

# Modern Demographic Methods in Epidemiology with R

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## R Packages

- ▶ The capabilities of R can be extended using “packages”.
- ▶ Distributed over the Internet *via* **CRAN**: (the **C**omprehensive **R** **A**rchive **N**etwork) and can be downloaded directly from an R session.
- ▶ There is an R package developed during the annual course on “Statistical Practice in Epidemiology using R, called “Epi”.
- ▶ Contains special functions for epidemiologists and some data sets that .
- ▶ There are 5,825 other user contributed packages on CRAN.

Introducing R (Data)

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## Introducing R

Modern Demographic  
Methods in Epidemiology  
with R  
23 November 2015  
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Data

## Objects and functions

R allows you to build powerful procedures from simple building blocks. These building blocks are **objects** and **functions**.

- ▶ All data in R is represented by **objects**, for example:
  - ▶ A dataset (called data frame in R)
  - ▶ A vector of numbers
  - ▶ The result of fitting a model to data
- ▶ You, the user, call **functions**
- ▶ Functions act on objects to create **new objects**:
- ▶ Using `glm` on a dataframe (an object) produces a fitted model (another object).

Introducing R (Data)

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## The best way to learn R

- ▶ The best way to learn R is to use it!
- ▶ This is a very short introduction before you sit down in front of a computer.
- ▶ R is a little different from other packages for statistical analysis.
- ▶ These differences make R very powerful, but for a new user they can sometimes be confusing.
- ▶ Our first job is to help you up the initial learning curve so that you can be comfortable with R.

Introducing R (Data)

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## Because all is functions. . .

- ▶ You will always (almost) use parentheses:  

```
> res <- FUN( x, y )
```
- ▶ . . . which is pronounced
- ▶ `res` **gets** (“<-”) **FUN of** `x,y` (“(x,y)”)

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## Nothing is lost or hidden

- ▶ Statistical software provides “canned” procedures to address common statistical problems.
- ▶ Canned procedures are useful for routine analysis, but they are also limiting.
  - ▶ You can only do what the programmer lets you do.
- ▶ In R, the results of statistical calculations are always accessible.
  - ▶ You can use them for further calculations.
  - ▶ You can always see how the calculations were done.

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## Vectors

One of the simplest objects in R is a sequence of numbers, called a **vector**.

You can create a vector in R with the collection (`c`) function:

```
> c(1,3,2)
[1] 1 3 2
```

You can save the results of any calculation using the left arrow:

```
> x <- c(1,3,2)
> x
[1] 1 3 2
```

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## The workspace

- ▶ Every time you use `<-`, you create a new object in the **workspace** (or overwrite an old one).
- ▶ A list of objects in the workspace can be seen with the `objects` function (synonym: `ls()`):  

```
> objects()
[1] "a" "aa" "acz2" "alpha" "b"
[6] "bar" "bb" "bdendo" "beta" "cc"
[11] "Col"
```
- ▶ In `Epi` is a function `lls()` that gives a bit more information on the objects.
- ▶ The workspace is held entirely in (volatile) computer memory and will be lost at the end of the session unless you explicitly save it.

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## Rectangular Data

Rectangular data sets are common to most statistical packages

"id"	"visit"	"time"	"status"
1	1	0.0	0
1	2	1.5	0
2	1	0.0	0
2	2	1.1	0
2	3	2.3	1

Columns represent variables.  
Rows represent individual records.

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## Working Directory

Every R session has a **current working directory**, which is the location on the hard disk where files are saved, and the default location from which files are read into R.

- ▶ `getwd()` Prints the current working directory
- ▶ `setwd("c:/Users/Martyn/Project")` sets the current working directory.
- ▶ You may also use a Graphical User Interface (GUI) to change directory.

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## The world is not a rectangle!

- ▶ Most statistical packages used by epidemiologists assume that **all data** can be represented as a rectangular data set.
- ▶ R allows a much richer set of data structures, represented by *objects* of different *classes*.
- ▶ Rectangular data sets are just one type of object that may be in your workspace. This class of object is called a *data frame*.

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## Ending an R session

- ▶ To end an R session, call the `quit()` function
  - ▶ Every time you want to do something in R, you call a function.
- ▶ You will be asked "Save workspace image?"
  - `Yes` saves the workspace to the file `".RData"` in your current working directory. It will be automatically loaded into R the next time you start an R session.
  - `No` does not save the workspace.
  - `Cancel` continues the current R session without saving anything.
- ▶ It is recommended you just say "No".

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## Data Frames

Each column of a data frame is a variable.

Variables may be of different types:

- ▶ **vectors:**
  - ▶ **numeric:** `c(1,2,3)`
  - ▶ **character:**  
`c("John","Paul","George","Ringo")`
  - ▶ **logical:** `c(FALSE,FALSE,TRUE)`
- ▶ **factors:**  
`factor(c("low","medium","high","low","low"))`

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## Always start with a clean workspace

Keeping objects in your workspace from one session to another can be dangerous:

- ▶ You forget how they were made.
- ▶ You cannot easily recreate them if your data changes.
- ▶ They may not even be from the same project

It is almost always best to start with an empty workspace and use a script file to create the objects you need from scratch.

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## Building your own data frame

Data frames can be constructed from a list of vectors

```
> mydata <- data.frame(x=c(3,6,7),f=c("a","b","a"))
> mydata
  x f
1 3 a
2 6 b
3 7 a
```

Character vectors are automatically converted to factors.

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## Inspecting data frames

Most data frames are too large to inspect by printing them to the screen, so use:

- ▶ `names` returns a vector of variable names.
  - ▶ You can use `sort(names(x))` to get them in alphabetical order.
- ▶ `head` prints the first few lines, and `tail...`
- ▶ `str` prints a brief overview of the **structure** of the data frame. Can be used on any object.
- ▶ `summary` prints a more comprehensive summary
  - ▶ Quantiles for numeric variables
  - ▶ Tables for factors

## Some useful arguments to `read.table`

- ▶ `header = TRUE` if first line contains variable names
- ▶ `sep=","` if values are comma-separated instead of being space-delimited.
- ▶ `as.is = TRUE` to stop strings being converted to factors
- ▶ `na.strings = "99"` to denote that 99 means "missing". Default values are:
  - ▶ `NA` "Not Available"
  - ▶ `NaN` "Not a Number"
- ▶ For comma-separated files there is `coderead.csv`

## Extracting values from a data frame

Use square brackets to take **subsets** of a data frame

- ▶ `mydata[1,2]`. The value in row 1, column 2.
- ▶ `mydata[1,]`. The whole of the first row.
- ▶ `mydata[,2]`. The whole of the second column.

You can also extract a column from a data frame by name:

- ▶ `mydata$age`. The column, or variable, named "age"
- ▶ `mydata[, "age"]`. The same.

## Reading Binary Data

- ▶ R can read in data in binary (non-text) format from other statistical systems using the foreign extension package.
- ▶ R is an open source project, and relies on the format for binary files to be well-documented.
- ▶ Example: SAS XPORT format has been adopted as a data exchange standard by the US Food and Drug Administration. SAS CPORT format remains a proprietary format.

## Importing data

- ▶ R has good facilities for importing data from other applications:
  - ▶ `read.dta` for reading Stata datasets.
  - ▶ `read.spss` for reading SPSS datasets.
  - ▶ `read.xport` and `read.ssd` for reading SAS-datasets.

## Some functions in the foreign package

- ▶ `read.dta` for Stata (also `write.dta`)
- ▶ `read.xport` for SAS XPORT format (not CPORT)
- ▶ `read.epiinfo` for EPIINFO
- ▶ `read.mtp` for MiniTab Portable Worksheet
- ▶ `read.spss` for SPSS

See the "R Data Import/Export manual" for more details. `RShowDoc("R-data")`

## Reading Text Files

The function `read.table` reads data from a text file and returns a data frame.

- ▶ `mydata <- read.table("myfile")`
- ▶ `myfile` could be
  - ▶ A file in the **current working directory**: `fem.dat`
  - ▶ A path to a file: `c:/rex/fem.dat`
  - ▶ A URL:  
`http://BendixCarstensen.com/AdvCoh/Scot-2014/data/bogus.txt`
- ▶ Note: `myfile` must be enclosed in quotes.

`write.table` does the opposite.

R uses a forward slash / for file paths. If you want to use backslash, you have to double it:

## Accessing databases systems

Microsoft **Access**:

```
> library(RODBC)
> ch <- odbcConnectAccess("../data/theData.mdb")
> bd <- sqlFetch(ch, "aTable" )
```

Microsoft **Excel**:

```
> library(RODBC)
> cnc <- odbcConnectExcel(paste("../theXel.xls", sep=""))
> sht <- sqlFetch(cnc, "theSheet" )
> close(cnc)
```

Other databases

```
> ?odbcConnect
```

## Summary - data

- ▶ You can use a data frame to organize your variables
- ▶ You can extract variables from a data frame using `$`.
- ▶ You can extract variables and observation using indexing `[,]`
- ▶ You can read in data using
  - ▶ `read.table`
  - ▶ tailored function from the `foreign` package
  - ▶ database interface from the `RODBC` package

## Objects

- ▶ The simplest object type is *vector*
- ▶ Modes: numeric, integer, character, generic (list)
- ▶ Operations are vectorized: you can add entire vectors with `a + b`
- ▶ Recycling of objects: If the lengths don't match, the shorter vector is reused

## Summary - when it goes wrong

When something is fishy with an object `obj`, try to find out what you (accidentally) got, by using:

```
> lls()
> str( obj )
> dim( obj )
> length( obj )
> names( obj )
> head( obj )
> class( obj )
> mode( obj )
```

## R expressions

```
x <- rnorm(10, mean=20, sd=5)
m <- mean(x)
sum((x - m)^2)
```

- ▶ Object **names**
- ▶ Explicit **constants**
- ▶ Arithmetic **operators**
- ▶ **Function calls**
- ▶ **Assignment** of results to names

## R language

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lang

## Function calls

Lots of things you do with R involve calling functions.  
For instance

```
mean(x, na.rm=TRUE)
```

The important parts of this are

- ▶ The **name** of the function
- ▶ **Arguments**: input to the function
- ▶ Sometimes, we have **named arguments**

## Language

- ▶ R is a programming language – also on the command line
- ▶ (This means that there are *syntax rules*)
- ▶ Print an object by typing its name
- ▶ Evaluate an expression by entering it on the command line
- ▶ Call a function, giving the arguments in parentheses – possibly empty
- ▶ Notice `ls` vs. `ls()`

## Function arguments

```
rnorm(10, mean=m, sd=s)
hist(x, main="My histogram")
mean(log(x + 1))
```

Items which may appear as arguments:

- ▶ **Names** of an R objects
- ▶ Explicit **constants**
- ▶ **Return values** from another function call or expression
- ▶ Some arguments have their *default values*.
- ▶ Use `help(function)` or `args(function)` to see the **arguments** (and their order and default values) that can be given to any function.

## Creating simple functions

```
logit <- function(p) log(p/(1-p))
logit(0.5)
```

```
simpsum <-
function(x, dec=5)
{
  # produces mean and SD of a variable
  # default value for dec is 5
  round(c(mean=mean(x),sd=sd(x)),dec)
}
```

```
x <- rnorm(100)
simpsum(x)
simpsum(x,2)
```

R language (1ang)

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## The workspace

- ▶ The *global environment* contains R objects created on the command line.
- ▶ There is an additional *search path* of loaded packages and attached data frames.
- ▶ When you request an object by name, R looks first in the global environment, and if it doesn't find it there, it continues along the search path.
- ▶ The search path is maintained by `library()`, `attach()`, and `detach()`
- ▶ List the search path by `search()`
- ▶ Notice that objects in the global environment may mask objects in packages and attached data frames

R language (1ang)

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## Indexing

- ▶ R has several useful indexing mechanisms:
- ▶ `a[5]` single element
- ▶ `a[5:7]` several elements
- ▶ `a[-6]` all except the 6th
- ▶ `a[c(1,1,2,1,2)]` some elements repeated
- ▶ `a[b>200]` logical index
- ▶ `a[well]` indexing by name

R language (1ang)

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## Data manipulation and `with`

```
bmi <- with(stud, weight/(height/100)^2)
```

uses variables `weight` and `height` in the data frame `stud` (not the variables with the same name in the workspace), but creates the variable `bmi` in the global environment (not in the data frame).

