Biostatistics (MATH11230)

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We will start by reproducing the plots we have in slides 7 and 9 of the lectures (sampling distributions of \widehat{OR} and \widehat{OR}). We will also make the corresponding plots for the sampling distributions of \widehat{RR} and \widehat{OR} . The key is to remember that, under a cohort design, the entry a in the 2×2 contingency table (slide 3) follows a binomial distribution whose parameters are the number of exposed individuals and probability of disease given exposure. Similarly, the entry c (in the same contingency table) also follows a binomial distribution whose parameters are, in turn, the number of individuals who are not exposed and the probability of disease given no exposure. We will be assuming that $p_1 = \Pr(D \mid E) = 0.2$ and that $p_2 = \Pr(D \mid \text{not } E) = 0.2$ and, further, that we have 50 individuals in the exposed group and another 50 individuals in the unexposed group. To make the results reproducible, I will be fixing the seed. The code is as follows.

```
n_exp <- 50
n_unexp <- 50
p1 <- 0.2
p2 < -0.2
nsim <- 1000
OR <- RR <- numeric(nsim)
set.seed(123)
for(i in 1:nsim){
  a <- rbinom(1, n_exp, p1)
  c <- rbinom(1, n_unexp, p2)</pre>
  b <- n_exp - a
  d <- n unexp - c
  OR[i] \leftarrow (a*d)/(b*c)
  RR[i] \leftarrow (a/(a+b))/(c/(c+d))
}
df_OR <- data.frame("Mean" = c(mean(OR), mean(log(OR))),</pre>
                  "Median" = c(median(OR), median(log(OR))),
                  "Min" = c(min(OR), min(log(OR))),
                  "Max" = c(max(OR), max(log(OR)))
rownames(df_OR) <- c("OR", "log OR")</pre>
knitr::kable(df_OR, escape = FALSE, digits = 3,
              caption = "Summary statistics of the sampling distributions of the OR and log OR")
```

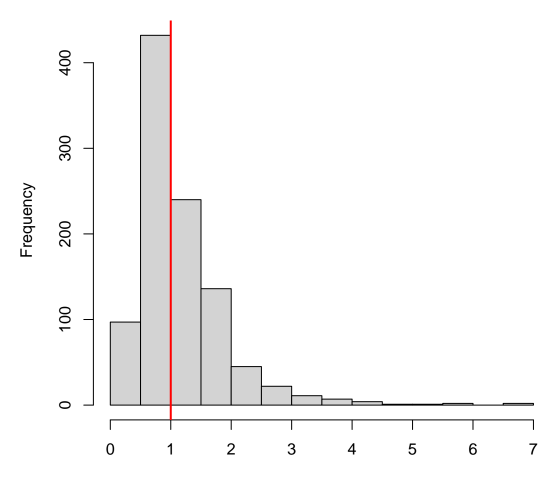
Table 1: Summary statistics of the sampling distributions of the OR and log OR

	Mean	Median	Min	Max
OR	1.193	1	0.202	6.769

	Mean	Median	Min	Max
$\log OR$	0.023	0	-1.599	1.912

```
hist(OR, nclass = 20, xlab = "Estimated odds ratio",
     ylab = "Frequency", main = "Sampling distribution estimated OR")
abline(v = 1, lwd = 2, col = "red")
```

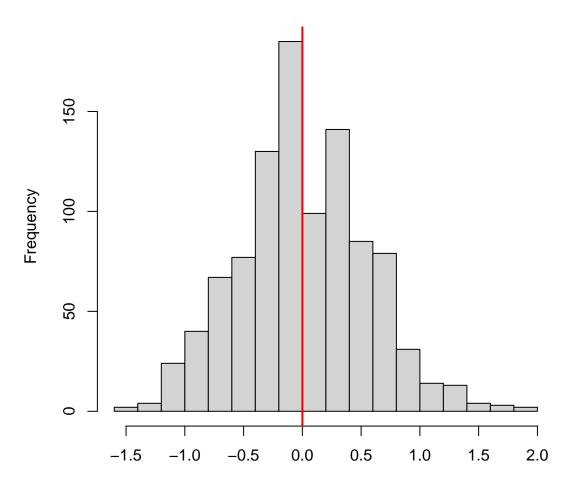
Sampling distribution estimated OR



Estimated odds ratio

```
hist(log(OR), nclass = 20, xlab = "Estimated log odds ratio",
     ylab = "Frequency", main = "Sampling distribution estimated log OR")
abline(v = 0, lwd = 2, col = "red")
```

