Epidemiologic Data Analysis using R Part 3: Basic analysis of rates

Janne Pitkäniemi (Esa Läärä)

Finnish Cancer Registry, Finland, <janne.pitkaniemi@cancer.fi>
(University of Oulu, Finland, <esa.laara@oulu.fi>)

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- 1. Person-time data, hazard and incidence rates,
- 2. Comparative parameters of rates and their estimation,
- 3. Poisson regression models and comparative parameters,
- 4. Adjustment for confounding and evaluation of modification by Poisson regression,
- Goodness-of-fit evaluation.

Main R functions covered:

- ▶ glm()
- ▶ tools for extracting results from a glm model object

Person-time data and incidence rates

Summarized data on outcome from cohort study, in which two exposure groups, as to binary risk factor X, have been followed-up over individually variable times.

Exposure to	Number of	Person-
risk factor	cases	time
yes	D_1	Y_1
no	D_0	Y_0
total	D_+	Y_{+}

Empirical **incidence rates** by exposure group:

$$I_1 = D_1/Y_1, \qquad I_0 = D_0/Y_0.$$

These provide estimates for the true **hazards** (or hazard rates) λ_1 and λ_0 assumed constant within exposure categories.

Hazards and their comparison

Parameters of interest:

(i) hazard ratio

$$\rho = \frac{\lambda_1}{\lambda_0} = \frac{\text{hazard among exposed}}{\text{hazard among unexposed}}$$

(ii) hazard difference

$$\delta = \lambda_1 - \lambda_0$$

Null hypothesis $H_0: \rho = 1 \Leftrightarrow \delta = 0$ \Leftrightarrow exposure has no effect.

Estimation of hazard ratio

Point estimator of true hazard ratio ρ : empirical incidence rate ratio (IR)

$$\widehat{\rho} = IR = \frac{I_1}{I_0} = \frac{D_1/Y_1}{D_0/Y_0} = \frac{D_1/D_0}{Y_1/Y_0}.$$

NB. The last form is particularly useful:

= exposure odds ratio (EOR).

Standard error of log(IR), 95% error factor & 95% CI for ρ :

SEL =
$$\sqrt{\frac{1}{D_1} + \frac{1}{D_0}}$$

EF = $\exp\{1.96 \times \text{SEL}\}$
CI = $[IR/EF, IR \times EF]$.

NB. Random error depends inversely on numbers of cases.

Estimation of hazard difference

Point estimator of true hazard difference δ : empirical incidence rate difference (ID)

$$\hat{\delta} = ID = I_1 - I_0 = \frac{D_1}{Y_1} - \frac{D_0}{Y_0}$$

Standard error of ID, 95% error margin & 95% CI for δ :

$$SE = \sqrt{\frac{l_1^2}{D_1} + \frac{l_0^2}{D_0}}$$

$$\rm EM = 1.96 \times SE$$

$$CI = [ID - EM, ID + EM].$$

NB. Random error again depends inversely on no. of cases.

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Example. British doctors' study (Doll & Hill '66)

CHD mortality in males by smoking and age. Cases (D), person-years (Y), and mortality rates $(I \text{ per } 10^4 \text{ y})$.

		Smokers		N	Non-smokers			
Age(y)	D	D Y I			Y	1		
35-44	32	52407	6	2	18790	1		
45-54	104	43248	24	12	10673	11		
55-64	206	28612	72	28	5710	49		
65-74	186	12663	147	28	2585	108		
75-84	102	5317	192	31	1462	212		
Total	630	142247	44	101	39220	26		

Example (cont'd).

Crude incidence rates:

$$I_1 = 630/142247 \text{ y} = 44.3 \text{ per } 10^4 \text{ y, and}$$

 $I_0 = 101/39220 \text{ y} = 25.8 \text{ per } 10^4 \text{ y.}$

Crude estimate of overall hazard ratio ρ with SE, etc.

$$\widehat{\rho} = IR = \frac{44.3}{25.8} = 1.72$$

SEL =
$$\sqrt{\frac{1}{630} + \frac{1}{101}} = 0.1072$$

$$EF = exp(1.96 \times 0.1072) = 1.23$$

95% CI for ρ : [1.72/1.23, 1.72 × 1.23] = [**1.39**, **2.12**] Two-tailed P < 0.001.

