
-rw-r--r--@ 1 vickey staff
                             117644075 Oct 3 09:08 microbiologyevents.csv.gz
-rw-r--r--@ 1 vickey
                     staff
-rw-r--r--@ 1 vickey
                              44069351 Oct 3 09:08 omr.csv.gz
                     staff
-rw-r--r--@ 1 vickey
                               2835586 Apr 12 2024 patients.csv.gz
                     staff
                             525708076 Apr 12 2024 pharmacy.csv.gz
-rw-r--r--@ 1 vickey
                     staff
                             666594177 Apr 12
                                              2024 poe.csv.gz
-rw-r--r-0 1 vickey staff
-rw-r--r-0 1 vickey staff
                              55267894 Apr 12 2024 poe_detail.csv.gz
-rw-r--r--@ 1 vickey staff
                             606298611 Apr 12 2024 prescriptions.csv.gz
                               7777324 Apr 12 2024 procedures_icd.csv.gz
-rw-r--r-0 1 vickey staff
-rw-r--r-0 1 vickey staff
                                127330 Apr 12 2024 provider.csv.gz
                               8569241 Apr 12 2024 services.csv.gz
-rw-r--r-0 1 vickey staff
-rw-r--r-0 1 vickey staff
                              46185771 Oct 3 09:08 transfers.csv.gz
total 8506784
                                              2024 caregiver.csv.gz
-rw-r--r-0 1 vickey staff
                                 41566 Apr 12
-rw-r--r-0 1 vickey staff
                            3502392765 Apr 12
                                              2024 chartevents.csv.gz
-rw-r--r-0 1 vickey staff
                                 58741 Apr 12 2024 d_items.csv.gz
-rw-r--r-0 1 vickey staff
                              63481196 Apr 12 2024 datetimeevents.csv.gz
-rw-r--r-0 1 vickey staff
                               3342355 Oct 3 07:36 icustays.csv.gz
-rw-r--r--@ 1 vickey
                             311642048 Apr 12 2024 ingredientevents.csv.gz
                     staff
-rw-r--r-0 1 vickey staff
                             401088206 Apr 12 2024 inputevents.csv.gz
                              49307639 Apr 12 2024 outputevents.csv.gz
-rw-r--r--@ 1 vickey
                     staff
                              24096834 Apr 12 2024 procedureevents.csv.gz
-rw-r--r-0 1 vickey staff
```

3. Briefly describe what Bash commands zcat, zless, zmore, and zgrep do.

Solution

• zcat is used to display the contents of a compressed file. It is equivalent to cat command for uncompressed files.

```
zcat < ~/mimic/icu/icustays.csv.gz | head</pre>
```

subject_id,hadm_id,stay_id,first_careunit,last_careunit,intime,outtime,los
10000032,29079034,39553978,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10000690,25860671,37081114,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10000980,26913865,39765666,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10001217,24597018,37067082,Surgical Intensive Care Unit (SICU),Surgical Intensive Care Unit
10001217,27703517,34592300,Surgical Intensive Care Unit (SICU),Surgical Intensive Care Unit
10001725,25563031,31205490,Medical/Surgical Intensive Care Unit (MICU/SICU),Medical/Surgical
10001843,26133978,39698942,Medical/Surgical Intensive Care Unit (MICU/SICU),Medical/Surgical
10001884,26184834,37510196,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10002013,23581541,39060235,Cardiac Vascular Intensive Care Unit (CVICU),Cardiac Vascular Intensive Care Unit (CVICU)

• zless is used to display the contents of a compressed file one screen at a time. It is equivalent to less command for uncompressed files.

zless ~/mimic/hosp/d_hcpcs.csv.gz | head

```
code,category,long_description,short_description
00000,,,Invalid Code
0001F,2,,Composite measures
0002F,2,,Composite measures
0003F,2,,Composite measures
0004F,2,,Composite measures
0005F,2,,Composite measures
0006F,2,,Composite measures
0007F,2,,Composite measures
0007F,2,,Composite measures
0008F,2,,Composite measures
```

• zmore is used to display the contents of a compressed file one screen at a time. It is equivalent to more command for uncompressed files.

zmore ~/mimic/hosp/d_hcpcs.csv.gz | head

