IS5 in R: The Standard Deviation as a Ruler and the Normal Model (Chapter 5)

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Introduction and background

This document is intended to help describe how to undertake analyses introduced as examples in the Fifth Edition of *Intro Stats* (2018) by De Veaux, Velleman, and Bock. More information about the book can be found at http://wps.aw.com/aw_deveaux_stats_series. This file as well as the associated R Markdown reproducible analysis source file used to create it can be found at http://nhorton.people.amherst.edu/is5.

This work leverages initiatives undertaken by Project MOSAIC (http://www.mosaic-web.org), an NSF-funded effort to improve the teaching of statistics, calculus, science and computing in the undergraduate curriculum. In particular, we utilize the mosaic package, which was written to simplify the use of R for introductory statistics courses. A short summary of the R needed to teach introductory statistics can be found in the mosaic package vignettes (http://cran.r-project.org/web/packages/mosaic). A paper describing the mosaic approach was published in the R Journal: https://journal.r-project.org/archive/2017/RJ-2017-024.

Chapter 5: The Standard Deviation as a Ruler and the Normal Model

```
library(mosaic)
library(readr)
library(janitor)
WomenHeptathlon2016 <-
  read csv("http://nhorton.people.amherst.edu/is5/data/Womens Heptathlon 2016.csv") %>%
  clean names()
## Parsed with column specification:
## cols(
##
     `First Name` = col_character(),
##
     `Last Name` = col_character(),
##
     200m = col_double(),
##
     LongJump = col_double(),
##
     `800m` = col_double(),
##
     HighJump = col_double(),
     `100m.hurdles` = col_double(),
##
##
     Javelin = col_double(),
##
     ShotPut = col_double()
## )
```

By default, read_csv() prints the variable names. These messages can be suppressed using the message = FALSE code chunk option to save space and improve readability.

Here we use the clean_names() function from the janitor package to sanitize the names of the columns (which would otherwise contain special characters or whitespace).

```
# page 123
favstats(~ long_jump, data = WomenHeptathlon2016)
## min Q1 median Q3 max mean sd n missing
```

```
## 5.51 6.08 6.19 6.31 6.58 6.169655 0.2474655 29
favstats(~ x200m, data = WomenHeptathlon2016)
##
             Q1 median
      min
                          Q3 max
                                       mean
                                                   sd n missing
                 24.6 24.99 26.32 24.58207 0.6544975 29
   23.26 24.12
with(WomenHeptathlon2016, stem(x200m))
##
##
     The decimal point is at the |
##
     23 | 3
##
##
     23 | 589
##
     24 | 011123334
##
     24 | 5667789
     25 | 00112444
##
##
     25 I
     26 | 3
##
with(WomenHeptathlon2016, stem(long_jump))
##
##
     The decimal point is 1 digit(s) to the left of the |
##
##
     54 | 1
     56 | 2
##
     58 | 181
##
##
     60 | 0588002569
    62 | 023501145
##
     64 | 38158
##
Section 5.1: Using the Standard Deviation to Standardize Values
filter(WomenHeptathlon2016, last_name == "Thiam") %>%
data.frame()
    first_name last_name x200m long_jump x800m high_jump x100m_hurdles
## 1 Nafissatou
                   Thiam 25.1
                                     6.58 136.54
                                                      1.98
    javelin shot_put
      53.13
               14.91
\# calculate z-score with mean and sd from favstats
(6.58 - 6.17)/.247 # long jump
## [1] 1.659919
filter(WomenHeptathlon2016, last_name == "Johnson-Thompson") %>%
data.frame()
##
    first name
                       last_name x200m long_jump x800m high_jump
      Katarina Johnson-Thompson 23.26
                                        6.51 130.47
   x100m_hurdles javelin shot_put
## 1
             13.48
                   36.36
                              11.68
```

data.frame() converts an object into a data frame.

