

IPS9 in R: Inference for Categorical Data (Chapter 9)

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Introduction and background

These documents are intended to help describe how to undertake analyses introduced as examples in the Ninth Edition of *Introduction to the Practice of Statistics* (2017) by Moore, McCabe, and Craig.

More information about the book can be found [here](#). The data used in these documents can be found under Data Sets in the Student Site. This file as well as the associated R Markdown reproducible analysis source file used to create it can be found at <https://nhorton.people.amherst.edu/ips9/>.

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 9: Inference for Categorical Data

This file replicates the analyses from Chapter 9: Inference for Categorical Data.

First, load the packages that will be needed for this document:

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

Section 9.1: Inference for two-way tables

To recreate the dataset that was used in Example 9.1, we will use a combination of several `do()` calls and `rbind()`. This will allow us to create the observations with the specific attributes based on the counts that appear in the table.

We recreate it like this.

```
#Ex9.1
Instag <- rbind(
  do(298) * data.frame(Sex = "Men",   User = "No"),
  do(209) * data.frame(Sex = "Women", User = "No"),
  do(234) * data.frame(Sex = "Men",   User = "Yes"),
  do(328) * data.frame(Sex = "Women", User = "Yes")
)
head(Instag)
```

```
##   Sex User .row .index
## 1 Men   No    1      1
## 2 Men   No    1      2
## 3 Men   No    1      3
## 4 Men   No    1      4
## 5 Men   No    1      5
```

```
## 6 Men    No    1    6
```

We take a small peek of the dataset with the `head()` function that returns the first few observations from a given dataset. Some useful columns were returned with our dataset. You don't have to worry about them now.

To create the summary table in Example 9.1 we will have to compute some intermediate steps.

To start, we first get the total counts for each sex.

```
#Ex9.1
Combined_Sex <- Instag %>%
  group_by(Sex) %>%
  summarize(n = n())
Combined_Sex
```

```
## # A tibble: 2 x 2
##   Sex      n
##   <fct> <int>
## 1 Men    532
## 2 Women  537
```

This will be the first column of our table. Then we get a dataframe with only the counts of those who have a value of "Yes" for User. This will be the second column.

```
#Ex9.1
YesUsers <- Instag %>%
  group_by(User, Sex) %>%
  summarize(n = n()) %>%
  filter(User == "Yes")
YesUsers
```

```
## # A tibble: 2 x 3
## # Groups:   User [1]
##   User Sex      n
##   <fct> <fct> <int>
## 1 Yes  Men    234
## 2 Yes  Women  328
```

And finally, we use a `mutate()` call to create a variable derived from other variables. This will create the

$$\hat{p} = X/n$$

column.

```
#Ex9.1
Ex8.11Table <- Combined_Sex %>%
  left_join(YesUsers, by = "Sex") %>%
  select(Sex, n = n.x, X = n.y) %>%
  mutate(`p_hat` = X/n` = X/n)
Ex8.11Table
```

```
## # A tibble: 2 x 4
##   Sex      n      X `p_hat` = X/n`
##   <fct> <int> <int>      <dbl>
## 1 Men    532   234      0.440
## 2 Women  537   328      0.611
```

And we then have the table with the percentages of people who are users over the total number of people of the same sex!

Now take look at Example 9.2 in page 526. To recreate that table of counts we simply have to call the `tally()` function and it will make the 2-way table for us. We call it like this:

```
#Ex9.2
tally(~ User + Sex, data = Instag, margins = TRUE)
```

```
##           Sex
## User      Men Women Total
##  No       298   209   507
##  Yes      234   328   562
##  Total   532   537  1069
```

The `margins = TRUE` optio makes sure that `tally()` outputs the convenient Total columns just like in page 527!