Poisson regression model for rate ratio

▶ Random part: Number of cases in exposure group j = 0, 1

$$D_j \sim \mathsf{Poisson}(\lambda_j Y_j),$$

where $\mu_j = \lambda_j Y_j = expected number$ of cases.

Systematic part & link function: linear predictor $\alpha + \beta X_j$ with logarithmic (log) link

$$\log(\lambda_i) = \alpha + \beta X_i,$$

equivalently on the original hazard scale:

$$\lambda_i = \exp(\alpha + \beta X_i).$$

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Poisson model for rate ratio (cont'd)

Interpretation,

$$X_j = egin{cases} 1 & ext{if exposed } (j=1), \ 0 & ext{if unexposed } (j=0), \ lpha = \log(\lambda_0), ext{ log-baseline rate,} \ eta = \log(
ho) = \log(\lambda_1/\lambda_0), ext{ logarithm of true hazard ratio,} \ e^eta =
ho = ext{true hazard ratio.} \end{cases}$$

Special case of generalized linear models!

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Example. Crude analysis of CHD mortality in R

A ready data frame contains

four variables:

age = age group - a factor with 5 levels,
$$smok = smoking$$
: 1 = "yes", 0 = "no", $d = number of cases$, $y = person-years$.

▶ 10 observations (one for each age-smoking combination).

Example. Analysis of CHD rates (cont'd)

```
> data.frame(age, smok, d, y, rate)
     age smok
               d
                     y rate
              32 52407
 1 35-44
                         6.1
 2 35-44
               2 18790
                         1.1
 3 45-54
           1 104 43248
 4 45-54
           0 12 10673 11.2
 5 55-64
           1 206 28612 72.0
6 55-64
           0 28 5710 49.0
           1 186 12663 146.9
 7 65-74
 8 65-74
           0 28 2585 108.3
9 75-84
           1 102 5317 191.8
10 75-84
           0 31 1462 212.0
```

Fitting Poisson model for crude rate ratio

Poisson model with log-link (default) for crude rates

```
> m1 <- glm(d/y ~ smok, family=poisson(), weights=y)</pre>
```

Output (shortened) using summary() method

> summary(m1)
Coefficients:

Null deviance: 935.07 on 9 degrees of freedom Residual deviance: 905.98 on 8 degrees of freedom

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Fitting crude rate ratio model (cont'd)

Main results:

$$\widehat{\alpha} = -5.96 = \log(25.8/10^4 \text{ y}), \quad (SE = 0.10),$$

 $\widehat{\beta} = 0.54 = \log(1.72), \quad (SE = 0.11)$

Function ci.lin() transforms results to ratio scale

Compare the results with those obtained above using simple estimation & SE formulas.

Stratified analysis

Stratification of cohort data with person-time

– at each level k of covariate Z results are summarized:

	Exposure to risk factor	Number of cases	Person- time
٠	yes	D_{1k}	Y_{1k}
	no	D_{0k}	Y_{0k}
	Total	D_{+k}	Y_{+k}

Stratum-specific rates by exposure group:

$$I_{1k} = \frac{D_{1k}}{Y_{1k}}, \qquad I_{0k} = \frac{D_{0k}}{Y_{0k}}.$$

Fitting crude rate ratio model (cont'd)

The Poisson model above can also be fitted as follows:

Here offset refers to the logarithm of person-years y in formula for expected numbers of cases $\mu_i = \lambda_i \times Y_i$:

$$\log(\mu_j) = \log(\lambda_j Y_j) = \log(Y_j) + \log(\lambda_j)$$

=
$$\log(Y_i) + \alpha + \beta X_i,$$

 $log(Y_j)$ is an **offset** term in the linear predictor, meaning that it has a fixed value 1 for the regression coefficient.

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Stratum-specific comparisons

Let λ_{jk} be true rate for exposure group j (j = 0, 1) and stratum k (k = 0, ..., K). Let also

$$\rho_k = \frac{\lambda_{1k}}{\lambda_{0k}}, \qquad \delta_k = \lambda_{1k} - \lambda_{0k}$$

be the rate ratios and rate differences between the exposure groups in stratum k.

Two simple models assuming homogeneity:

- common rate ratio: $\rho_k = \rho$ for all k,
- common rate difference: $\delta_k = \delta$ for all k.

