

STK4900/9900 - Lecture 6

Program

1. Binary data and proportions
 2. Comparing two proportions
 3. Contingency tables
 4. Excess risk, relative risk, and odds ratio
 5. Logistic regression with one predictor
- Section 3.4
 - Section 6.1
 - Supplementary material on proportions and contingency tables (cf. your introductory statistics textbook)

Binary data and proportions

In the first part of the course, we considered the situation where the outcome was numerical

We will now consider the situation where the outcome is a binary variable (coded as 0 or 1)

Example: Opinion polls

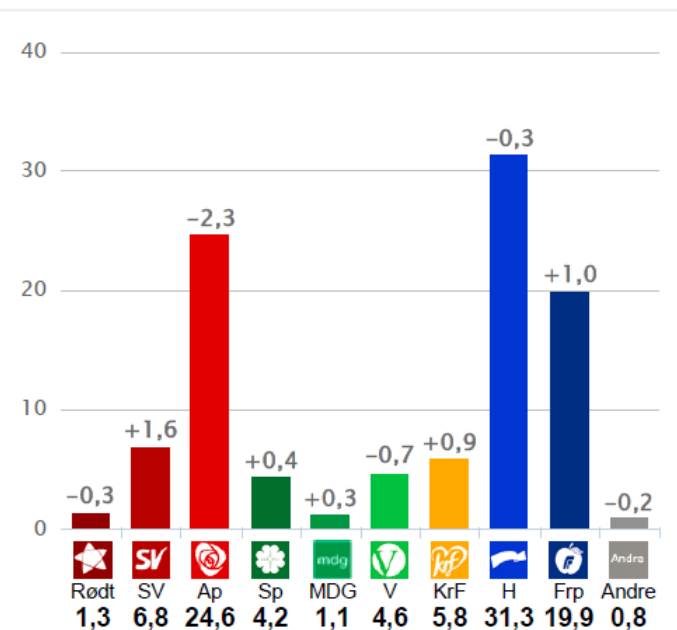
In April 2013 Norstat asked $n = 968$ individuals which party they would support if there had been election to the parliament tomorrow

303 would have voted Høyre

Høyre's support on the opinion poll is

$$\frac{303}{968} = 0.313 = 31.3\%$$

Siste måling - Norstat/Vårt Land, 2. apr 2013



In general we have a sample of binary data y_1, y_2, \dots, y_n from a population

Here $y_i = 1$ if subject i has a certain property (e.g. vote Høyre), while $y_i = 0$ otherwise

We let $p = P(y_i = 1)$

Then p is the proportion in the population who has the property

We may estimate p by the sample proportion:

$$\hat{p} = \frac{\sum_{i=1}^n y_i}{n} = \frac{\#(y_i = 1)}{n}$$

Standard error: $se(\hat{p}) = \sqrt{\frac{\hat{p}(1 - \hat{p})}{n}}$

In the example, the standard error becomes

$$se(\hat{p}) = \sqrt{\frac{0.313(1 - 0.313)}{968}} = 0.0149$$

One may show that \hat{p} is approximately normally distributed (cf. the central limit theorem)

95% confidence interval for the population proportion p :

$$\hat{p} \pm 1.96 \cdot se(\hat{p})$$

In the example a 95% confidence interval becomes:

$$0.313 \pm 1.96 \cdot 0.0149$$

i.e.

$$0.313 \pm 0.029$$

Thus our estimate of Høyre's support is 31.3% with a "margin of error" of $\pm 2.9\%$

Comparing two proportions

Assume that we have a random sample of binary data from each of two populations, and that the two samples are independent

Example: "Divorce" among seagulls

Kittiwake (krykkje) is a seagull whose mating behavior is basically monogamous, but some couples do not reunite the next breeding season ("divorce")

Does the "divorce rate" depend on whether breeding was successful or not?