To create a new variable in the data frame, you can use:

```
stud$bmi <- with( stud, weight/(height/100)^2 )
```

R language (1ang)

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## Lists

- ▶ Lists are vectors where the elements can have different types
- ▶ Functions often return lists
- ▶ `lst <- list(A=rnorm(5),B="hello",K=12)`
- ▶ Special indexing:
- ▶ `lst$A`
- ▶ `lst[1:2]` a list with first two first elements (`A` and `B` — NB: single brackets)
- ▶ `lst[1]` a list of length 1 which is the first element (codeA — NB: single brackets)
- ▶ `lst[[1]]` first element (NB: double brackets) — a vector of length 5.

R language (1ang)

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## Constructors

- ▶ Matrices and arrays, constructed by the (surprise) `matrix` and `array` functions.
- ▶ You can extract and set names with `names(x)`; for matrices and data frames also `colnames(x)` and `rownames(x)`
- ▶ You can also construct a matrix from its columns using `cbind`, whereas joining two matrices with equal no of columns (with the same column names) can be done using `rbind`.

R language (1ang)

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## Classes, generic functions

- ▶ R objects have *classes*
- ▶ Functions can behave differently depending on the class of an object
- ▶ E.g. `summary(x)` or `print(x)` does different things if `x` is numeric, a factor, or a linear model fit

R language (1ang)

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## Factors (class variables)

- ▶ Factors are used to describe groupings.
- ▶ Basically, these are just integer codes plus a set of names for the *levels*
- ▶ They have class "`factor`" making them (a) print nicely and (b) maintain consistency
- ▶ A factor can also be *ordered* (class "`ordered`"), signifying that there is a natural sort order on the levels
- ▶ In model specifications, factors play a fundamental role by indicating that a variable should be treated as a classification rather than as a quantitative variable (similar to a `CLASS` statement in SAS)

R language (1ang)

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## The factor function

- ▶ This is typically used when `read.table` gets it wrong,
- ▶ e.g. group codes read as numeric
- ▶ or read as factors, but with levels in the wrong order (e.g. `c("rare", "medium", "well-done")` sorted alphabetically.)
- ▶ Notice that there is a slightly confusing use of `levels` and `labels` arguments:
  - ▶ `levels` are the value codes *on input*
  - ▶ `labels` are the value codes *on output* (and becomes the levels of the resulting factor)
  - ▶ The levels of a factor is shown by the `levels()` function.

R language (lang)

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## Basic graphics

Similarly, the `plot(x,y)` produces:

- ▶ a scatter plot of `x` is a numeric variable
- ▶ a bar plot of category frequencies, when `x` is a factor variable

R language (lang)

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## Working with Dates

- ▶ Dates are usually read as character or factor variables
  - ▶ Use the `as.Date` function to convert them to objects of class "Date"
  - ▶ If data are not in the default format (`yyyy-mm-dd`) you need to supply a format specification
- ```
> as.Date("11/3-1959",format="%d/%m-%Y")
[1] "1959-03-11"
```

R language (lang)

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## Basic graphics

Examples:

```
x <- c(0,1,2,1,2,2,1,1,3,3)
plot(x)
plot(factor(x))
plot(ts(x)) # ts() defines x as time series
y <- c(0,1,3,1,2,1,0,1,4,3)
plot(x,y)
plot(factor(x),y)
```

R language (lang)

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## Working with Dates

- ▶ Computing the differences between `Date` objects gives an object of class "difftime", which is number of days between the two dates:
- ```
> as.numeric(as.Date("2007-5-25")-
             as.Date("1959-3-11"),"days")
[1] 17607
```
- ▶ In the `Epi` package is a function that converts dates to calendar years with decimals:
- ```
> as.Date("1952-07-14")
[1] "1952-07-14"
> cal.yr( as.Date("1952-07-14") )
[1] 1952.533
attr(,"class")
[1] "cal.yr" "numeric"
```

R language (lang)

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## Basic graphics

More simple plots:

- ▶ `hist(x)` produces a histogram
- ▶ `barplot(x)` produces a bar plot (useful when `x` contains counts – often one uses `barplot(table(x))`)
- ▶ `boxplot(y ~ x)` produces a box plot of `y` by levels of a (factor) variable `x`.

R language (lang)

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## Basic graphics

The `plot()` function is a generic function, producing different plots for different types of arguments. For instance, `plot(x)` produces:

- ▶ a plot of observation index against the observations, when `x` is a numeric variable
- ▶ a bar plot of category frequencies, when `x` is a factor variable
- ▶ a time series plot (interconnected observations) when `x` is a time series
- ▶ a set of diagnostic plots, when `x` is a fitted regression model
- ▶ ...

R language (lang)

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## Rates and Survival

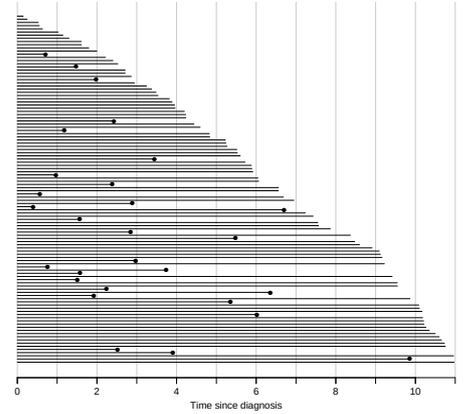
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surv-rate

## Survival data

Persons enter the study at some date.  
Persons exit at a later date, either dead or alive.  
Observation:  
Actual time span to death ("event")  
or  
Some time alive ("at least this long")

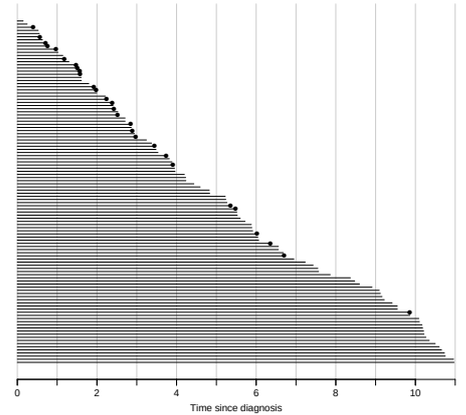
Timescale  
changed to  
"Time since  
diagnosis".



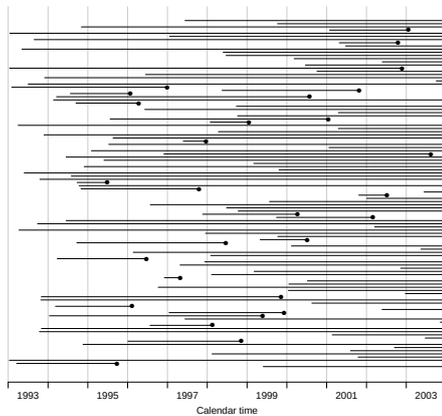
## Examples of time-to-event measurements

- ▶ Time from diagnosis of cancer to death.
- ▶ Time from randomisation to death in a cancer clinical trial
- ▶ Time from HIV infection to AIDS.
- ▶ Time from marriage to 1st child birth.
- ▶ Time from marriage to divorce.
- ▶ Time to re-offending after being released from jail

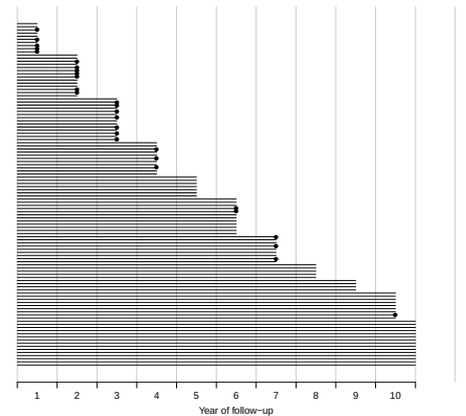
Patients  
ordered by  
survival  
time.



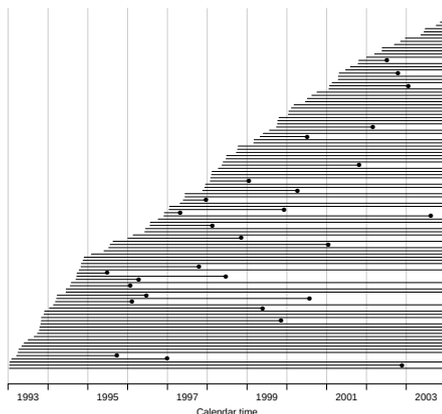
Each line a  
person  
  
Each blob a  
death  
  
Study ended  
at 31 Dec.  
2003



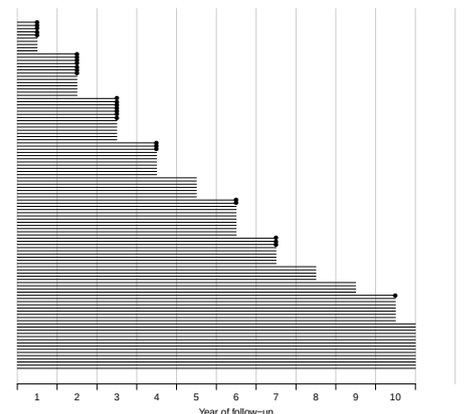
Survival  
times  
grouped into  
bands of  
survival.



Ordered by  
date of entry  
  
Most likely  
the order in  
your  
database.



Patients  
ordered by  
survival  
status within  
each band.



## Survival after Cervix cancer

| Year | Stage I |   |    | Stage II |    |    |
|------|---------|---|----|----------|----|----|
|      | N       | D | L  | N        | D  | L  |
| 1    | 110     | 5 | 5  | 234      | 24 | 3  |
| 2    | 100     | 7 | 7  | 207      | 27 | 11 |
| 3    | 86      | 7 | 7  | 169      | 31 | 9  |
| 4    | 72      | 3 | 8  | 129      | 17 | 7  |
| 5    | 61      | 0 | 7  | 105      | 7  | 13 |
| 6    | 54      | 2 | 10 | 85       | 6  | 6  |
| 7    | 42      | 3 | 6  | 73       | 5  | 6  |
| 8    | 33      | 0 | 5  | 62       | 3  | 10 |
| 9    | 28      | 0 | 4  | 49       | 2  | 13 |
| 10   | 24      | 1 | 8  | 34       | 4  | 6  |

Estimated risk in year 1 for Stage I women is  $5/107.5 = 0.0465$

Estimated 1 year survival is  $1 - 0.0465 = 0.9535$

Life-table estimator.

## Rate and survival

$$S(t) = \exp\left(-\int_0^t \lambda(s) ds\right) \quad \lambda(t) = \frac{S'(t)}{S(t)}$$

Survival is a *cumulative* measure, the rate is an *instantaneous* measure.

**Note:** A cumulative measure requires an origin!  
... it is always survival **since** some timepoint.

## Survival function

Persons enter at time 0:

Date of birth, date of randomization, date of diagnosis.

How long do they survive?

Survival time  $T$  — a stochastic variable.

Distribution is characterized by the survival function:

$$\begin{aligned} S(t) &= P\{\text{survival at least till } t\} \\ &= P\{T > t\} = 1 - P\{T \leq t\} = 1 - F(t) \end{aligned}$$

$F(t)$  is the cumulative risk of death before time  $t$ .