```
code, category, long_description, short_description
00000,,,Invalid Code
0001F,2,,Composite measures
0002F,2,,Composite measures
0003F,2,,Composite measures
0004F,2,,Composite measures
0005F,2,,Composite measures
0006F,2,,Composite measures
0007F,2,,Composite measures
0007F,2,,Composite measures
0008F,2,,Composite measures
```

• zgrep is used to search the contents of a compressed file using regular expressions. It is equivalent to grep command for uncompressed files.

```
zgrep -c "99291" ~/mimic/hosp/services.csv.gz
```

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4. (Looping in Bash) What's the output of the following bash script?

Solution

```
for datafile in ~/mimic/hosp/{a,l,pa}*.gz
do
   ls -l $datafile
done
```

The output of this bash is displaying all the file names statring with a, 1, and pa in the hosp folder.

Display the number of lines in each data file using a similar loop. (Hint: combine linux commands zcat < and wc -1.)

Solution:

```
for datafile in ~/mimic/hosp/{a,l,pa}*.gz
do
    zcat< $datafile | wc -l
done</pre>
```

546029 158374765 364628

5. Display the first few lines of admissions.csv.gz. How many rows are in this data file, excluding the header line? Each hadm_id identifies a hospitalization. How many hospitalizations are in this data file? How many unique patients (identified by subject_id) are in this data file? Do they match the number of patients listed in the patients.csv.gz file? (Hint: combine Linux commands zcat <, head/tail, awk, sort, uniq, wc, and so on.)

```
# display the first few lines
zcat < ~/mimic/hosp/admissions.csv.gz | head</pre>
```

```
# count the number of rows (excluding the header line)
zcat < ~/mimic/hosp/admissions.csv.gz |tail -n +2 | wc -l

546028

# count the number of unique hospitalizations
zcat < ~/mimic/hosp/admissions.csv.gz | tail -n +2 | awk -F, '{print $2}' | sort | uniq | wc</pre>
```

546028

```
# count the number of unique patients zcat < \sim/mimic/hosp/admissions.csv.gz | tail -n +2 | awk -F, '{print $1}' | sort | uniq | wc
```

223452

```
# display the first few lines of patients.csv.gz
zcat < ~/mimic/hosp/patients.csv.gz | head
# count the number of unique patients in patients.csv.gz
zcat < ~/mimic/hosp/patients.csv.gz | tail -n +2 | awk -F, '{print $1}' | sort | uniq | wc -1
subject_id,gender,anchor_age,anchor_year,anchor_year_group,dod
10000032,F,52,2180,2014 - 2016,2180-09-09
10000048,F,23,2126,2008 - 2010,</pre>
```

```
10000048,F,23,2126,2008 - 2010,

10000058,F,33,2168,2020 - 2022,

10000068,F,19,2160,2008 - 2010,

10000084,M,72,2160,2017 - 2019,2161-02-13

10000102,F,27,2136,2008 - 2010,

10000108,M,25,2163,2014 - 2016,

10000115,M,24,2154,2017 - 2019,

10000117,F,48,2174,2008 - 2010,

364627
```

Therefore the number of unique patients in admissions.csv.gz and patients.csv.gz are not the same.

6. What are the possible values taken by each of the variable admission_type, admission_location, insurance, and ethnicity? Also report the count for each unique value of these variables in decreasing order. (Hint: combine Linux commands zcat, head/tail, awk, uniq -c, wc, sort, and so on; skip the header line.)

Solution:

display the first few lines of admissions.csv.gz

zcat < ~/mimic/hosp/admissions.csv.gz | head</pre>

56227 TRANSFER FROM HOSPITAL 42365 WALK-IN/SELF REFERRAL

6317 TRANSFER FROM SKILLED NURSING FACILITY 5837 INTERNAL TRANSFER TO OR FROM PSYCH

12965 CLINIC REFERRAL 8518 PROCEDURE SITE

