Chi-Square Tests

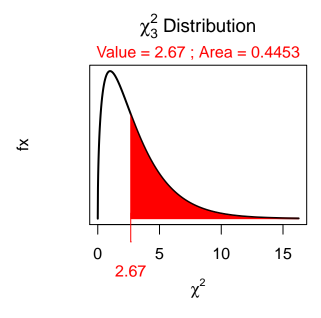
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First Commands

> library(NCStats)

Chi-Square Distribution Calculations

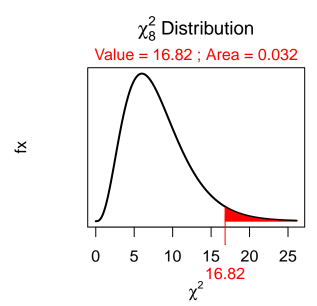
An example of computing the p-value $\chi^2=2.67$ and df=3.



[1] 0.4453496

An example of computing the p-value if $\chi^2=16.82$ and df=8.

```
> ( distrib(16.82,distrib="chisq",df=8,lower.tail=FALSE) )
```



[1] 0.03203895

Chi-Square Test

From Summarized Observed Table

When Chinook Salmon (*Oncorhynchus tshawytscha*) were first introduced to Lake Superior there was concern that they would compete with native Lake Trout (*Salvelinus namaycush*) for Cisco (*Coregonus artedi*). Preliminarily, fisheries biologists classified the dominant food items (Cisco, Smelt (*Osmerus mordax*) (another type of fish), or *Mysis* (an invertebrate)) in the diets of 50 Lake Trout and 40 Chinook Salmon. They found that 32, 10, and 8 Lake Trout diets were dominated by Cisco, Smelt, and *Mysis*, respectively. Similarly, 18, 18, and 4 Chinook Salmon diets were dominated by Cisco, Smelt, and *Mysis*, respectively. Test (at the 10% level) if the distribution of dominant food items differs at the 5% level?

```
> chi1 <- chisq.test(obs,correct=FALSE)</pre>
> chi1$expected
                  Cisco
                           Smelt
                                    Mysis
Lake Trout
               27.77778 15.55556 6.666667
Chinook Salmon 22.22222 12.44444 5.333333
> chi1$expected >= 5
               Cisco Smelt Mysis
Lake Trout
                TRUE TRUE TRUE
Chinook Salmon TRUE TRUE TRUE
> chi1
Pearson's Chi-squared test with obs
X-squared = 6.5083, df = 2, p-value = 0.03861
> chi1$residuals
                    Cisco
                              Smelt
                                         Mysis
                0.8011103 -1.408590 0.5163978
Lake Trout
Chinook Salmon -0.8956686 1.574852 -0.5773503
> percTable(obs,margin=1,digits=1)
               Cisco Smelt Mysis Sum
Lake Trout
                              16 100
                  64
                        20
Chinook Salmon
                  45
                        45
                              10 100
> ( obs2 <- obs[,-2] )
               Cisco Mysis
Lake Trout
                  32
                         8
Chinook Salmon
                  18
> ( chi2 <- chisq.test(obs2,correct=FALSE) )</pre>
Warning in chisq.test(obs2, correct = FALSE): Chi-squared approximation may be incorrect
Pearson's Chi-squared test with obs2
X-squared = 0.0301, df = 1, p-value = 0.8624
```

From Raw Data

The General Social Survey (GSS) is a nationwide survey that has been administered since 1972 to gather data on contemporary American society in an attempt to monitor and explain trends in attitudes, behaviors, and attributes of American society. One part of that survey asked respondents to state their opinion on how true the following statement was – "All radioactivity is made by humans." Respondents were also categorized by their highest educational degree. The results from this portion of the GSS is in SciTest1.csv. Use these data to determine, at the 5% level, if the distribution of responses to this statement differs among levels of education.

```
> setwd("C:/aaaWork/Web/GitHub/NCMTH107/resources/class/HOs")
> ST1 <- read.csv("SciTest1.csv")</pre>
> str(ST1)
'data.frame':
                2549 obs. of 2 variables:
 $ degree : Factor w/ 5 levels "bach", "grad",..: 5 5 5 5 5 5 5 5 5 5 5 ...
 $ scitest: Factor w/ 4 levels "def.not","def.true",..: 2 2 2 2 2 2 2 2 2 ...
> levels(ST1$degree)
[1] "bach" "grad"
                             "jc"
                                     "lt.hs"
> ST1$fdegree <- factor(ST1$degree,levels=c("lt.hs","hs","jc","bach","grad"))
> levels(ST1$scitest)
[1] "def.not"
                             "prob.not"
                "def.true"
                                         "prob.true"
> ST1$fscitest <- factor(ST1$scitest,levels=c("def.not","prob.not","prob.true","def.true"))
> ( freq.tbl <- xtabs(~fdegree+fscitest,data=ST1) )</pre>
       fscitest
fdegree def.not prob.not prob.true def.true
  lt.hs
             52
                     112
                                155
                                          70
            366
                     451
                                437
                                         114
  hs
  jс
             60
                      44
                                 36
                                           9
  bach
            214
                      135
                                 78
                                          12
            123
                      57
                                 18
  grad
> ST1.chi <- chisq.test(freq.tbl,correct=FALSE)
> ST1.chi$expected
       fscitest
fdegree
          def.not prob.not prob.true def.true
  lt.hs 124.37623 121.93448 110.48882
                                        32.20047
        437.39506 428.80816 388.55708 113.23970
  hs
  jс
         47.64025 46.70498 42.32091 12.33386
  bach 140.36289 137.60730 124.69047
                                        36.33935
         65.22558 63.94508 57.94272
  grad
                                       16.88662
```

