# IS5 in R: Testing Hypotheses (Chapter 15)

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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 15: Testing Hypotheses

```
library(mosaic)
library(readr)
library(janitor)
```

Section 15.1: Hypotheses

Section 15.2: P-Values

Section 15.3: The Reasoning of Hypothesis Testing

### Example 15.5: Finding A P-Value

```
n <- 90
x <- 61
p <- .8
phat <- x/n
sdphat <- ((p * (1 - p))/n)^.5
z <- (phat - p)/sdphat
pnorm(z)</pre>
```

```
## [1] 0.00187324
```

```
# Or, without calculating the z-score:
pnorm(q = phat, mean = p, sd = sdphat)
```

## [1] 0.00187324

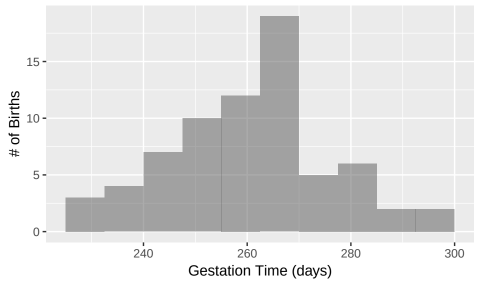
#### Section 15.4: A Hypothesis Test for the Mean

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

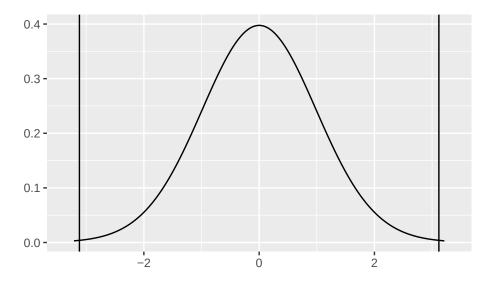
```
## Parsed with column specification:
## cols(
## Gestation = col_integer(),
## Time = col_character()
## )
```

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.

```
# 2. Model (page 482)
gf_histogram(~ Gestation, data = GestationTime, binwidth = 7.5, center = 3.75) %>%
gf_labs(x = "Gestation Time (days)", y = "# of Births")
```



```
# 3. Mechanics
gf_dist(dist = "t", df = 69) %>%
gf_vline(xintercept = -3.118) %>%
    gf_vline(xintercept = 3.118) %>%
gf_labs(x = "", y = "") +
    xlim(-3.347, 3.347)
```



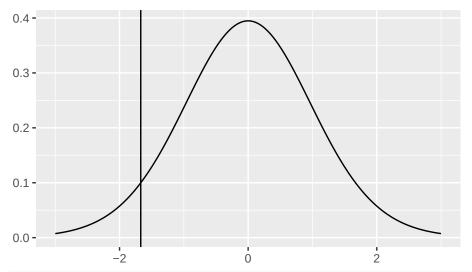
# Step-By-Step Example: A One-Sample t-Test for the Mean

gf\_vline(xintercept = -1.67) %>%

```
# page 485
Sleep <- read_csv("http://nhorton.people.amherst.edu/is5/data/Sleep.csv")</pre>
## Parsed with column specification:
## cols(
     Sleep = col_integer()
## )
# Plan
favstats(~ Sleep, data = Sleep)
## min Q1 median Q3 max mean
                                     sd n missing
                7 7 8 6.64 1.075484 25
      4 6
gf_histogram(~ Sleep, data = Sleep, binwidth = 1) %>%
 gf_labs(x = "Hours of Sleep", y = "")
8 -
6 -
4 -
2 -
0 -
                      5
                                                         8
                           Hours of Sleep
gf_dist(dist = "t", df = 24) %>%
```

```
gf_labs(x = "", y = "") + xlim(-3, 3)
```

## Warning: Removed 674 rows containing missing values (geom\_path).



```
# Mechanics
n <- 25
mean <- 7.0
df <- 24
y <- 6.64
s <- 1.075
sey <- s/(n^.5)
t <- (y - mean)/sey # t-statistic
pt(q = t, df = df) # p-value</pre>
```

