Measures of disease frequency and effects Analysis of epidemiological data

Esa Läärä

University of Oulu, Finland esa.laara@oulu.fi http://stat.oulu.fi/laara

Bendix Carstensen

Steno Diabetes Center, Denmark
& Department of Biostatistics, University of Copenhagen
bxc@steno.dk www.biostat.ku.dk/~bxc

Nordic Summer School in Cancer Epidemiology August 2011, Danish Cancer Society, Copenhagen www.biostat.ku.dk/~bxc/NSCE

Outline

Introduction

Basic concepts

Frequency

Comparison

Age, period, etc.

Standardization

Survival

Conclusion

Introduction

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Key references

IS: dos Santos Silva, I. (1999).
 Cancer Epidemiology: Principles and Methods. International Agency for Research on Cancer (IARC), Lyon.

B&D: Breslow, N.E., Day, N.E. (1987).
Statistical Methods in Cancer Research Volume II
The Design and Analysis of Cohort Studies.
IARC Scientific Publications No. 82, IARC, Lyon.

C&H: Clayton, D., Hills, M. (1993). Statistical Models in Epidemiology. OUP, Oxford.

Internet resources on cancer statistics

NORDCAN : Cancer Incidence and Mortality in the Nordic Countries, Version 4.0. Association of Nordic Cancer Registries, Danish Cancer Society, 2002. http://www-dep.iarc.fr/nordcan.htm NORDCAN is a graphical package providing data on the incidence of, and mortality from 40 major cancers for 80 regions of the Nordic countries (Denmark, Finland, Iceland, Norway and Sweden). Using NORDCAN, these data can be presented as a variety of tables and graphs that can be easily exported or printed. NORDCAN allows countries and cancer sites to be grouped and compared as desired

GLOBOCAN 2008 : Cancer Incidence and Mortality Worldwide in 2008 http://globocan.iarc.fr/

Basic Concepts

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

What is Epidemiology?

Some textbook definitions of epidemiology: Greek: *epi* = upon, *demos* = people

- "study of the distribution and determinants of disease frequency in man" (MacMahon and Pugh, 1970)
- "study of the distribution and determinants of health related states and events in specified populations,..." (Last (ed.) Dictionary of Epidemiology, 2000)
- "discipline on principles of occurrence research in medicine" (Miettinen, 1985)

Different epidemiologies

descriptive epidemiology

— monitoring & surveillance of diseases for planning of health services

- a major activity of cancer registries.
- etiologic or "analytic" epidemiology
 study of cause-effect relationships
- disease epidemiologies e.g. of cancer, cardiovascular diseases, infectious diseases, musculoskeletal disorders, mental health, ...
- determinant-based epidemiologies e.g. occupational epidemiology, nutritional epidemiology, ...
- clinical epidemiology study of diagnosis, prognosis and effectiveness of therapies in patient populations
 - basis of evidence-based medicine

Frequency (from Webster's Dictionary)

Etymology: *frequentia* = assembly, multitude, crowd.

- 1. Also, **frequency**. the state or fact of being frequent; frequent occurrence. We are alarmed by the frequency of fires in the neighborhood.
- 2. Rate of occurrence: The doctor has increased the frequency of his visits.
- 3. *Physics*: number of periods or ... regularly occurring events ... of any given kind in unit of time, usually in one second.
- 4. *Math*: the number of times a value recurs in a unit change of the independent variable of a given function.
- 5. *Statistics*: the number of items occurring in a given category. Cf. **relative frequency**.

Meanings 2 and 5 are both relevant in epidemiology.

But what is "rate" and "occurrence"?

Cancer i Norden 1997 (NORDCAN)

Frequency of cancer (all sites excl. non-melanoma skin) in Nordic male populations expressed by different measures:

| | New cases | Crude rate | ASR (World) | Cumul. risk | SIR |
|---------|--------------|---------------|----------------|----------------|-----------|
| Denmark | 11,787 | 452 | 281 | 27.8 | 104 |
| Finland | 10,058 | <u>401</u> | 269 | 26.5 | 101 |
| Iceland | <u>633</u> | 464 | 347 | 32.6 | 132 |
| Norway | 10,246 | 469 | 294 | 29.4 | 109 |
| Sweden | 19 908 | 455 | <u>249</u> | <u>25.4</u> | <u>93</u> |

- Where is the frequency truly highest, where lowest?
- What do these measures mean?

Questions on frequency & occurrence

How many women in Denmark:

- ► are carriers of breast cancer today? prevalence
- ▶ will contract a new breast ca. during 2007? incidence
- die from breast ca. in 2007? mortality
- will be alive after 5 years since diagnosis among those getting breast ca. in 2007? — survival
- ▶ are cured from breast cancer during 2007? cure

Questions on frequency & occurrence

- What is the relative frequency or/and rate of occurrence of these states and events?
- How great are the **risks** of these events?
- Is the frequency/occurrence/risk of breast cancer greater among nulliparous than parous women?
- What are the excess and relative risks for nulliparous compared to parous women?
- What is the dose-response relationship between occupational exposure to crystalline silica and the risk of getting lung cancer in terms of level and length of exposure?

What is risk?

What do we mean by "risk of disease S"?

- (a) probability of getting S during a given risk period \rightarrow incidence probability,
- (b) rate of change of that probability \rightarrow hazard or intensity,
- (c) **probability** of *carrying* S at a given *time point* \rightarrow **prevalence** probability.

Most commonly meaning (a) is attached with risk. **NB:** "Risk" should not be used in the meaning of **risk factor** However, in **risk assessment** literature: "hazard" is often used in that meaning. In statistics, though, hazard refers to notion (b): change of probability per unit time.

Risks are conditional probabilities

- There are no "absolute risks".
- > All risks are conditional on a multitude of factors, like
 - length of risk period (e.g. next week or lifetime),
 - age and gender,
 - genetic constitution,
 - health behaviour & environmental exposures.
- In principle each individual has a "personal" value for the risk of given disease in any defined risk period, depending on his/her own risk factor profile.
- > Yet, these individual risks are latent and unmeasurable.
- Average risks of disease in large groups sharing common characteristics (like gender, age, smoking status) are estimable from appropriate epidemiologic studies by pertinent measures of occurrence.

Types of epidemiologic studies

Can crudely be classified along the following axes:

- study question: descriptive \leftrightarrow causal
- ► study unit: individual ↔ aggregate (ecological study)
- ► allocation of exposure: experimental ↔ observational
- *population*: closed (cohort) \leftrightarrow open (dynamic)
- ► *dimensionality*: cross-sectional ↔ longitudinal
- ► timing of observations: concurrent ↔ historical ("pro-" vs. "retrospective")
- ► sampling of exposure data: cohort ↔ case-control

Focus in this course: *observational*, and *longitudinal cohort and case-control* studies.

Descriptive and causal questions

Descriptive: What is the occurrence of outcome C in different population groups.

— Medical demography

Descriptive (II) — groups defined *e.g.* by exposure to a determinant or risk factor X?

Causal (also etiological or "analytical"): What is the occurrence of outcome C in a population exposed to risk factor X as compared to ... what the occurrence in the same population would have been, if not exposed?

N.B.: Causal question — *counterfactual conditional*! Challenge: How to find a *comparable* group of unexposed?

Experimental and observational studies

Allocation of exposure in etiologic studies?