Only one of these can in principle hold. However, almost always neither homogeneity assumption is exactly true.

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Example (cont'd).

- ▶ Both types of comparative parameter, rate ratios ρ_k and rate differences δ_k appear heterogenous, because
 - ▶ ID increases by age at least up to 75 y,
 - ▶ IR decreases by age.
- ▶ Part of this observed heterogeneity may be due to random variation.
- ➤ Yet, any single-parameter comparison by common rate ratio or rate difference may not adequately capture the joint pattern of true rates.
- ⇒ Effect modification must be evaluated.

Example. British male doctors (cont'd)

CHD mortality rates (per 10^4 y) and numbers of cases (D) by age and cigarette smoking.

Mortality rate differences (ID) and ratios (IR) in age strata.

	Smokers		Non-s	Non-smokers		
Age (y)	rate	(D)	rate	(D)	ID	IR
35-44	6.1	(32)	1.1	(2)	5	5.7
45-54	24	(104)	11	(12)	13	2.1
55-64	72	(206)	49	(28)	23	1.5
65-74	147	(186)	108	(28)	39	1.4
75-84	192	(102)	212	(31)	-20	0.9
Crude	44	(630)	26	(101)	18	1.7

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Rate ratio adjustment by Poisson model

Define Poisson regression model for

- one binary exposure variable X and
- one categorical (polytomous) factor Z.

Random part: No. of cases in exposure group j (j=0,1) and covariate level k ($k=1,\ldots,K$) is $D_{jk}\sim \mathsf{Poisson}(\lambda_{jk}Y_{jk})$

Systematic part: $\log(\lambda_{ik}) = \alpha + \beta X_i + \gamma_k$, where X_i is (0/1),

$$\alpha = \log(\lambda_{01}) = \text{log-baseline rate,}$$

$$\gamma_k = \log(\lambda_{jk}/\lambda_{j1}),$$

$$\beta = \log(\rho) = \log(\lambda_{1k}/\lambda_{0k}),$$

 $e^{\beta} = \rho = \text{true rate ratio for the effect of exposure to } X.$

How do we read this?

Implications of model definition

- ▶ homogeneity of true rate ratio $\rho_k = \rho$ for X across levels of Z is assumed,
- ▶ inclusion of Z leads to adjustment for Z in estimating the common effect of X,
- e^{γ_k} = rate ratio for level k of Z vs. level 1 is the same in both exposure groups (j=0,1) \Rightarrow homogeneity of the effect of Z is assumed, too.
- ▶ level k = 1 is chosen as the *reference* level for Z (like "unexposed" is reference for X),
- ▶ before model fitting, binary *indicator variables* Z_k for levels k = 1, ..., K of Z must be defined:

$$Z_k = \begin{cases} 1, & \text{if observation belongs to level } k, \\ 0, & \text{otherwise.} \end{cases}$$

Example. CHD in British doctors (cont'd)

Factor age has 5 levels.

Indicator variables for each age level are generated in R when defining the model, and the following model matrix is returned.

> cbind(data.frame(age), model.matrix(m2))

	age	(I't)	age45-54	age55-64	age65-74	age75-84	smok
	1 35-44	1	0	0	0	0	1
:	2 35-44	1	0	0	0	0	0
;	3 45-54	1	1	0	0	0	1
	4 45-54	1	1	0	0	0	0
	5 55-64	1	0	1	0	0	1
(55-64	1	0	1	0	0	0
•	7 65-74	1	0	0	1	0	1
;	8 65-74	1	0	0	1	0	0
!	9 75-84	1	0	0	0	1	1
1	75-84	1	0	0	0	1	0

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Fitting the adjustment model

Define a new model object m2 with age and smok

```
> m2 <- glm( d/y ~ age + smok,
+ family=poisson(link=log), weights=y)}
> summary(m2)
```

Coefficients:

	Estimate	Std. Error	z value	Pr(> z)
(Intercept)	-7.9193	0.1917	-41.301	< 2e-16
age 45-54	1.4840	0.1951	7.607	2.81e-14
age 55-64	2.6275	0.1837	14.302	< 2e-16
age 65-74	3.3505	0.1848	18.132	< 2e-16
age 75-84	3.7001	0.1922	19.251	< 2e-16
smok	0.3545	0.1074	3.302	0.00096