769 kittiwake pair-bonds were studied over two breeding seasons

Of the 160 couples that had not successful breeding the first season, 100 divorced

Of the 609 couples that were successful, 175 divorced



Population 1:

Population proportion: p_1

Sample size: n_1

Sample proportion: \hat{p}_1

Population 2:

Population proportion: p_2

Sample size: n_2

Sample proportion: \hat{p}_2

We are interested in estimating $p_1 - p_2$ and testing $H_0 : p_1 = p_2$

95% confidence interval for $p_1 - p_2$:

$$\hat{p}_1 - \hat{p}_2 \pm 1.96 \cdot se(\hat{p}_1 - \hat{p}_2)$$

where

$$se(\hat{p}_1 - \hat{p}_2) = \sqrt{se(\hat{p}_1)^2 + se(\hat{p}_2)^2} = \sqrt{\frac{\hat{p}_1(1 - \hat{p}_1)}{n_1} + \frac{\hat{p}_2(1 - \hat{p}_2)}{n_2}}$$

In the example:

Unsuccessful (population 1):

Sample size: $n_1 = 160$

Sample proportion: $\hat{p}_1 = \frac{100}{160} = 0.625$

Successful (population 2):

Sample size: $n_2 = 609$

Sample proportion: $\hat{p}_2 = \frac{175}{609} = 0.287$

We obtain:

$$\hat{p}_1 - \hat{p}_2 = 0.625 - 0.287 = 0.338$$

$$se(\hat{p}_1 - \hat{p}_2) = 0.0424$$

95% confidence interval:

$$0.338 \pm 1.96 \cdot 0.0424 \quad \text{i.e.} \quad 0.338 \pm 0.083$$

We then consider testing the null hypothesis $H_0 : p_1 = p_2$ versus the (two-sided) alternative $H_A : p_1 \neq p_2$

Test statistic:

$$z = \frac{\hat{p}_1 - \hat{p}_2}{se_0(\hat{p}_1 - \hat{p}_2)}$$

Here $se_0(\hat{p}_1 - \hat{p}_2)$ is the estimated standard error under the null hypothesis, obtained by using the sample proportion \hat{p} in the two samples combined:

$$se_0(\hat{p}_1 - \hat{p}_2) = \sqrt{\frac{\hat{p}(1 - \hat{p})}{n_1} + \frac{\hat{p}(1 - \hat{p})}{n_2}}$$

We reject H_0 for large values of $|z|$

Under H_0 the test statistic is approximately standard normal

P-value (two-sided): $P = 2 P(Z > |z|)$ where Z is standard normal

In the example:

Unsuccessful (population 1):

$$n_1 = 160 \quad \hat{p}_1 = \frac{100}{160} = 0.625$$

Successful (population 2):

$$n_2 = 609 \quad \hat{p}_2 = \frac{175}{609} = 0.287$$

We obtain:

$$\hat{p} = \frac{100 + 175}{160 + 609} = 0.358$$

$$se_0(\hat{p}_1 - \hat{p}_2) = 0.0426$$

The test statistic takes the value

$$z = \frac{0.625 - 0.287}{0.0426} = 7.9$$

which is highly significant

2x2 tables

It is common to summarize the situation with two binary samples in a 2x2 table. For the example we have the 2x2 table:

	divorced	not divorced	Total
Unsuccessful	100	60	160
Successful	175	434	609
Total	275	494	769

An alternative way of formulating the test for the null hypothesis of no difference between the populations (cf. slide 8), is to compare the observed numbers in the table (denoted O's) with the corresponding expected numbers if the null hypothesis is true (denoted E's)

If there is no difference between the two groups we would (e.g.) expect

$$160 \cdot \frac{275}{769} = 57.2$$

divorces among the unsuccessful couples

Expected numbers:

	divorced	not divorced	Total
Unsuccessful	57.2	102.8	160
Successful	217.8	391.2	609
Total	275	494	769

Test statistic:
$$\chi^2 = \sum_{\text{all cells}} \frac{(O - E)^2}{E}$$

We reject H_0 for large values of χ^2

Under H_0 the test statistic is approximately chi-squared distributed with 1 degree of freedom (df) provided that all E's are at least 5

P-value: $P(\chi^2 \geq \chi_{\text{obs}}^2)$ where χ^2 is chi-squared distributed with 1 df

One may show that $\chi^2 = z^2$ so this is a reformulation of the test on slide 8

