### Faculty of Health Sciences



# Day 5: binary responses and $2\times 2$ tables

Basic Statistics for health researchers

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# Outline/Intended Learning Outcomes (ILOs)

#### **Preliminaries**

ILO: calculate 95% Cls for population proportions

ILO: distinguish between exact and approximate (asymptotic) 95% CIs

#### Group comparison

ILO: to define a suitable association measure and compute its 95% C

ILO: to (correctly) use the  $\chi^2$  test and Fisher's test

#### Sample size and power calculation

ILO: to identify why and how to make power and sample size calculations

ILO: to analyse their strengths and limitations

#### Confounding

ILO: to exemplify confounding and its potential to be misleading

ILO: to name two commonly used remedies

#### Cohort vs case-control study

ILO: to differentiate the cohort and case-control designs

ILO: to restate which association measure(s) can be used for each design

#### Screening: jargon

ILO: to recognize some jargon

#### Paired binary data (if time allows)

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### Binary outcome

$$Y = \begin{cases} 1 & \text{event / positive / disease} \\ 0 & \text{no event / negative / non-disease} \end{cases}$$



## Binary outcome

$$Y = \begin{cases} 1 & \text{event / positive / disease} \\ 0 & \text{no event / negative / non-disease} \end{cases}$$

#### **Parameters**

Prevalence: proportion of the population with event at fixed time point.

How many have the disease right now?

► Risk: probability that event occurs in given time period:

How likely will a subject acquire the disease within 1-year?



### Statistical inference

### Estimating risks and prevalence

$$\widehat{p} = \text{Relative frequency} = \frac{\text{Number of events}}{\text{Number of subjects}} = \frac{x}{n}$$

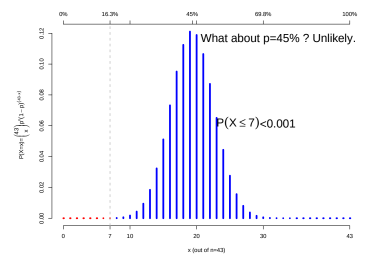
Confidence limits: normal approximation ("large"  $n^1$ )

$$\left[\widehat{p} - 1.96\sqrt{\frac{\widehat{p}(1-\widehat{p})}{n}}; \widehat{p} + 1.96\sqrt{\frac{\widehat{p}(1-\widehat{p})}{n}}\right]$$

Confidence limits: "exact" (any n)

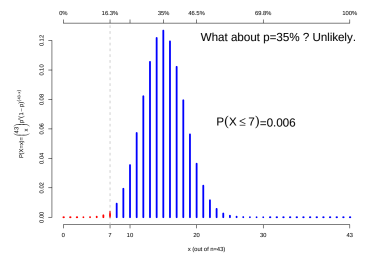






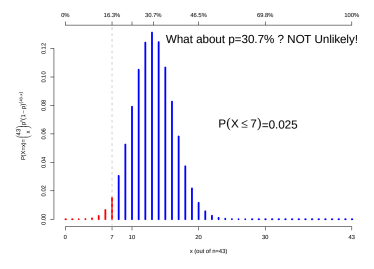






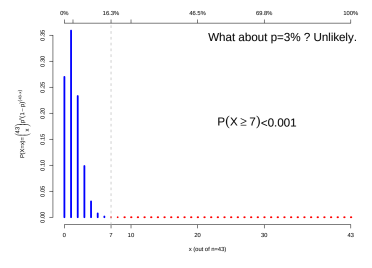






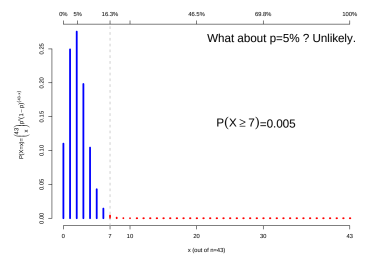






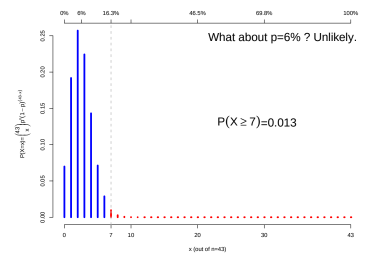






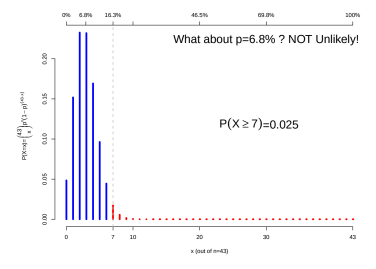






 $\blacktriangleright$  x=7 and n=43 leads to  $\hat{p}=16.3\%$  and 95% CI= [6.8;30.7].