- **Experimental**: Exposure controlled by investigators, its levels being **randomized** among the study subjects.
 - + **Comparability** of exposure groups.
 - + Feasible in clinical and preventive trials.
 - Ethically impossible for hazardous exposures.
- **Observational**: Exposure imposed by the own behaviour of the subjects themselves & and by their environment.
 - Possibility of confounding: due to other determinants of the outcome, correlated with exposure.
 - * Challenges: **Valid**: and **efficient** non-randomized design and statistical analysis.

Experimental and observational studies

Allocation of exposure or risk factor in causal studies?

Experimental (Intervention trial): Exposure is controlled by investigators; its levels are allocated among recruited subjects by **randomization**,

 \Rightarrow **comparability** of exposure groups.

Observational: Exposure imposed by own behaviour of study subjects and/or by their environment,

 \Rightarrow possibility of **confounding** due to other determinants.

Time dimensionality of a study

Cross-sectional: Outcome status and its prevalence in population at given time point are studied, e.g.
number of Danish citizens living with existing cancer on 13 August 2007.

Longitudinal: *Change* in health status, like the *incidence* of new cases over a *time period* is of interest, *e.g.*

 number of Danish citizens getting a new cancer diagnosed during year 2007.

Causal question \longrightarrow longitudinal study preferred.

Study population & study base

Types of study population & its membership defined

- closed cohort: members taken by certain event, e.g.
 - 1. birth cohort, people born during same year,
 - 2. workers employed by Carlsberg brewery during 1970's, followed up since then, even after retirement
- open dynamic: defined by changeable status, e.g.
 - 1. citizens of Copenhagen, currently resident;
 - 2. *catchment population* of the Oncological Clinic at Rigshospitalet (CPH),
- **Study base** = study population \times its experience in time.

Study base (SB): population experience

Cross-sectional: SB = study population at a*time point*,Longitudinal: SB comprises follow-up times of individuals in the study population over a given *period*. Cohort: Follow-up time = period from entry until a single exit at which outcome or censoring occurs. Dynamic: Follow-up time consists of possibly several periods of membership since the first entry until the final exit. Follow-up calculation

- complicated.
- Approximation by mid-population.

R and how we use it Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

What is R?

- A practical calculator:
 - You can see what you compute
 - …and change easily to do similar calculations.
- A statistical program.
- An environment for data analysis and graphics.
- Free.
- Runs on any computer.
- Updated every 6 months.

A simple calculator

R lets you enter simple arithmetic and giver you back the ansver straightaway:

```
> 5+8
[1] 13
> sqrt( 1/12 + 1/17 )
[1] 0.3770370
> exp( 1.96 * sqrt( 1/12 + 1/17 ) )
[1] 2.093825
> D0 <- 12
> D1 <- 17
> exp( 1.96 * sqrt( 1/D0 + 1/D1 ) )
[1] 2.093825
```

Handy in daily life too.

A smart calculator

Case-control study of MI:

| | Men | | Women | | |
|------------------------|------|------|-------|------|------|
| PA index | Case | Cont | | Case | Cont |
| 2500+ kcals | 141 | 208 | | 49 | 58 |
| $< 2500 {\rm \ kcals}$ | 144 | 112 | | 32 | 45 |
| Total | 285 | 320 | | 81 | 103 |

> (141/208)/(144/112)
[1] 0.5272436
> (49/58)/(32/45)
[1] 1.188039

A smart calculator

> D1 <- c(141, 49) > D0 <- c(144, 32) > H1 <- c(208, 58) > H0 <- c(112, 45) > OR <- (D1/D0)/(H1/H0) > OR [1] 0.5272436 1.1880388

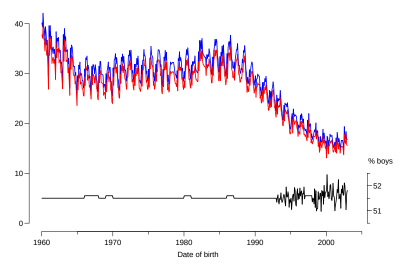
Things done in parallel for the two exposure groups.

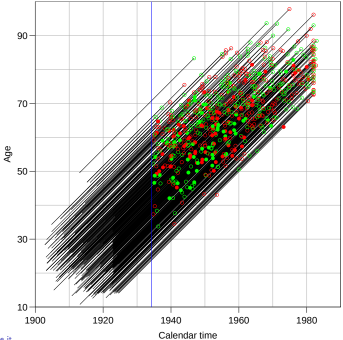
R for epidemiology

Versatile graphics:

- Simple graphs easy
- Complicated graphs possible
- You can add things to a graph
- Interactive graphs:
 - Put things on with the mouse
 - Identify points with the mouse

Monthly no. births in Ukraine (1000s)





Getting your graphs out

You can save graphs to disk and later fetch them into your documents in almost any format you like:

(.eps, .pdf, .emf, .bmp, .png).

You can choose to save graphs from the screen or to write directly to a file.

Tools for anything!

- More than 1500 add-on packages.
- Several packages for epidemiology:
 - Epi: Mostly chronic disease epidemiology:
 - Cohort studies, split follow-up time
 - Lexis diagram, sevral timescales
 - Multistate model support
 - Advanced tabulation
 - Parameter reporting
 - epicalc: For a book by Virasakdi Chongsuvivatwong.
 - epitools: Mostly infectious diseases.
 - epiR: Leaning towards veterinay epidemiology.
- Install and update packages from within R.

Versatility is paid by steep learning curve

Command line interface:

- You must write commands
- You must know what they are called
- Easy to repeat analyses, because you always have a script of what you did.
- There is a simple editor built into R.
- A good workbook introduction is: www.mhills.pwp.blueyonder.co.uk/Rwork_book. html
- Many other introductions to R on the R homepage.

R in this course

- Only use R as a simple calculator.
- No need for for a lot of fancy stuff.
- The script editor (we will show you what that is) will help you keep your solutions for future reference.
- A short recap of exercises tomorrow morning, and tomorrow afternoon.
- After the course, solutions to all exercises will be provided.

Frequency measures Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Measuring frequency: Cases, population, time

Quantification of the occurence of disease (or any other health-related state or event) requires specification of:

 what is meant by a case, i.e., an individual in a population who has or gets the disease (more generally: possesses the state or undergoes the event of interest).

 \Rightarrow challenge to accurate diagnosis and classification!

- 2. the **population** from which the cases originate.
- 3. the **time point** or **period** of observation.

Types of occurrence measures

- ► Longitudinal **incidence** measures.
- Cross-sectional prevalence measures.

General form of frequency or occurrence measures

numerator

denominator

Numerator: number of cases observed in the population

- at a certain time point or during a specified period.

Denominator: generally proportional to the size of the population from which the cases emerge.

Numerator and denominator must cover the same population.

Prevalence

Prevalence:

Point prevalence, is the proportion of existing cases (old and new) in a population at a single point of time.

 $P = \frac{\text{No. of existing cases in a population at one point of time}}{\text{No. of people in the population at the same point of time}}$

This measure is called point prevalence, because it refers to a single point in time. It is often referred to simply as prevalence.

Incidence measures

Incidence proportion (Q) over a fixed *risk period*:

 $Q = \frac{\text{number of incident (new) cases during period}}{\text{size of pop'n at risk at start of the period}}$

Also called **cumulative incidence** or **cumulative risk** (*e.g.* by **IS**).

Indidence rate (I) over a defined observation period:

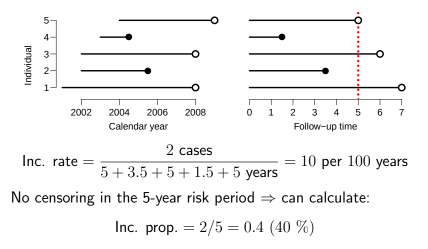
 $I = \frac{\text{number of incident (new) cases during period}}{\text{sum of follow-up times of pop'n at risk}}$

Also called incidence density or hazard.

