

## Week 1 Part 3

This file consists of exercises for the course

*HarvardX PH525.1x Statistics and R*

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*dplyr*

*Exercises #1*

Read in the `msleep_ggplot2.csv` file with the function `read.csv()` and use the function `class()` to determine what type of object is returned.

```
msleep <- read.csv('https://raw.githubusercontent.com/genomicsclass/dagdata/master/inst/extdata/msleep_ggplot2.csv')
class(msleep)

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

*Exercises #2*

Now use the `filter()` function to select only the primates.

How many animals in the table are primates? Hint: the `nrow()` function gives you the number of rows of a data frame or matrix.

```
f_primates <- dplyr::filter(msleep, order == 'Primates')
nrow(f_primates)

## [1] 12
```

*Exercises #3*

What is the class of the object you obtain after subsetting the table to only include primates?

```
class(f_primates)
```

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

*Exercises #4*

Now use the `select()` function to extract the sleep (total) for the primates.

What class is this object? Hint: use `%>%` to pipe the results of the `filter()` function to `select()`.

```
library(magrittr)

s_primates <- dplyr::filter(msleep, order == 'Primates') %>%
  dplyr::select(sleep_total)

class(s_primates)

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

### Exercises #5

Now we want to calculate the average amount of sleep for primates (the average of the numbers computed above). One challenge is that the `mean()` function requires a vector so, if we simply apply it to the output above, we get an error. Look at the help file for `unlist()` and use it to compute the desired average.

What is the average amount of sleep for primates?

```
s_primates <- unlist(s_primates)
mean(s_primates)
```

```
## [1] 10.5
```

### Exercises #6

For the last exercise, we could also use the `dplyr` `summarize()` function. We have not introduced this function, but you can read the help file and repeat exercise 5, this time using just `filter()` and `summarize()` to get the answer.

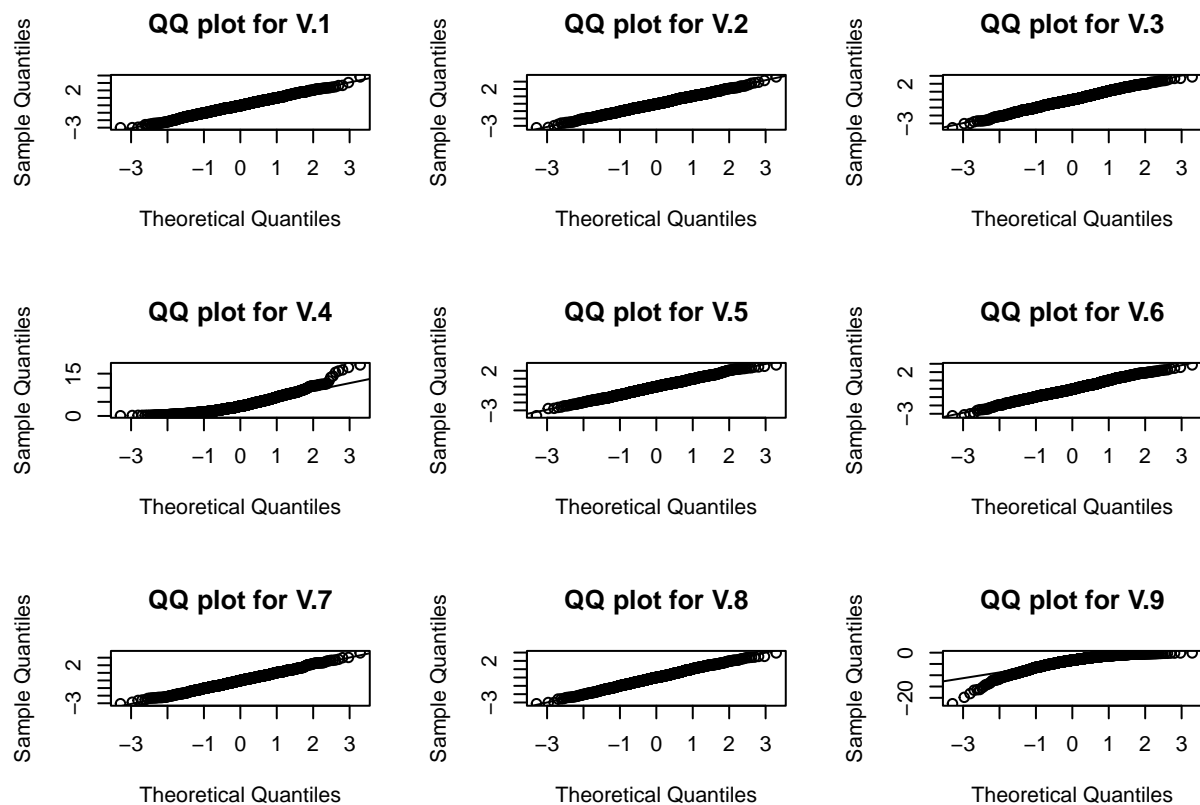
What is the average amount of sleep for primates calculated by `summarize()`

```
avg_sleep <- dplyr::filter(msleep,order=='Primates') %>%
  dplyr::summarise(avg_sleep=mean(sleep_total))
```

```
dat <- get(load(url("http://courses.edx.org/c4x/HarvardX/PH525.1x/asset/skew.RData")))
dim(dat)
```

```
## [1] 1000    9
```

```
par(mfrow = c(3,3))
for (k in 1:9) {
  x <- dat[,k]
  qqnorm(x, main=paste0('QQ plot for V.',k,sep=''))
  qqline(x)
}
```



