# Statistical Methods in Cancer Epidemiology using R

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Lecture 2

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### Contents

- Binary outcomes and proportions
- Comparative parameters of risks and their estimation
- Binomial regression models and comparative parameters
- Adjustment for confounding and evaluation of modification by binomial regression

#### Main R functions covered:

- twoby2() (Epi package)
- ▶ glm()
- ci.lin() (Epi package)

# Outcomes in epidemiologic research

Epidemiologic studies address the occurrence of diseases and other health related phenomena:

(a) cross-sectional: **prevalence** of diseases,

(b) longitudinal: disease incidence, and mortality

Often we want to compare the prevalence or incidence of disease between two groups defined by a binary  $risk\ factor\ X$ 

 $\triangleright$  X=1: exposed X=0: unexposed

### Types of outcome variables

- ightharpoonup Binary (0/1) variables at individual level
  - disease status at a time point
  - change of status, event or transition ({e.g.} from healthy to diseased)
- Proportions at group level
  - prevalence
  - incidence proportion or cumulative incidence,
- Rates of events
  - incidence or mortality rate (per 1000 y)
  - car accidents (per million km)
- Time to event
  - survival time (often censored)

# Incidence and prevalence proportions}

▶ Incidence proportion (R) of a binary (0/1) outcome (disease, death etc.) over a fixed risk period is defined

$$R = \frac{D}{N} = \frac{\text{number of new cases during period}}{\text{size of population-at-risk at start}}$$

Also called {cumulative incidence} (or even "risk").\ NB.

This formula requires complete follow-up, i.e. no {censorings}, and absence of {competing risks}.

Prevalence (proportion) P of disease at time point t

$$P = \frac{\text{no. of existing cases at t}}{\text{total population size at t}}.$$

### Two-group comparison

- ▶ Binary risk factor *X*: exposed vs. unexposed.
- Summarizy results from cohort study with fixed risk period and no losses:

Exposure	Cases	Non-cases	Group size
yes	$D_1$	$C_1$	$N_1$
no	$D_0$	$C_0$	$N_0$
total	$D_+$	<i>C</i> <sub>+</sub>	$N_+$

▶ Incidence proportions in the two exposure groups

$$R_1 = \frac{D_1}{N_1}, \qquad R_0 = \frac{D_0}{N_0}.$$

► These are crude *estimates* of the true *risks*  $\pi_1$ , and  $\pi_0$  of outcome in the two exposure categories.

### Example: Observational clinical study

Treatment failure in two types of operation for renal calculi (Charig et al. 1986. BMJ 292: 879-882)

- ► OS = open surgery (invasive)
- ► PN = percutaneous nephrolithotomy

Treatment	Failure	Success	Patients	Failure-%
$group\;(j)$	$(D_j)$	$(C_j)$	$(N_j)$	$(R_j)$
OS $(j = 1)$	77	273	350	22.0
PN $(j = 0)$	60	290	350	17.1

Crude incidence proportions of treatment failure:

$$R_1 = 77/350 = 22.0\%, \qquad R_0 = 60/350 = 17.1\%$$

### Risks and their comparative parameters

The **risk** or **probability** of binary outcome (e.g. new case of disease) in the exposed  $\pi_1$  and in the unexposed  $\pi_0$  as to binary risk factor X (values 1 and 0) are typically compared by

- risk difference  $\theta = \pi_1 \pi_0$
- lacktriangledown risk ratio  $\phi=\pi_1/\pi_0$
- odds ratio (risk odds ratio)

$$\psi = \frac{\pi_1/(1-\pi_1)}{\pi_0/(1-\pi_0)}$$

The odds ratio is close to the risk ratio when the risks are small (less than 0.1 – the rare-disease assumption).