Turn your attention to Example 9.3 now. After creating the dataset from the counts, we can use a similar `tally()` call to recreate the table and verify that our method to create the dataset is in fact accurate.

```
#Ex9.3
Vaccine <- rbind(
  do(729) * data.frame(Required = "Yes", Party = "Democratic"),
  do(479) * data.frame(Required = "Yes", Party = "Republican"),
  do(230) * data.frame(Required = "No", Party = "Democratic"),
  do(258) * data.frame(Required = "No", Party = "Republican")
)

tally(~ Required + Party, data = Vaccine, margins = TRUE)
```

```
##           Party
## Required Democratic Republican Total
##  Yes              729          479  1208
##  No              230          258   488
##  Total          959          737  1696
```

Now we continue to explore our 2 way tables. In Example 9.5 we can see the marginal distribution of our Vaccine tables across political party preference. We recreate it with a call to `tally()` but this time we will use a new parameter too.

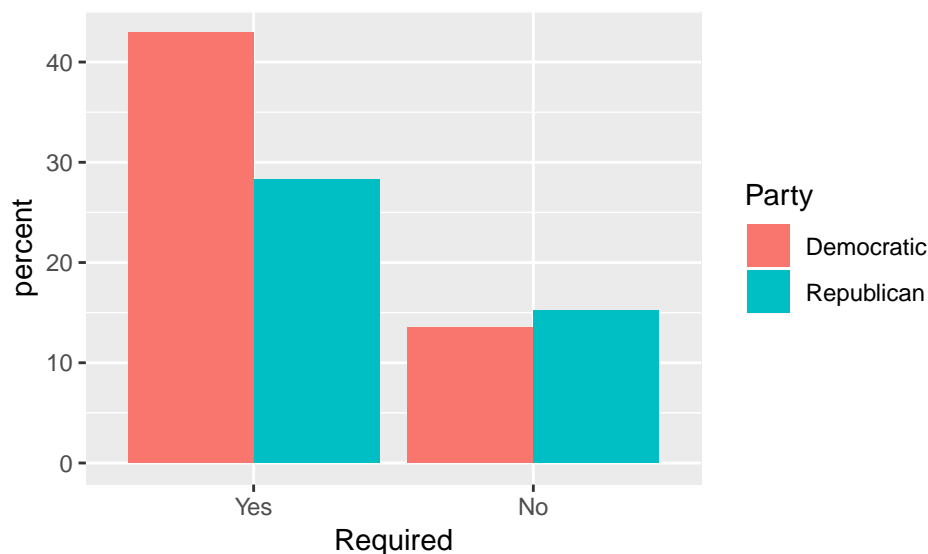
```
#Ex9.5
tally(Required ~ Party, data = Vaccine, margins = TRUE, format = "percent")
```

```
##           Party
## Required Democratic Republican
##  Yes       76.01668  64.99322
##  No       23.98332  35.00678
##  Total  100.00000  100.00000
```

The `format = "percent"` will nicely output the results in percentage form!

The output from `tally()` is good enough. However, a picture is worth a thousand words. Let's try to create a bar graph out of the Vaccine dataset.

```
#Ex9.6
gf_percents(~ Required, data = Vaccine, fill = ~ Party, position = "dodge")
```

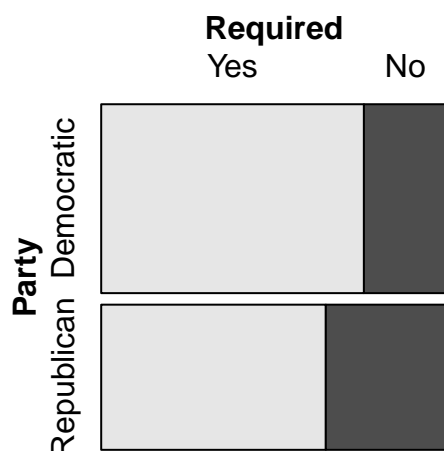


With the help of `gf_percents()` we can plot the percentage of each group (e.g. Democratic & “No”) and compare them. This is a useful way to draw insights from two variables at once.

Note: This is not an equivalent bar graph but still provides the same useful information. The original bar graph in page 530 graphs the percentages across political party (i.e. adding both columns belonging to a party will give 100%).