```
subject_id, hadm_id, admittime, dischtime, deathtime, admission_type, admit_provider_id, admission_
10000032,22595853,2180-05-06 22:23:00,2180-05-07 17:15:00,,URGENT,P49AFC,TRANSFER FROM HOSPI
10000032,22841357,2180-06-26 18:27:00,2180-06-27 18:49:00,,EW EMER.,P784FA,EMERGENCY ROOM,HO
10000032,25742920,2180-08-05 23:44:00,2180-08-07 17:50:00,,EW EMER.,P19UTS,EMERGENCY ROOM,HO
10000032,29079034,2180-07-23 12:35:00,2180-07-25 17:55:00,,EW EMER.,PO60TX,EMERGENCY ROOM,HO
10000068,25022803,2160-03-03 23:16:00,2160-03-04 06:26:00,,EU OBSERVATION,P39NWO,EMERGENCY R
10000084,23052089,2160-11-21 01:56:00,2160-11-25 14:52:00,,EW EMER.,P42H7G,WALK-IN/SELF REFE
10000084,29888819,2160-12-28 05:11:00,2160-12-28 16:07:00,,EU OBSERVATION,P35NE4,PHYSICIAN R
10000108,27250926,2163-09-27 23:17:00,2163-09-28 09:04:00,EU OBSERVATION,P40JML,EMERGENCY R
10000117,22927623,2181-11-15 02:05:00,2181-11-15 14:52:00,,EU OBSERVATION,P47EY8,EMERGENCY R
# count the number of unique values for admission_type
zcat < ~/mimic/hosp/admissions.csv.gz | tail -n +2 | awk -F, '{print $6}' | sort | uniq -c |
177459 EW EMER.
119456 EU OBSERVATION
84437 OBSERVATION ADMIT
54929 URGENT
42898 SURGICAL SAME DAY ADMISSION
24551 DIRECT OBSERVATION
21973 DIRECT EMER.
13130 ELECTIVE
7195 AMBULATORY OBSERVATION
# count the number of unique values for admission_location
zcat < ~/mimic/hosp/admissions.csv.gz | tail -n +2 | awk -F, '{print $8}' | sort | uniq -c |</pre>
244179 EMERGENCY ROOM
163228 PHYSICIAN REFERRAL
```

```
5734 PACU
 402 INFORMATION NOT AVAILABLE
 255 AMBULATORY SURGERY TRANSFER
# count the number of unique values for insurance
zcat < ~/mimic/hosp/admissions.csv.gz | tail -n +2 | awk -F, '{print $10}' | sort | uniq -c</pre>
244576 Medicare
173399 Private
104229 Medicaid
14006 Other
9355
463 No charge
# count the number of unique values
zcat < ~/mimic/hosp/admissions.csv.gz | tail -n +2 | awk -F, '{print $13}' | sort | uniq -c</pre>
336538 WHITE
75482 BLACK/AFRICAN AMERICAN
19788 OTHER
13972 WHITE - OTHER EUROPEAN
13870 UNKNOWN
10903 HISPANIC/LATINO - PUERTO RICAN
8287 HISPANIC OR LATINO
7809 ASIAN
7644 ASIAN - CHINESE
6597 WHITE - RUSSIAN
6205 BLACK/CAPE VERDEAN
6070 HISPANIC/LATINO - DOMINICAN
3875 BLACK/CARIBBEAN ISLAND
3495 BLACK/AFRICAN
3478 UNABLE TO OBTAIN
2162 PATIENT DECLINED TO ANSWER
2082 PORTUGUESE
1973 ASIAN - SOUTH EAST ASIAN
1886 WHITE - EASTERN EUROPEAN
1858 HISPANIC/LATINO - GUATEMALAN
1661 ASIAN - ASIAN INDIAN
1526 WHITE - BRAZILIAN
1320 HISPANIC/LATINO - SALVADORAN
```