Residual deviance: 12.132 on 4 degrees of freedom

Fitting adjustment model (cont'd)

Results on the ratio scale

> round(ci.lin	(m2, Exp=T)	[, 5:7]	, 4)
	<pre>exp(Est.)</pre>	2.5%	97.5%
(Intercept)	0.0004	0.0002	0.0005
age 45-54	4.4106	3.0091	6.4648
age 55-64	13.8392	9.6546	19.8375
age 65-74	28.5168	19.8524	40.9627
age 75-84	40.4512	27.7541	58.9571
smok	1.4255	1.1550	1.7594

 \Rightarrow Age-adjusted rate ratio [95% CI] for smoking:

$$\widehat{\rho} = 1.43 [1.16, 1.76]$$

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Fitted values & residuals

From the estimated coefficients we can calculate **fitted** linear predictors $\widehat{\eta}_{jk}$, hazards $\widehat{\lambda}_{jk}$ and numbers $\widehat{\mu}_{jk}$:

$$\widehat{\eta}_{jk} = \widehat{\alpha} + \widehat{\beta}x + \widehat{\gamma}_{k}
\widehat{\lambda}_{jk} = \exp(\widehat{\eta}_{jk}), \quad \widehat{\mu}_{jk} = \widehat{\lambda}_{jk}Y_{jk}$$

In R the two first can be extracted directly from the fitted model object m2:

```
> fit.eta <- m2$linear.predictor
> fit.rate <- fitted(m2); fit.mu <- y*fit.rate</pre>
```

NB. If count d is declared as response with log(y) as offset, then fitted() returns the fitted numbers of cases $\widehat{\mu}_{ik}$.

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Example. Fitted values & residuals

> data.frame(age,smok,rate,fit.rate,d,fit.mu,res.D,res.P)

	age	smok	rate	fit.rate	d	fit.mu	${\tt res.D}$	res.P	
1	35-44	1	6.1	5.2	32	27.3	0.90	0.93	
2	35-44	0	1.1	3.6	2	6.8	-2.18	-1.85	
3	45-54	1	24.0	22.9	104	99.0	0.51	0.51	
4	45-54	0	11.2	16.0	12	17.1	-1.31	-1.24	
5	55-64	1	72.0	71.7	206	205.1	0.05	0.05	
6	55-64	0	49.0	50.3	28	28.7	-0.14	-0.14	
7	65-74	1	146.9	147.8	186	187.2	-0.09	-0.09	
8	65-74	0	108.3	103.7	28	26.8	0.23	0.23	
9	75-84	1	191.8	209.7	102	111.5	-0.91	-0.90	
10	75-84	0	212.0	147.1	31	21.5	1.92	2.05	

NB! Fitted rate ratios between smokers and non-smokers:

$$\frac{5.2}{3.6} = \dots = \frac{209.7}{147.1} = 1.43$$
 at each age level

Fitted values & residuals (cont'd)

Deviance residual for cell *jk* (resid(m2) in R):

$$d_{jk} = \operatorname{sign}(Y_{jk} - \widehat{\mu}_{jk}) \times \sqrt{2\left\{Y_{jk} \log\left(\frac{Y_{jk}}{\widehat{\mu}_{jk}}\right) - (Y_{jk} - \widehat{\mu}_{jk})\right\}}$$

Pearson residual (resid(m2, type="pearson")):

$$r_{jk} = \frac{Y_{jk} - \widehat{\mu}_{jk}}{\sqrt{\widehat{\mu}_{jk}}}.$$

Small value of either residual

 \rightarrow consistency of observation with model.

"Large" (in absolute value) residual

 \rightarrow lack of fit for that cell.

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Summary goodness-of-fit statistics

- ▶ **Deviance** (Dev) = sum of squares of deviance residuals,
- **Pearson's** X^2 (chi-square) = s.s. of Pearson residuals

Assume that

- ▶ the model is "true", and
- ▶ the data are not *sparse*, i.e. numbers of cases in each line of the (grouped) data matrix are sufficiently big.