Another way that we can visualize two categorical variables is to create a mosaic plot. We will use the `vcd` package’s `mosaic()` function to plot the mosaic plot. Note the call resembles the same syntax of the `tally()` commands we made earlier.

```
#Ex9.7
vcd::mosaic(Required ~ Party, data = Vaccine)
```



Having multiple ways to visualize variables will help you analyze your data more thoroughly and communicate your findings in a more intuitive way.

In Example 9.7 we are interested in getting the expected counts of our Vaccine data. In R you can take advantage of the `xchisq.test()` function and get the relevant output like this:

#Ex9.8

```
chiSqVaccine <- xchisq.test(tally(Required ~ Party, data = Vaccine), correct = FALSE)
```

```
##
##  Pearson's Chi-squared test
##
## data:  x
## X-squared = 24.709, df = 1, p-value = 6.666e-07
##
##      729      479
## (683.06) (524.94)
## [ 3.09] [ 4.02]
## < 1.76> <-2.01>
##
##      230      258
## (275.94) (212.06)
## [ 7.65] [ 9.95]
## <-2.77> < 3.15>
##
## key:
##  observed
##  (expected)
## [contribution to X-squared]
## <Pearson residual>
```

```
with(chiSqVaccine, expected)
```

```
##           Party
## Required Democratic Republican
##      Yes      683.0613      524.9387
##      No       275.9387      212.0613
```

To understand what is going on in this code, break it down into its components. We are creating a variable called `chiSqVaccine` and we are assigning the output of the `xchisq.test()` call. The object stored in our variable will contain several useful fields as we will see. The first one is the expected values. To extract it from the object we use the `with()` function.

Note: We specify the `correct = FALSE` option to match the book's table. This option specifies that there should be no continuity correction applied to our test. You can see how the output changes by removing that option.

In a manner similar to the one above, we can get the observed counts we calculated with `tally()` before. We just retrieve the relevant field from our object with the `with()` function again.

#Ex9.8

```
with(chiSqVaccine, observed)
```

```
##           Party
## Required Democratic Republican
##      Yes      729      479
##      No       230      258
```

To see the output of the Chi-Square test discussed in Example 9.8 we just need to print the object we stored in our variable earlier.

#Ex9.8

```
chiSqVaccine
```

```
##
## Pearson's Chi-squared test
##
## data:  x
## X-squared = 24.709, df = 1, p-value = 6.666e-07
```

All this useful features are already built into how R's `xchisq.test()` function works.

Note: There is an error in the

$$\chi^2$$

value in the book. While it showed the correct machine output, it specified the wrong

$$\chi^2$$

squared value.

We continue with Example 9.9 in page 537.

```
#Ex9.9
Health <- rbind(
  do(69) * data.frame(PhysAct = "Low", FruitConsumption = "Low"),
  do(206) * data.frame(PhysAct = "Moderate", FruitConsumption = "Low"),
  do(294) * data.frame(PhysAct = "Vigorous", FruitConsumption = "Low"),
  do(25) * data.frame(PhysAct = "Low", FruitConsumption = "Medium"),
  do(126) * data.frame(PhysAct = "Moderate", FruitConsumption = "Medium"),
  do(170) * data.frame(PhysAct = "Vigorous", FruitConsumption = "Medium"),
  do(14) * data.frame(PhysAct = "Low", FruitConsumption = "High"),
  do(111) * data.frame(PhysAct = "Moderate", FruitConsumption = "High"),
  do(169) * data.frame(PhysAct = "Vigorous", FruitConsumption = "High")
)
```

You should already know what is happening in the code chunk above. We will store the dataset into a variable called `Health`.