Later we will provide a more precise mathematical definition of the concepts.

Example: Follow-up of a small cohort

- $\circ =$ exit with censoring; outcome not observed,
- = exit with outcome event (disease onset) observed



Properties of incidence proportion

- Dimensionless quantity ranging from 0 to 1 (0% to 100%) = relative frequency,
- Estimates the average theoretical **risk** or probability of the outcome occurring during the risk period, in the **population at risk** — *i.e.* among those who are still free from the outcome at the start of the period,
- Simple formula valid when the follow-up time is fixed & equals the risk period, and when there are no competing events or censoring (see below),
- Competing events & censoring ⇒ Calculations need to be corrected using special methods of survival analysis.

Properties of incidence rate

- Like a frequency quantity in physics; it is a scaled quantity; it is measured in time⁻¹: cases/1000 Y, say.
- Estimates the average underlying intensity or hazard rate of the outcome in a population,
- Estimation accurate in the constant hazard model,
- Calculation straightforward also with competing events and censored observations.
- ► Hazard depends on age (& other time variables) ⇒ rates specific to age group etc. needed,
- Incidence proportions can be estimated from rates. In the constant hazard model with no competing risks:

$$Q = 1 - \exp(-I \times \Delta) \approx I \times \Delta$$

(we shall return to the derivation of this).

Competing events and censoring

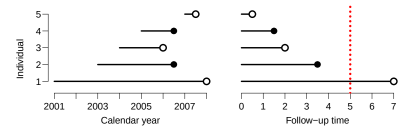
The outcome event of interest (*e.g.* onset of disease) is not always observed for all subjects during the chosen risk period.

- ► Some subjects die (from other causes) before the event. ⇒ Death is a competing event after which the outcome cannot occur any more.
- Others emigrate and escape national disease registration, or the whole study is closed "now", which prematurely interrupts the follow-up of some individuals,

 \Rightarrow censoring, withdrawal, or loss to follow-up

Competing events and censorings require special statistical treatment in incidence and risk calculations.

Follow-up of another small cohort



Two censored observations \Rightarrow can calculate the rate:

$$I=2/12.5$$
 y $=16$ per 100 years

but the 5-year Q is no more 2/5 ! However, under constant rate model

$$Q = 1 - \exp(-5 \times 2/12.5) = 0.55$$

Person-years in dynamic populations

With dynamic study population individual follow-up times are always variable and impossible to measure accurately.

Common approximation – **mid-population** principle:

- ► Let the population size be N_{t-1} at start and N_t at the end of the observation period t with length L_t years,
- Mid-population for the period: $\bar{N}_t = \frac{1}{2} \times (N_{t-1} + N_t)$.
- Approximate person-years: $Y_t \approx \bar{N}_t \times L_t$.

NB. The actual study population often contains also some already affected, who thus do not belong to the population at risk. With rare outcomes the influence of this is small.

Male person-years in Finland 1991-95

Total male population (1000s) on 31 December by year:

| 1990 | 1991 | 1992 | 1993 | 1994 | 1995 |
|------|------|------|------|------|------|
| 2431 | 2443 | 2457 | 2470 | 2482 | 2492 |

Approximate person-years (1000s):

1992:
$$\frac{1}{2} \times (2443 + 2457) \times 1 = 2450$$
1993-94: $\frac{1}{2} \times (2457 + 2482) \times 2 = 4937$ 1991-95: $\frac{1}{2} \times (2431 + 2492) \times 5 = 12307.5$

Relationships between incidence measures

With constant incidence rate over risk period (length = Δ), incidence proportion Q and rate I are related:

$$Q = 1 - \exp(-I \times \Delta) \approx I \times \Delta$$
$$I = -\log(1 - Q)/\Delta \approx Q/\Delta,$$

The approximations are good when

- ▶ the incidence proportion is "small" (under 10 %).
 - incidence rate (I) is small
 - the risk period (Δ) is small

Mortality

Cause-specific mortality from disease C is described by **mortality rate** (and proportion), defined like I (and Q), but

- ► cases are only *deaths* from cause *C*, and
- follow-up is extended until death (from *any* acuse) or censoring

The cumulative risk of death from a given cause (cause-specific mortality proportion/risk) requires correction for *competing events*. **Total mortality**: cases are deaths from any cause. Mortality depends on the incidence and the **prognosis** or fatality of the disease, *i.e.* the **survival** of those affected.

Theoretical concepts behind incidences

Analysis of incidences

= analysis of time to event or failure time or survival data.

Mathematical concepts:

$$\begin{array}{lll} T &=& {\rm time \ to \ outcome \ event - random \ variable,} \\ S(t) &=& P(T>t) = \ {\rm survival \ function \ of \ } T, \\ &=& {\rm probability \ of \ avoiding \ the \ event \ up \ to \ given \ time \ } t, \\ \lambda(t) &=& -S'(t)/S(t) = \ {\rm intensity \ or \ hazard \ function,} \\ \Lambda(t) &=& \int_0^t \lambda(u) du = -\log S(t) = {\rm cumulative \ hazard,} \\ F(t) &=& 1 - S(t) = 1 - \exp\{-\Lambda(t)\} = {\rm risk \ function} \\ &=& {\rm probability \ of \ the \ outcome \ to \ occur \ before \ t} \end{array}$$

Intensity or hazard function

Can be viewed as theoretical incidence rate. Formally:

$$\lambda(t) = \lim_{\Delta \to 0} \frac{P(t < T \le t + \Delta \mid T > t)}{\Delta}$$

 \approx Probability of outcome event occurring in a short risk period $]t, t + \Delta]$, given "survival" or avoidance of the event up to the start t, divided by the period length — "risk per time".

This is equivalent to saying that over a short interval

risk $\,pprox\,$ intensity $\, imes\,$ length of interval

$$\text{ or } \qquad P(t < T \leq t + \Delta \mid T > t) \; \approx \; \lambda(t) \times \Delta.$$

Exponential survival times (constant hazard)

Simplest probability model for time to event: **Exponential distribution**, $Exp(\lambda)$, in which

rate $\lambda(t) = \lambda$ (constant) \Rightarrow risk over $]0, t] = 1 - \exp(-\lambda t)$

Analysis of event data of n individuals. For subject i let

 y_i = time to event or censoring, total: $Y = \sum y_i$ d_i = 1/0-indicator for observing event, total: $D = \sum d_i$

 $Exp(\lambda) \mod \Rightarrow$ Likelihood function of λ is equivalent to that when number of cases D is *Poisson*-distributed (Analysis part of the course)

Basic statistical analysis of empirical rates

Asymptotic statistical inference based on likelihood:

• Maximum likelihood estimator (MLE) of λ is

 $\widehat{\lambda} = \frac{D}{Y} = \frac{\text{number of cases}}{\text{total person-time}} = I, \text{ empirical incidence rate!}$

- Standard error of the empirical rate is I/\sqrt{D}
- $\Rightarrow\,$ The more cases, the greater is precision in rate!
 - Approximate confidence interval for "true" rate λ:

estimator $\pm 1.96 \times$ standard error

More about these issues in the analysis lectures.