Sampling distribution estimated log OR



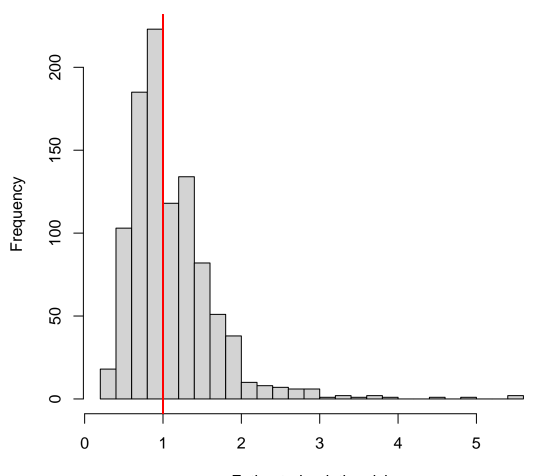
Estimated log odds ratio

Table 2: Summary statistics of the sampling distributions of the RR and \log RR

	Mean	Median	Min	Max
RR	1.129	1	0.250	5.500
$\log RR$	0.019	0	-1.386	1.705

```
hist(RR, nclass = 20, xlab = "Estimated relative risk",
    ylab = "Frequency", main = "Sampling distribution estimated RR")
```

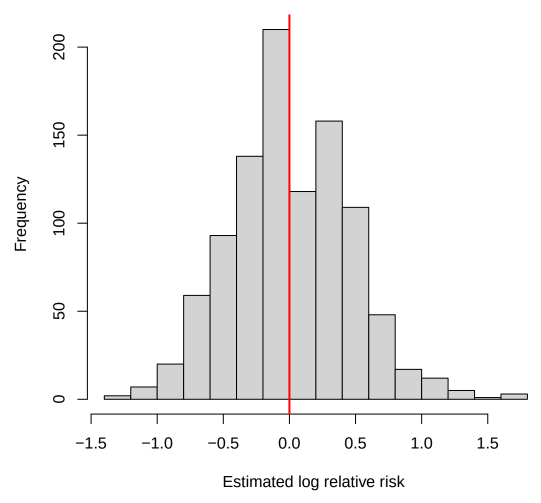
Sampling distribution estimated RR



Estimated relative risk

```
hist(log(RR), nclass = 20, xlab = "Estimated log relative risk ",
    ylab = "Frequency", main = "Sampling distribution estimated log RR")
abline(v = 0, lwd = 2, col = "red")
```

Sampling distribution estimated log RR



I now illustrate the usage of the epiR and epitools packages which, among many things, have functions to compute estimates and CIs for different measures of association. We will use the data from the example in slide 17.

```
require(epiR)
data \leftarrow c(62, 76, 5, 55)
epi.2by2(data, method = "case.control", conf.level = 0.95, units = 100,
         interpret = TRUE, outcome = "as.columns")
##
                              Outcome -
                                              Total
                Outcome +
                                                                           Odds
## Exposed +
                        62
                                     76
                                                138
                                                           0.82 (0.59 to 1.12)
## Exposed -
                        5
                                     55
                                                60
                                                           0.09 (0.02 to 0.20)
## Total
                        67
                                    131
                                                198
                                                           0.51 (0.37 to 0.68)
##
## Point estimates and 95% CIs:
                                                    8.97 (3.38, 23.79)
## Exposure odds ratio
## Attrib fraction (est) in the exposed (%)
                                                   88.75 (69.69, 96.69)
## Attrib fraction (est) in the population (%)
                                                   82.23 (57.71, 92.53)
## Uncorrected chi2 test that OR = 1: chi2(1) = 25.013 Pr>chi2 = <0.001
```

```
## Fisher exact test that OR = 1: Pr>chi2 = <0.001
## Wald confidence limits
## CI: confidence interval
## Measures of association strength:
## The exposure odds among cases was 8.97 (95% CI 3.38 to 23.79) times the exposure odds among control
##
## Measures of effect in the exposed:
## 88.7% of outcomes in the exposed were attributable to exposure (95% CI 69.7% to 96.7%).
##
## Measures of effect in the population:
## 82.2% of outcomes in the population were attributable to exposure (95% CI 57.7% to 92.5%).
require(epitools)
fit <- oddsratio(data, method = "wald", conf = 0.95, correct = FALSE)</pre>
```