Now we recreate the table in page 537 as follows:

```
#Ex9.9
tally(~ FruitConsumption + PhysAct, data = Health, margins = TRUE)
```

```
##              PhysAct
## FruitConsumption  Low Moderate Vigorous Total
##              Low    69      206      294    569
##              Medium  25      126      170    321
##              High   14      111      169    294
##              Total  108     443     633   1184
```

The table above is the 2 way table of counts from the `Health` data. To get the percentages instead we use the `format` parameter.

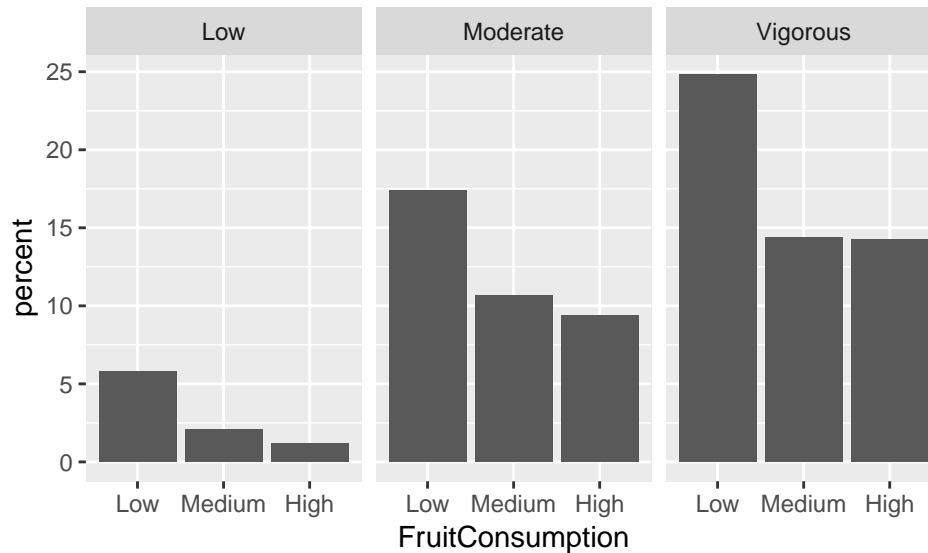
```
#Ex9.10
tally(~ FruitConsumption + PhysAct, data = Health, margins = TRUE, format = "percent")
```

```
##              PhysAct
## FruitConsumption  Low Moderate Vigorous Total
##              Low    5.827703 17.398649 24.831081 48.057432
##              Medium  2.111486 10.641892 14.358108 27.111486
##              High   1.182432  9.375000 14.273649 24.831081
##              Total   9.121622 37.415541 53.462838 100.000000
```

Again, visualizations will trump tables in all appropriate cases. We will create a faceted bar graph.

#Fig9.7

```
gf_percents(~ FruitConsumption | PhysAct, data = Health)
```



Note: These are not equivalent bar graphs to Figure 9.7, but they still provide the same useful information.

Now, let's get the expected counts for our Health data.

#Ex9.11

```
chiSqHealth <- xchisq.test(tally(FruitConsumption ~ PhysAct, data = Health), correct = FALSE)
```

```
##
##  Pearson's Chi-squared test
##
## data:  x
## X-squared = 14.152, df = 4, p-value = 0.006824
##
##      69      206      294
## ( 51.90) (212.89) (304.20)
## [5.6325] [0.2233] [0.3422]
## < 2.373> <-0.473> <-0.585>
##
##      25      126      170
## ( 29.28) (120.10) (171.62)
## [0.6257] [0.2895] [0.0152]
## <-0.791> < 0.538> <-0.123>
##
##      14      111      169
## ( 26.82) (110.00) (157.18)
## [6.1262] [0.0091] [0.8888]
## <-2.475> < 0.095> < 0.943>
##
## key:
##  observed
##  (expected)
##  [contribution to X-squared]
```

```
## <Pearson residual>
```

```
with(chiSqHealth, expected)
```

```
##               PhysAct
## FruitConsumption  Low Moderate Vigorous
##           Low  51.90203 212.8944 304.2035
##           Medium 29.28041 120.1039 171.6157
##           High   26.81757 110.0017 157.1807
```

And our observed counts...

```
#Ex9.11
```

```
with(chiSqHealth, observed)
```

```
##               PhysAct
## FruitConsumption  Low Moderate Vigorous
##           Low      69      206      294
##           Medium   25      126      170
##           High     14      111      169
```

And finally our

χ^2

statistic.

```
#Ex9.11
```

```
chiSqHealth
```

```
##
## Pearson's Chi-squared test
##
## data:  x
## X-squared = 14.152, df = 4, p-value = 0.006824
```

Remember these are all possible thans to the functionality of the `xchisq.test()` function.