# Odds and Odds Ratio (OR)

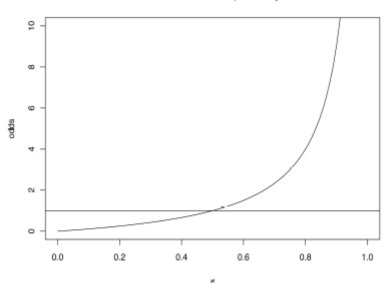
The **odds** ( $\Omega$ ) is the probability of binary outcome  $P(Y=1)=\pi$  divided by the the probability of binary outcome  $P(Y=0)=1-\pi$ .

$$\Omega = \frac{\pi}{1-\pi}$$

- ▶ Odds of 2.5 means that the probability of Y=1 (success) is two and half times higher than the probability of Y=0 (failure)
- Odds 0.5 means that success probability of success is 50% of the probability of failure
- Odds of 1 implies that probability of both outcomes 0.5 (equal)

# Probability and odds

### Odds as function of probability



### Risks and comparative parameters estimated

The risks  $\pi_1$  and  $\pi_0$  are estimated by empirical incidence proportions  $R_1 = D_1/N_1$ , and  $R_0 = D_0/N_0$ .

Crude estimates of comparative parameters

- ▶ incidence proportion difference  $RD = R_1 R_0$
- ▶ incidence proportion ratio  $RR = R_1/R_0$
- incidence odds ratio

$$OR = \frac{R_1/(1-R_1)}{R_0/(1-R_0)}$$

**NB**. To remove *confounding*, the estimated must be adjusted for relevant *confounders*.

# Example: OS vs. PN (cont'd)

Crude estimates of true risk difference  $\theta$ , risk ratio  $\phi$ , and  $\setminus$  odds ratio  $\psi$  between OS and PN:

RD = 
$$\frac{77}{350} - \frac{60}{350} = 0.22 - 0.171 = +0.049 (+4.9\%)$$
  
RR =  $\frac{77/350}{60/350} = \frac{77/60}{350/350} = \frac{0.22}{0.171} = 1.283$   
OR =  $\frac{77/273}{60/290} = \frac{0.22/(1 - 0.22)}{0.171/(1 - 0.171)} = 1.363$ 

PN appears more successful than OS.

Is this (a) true, (b) due to bias, or (c) due to chance?

# Example: OS vs. PN (cont'd)

Standard error of log(RR), 95% error factor (EF) of RR, and 95% CI for true risk ratio  $\phi$ :

SEL = 
$$\sqrt{\frac{1}{73} + \frac{1}{60} - \frac{1}{350} - \frac{1}{350}}$$
  
= **0.1547**  
EF =  $\exp\{1.96 \times 0.1547\}$   
= **1.3543**  
CI = [1.2833/1.3543, 1.2833 × 1.3543]  
= [**0.9476, 1.7380**].

### Estimating comparative parameters in R

- ► A multitude of R functions in several packages are readily available for point estimation and CI calculation using either "exact" or/and various approximative methods.
- We shall here demonstrate the use of function twoby2() in the Epi package. It applies the simple Wald approximations as described above, but for
  - risk difference: the Newcombe method is used, and
  - odds ratio: the exact conditional method is also available.
- ► Hence, similar results are expected as obtained above.

### Use of function twoby2()

Loading the Epi package:

### library(Epi)

▶ Reading the counts of the 2 x 2-table into a matrix:

```
counts <- c(77, 273, 60, 290)
tab <- matrix( counts, nrow=2, byrow=T)</pre>
```

▶ Viewing the contents of the matrix/table:

#### tab

```
[,1] [,2]
[1,] 77 273
[2,] 60 290
```

### Make 2 by 2 table

twoby2(tab)

Calling the function with tab as its argument:

```
2 by 2 table analysis:
Outcome · Col 1
Comparing: Row 1 vs. Row 2
     Col 1 Col 2 P(Col 1) 95% conf. interval
                0.2200 0.1797 0.2664
       77 273
Row 1
Row 2
        60 290
                0.1714 0.1355 0.2146
                               95% conf. interval
           Relative Risk: 1.2833
                                  0.9476 1.7380
        Sample Odds Ratio: 1.3632 0.9362 1.9851
Conditional MLE Odds Ratio: 1.3626 0.9206 2.0237
   Probability difference: 0.0486 -0.0103 0.1071
           Exact P-value: 0.1272
       Asymptotic P-value: 0.1061
```

### Analyses based on binary regression model

Crude estimates and CIs for the comparative parameters can also be obtained by fitting appropriate **binary regression models** for the numbers  $D_j$  or proportions  $R_j$ .