R commands:

```
kittiwake=matrix(c(100,175,60,434),nrow=2)
dimnames(kittiwake)=list(c("unsuccessful","successful"),c("divorced","not_divorced"))
kittiwake
chisq.test(kittiwake,correct=F)$expected
prop.test(kittiwake,correct=F)
```

R output (edited):

	divorced	not_divorced
unsuccessful	100	60
successful	175	434

	divorced	not_divorced
unsuccessful	57.217	102.783
successful	217.783	391.217

X-squared = 62.8813, df = 1, p-value = 2.196e-15

alternative hypothesis: two.sided

95 percent confidence interval:

0.25446 0.42082

sample estimates:

prop 1	prop 2
0.62500	0.28736

Contingency tables

The chi-squared test may be extended to contingency tables of higher order

Example: Blood pressure

Blood pressure of 92 teenagers according to the blood pressure of their fathers:

		<u>Child's blood pressure</u>			Total
		Lower third	Middle third	Upper third	
<u>Father's blood pressure</u>	Lower third	14	11	8	33
	Middle third	11	11	9	31
	Upper third	6	10	12	28
	Total	31	32	29	92

Does the blood pressure of the children depend on the blood pressure of their fathers?

We will test the null hypothesis that there is no difference between the groups (in the example, that the blood pressure of the children does not depend on the blood pressure of their fathers)

Expected numbers (E's) are computed as for 2x2 tables

Test statistic:
$$\chi^2 = \sum_{\text{all cells}} \frac{(O - E)^2}{E}$$

We reject H_0 for large values of χ^2

Under H_0 the test statistic is approximately chi-squared distributed with $df = (\#rows - 1) \cdot (\#columns - 1)$ provided that all E's are at least 5

In the example we have $df = (3 - 1) \cdot (3 - 1) = 4$

R commands:

```
bloodpr=matrix(c(14,11,6,11,11,10,8,9,12),nrow=3)
dimnames(bloodpr)=list(c("F.low","F.middle","F.upper"), c("C.low","C.middle","C.upper"))
```

`bloodpr`

```
chisq.test(bloodpr,correct=F)$expected
```

```
chisq.test(bloodpr,correct=F)
```

R output (edited):

	C.low	C.middle	C.upper
F.low	14	11	8
F.middle	11	11	9
F.upper	6	10	12

	C.low	C.middle	C.upper
F.low	11.120	11.478	10.402
F.middle	10.446	10.783	9.772
F.upper	9.435	9.739	8.826

Pearson's Chi-squared test

X-squared = 3.814, df = 4, p-value = 0.432

Risk measures

Assume that we have a random sample of binary data from each of two populations, and that the two samples are independent

We assume that population 1 corresponds to an "exposed" population (specified by $x = 1$) and that population 2 corresponds to an "unexposed" population (specified by $x = 0$)

Example: "Divorce" among seagulls

	divorced ($y=1$)	not divorced ($y=0$)	Total
Unsuccessful ($x=1$)	100	60	160
Successful ($x=0$)	175	434	609
Total	275	494	769

Population 1: $p_1 = p(1) = P(y = 1 | x = 1)$

Population 2: $p_2 = p(0) = P(y = 1 | x = 0)$

On slide 6 we used the *excess risk*

$$ER = p(1) - p(0)$$

to measure the effect of the "exposure"

An alternative would be to use the *relative risk* given by:

$$RR = \frac{p(1)}{p(0)}$$

In the example, estimates of these two measures of risk are given by (cf. slide 7)

$$ER = \hat{p}(1) - \hat{p}(0) = 0.625 - 0.287 = 0.338$$

$$RR = \frac{\hat{p}(1)}{\hat{p}(0)} = \frac{0.625}{0.287} = 2.18$$

A third risk measure is based on the concept of *odds*, so we will first discuss this concept

Assume that an event has a probability p of occurring

Then the odds for the event is

$$\text{odds} = \frac{p}{1-p}$$

The odds is one if the probability that an event will happen is equal to the probability that it will not happen, cf. the expression "a fifty-fifty chance"

When you throw a die, the odds that it will face six is 1 : 5 (i.e. it is five times more likely that it will not face six than it will face six)

