

IS5 in R: Paired Samples and Blocks (Chapter 18)

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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 18: Paired Samples and Blocks

```
library(mosaic)
library(readr)
library(janitor)
library(tidyr) # for the gather() function
Dexterity <- read_csv("http://nhorton.people.amherst.edu/is5/data/Dexterity.csv") %>%
  clean_names()
```

```
## Warning: Duplicated column names deduplicated: 'Dominant' =>
## 'Dominant_1' [6]
```

```
## Parsed with column specification:
## cols(
##   `Age(months)` = col_integer(),
##   Gender = col_character(),
##   `Dominant Hand` = col_character(),
##   Dominant = col_double(),
##   `non-dominant` = col_double(),
##   Dominant_1 = col_double(),
##   `Non-dominant` = 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).

```
Dexterity %>%
  select(age_months, dominant_1, non_dominant_2, gender) %>%
  head(n = 7)
```

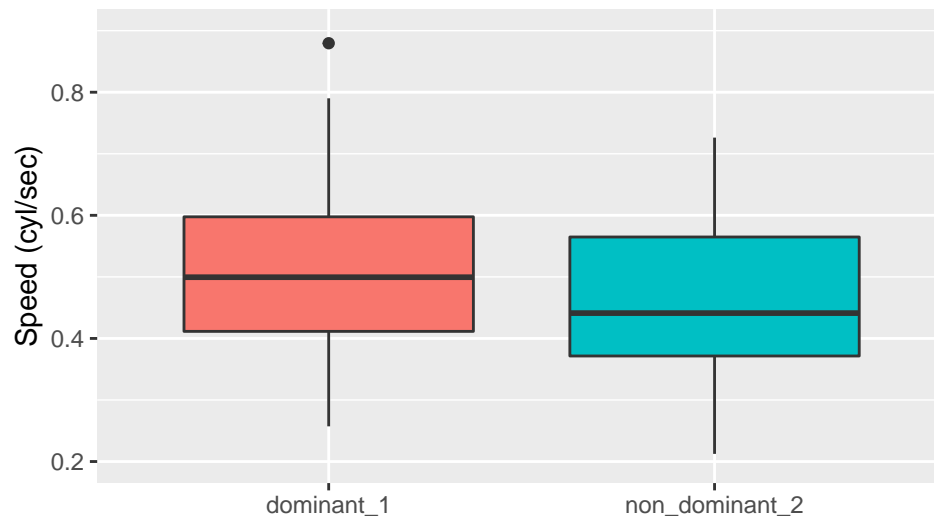
```
## # A tibble: 7 x 4
##   age_months dominant_1 non_dominant_2 gender
```

##	<int>	<dbl>	<dbl>	<chr>
## 1	117	0.353	0.216	male
## 2	101	0.257	0.343	male
## 3	135	0.537	0.497	male
## 4	119	0.444	0.496	male
## 5	124	0.483	0.388	female
## 6	127	0.524	0.422	female
## 7	101	0.455	0.381	male

Section 18.1: Paired Data

```
# Figure 18.1
Dexterity %>%
  select(dominant_1, non_dominant_2) %>%
  gather(key = hand_type, value = speed, dominant_1, non_dominant_2) %>%
  gf_boxplot(speed ~ hand_type, fill = ~ hand_type) %>%
  gf_labs(x = "", y = "Speed (cyl/sec)") +
  ylim(.2, .9) +
  guides(fill = FALSE)
```

```
## Warning: Removed 1 rows containing non-finite values (stat_boxplot).
```



Example 18.1: Identifying Paired Data

```
# page 586
# Create the data set
WorkWeek <- rbind(
  data.frame(name = "Jeff", fiveday = 2798, fourday = 2914),
  data.frame(name = "Betty", fiveday = 7724, fourday = 6112),
  data.frame(name = "Roger", fiveday = 7505, fourday = 6177),
  data.frame(name = "Tom", fiveday = 838, fourday = 1102),
  data.frame(name = "Aimee", fiveday = 4592, fourday = 3281),
  data.frame(name = "Greg", fiveday = 8107, fourday = 4997),
  data.frame(name = "Larry G.", fiveday = 1228, fourday = 1695),
  data.frame(name = "Tad", fiveday = 8718, fourday = 6606),
  data.frame(name = "Larry M.", fiveday = 1097, fourday = 1063),
```