Prevalence measures

Point prevalence or simply **prevalence** P of a health state C in a population at a given time point t is defined

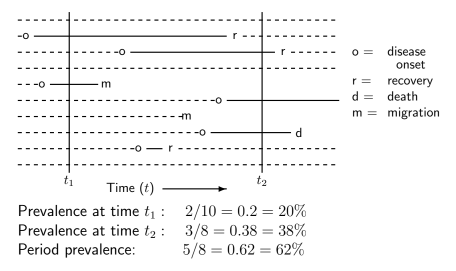
 $P = \frac{\text{number of existing or prevalent cases of } C}{\text{size of the whole population}}$

This is calculable from a cross-sectional study base.

Period prevalence for period from t_1 to t_2 is like P but

- numerator refers to all cases prevalent already at t₁ plus new cases occurring during the period, and
- denominator is the population size at t_2 .

Example 4.1 (IS: p. 59)



Relationships between measures

Point prevalence of C at given time point t depends on

- *incidence* of new cases of C before t
- duration of C, depending in turn on the probability of cure or recovery from C or survival of those affected.

Stationary ("stable") population: prevalence (P), incidence (I), and average duration (\overline{d}) of C are related:

$$P = \frac{I \times \bar{d}}{I \times \bar{d} + 1} \quad \approx \quad I \times \bar{d}$$

 $\mathsf{prevalence} = \mathsf{incidence} \times \mathsf{duration}$

The approximation works well, when P < 0.1 (10%).

Prevalence of cancer?

Difficult to ascertain, whether and when a cancer is cured.

- \Rightarrow Existing or prevalent cancer case problematic to define.
- Cancer registry practice: Prevalence of cancer C at time point t in the target population refers to the

number & proportion of population members who

- \blacktriangleright are alive and resident in the population at t, and
- ▶ have a record of incident cancer C diagnosed before t.

Often further classified by years since diagnosis.

Example: Liver and testis cancer

Crude comparison of incidence, mortality and prevalence in the male population of Finland 1999

| | Liver | Testis |
|--|----------------------|-------------------------|
| No. of new cases during 1999 | 119 | 103 |
| No. of deaths during 1999 | 123 | 8 |
| No. of prevalent cases 1.1.2000 | 120 | 1337 |
| – " – diagnosed < 1 y ago – " – diagnosed 1-< 5 y ago – " – diagnosed 5-< 10 y ago – " – diagnosed > 10 y ago | 36 53 17 14 | 97 291 304 642 |

Comparative measures

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Relative and absolute comparisons (IS: Ch 5.2)

Quantification of the **association** between a determinant (risk factor or exposure) and an outcome (disease) is based on **comparison of occurrence** between the *index* ("exposed") and the *reference* ("unexposed") groups or populations by

- relative measures (ratio)
- absolute measures (difference)

In causal studies these are used to estimate the **causal effect** of the exposure factor on the disease risk.

 \Rightarrow comparative measures \approx effect measures

Relative comparative measures

Generic name "**relative risk**" RR comparing occurrences between exposed (1) and unexposed (0) groups can be

- incidence rate ratio I_1/I_0 ,
- incidence proportion ratio Q_1/Q_0 ,
- incidence odds ratio $[Q_1/(1-Q_1)]/[Q_0/(1-Q_0)]$,
- prevalence ratio P_1/P_0 , or
- prevalence odds ratio $[P_1/(1-P_1)]/[P_0/(1-P_0)]$,

depending on study base and details of its design.

Absolute comparative measures

Generic "excess risk" btw exposed and unexposed can be

- incidence rate difference $I_1 I_0$,
- incidence proportion difference $Q_1 Q_0$,
- prevalence difference $P_1 P_0$.

Use of relative and absolute comparisons

Ratio – describes the *biological strength* of the exposure Difference – informs about its *public health importance*.

Example: (IS, Table 5.2, p.97)

Relative and absolute comparisons between the exposed and the unexposed to risk factor X in two diseases.

| | Disease A | Disease B |
|---|-----------|-----------|
| Incidence rate among exposed ^a | 20 | 80 |
| Incidence rate among unexposed a | 5 | 40 |
| Rate ratio | 4.0 | 2.0 |
| Rate difference ^a | 15 | 40 |

^a Rates per 100 000 pyrs.

Factor X has a stronger biological potency for disease A, but it has a greater public health importance for disease B.

Ratio measures in "rare diseases" (IS: Ex 5.13)

| | | | Exposure | | | | |
|-------------------|-----------------------|--|----------|---|----|-----|--------|
| | | | Y | es | No | 0 | _ |
| No. initi | No. initially at risk | | | 4 000 | 16 | 000 | - |
| Deaths | Deaths | | | 30 | | 60 | |
| Person-y | Person-years at risk | | | 7 970 | 31 | 940 | |
| | | | | | | | - |
| Inc. prop'n ratio | = | $\frac{30/4000}{60/16000}$ | = | $rac{7.5}{3.75}$ per $rac{1}{2}$ | | = | 2.0000 |
| Inc. rate ratio | = | $\frac{30/7970~{\rm y}}{60/31940~{\rm y}}$ | = | $\frac{3.76 \text{ per}}{1.88 \text{ per}}$ | | = | 2.0038 |
| | = | $\frac{0.00756}{0.00376}$ | = | 2.00 | 76 | | |

Attributable fraction

Combine absolute and relative comparisons.

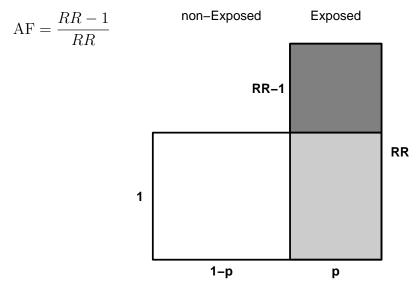
When incidence is higher for the exposed, we can calculate

Excess fraction,
$$\mathsf{EF} = rac{Q_1 - Q_0}{Q_1} = rac{\mathsf{RR} - 1}{\mathsf{RR}}$$

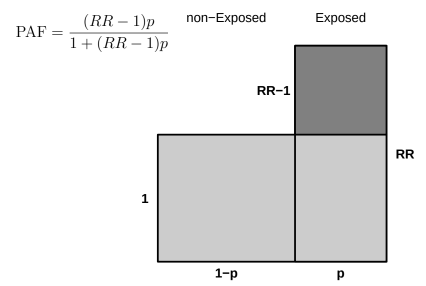
also called attributable fraction, AF or attributable risk.

EF Estimates the fraction out of all new cases among those exposed, which are "caused" by the exposure itself, and which thus could be "avoided" if the exposure were absent

Attributable fraction, AF



Population attributable fraction, PAF



Population attributable fraction

If we instead ask:

"How large a fraction of **all** cases would be prevented if exposure was abolished?".

Depends on the fraction of the population which is exposed

$$PAF = \frac{(RR - 1)p}{1 + (RR - 1)p}$$

PAF Estimates the fraction out of all new cases, which are "caused" by the exposure itself, and which thus could be "avoided" if the exposure were absent.

AF is a "biological" measure. PAF is a "population level" measure.

Measures of potential impact (cont'd)

When the exposed have a lower incidence, we can calculate

Preventive fraction,
$$\mathsf{PF} = rac{Q_0 - Q_1}{Q_0} = 1 - \mathsf{RR}$$

also called **relative risk reduction** = percentage of cases prevented among the exposed due to the exposure.

Used to evaluate the relative effect of a preventive intervention (exposed) vs. no intervention (unexposed).