> ST1.chi\$expected >= 5

```
fscitest
fdegree def.not prob.not prob.true def.true
  lt.hs
           TRUE
                     TRUE
                                TRUE
                                          TRUE
  hs
           TRUE
                     TRUE
                                TRUE
                                          TRUE
           TRUE
                     TRUE
                                TRUE
                                          TRUE
  jс
                     TRUE
           TRUE
                                TRUE
                                          TRUE
  bach
           TRUE
                     TRUE
                                TRUE
  grad
                                          TRUE
```

```
> all(chi1$expected >= 5)
```

[1] TRUE

```
> ST1.chi
```

```
Pearson's Chi-squared test with freq.tbl X-squared = 288.2331, df = 12, p-value < 2.2e-16
```

> ST1.chi\$residuals

fscitest

```
fdegree
           def.not
                     prob.not
                               prob.true
                                            def.true
  lt.hs -6.4897392 -0.8996675
                               4.2345762
                                          6.6612431
        -3.4137460
                   1.0716718
                               2.4575530
                                          0.0714471
         1.7906992 -0.3958064 -0.9716327 -0.9492868
  jс
  bach
         6.2154218 -0.2222643 -4.1813025 -4.0375730
  grad
         7.1536326 -0.8685073 -5.2473280 -2.6492428
```

```
> percTable(freq.tbl,margin=1,digits=1)
```

fscitest

```
fdegree def.not prob.not prob.true def.true
                                                 Sum
  lt.hs
           13.4
                     28.8
                                39.8
                                         18.0 100.0
           26.8
                     33.0
  hs
                                31.9
                                          8.3 100.0
           40.3
  jс
                     29.5
                                24.2
                                           6.0 100.0
            48.7
                     30.8
                                17.8
                                          2.7 100.0
  bach
           60.3
                     27.9
                                 8.8
                                          2.9 99.9
  grad
```

Goodness-of-Fit Test

Four-o'clocks (*Mirabilis jalapa*) are plants native to tropical America. Individual four-o'clocks can have red, white, or pink flowers. Flower color in this species is thought to be controlled by a single gene locus with two alleles experssing incomplete dominance, so that heterozygotes are pink-flowered, while homozygotes for one allele are white-flowered and homozygotes for the other allele are red-flowered (see this). According to Mendelian genetic principles, self-pollination of pink-flowered plants should produce progeny that have red, pink, and white flowers in a 1:2:1 ratio. A horticulturist allowed several pink-flowered plants to self-pollinate and produce 240 progeny with 55 that were red-flowered, 132 that were pink-flowered, and 53 that were white-flowered. Use the results to determine, at the 5% level, if the theoretical 1:2:1 ratio is upheld with these data.

```
> obs <- c(red=55,pink=132,white=53)
> exp.p <- c(red=1/4,pink=2/4,white=1/4)
> chi1 <- chisq.test(obs,p=exp.p,rescale=TRUE,correct=FALSE)
> chi1$expected

red pink white
    60    120    60
> chi1$observed

red pink white
    55    132    53
> chi1

Chi-squared test for given probabilities with obs
X-squared = 2.4333, df = 2, p-value = 0.2962
> gofCI(chi1,digits=3)

p.obs p.LCI p.UCI p.exp
```

red 0.229 0.181 0.286 0.25 pink 0.550 0.487 0.612 0.50 white 0.221 0.173 0.277 0.25