```
data.frame(name = "Leslie", fiveday = 8089, fourday = 6392),
data.frame(name = "Lee", fiveday = 3807, fourday = 3362)
)
WorkWeek
```

```
##      name fiveday fourday
## 1    Jeff   2798   2914
## 2    Betty  7724   6112
## 3    Roger  7505   6177
## 4     Tom   838    1102
## 5    Aimee  4592   3281
## 6     Greg  8107   4997
## 7  Larry G.  1228   1695
## 8     Tad   8718   6606
## 9  Larry M.  1097   1063
## 10  Leslie  8089   6392
## 11    Lee   3807   3362
```

Looking at pairwise differences in Dexterity.

```
Dexterity %>%
  select(dominant_1, non_dominant_2) %>%
  mutate(difference = dominant_1 - non_dominant_2) %>%
  head(n = 18)
```

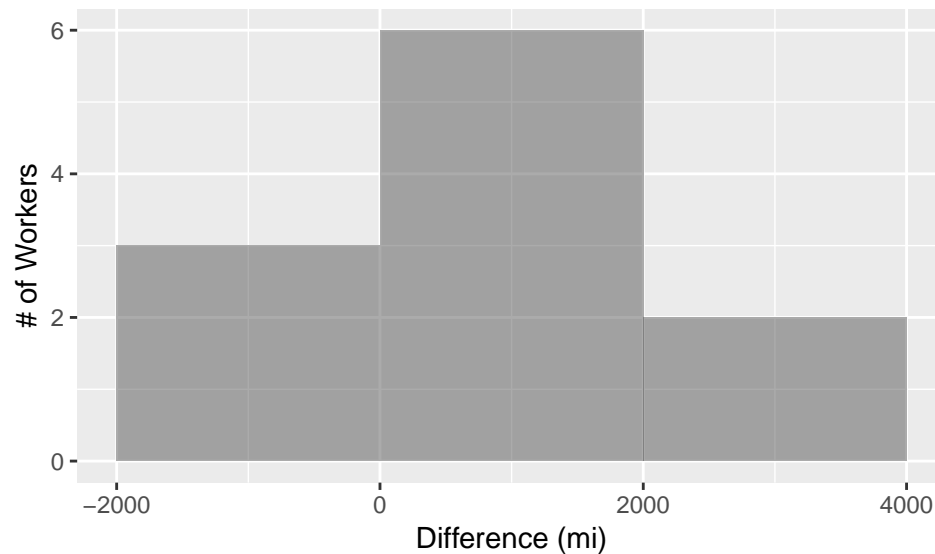
```
## # A tibble: 18 x 3
##   dominant_1 non_dominant_2 difference
##   <dbl>         <dbl>         <dbl>
## 1     0.353         0.216         0.137
## 2     0.257         0.343        -0.0863
## 3     0.537         0.497         0.0392
## 4     0.444         0.496        -0.0524
## 5     0.483         0.388         0.0947
## 6     0.524         0.422         0.102
## 7     0.455         0.381         0.0742
## 8     0.394         0.403        -0.00904
## 9     0.451         0.328         0.124
## 10    0.527         0.271         0.256
## 11    0.565         0.415         0.149
## 12    0.653         0.298         0.355
## 13    0.421         0.337         0.0833
## 14    0.320         0.233         0.0872
## 15    0.344         0.241         0.102
## 16    0.428         0.612        -0.184
## 17    0.556         0.521         0.0357
## 18    0.465         0.411         0.0543
```

Section 18.2: The Paired *t*-Test

Example 18.2: Checking Assumptions and Conditions

```
# page 588
WorkWeek <- WorkWeek %>%
  mutate(difference = fiveday - fourday)
```

```
gf_histogram(~ difference, data = WorkWeek, binwidth = 2000, center = 1000) %>%
  gf_labs(x = "Difference (mi)", y = "# of Workers")
```



Example 18.3: Doing a Paired t -Test