## [1] 0.05351625

#### Section 15.5: Intervals and Tests

```
7.5 - 2.5 - 0.0 - 97 98 99 100 Temp
```

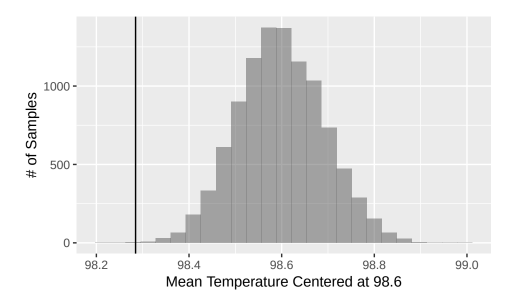
```
# Confidence interval
y <- mean(~ Temp, data = Temperatures)</pre>
У
## [1] 98.28462
s <- sd(~ Temp, data = Temperatures)</pre>
## [1] 0.6823789
n <- nrow(Temperatures)</pre>
## [1] 52
tstats <- qt(df = n - 1, p = c(.005, .995))
tstats
## [1] -2.675722 2.675722
y + (tstats * (s/(n^{\cdot}.5)))
## [1] 98.03141 98.53782
# Hypothesis test
mu <- 98.6
t <- (y - mu)/(s/(n^{.5}))
## [1] -3.332856
2 * pt(q = t, df = n - 1) # two sided test
```

# Random Matters: Bootstrap Hypothesis Tests and Intervals

## [1] 0.001605849

```
numsamp <- 10000
```

```
# What does do() do?
mean(~ Temp, data = resample(Temperatures)) # Mean of one random resample
## [1] 98.31538
mean(~ Temp, data = resample(Temperatures)) # Mean of another random resample
## [1] 98.30385
do(2) * mean(~ Temp, data = resample(Temperatures)) # Calculates means of two resamples
##
         mean
## 1 98.22692
## 2 98.37500
# We will use do() a numsamp number of times
resampletemps <- do(numsamp) * mean(~ Temp, data = resample(Temperatures))
For more information about resample(), refer to the resample vignette in mosaic.
gf_histogram(~ mean, data = resampletemps) %>%
  gf_labs(x = "Mean Temperature", y = "# of Samples")
   1000 -
of Samples
    500 -
      0 -
                                      98.3
                                                     98.5
         97.9
                        98.1
                                                                    98.7
                              Mean Temperature
qdata(~ mean, p = c(.005, .995), data = resampletemps) # reject null hypothesis
##
         quantile
## 0.5% 98.04808 0.005
## 99.5% 98.52886 0.995
# Making a model-centric distribution
Temperatures2 <- Temperatures %>%
  mutate(Temp = Temp + .315)
resampletemps2 <- do(numsamp) * mean(~ Temp, data = resample(Temperatures2))</pre>
gf_histogram(~ mean, data = resampletemps2) %>%
  gf_vline(xintercept = mean(~ Temp, data = Temperatures)) %>%
  gf_labs(x = "Mean Temperature Centered at 98.6", y = "# of Samples")
```



## Step-By-Step Example: Tests and Intervals

```
# Creating the data set
Baseball <- rbind(</pre>
  do(1308) * (winner = "HOME"),
  do(2431 - 1308) * (winner = "AWAY")
  rename(winner = result)
# Mechanics (page 490)
n <- nrow(Baseball)</pre>
p < -.5
phat <- Baseball %>%
  filter(winner == "HOME") %>%
  nrow()/n
phat
## [1] 0.5380502
sdphat <- ((p * (1 - p))/n)^.5
sdphat
## [1] 0.01014092
z <- (phat - p)/sdphat # z-value
## [1] 3.752142
1 - pnorm(z) # p-value
## [1] 8.76651e-05
# Or, without calculating the z-score:
1 - pnorm(q = phat, mean = p, sd = sdphat)
## [1] 8.76651e-05
# Mechanics (page 491)
sep <- ((phat * (1 - phat))/n)^.5
```

```
## [1] 0.01011152
me <- 1.96 * sep
phat - me # lower bound of 95% confidence
## [1] 0.5182316
phat + me # upper bound of 95% confidence
## [1] 0.5578688</pre>
```

Section 15.6: P-Values and Decisions: What to Tell About a Hypothesis Test