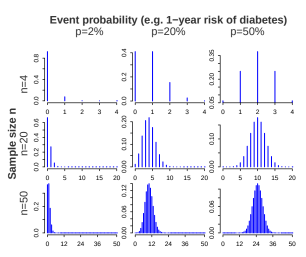




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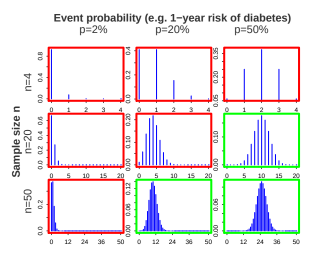
# Normal approximation



▶ Binomial distribution:  $P(X = x) = \binom{N}{x} p^x (1-p)^{N-x}$ 



## Normal approximation



<sup>• &</sup>quot;good"approximation if  $np \ge 5$  and  $n(1-p) \ge 5$  (green boxes).





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# Case: clinical trial on Dalteparin <sup>3</sup>

- ▶ Placebo (n = 42)
- ▶ Dalteparin (n = 43)



#### Outcome

tcome:				
	Category <sup>2</sup>	Label		
	intact skin	healed		
	decreased ulcer area $\geq 50\%$	improved		
	increased ulcer area $\geq 50\%$	impaired		
	decreased or increased ulcer area $< 50\%$	unchanged		
	amputation above/below ankle	amputation		

Research question: Does Dalteparin improve the outcome, when injected once daily until ulcer healing or for a maximum of 6 months?



<sup>&</sup>lt;sup>2</sup>mutually exclusive.

<sup>&</sup>lt;sup>8/60</sup> <sup>3</sup>Kalani et al. *Diabetes Care* **26**: 2575-2580, 2003

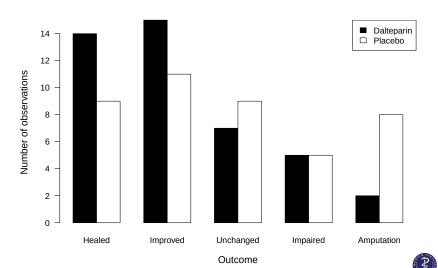
# Frequency table

	Dalteparin	Placebo
Healed	14 (33%)	9 (21%)
Improved	15 (35%)	11 (26%)
Unchanged	7 (16%)	9 (21%)
Impaired	5 (12%)	5 (12%)
Amputation	2 (5%)	8 (19%)
total (100%)	43	42

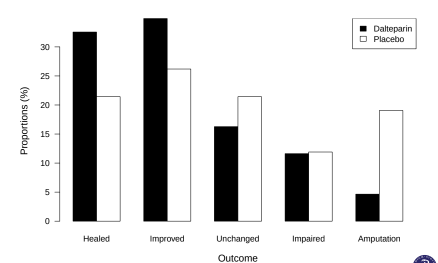
- ► Summarizes the outcome data.
- ► Prepare/Format data for analyzes.



# Barplot (frequencies)



# Barplot (proportions<sup>4</sup>)



# Here we pool the outcome categories as follows

Category	Dichotomized outcome
intact skin	better
ulcer area decreased $\geq 50\%$	
decreased or increased ulcer area $< 50\%$	
increased ulcer area $\geq 50\%$	worse
amputation above/below ankle	

**Important:** this dichotomization should be prespecified (i.e. decision made before seeing the data).  $^{5}$ 

<sup>&</sup>lt;sup>5</sup> For an illustration of why prespecification matters, see e.g. Austin & Goldwasser. "Pisces did not have increased heart failure: data-driven comparisons of binary proportions between levels of a categorical variable can result in incorrect statistical significant levels." Journal of clinical epidemiology 61.3 (2008): 295-300.

### Group comparison

### Placebo group

Risk of worse outcome 
$$=\frac{22}{42}=\widehat{p}_1$$

### Dalteparin group

Risk of worse outcome 
$$=\frac{14}{43}=\widehat{p}_2$$



### Group comparison

### Placebo group

Risk of worse outcome = 
$$\frac{22}{42} = \hat{p}_1$$

### Dalteparin group

Risk of worse outcome = 
$$\frac{14}{43} = \hat{p}_2$$

### Association measures<sup>6</sup>

Relative risk: 
$$\frac{\widehat{p}_1}{\widehat{p}_2}$$
 Odds ratio:  $\frac{\widehat{p}_1}{\frac{1-\widehat{p}_1}{\widehat{p}_2}}$  Risk difference:  $\widehat{p}_1-\widehat{p}_2$ 

$$\frac{1-\widehat{p_1}}{\widehat{p_2}}$$

$$1-\widehat{p_2}$$



<sup>&</sup>lt;sup>6</sup>whenever possible, we prefer using risk ratios or risk differences to odds ratios. They are often better understood and easier to communicate!