## Observed survival and rate

- **Survival studies:** Observation of (right censored) survival time:

$$X = \min(T, Z), \quad \delta = 1\{X = T\}$$

— sometimes conditional on  $T > t_0$   
(left truncation, delayed entry).

- **Epidemiological studies:**  
Observation of (components of) a rate:

$$D/Y$$

$D$ : no. events,  $Y$  no of person-years, in a prespecified time-frame.

## Intensity or rate

$$P\{\text{event in } (t, t+h] \mid \text{alive at } t\} / h$$

$$= \frac{F(t+h) - F(t)}{S(t) \times h}$$

$$= -\frac{S(t+h) - S(t)}{S(t)h} \xrightarrow{h \rightarrow 0} -\frac{d \log S(t)}{dt}$$

$$= \lambda(t)$$

This is the **intensity** or **hazard function** for the distribution. Characterizes the survival distribution as does  $f$  or  $F$ .

Theoretical counterpart of a **rate**.

## Empirical rates for individuals

- At the *individual* level we introduce the **empirical rate**:  $(d, y)$ ,  
— number of events ( $d \in \{0, 1\}$ ) during  $y$  risk time.
- A person contributes several observations of  $(d, y)$ , with associated covariate values.
- Empirical rates are **responses** in survival analysis.
- The timescale  $t$  is a **covariate** — varies within each individual:  
 $t$ : age, time since diagnosis, calendar time.
- Don't confuse with  $y$  — difference between two points on **any** timescale we may choose.

## Relationships

$$-\frac{d \log S(t)}{dt} = \lambda(t)$$

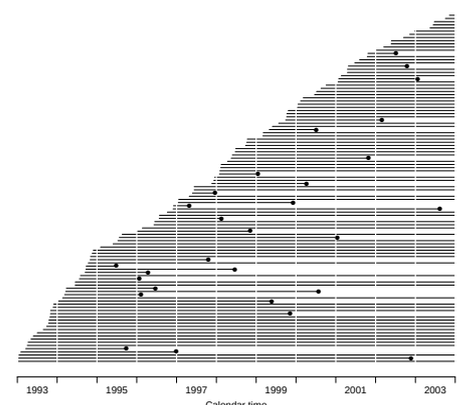
⇕

$$S(t) = \exp\left(-\int_0^t \lambda(u) du\right) = \exp(-\Lambda(t))$$

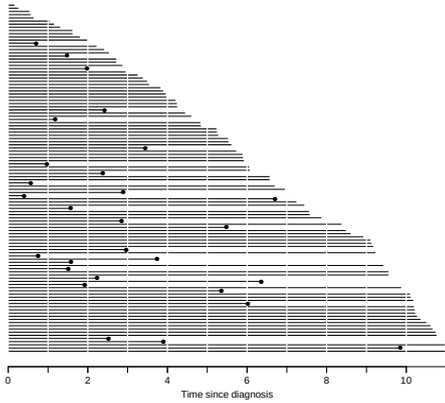
$\Lambda(t) = \int_0^t \lambda(s) ds$  is called the **integrated intensity**. **Not** an intensity, it is dimensionless.

$$\lambda(t) = -\frac{d \log(S(t))}{dt} = -\frac{S'(t)}{S(t)} = \frac{F'(t)}{1 - F(t)} = \frac{f(t)}{S(t)}$$

Empirical rates by calendar time.



Empirical rates by time since diagnosis.



## Likelihood for follow-up of many persons

Adding empirical rates over the follow-up of persons:

$$D = \sum d \quad Y = \sum y \Rightarrow D \log(\lambda) - \lambda Y$$

- ▶ Persons are assumed independent
- ▶ Contribution from the same person are **conditionally** independent, hence give separate contributions to the log-likelihood.
- ▶ Therefore equivalent to likelihood for independent Poisson variates
- ▶ No need to correct for dependent observations; the likelihood is a product.

## Statistical inference: Likelihood

Two things needed:

- ▶ **Data** — what did we actually observe  
Follow-up for each person:  
Entry time, exit time, exit status, covariates
- ▶ **Model** — how was data generated  
Rates as a function of time:  
Probability machinery that generated data

**Likelihood** is the probability of observing the **data**, assuming the **model** is correct.

**Maximum likelihood** estimation is choosing **parameters** of the model that makes the likelihood maximal.

## Likelihood

Probability of the data and the parameter:

Assuming the rate (intensity) is constant,  $\lambda$ , the probability of observing 7 deaths in the course of 500 person-years:

$$\begin{aligned} P\{D = 7, Y = 500 | \lambda\} &= \lambda^D e^{-\lambda Y} \times K \\ &= \lambda^7 e^{-\lambda 500} \times K \\ &= L(\lambda | \text{data}) \end{aligned}$$

Best guess of  $\lambda$  is where this function is as large as possible.

Confidence interval is where it is not too far from the maximum

## Likelihood from one person

The likelihood from several empirical rates from one individual is a product of conditional probabilities:

$$\begin{aligned} P\{\text{event at } t_4 | t_0\} &= P\{\text{survive}(t_0, t_1) | \text{alive at } t_0\} \times \\ &P\{\text{survive}(t_1, t_2) | \text{alive at } t_1\} \times \\ &P\{\text{survive}(t_2, t_3) | \text{alive at } t_2\} \times \\ &P\{\text{event at } t_4 | \text{alive at } t_3\} \end{aligned}$$

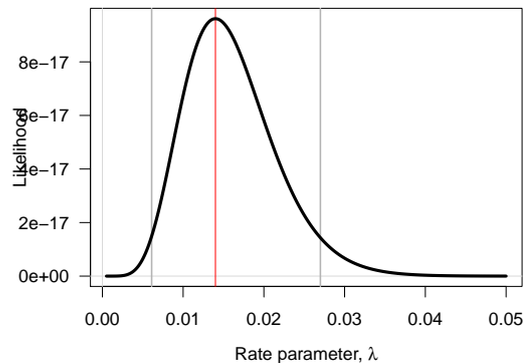
Log-likelihood from one individual is a sum of terms.

Each term refers to one empirical rate ( $d, y$ )

—  $y = t_i - t_{i-1}$  and mostly  $d = 0$ .

$t_i$  is the timescale (covariate).

## Likelihood function



## Poisson likelihood

The likelihood contributions from follow-up of **one** individual:

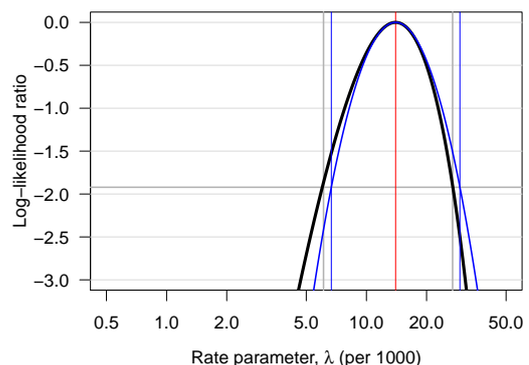
$$d_t \log(\lambda(t)) - \lambda(t)y_t, \quad t = t_1, \dots, t_n$$

is also the log-likelihood from several independent Poisson observations with mean  $\lambda(t)y_t$ , i.e. log-mean  $\log(\lambda(t)) + \log(y_t)$

Analysis of the rates, ( $\lambda$ ) can be based on a Poisson model with log-link applied to empirical rates where:

- ▶  $d$  is the response variable.
- ▶  $\log(\lambda)$  is modelled by covariates
- ▶  $\log(y)$  is the offset variable.

## Likelihood function



## Confidence interval for a rate

A 95% confidence interval for the log of a rate is:

$$\hat{\theta} \pm 1.96/\sqrt{D} = \log(\lambda) \pm 1.96/\sqrt{D}$$

Take the exponential to get the confidence interval for the rate:

$$\lambda \div \underbrace{\exp(1.96/\sqrt{D})}_{\text{error factor, erf}}$$

## Example using R

Poisson likelihood, for one rate, based on 17 events in 843.7 PY:

```
library( Epi )
D <- 17 ; Y <- 843.7
m1 <- glm( D ~ 1, offset=log(Y/1000), family=poisson)
ci.exp( m1 )
      exp(Est.)    2.5%    97.5%
(Intercept) 20.14934 12.52605 32.41213
```

Poisson likelihood, two rates, or one rate and RR:

```
D <- c(17,28) ; Y <- c(843.7,632.3) ; gg <- factor(0:1)
m2 <- glm( D ~ gg, offset=log(Y/1000), family=poisson)
ci.exp( m2 )
      exp(Est.)    2.5%    97.5%
(Intercept) 20.149342 12.526051 32.412130
gg1         2.197728  1.202971  4.015068
```

## Example

Suppose we have 17 deaths during 843.6 years of follow-up.

The rate is computed as:

$$\hat{\lambda} = D/Y = 17/843.7 = 0.0201 = 20.1 \text{ per } 1000 \text{ years}$$

The confidence interval is computed as:

$$\hat{\lambda} \div \text{erf} = 20.1 \div \exp(1.96/\sqrt{D}) = (12.5, 32.4)$$

per 1000 person-years.

## Example using R

Poisson likelihood, two rates, or one rate and RR:

```
D <- c(17,28) ; Y <- c(843.7,632.3) ; gg <- factor(0:1)
m2 <- glm( D ~ gg, offset=log(Y/1000), family=poisson)
ci.exp( m2 )
      exp(Est.)    2.5%    97.5%
(Intercept) 20.149342 12.526051 32.412130
gg1         2.197728  1.202971  4.015068
m3 <- glm( D ~ gg - 1, offset=log(Y/1000), family=poisson)
ci.exp( m3 )
      exp(Est.)    2.5%    97.5%
gg0 20.14934 12.52605 32.41213
gg1 44.28278 30.57545 64.13525
```

You do it!

## Ratio of two rates

If we have observations two rates  $\lambda_1$  and  $\lambda_0$ , based on  $(D_1, Y_1)$  and  $(D_0, Y_0)$ , the variance of the difference of the log-rates, the  $\log(\text{RR})$ , is:

$$\begin{aligned} \text{var}(\log(\text{RR})) &= \text{var}(\log(\lambda_1/\lambda_0)) \\ &= \text{var}(\log(\lambda_1)) + \text{var}(\log(\lambda_0)) \\ &= 1/D_1 + 1/D_0 \end{aligned}$$

As before a 95% c.i. for the RR is then:

$$\text{RR} \div \underbrace{\exp\left(1.96\sqrt{\frac{1}{D_1} + \frac{1}{D_0}}\right)}_{\text{error factor}}$$

## Representation of follow-up data

Modern Demographic  
Methods in Epidemiology  
with R

23 November 2015

University of Melbourne

<http://BendixCarstensen/AdvCoh/Melb-2015>

time-split

## Example

Suppose we in group 0 have 17 deaths during 843.6 years of follow-up in one group, and in group 1 have 28 deaths during 632.3 years.

The rate-ratio is computed as:

$$\begin{aligned} \text{RR} &= \hat{\lambda}_1/\hat{\lambda}_0 = (D_1/Y_1)/(D_0/Y_0) \\ &= (28/632.3)/(17/843.7) = 0.0443/0.0201 = 2.19 \end{aligned}$$

The 95% confidence interval is computed as:

$$\begin{aligned} \hat{\text{RR}} \div \text{erf} &= 2.198 \div \exp(1.96\sqrt{1/17 + 1/28}) \\ &= 2.198 \div 1.837 = (1.20, 4.02) \end{aligned}$$

## Follow-up and rates

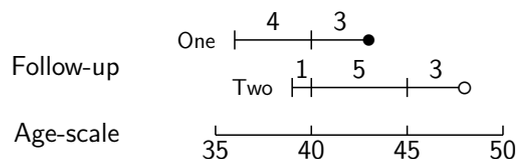
- ▶ Follow-up studies:
  - ▶  $D$  — events, deaths
  - ▶  $Y$  — person-years
  - ▶  $\lambda = D/Y$  rates
- ▶ Rates differ between persons.
- ▶ Rates differ **within** persons:
  - ▶ By age
  - ▶ By calendar time
  - ▶ By disease duration
  - ▶ ...
- ▶ Multiple timescales.
- ▶ Multiple states (little boxes — later)

## Stratification by age

If follow-up is rather short, age at entry is OK for age-stratification.