Effect of smoking on mortality by cause

(**IS**: Example 5.14, p. 98)

| Underlying cause of death | Never smoked regularly Rate ^b | Current cigarette smoker Rate ^b | Rate ratio | Rate differ- ence ^b | Excess fraction (%) |
|---------------------------------|---|---|---------------|--------------------------------------|------------------------------------|
| | (1) | (2) | (2)/(1) | (2) - (1) | $\frac{(2) - (1)}{(2)} \times 100$ |
| Cancer | | | | | |
| All sites | 305 | 656 | 2.2 | 351 | 54 |
| Lung | 14 | 209 | 14.9 | 195 | 93 |
| Oesophagus | 4 | 30 | 7.5 | 26 | 87 |
| Bladder | 13 | 30 | 2.3 | 17 | 57 |
| Respiratory diseases | | | | | |
| (except cancer) | 107 | 313 | 2.9 | 206 | 66 |
| Vascular diseases | 1037 | 1643 | 1.6 | 606 | 37 |
| All causes | 1706 | 3038 | 1.8 | 1332 | 44 |

^a Data from Doll et al., 1994a.

^b Age-adjusted rates per 100 000 pyrs.

Time scales

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Incidence by age, calendar year, and other time variables

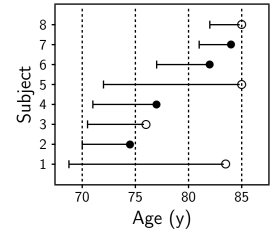
Incidence can be studied on various time scales, e.g.:

| Time scale | Origin (date of:) |
|---------------------|-------------------|
| age | birth |
| exposure time | first exposure |
| follow-up time | entry to study |
| duration of disease | diagnosis |

Age is usully the strongest time-dependent determinant of health outcomes.

Age is also often correlated with duration of "chronic" exposure (e.g. years of smoking).

Follow-up of a geriatric cohort



Overall rate: 4 cases/53.5 person-years = 7.5 per 100 y Hides the fact that the "true" rate probably varies by age, being higher among the old.

Person-years and cases in agebands: age-specific rates

| Subject | 70-74 | 75-79 | 80-84 | Total |
|---------------------|-------|---------|-------|-------|
| 1 | 5.0 | 5.0 | 3.5 | 13.5 |
| 2 | 4.5 | - | - | 4.5 |
| 3 | 4.5 | 1.0 | - | 5.5 |
| 4 | 4.0 | 2.0 | - | 6.0 |
| 5 | 3.0 | 5.0 | 5.0 | 13.0 |
| 6 | - | 3.0 | 2.0 | 5.0 |
| 7 | - | - | 3.0 | 3.0 |
| 8 | - | - | 3.0 | 3.0 |
| Sum of person-years | 21.0 | 16.0 | 16.5 | 53.5 |
| Cases | 1 | 1 | 2 | 4 |
| Rate (/100 y) | 4.8 | 6.2 | 12.1 | 7.5 |
| | Age- | overall | | |

Lung cancer incidence rates in Finland by age, period and cohort

| Calendar | Age group (y) | | | | | | | | | |
|----------|---------------|-------|-------|-------|-------|-------|-------|-------|-------|-----|
| period | 40-44 | 45-49 | 50-54 | 55-59 | 60-64 | 65-69 | 70-74 | 75-79 | 80-84 | 85+ |
| 1953-57 | 21 | 61 | 119 | 209 | 276 | 340 | 295 | 279 | 193 | 93 |
| 1958-62 | 22 | 65 | 135 | 243 | 360 | 405 | 429 | 368 | 265 | 224 |
| 1963-67 | 24 | 61 | 143 | 258 | 395 | 487 | 509 | 479 | 430 | 280 |
| 1968-72 | 21 | 61 | 134 | 278 | 424 | 529 | 614 | 563 | 471 | 358 |
| 1973-77 | 16 | 50 | 134 | 251 | 413 | 541 | 629 | 580 | 490 | 392 |
| 1978-82 | 13 | 36 | 115 | 234 | 369 | 514 | 621 | 653 | 593 | 442 |
| 1983-87 | 11 | 31 | 74 | 186 | 347 | 450 | 566 | 635 | 592 | 447 |
| 1988-92 | 9 | 25 | 57 | 128 | 262 | 411 | 506 | 507 | 471 | 441 |
| 1993-97 | 7 | 22 | 48 | 106 | 188 | 329 | 467 | 533 | 487 | 367 |
| 1998-02 | 5 | 14 | 46 | 77 | 150 | 239 | 358 | 445 | 396 | 346 |

- Rows: age-incidence pattern in different calendar periods.
- Columns: Trends of age-specific rates over calendar time.
- Diagonals: age-incidence pattern in birth cohorts.

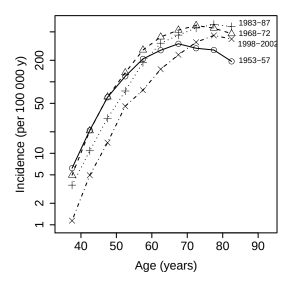
Incidence by age, calendar time & birth cohort

 Secular trends of specific and adjusted rates show, how the "cancer burden" has developed over periods of calendar time.

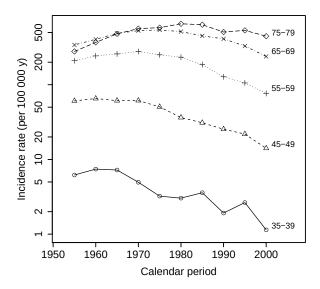
Birth cohort = people born during the same limited time interval, *e.g.* single calendar year, or 5 years period.

- Analysis of rates by birth cohort reveals, how the level of incidence (or mortality) differs between successive generations.
- Often more informative about "true" age-incidence pattern than age-specific incidences of single calendar period.

Age-incidence curves by period (rows)



Time trends by age (columns)

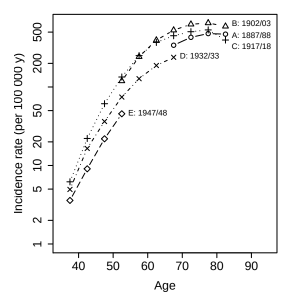


Age-specific rates by birth cohort

| Calendar | | | | Age gr | oup (y) | | | | |
|----------|-------|-------|--------|--------|---------|-------|-------|-------|---|
| period | 40-44 | 45-49 | 50-54 | 55-59 | 60-64 | 65-69 | 70-74 | 75-79 | |
| 1953-57 | 21 | 61 | 119 | 209 | 276 | 340 | 295 | 279 | |
| 1958-62 | 22 | 65 | 135 | 243 | 360 | 405 | 429 | 368 | |
| 1963-67 | 24 | 61 | 143 | 258 | 395 | 487 | 509 | 479 | А |
| 1968-72 | 21 | 61 | 134 | 278 | 424 | 529 | 614 | 563 | |
| 1973-77 | 16 | 50 | 134 | 251 | 413 | 541 | 629 | 580 | |
| 1978-82 | 13 | 36 | 115 | 234 | 369 | 514 | 621 | 653 | В |
| 1983-87 | 11 | 31 | 74 | 186 | 347 | 450 | 566 | 635 | |
| 1988-92 | 9 | 25 | 57 | 128 | 262 | 411 | 506 | 507 | |
| 1993-97 | 7 | 22 | 48 | 106 | 188 | 329 | 467 | 533 | С |
| 1998-02 | 5 | 14 | 46 | 77 | 150 | 239 | 358 | 445 | |
| | | | E: 194 | 47/48 | | D: 19 | 32/33 | | |

A = synthetic cohort born around 1887/88, B: 1902/03, C: 1917/18

Age-incidence curves in 5 birth cohorts



Split of follow-up by age and period

Incidence of (or mortality from) disease C in special study cohort (e.g. occupational group, users of certain medicine)

 $\rightarrow\,$ often compared to incidence in a $\it reference$ or "general" population

Appropriate adjustment for age and calendar time needed in this, *e.g.* by comparing *observed* to *expected* cases with SIR (see p. 70-71).