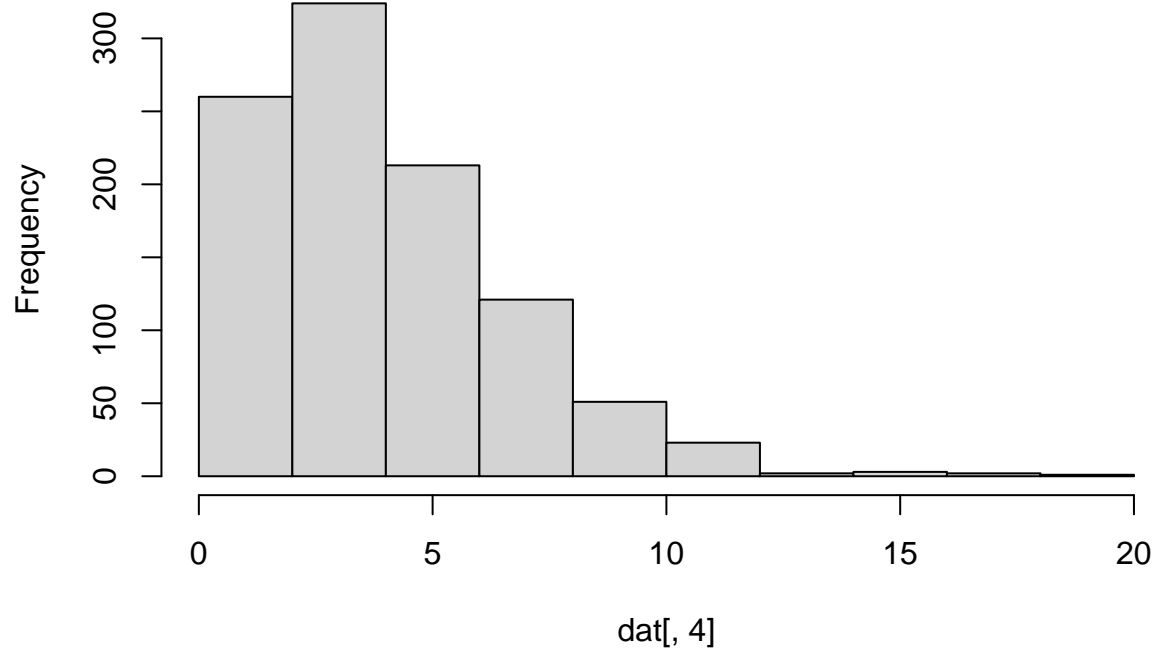
Identify the two columns which are skewed.

Examine each of these two columns using a histogram. Note which column has “positive skew”, in other words the histogram shows a long tail to the right (toward larger values). Note which column has “negative skew”, that is, a long tail to the left (toward smaller values). Note that positive skew looks like an up-shaping curve in a `qqnorm()` plot, while negative skew looks like a down-shaping curve.

You can use the following line to reset your graph to just show one at a time: `par(mfrow=c(1,1))`

```
hist(dat[,4])
```

**Histogram of dat[, 4]**



```
hist(dat[,9])
```

**Histogram of dat[, 9]**