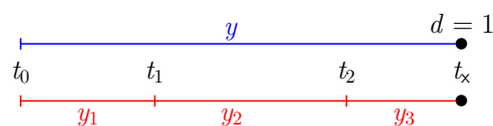
If follow-up is long, use stratification by categories of **current age**, both for:

No. of events,  $D$ , and Risk time,  $Y$ .



Representation of follow-up data (time-split)

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Probability

log-Likelihood

$P(\text{event at } t_x | \text{entry } t_0)$

$1 \log(\lambda) - \lambda y$

$= P(\text{surv } t_0 \rightarrow t_1 | \text{entry } t_0)$

$= 0 \log(\lambda) - \lambda y_1$

$\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1)$

$+ 0 \log(\lambda) - \lambda y_2$

$\times P(\text{event at } t_x | \text{entry } t_2)$

$+ 1 \log(\lambda) - \lambda y_3$

Representation of follow-up data (time-split)

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## Representation of follow-up data

A cohort or follow-up study records:

**Events** and **Risk time**.

The outcome is thus **bivariate**:  $(d, y)$

Follow-up **data** for each individual must therefore have (at least) three variables:

|                |       |                 |
|----------------|-------|-----------------|
| Date of entry  | entry | date variable   |
| Date of exit   | exit  | date variable   |
| Status at exit | fail  | indicator (0/1) |

Specific for each **type** of outcome.

Representation of follow-up data (time-split)

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## Dividing time into bands:

If we want to put  $D$  and  $Y$  into intervals on the timescale we must know:

**Origin:** The date where the time scale is 0:

- ▶ Age — 0 at date of birth
- ▶ Disease duration — 0 at date of diagnosis
- ▶ Occupation exposure — 0 at date of hire

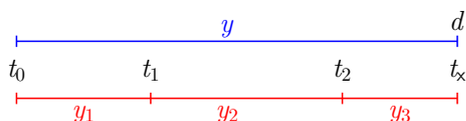
**Intervals:** How should it be subdivided:

- ▶ 1-year classes? 5-year classes?
- ▶ Equal length?

**Aim:** Separate rate in each interval

Representation of follow-up data (time-split)

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Probability

log-Likelihood

$P(d \text{ at } t_x | \text{entry } t_0)$

$d \log(\lambda) - \lambda y$

$= P(\text{surv } t_0 \rightarrow t_1 | \text{entry } t_0)$

$= 0 \log(\lambda) - \lambda y_1$

$\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1)$

$+ 0 \log(\lambda) - \lambda y_2$

$\times P(d \text{ at } t_x | \text{entry } t_2)$

$+ d \log(\lambda) - \lambda y_3$

Representation of follow-up data (time-split)

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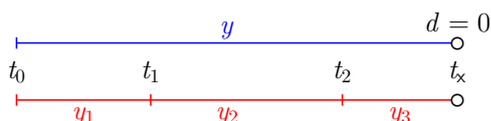
## Example: cohort with 3 persons:

| Id | Bdate      | Entry      | Exit       | St |
|----|------------|------------|------------|----|
| 1  | 14/07/1952 | 04/08/1965 | 27/06/1997 | 1  |
| 2  | 01/04/1954 | 08/09/1972 | 23/05/1995 | 0  |
| 3  | 10/06/1987 | 23/12/1991 | 24/07/1998 | 1  |

- ▶ Age bands: 10-years intervals of current age.
- ▶ Split  $Y$  for every subject accordingly
- ▶ Treat each segment as a separate unit of observation.
- ▶ Keep track of exit status in each interval.

Representation of follow-up data (time-split)

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Probability

log-Likelihood

$P(\text{surv } t_0 \rightarrow t_x | \text{entry } t_0)$

$0 \log(\lambda) - \lambda y$

$= P(\text{surv } t_0 \rightarrow t_1 | \text{entry } t_0)$

$= 0 \log(\lambda) - \lambda y_1$

$\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1)$

$+ 0 \log(\lambda) - \lambda y_2$

$\times P(\text{surv } t_2 \rightarrow t_x | \text{entry } t_2)$

$+ 0 \log(\lambda) - \lambda y_3$

Representation of follow-up data (time-split)

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## Splitting the follow up

|                 | subj. 1 | subj. 2 | subj. 3 |
|-----------------|---------|---------|---------|
| Age at Entry:   | 13.06   | 18.44   | 4.54    |
| Age at eXit:    | 44.95   | 41.14   | 11.12   |
| Status at exit: | Dead    | Alive   | Dead    |
| <hr/>           |         |         |         |
| $Y$             | 31.89   | 22.70   | 6.58    |
| $D$             | 1       | 0       | 1       |

Representation of follow-up data (time-split)

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| Age    | subj. 1 |   | subj. 2 |   | subj. 3 |   | $\sum$ |   |
|--------|---------|---|---------|---|---------|---|--------|---|
|        | Y       | D | Y       | D | Y       | D | Y      | D |
| 0-     | 0.00    | 0 | 0.00    | 0 | 5.46    | 0 | 5.46   | 0 |
| 10-    | 6.94    | 0 | 1.56    | 0 | 1.12    | 1 | 8.62   | 1 |
| 20-    | 10.00   | 0 | 10.00   | 0 | 0.00    | 0 | 20.00  | 0 |
| 30-    | 10.00   | 0 | 10.00   | 0 | 0.00    | 0 | 20.00  | 0 |
| 40-    | 4.95    | 1 | 1.14    | 0 | 0.00    | 0 | 6.09   | 1 |
| $\sum$ | 31.89   | 1 | 22.70   | 0 | 6.58    | 1 | 60.17  | 2 |

Representation of follow-up data (time-split)

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## Follow-up data in Epi — Lexis objects

A follow-up study:

```
> round( th, 2 )
      id sex birthdat contrast injecdat volume exitdat ex
1     1  2  1916.61         1  1938.79     22 1976.79
2    640  2  1896.23         1  1945.77     20 1964.37
3   3425  1  1886.97         2  1955.18      0 1956.59
4   4017  2  1936.81         2  1957.61      0 1992.14
...
```

Timescales of interest:

- ▶ Age
- ▶ Calendar time
- ▶ Time since injection

Representation of follow-up data (time-split)

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## Splitting the follow-up

| id | Bdate      | Entry      | Exit       | St | risk    | int |
|----|------------|------------|------------|----|---------|-----|
| 1  | 14/07/1952 | 03/08/1965 | 14/07/1972 | 0  | 6.9432  | 10  |
| 1  | 14/07/1952 | 14/07/1972 | 14/07/1982 | 0  | 10.0000 | 20  |
| 1  | 14/07/1952 | 14/07/1982 | 14/07/1992 | 0  | 10.0000 | 30  |
| 1  | 14/07/1952 | 14/07/1992 | 27/06/1997 | 1  | 4.9528  | 40  |
| 2  | 01/04/1954 | 08/09/1972 | 01/04/1974 | 0  | 1.5606  | 10  |
| 2  | 01/04/1954 | 01/04/1974 | 31/03/1984 | 0  | 10.0000 | 20  |
| 2  | 01/04/1954 | 31/03/1984 | 01/04/1994 | 0  | 10.0000 | 30  |
| 2  | 01/04/1954 | 01/04/1994 | 23/05/1995 | 0  | 1.1417  | 40  |
| 3  | 10/06/1987 | 23/12/1991 | 09/06/1997 | 0  | 5.4634  | 0   |
| 3  | 10/06/1987 | 09/06/1997 | 24/07/1998 | 1  | 1.1211  | 10  |

Keeping track of calendar time too?

Representation of follow-up data (time-split)

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## Definition of Lexis object

```
> thL <- Lexis( entry = list( age = injecdat-birthdat,
+                             per = injecdat,
+                             tfi = 0 ),
+               exit = list( per = exitdat ),
+               exit.status = as.numeric(exitstat==1),
+               data = th )
```

**entry** is defined on **three** timescales,  
but **exit** is only defined on **one** timescale:  
**Follow-up time** is the same on all timescales:

`exitdat - injecdat`

Representation of follow-up data (time-split)

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## Timescales

- ▶ A timescale is a variable that varies **deterministically** *within* each person during follow-up:
  - ▶ Age
  - ▶ Calendar time
  - ▶ Time since treatment
  - ▶ Time since relapse
- ▶ All timescales advance at the same pace (1 year per year ...)
- ▶ Note: Cumulative exposure is **not** a timescale.

Representation of follow-up data (time-split)

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## The looks of a Lexis object

```
> thL[,1:9]
      age  per tfi lex.dur lex.Cst lex.Xst lex.id
1  22.18 1938.79  0  37.99      0      1      1
2  49.54 1945.77  0  18.59      0      1      2
3  68.20 1955.18  0   1.40      0      1      3
4  20.80 1957.61  0  34.52      0      0      4
...
```

```
> summary( thL )
Transitions:
To
From 0 1 Records: Events: Risk time: Persons:
    0 3 20         23      20      512.59      23
```

Representation of follow-up data (time-split)

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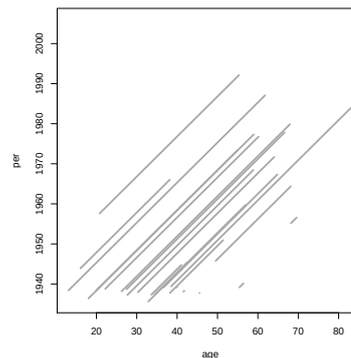
## Follow-up on several timescales

- ▶ The risk-time is the same on all timescales
- ▶ Only need the entry point on each time scale:
  - ▶ Age at entry.
  - ▶ Date of entry.
  - ▶ Time since treatment at entry. — if time of treatment is the entry, this is 0 for all.