We then return to the situation with two populations:

$$\text{Population 1: } p(1) = P(y = 1 | x = 1)$$

$$\text{Population 2: } p(0) = P(y = 1 | x = 0)$$

The odds for the two populations are:

$$\text{Population 1: } \frac{p(1)}{1 - p(1)} \quad \text{Population 2: } \frac{p(0)}{1 - p(0)}$$

Then a third risk measure is the *odds ratio*

$$OR = \frac{p(1)/[1 - p(1)]}{p(0)/[1 - p(0)]}$$

In the example, an estimate for the odds ratio becomes (cf. slide 7)

$$OR = \frac{0.625/(1 - 0.625)}{0.287/(1 - 0.287)} = \frac{1.667}{0.403} = 4.14$$

Logistic regression with one predictor

When discussing logistic regression, we will to a large extent use the WCGS study for illustration

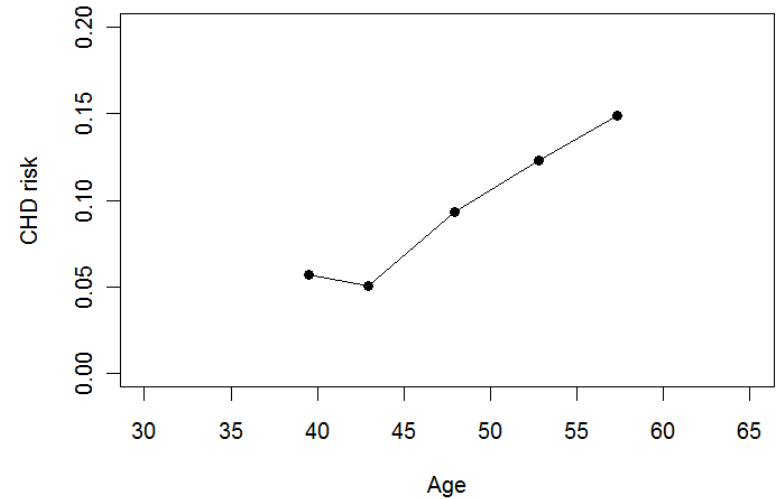
WCGS is a large epidemiological study designed to study risk factors for coronary heart disease (CHD) among middle-aged men

The men were followed for 10 years, and for each man it was recorded if he developed CHD ($y=1$) or not ($y=0$) over the course of the study

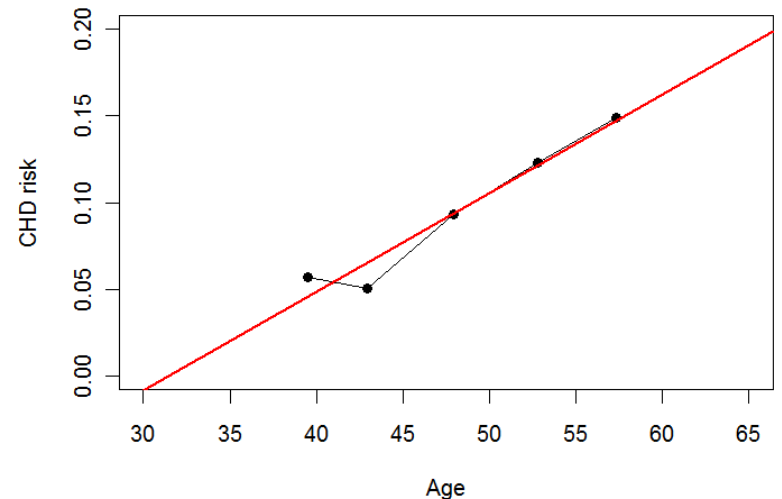
How does the age (at entry to the study) affect the risk (probability) of developing CHD?