 $\Rightarrow\,$ Cases and person-years in the study cohort must be split by more than one time scale (age).

Example of follow-up

| Entry and exit dates for a small cohort of four subjects | | | | | | | | | |
|--|------------------------------|-----------------------------------|---|---|--|--|--|--|--|
| Born | Entry | Exit | Age at entry | Outcome | | | | | |
| 1904 | 1943 | 1952 | 39 | Migrated | | | | | |
| 1924 | 1948 | 1955 | 24 | Disease C | | | | | |
| 1914 | 1945 | 1961 | 31 | Study ends | | | | | |
| 1920 | 1948 | 1956 | 28 | Unrelated death | | | | | |
| | Born 1904 1924 1914 | BornEntry190419431924194819141945 | BornEntryExit190419431952192419481955191419451961 | BornEntryExitAge at entry190419431952391924194819552419141945196131 | | | | | |

·· · · · · ı. c c ı. 1.1

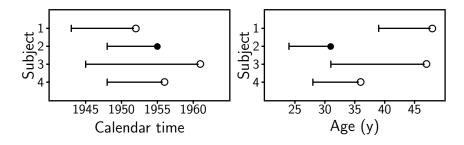
Subject 1: Follow-up time spent in each ageband

| Age band | Date in | Date out | Time (years) |
|----------|---------|----------|--------------|
| 35–39 | 1943 | 1944 | 1 |
| 40–44 | 1944 | 1949 | 5 |
| 45–49 | 1949 | 1952 | 3 |

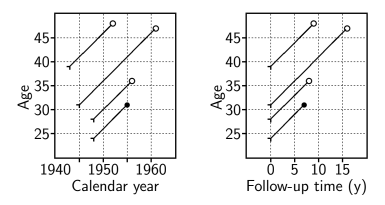
Follow-up of cohort members by calendar time and age

entry

- exit because of disease onset (outcome of interest)
- exit due to other reason (censoring)



Follow-up in Lexis-diagrams — by age and period



Follow-up lines run diagonally through different ages and calendar periods.

Standardization

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Crude & adjusted rates

- Incidence of most cancers (and many other diseases) increases strongly by age in all populations.
 - \Rightarrow Most of the caseload comes from older age groups.
- Crude incidence rate is a rate in which:
 - numerator = sum of age-specific numbers of cases,
 - denominator = sum of age-specific person-years.
- This is generally a poor **summary measure**.
- Comparisons of crude incidences between populations can be very misleading, when the age structures differ.
- Solution: Standardization.

Stomach cancer in Cali and Birmingham (IS, Table 4.2, p. 71)

| | | Cali | | | | | |
|----------------------|----------------------|---------------------------------------|-------------------------------------|----------------------|---------------------------------------|-------------------------------------|----------------------|
| | No. of Male | Male Popu- | Inci- Rate | No. of Male | Male Popu- | Inci- Rate | |
| Age (y) | cases 1982 -86 | lation 1984 (10 ³ s) | (/10 ⁵ y) 1982 -86 | cases 1983 -86 | lation 1985 (10 ³ s) | (/10 ⁵ y) 1983 -86 | Rate ratio |
| 0–44 45-64 65+ | 39 266 315 | 524.2 76.3 22.4 | 1.5 69.7 281.3 | 79 1037 2352 | 1 683.6 581.5 291.1 | 1.2 44.6 202.0 | 1.25 1.56 1.39 |
| Total | 620 | 622.9 | 19.9 | 3468 | 2 556.2 | 33.9 | 0.59 |

In each age group Cali has a higher incidence but the crude incidence is higher in Birmingham. *Is there a paradox*?

Comparison of age structures (IS, Tables 4.3,4.4)

| | (| % of male population | | | | | | |
|----------|------|----------------------|---------|--------|--|--|--|--|
| Age | Cali | B'ham | Finland | World | | | | |
| (years) | 1984 | 1985 | 1999 | Stand. | | | | |
| 0–44 | 84 | 66 | 61 | 74 | | | | |
| 45–64 | 12 | 23 | 27 | 19 | | | | |
| 65+ | 4 | 11 | 12 | 7 | | | | |
| All ages | 100 | 100 | 100 | 100 | | | | |

- The fraction of old men greater in Birmingham than in Cali. \Rightarrow The crude rates are **confounded** by age.
- \Rightarrow Any summary rate must be **adjusted for age**.

Age-adjustment by standardisation

Age-standardised incidence rate (ASR):

$$\mathsf{ASR} = \sum_{k=1}^{K} \mathsf{weight}_k imes \mathsf{rate}_k \ / \ \mathsf{sum} \ \mathsf{of} \ \mathsf{weights}$$

- = Weighted average of age-specific rates over the age-groups $k = 1, \ldots, K$.
- Weights describe age distribution of some standard population.
- Standard population can be real (*e.g.* one of the populations under comparison, or their average) or fictitious (*e.g.* World Standard Population, WSP)

Some standard populations:

| Age group (years) | African | World | European | Truncated |
|-------------------|---------|---------|----------|-----------|
| 0 | 2 000 | 2 400 | 1 600 | - |
| 1-4 | 8 000 | 9 600 | 6 400 | - |
| 5–9 | 10 000 | 10 000 | 7 000 | - |
| 10-14 | 10 000 | 9 000 | 7 000 | - |
| 15–19 | 10 000 | 9 000 | 7 000 | - |
| 20–24 | 10 000 | 8 000 | 7 000 | - |
| 25–29 | 10 000 | 8 000 | 7 000 | - |
| 30–34 | 10 000 | 6 000 | 7 000 | - |
| 35–39 | 10 000 | 6 000 | 7 000 | 6 000 |
| 40–44 | 5 000 | 6 000 | 7 000 | 6 000 |
| 45–49 | 5 000 | 6 000 | 7 000 | 6 000 |
| 50–54 | 3 000 | 5 000 | 7 000 | 5 000 |
| 55–59 | 2 000 | 4 000 | 6 000 | 4 000 |
| 60–64 | 2 000 | 4 000 | 5 000 | 4 000 |
| 65–69 | 1 000 | 3 000 | 4 000 | - |
| 70–74 | 1 000 | 2 000 | 3 000 | - |
| 75–79 | 500 | 1 000 | 2 000 | - |
| 80–84 | 300 | 500 | 1 000 | - |
| 85+ | 200 | 500 | 1 000 | _ |
| Total | 100 000 | 100 000 | 100 000 | 31 000 |

Stomach cancer in Cali & B'ham

Age-standardized rates by the World Standard Population:

| | | Cali | Bi | rmingham |
|----------------------|----------------------------|--|----------------------------|---|
| Age | $Rate^a$ | Weight | $Rate^a$ | Weight |
| 0–44 45–64 65+ | 1.5 × 69.7 × 281.3 × | $\begin{array}{c} 0.74 = \ 1.11 \\ 0.19 = 13.24 \\ 0.07 = 19.69 \end{array}$ | 1.2 × 44.6 × 202.0 × | $\begin{array}{r} 0.74 = \ 0.89 \\ 0.19 = \ 8.47 \\ 0.07 = 14.14 \end{array}$ |
| Age-sta | ndardised | rate 34.04 | | 23.50 |

ASR in Cali higher – coherent with the age-specific rates. Summary rate ratio estimate: **standardized rate ratio**

$$SRR = 34.0/23.5 = 1.44$$

Known as **comparative mortality figure (CMF)** when the outcome is death (from specific cause C or all causes).