- ▶ Response variable in analysis of rates:

$(d, y)$  (event, duration)

- ▶ Covariates in analysis of rates:
  - ▶ timescales
  - ▶ other (fixed) measurements



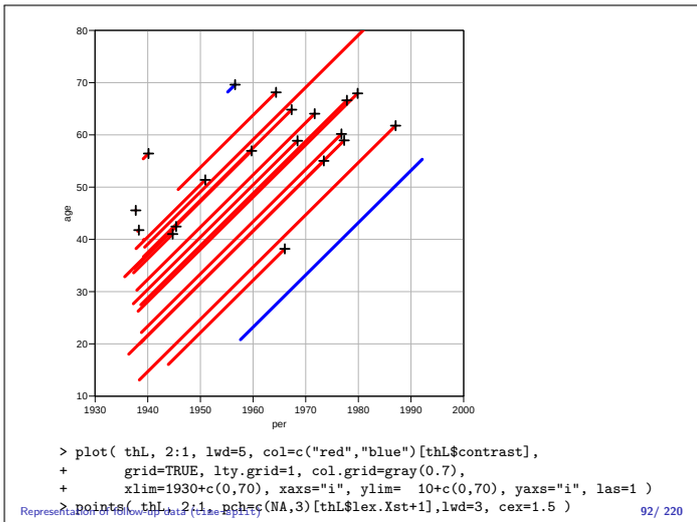
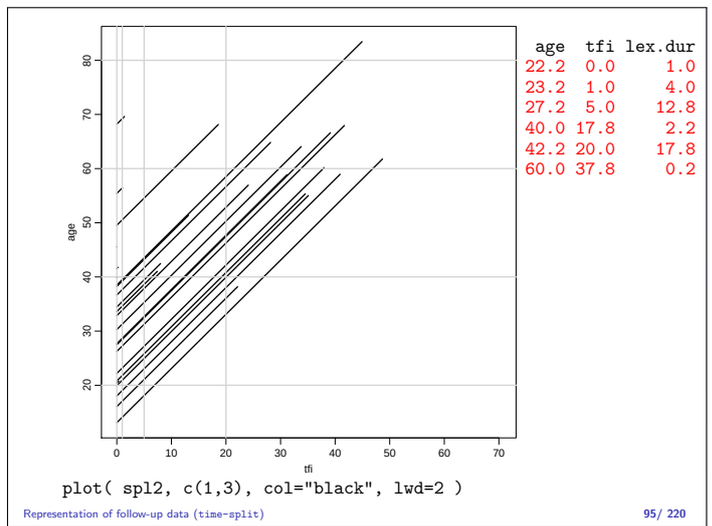
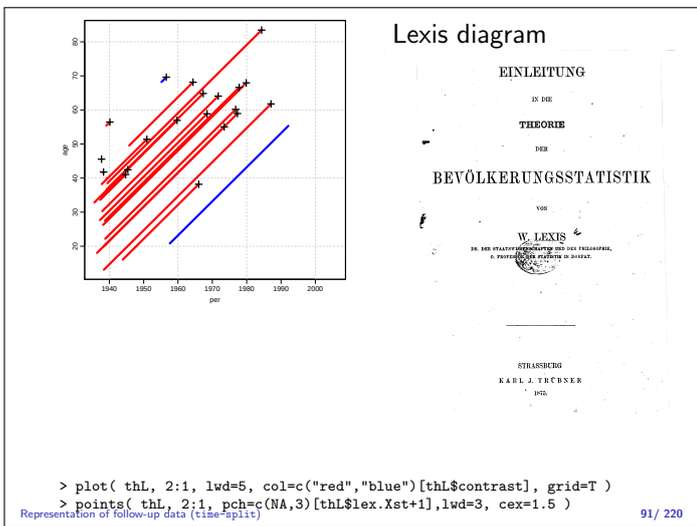
```
> plot( thL, lwd=3 )
```

Representation of follow-up data (time-split)

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Representation of follow-up data (time-split)

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### Likelihood for a piecewise constant rate

- ▶ This setup is for a situation where it is assumed that rates are constant in each of the intervals.
- ▶ Each observation in the dataset contributes a term to a "Poisson" likelihood.
- ▶ Models can include fixed covariates, as well as the timescales (the left end-points of the intervals) as continuous variables.
- ▶ Rates are assumed to vary by timescales:
  - ▶ continuously
  - ▶ non-linearly
- ▶ Rates can vary along several timescales simultaneously.

Representation of follow-up data (time-split)

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### Splitting follow-up time

```
> spl1 <- splitLexis( thL, breaks=seq(0,100,20),
+ time.scale="age" )
> round(spl1,1)
  age per tfi lex.dur lex.Cst lex.Xst id sex birthdat con
1 22.2 1938.8 0.0 17.8 0 0 1 2 1916.6
2 40.0 1956.6 17.8 20.0 0 0 1 2 1916.6
3 60.0 1976.6 37.8 0.2 0 1 1 2 1916.6
4 49.5 1945.8 0.0 10.5 0 0 640 2 1896.2
5 60.0 1956.2 10.5 8.1 0 1 640 2 1896.2
6 68.2 1955.2 0.0 1.4 0 1 3425 1 1887.0
7 20.8 1957.6 0.0 19.2 0 0 4017 2 1936.8
8 40.0 1976.8 19.2 15.3 0 0 4017 2 1936.8
...
```

Representation of follow-up data (time-split)

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### Where is $(d_{pi}, y_{pi})$ in the split data?

Likelihood is  $d_{pi} \log(\lambda_{pi}) - \lambda_{pi} y_{pi}$

```
> round( spl2, 1 )
  lex.id age per tfi lex.dur lex.Cst lex.Xst id sex birt
1 1 22.2 1938.8 0.0 1.0 0 0 1 2 19
2 1 23.2 1939.8 1.0 4.0 0 0 1 2 19
3 1 27.2 1943.8 5.0 12.8 0 0 1 2 19
4 1 40.0 1956.6 17.8 2.2 0 0 1 2 19
5 1 42.2 1958.8 20.0 17.8 0 0 1 2 19
6 1 60.0 1976.6 37.8 0.2 0 1 1 2 19
7 2 49.5 1945.8 0.0 1.0 0 0 640 2 18
8 2 50.5 1946.8 1.0 4.0 0 0 640 2 18
9 2 54.5 1950.8 5.0 5.5 0 0 640 2 18
10 2 60.0 1956.2 10.5 8.1 0 1 640 2 18
...
```

— and what are **covariates** for the rates?

Representation of follow-up data (time-split)

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### Split on another timescale

```
> spl2 <- splitLexis( spl1, time.scale="tfi",
+ breaks=c(0,1,5,20,100) )
> round( spl2, 1 )
  lex.id age per tfi lex.dur lex.Cst lex.Xst id sex birt
1 1 22.2 1938.8 0.0 1.0 0 0 1 2 19
2 1 23.2 1939.8 1.0 4.0 0 0 1 2 19
3 1 27.2 1943.8 5.0 12.8 0 0 1 2 19
4 1 40.0 1956.6 17.8 2.2 0 0 1 2 19
5 1 42.2 1958.8 20.0 17.8 0 0 1 2 19
6 1 60.0 1976.6 37.8 0.2 0 1 1 2 19
7 2 49.5 1945.8 0.0 1.0 0 0 640 2 18
8 2 50.5 1946.8 1.0 4.0 0 0 640 2 18
9 2 54.5 1950.8 5.0 5.5 0 0 640 2 18
10 2 60.0 1956.2 10.5 8.1 0 1 640 2 18
11 3 68.2 1955.2 0.0 1.0 0 0 3425 1 18
12 3 69.2 1956.2 1.0 0.4 0 1 3425 1 18
13 4 20.8 1957.6 0.0 1.0 0 0 4017 2 19
14 4 21.8 1958.6 1.0 4.0 0 0 4017 2 19
15 4 25.8 1962.6 5.0 14.2 0 0 4017 2 19
16 4 40.0 1976.8 19.2 0.8 0 0 4017 2 19
17 4 40.8 1977.6 20.0 14.5 0 0 4017 2 19
...
```

Representation of follow-up data (time-split)

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### Analysis of results

- ▶  $d_{pi}$  — events in the variable: lex.Xst:  
In the model as response: lex.Xst==1
- ▶  $y_{pi}$  — risk time: lex.dur (duration):  
In the model as offset  $\log(y)$ ,  $\log(\text{lex.dur})$ .
- ▶ Covariates are:
  - ▶ timescales (age, period, time in study)
  - ▶ other variables for this person (constant or assumed constant in each interval).
- ▶ Model rates using the covariates in glm:  
— no difference between time-scales and other covariates.

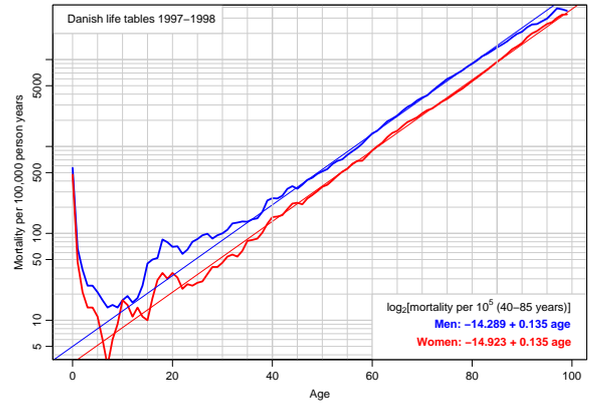
Representation of follow-up data (time-split)

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# Classical estimators: Lifetable

Modern Demographic  
Methods in Epidemiology  
with R  
23 November 2015  
University of Melbourne  
<http://BendixCarstensen/AdvCoh/Melb-2015>

1tab



Classical estimators: Lifetable (1tab)

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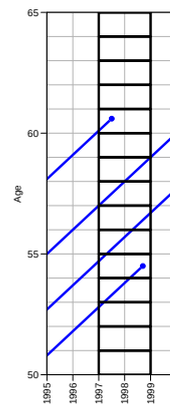
## Survival analysis

- ▶ Response variable: Time to event,  $T$
- ▶ Censoring time,  $Z$
- ▶ We observe  $(\min(T, Z), \delta = 1\{T < Z\})$ .
- ▶ This gives time a special status, and mixes the response variable (risk)time with the covariate time(scale).
- ▶ Originates from clinical trials where everyone enters at time 0, and therefore  $Y = T - 0 = T$

Classical estimators: Lifetable (1tab)

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## Observations for the lifetable



Life table is based on person-years and deaths accumulated in a short period.

Age-specific rates — cross-sectional!

Survival function:

$$S(t) = e^{-\int_0^t \lambda(a) da} = e^{-\sum_0^t \lambda(a)}$$

— assumes stability of rates to be interpretable for actual persons.

Classical estimators: Lifetable (1tab)

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## The life table method

The simplest analysis is by the “life-table method”:

| interval | alive | dead  | cens. |                        |
|----------|-------|-------|-------|------------------------|
| $i$      | $n_i$ | $d_i$ | $l_i$ | $p_i$                  |
| 1        | 77    | 5     | 2     | $5/(77 - 2/2) = 0.066$ |
| 2        | 70    | 7     | 4     | $7/(70 - 4/2) = 0.103$ |
| 3        | 59    | 8     | 1     | $8/(59 - 1/2) = 0.137$ |

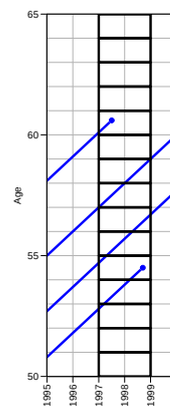
$$p_i = P\{\text{death in interval } i\} = 1 - d_i/(n_i - l_i/2)$$

$$S(t) = (1 - p_1) \times \dots \times (1 - p_t)$$

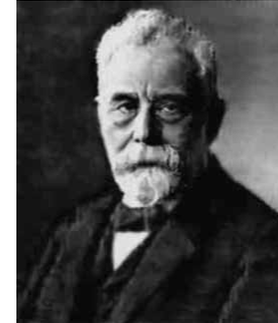
Classical estimators: Lifetable (1tab)

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## Observations for the lifetable



This is a Lexis diagram.