```
nwork <- nrow(WorkWeek)
nwork # number of pairs
```

```
## [1] 11
```

```
dwork <- mean(~ difference, data = WorkWeek)
dwork # mean of differences
```

```
## [1] 982
```

```
swork <- sd(~ difference, data = WorkWeek)
swork # SD of differences
```

```
## [1] 1139.568
```

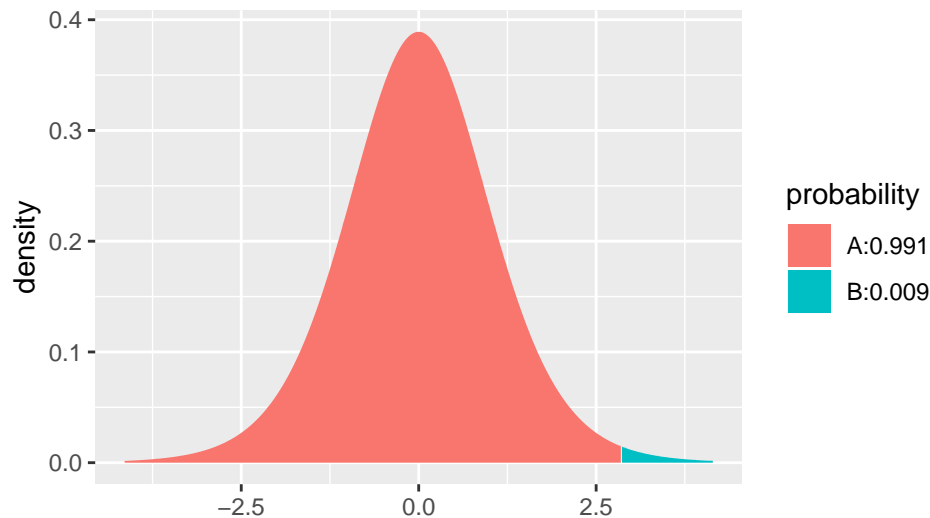
```
sework <- swork/(nwork^.5)
sework # SE of differences
```

```
## [1] 343.5928
```

```
twork <- (dwork - 0)/sework
twork # t stat
```

```
## [1] 2.858034
```

```
2 * xpt(twork, df = nwork - 1, lower.tail = FALSE)
```



```
## [1] 0.01701413
```

The `xpt()` function finds the p-value and plots it on a graph to visualize it. Here, the visualization shows a one-sided test, but in the book, it is two sided.

Section 18.3: Confidence Intervals for Matched Pairs

```
Couples <- read_csv("http://nhorton.people.amherst.edu/is5/data/Couples.csv") %>%
  filter(wAge != "*") %>%
  mutate(wAge = as.numeric(wAge))
```

```
## Parsed with column specification:
## cols(
##   Names = col_character(),
##   wAge = col_character(),
##   hAge = col_integer(),
##   wHeight = col_integer(),
##   hHeight = col_integer()
## )
```

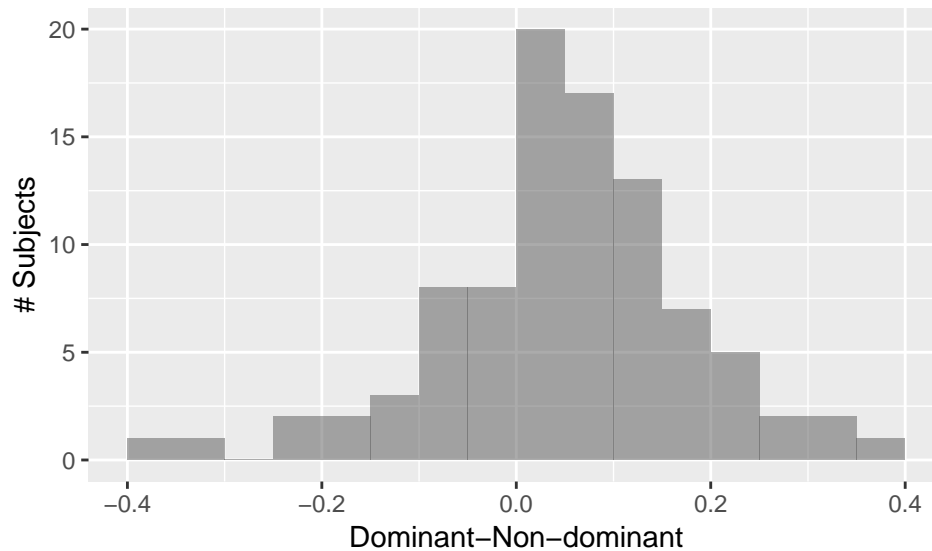
table on page 592