```
1247 AMERICAN INDIAN/ALASKA NATIVE
920 HISPANIC/LATINO - COLUMBIAN
883 HISPANIC/LATINO - MEXICAN
774 SOUTH AMERICAN
725 HISPANIC/LATINO - HONDURAN
664 ASIAN - KOREAN
641 HISPANIC/LATINO - CUBAN
603 HISPANIC/LATINO - CENTRAL AMERICAN
596 MULTIPLE RACE/ETHNICITY
494 NATIVE HAWAIIAN OR OTHER PACIFIC ISLANDER
```

7. The icusays.csv.gz file contains all the ICU stays during the study period. How many ICU stays, identified by stay_id, are in this data file? How many unique patients, identified by subject_id, are in this data file?

Solution:

```
# display the first few lines
zcat < ~/mimic/icu/icustays.csv.gz | head
# the number of ICU stays
zcat < ~/mimic/icu/icustays.csv.gz | tail -n +2 | awk -F, '{print $3}' | sort | uniq | wc -l
# the number of unique patients
zcat < ~/mimic/icu/icustays.csv.gz | tail -n +2 | awk -F, '{print $1}' | sort | uniq | wc -l</pre>
```

subject_id,hadm_id,stay_id,first_careunit,last_careunit,intime,outtime,los
10000032,29079034,39553978,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10000690,25860671,37081114,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10000980,26913865,39765666,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10001217,24597018,37067082,Surgical Intensive Care Unit (SICU),Surgical Intensive Care Unit
10001217,27703517,34592300,Surgical Intensive Care Unit (SICU),Surgical Intensive Care Unit
10001725,25563031,31205490,Medical/Surgical Intensive Care Unit (MICU/SICU),Medical/Surgical
10001843,26133978,39698942,Medical/Surgical Intensive Care Unit (MICU/SICU),Medical/Surgical
10001884,26184834,37510196,Medical Intensive Care Unit (MICU),Medical Intensive Care Unit (M
10002013,23581541,39060235,Cardiac Vascular Intensive Care Unit (CVICU),Cardiac Vascular Intensive
94458
65366

The number of ICU stays is 94458 and the number of unique patients is 65366 in this data file.

8. To compress, or not to compress. That's the question. Let's focus on the big data file labevents.csv.gz. Compare compressed gz file size to the uncompressed file size. Compare the run times of zcat < ~/mimic/labevents.csv.gz | wc -1 versus

wc -l labevents.csv. Discuss the trade off between storage and speed for big data files. (Hint: gzip -dk < FILENAME.gz > ./FILENAME. Remember to delete the large labevents.csv file after the exercise.)

Solution:

```
# compare compressed and uncompressed file size
ls -lh ~/mimic/hosp/labevents.csv.gz
# Uncompress the file temporarily and check its size
gzip -dk < ~/mimic/hosp/labevents.csv.gz > ./labevents.csv
ls -lh labevents.csv
-rw-r--r-@ 1 vickey staff 2.4G Oct 3 09:08 /Users/vickey/mimic/hosp/labevents.csv.gz
-rw-r--r--@ 1 vickey staff
                               17G Jan 24 14:29 labevents.csv
# compare run times
time zcat < ~/mimic/hosp/labevents.csv.gz | wc -1
time wc -1 ./labevents.csv
 158374765
real
        0m34.502s
        0m49.585s
user
```

sys 0m2.878s 158374765 ./labevents.csv real 0m37.933s user 0m33.418s sys 0m4.137s

The compressed file size is smaller than the uncompressed file size. The run time of zcat < ~/mimic/labevents.csv.gz | wc -l is faster than wc -l ~/mimic/labevents.csv. The trade off between storage and speed for big data files is that compressed files save storage space but take longer time to read and write, while uncompressed files take more storage space but are faster to read and write.

Q4. Who's popular in Price and Prejudice

1. You and your friend just have finished reading *Pride and Prejudice* by Jane Austen. Among the four main characters in the book, Elizabeth, Jane, Lydia, and Darcy, your friend thinks that Darcy was the most mentioned. You, however, are certain it was

Elizabeth. Obtain the full text of the novel from http://www.gutenberg.org/cache/epub/42671/pg42671.txt and save to your local folder.