Classical estimators: Lifetable (1tab)

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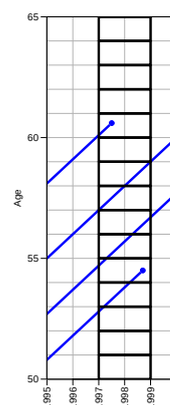
## Population life table, DK 1997-98

| $a$ | Men     |              |                    | Women   |              |                    |
|-----|---------|--------------|--------------------|---------|--------------|--------------------|
|     | $S(a)$  | $\lambda(a)$ | $E[\ell_{res}(a)]$ | $S(a)$  | $\lambda(a)$ | $E[\ell_{res}(a)]$ |
| 0   | 1.00000 | 567          | 73.68              | 1.00000 | 474          | 78.65              |
| 1   | 0.99433 | 67           | 73.10              | 0.99526 | 47           | 78.02              |
| 2   | 0.99366 | 38           | 72.15              | 0.99479 | 21           | 77.06              |
| 3   | 0.99329 | 25           | 71.18              | 0.99458 | 14           | 76.08              |
| 4   | 0.99304 | 25           | 70.19              | 0.99444 | 14           | 75.09              |
| 5   | 0.99279 | 21           | 69.21              | 0.99430 | 11           | 74.10              |
| 6   | 0.99258 | 17           | 68.23              | 0.99419 | 6            | 73.11              |
| 7   | 0.99242 | 14           | 67.24              | 0.99413 | 3            | 72.11              |
| 8   | 0.99227 | 15           | 66.25              | 0.99410 | 6            | 71.11              |
| 9   | 0.99213 | 14           | 65.26              | 0.99404 | 9            | 70.12              |
| 10  | 0.99199 | 17           | 64.26              | 0.99395 | 17           | 69.12              |
| 11  | 0.99181 | 19           | 63.28              | 0.99378 | 15           | 68.14              |
| 12  | 0.99162 | 16           | 62.29              | 0.99363 | 11           | 67.15              |
| 13  | 0.99147 | 18           | 61.30              | 0.99352 | 14           | 66.15              |
| 14  | 0.99129 | 25           | 60.31              | 0.99338 | 11           | 65.16              |
| 15  | 0.99104 | 45           | 59.32              | 0.99327 | 10           | 64.17              |
| 16  | 0.99059 | 50           | 58.35              | 0.99317 | 18           | 63.18              |
| 17  | 0.99009 | 52           | 57.38              | 0.99299 | 29           | 62.19              |
| 18  | 0.98957 | 85           | 56.41              | 0.99270 | 35           | 61.21              |
| 19  | 0.98873 | 79           | 55.46              | 0.99235 | 30           | 60.23              |
| 20  | 0.98795 | 70           | 54.50              | 0.99205 | 35           | 59.24              |
| 21  | 0.98726 | 71           | 53.54              | 0.99170 | 31           | 58.27              |

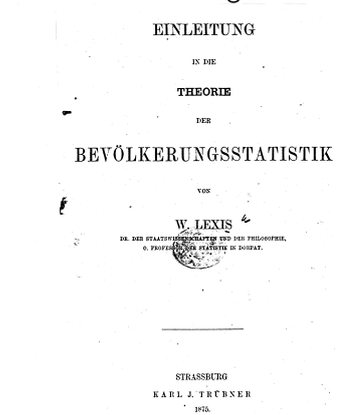
Classical estimators: Lifetable (1tab)

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## Observations for the lifetable



This is a Lexis diagram.



Classical estimators: Lifetable (1tab)

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## Life table approach

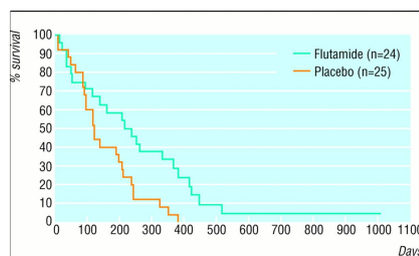
individual.

- ▶ The **population** experience:
  - $D$ : Deaths (events).
  - $Y$ : Person-years (risk time).
- ▶ The classical lifetable analysis compiles these for prespecified intervals of age, and computes age-specific mortality **rates**.
- ▶ Data are collected crosssectionally, but interpreted longitudinally.
- ▶ The **rates** are the basic building blocks — used for construction of:
  - ▶ RRs
  - ▶ cumulative measures (survival and risk)

Classical estimators: Lifetable (1tab)

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## Example of KM Survival Curve from BMJ



BMJ 1998;316:1935-1938

Kaplan-Meier curve from an RCT of patients with pancreatic cancer

Classical estimators: Kaplan-Meier (km-na)

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## Summary

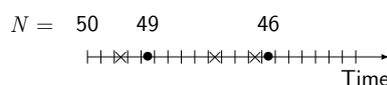
- ▶ Follow-up studies observe time to event
- ▶ — in the form of **empirical rates**,  $(d, y)$  for small interval
- ▶ each interval (empirical rate) has covariates attached
- ▶ each interval contribute  $d \log(\lambda) - \lambda y$
- ▶ — like a Poisson observation  $d$  with mean  $\lambda y$
- ▶ identical covariates: pool observations to  $D = \sum D, Y = \sum y$
- ▶ — like a Poisson observation  $D$  with mean  $\lambda Y$
- ▶ the result is an **estimate** of the rate  $\lambda$
- ▶ from a **model** where rates are constant within intervals — but varies between intervals.

Classical estimators: Lifetable (1tab)

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## Kaplan-Meier method illustrated

(● = failure and × = censored):



- ▶ Steps caused by multiplying by  $(1 - 1/49)$  and  $(1 - 1/46)$  respectively
- ▶ Late entry can also be dealt with

Classical estimators: Kaplan-Meier (km-na)

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## Classical estimators: Kaplan-Meier

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km-na

## Using R: Surv()

```
library( survival )
data( lung )
head( lung, 3 )
inst time status age sex ph.ecog ph.karno pat.karno meal.cal w
1 3 306 2 74 1 1 90 100 1175
2 3 455 2 68 1 0 90 90 1225
3 3 1010 1 56 1 0 90 90 NA
with( lung, Surv( time, status==2 ) )[1:10]
[1] 306 455 1010+ 210 883 1022+ 310 361 218 166
( s.km <- survfit( Surv( time, status==2 ) ~ 1, data=lung ) )
Call: survfit(formula = Surv(time, status == 2) ~ 1, data = lung

      n events median 0.95LCL 0.95UCL
228 165 310 285 363

plot( s.km )
abline( v=310, h=0.5, col="red" )
```

Classical estimators: Kaplan-Meier (km-na)

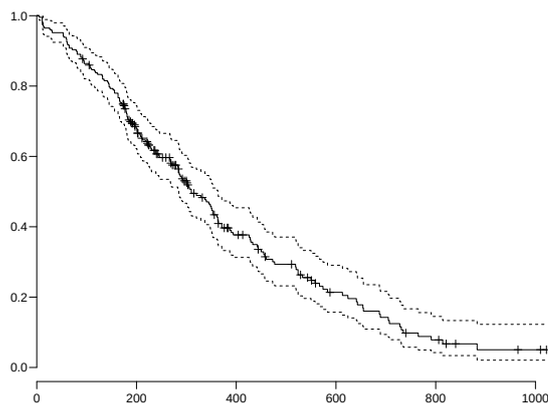
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## The Kaplan-Meier Method

- ▶ The most common method of estimating the survival function.
- ▶ A non-parametric method.
- ▶ Divides time into small intervals where the intervals are defined by the unique times of failure (death).
- ▶ Based on conditional probabilities as we are interested in the probability a subject surviving the next time interval given that they have survived so far.

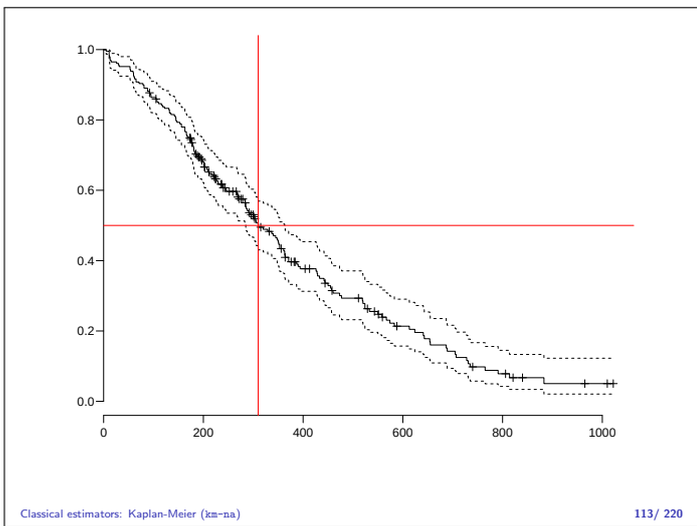
Classical estimators: Kaplan-Meier (km-na)

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Classical estimators: Kaplan-Meier (km-na)

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## Proportional Hazards model

- ▶ The baseline hazard rate,  $\lambda_0(t)$ , is the hazard rate when all the covariates are 0.
- ▶ The form of the above equation means that covariates act **multiplicatively** on the baseline hazard rate.
- ▶ Time is a covariate (albeit with special status).
- ▶ The baseline hazard is a function of time and thus varies with time.
- ▶ No assumption about the shape of the underlying hazard function.
- ▶ — but you will never see the shape . . .

The Cox model (cox)

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## The Cox model

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cox

## Interpreting Regression Coefficients

- ▶ If  $x_j$  is binary  $\exp(\beta_j)$  is the estimated hazard ratio for subjects corresponding to  $x_j = 1$  compared to those where  $x_j = 0$ .
- ▶ If  $x_j$  is continuous  $\exp(\beta_j)$  is the estimated increase/decrease in the hazard rate for a unit change in  $x_j$ .
- ▶ With more than one covariate interpretation is similar, i.e.  $\exp(\beta_j)$  is the hazard ratio for subjects who **only** differ with respect to covariate  $x_j$ .

The Cox model (cox)

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## The proportional hazards model

$$\lambda(t, x) = \lambda_0(t) \times \exp(x'\beta)$$

A model for the rate as a function of  $t$  and  $x$ .

The covariate  $t$  has a special status:

- ▶ Computationally, because all individuals contribute to (some of) the range of  $t$ .
- ▶ Conceptually it is less clear —  $t$  is but a covariate that varies **within** each individual.

The Cox model (cox)

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## Fitting a Cox- model in R

```
library( survival )
data(bladder)
bladder <- subset( bladder, enum<2 )
head( bladder)
  id rx number size stop event enum
1  1  1     1   3     1     0    1
5  2  1     2   1     4     0    1
9  3  1     1   1     7     0    1
13 4  1     5   1    10     0    1
17 5  1     4   1     6     1    1
21 6  1     1   1    14     0    1
```

The Cox model (cox)

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## Cox-likelihood

The partial likelihood for the regression parameters:

$$\ell(\beta) = \sum_{\text{death times}} \log \left( \frac{e^{x_{\text{death}}\beta}}{\sum_{i \in \mathcal{R}_t} e^{x_i\beta}} \right)$$

- ▶ This is David Cox's invention.
- ▶ Extremely efficient from a computational point of view.
- ▶ The baseline hazard is bypassed (profiled out).

The Cox model (cox)

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## Fitting a Cox-model in R

```
c0 <- coxph( Surv(stop,event) ~ number + size, data=bladder )
c0
Call:
coxph(formula = Surv(stop, event) ~ number + size, data = bladder)

      coef exp(coef) se(coef)      z      p
number 0.2049   1.2274   0.0704  2.91 0.0036
size    0.0613   1.0633   0.1033  0.59 0.5525

Likelihood ratio test=7.04 on 2 df, p=0.0296
n= 85, number of events= 47
```

The Cox model (cox)

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## Plotting the base survival in R

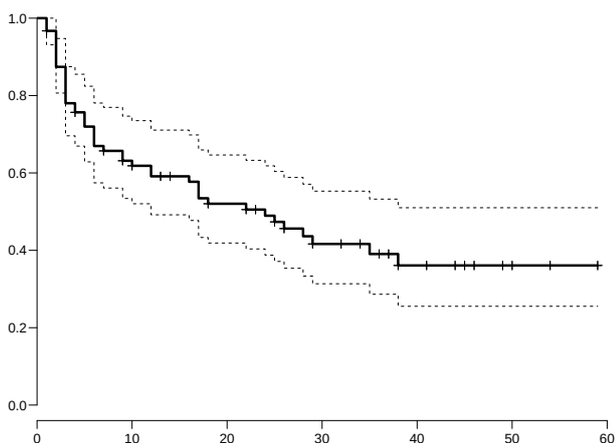
```
plot(survfit(c0))
lines(survfit(c0), conf.int=F, lwd=3)
```

The `plot.coxph` plots the survival curve for a person with an average covariate value

- which is **not** the average survival for the population considered...
- and not necessarily meaningful

## Who needs the Cox-model anyway?

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## The proportional hazards model

$$\lambda(t, x) = \lambda_0(t) \times \exp(x'\beta)$$

A model for the rate as a function of  $t$  and  $x$ .