Under these assumptions

- residuals d_{ik} and r_{ik} all approximately Normal(0,1),
- ▶ Both Dev and X^2 are approximately χ^2 -distributed with degrees of freedom (df) being equal to the residual df = no. of obs no. of coeff's
- ▶ Expected value of these statistics = residual df.

Goodness of fit (cont'd)

In our example

- "large" residuals in extreme age groups,
- \triangleright X^2 and Dev 2.8 to 3 times higher than df.
- \rightarrow Some lack of fit with assumed model.

(*P*-values:
$$P(X^2 > 11.2) = 0.025$$
, and $P(Dev > 12.1) = 0.017$)

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Possible causes for lack of fit

- ► Unrealistic random part (Poisson)?
- Misspecified systematic part?
 - Wrong functional form for effects?
 - Missing important risk factors?
 - Failure to take into account effect modification or "interaction".

In this example the last possibility is very likely: the rate ratios are clearly heterogenous.

Yet, inclusion of 4 interaction parameters would lead to a saturated model.

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Continuous covariates

Stratified analysis is adequate with

- qualitative exposure factor with few levels,
- few qualitative covariates with few levels.

Effects of continous factors can often be described by smooth functions with small number of parameters.

Stratification requires categorization of continuous variables and level specific parameters

 \Rightarrow loss of efficiency.

Continuous covariates (cont'd)

Regression models can accommodate functions of continuous variables

- + more parsimonious models,
- + greater flexibility in modelling,
- more opportunities for misspecification, i.e. creation of "wrong" models.

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Example (cont'd)

Treat age as continuous variable taking midpoint of each ageband (40, 50, ..., 80) as quantitative age value.

Include linear and quadratic term of age applying

- centering at 60 years, and
- ▶ scaling, *i.e.* division by 10 years

This is achieved by

```
> A.L <- as.numeric(age)-3
> A.Q <- A.L*A.L - 2</pre>
```

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Example (cont'd)

Observed and fitted rates & fitted rate ratios between smokers and non-smokers in each ageband

```
age smok rate fit.rate
                             d fit.mu res.D res.P
1 35-44
               6.1
                       5.6 32
                                 29.3 0.44 0.44
  35-44
           0 1.1
                       1.8
                                  3.4 - 0.83 - 0.77
  45-54
           1 24.0
                      24.7 104 106.8 -0.27 -0.27
4 45-54
           0 11.2
                      10.8 12
                                11.5 0.13 0.13
  55-64
           1 72.0
                     72.8 206 208.3 -0.15 -0.15
  55-64
           0 49.0
                      43.3 28
                                24.7 0.64 0.65
  65-74
           1 146.9
                     144.4 186 182.9 0.23 0.23
  65-74
           0 108.3
                     116.9 28
                                30.2 -0.41 -0.41
9 75-84
           1 191.8
                     192.9 102 102.6 -0.06 -0.06
10 75-84
           0 212.0
                     212.5 31
                               31.1 -0.01 -0.01
```

Residual Dev = 1.64 df = 5; The fit is excellent!

Example (cont'd)

Model with linear and quadratic terms for age, and interaction between linear term of age and smoking:

```
> m3 \leftarrow glm(d/y \sim A.L + A.Q + smok + A.L:smok,
            family=poisson(), weights=y)
> round(ci.lin(m3, Exp=T)[, -(3:4)], 4)
           Estimate StdErr exp(Est.)
                                        2.5% 97.5%
(Intercept) -5.8368 0.1213
                               0.0029 0.0023 0.0037
A.L
             1.1904 0.0923
                               3.2885 2.7441 3.9408
A.Q
            -0.1977 0.0274
                               0.8206 0.7778 0.8659
smok
             0.5183 0.1262
                               1.6792 1.3112 2.1505
            -0.3075 0.0970
                               0.7352 0.6079 0.8893
A.L:smok
```

Interpretation of parameters?

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Some closing remarks

- Hazard differences can also be estimated by Poisson-modelling, because link='identity' can be coupled with family=poisson in glm().
- ▶ Poisson models can also be fitted on ungrouped data. Units of observations may even be shorter intervals of total individual follow-up times:
 - ► Each member of the study population can contribute several separate time-slots as observational units.
 - ▶ More about time-splitting in the next lectures.
- ► With continuous covariates it may be difficult to keep hazards positive when the *additive hazards* model with link='identity' is attempted to be fitted.

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