```
wget -nc http://www.gutenberg.org/cache/epub/42671/pg42671.txt
```

Explain what wget -nc does. Do not put this text file pg42671.txt in Git. Complete the following loop to tabulate the number of times each of the four characters is mentioned using Linux commands.

Solution: wget -nc is used to download the file only if it does not exist in the current directory.

```
wget -nc http://www.gutenberg.org/cache/epub/42671/pg42671.txt

for char in Elizabeth Jane Lydia Darcy
do
    echo $char:
    # some bash commands here
    grep -oi "$char" pg42671.txt | wc -l
done
```

2. What's the difference between the following two commands?

```
echo 'hello, world' > test1.txt
```

and

```
echo 'hello, world' >> test2.txt
```

Solution: The difference of the two commands is that the first command echo 'hello, world' > test1.txt writes the output to a file test1.txt and overwrites the file if it already exists, while the second command echo 'hello, world' >> test2.txt appends the output to a file test2.txt if the file already exists. test2.txt will have the content hello, world twice if I run the second command twice.

3. Using your favorite text editor (e.g., vi), type the following and save the file as middle.sh:

```
#!/bin/sh
# Select lines from the middle of a file.
# Usage: bash middle.sh filename end_line num_lines
head -n "$2" "$1" | tail -n "$3"
```

Using chmod to make the file executable by the owner, and run

chmod +x middle.sh

```
./middle.sh pg42671.txt 20 5
```

Explain the output. Explain the meaning of "\$1", "\$2", and "\$3" in this shell script. Why do we need the first line of the shell script?

Solution: The meaning of "\$1", "\$2", and "\$3" in this shell script is the first, second, and third argument passed to the script, respectively. The first line of the shell script #!/bin/sh is called a shebang line, which tells the system which interpreter to use to execute the script. In this case, it tells the system to use the sh interpreter to execute the script.

Q5. More fun with Linux

Try following commands in Bash and interpret the results: cal, cal 2025, cal 9 1752 (anything unusual?), date, hostname, arch, uname -a, uptime, who am i, who, w, id, last | head, echo {con,pre}{sent,fer}{s,ed}, time sleep 5, history | tail.

Solution

cal

cal 2025

2025 January February March Su Mo Tu We Th Fr Sa Su Mo Tu We Th Fr Sa Su Mo Tu We Th Fr Sa 2 3 1 7 8 9 10 11 5 6 7 3 4 5 6 12 13 14 15 16 17 18 9 10 11 12 13 14 15 9 10 11 12 13 14 15