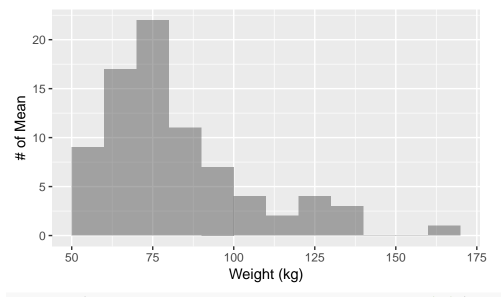
Section 5.2: Shifting and Scaling

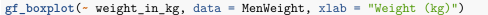
Shifting to Adjust the Center

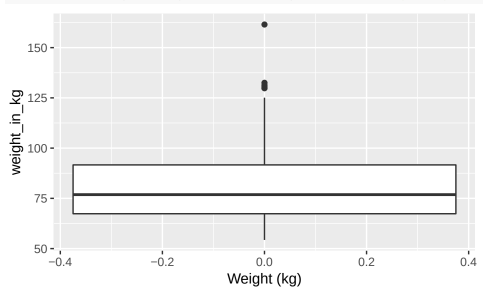
```
MenWeight <- read_csv("http://nhorton.people.amherst.edu/is5/data/Mens_Weights.csv") %>%
    clean_names()
```

```
## Parsed with column specification:
## cols(
## `Weight in kg` = col_double(),
## `Weight in pounds` = col_double()
## )
```

```
# Figure 5.2, page 125
gf_histogram(~ weight_in_kg, data = MenWeight, binwidth = 10, center = 5) %>%
gf_labs(x = "Weight (kg)", y = "# of Mean")
```







```
favstats(~ weight_in_kg, data = MenWeight)
            Q1 median
                          QЗ
                               max
                                        mean
                                                    sd n missing
## 54.3 67.35 76.85 91.65 161.5 82.35625 22.26881 80
# Figure 5.3
gf_histogram(~ (weight_in_kg - 74), data = MenWeight, binwidth = 10) %>%
gf_labs(x = "Kg Above Recommended Weight", y = "# of Men")
   20 -
   15 -
# of Men
   10-
    5 -
    0 -
                                 25
                                             50
                                                          75
       -<del>2</del>5
                                                                      100
                      Kg Above Recommended Weight
```

Rescaling to Adjust the Scale

95.7

68.9

60.3

60.4

69.7

2

3

4

5

6

```
favstats(~ weight_in_kg, data = MenWeight)
    min
            Q1 median
                         QЗ
                              max
                                      mean
                                                 sd n missing
## 54.3 67.35 76.85 91.65 161.5 82.35625 22.26881 80
favstats(~ weight_in_pounds, data = MenWeight)
##
       min
               Q1 median
                             QЗ
                                  max
                                          mean
                                                     sd n missing
## 119.46 148.17 169.07 201.63 355.3 181.1838 48.99137 80
library(tidyr) # for gather() function
# What does gather() do?
MenWeight %>%
  head() # There are two variables: weight_in_kg and weight_in_pounds. Each observation has a value for
## # A tibble: 6 x 2
##
    weight_in_kg weight_in_pounds
##
            <dbl>
                             <dbl>
            107.
                              236.
## 1
```

211.

152.

133.

133.

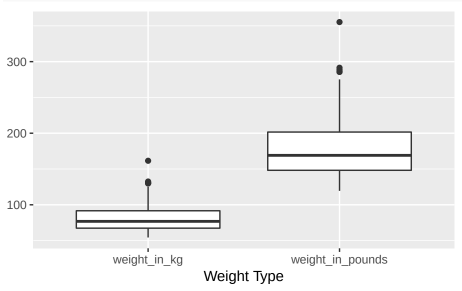
153.

```
nrow(MenWeight)
## [1] 80
MenGather <- MenWeight %>%
  gather(key = weighttype, value = weight, weight_in_kg, weight_in_pounds)
MenGather %>%
  head() # The two variables are weighttype and weight, weighttype is a categorical variable that is ei
## # A tibble: 6 x 2
##
     weighttype weight
##
     <chr>
                   <dbl>
## 1 weight_in_kg 107.
## 2 weight_in_kg
                   95.7
## 3 weight_in_kg
                    68.9
                    60.3
## 4 weight_in_kg
## 5 weight_in_kg
                    60.4
## 6 weight_in_kg
                    69.7
nrow(MenGather) # Each observation from before is now two rows
```

[1] 160

Here we use the gather() function to transform the dataset into the needed format, which can be seen with the head() function.

```
MenGather %>%
  gf_boxplot(weight ~ weighttype) %>%
  gf_labs(x = "Weight Type", y = "")
```



We see the use of goal(Y ~ X) as an example of the general modeling language for two variables in the mosaic package.

