# Bios 6301: Assignment 8

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Due Tuesday, 16 November, 1:00 PM

 $5^{n=day}$  points taken off for each day late.

30 points total.

Submit a single knitr file (named homework8.rmd), along with a valid PDF output file. Inside the file, clearly indicate which parts of your responses go with which problems (you may use the original homework document as a template). Add your name as author to the file's metadata section. Raw R code/output or word processor files are not acceptable.

Failure to name file homework8.rmd or include author name may result in 5 points taken off.

#### Question 1

#### 15 points

Install the readxl package and run the following

```
library(readxl)
fn <- 'icd10.xlsx'
if(file.access(fn, mode = 4) == -1) {
    url <- "https://www.cdc.gov/nhsn/xls/icd10-pcs-pcm-nhsn-opc.xlsx"
    download.file(url, destfile = fn, mode = 'wb')
}
dat <- readxl::read_excel(fn, sheet = 2)</pre>
```

1. Show the class of dat. (1 point)

class(dat)

```
## [1] "tbl df" "tbl" "data.frame"
```

2. Show the methods available for objects of the given class (if there are multiple classes, show methods for all classes). (3 points)

```
methods(class = 'data.frame')
```

```
[[<-
                                                                   $<-
    [1] [
                       [6] aggregate
                       anyDuplicated anyNA
                                                    as.data.frame as.list
## [11] as.matrix
                      by
                                      cbind
                                                    coerce
                                                                   dim
## [16] dimnames
                       dimnames<-
                                     droplevels
                                                    duplicated
                                                                   edit
## [21] format
                       formula
                                     head
                                                    initialize
                                                                   is.na
## [26] Math
                       merge
                                     na.exclude
                                                    na.omit
                                                                   Ops
                                                                   row.names
## [31] plot
                      print
                                     prompt
                                                    rbind
## [36] row.names<-
                       rowsum
                                      show
                                                    slotsFromS3
                                                                   split
## [41] split<-
                       stack
                                     str
                                                    subset
                                                                   summary
## [46] Summary
                                     tail
                                                    transform
                                                                   type.convert
## [51] unique
                                     within
                       unstack
                                                    xtfrm
## see '?methods' for accessing help and source code
```

```
methods(class = 'tbl')
##
    [1] [[<-
                      [<-
                                   $<-
                                                                           initialize
                                                 coerce
                                                              format
##
    [7] Ops
                      print
                                   show
                                                 slotsFromS3
## see '?methods' for accessing help and source code
methods(class = 'tbl_df')
                                        [[<-
    [1] [
##
    [6] $<-
                        as.data.frame coerce
                                                       initialize
                                                                       names<-
## [11] Ops
                        row.names<-
                                        show
                                                       slotsFromS3
                                                                       str
## see '?methods' for accessing help and source code
  3. If you call print(dat), what print method is being dispatched? (1 point)
The print method will dispatch the print under tbl.
  4. Set the class of dat to be a data.frame. (1 point)
class(dat) <- c('data.frame')</pre>
  5. If you call print(dat) again, what print method is being dispatched? (1 point)
This time, the print method will dispatch the print under data.frame.
Define a new generic function nUnique with the code below.
nUnique <- function(x) {
    UseMethod('nUnique')
  6. Write a default method for nUnique to count the number of unique values in an element. (2 points)
nUnique <- function(x) {</pre>
  length(unique(x))
  7. Check your function (2 points)
nUnique(letters) # should return 26
nUnique(sample(10, 100, replace = TRUE)) # should return 10 (probably)
  8. Write a data frame method for nUnique to operate on data frame objects. This version should return
     counts for each column in a data.frame. (2 points)
nUnique.data.frame <- function(x) {</pre>
  output <- matrix(NA, nrow = 1, ncol = ncol(x))</pre>
  for(i in seq(ncol(x))){
    output[,i] <- length(unique(x[,i]))</pre>
  colnames(output) <- names(x)</pre>
  output
}
  9. Check your function (2 points)
nUnique.data.frame(dat)
```

#### Question 2

15 points

Programming with classes. The following function will generate random patient information.