Section 9.2: Goodness of fit

We will be using data of the ACT from six different states. We recreate our dataset from the counts.

```
#Ex9.13
```

```
ACT <- rbind(
  do(167) * data.frame(State = "AZ", label = 1),
  do(257) * data.frame(State = "CA", label = 2),
  do(257) * data.frame(State = "HI", label = 3),
  do(297) * data.frame(State = "IN", label = 4),
  do(107) * data.frame(State = "NV", label = 5),
  do(482) * data.frame(State = "OH", label = 6)
)
```

To get a sense of the number of participants in the study (pg 546) we can quickly do a `tally()` call on the State column.

```
#Ex9.13
```

```
tally(~ State, data = ACT, margins = TRUE)
```

```
## State
##   AZ   CA   HI   IN   NV   OH Total
##  167  257  257  297  107  482 1567
```


We will now import the population proportions from a csv file. We will use these values to see how close our sample counts are to the population values.

#Ex9.13

```
ACTPopProp <- read_csv("https://nhorton.people.amherst.edu/ips9/data/chapter09/EG09-13ACT.csv")
ACTPopProp
```

```
## # A tibble: 6 x 4
##   State Label Count  Prob
##   <chr> <int> <dbl> <dbl>
## 1 AZ      1    167 0.105
## 2 CA      2    257 0.172
## 3 HI      3    257 0.164
## 4 IN      4    297 0.188
## 5 NV      5    107 0.07
## 6 OH      6    482 0.301
```

We will use the same `xchisq.test()` function from before. It is important to note the new behavior we expect from the function when we provide a vector of the population proportions. The expected counts and the

$$\chi^2$$

value will depend on this new parameter.

#Ex9.13

```
chisqACT <- xchisq.test(tally(~ State, data = ACT), p = c(0.105, .172, .164, .188, .07, .301), correct = FALSE)
```

```
##
## Chi-squared test for given probabilities
##
## data:  x
## X-squared = 0.93084, df = 5, p-value = 0.9679
##
##      167      257      257      297      107      482
## (164.53) (269.52) (256.99) (294.60) (109.69) (471.67)
## [3.7e-02] [5.8e-01] [5.6e-07] [2.0e-02] [6.6e-02] [2.3e-01]
## < 0.19217> <-0.76286> < 0.00075> < 0.14006> <-0.25684> < 0.47578>
##
## key:
## observed
## (expected)
## [contribution to X-squared]
## <Pearson residual>
```

Now that we have saved our object, we can access the expected counts and the test statistic just like before.

#Ex9.13

```
with(chisqACT, expected)
```

```
##      AZ      CA      HI      IN      NV      OH
## 164.535 269.524 256.988 294.596 109.690 471.667
```

#Ex9.14

```
chisqACT
```

```
##
## Chi-squared test for given probabilities
##
## data:  x
```

```
## X-squared = 0.93084, df = 5, p-value = 0.9679
```

Another example of a field included in the return value of the `xchisq.test()` is the `residuals` field. Let's take a look.

```
#Ex9.15
```

```
with(chisqACT, residuals)
```

```
## State
```

```
##           AZ           CA           HI           IN           NV
## 0.1921709671 -0.7628591106 0.0007485569 0.1400622312 -0.2568436078
##           OH
## 0.4757827403
```

And just like that, let R help you with most of the computations when you are analyzing your categorical variables.