Special cases of generalized linear models (GLM) with

- (i) random part:  $D_j$  is assumed to obey the binomial distribution or  $\{family\}$  with index  $N_j$  and probability  $\pi_j$ ,
- (ii) systematic part: linear predictor  $\eta_j = \alpha + \beta X_j$ , in which  $X_j = 0$  for unexposed and  $X_j = 1$  for exposed,
- **| (iii) link function**: g(.) that connects the probability  $\pi_j$  and the systematic part  $\eta_j$  by:

$$g(\pi_j) = \eta_j = \alpha + \beta X_j$$

# Link functions and comparative parameters

General model:  $g(\pi_j) = \alpha + \beta X_j$  for the risks by binary X

- ▶ identity link:  $g(\pi_j) = \pi_j = \alpha + \beta X_j$ :\\\  $\Rightarrow \beta = \pi_1 \pi_0 = \theta$ , \\\  $= \frac{\text{risk difference}}{1}$  btw  $X_j = 1$  and  $X_j = 0$
- ▶ **logit** link:  $g(\pi_j) = \log[\pi_j/(1 \pi_j)]$

$$\Leftrightarrow \quad \pi_j = \frac{1}{1 + \exp\{-(\alpha + \beta X_i)\}} = \exp(\alpha + \beta X_j)$$

# Logit model for odds ratio

Substituting logit function for g(.) and values of  $X_j$  we get

$$\begin{array}{lcl} \log \left( \frac{\pi_0}{1-\pi_0} \right) & = & \alpha = \text{baseline logit} \\ \log \left( \frac{\pi_1}{1-\pi_1} \right) & = & \alpha + \beta. \end{array}$$

This implies

$$\begin{array}{lcl} \pi_0 & = & \frac{1}{1+\exp(-\alpha)}, & \pi_1 = \frac{1}{1+\exp\{-(\alpha+\beta)\}}, \\ \\ \beta & = & \log\left\{\frac{\pi_1/(1-\pi_1)}{\pi_0/(1-\pi_0)}\right\} \\ \\ e^\beta & = & \frac{\pi_1/(1-\pi_1)}{\pi_0/(1-\pi_0)} = \psi = \underline{\text{odds ratio}} \text{ btw exp'd and unexp'd}. \end{array}$$

# Fitting binary regression models in R

### Function glm()

estimation method: maximum likelihood (ML),computation algorithm: IWLS.

### Key arguments of glm():

- ightharpoonup model formula: *response*  $\sim$  expression of regressors
- weights = group sizes  $N_j$  when proportions  $R_j$  are given as the response (outcome) variable,
- family = binomial(link = 'log'), if risk ratio, family = binomial(link = 'logit'), if odds ratio, family = binomial(link = 'identity'), if risk diff'ce is the parameter of interest.

# Example - Colorectal screenning RHS study

	All		Males		Females	
	Screening	Control	Screening	Control	Screening	Control
Number of persons	180 210	180 282	89 712	89 807	90 498	90 475
Person-years	805 480	805 693	395 614	395 851	409 866	409 843
Deaths						
All causes	8000	7963	5486	5453	2514	2510
Colorectal cancer	170	164	93	106	77	58
Patients with colorectal cancer						
Number of patients	903	811	525	477	378	334
Incidence rate per 100 000 person-years	112.4	100.9	133.1	120.8	92.4	81.7
Person-years	2285	1805	1318	1016	967	790
Deaths	202	190	123	127	79	63
Expected number of deaths*	25.0	21.1	18.6	15.7	6.4	5.5
Excess number of deaths†	177.0	168.9	104.4	111.3	72.6	57.5
CRC incidence rate ratio (95% CI)	1.11 (1.01 to 1.23)	1	1.10 (0.97 to 1.25)	1	1.13 (0.98 to 1.31)	1
Mortality rates per 100 000						
All causes	993	988	1387	1378	613	612
Non-colorectal cancer causes	972	968	1363	1351	595	598
Colorectal cancer	21.1	20.4	23.5	26.8	18.8	14.2
Excess mortality due to CRC	21.7	21.0	26.4	28.1	17.7	14.0
Mortality rate ratios						
All causes	1.00 (0.97 to 1.04)	1	1.01 (0.97 to 1.05)	1	1.00 (0.95 to 1.06)	1
Non-colorectal cancer causes	1.00 (0.97 to 1.04)	1	1.01 (0.97 to 1.05)	1	0.99 (0.94 to 1.05)	1
Colorectal cancer	1.04 (0.84 to 1.28)	1	0.88 (0.66 to 1.16)	1	1.33 (0.94 to 1.87)	1
Excess mortality due to CRC	1.05 (0.84 to 1.30)	1	0.94 (0.71 to 1.24)	1	1.26 (0.88 to 1.80)	1