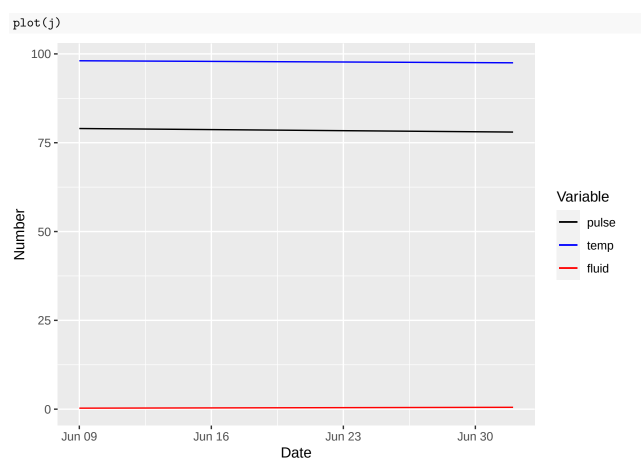
```
makePatient <- function() {
  vowel <- grep("[aeiou]", letters)
  cons <- grep("[^aeiou]", letters)
  name <- paste(sample(LETTERS[cons], 1), sample(letters[vowel], 1), sample(letters[cons], 1), sep='')
  gender <- factor(sample(0:1, 1), levels=0:1, labels=c('female', 'male'))
  dob <- as.Date(sample(7500, 1), origin="1970-01-01")
  n <- sample(6, 1)
  doa <- as.Date(sample(1500, n), origin="2010-01-01")
  pulse <- round(rnorm(n, 80, 10))
  temp <- round(rnorm(n, 98.4, 0.3), 2)
  fluid <- round(runif(n), 2)
  list(name, gender, dob, doa, pulse, temp, fluid)
}</pre>
```

1. Create an S3 class medicalRecord for objects that are a list with the named elements name, gender, date\_of\_birth, date\_of\_admission, pulse, temperature, fluid\_intake. Note that an individual patient may have multiple measurements for some measurements. Set the RNG seed to 8 and create a medical record by taking the output of makePatient. Print the medical record, and print the class of the medical record. (5 points)

```
set.seed(8)
j <- makePatient()</pre>
class(j) <- 'medicalRecord'</pre>
print(j)
## [[1]]
## [1] "Yes"
##
## [[2]]
## [1] male
## Levels: female male
## [[3]]
## [1] "1977-05-03"
##
## [[4]]
## [1] "2013-06-09" "2013-07-02"
##
## [[5]]
## [1] 79 78
##
## [[6]]
## [1] 98.07 97.50
##
## [[7]]
## [1] 0.28 0.52
## attr(,"class")
## [1] "medicalRecord"
```

2. Write a medicalRecord method for the generic function mean, which returns averages for pulse, temperature and fluids. Also write a medicalRecord method for print, which employs some nice formatting, perhaps arranging measurements by date, and plot, that generates a composite plot of measurements over time. Call each function for the medical record created in part 1. (5 points)

```
mean.medicalRecord <- function(x){</pre>
  output <- matrix(NA, nrow = 1, ncol = 3)</pre>
  x \leftarrow x[c(5,6,7)]
  for(i in 1:3){
    output[,i] <- sapply(x[i], mean, na.rm = TRUE)</pre>
  colnames(output) <- c("pulse", "temperature", "fluids")</pre>
  output
}
print.medicalRecord <- function(x){</pre>
  if(x[2] == 2) \{gender = "male"\}
  else{gender = "female"}
  dob = as.Date(as.integer(as.character.POSIXt(x[3])), origin="2010-01-01")
  four = sapply(x[4], as.character.POSIXt)
  four = sort(four)
  four = paste(four, collapse = ", ")
  cat(sprintf("name: %s\ngender: %s\ndoa: %s\npulse: %s\ntemp: %s\nfluid: %s",
              x[1], gender, dob, four, substr(x[5],3,nchar(x[5]) - 1), substr(x[6],3,nchar(x[6]) - 1),
}
library(ggplot2)
plot.medicalRecord <- function(x){</pre>
  dates = sapply(x[4], as.character.POSIXt)
  dates = sapply(dates, as.Date)
  data <- data.frame(</pre>
    doa = as.Date(dates, origin="2010-01-01"),
    pulse = x[5],
    temp = x[6],
    fluid = x[7]
  colnames(data) <- c("doa", "pulse", "temp", "fluid")</pre>
  ggplot(data) + geom_line(aes(x = doa, y = pulse, color = "black")) + geom_line(aes(x = doa, y = temp,
                           breaks = c("black", "blue", "red"),
                           labels = c("pulse", "temp", "fluid"),
                           guide = "legend") + xlab("Date") + ylab("Number")
}
mean(j)
        pulse temperature fluids
## [1,] 78.5
                   97.785
print(j)
## name: Yes
## gender: male
## dob: 2017-05-03
## doa: 2013-06-09, 2013-07-02
## pulse: 79, 78
## temp: 98.07, 97.5
## fluid: 0.28, 0.52
```