The covariate  $t$  has a special status:

- ▶ Computationally, because all individuals contribute to (some of) the range of  $t$ .
- ▶ Conceptually it is less clear —  $t$  is but a covariate that varies within individual.

## Plotting the base survival in R

You can plot the survival curve for specific values of the covariates, using the `newdata=` argument:

```
plot(survfit(c0))
lines(survfit(c0), conf.int=F, lwd=3)
lines(survfit(c0, newdata=data.frame(number=1, size=1)),
      lwd=2, col="limegreen")
text(par("usr")[2]*0.98, 1.00, "number=1,size=1",
     col="limegreen", font=2, adj=1)
```

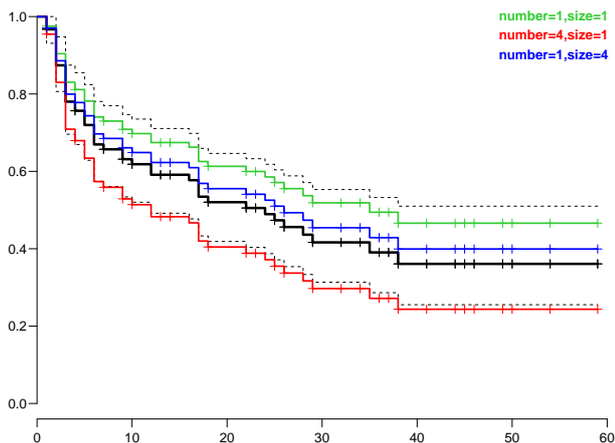
## Cox-likelihood

The (partial) log-likelihood for the regression parameters:

$$\ell(\beta) = \sum_{\text{death times}} \log \left( \frac{e^{\eta_{\text{death}}}}{\sum_{i \in \mathcal{R}_t} e^{\eta_i}} \right)$$

is also a **profile likelihood** in the model where observation time has been subdivided in small pieces (empirical rates) and each small piece provided with its own parameter:

$$\log(\lambda(t, x)) = \log(\lambda_0(t)) + x'\beta = \alpha_t + \eta$$



## The Cox-likelihood as profile likelihood

- ▶ Regression parameters describing the effect of covariates (other than the chosen underlying time scale).
- ▶ One parameter per death time to describe the effect of time (i.e. the chosen timescale).

$$\log(\lambda(t, x_i)) = \log(\lambda_0(t)) + \beta_1 x_{1i} + \dots + \beta_p x_{pi} = \alpha_t +$$

- ▶ Profile likelihood:
  - ▶ Derive estimates of  $\alpha_t$  as function of data and  $\beta$ s
  - ▶ Insert in likelihood, now only a function of data and  $\beta$ s
  - ▶ Turns out to be Cox's partial likelihood

- ▶ Suppose the time scale has been divided into small intervals with at most one death in each.
- ▶ Assume w.l.o.g. the  $y_i$  in the empirical rates all are 1.
- ▶ Log-likelihood contributions that contain information on a specific time-scale parameter  $\alpha_t$  will be from:
  - ▶ the (only) empirical rate (1, 1) with the death at time  $t$ .
  - ▶ all other empirical rates (0, 1) from those who were at risk at time  $t$ .

## Sensible modelling

Replace the  $\alpha_{t_i}$  by a parametric function  $f(t)$  with a limited number of parameters, for example:

- ▶ Piecewise constant
- ▶ Splines (linear, quadratic or cubic)
- ▶ Fractional polynomials

Use Poisson modelling software on a dataset of empirical rates for small intervals ( $y_i$ ).

Note: There is one contribution from each person at risk to this part of the log-likelihood:

$$\begin{aligned} \ell_t(\alpha_t, \beta) &= \sum_{i \in \mathcal{R}_t} d_i \log(\lambda_i(t)) - \lambda_i(t) y_i \\ &= \sum_{i \in \mathcal{R}_t} \{d_i(\alpha_t + \eta_i) - e^{\alpha_t + \eta_i}\} \\ &= \alpha_t + \eta_{\text{death}} - e^{\alpha_t} \sum_{i \in \mathcal{R}_t} e^{\eta_i} \end{aligned}$$

where  $\eta_{\text{death}}$  is the linear predictor for the person that died.

## Splitting the dataset

- ▶ The Poisson approach needs a dataset of empirical rates with small values of  $y$ .
- ▶ Larger than the original: each individual contributes many empirical rates. From each empirical rate we get:
  - ▶ Poisson-response  $d$
  - ▶ Risk time  $y$
  - ▶ Covariate value for the timescale (time since entry, current age, current date, ...)
  - ▶ other covariates

The derivative w.r.t.  $\alpha_t$  is:

$$D_{\alpha_t} \ell(\alpha_t, \beta) = 1 - e^{\alpha_t} \sum_{i \in \mathcal{R}_t} e^{\eta_i} = 0 \quad \Leftrightarrow \quad e^{\alpha_t} = \frac{1}{\sum_{i \in \mathcal{R}_t} e^{\eta_i}}$$

If this estimate is fed back into the log-likelihood for  $\alpha_t$ , we get the **profile likelihood** (with  $\alpha_t$  "profiled out"):

$$\log \left( \frac{1}{\sum_{i \in \mathcal{R}_t} e^{\eta_i}} \right) + \eta_{\text{death}} - 1 = \log \left( \frac{e^{\eta_{\text{death}}}}{\sum_{i \in \mathcal{R}_t} e^{\eta_i}} \right) - 1$$

which is the same as the contribution from time  $t$  to Cox's partial likelihood.

## Example: Mayo Clinic lung cancer

```
> library( survival ) ; library( Epi )
> data( lung )
> head( lung )
  inst time status age sex ph.ecog ph.karno pat.karno meal.cal w
1     3  306     2  74  1     1     90     100    1175
2     3  455     2  68  1     0     90     90    1225
3     3 1010     1  56  1     0     90     90     NA
4     5  210     2  57  1     1     90     60    1150
5     1  883     2  60  1     0    100     90     NA
6    12 1022     1  74  1     1     50     80     513
> Lx <- Lexis( exit=list( tfd=time), exit.status=(status==2), da
NOTE: entry is assumed to be 0 on the tfd timescale.
> summary( Lx, scale=365.25 )
Transitions:
  To
From FALSE TRUE Records: Events: Risk time: Persons:
FALSE  63 165    228    165    190.54    228
> Sx <- splitLexis( Lx, "tfd", breaks=c(0,unique(Lx$time)) )
> summary( Sx, scale=365.25 )
Transitions:
  To
From FALSE TRUE Records: Events: Risk time: Persons:
FALSE 1985 165    20022    165    190.54    228
```

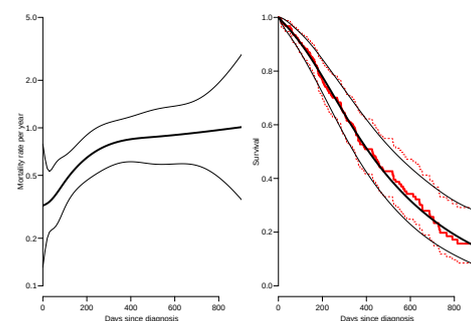
## What the Cox-model really is

Taking the life-table approach *ad absurdum* by:

- ▶ dividing time very finely,
- ▶ modelling one covariate, the time-scale, with one parameter per distinct value,
- ▶ profiling these parameters out and maximizing the profile likelihood,
- ▶ regression parameters are the same as in the full model with all the interval-specific parameters
- ▶ Subsequently, one may recover the effect of the timescale by smoothing an estimate of the cumulative sum of these.

## Mayo clinic lung cancer data

Smoothing by natural splines with 5 parameters, knots at 0, 25, 100, 500, 1000 days:



# Multiple time scales and continuous rates

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crv-mod

## Linear effects in glm

```
> nd <- data.frame( A=15:60, Y=10^-5 )
> pr <- ci.pred( ml, newdata=nd )
> head( pr )
      Estimate      2.5%      97.5%
1 6.170105 5.991630 6.353896
2 6.204034 6.028525 6.384652
3 6.238149 6.065547 6.415662
4 6.272452 6.102689 6.446937
5 6.306943 6.139944 6.478485
6 6.341624 6.177301 6.510319
> matplot( nd$A, pr,
+         type="l", lty=1, lwd=c(3,1,1), col="black", log="y" )
```

Multiple time scales and continuous rates (crv-mod)

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## Testis cancer

Testis cancer in Denmark:

```
> options( show.signif.stars=FALSE )
> library( Epi )
> data( testisDK )
> str( testisDK )
'data.frame': 4860 obs. of 4 variables:
 $ A: num 0 1 2 3 4 5 6 7 8 9 ...
 $ P: num 1943 1943 1943 1943 1943 ...
 $ D: num 1 1 0 1 0 0 0 0 0 0 ...
 $ Y: num 39650 36943 34588 33267 32614 ...
> head( testisDK )
  A   P   D     Y
1 0 1943 1 39649.50
2 1 1943 1 36942.83
3 2 1943 0 34588.33
4 3 1943 1 33267.00
5 4 1943 0 32614.00
6 5 1943 0 32020.33
```

Multiple time scales and continuous rates (crv-mod)

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## Linear effects in glm

```
> round( ci.lin( ml ), 4 )
      Estimate StdErr      z P      2.5%      97.5%
(Intercept) -9.7755 0.0207 -472.3164 0 -9.8160 -9.7349
A            0.0055 0.0005  11.3926 0  0.0045  0.0064
> CI <- cbind( 1, nd$A )
> head( CI )
      [,1] [,2]
[1,]    1  15
[2,]    1  16
[3,]    1  17
[4,]    1  18
[5,]    1  19
[6,]    1  20
> matplot( nd$A, ci.exp( ml, ctr.mat=CI ),
+         type="l", lty=1, lwd=c(3,1,1), col="black", log="y" )
```

Multiple time scales and continuous rates (crv-mod)

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## Cases, PY and rates

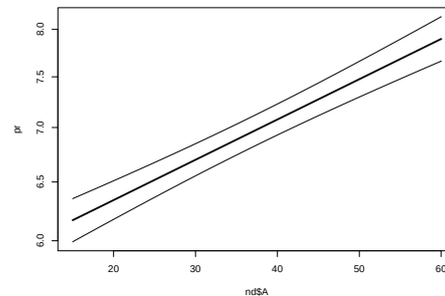
```
> stat.table( list(A=floor(A/10)*10,
+                 P=floor(P/10)*10),
+            list( D=sum(D),
+                 Y=sum(Y/1000),
+                 rate=rate(D,Y,10^-5) ),
+            margins=TRUE, data=testisDK )
```

| A  | P       |         |         |         |         |         |
|----|---------|---------|---------|---------|---------|---------|
|    | 1940    | 1950    | 1960    | 1970    | 1980    | 1990    |
| 0  | 10.00   | 7.00    | 16.00   | 18.00   | 9.00    | 10.00   |
|    | 2604.66 | 4037.31 | 3884.97 | 3820.88 | 3070.87 | 2165.54 |
|    | 0.38    | 0.17    | 0.41    | 0.47    | 0.29    | 0.46    |
| 10 | 13.00   | 27.00   | 37.00   | 72.00   | 97.00   | 75.00   |
|    | 2135.73 | 3505.19 | 4004.13 | 3906.08 | 3847.40 | 2260.97 |
|    | 0.61    | 0.77    | 0.92    | 1.84    | 2.52    | 3.32    |
| 20 | 124.00  | 221.00  | 280.00  | 535.00  | 724.00  | 557.00  |
|    | 2225.55 | 2923.22 | 3401.65 | 4028.57 | 3941.18 | 2824.58 |
|    | 5.57    | 7.56    | 8.23    | 13.28   | 18.37   | 19.72   |
| 30 | 149.00  | 288.00  | 377.00  | 624.00  | 771.00  | 744.00  |