### Example - data to R

### CRC parient mortality between arms

- two observations (one for each treatment group),
- three variables:
  - ▶ treat = arm, with values 1 = Screen, 0 = control,
  - ightharpoonup d = number of CRC (patient) deaths  $D_i$ ,
  - npat = number of patients  $N_j$ ,
- variable vectors defined:

```
treat <- c(0, 1)
fail <- c(190, 202)
npat <- c(811, 903)
prop <- fail/npat
c(prop*100,prop[2]/prop[1])</pre>
```

[1] 23.4278668 22.3698782 0.9548406

### Estimation of risk ratio

▶ Defining the *model object*:

```
RRmodel <- glm( prop ~ treat, family=binomial(link='log'),</pre>
```

Estimation results extracted by function ci.lin() in Epi
(two columns of the whole output omitted for clarity)

```
library(Epi)
round( ci.lin(RRmodel, Exp=T)[, -(3:4)], 4)
```

```
Estimate StdErr exp(Est.) 2.5% 97.5% (Intercept) -1.4512 0.0635 0.2343 0.2069 0.2653 treat -0.0462 0.0887 0.9548 0.8024 1.1362
```

- ▶ The estimate of  $\beta$  is  $\widehat{\beta} = -0.0462 = \log(0.9548)$ , and that of mortality (ratio) ratio  $\phi$  is  $RR = \exp(-0.0462) = 0.9548$ .
- ► Estimate of  $\alpha$  is  $\hat{\alpha} = -1.4512 = \log(0.2342787) = \log(R_0)$ .

### Estimation of odds ratio

```
round( ci.lin(ORmodel, Exp=T)[, -(3:4)], 4)
```

```
Estimate StdErr exp(Est.) 2.5% 97.5% (Intercept) -1.1843 0.0829 0.3060 0.2601 0.3599 treat -0.0599 0.1151 0.9418 0.7516 1.1802
```

- ► The estimate of  $\beta$  is  $\widehat{\beta} = -0.0599 = \log(0.9418)$ , and the estimated  $\psi$  is OR = exp(-0.0599) = 0.9418.
- ▶ The estimate of  $\alpha$  is  $\hat{\alpha} 1.1843 = \log(0.3060)$ , in which 0.3060 = 0.2342787/(1 0.2342787) is the estimated baseline odds  $R_0/(1 R_0)$ .

### Estimation of risk difference

```
RDmodel <- glm( prop ~ treat,
  fam=binomial(link='identity'), w=npat)</pre>
```

```
round( ci.lin(RDmodel), 3)

Estimate StdErr z P 2.5% 97.5%
(Intercept) 0.234 0.015 15.752 0.000 0.205 0.263
treat -0.011 0.020 -0.520 0.603 -0.050 0.029
```

- Again, same results obtained as with *e.g.* twoby2(), although CIs are here based on Wald statistic.
- ▶ NB. Fitting binomial model with this link easily fails with more complicated models, especially involving continuous variables.

### Same with twoby2

```
counts<-c(fail[2],npat[2]-fail[2],fail[1],npat[1]-fail[1])
counts</pre>
```

#### [1] 202 701 190 621

```
twoby2(matrix( counts, nrow=2, byrow=T))
2 by 2 table analysis:
```

Outcome : Col 1

Comparing: Row 1 vs. Row 2

```
Col 1 Col 2 P(Col 1) 95% conf. interval
Row 1 202 701 0.2237 0.1977 0.2520
Row 2 190 621 0.2343 0.2064 0.2647
```

```
| 95% conf. interval
| Relative Risk: 0.9548 | 0.8024 | 1.1362
| Sample Odds Ratio: 0.9418 | 0.7516 | 1.1802
| Conditional MLE Odds Ratio: 0.9419 | 0.7468 | 1.1881
| Probability difference: -0.0106 | -0.0505 | 0.0291
```

Exact P-value: 0.6048 Asymptotic P-value: 0.6026