Age group (mean)	35-40 (39.5)	41-45 (42.9)	46-50 (47.9)	51-55 (52.8)	56-60 (57.3)
# Total	543	1091	750	528	242
# CHD	31	55	70	65	36
% CHD	5.7 %	5.0 %	9.3 %	12.3 %	14.9 %

The figure shows a the observed proportion with CHD plotted versus the mean age in each age group



A least square fit to the observed proportions give the fitted line



This least squares line may give an all right description of the observed proportions, but there are in general problem with using linear regression for binary data and proportions

In general we have data $(x_1, y_1), \dots, (x_n, y_n)$

Here y_i is a binary outcome (0 or 1) for subject i and x_i is a predictor for the subject (which may be binary or numerical)

In the WCGS study, $y_i = 1$ if man number i developed CHD during the course of the study, $y_i = 0$ if not, and x_i may be his age (at entry to the study)

In general we let

$$p(x) = E(y | x) = P(y = 1 | x)$$

We want a model that specifies a relation between $p(x)$ and x

One option would be a linear model:

$$p(x) = \beta_0 + \beta_1 x$$

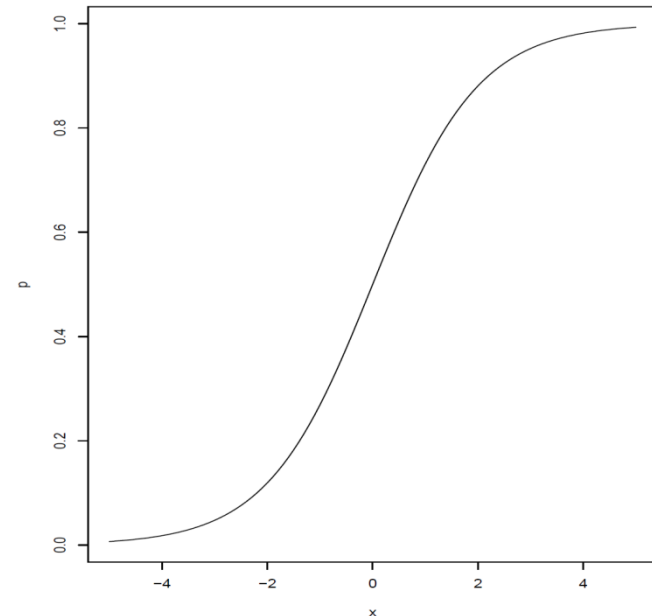
This is an **additive risk model**, which may be useful in some situations

However, it is a main problem with the additive risk model that it may give impossible values for the probabilities (negative or above 1)

To avoid this problem it is common to consider the **logistic regression model** given by

$$p(x) = \frac{\exp(\beta_0 + \beta_1 x)}{1 + \exp(\beta_0 + \beta_1 x)}$$

This gives a "S-shaped" relation between $p(x)$ and x

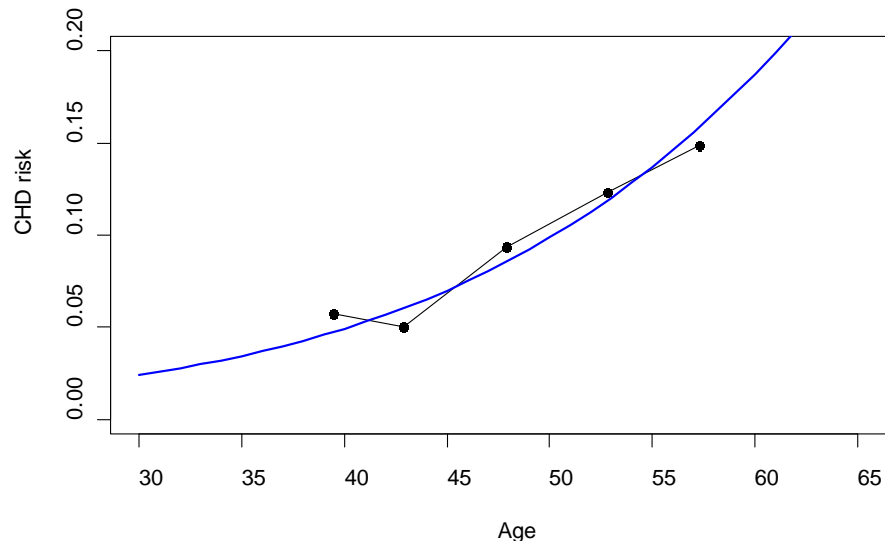


If we fit a logistic regression model for the WCGS data using the mean age in each age group as a numeric covariate, we get

$$\hat{\beta}_0 = -5.947 \quad \text{and} \quad \hat{\beta}_1 = 0.0747$$

This gives the fitted model

$$\hat{p}(\text{age}) = \frac{\exp(-5.947 + 0.0747 \cdot \text{age})}{1 + \exp(-5.947 + 0.0747 \cdot \text{age})}$$