3. Create a further class for a cohort (group) of patients, and write methods for mean and print which, when applied to a cohort, apply mean or print to each patient contained in the cohort. Hint: think of this as a "container" for patients. Reset the RNG seed to 8 and create a cohort of ten patients, then show the output for mean and print. (5 points)

```
set.seed(8)
cohort <- function(n){</pre>
  result = makePatient()
  for(i in 1:(n-1)){
    result = c(result, makePatient())
  }
  result
}
j2 <- cohort(10)
class(j2) <- 'cohortRecord'</pre>
mean.cohortRecord <- function(x){</pre>
  output \leftarrow matrix(NA, nrow = length(x)/7, ncol = 3)
  pulse = x[seq(5, length(x), 7)]
  temp = x[seq(6, length(x), 7)]
  fluids = x[seq(7, length(x), 7)]
  for(i in 1:as.integer(length(x)/7)){
    output[i,1] <- sapply(pulse[i], mean, na.rm = TRUE)</pre>
    output[i,2] <- sapply(temp[i], mean, na.rm = TRUE)</pre>
    output[i,3] <- sapply(fluids[i], mean, na.rm = TRUE)</pre>
```

```
colnames(output) <- c("pulse", "temperature", "fluids")</pre>
  output
}
print.cohortRecord <- function(x){</pre>
  count = 1
  for(i in 1:length(x)){
   if(count ==1){
      print(paste("name: ", x[i]))
      count = count + 1
   } else if (count ==2) {
      if(x[i] == 2) {print("gender: male ")}
      else{print("gender: female ")}
      count = count + 1
   } else if (count ==3){
      dob = as.Date(as.integer(as.character.POSIXt(x[i])), origin="2010-01-01")
      print(paste("dob: ", dob))
      count = count + 1
   } else if (count ==4){
      four = sapply(x[i], as.character.POSIXt)
      four = sort(four)
      four = paste(four, collapse = ", ")
      print(paste("doa: ", four))
      count = count + 1
   } else if (count ==5){
      print(paste("pulse: ", substr(x[i],3,nchar(x[i]) - 1)))
      count = count + 1
   } else if (count ==6){
      print(paste("temp: ", substr(x[i],3,nchar(x[i]) - 1)))
      count = count + 1
   } else if (count ==7){
      print(paste("fluid: ", substr(x[i],3,nchar(x[i]) - 1)))
      count = 1
      cat("\n")
   }
 }
}
mean.cohortRecord(j2)
##
                                 fluids
            pulse temperature
## [1,] 78.50000
                     97.78500 0.4000000
## [2,] 86.33333
                     98.39667 0.4133333
## [3,] 77.00000
                     98.64750 0.5200000
## [4,] 83.16667
                     98.48500 0.2966667
## [5,] 83.50000
                     98.45000 0.4525000
## [6,] 84.40000
                     98.48400 0.5220000
## [7,] 76.50000
                     98.38000 0.3975000
## [8,] 75.00000
                     98.36750 0.5225000
## [9,] 73.00000
                     98.36000 0.1500000
```