# 2x2 contingency table

### Response

Exposure –

	yes	no	total
yes	а	b	a+b
no	С	d	c+d
total	a+c	b+d	N

### Risk estimates

$$\widehat{p}_1 = \frac{a}{a+b}$$
  $\widehat{p}_2 = \frac{c}{c+d}$ 



### Relative risk

# $\widehat{RR} = \frac{a/(a+b)}{c/(c+d)}$

Exposure

Response				
	yes	no	total	
yes	а	b	a+b	
no	С	d	c+d	
total	a+c	b+d	N	

Standard error of  $\log(\widehat{RR})$  and confidence interval of RR  $^7$ 

$$\widehat{\sigma} = \sqrt{\frac{1}{a} - \frac{1}{a+b} + \frac{1}{c} - \frac{1}{c+d}}$$

$$CI_{95\%} = \left[\widehat{RR} \cdot \exp(-1.96\,\widehat{\sigma}) \; ; \; \widehat{RR} \cdot \exp(1.96\,\widehat{\sigma})\right]$$



# Relative risk: placebo versus dalteparin

$$\widehat{RR} = \frac{22/42}{14/43} = 1.609$$

Treatment

	Outcome			
	worse	better	total	
placebo	22	20	42	
dalteparin	14	29	43	
total	36	49	85	

Standard error of  $\log(\widehat{RR})$  and confidence interval

$$\hat{\sigma} = \sqrt{\frac{1}{22} - \frac{1}{42} + \frac{1}{14} - \frac{1}{43}} = 0.264$$

$$CI_{95\%} = [0.959; 2.7]$$
 (does include 1)



### Risk difference

$$\widehat{\Delta} = \frac{a}{a+b} - \frac{c}{c+d}$$

Exposure

Response				
	yes	no	total	
yes	а	b	a+b	
no	С	d	c+d	
total	a+c	b+d	N	

Standard error of  $\widehat{\Delta}$  and confidence interval <sup>8</sup>

$$\widehat{\sigma} = \sqrt{ab/(a+b)^3 + cd/(c+d)^3}$$

$$CI_{95\%} = \left[ \widehat{\Delta} - 1.96 \,\widehat{\sigma} \, ; \, \widehat{\Delta} - 1.96 \,\widehat{\sigma} \right]$$



# Risk difference: placebo versus dalteparin

Treatment

$\widehat{\Lambda}$		22		14		0.100
Δ	=	$\overline{42}$	_	$\overline{43}$	=	0.198

	worse	better	total
placebo	22	20	42
dalteparin	14	29	43
total	36	49	85

Outcome

Standard error of  $\widehat{\Delta}$  and confidence interval

$$\widehat{\sigma} = \sqrt{22 \cdot 20/42^3 + 14 \cdot 29/43^3} = 0.105$$

 $CI_{95\%} = [-0.008 ; 0.404]$  (does include 0)



# Odds Ratio (OR)

#### Concept needed for

- case-control studies
- logistic regression

Odds: are defined as "risk of event divided by risk of no event"

$$\mathsf{odds} = p/(1-p) \ ,$$

and the risk can be computed back from the odds,  $p={\rm odds}/(1+{\rm odds})$ . Odds are difficult to interpret, but if risks are small, then risks  $\approx$  odds.



The Odds ratio (OR) is defined as the ratio of the odds,

$$OR = \frac{\mathsf{odds}_1}{\mathsf{odds}_2} = \frac{p_1/(1-p_1)}{p_2/(1-p_2)} \ .$$

OR are difficult to interpret, but from the equation...

$$RR = \frac{OR}{\left\{1 - p_2\right\} + p_2 OR},$$

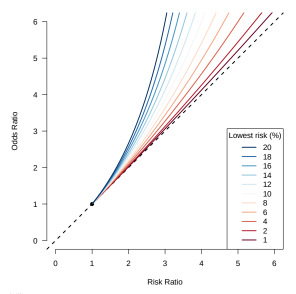
...we can first conclude:

- $ightharpoonup OR > 1 \Leftrightarrow RR > 1$
- $ightharpoonup OR = 1 \Leftrightarrow RR = 1$
- $ightharpoonup OR < 1 \Leftrightarrow RR < 1$

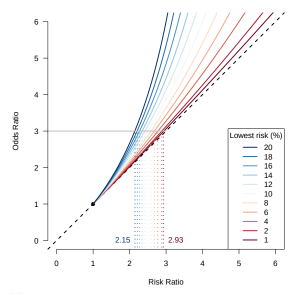
...and further conclude that

- ▶ the OR is sufficient to deduce whether a risk increases or decreases.
- ▶ if  $p_2$  is small (e.g. rare disease), then  $OR \approx RR$ .