The method for estimating the parameters of a logistic regression model will be described in Lecture 7

The logistic model may alternatively be given in terms of the odds:

$$\frac{p(x)}{1-p(x)} = \exp(\beta_0 + \beta_1 x)$$

If we consider two subjects with covariate values $x + \Delta$ and x , respectively, their odds ratio becomes

$$\frac{p(x + \Delta)/[1 - p(x + \Delta)]}{p(x)/[1 - p(x)]} = \frac{\exp(\beta_0 + \beta_1 (x + \Delta))}{\exp(\beta_0 + \beta_1 x)} = \exp(\beta_1 \Delta)$$

In particular e^{β_1} is the odds ratio corresponding to one unit's increase in the value of the covariate

In the WCGS study the odds ratio for one year increase in age is $e^{0.0747} = 1.078$ while the odds ratio for a ten-year increase is $e^{0.0747 \cdot 10} = 2.11$

(The numbers deviate slightly from those on pp 162-163 in the text book, since we have used mean age for each age group in this illustration; cf. the exercises for the results when actual age is used.)

R commands for logistic regression

Binary CHD data with mean age in each age group as covariate

R commands:

```
wcgs=read.table("http://www.uio.no/studier/emner/matnat/math/STK4900/v13/wcgs.txt",
               sep="\t",header=T,na.strings=".")
wcgs$agem=39.5*(wcgs$agec==0)+42.9*(wcgs$agec==1)+47.9*(wcgs$agec==2)+
          52.8*(wcgs$agec==3)+57.3*(wcgs$agec==4)
attach(wcgs)
cbind(chd69, agem)
```

R output of binary CHD data (edited):

chd69	agem	chd69	agem	chd69	agem
[1,]	0 47.9	[10,]	0 42.9	[3145,]	0 42.9
[2,]	0 52.8	[11,]	0 57.3	[3146,]	0 42.9
[3,]	0 57.3	[12,]	0 52.8	[3147,]	0 52.8
[4,]	0 52.8	[13,]	0 47.9	[3148,]	0 42.9
[5,]	0 42.9	[14,]	1 39.5	[3149,]	0 42.9
[6,]	0 47.9	[15,]	0 47.9	[3150,]	0 47.9
[7,]	0 39.5	[16,]	0 52.8	[3151,]	0 42.9
[8,]	0 42.9	[17,]	0 42.9	[3152,]	0 52.8
[9,]	0 47.9	[18,]	0 57.3	[3153,]	0 52.8
		[19,]	0 42.9	[3154,]	0 47.9

When we use the mean age in each age group as covariate, all information is summarized in the table

Age group (mean)	35-40 (39.5)	41-45 (42.9)	46-50 (47.9)	51-55 (52.8)	56-60 (57.3)
# Total	543	1091	750	528	242
# CHD	31	55	70	65	36

As an alternative to using the individual binary data, we may therefore use the grouped data given in the table

R commands:

```
chd.grouped=read.table("http://www.uio.no/studier/emner/matnat/math/STK4900/v13/chd_grouped.txt ",  
                        header=T)
```

```
chd.grouped
```

R output of grouped CHD data:

```
no  chd  agem  
543  31  39.5  
1091  55  42.9  
750  70  47.9  
528  65  52.8  
242  36  57.3
```

We may fit the logistic regression model using the individual binary data or by using the grouped data

R commands for binary data:

```
fit.binary=glm(chd69~agem, data=wcgs,family=binomial)
summary(fit.binary)
```

R commands for grouped data:

```
fit.grouped=glm(cbind(chd,no-chd)~agem, data=chd.grouped, family=binomial)
summary(fit.grouped)
```

The two ways of fitting the logistic regression model give the same estimates and standard errors:

R output (edited):

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-5.9466	0.5616	-10.588	< 2e-16
agem	0.0747	0.0116	6.445	1.15e-10

(Other parts of the R output will differ, as we will discuss in Lecture 7)