Cumulative rate and cumulative risk

- Choice of standard population weights somewhat arbitrary.
- Alternative and perhaps more "natural" method for age-adjustment is provided by:

$$\textbf{Cumulative rate} = \sum_{k=1}^{K} \mathsf{width}_k \times \mathsf{rate}_k$$

- ► Weigths are widths of the agebands to be included: Cumulative risk = 1-exp(-cumul. rate) ≈ cumul. rate
- Usually calculated up to 65 or 75 years with 5-year agebands.
- These estimate the average risk in the population to get the disease by 65 or 75 years given survival until then.
- The competing acuses of exit (death) is **not** taken into account.

Stomach cancer in Cali & B'ham

From age-specific rates of Table 4.2. the cumulative rates up to 65 years and their ratio are

Cali:
$$45 \ y \times \frac{1.5}{10^5 y} + 20 \ y \times \frac{69.7}{10^5 y} = 0.0146 = \mathbf{1.46} \text{ per } 100$$

B'ham: $45 \ y \times \frac{1.2}{10^5 y} + 20 \ y \times \frac{44.6}{10^5 y} = 0.0095 = \mathbf{0.95} \text{ per } 100$
ratio: $1.46/0.95 = \mathbf{1.54}$

Cumulative risks (inc. proportions) & their ratio up to 65 y:

Cali:
$$1 - \exp(-0.0146) = 0.0145 = 1.45\%$$

B'ham: $1 - \exp(-0.0095) = 0.0094 = 0.94\%$
ratio: $1.45/0.94 = 1.54$

Cumulative measures in 5-y groups

| | Age-group (years) | Incidence rate (per 100 000 pyrs) | |
|---------|----------------------------------|---|-------------------------------|
| | 0–4,, 15–19 | 0.0 | |
| | 20-24, 25-29 | 0.1 | |
| | 30–34 | 0.9 | |
| | 35–39 | 3.5 | |
| | 40–44 | 6.7 | |
| | 45–49 | 14.5 | |
| | 50–54 | 26.8 | |
| | 55–59 | 52.6 | |
| | 60–64 | 87.2 | |
| | 65–69 | 141.7 | |
| | 70–74 | 190.8 | |
| | Sum | 524.9 | - |
| | Cum. rate 0-75 y = | $= 5 \text{ y} \times \frac{524.9}{10^5 \text{ y}} = 0.0$ | 262 = 2.6% |
| Standar | diz Gilu m. risk 0-75 y = | $= 1 - \exp(-0.0262)$ | $0=0.0259={f 2.6\%}.$ 84/ 102 |

Observed and expected cases

- Suppose O cases are observed in an index population of interest (*e.g.* an occupational cohort) during its follow-up over a lengthy calendar period.
- Question: What would be the expected number of cases E, if the age- and period-specific rates of a reference population for comparison were valid for the index population?
- The ratio "observed/expected" estimates of the "true" rate ratio between the index and the reference populations jointly adjusted for age and period.

Standardized incidence ratio, SIR

Let λ_{kl} = incidence rate in a Lexis-diagram cell defined by ageband k and period l in the reference population. Hence,

expected number
$$(E) = \sum_{k=1}^{K} \sum_{l=1}^{L} \lambda_{kl} \times Y_{kl},$$

where Y_{kl} is the person-years in cell kl of the index population. The **standardised incidence ratio** (SIR) is defined

$$\mathsf{SIR} = \frac{O}{E}$$

When the outcome is death, this measure is called **standardized mortality ratio**, SMR.

SIR for Cali with Birmingham as reference

Total person-years at risk and expected number of cases in Cali 1982-86 based on age-specific rates in Birmingham (**IS**: Fig. 4.9, p. 74)

| Age | Person-years | Expected cases in Cali |
|----------|---------------------------------|--|
| 0–44 | 524 220×5= 2 621 100 | $0.000012 \times 2\ 621\ 100 =\ 31.45$ |
| 45–64 | 76 304×5= 381 520 | $0.000446 \times 381520 = 170.15$ |
| 65+ | $22 \ 398 \times 5 = 111 \ 990$ | $0.002020 \times 111\ 990 = 226.00$ |
| All ages | =3 114 610 | Total expected (E) 427.82 |

Total observed number O = 620. Standardised incidence ratio:

$$\mathsf{SIR} = \frac{O}{E} = \frac{620}{427.8} = 1.45 \quad (\text{or } 145 \text{ per } 100)$$

Crude and adjusted measures

(IS: Table 4.6, p. 78, extended)

| | Cali, 1982-86 | B'ham, 1983-86 | Rate ratio |
|---|------------------|-------------------|---------------|
| Crude rates ($/10^5$ y) | 19.9 | 33.9 | 0.59 |
| ASR $(/10^5 \text{ y})^B$ with 3 broad age groups | 48.0 | 33.9 | 1.42 |
| ASR $(/10^5 \text{ y})^C$ –"– | 19.9 | 14.4 | 1.38 |
| ASR $(/10^5 \text{ y})^W$ -"- | 34.0 | 23.5 | 1.44 |
| Cum. rate < 65 y (per 1000) –"– | 14.6 | 9.5 | 1.54 |
| ASR $(/10^5 \text{ y})^W$ with 18 5-year age groups | 36.3 | 21.2 | 1.71 |
| Cum. rate < 75 y (per 1000) –"– | 46.0 | 26.0 | 1.77 |

Standard population: B Birmingham 1985, C Cali 1985, W World SP

NB: The ratios of age-adjusted rates appear less dependent on the choice of standard weights than on the coarseness of age grouping. 5-year age groups are preferred.

Survival

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Survival analysis

The **prognosis** of cancer patients: what is their chance to **survive** 1 year, 5 years etc. after diagnosis?

Survival analysis: In principle like incidence analysis but

- population at risk = patients with cancer,
- basic time variable = time since the date of diagnosis, at which the follow-up starts,
- outcome event of interest = death,
- measures and methods used somewhat different from those used in incidence analysis.

Follow-up of 8 out of 40 breast cancer patients (from IS, table 12.1., p. 264)

| No. | Age (y) | Sta- ge ^a | Date of diag- nosis | Date at end of follow -up | Vital status at end of follow -up | Cause of death ^c | Full years from diagn's up to end of follow -up | Days from diagn's up to end of follow -up |
|-----|------------|-------------------------|---------------------------|---------------------------------------|--|-----------------------------------|--|---|
| 1 | 39 | 1 | 01/02/89 | 23/10/92 | А | _ | 3 | 1360 |
| 3 | 56 | 2 | 16/04/89 | 05/09/89 | D | BC | 0 | 142 |
| 5 | 62 | 2 | 12/06/89 | 28/12/95 | Α | - | 6 | 2390 |
| 15 | 60 | 2 | 03/08/90 | 27/11/94 | A | - | 4 | 1577 |
| 22 | 64 | 2 | 17/02/91 | 06/09/94 | D | 0 | 3 | 1297 |
| 25 | 42 | 2 | 20/06/91 | 15/03/92 | D | BC | 0 | 269 |
| 30 | 77 | 1 | 05/05/92 | 10/05/95 | Α | - | 3 | 1100 |
| 37 | 45 | 1 | 11/05/93 | 07/02/94 | D | BC | 0 | 272 |