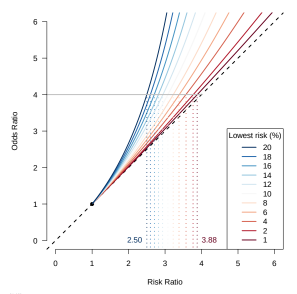




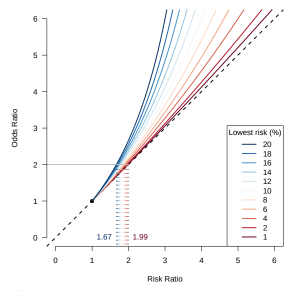














### Odds ratio

 $\widehat{OR} = \frac{\frac{a/(a+b)}{b/(a+b)}}{\frac{c/(c+d)}{d/(c+d)}} = \frac{a \cdot d}{b \cdot c}$ 

Exposure

Response				
	yes	no	total	
yes	a	b	a+b	
no	С	d	c+d	
total	a+c	b+d	N	

Standard error of  $\log(\widehat{OR})$  and confidence interval<sup>9</sup>

$$\widehat{\sigma} = \sqrt{\frac{1}{a} + \frac{1}{b} + \frac{1}{c} + \frac{1}{d}}$$

$$CI_{95\%} = \left[\widehat{OR} \cdot \exp(-1.96\,\widehat{\sigma}); \widehat{OR} \cdot \exp(1.96\,\widehat{\sigma})\right]$$



<sup>&</sup>lt;sup>22/60</sup> <sup>9</sup>This method is "good enough" with "large enough" sample sizes.

# Odds ratio: placebo versus dalteparin

# $\widehat{OR} = \frac{22 \cdot 29}{14 \cdot 20} = 2.279$

Treatment

	worse	better	total
placebo	22	20	42
dalteparin	14	29	43
total	36	49	85

Outcome

Standard error of  $\log(\widehat{OR})$  and confidence interval

$$\widehat{\sigma} = \sqrt{\frac{1}{22} + \frac{1}{20} + \frac{1}{14} + \frac{1}{29}} = 0.449$$

$$CI_{95\%} = [0.946; 5.491]$$
 (does include 1)



## Reporting results

The relative risk (of worsening) of group 1 (Dalteparin) versus group 2 (Placebo) is estimated as

$$RR = \frac{14/43}{22/42} = 0.622$$

## Equivalent statements:

- The risk in group 1 is reduced by a factor 0.622 compared to group 2.
- ► The risk in group 1 is 37.8% lower than in group 2.10
- ► The risk in group 2 is 1.609 times higher than in group 1.11
- ► The risk in group 2 is 60.9% higher than in group 1.



<sup>&</sup>lt;sup>10</sup>because 1-0.622=0.378

## Testing independence in a randomized clinical trial

Null hypothesis  $H_0$ : the treatment has no effect.

$$\begin{array}{lll} \operatorname{Prob}(\operatorname{worse\ given\ dalteparin}) = \operatorname{Prob}(\operatorname{worse\ given\ placebo}) \\ \Leftrightarrow & p_1 - p_2 = 0 & (\operatorname{Difference} = 0) \\ \Leftrightarrow & \frac{p_1}{p_2} = 1 & (\operatorname{Relative\ risk} = 1) \\ \Leftrightarrow & \frac{p_1/(1-p_1)}{p_2/(1-p_2)} = 1 & (\operatorname{Odds\ ratio} = 1) \end{array}$$

Popular tests of independence between the treatment group and the outcome groups:

- $\triangleright \chi^2$  test (normal approximation)<sup>12</sup>
- ► Fisher's exact test: recommended as the default choice! 13

<sup>&</sup>lt;sup>12</sup>This method is "good enough" with "large enough" sample sizes.





# The $\chi^2$ test statistic

$$\chi^2 = \sum \frac{(\text{observed counts} - \text{expected counts})^2}{\text{expected counts}}$$

#### Observed counts

Exposure

Response									
	yes	no	total						
yes	a	b	a+b						
no	С	d	c+d						
total	a+c	b+d	N						

### **Expected counts**

The expected counts are calculated under the null hypothesis.

Exposure

	yes	no	total
yes	(a+b)(a+c)/N	(a+b)(b+d)/N	a+b
no	(c+d)(a+c)/N	(c+d)(b+d)/N	c+d
total	a+c	b+d	N

Response

Rule of thumb: a valid analysis requires that all expected counts are  $\geq 5$ .

under the null hypothesis the groups are identical, hence data can be merged into a single group

### Test results

#### Null hypothesis:

dalteparin treatment has no effect for chronic foot ulcers.

Test	p-value
Fisher's exact test	0.0808
Pearson's $\chi^2$ test	0.0644
Pearson's $\chi^2$ test with Yates' continuity correction <sup>14</sup>	0.1032

#### R code:

```
tab <- rbind(c(22,20),c(14,29))
fisher.test(tab)  # always works (default choice!)
chisq.test(tab,correct=FALSE)  # fine with large samples
chisq.test(tab,correct=TRUE)  # no longer useful</pre>
```

<sup>&</sup>lt;sup>14</sup>Expected to be more precise than the usual Pearson's  $\chi^2$  test when the sample size is very small. **NOT RECOMMENDED**, with small sample sizes, use Fisher's test instead.