```
    19
    20
    21
    22
    23
    24
    25
    16
    17
    18
    19
    20
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    22
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    18
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    31
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    28
    29
```

April									1	¶ay			June							
Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa	Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa	Su	Мо	Tu	We	Th	Fr	Sa
		1	2	3	4	5					1	2	3	1	2	3	4	5	6	7
6	7	8	9	10	11	12	4	5	6	7	8	9	10	8	9	10	11	12	13	14
13	14	15	16	17	18	19	11	12	13	14	15	16	17	15	16	17	18	19	20	21
20	21	22	23	24	25	26	18	19	20	21	22	23	24	22	23	24	25	26	27	28
27	28	29	30				25	26	27	28	29	30	31	29	30					

July									Αι	ıgus	st		September							
Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa	Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa	Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa
		1	2	3	4	5						1	2		1	2	3	4	5	6
6	7	8	9	10	11	12	3	4	5	6	7	8	9	7	8	9	10	11	12	13
13	14	15	16	17	18	19	10	11	12	13	14	15	16	14	15	16	17	18	19	20
20	21	22	23	24	25	26	17	18	19	20	21	22	23	21	22	23	24	25	26	27
27	28	29	30	31			24	25	26	27	28	29	30	28	29	30				
							31													

October									Nov	zem l	oer		December							
Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa	Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa	Su	Мо	Tu	We	Th	$\operatorname{\mathtt{Fr}}$	Sa
			1	2	3	4							1		1	2	3	4	5	6
5	6	7	8	9	10	11	2	3	4	5	6	7	8	7	8	9	10	11	12	13
12	13	14	15	16	17	18	9	10	11	12	13	14	15	14	15	16	17	18	19	20
19	20	21	22	23	24	25	16	17	18	19	20	21	22	21	22	23	24	25	26	27
26	27	28	29	30	31		23	24	25	26	27	28	29	28	29	30	31			
							30													

cal 9 1752

Unusual thing: The days from 3rd to 13th on Sepetember of 1752 in the calnedar are missing.

date

Fri Jan 24 14:30:27 PST 2025

hostname

MacBookPro.lan

arch

i386

uname -a

Darwin MacBookPro.lan 24.1.0 Darwin Kernel Version 24.1.0: Thu Oct 10 21:02:27 PDT 2024; roo

uptime

14:30 up 11:27, 1 user, load averages: 2.68 2.80 2.62

who am i

vickey Jan 24 14:30

who

vickey console Jan 24 03:04

W

14:30 up 11:27, 1 user, load averages: 2.68 2.80 2.62

USER TTY FROM LOGIN@ IDLE WHAT vickey console - 3:04 11:25 -

```
id
```

uid=501(vickey) gid=20(staff) groups=20(staff),101(access_bpf),12(everyone),61(localaccounts

```
last | head
```

wtmp begins Fri Jan 24 14:30:27 PST 2025

```
echo {con,pre}{sent,fer}{s,ed}
```

consents consented confers confered presents presented prefers prefered

```
time sleep 5
```

```
real 0m5.015s
user 0m0.003s
sys 0m0.005s
```

history | tail

9 brew install wget 10 history | tail 11 git branch 12 git check out main 13 git checkout main 14 ls 15 git branch 16 git checkout main 17 git merge develop 18 history | tail

Q6. Book

- 1. Git clone the repository https://github.com/christophergandrud/Rep-Res-Book for the book Reproducible Research with R and RStudio to your local machine. Do **not** put this repository within your homework repository biostat-203b-2025-winter.
- 2. Open the project by clicking rep-res-3rd-edition.Rproj and compile the book by clicking Build Book in the Build panel of RStudio. (Hint: I was able to build git_book and epub_book directly. For pdf_book, I needed to add a line \usepackage{hyperref} to the file Rep-Res-Book/rep-res-3rd-edition/latex/preabmle.tex.)

The point of this exercise is (1) to obtain the book for free and (2) to see an example how a complicated project such as a book can be organized in a reproducible way. Use sudo apt install PKGNAME to install required Ubuntu packages and tlmgr install PKGNAME to install missing TexLive packages.

For grading purpose, include a screenshot of Section 4.1.5 of the book here.

Solution: Here is the screenshot of Section 4.1.5 of the book.

4.1.5 Spaces in directory and file names

It is good practice to avoid putting spaces in your file and directory names. For example, I called the example project parent directory in Figure 4.1 "example-project" rather than "Example Project". Spaces in file and directory names can sometimes create problems for computer programs trying to read the file path. The program may believe that the space indicates that the path name has ended. To make multi-word names easily readable without using spaces, adopt a consistent naming convention.

One approach is to use a convention that contrasts with the R object naming convention you are using. A contrasting convention helps make it clear if something is an R object or a file name. For example, if we adopt the underscore method for R object names used in Chapter 3 (e.g. health_data) we could use hyphens (-) to separate words in file names. For example: example-source.R. This is sometimes called kebab-case.