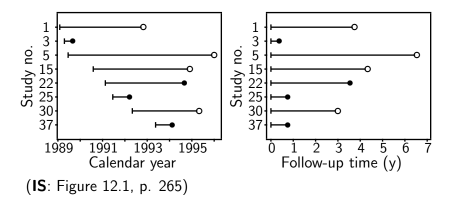
 a 1 = absence of regional lymph node involment and metastases

2 = involvment of regional lymph node and/or presence of metastases

^b A = alive; D = dead; ^c BC = breast cancer; O = other causes

Follow-up of breast cancer patients (cont'd)

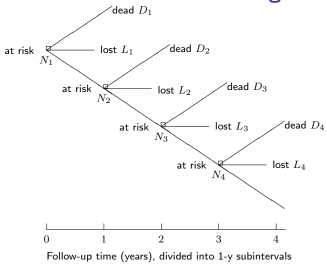
entry = diagnosis; • exit = death; \circ exit = censoring



Life table or "actuarial" method

- (1) Divide the follow-up time into subintervals k = 1, ..., K; usually each with 1 year width.
- (2) Tabulate from original data for each interval
 - $N_k =$ size of the **risk set**, *i.e.* the no. of subjects still alive and under follow-up at the start of interval,
 - $D_k =$ no. of **cases**, *i.e.* deaths observed in the interval,
 - $L_k =$ no. of **losses**, *i.e.* individuals **censored** during the interval before being observed to die.

Life table items in a tree diagram



 N_k = population at risk at the start of the kth subinterval D_k = no. of deaths, L_k = no. of losses or censorings in interval k

Life table items for breast ca. patients

(IS: Table 12.2., p. 273, first 4 columns)

| Inter- val | Years since | No. at start of | No. of deaths | No. of losses |
|---------------|----------------|--------------------|------------------|------------------|
| | diagnosis | interval | | |
| (k) | | (N_k) | (D_k) | (L_k) |
| 1 | 0 - < 1 | 40 | 7 | 0 |
| 2 | 1 - < 2 | 33 | 3 | 6 |
| 3 | 2 – < 3 | 24 | 4 | 3 |
| 4 | 3 - < 4 | 17 | 4 | 4 |
| 5 | 4 - < 5 | 9 | 2 | 3 |
| 6 | 5 - < 6 | 4 | 1 | 2 |
| 7 | 6-<7 | 1 | 0 | 1 |
| Total | | | 21 | 19 |

Life table calculations (cont'd)

(3) Calculate and tabulate for each interval

 $N'_k = N_k - L_k/2 =$ corrected size of the risk set, or "effective denominator" at start of the interval,

 $q_k = D_k/N'_k$ = estimated conditional probability of dying during the interval given survival up to its start,

 $p_k = 1 - q_k = \text{conditional survival proportion over the int'l,}$

 $S_k = p_1 \times \cdots \times p_k =$ cumulative survival proportion from date of diagnosis until the end of the *k*th interval

= estimate of **survival probability** up to this time point.

Follow-up of breast ca. patients (cont'd)

Actuarial life table completed (IS, table 12.2, p. 273)

| Inter- val | Years since dia- gnosis | No. at start of in- terval | No. of deaths | No. of losses | Effec- tive deno- minator | Cond'l prop'n of deaths during int'l | Survival prop'n over int'l | Cumul. survival; est'd survival prob'ty |
|---------------|----------------------------------|--|---------------------|---------------------|------------------------------------|--|-------------------------------------|---|
| (k) | | (N_k) | (D_k) | (L_k) | (N'_k) | (q_k) | (p_k) | (S_k) |
| 1 | 0- < 1 | 40 | 7 | 0 | 40.0 | 0.175 | 0.825 | 0.825 |
| 2 | 1 - < 2 | 33 | 3 | 6 | 30.0 | 0.100 | 0.900 | 0.743 |
| 3 | 2 - < 3 | 24 | 4 | 3 | 22.5 | 0.178 | 0.822 | 0.610 |
| 4 | 3 - < 4 | 17 | 4 | 4 | 15.0 | 0.267 | 0.733 | 0.447 |
| 5 | 4 - < 5 | 9 | 2 | 3 | 7.5 | 0.267 | 0.733 | 0.328 |
| 6 | 5 - < 6 | 4 | 1 | 2 | 3.0 | 0.333 | 0.667 | 0.219 |
| 7 | 6 - < 7 | 1 | 0 | 1 | 0.5 | 0.0 | 1.0 | 0.219 |

1-year survival probability is thus estimated 82.5% and 5-year probability 32.8%.

Comparison to previous measures and methods

Complement of survival proportion $Q_k = 1 - S_k$ is actually incidence proportion of deaths. It estimates cumulative risk of death from start of follow-up till end of kth interval.

"Actuarial" indidence rate in the kth interval:

$$I_k = \frac{\text{number of cases } (D_k)}{\text{approximate person-time}}$$

where the person-time is approximated by

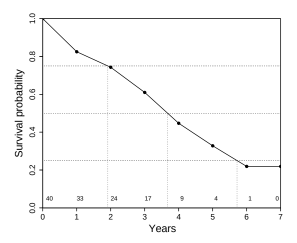
$$\left[N_k - \frac{1}{2}(D_k + L_k)\right] imes ext{ length of interval}$$

Survival curve and other measures

Line diagram of survival proportions through interval endpoints provides graphical estimates of interesting parameters of the survival time distribution, *e.g.*:

- median and quartiles: time points at which the curve crosses the 50%, 75%, and 25% levels
- ▶ mean residual lifetime: area under the curve, given that it decreases all the way down to the 0% level.
- **NB.** Often the curve ends at higher level than 0%, in which case some measures cannot be calculated.

Survical curve of breast ca. patients (IS: Fig 12.8)



Numbers above *x*-axis show the size of population at risk.

Survival

Cause-specific and relative survival

Cause-specific survival analysis:

- outcome event: death from the disease C itself that defines study population
- deaths from other causes \rightarrow losses.
- problem: ambiguity in cause of death.

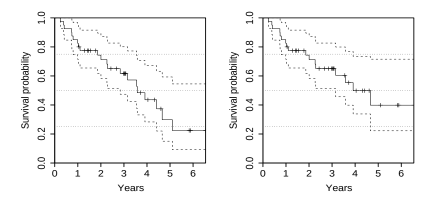
Relative survival: $S_k^{\text{rel}} = S_k^{\text{obs}}/S_k^{\text{exp}}$, i.e. ratio of

- observed survival proportion S_k^{obs}
 in the study population, and
- expected survival proportion S_k^{exp}
 based on age-specific mortalities in the reference (national) population. (See SIR!)

Breast Cancer patients (cont'd)

Overall and cause-specific (death from breast ca.) survival (**IS**: Fig 12.9 & 12.12, p. 271-3)

Kaplan-Meier curves – alternative to "actuarial":



Conclusion

Measures of Disease Occurrence

Bendix Carstensen & Esa Läärä

Nordic Summerschool of Cancer Epidemiology 15-26 August 2011 Copenhagen www.biostat.ku.dk/~bxc/NSCE

Conclusion

Measuring and comparing disease frequencies

- not a trivial task but
- demands expert skills in epidemiologic methods.

Major challenges:

- obtain the right denominator for each numerator,
- valid calculation of person-years,
- appropriate treatment of time and its various aspects,
- removal of confounding from comparisons.