## A note of caution

Because the (simple) formulas for the 95% CI (of the previous slides) are based on large sample size approximations, they are not necessarily consistent with the result of the Fisher's exact test, especially with "very small" sample sizes.

Example:

	event	no event
exposed	5	12
non-exposed	8	3

- $\widehat{p}_1 = 8/11 = 0.73, \quad \widehat{p}_2 = 5/17 = 0.29.$
- $\triangle$   $\hat{\Delta}$  = 0.43 (0.09; 0.77)
- $ightharpoonup \widehat{RR} = 2.47 \ (1.09 \ ; 5.62)$
- $\widehat{OR}$ = 6.40 (1.18; 34.61)
- ightharpoonup p-values from Fisher's exact test and Pearson's  $\chi^2$  (with and without Yates correction) are 0.051, 0.063 and 0.025, respectively.

Here the confidence intervals show a significant result, but not Fisher's test.



Advanced methods and software<sup>15</sup> are available to avoid running into this kind of inconsistency between hypothesis test and confidence intervals.

Fortunately, it is rare that we run into this problem.... and even rarer that it matters for the interpretation.



# Larger contigency tables (1/2)

If the table is not 2x2 but, e.g., 3x4 or 2x4, the  $\chi^2$  test and Fisher's exact test are testing an "ANOVA-like" null hypothesis similarly to what the F-test does to compare several means.

### First example:

	underweight	normal	overweight	obese
no SCD	9	51	20	8
SCD	23	61	3	1

#### R code:

fisher.test(table(d\$SCD,d\$BMIgroup))

returns a p-value <0.001, for the null hypothesis

 $H_0$ : "the prevalence of SCD is the same in all groups of BMI"

that is, "no association between BMI group and SCD".



# Larger contigency tables (2/2)

### Second example:

	underweight	normal	overweight	obese
age=[16, 25)	14	45	1	1
[25, 30)	3	25	3	1
[30, 67]	15	42	19	7

#### R code:

fisher.test(table(d\$ageGroup,d\$BMIgroup))
returns p-value=0.004, for the null hypothesis

 $H_0$ : "the prevalence of each BMI group is the same in all groups of age"

that is, "no association between BMI group and age".



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## Textbook formula ("large n" approximation)

$$n = \frac{\left\{z_{\alpha/2}\sqrt{2\bar{p}(1-\bar{p})} + z_{\beta}\sqrt{p_1(1-p_1) + p_2(1-p_2)}\right\}^2}{(p_1 - p_2)^2}$$

- $z_{\alpha/2} = -1.96$  for  $\alpha = 5\%.^{16}$
- $ightharpoonup z_{1-\beta} = 0.84$  and 1.28 for  $1 \beta = 80\%$  and 90%.
- $\bar{p} = (p_1 + p_2)/2.$
- ▶ *n*: number of observations in **each** group.

### Useful for computing:

- ▶ Sample size: n for given "guesses" of  $p_1$  and  $p_2$  and desired  $1 \beta$  and  $\alpha$ .
- ▶ Power for a given budget/sample size:  $1 \beta$  for "guesses" of  $p_1$  and  $p_2$  and desired n and  $\alpha$ .
- ▶ Least detectable difference (or ratio):  $\delta=p_1-p_2$  (or  $r=p_1/p_2$ ) for given n, "guess" of  $p_1$  and desired  $\alpha$  and minimal power  $1-\beta$ .



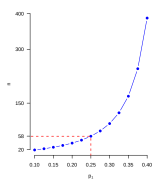
## Sample size calculation

### Standard software can be used, e.g. R:

```
power.prop.test(p1 = 0.25, p2 = 0.5, power=0.8)
```

Two-sample comparison of proportions power calculation

NOTE: n is number in \*each\* group



▶ n = 58 subjects needed in **each** group (i.e. 116 in total) to detect significant risk difference with a power of 80%, if the risks in the two groups are 25% and 50%.

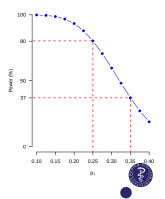
### Power calculation

**Example:** an initial calculation suggests n=58 subjects per group (i.e. 116 in total), for detecting a difference of 25% survival between the two groups, assuming 50% survival in the placebo group (with 80% power). But what does the power become if we were too optimistic with the expected treatment effect? E.g. what if the difference in survival probability is only 15%?

```
power.prop.test(n=58, p1 = 0.35, p2 = 0.5)
```

Two-sample comparison of proportions power calculation

NOTE: n is number in \*each\* group



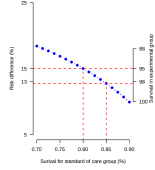
## Least detectable difference

Example: My grant can finance a total sample size of n=150 (i.e. 75 per group). What is the smallest survival difference that I can hope to show with a decent power (e.g. 80%), if I expect 80% survival in the "standard of care" (i.e. control) group? And if I expect 85% in the "standard of care" group?

```
power.prop.test(n=75, p1 = 0.8, power=0.8)
```