98.54000 0.1500000

**##** [10,] 77.00000

### print.cohortRecord(j2)

```
## [1] "name: Yes"
## [1] "gender: male "
## [1] "dob: 2017-05-03"
## [1] "doa: 2013-06-09, 2013-07-02"
## [1] "pulse: 79, 78"
## [1] "temp: 98.07, 97.5"
## [1] "fluid: 0.28, 0.52"
##
## [1] "name: Fal"
## [1] "gender: male "
## [1] "dob: 2028-05-24"
## [1] "doa: 2010-11-16, 2013-03-24, 2013-09-12"
## [1] "pulse: 76, 96, 87"
## [1] "temp: 98.23, 98.75, 98.21"
## [1] "fluid: 0.18, 0.96, 0.1"
## [1] "name: Zog"
## [1] "gender: male "
## [1] "dob: 2028-12-14"
## [1] "doa: 2010-02-24, 2013-03-25, 2013-07-29, 2013-10-27"
## [1] "pulse: 69, 75, 80, 84"
## [1] "temp: 98.49, 98.82, 98.74, 98.54"
## [1] "fluid: 0.81, 0.59, 0.28, 0.4"
##
## [1] "name: Yol"
## [1] "gender: male "
## [1] "dob: 2026-03-11"
## [1] "doa: 2010-02-22, 2011-12-27, 2012-03-10, 2012-11-26, 2013-03-24, 2014-01-28"
## [1] "pulse: 69, 78, 87, 84, 89, 92"
## [1] "temp: 98.29, 98.44, 98.78, 98.87, 98.27, 98.26"
## [1] "fluid: 0.03, 0.13, 0.12, 0.39, 0.97, 0.14"
##
## [1] "name: Yak"
## [1] "gender: female "
## [1] "dob: 2023-09-15"
## [1] "doa: 2011-07-19, 2012-04-07, 2012-07-11, 2012-08-30"
## [1] "pulse: 90, 88, 75, 81"
## [1] "temp: 98.58, 97.53, 98.58, 99.11"
## [1] "fluid: 0.26, 0.29, 0.6, 0.66"
##
## [1] "name: Gaf"
## [1] "gender: female "
## [1] "dob: 2018-04-27"
## [1] "doa: 2010-07-19, 2011-05-03, 2012-04-24, 2012-08-06, 2013-08-21"
## [1] "pulse: 89, 91, 77, 75, 90"
## [1] "temp: 98.32, 98.01, 98.96, 98.52, 98.61"
## [1] "fluid: 0.42, 0.47, 0.74, 0.62, 0.36"
##
## [1] "name: Kuw"
## [1] "gender: female "
## [1] "dob: 2020-11-07"
## [1] "doa: 2010-10-03, 2010-10-29, 2011-09-16, 2012-07-10"
```

```
## [1] "pulse: 72, 81, 71, 82"
## [1] "temp: 98.21, 98.17, 98.65, 98.49"
## [1] "fluid: 0.29, 0.93, 0.25, 0.12"
##
## [1] "name: Mav"
## [1] "gender: female "
## [1] "dob: 2029-07-16"
## [1] "doa: 2010-02-08, 2010-04-19, 2010-06-11, 2012-03-02"
## [1] "pulse: 63, 83, 66, 88"
## [1] "temp: 99.07, 98.45, 97.95, 98"
## [1] "fluid: 0.01, 0.79, 0.79, 0.5"
##
## [1] "name: Fel"
## [1] "gender: male "
## [1] "dob: 2025-08-16"
## [1] "doa: 2010-09-26, 2012-06-24"
## [1] "pulse: 65, 81"
## [1] "temp: 98.21, 98.51"
## [1] "fluid: 0.06, 0.24"
##
## [1] "name: Say"
## [1] "gender: female "
## [1] "dob: 2014-09-22"
## [1] "doa: 2010-03-14"
## [1] "pulse: "
## [1] "temp: .5"
## [1] "fluid: 1"
```