Shifting, Scaling, and the z-Scores

Section 5.3: Normal Models

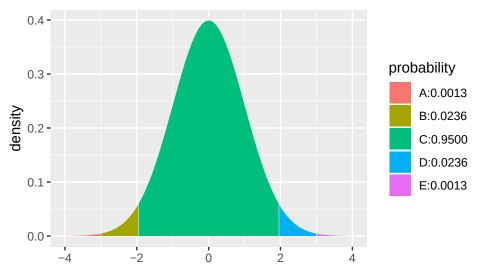
The 68-95-99.7 Rule

See display on page 129.

```
# Figure 5.6
# 1, 2 (1.96), and 3 SD's
xpnorm(c(-3, -1.96, -1, 1, 1.96, 3), mean = 0, sd = 1, verbose = FALSE)
   0.4 -
                                                             probability
                                                                  A:0.0013
   0.3 -
                                                                  B:0.0236
density
                                                                  C:0.1337
                                                                  D:0.6827
                                                                  E:0.1337
   0.1 -
                                                                  F:0.0236
                                                                  G:0.0013
   0.0 -
                   -2
                               0
                                           2
```

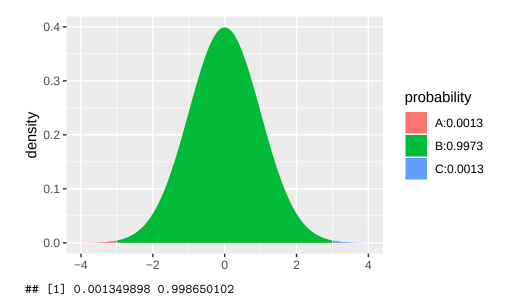
[1] 0.001349898 0.024997895 0.158655254 0.841344746 0.975002105 0.998650102

```
# 2 (1.96) and 3 SD's
xpnorm(c(-3, -1.96, 1.96, 3), mean = 0, sd = 1, verbose = FALSE)
```



[1] 0.001349898 0.024997895 0.975002105 0.998650102

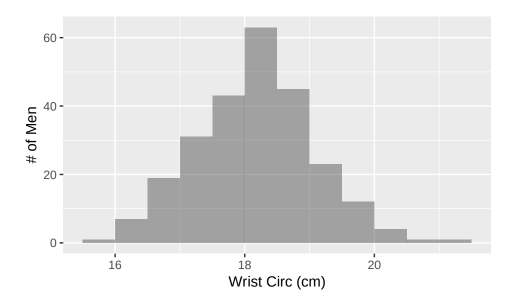
```
# 3 SD's
xpnorm(c(-3, 3), mean = 0, sd = 1, verbose = FALSE)
```



Example 5.4: Using the 68-95-99.7 Rule

```
BodyFat <- read_csv("http://nhorton.people.amherst.edu/is5/data/Bodyfat.csv")</pre>
```

```
## Parsed with column specification:
## cols(
     Density = col_double(),
##
     Pct.BF = col_double(),
##
##
     Age = col_integer(),
     Weight = col_double(),
##
##
     Height = col_double(),
##
     Neck = col_double(),
     Chest = col_double(),
##
     Abdomen = col_double(),
##
     Waist = col_double(),
##
##
     Hip = col_double(),
##
     Thigh = col_double(),
     Knee = col_double(),
##
##
     Ankle = col_double(),
##
     Bicep = col_double(),
##
     Forearm = col_double(),
##
     Wrist = col_double()
## )
gf_histogram(~ Wrist, data = BodyFat, binwidth = .5,
             center = -.25) %>%
 gf_labs(x = "Wrist Circ (cm)", y = "# of Men")
```