 ${\small {\tt Two-sample \ comparison \ of \ proportions \ power \ calculation}}$ 

NOTE: n is number in \*each\* group



**Note:** you need to supply a value for p1, not p2, otherwise the software is looking for a lower risk and it returns 0.72

# Digression: Tables also exist (for sample size calculation) 17

TABLE II—Sample sizes to detect a difference in two proportions,  $p_A$  and  $p_B$ , at a 5% significance level with 80% power

	$P_B$																			
PA	0.05	0-10	0.15	0.20	0.25	0.30	0.35	0.40	0.45	0.50	0.55	0.60	0.65	0.70	0.75	0.80	0.85	0.90	0.95	1.00
0.00	152	74	48	35	27	22	18 27	15 22	13	11	10	.8	7	6	6	5	4	4	3	2
0·05 0·10		435	141 686	76 199	49 100	36 62	43	32	18 25	15 20	12 16	11 14	11	10	8	7	6	5	4	4
0·15 0·20				906	250 1094	121 294	73 138	49 82	36 54	27 39	22 29	17 23	14 18	12 15	10 12	8 10	7 8	6 7	5 6	4 5
0·25 0·30						1251	329 1377	152 356	89 163	58 93	41 61	31 42	24 31	19 24	15 19	12 15	10 12	8 10	7 8	6
0.35							13	1471	376	170	96	62 97	43	31	24 31	18	14	11	9	7
0·40 0·45									1534	388 1565	173 392	173	62 96	42 61	41	23 29	17 22	14 16	11 12	10

▶ Here again we can see again n = 58 (as in a previous slide).

<sup>3</sup> 

<sup>37/</sup> $\pm$  Source: Campbell, Julious & Altman (1995). Estimating sample sizes for binary, ordered categorical, and continuous outcomes two group comparisons. BMJ, 311(7013), 1145-1148.

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## Confounding

"A simple definition of confounding is the confusion of effects. This definition implies that the effect of the exposure is mixed with the effect of another variable, leading to a bias." 18

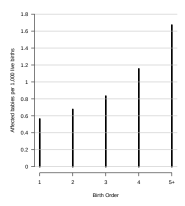
Failing to take a confounding variable into account can lead to a **false conclusion** that the outcome are in a **causal relationship** with the predictor variable.

Confounding variables are typically encountered in observational studies, but not in "ideal" randomized experiments.



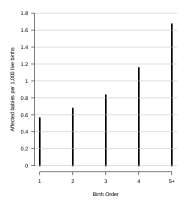
<sup>&</sup>lt;sup>39/60</sup>18 Rothman (2012), Epidemiology: an introduction.

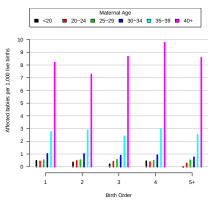
## Confounding example (birth order and risk of Down syndrome 19)





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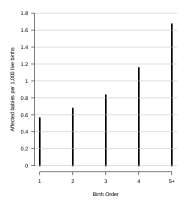


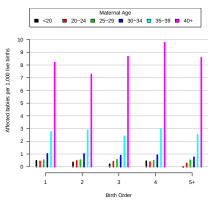




<sup>&</sup>lt;sup>40/40</sup>19Stark and Mantel (1966), J. Natl. Cancer Inst. 37(5) 687–698.

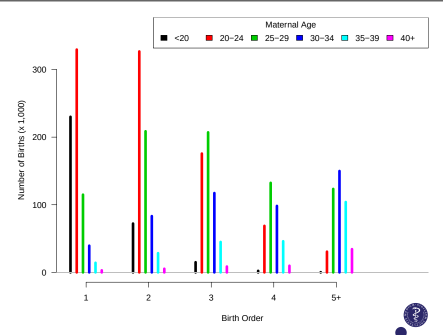
## Confounding example (birth order and risk of Down syndrome 19)







<sup>&</sup>lt;sup>40/40</sup>19Stark and Mantel (1966), J. Natl. Cancer Inst. 37(5) 687–698.



# When can association mean causation? (1/2)

We usually say that (statistical) association does not imply causation.

Confounding (among others things) can lead to non-causal associations.

There are however some exceptions, i.e., it exists some situations in which an association can be interpreted as a causal association. An example is the situation in which we analyze data from an "ideal" randomized experiments.



# When can association mean causation? (2/2)

This is because the randomization ensures that the two groups that we compare are similar with respect to everything except the intervention / treatment under study. Hence, if a difference in outcome is observed between the two groups, then we can be confident that this is the consequence of this unique difference in exposure / treatment.

In non-randomized (or non "ideally" randomized) experiments the two compared groups will usually differ with respect to more than one characteristic. This generates multiple plausible explanations for the observation of the difference in outcome – some causal and some non causal.



