Statistics 305/605: Introduction to Biostatistical Methods for Health Sciences

Demo for Chapter 15, part 4: Inference for Odds Ratios

Jinko Graham

2018-10-06

Testing whether OR = 1

▶ The chi-square test assesses the null hypothesis that OR = 1 (no association between exposure and disease) against the alternative hypothesis that $OR \neq 1$ (an association).

```
mydf <- data.frame(case=c(1350,7),control=c(1296,61)) # Doll and Hill's data
rownames(mydf) <- c("smoker", "non-smoker")</pre>
mydf
##
              case control
## smoker
              1350
                       1296
                         61
## non-smoker
chisq.test(mydf)
##
##
    Pearson's Chi-squared test with Yates' continuity correction
##
## data: mydf
## X-squared = 42.37, df = 1, p-value = 7.552e-11
```

Using R to get point and interval estimates of the OR.

- ► For a 2 × 2 table, the calculations are easy enough to do with a hand calculator or spreadsheet.
- Below we present some R code that illustrates data frame manipulation in R.
 - This will give us code that will generalize to the case of multiple exposures, as in the Doll and Hill dataset where smoking status has four levels.
 - ▶ If you are not interested in learning R, don't struggle to understand each line of code.
 - ▶ Instead focus on understanding the *purpose* of each line of code.
 - There will be similar code in assignment 2 and you will need to understand the steps to make simple modifications as necessary.

Add Odds of Case to Data Frame

```
## group case control total prcase odds
## 1 smoker 1350 1296 2646 0.5102041 1.0416667
## 2 non-smoker 7 61 68 0.1029412 0.1147541
```

- mutate() is used to create new variables from existing ones and add them to our data frame.
- In this example, the variables total, prcase and odds are created and added to mydf.
 - Notice that the calculation of odds can use the newly-created variable prcase.

Extract the Baseline Odds

As a baseline for comparison, we will use the group of non-smokers. Let's extract their estimated odds of lung cancer.

```
odds0 <- mydf[mydf$group=="non-smoker","odds"]
odds0</pre>
```

- ## [1] 0.1147541
 - We can grab elements of a dataframe by referencing the desired rows and columns inside square brackets.
 - ► E.G, mydf[1,2] will grab the element in the 1st row and 2nd column of mydf.
 - Above, we indicate the desired row with the logical condition mydf\$group=="non-smoker",
 - mydf\$group refers to the variable group in the data frame mydf.
 - Grab only the row for the non-smoking group.
 - ► The desired column has the variable name odds.

Extracting Other Baseline Data

Extract the number of non-smoker cases and controls analogously.

```
c <- mydf[mydf$group=="non-smoker","case"]

## [1] 7

d <- mydf[mydf$group=="non-smoker","control"]
d

## [1] 61</pre>
```

Add the SE for the log-OR

Add the CI

```
##
        group case control total prcase odds OR se.logOR lowerCI
       smoker 1350
                    1296 2646 0.510 1.042 9.077
                                                 0.401
                                                        4.137
## 1
## 2 non-smoker
             7
                      61
                           68 0.103 0.115 1.000
                                                 0.564 0.331
##
    upperCI
## 1 19.918
## 2 3.023
```

- ► The se.logOR and CI for the non-smokers are not defined because we are using the non-smokers as the baseline group in our calculations.
 - Technically, the se.logOR is 0 and the CI is exactly 1, by definition.
- ► As a result, we set these to be the missing data code NA in the baseline group:

```
mydf[mydf$group=="non-smoker",
        c("se.logOR", "lowerCI", "upperCI")] <-
        c(NA, NA, NA)
mydf</pre>
```

Notice how we referenced multiple columns at once with c("se.logOR", "lowerCI", "upperCI").

More Than Two Exposure Levels

▶ Doll and Hill's data with smokers classified by the average number of cigarettes per day:

		case	control	
Number of	25+	340	182	
cigarettes	15-24	445	408	
per day	1-14	565	706	
	0	7	61	

- ► Can use the last row with 0 cigs per day (unexposed) as a baseline group, and calculate our ORs for each level of exposure.
- ► Here is where the R code we wrote can pay off. We essentially repeat the code, but now refer to the baseline group as "0" instead of "non-smoker".

```
mydf <- data.frame(group=c("25+", "15-24", "1-14", "0"),
                    case=c(340,445,565,7),
                    control=c(182.408.706.61))
library(dplyr)
mydf <- mutate(mydf,total = case+control,</pre>
                prcase = case/total,odds = prcase/(1-prcase))
odds0 <- mydf[mydf$group=="0","odds"]</pre>
c <- mydf[mydf$group=="0","case"]</pre>
d <- mydf[mydf$group=="0","control"]</pre>
mydf <- mutate(mydf, OR=odds/odds0,
                se.logOR = sqrt(1/case + 1/control + 1/c + 1/d))
critval <- gnorm( 0.025, lower.tail=FALSE)</pre>
mydf <- mutate(mydf,
             lowerCI = round(exp(log(OR) - critval*se.logOR),3),
              upperCI = round(exp(log(OR) + critval*se.logOR),3))
mydf <- mutate(mydf, prcase=round(prcase,3), odds=round(odds,3),</pre>
                OR=round(OR,3), se.logOR=round(se.logOR,3))
mydf[mydf$group=="0",c("se.logOR", "lowerCI", "upperCI")] <-</pre>
  c(NA, NA, NA)
```

mydf

##		group	case	${\tt control}$	total	prcase	odds	OR	se.logOR	lowerCI	upperCI
##	1	25+	340	182	522	0.651	1.868	16.279	0.409	7.296	36.325
##	2	15-24	445	408	853	0.522	1.091	9.505	0.405	4.298	21.018
##	3	1-14	565	706	1271	0.445	0.800	6.974	0.403	3.165	15.365
##	4	0	7	61	68	0.103	0.115	1.000	NA	NA	NA