Random Matters

```
Starts on page 133.
```

```
read_csv("http://nhorton.people.amherst.edu/is5/data/Population_Commute_Times.csv") %>%
  clean_names()
## Parsed with column specification:
     Commute.Time = col_integer()
##
## )
gf_histogram(~ commute_time, data = Commute, binwidth = 10, center = 5) %>%
  gf_labs(x = "Commute Times (min)", y = "# of Employees")
   1000 -
    750 -
# of Employees
    500 -
    250 -
      0 -
                        50
                                     100
                                                                200
           0
                                                   150
                             Commute Times (min)
```

```
# What does do() do? XX Does this comment have to be here?
mean(~ commute_time, data = sample(Commute, size = 100)) # Mean of one random sample

## [1] 46.77
mean(~ commute_time, data = sample(Commute, size = 100)) # Mean of another random sample

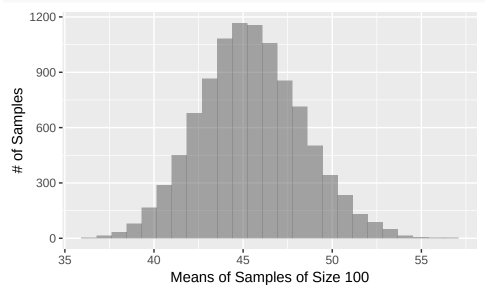
## [1] 42.07
do(2) * mean(~ commute_time, data = sample(Commute, size = 100)) # Carries out mean() twice

## mean
## 1 42.19
## 2 42.90

# For the visualization, we use do() 10,000 times
Commute_sample <- do(numsim) * mean(~ commute_time, data = sample(Commute, size = 100))

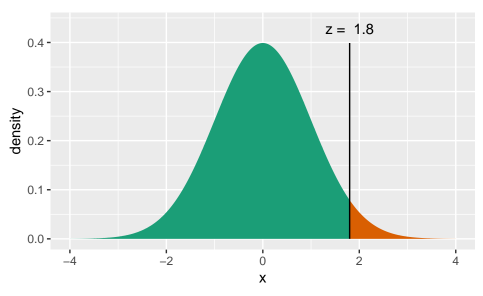
The do() function runs, 10,000 times, the mean and the sampling command on a random sample of 100.</pre>
```

```
gf_histogram(~ mean, data = Commute_sample) %>%
    gf_labs(x = "Means of Samples of Size 100", y = "# of Samples")
```



Section 5.4: Working with Normal Percentiles

```
xpnorm(1.8, mean = 0, sd = 1)
##
## If X ~ N(0, 1), then
## P(X <= 1.8) = P(Z <= 1.8) = 0.9641
## P(X > 1.8) = P(Z > 1.8) = 0.03593
##
```



[1] 0.9640697

The qnorm() function:

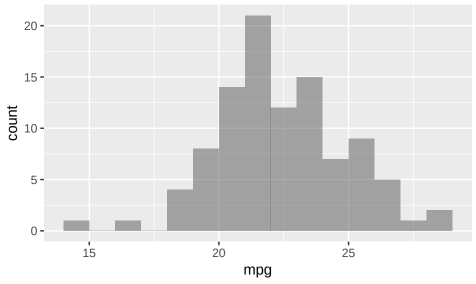
```
qnorm(0.964, mean = 500, sd = 100) # inverse of pnorm()
## [1] 679.9118
qnorm(0.964, mean = 0, sd = 1) # what is the z-score?
```

[1] 1.799118

See examples on pages 136-140.

Section 5.5: Normal Probability Plots

```
Nissan <- read_csv("http://nhorton.people.amherst.edu/is5/data/Nissan.csv")
## Parsed with column specification:
## cols(
## mpg = col_double()
## )
# Figure 5.10, page 141
gf_histogram(~ mpg, data = Nissan, binwidth = 1, center = .5)</pre>
```



gf_qq(~ mpg, data = Nissan, xlab = "Normal Scores") %>%
gf_qqline(linetype = "solid", color = "red")

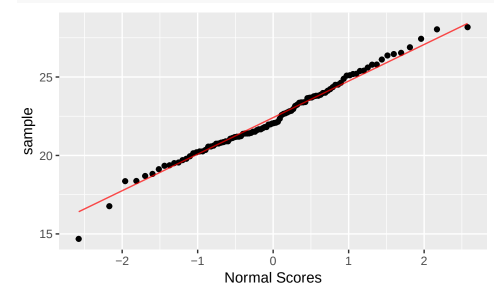


Figure 5.11
gf_histogram(~ weight_in_kg, data = MenWeight, xlab = "Weights", binwidth = 10, center = 5)