## Adjusted analysis

Suppose that in addition to the outcome and the exposure group a categorical confounder variable (e.g. gender) is measured for each individual.

### Subgroup analysis

Analyze 2x2 contingency tables separately in each strata defined by the confounder variable.

### ► Logistic regression (see Lecture 6)

To compute a "weighted" average of the subgroup analyses, assuming that the exposure-outcome association is the same in all subgroups. <sup>20</sup>.



<sup>&</sup>lt;sup>44/60</sup>20 Applicable also with continuous confounders.

# Outline/Intended Learning Outcomes (ILOs)

#### Preliminaries

ILO: calculate 95% Cls for population proportions

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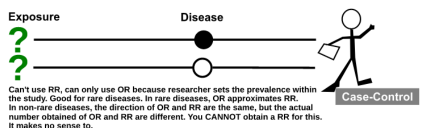
## Observational study design

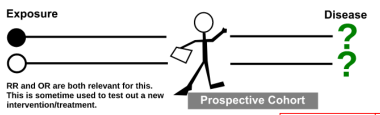
In a prospective **cohort study**, an outcome or disease-free study population is first identified by an exposure (e.g., onset of diabetes) or other inclusion criteria and followed in time until the disease or outcome of interest occurs.

**Case-control** studies identify subjects by outcome status at the outset of the investigation. First, subjects with outcome are identified and classified as **cases**. For each case a given number of controls (e.g., 4) are selected. A candidate **control** is a subject without the outcome but from the same source population.



## **Observational Study Designs: Case Control vs Cohort**





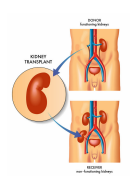


# Cohort study: example from Egerup et al. (2020) <sup>21</sup>

Research question: How larger is the 1-year risk of infection (leading to an hospitalization) among newborns of kidney-transplanted women?

Infection within first year of life

		yes	no	total
Kidney- transplanted mother	yes	26	98	124
	no	133	1098	1231
	total	159	1196	1355



The estimated risk ratio is  $\widehat{RR} = 1.94$  (Cl<sub>95%</sub> = [1.33; 2.83]).

<sup>&</sup>lt;sup>20</sup> Egerup et al. "Increased risk of neonatal complications and infections in children of kidney-transplanted women: A nationwide controlled cohort study." American Journal of Transplantation (2020).

DEPARTMENT OF BIOSTATISTIC:

## Case-control study: example of Frachon et al.<sup>22</sup>

Research question: Is the use of benfluorex associated with unexplained mitral regurgitation?





- ► Case study described in the movie "150 Milligrams" (2016) (The original title in French is "La fille de Brest")
- ► France's biggest modern health scandal



# Case-control study: example of Frachon et al.<sup>23</sup>

"unexplained"

mitral regurgitation

Benfluorex use

	yes	no	total
yes	19	3	24
no	8	51	59
total	27	54	81



Mitral Valve Regurgitation

$$\widehat{OR} = 40.4 \ (CI_{95\%} : [9.7; 168])$$

The number of controls (here 2 per case) is defined by the study design. Hence we cannot estimate risks as one minus the proportions of controls among exposed and non-exposed...

- ▶ The statistic  $\widehat{RR}$  depends also on the ratio between controls and cases and should **not** be used for measuring association in case-control studies.
- ▶ The statistic  $\widehat{OR}$  works.

<sup>50/6023</sup> 

# Why does $\widehat{OR}$ work? (1/2)

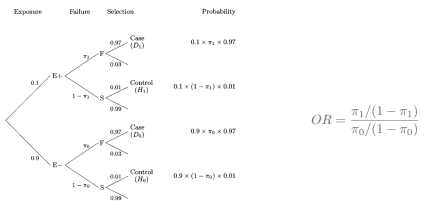


Fig. 16.1. The probability model in the study base.

- 97% of the cases are included in the case-control study and 1% of the "non cases" are selected as controls; all included "blinded" from exposure (i.e. before looking for the information on the exposure).
- Connection to notations of previous slides  $\pi_1 = p_1$  and  $\pi_0 = p_2$ .
- ► E="exposure", F="Fail", S="Survive", D="Disease", H="Healthy".
- source: "Statistical models in Epidemiology", by Clayton and Hills, page 155.



# Why does $\widehat{OR}$ work? (2/2)

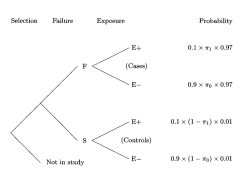


Fig. 16.2. The probability tree for the retrospective argument.

$$\widehat{OR} \approx \frac{\frac{0.1 \times \pi_1 \times 0.97}{0.1 \times (1 - \pi_1) \times 0.01}}{\frac{0.9 \times \pi_0 \times 0.97}{0.9 \times (1 - \pi_0) \times 0.01}}$$

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

source: "Statistical models in Epidemiology", by Clayton and Hills, page 156.



