IS5 in R: Sampling Distribution Models and Confidence Intervals for Proportions (Chapter 13)

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Introduction and background

sex = col_character(),

weight = col_integer(),

orig.id = col_integer(),
preemie = col_logical()

prenatalstart = col integer(),

##

##

##

)

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.

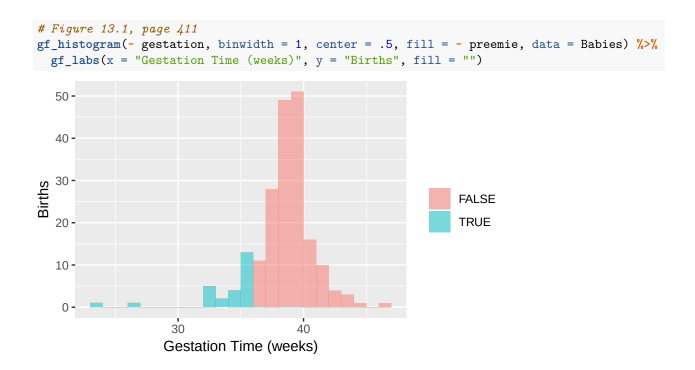
This work leverages initiatives undertaken by Project MOSAIC (http://www.mosaic-web.org), an NSF-funded effort to improve the teaching of statistics, calculus, science and computing in the undergraduate curriculum. In particular, we utilize the mosaic package, which was written to simplify the use of 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 13: Sampling Distribution Models and Confidence Intervals for Proportions

```
library(mosaic)
library(readr)
library(janitor)
Babies <- read_csv("http://nhorton.people.amherst.edu/is5/data/Babysamp_98.csv") %>%
  clean_names()
## Parsed with column specification:
## cols(
##
     MomAge = col_integer(),
     DadAge = col_integer(),
##
     MomEduc = col_integer(),
##
##
    MomMarital = col integer(),
##
     numlive = col integer(),
##
     dobmm = col_integer(),
##
     gestation = col_integer(),
```

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).



Section 13.1: The Sampling Distribution Model for a Proportion

The Normal Model

Section 13.2: When Does the Normal Model Work? Assumptions and Conditions

Random Matters: Does the Normal Model Always Work? Sampling Distributions for Other Statistics

```
# page 418
BodyFat <- read_csv("http://nhorton.people.amherst.edu/is5/data/Bodyfat.csv") %>%
  clean_names()
## 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()
```

```
## )
set.seed(3245) # For reproducibility
numsim <- 1000 # Number of samples
# What does do() do?
favstats (~ weight, data = sample (BodyFat, 10)) # favstats of one random sample of 10
##
               Q1 median
                               QЗ
                                     max
                                             mean
                                                        sd n missing
    148.25 166.25 176.5 192.125 247.25 183.725 29.88925 10
favstats(~ weight, data = sample(BodyFat, 10)) # favstats of another random sample
##
      min
                Q1 median
                                 03
                                       max
                                               mean
                                                         sd n missing
   127.5 154.9375
                      161 189.4375 216.25 168.725 27.3563 10
do(2) * favstats(~ weight, data = sample(BodyFat, 10)) # finds favstats twice
                  Q1 median
##
        min
                                    QЗ
                                           max
                                                  mean
                                                             sd n missing .row
## 1 125.00 168.7500 188.875 208.8750 241.75 186.400 33.96367 10
                                                                          0
## 2 156.75 167.4375 179.875 189.5625 224.50 182.625 21.03610 10
     .index
## 1
## 2
# For the visualization, we need 1,000 favstats
bodyfatsamples <- do(numsim) * favstats(~ weight, data = sample(BodyFat, 10))
Here, the do() function finds, 1,000 times, the favstats() of a random sample of 10 BodyFat weights.
bodyfatsamples <- bodyfatsamples %>%
  clean_names()
names(bodyfatsamples)
                  "q1"
                                        "q3"
    [1] "min"
                                                  "max"
                                                                       "sd"
##
                             "median"
                                                            "mean"
    [8] "n"
                   "missing" "row"
                                        "index"
gf_histogram(~ median, data = bodyfatsamples, binwidth = 3, center = 1.5) %>%
 gf_labs(x = "Medians", y = "# of Samples")
   120 -
    90 -
# of Samples
    60 -
    30 -
                             170
                                       180
                                                 190
                                                           200
                  160
        150
                                  Medians
```

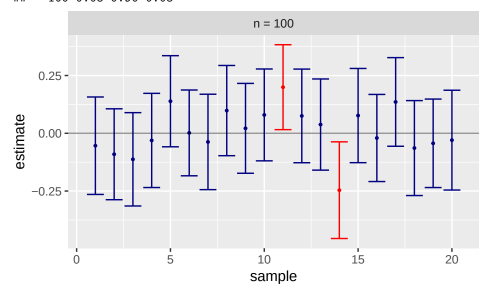
```
gf_histogram(~ sd^2, data = bodyfatsamples) %>%
  gf_labs(x = "Variance", y = "# of Samples")
   120 -
    90 -
# of Samples
    60 -
    30 -
     0
                        500
                                       1000
                                                       1500
                                                                       2000
                                     Variance
gf_histogram(~ min, data = bodyfatsamples, binwidth = 3, center = 1.5) %>%
  gf_labs(x = "Minimums", y = "# of Samples")
   120 -
# of Samples
    80 -
    40 -
     0 -
            120
                                 140
                                                      160
                                                                           180
                                     Minimums
```

Section 13.3: A Confidence Interval for a Proportion

Section 13.4: Interpreting Confidence Intervals: What Does 95% Confidence Really Mean?

```
# Here is a simulation of Figure 13.9 (page 422)
set.seed(118)
CIsim(n = 100, samples = 20) # We expect 19/20 intervals to cover the true mean
## Interval coverage:
## cover
## n Low Yes High
```

100 0.05 0.90 0.05



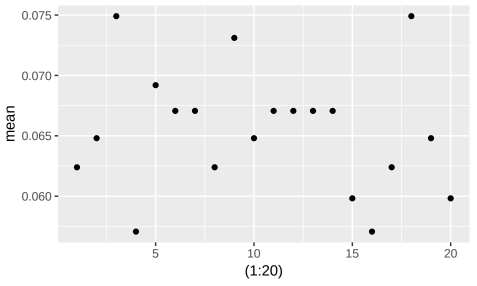
We expect 19 of the 20 intervals to cover the true mean, but since only 20 samples are drawn, there is more variability. Here, only 18 out of the 20 intervals cover the true mean.

To get the actual plot, the code is more complicated.

```
findingpoints <- function(sampsize, numsamp) {
   set.seed(2461)
   CIdata <- do(numsamp) * t.test(~ preemie, data = sample(Babies, size = sampsize))
   CIdata <- CIdata %>%
      select(lower, upper) %>%
      mutate(mean = (upper - lower)/2)
}

ConfData <- findingpoints(sampsize = 100, numsamp = 20)

gf_point(mean ~ (1:20), data = ConfData) #%>%
```



gf_line

Section 13.5: Margin of Error: Certainty vs. Precision

Section 13.6: Choosing the Sample Size