```
Couples %>%
  select(wAge, hAge) %>%
  mutate(difference = hAge - wAge) %>%
  head(n = 7)
```

```
## # A tibble: 7 x 3
##   wAge  hAge difference
##   <dbl> <int>     <dbl>
## 1    43    49         6
## 2    28    25        -3
## 3    30    40        10
## 4    57    52        -5
## 5    52    58         6
## 6    27    32         5
## 7    52    43        -9
```

Step-By-Step Example: A Paired t -Interval

```
DexData <- Dexterity %>%
  select(dominant_1, non_dominant_2) %>%
  mutate(difference = dominant_1 - non_dominant_2) %>%
  filter(dominant_1 < 1)
# For some reason, the book has removed one observation where dominant_1 = 1,
# but has kept the count of children at 93 instead of 92
gf_histogram(~ difference, data = DexData, binwidth = .05, center = .025) %>%
  gf_labs(x = "Dominant-Non-dominant", y = "# Subjects")
```



```
# Mechanics
ndex <- nrow(DexData) + 1 # the book kept n at 93 for some reason
ndex # number of pairs (children)
```

```
## [1] 93
```

```
ddex <- mean(~ difference, data = DexData)
ddex # mean difference
```

```
## [1] 0.05148209
```

```
sdex <- sd(~ difference, data = DexData)
sdex # standard deviation of the differences
```

```
## [1] 0.1298746
```

```
sedex <- sdex/(ndex^.5)
sedex # standard error of the differences
```

```
## [1] 0.01346736
```

```
df <- ndex - 1
df
```

```
## [1] 92
```

```
tstats <- qt(p = c(.025, .975), df = df)
tstats
```

```
## [1] -1.986086 1.986086
```

```

medex <- tstats * sedex
medex # margin of error of the differences

## [1] -0.02674735  0.02674735
ddex + medex

## [1] 0.02473474 0.07822943
# Or, if you don't want to go through all those calculations:
t.test(~ difference, data = DexData, df = df)

##
## One Sample t-test
##
## data: difference
## t = 3.8021, df = 91, p-value = 0.0002592
## alternative hypothesis: true mean is not equal to 0
## 95 percent confidence interval:
##  0.02458583 0.07837834
## sample estimates:
## mean of x
## 0.05148209

```

Effect Size

Example 18.4: Looking at Effect Size with a Paired t Confidence Interval

```

tstats <- qt(p = c(.025, .975), df = nwork - 1)
tstats

## [1] -2.228139  2.228139
me <- tstats * sework
me # margin of error

## [1] -765.5724  765.5724
dwork + me # confidence interval

## [1]  216.4276 1747.5724

```

Section 18.4: Blocking

What's Independent?

Random Matters: A Bootstrapped Paired Data Confidence Interval and Hypothesis Test

```

set.seed(2345)
numsim <- 5000

# What does do() do?
mean(~ difference, data = resample(DexData)) # One mean of a random resample

## [1] 0.04257654

```

```
mean(~ difference, data = resample(DexData)) # Another mean of a random resample
```

```
## [1] 0.04985414
```

```
do(2) * mean(~ difference, data = resample(DexData)) # Calculates two means
```

```
##          mean
## 1 0.03828900
## 2 0.02499735
```

```
# We need numsim means
```

```
DexBoots <- do(numsim) * mean(~ difference, data = resample(DexData))
```

For more information about `resample()`, refer to the `resample` vignette in `mosaic`.

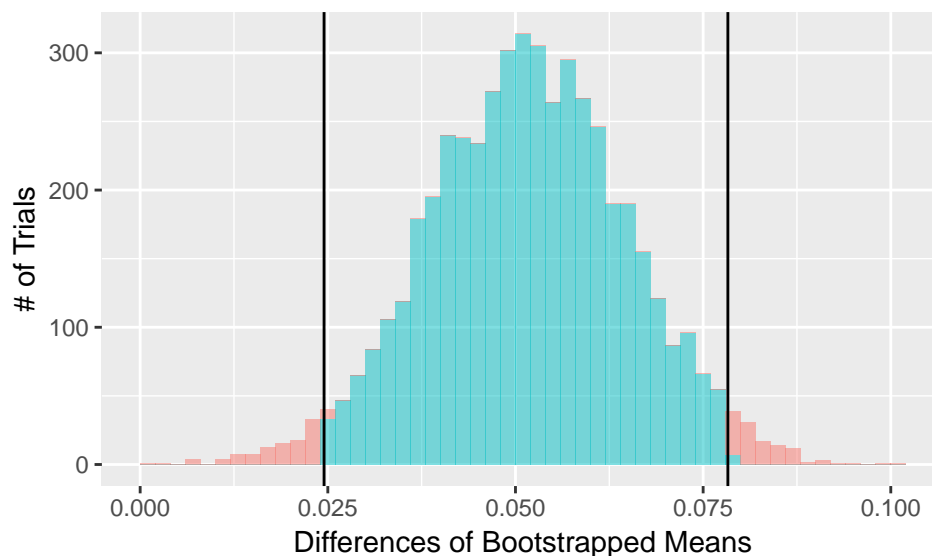
```
qdata(~ mean, p = c(.025, .975), data = DexBoots)
```