# Why does $\widehat{OR}$ work? (2/2)

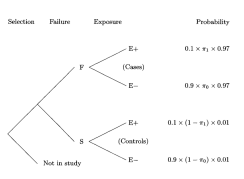


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$$= \frac{\pi_1 / (1 - \pi_1)}{\pi_0 / (1 - \pi_0)}$$

#### but

$$\widehat{RR} \approx \frac{\frac{0.1 \times \pi_1 \times 0.97}{0.1 \times \pi_1 \times 0.97 + 0.1 \times (1 - \pi_1) \times 0.01}}{\frac{0.9 \times \pi_0 \times 0.97}{0.9 \times \pi_0 \times 0.97 + 0.9 \times (1 - \pi_0) \times 0.01}}$$

$$= \frac{\pi_1 / (\pi_1 \times 0.97 + (1 - \pi_1) \times 0.01)}{\pi_0 / (\pi_0 \times 0.97 + (1 - \pi_0) \times 0.01)}$$

$$\neq \frac{\pi_1}{\pi_0}$$



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# Medical test / screening: jargon

- Y: Outcome (disease status) E.g. prostate cancer
- $X \colon \text{Test result (biomarker)}. \ \text{E.g.} \ X = \left\{ \begin{array}{ll} 1 & \text{positive if PSA} > 4.0\,\text{ng/mL} \\ 0 & \text{negative if PSA} \leq 4.0\,\text{ng/mL} \end{array} \right.$

$$egin{array}{cccc} Y=1 & Y=0 \\ X=1 & {\sf True\ positive} & {\sf False\ positive} \\ X=0 & {\sf False\ negative} & {\sf True\ negative} \\ \end{array}$$

- ▶ True positive rate (sensitivity):  $P(X = 1 \mid Y = 1)$
- ▶ True negative rate (specificity):  $P(X = 0 \mid Y = 0)$
- ▶ False positive rate (1-specificity):  $P(X = 1 \mid Y = 0)$
- ▶ The positive predictive value:  $P(Y = 1 \mid X = 1)$
- ▶ The negative predictive value:  $P(Y = 0 \mid X = 0)$



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## When do we typically meet paired binary data?

### Comparison of diagnostic tests

Example: compare sensitivity (i.e. True Positive Rate) of two diagnostic tests based on either Method 1 (e.g. Blood culture) or Method 2 (e.g. PCR: Polymerase Chain Reaction) using the the same blood samples (i.e. same patients).

#### Crossover clinical trials

Example: compare two sedatives, w.r.t. proportions of side effects (e.g. not waking when fire alarm rings), each drug is given to each patient one evening (two evenings separated by one week). The same patients receive the two drugs.



## Why does pairing matter?

### Comparison of diagnostic tests

Example (cont'): blood samples of "heavily" infected patients are easier to test positive than those of "mildly" infected patients. Hence, if one test is positive, the chance that the second test is positive is higher than expected in average.

#### Crossover clinical trials

Example (cont'): some people sleep better than others. Some will never wake no matter what. Others are bad sleepers and will always wake. Hence, if a subject wakes the first night, the chance that he/she wakes up the second night is higher than expected in average.

**Take home message:** we expect less variability between two observations from the same patient than between two observations from two different patients. Appropriate statistical analysis will recognize this smaller variability. Less variability implies less random variation, which further implies more certainty, that is, narrower 95% CI and smaller p-values (than if the pairing was "wrongly" ignored).

## How are paired data often presented?

► Comparison of diagnostic tests<sup>24</sup>

Example (cont'):

		PCR-	-test
		Negative	Positive
BC-test	Negative	1	19
DC-test	Positive	2	2

DCD toot

#### **Remarks:**

- 1. This 2 by 2 table shows the pairing (and the raw data).
- If the sensitivity of the two diagnostic tests are equally good, we expect (approx.) the same counts in the "upper right" and "lower left" cells

## Which statistical method with paired binary data?

- ► For p-value computation, we often use a McNemar's test
- Modern software can compute an "exact" version of the McNemar's test.
- ► An exact confidence interval can be computed for each of the two compared specificities (as seen in the first slides of the lecture)<sup>25</sup>



## Which R code and conclusions?

```
library(exact2x2)  # load a useful package
tab <- rbind(c(1,19),c(2,2))  # 2 by 2 table
mcnemar.exact(tab)  # exact McNemar test
binom.test(x=sum(tab[,2]),n=sum(tab))  # sensitivity for PCR-test (95%-CI)
binom.test(x=sum(tab[2,]),n=sum(tab))  # sensitivity for BC-test (95%-CI)</pre>
```

#### **Conclusions:**

The sensitivity of the PCR test (88%, 95%-CI=[68,97]) was found significantly higher than that of the blood culture test (17%, 95%-CI=[5,37]) among patients with deep-seated candidiasis (p-value<0.001).