```
##          quantile      p
## 2.5%  0.02515483 0.025
## 97.5% 0.07794129 0.975
```

```
DexBoots <- DexBoots %>%
  mutate(interval = ifelse(mean > .0245 & mean < .0783, "Within 95% Confidence",
                           "Outside 95% Confidence"))
```

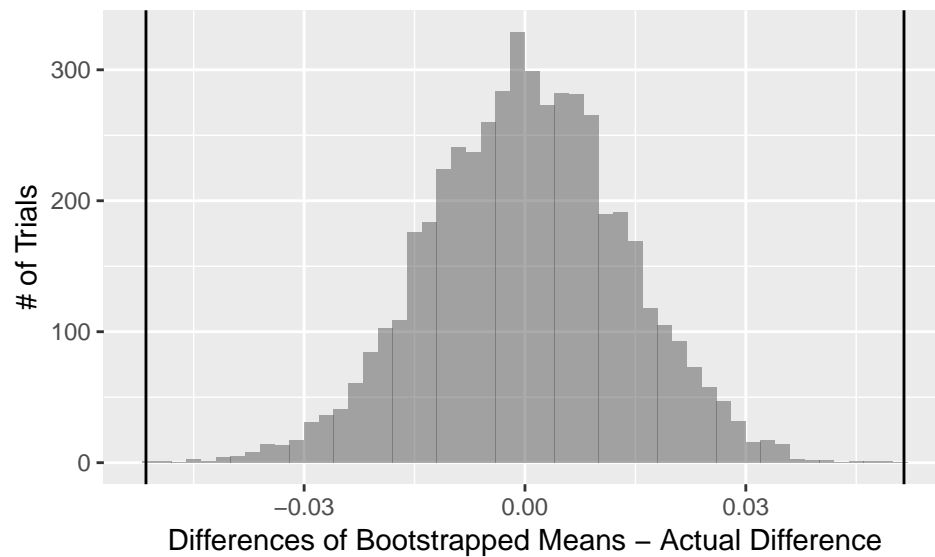
```
# Figure 18.4, page 597
```

```
gf_histogram(~ mean, fill = ~ interval, data = DexBoots, binwidth = .002, center = .001) %>%
  gf_vline(xintercept = .0245) %>%
  gf_vline(xintercept = .0783) %>%
  gf_labs(x = "Differences of Bootstrapped Means", y = "# of Trials") +
  guides(fill = FALSE)
```



```
# Figure 18.5
```

```
gf_histogram(~ (mean - ddex), data = DexBoots, binwidth = .002, center = .001) %>%
  gf_vline(xintercept = ddex) %>%
  gf_vline(xintercept = -ddex) %>%
  gf_labs(x = "Differences of Bootstrapped Means - Actual Difference", y = "# of Trials")
```

```
favstats(~ (mean - ddex), data = DexBoots)
```

```
##          min          Q1          median          Q3          max          mean
## -0.05087679 -0.008911368 0.000258236 0.009129865 0.04873482 0.0002299513
##          sd      n missing
## 0.01336809 5000          0
```

With `favstats()`, we can see that our minimum is within the interval, but our maximum isn't.

```
DexBoots %>%
  filter((mean - ddex) > ddex)
```

```
## [1] mean      interval
## <0 rows> (or 0-length row.names)
```

Like the book, there is one instance (out of 5,000), so we estimate the P-value as 1/5,000 (the book says 50,000, which is incorrect), or .0002.