Multiple time scales and continuous rates (crv-mod)

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## Linear effects in glm



```
> matplot( nd$A, pr,
+         type="l", lty=1, lwd=c(3,1,1), col="black", log="y" )
```

Multiple time scales and continuous rates (crv-mod)

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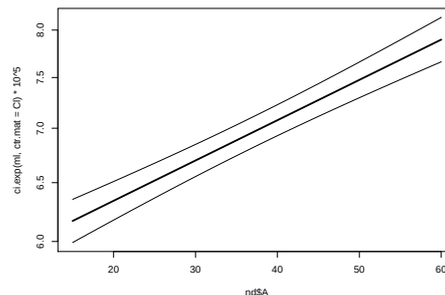
## Linear effects in glm

How do rates depend on age?

```
> ml <- glm( D ~ A, offset=log(Y), family=poisson, data=testisDK )
> round( ci.lin( ml ), 4 )
      Estimate StdErr      z P      2.5%      97.5%
(Intercept) -9.7755 0.0207 -472.3164 0 -9.8160 -9.7349
A            0.0055 0.0005  11.3926 0  0.0045  0.0064
> round( ci.exp( ml ), 4 )
      exp(Est.)      2.5%      97.5%
(Intercept) 0.0001 0.0001 0.0001
A            1.0055 1.0046 1.0064
```

Linear increase of log-rates by age

## Linear effects in glm



```
> matplot( nd$A, ci.exp( ml, ctr.mat=CI ) * 10^-5,
+         type="l", lty=1, lwd=c(3,1,1), col="black", log="y" )
```

Multiple time scales and continuous rates (crv-mod)

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Multiple time scales and continuous rates (crv-mod)

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## Quadratic effects in glm

How do rates depend on age?

```
> mq <- glm( D ~ A + I(A^2),
+           offset=log(Y), family=poisson, data=testisDK )
> round( ci.lin( mq ), 4 )
```

|             | Estimate | StdErr | z         | P | 2.5%     | 97.5%    |
|-------------|----------|--------|-----------|---|----------|----------|
| (Intercept) | -12.3656 | 0.0596 | -207.3611 | 0 | -12.4825 | -12.2487 |
| A           | 0.1806   | 0.0033 | 54.8290   | 0 | 0.1741   | 0.1871   |
| I(A^2)      | -0.0023  | 0.0000 | -53.7006  | 0 | -0.0024  | -0.0022  |

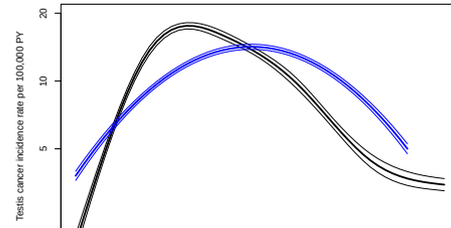
```
> round( ci.exp( mq ), 4 )
```

|             | exp(Est.) | 2.5%   | 97.5%  |
|-------------|-----------|--------|--------|
| (Intercept) | 0.0000    | 0.0000 | 0.0000 |
| A           | 1.1979    | 1.1902 | 1.2057 |
| I(A^2)      | 0.9977    | 0.9976 | 0.9978 |

Multiple time scales and continuous rates (crv-mod)

142/ 220

## Spline effects in glm



```
> matplot( aa, ci.exp( msp, ctr.mat=cbind(1,As) ) * 10^5,
+         log="y", xlab="Age", ylab="Testis cancer incidence rate per 100,000 PY in 1970",
+         type="l", lty=1, lwd=c(3,1,1), col="black", ylim=c(2,20) )
> matlines( nd$A, ci.exp( mq, ctr.mat=Cq ) * 10^5,
+         type="l", lty=1, lwd=c(3,1,1), col="blue" )
```

Multiple time scales and continuous rates (crv-mod)

146/ 220

## Quadratic effect in glm

```
> round( ci.lin( mq ), 4 )
```

|             | Estimate | StdErr | z         | P | 2.5%     | 97.5%    |
|-------------|----------|--------|-----------|---|----------|----------|
| (Intercept) | -12.3656 | 0.0596 | -207.3611 | 0 | -12.4825 | -12.2487 |
| A           | 0.1806   | 0.0033 | 54.8290   | 0 | 0.1741   | 0.1871   |
| I(A^2)      | -0.0023  | 0.0000 | -53.7006  | 0 | -0.0024  | -0.0022  |

```
> Cq <- cbind( 1, 15:60, (15:60)^2 )
> head( Cq, 4 )
```

|      | [,1] | [,2] | [,3] |
|------|------|------|------|
| [1,] | 1    | 15   | 225  |
| [2,] | 1    | 16   | 256  |
| [3,] | 1    | 17   | 289  |
| [4,] | 1    | 18   | 324  |

```
> matplot( nd$A, ci.exp( mq, ctr.mat=Cq ) * 10^5,
+         type="l", lty=1, lwd=c(3,1,1), col="black", log="y" )
```

Multiple time scales and continuous rates (crv-mod)

143/ 220

## Adding a linear period effect

```
> msp <- glm( D ~ Ns(A,knots=seq(15,65,10)) + P,
+           offset=log(Y), family=poisson, data=testisDK )
> round( ci.lin( msp ), 3 )
```

|                                 | Estimate | StdErr | z       | P     |
|---------------------------------|----------|--------|---------|-------|
| (Intercept)                     | -58.105  | 1.444  | -40.229 | 0.000 |
| Ns(A, knots = seq(15, 65, 10))1 | 2.120    | 0.057  | 37.444  | 0.000 |
| Ns(A, knots = seq(15, 65, 10))2 | 1.700    | 0.068  | 25.157  | 0.000 |
| Ns(A, knots = seq(15, 65, 10))3 | 0.007    | 0.060  | 0.110   | 0.913 |
| Ns(A, knots = seq(15, 65, 10))4 | 2.596    | 0.097  | 26.631  | 0.000 |
| Ns(A, knots = seq(15, 65, 10))5 | -0.780   | 0.042  | -18.748 | 0.000 |
| P                               | 0.024    | 0.001  | 32.761  | 0.000 |

```
> Ca <- cbind( 1, Ns( aa, knots=seq(15,65,10) ), 1970 )
> head( Ca )
```

|      | 1 | 2            | 3 | 4           | 5          |             |
|------|---|--------------|---|-------------|------------|-------------|
| [1,] | 1 | 0.0000000000 | 0 | 0.00000000  | 0.00000000 | 1970        |
| [2,] | 1 | 0.0001666667 | 0 | -0.02527011 | 0.07581034 | -0.05054022 |
| [3,] | 1 | 0.0013333333 | 0 | -0.05003313 | 0.15009940 | -0.10006626 |
| [4,] | 1 | 0.0045000000 | 0 | -0.07378197 | 0.22134590 | -0.14756393 |
| [5,] | 1 | 0.0106666667 | 0 | -0.09600952 | 0.28802857 | -0.19201905 |
| [6,] | 1 | 0.0208333333 | 0 | -0.11620871 | 0.34862613 | -0.23241742 |

```
> matplot( aa, ci.exp( msp, ctr.mat=Ca ) * 10^5,
+         log="y", xlab="Age", ylab="Testis cancer incidence rate per 100,000 PY in 1970",
+         type="l", lty=1, lwd=c(3,1,1), col="black", ylim=c(2,20) )
```

Multiple time scales and continuous rates (crv-mod)

147/ 220

## Quadratic effect in glm

```
> matplot( nd$A, ci.exp( mq, ctr.mat=Cq ) * 10^5,
+         type="l", lty=1, lwd=c(3,1,1), col="black", log="y" )
> matlines( nd$A, ci.exp( ml, ctr.mat=C1 ) * 10^5,
+         type="l", lty=1, lwd=c(3,1,1), col="blue" )
```

Multiple time scales and continuous rates (crv-mod)

144/ 220

## Adding a linear period effect

```
> matplot( aa, ci.exp( msp, ctr.mat=Ca ) * 10^5,
+         log="y", xlab="Age",
+         ylab="Testis cancer incidence rate per 100,000 PY in 1970",
+         type="l", lty=1, lwd=c(3,1,1), col="black", ylim=c(2,20) )
```

Multiple time scales and continuous rates (crv-mod)

148/ 220

## Spline effects in glm

```
> library( splines )
> ms <- glm( D ~ Ns(A,knots=seq(15,65,10)),
+           offset=log(Y), family=poisson, data=testisDK )
> round( ci.exp( ms ), 3 )
```

|                                 | exp(Est.) | 2.5%   | 97.5%  |
|---------------------------------|-----------|--------|--------|
| (Intercept)                     | 0.000     | 0.000  | 0.000  |
| Ns(A, knots = seq(15, 65, 10))1 | 8.548     | 7.650  | 9.551  |
| Ns(A, knots = seq(15, 65, 10))2 | 5.706     | 4.998  | 6.514  |
| Ns(A, knots = seq(15, 65, 10))3 | 1.002     | 0.890  | 1.128  |
| Ns(A, knots = seq(15, 65, 10))4 | 14.402    | 11.896 | 17.436 |
| Ns(A, knots = seq(15, 65, 10))5 | 0.466     | 0.429  | 0.505  |

```
> aa <- 15:65
> As <- Ns( aa, knots=seq(15,65,10) )
> head( As )
```

|      | 1            | 2 | 3           | 4          | 5           |
|------|--------------|---|-------------|------------|-------------|
| [1,] | 0.0000000000 | 0 | 0.00000000  | 0.00000000 | 0.00000000  |
| [2,] | 0.0001666667 | 0 | -0.02527011 | 0.07581034 | -0.05054022 |
| [3,] | 0.0013333333 | 0 | -0.05003313 | 0.15009940 | -0.10006626 |
| [4,] | 0.0045000000 | 0 | -0.07378197 | 0.22134590 | -0.14756393 |
| [5,] | 0.0106666667 | 0 | -0.09600952 | 0.28802857 | -0.19201905 |
| [6,] | 0.0208333333 | 0 | -0.11620871 | 0.34862613 | -0.23241742 |

Multiple time scales and continuous rates (crv-mod)

145/ 220

## Adding a linear period effect

```
> matplot( aa, ci.exp( msp, ctr.mat=Ca ) * 10^5,
+         log="y", xlab="Age",
+         ylab="Testis cancer incidence rate per 100,000 PY in 1970",
+         type="l", lty=1, lwd=c(3,1,1), col="black", ylim=c(2,20) )
> matlines( nd$A, ci.pred( ms, newdata=nd ),
+         type="l", lty=1, lwd=c(3,1,1), col="blue" )
```

Multiple time scales and continuous rates (crv-mod)

149/ 220

## The period effect

```
> round( ci.lin( msp ), 3 )
      Estimate StdErr      z      P
(Intercept) -58.105  1.444 -40.229 0.000 -6
Ns(A, knots = seq(15, 65, 10))1  2.120  0.057 37.444 0.000
Ns(A, knots = seq(15, 65, 10))2  1.700  0.068 25.157 0.000
Ns(A, knots = seq(15, 65, 10))3  0.007  0.060  0.110 0.913
Ns(A, knots = seq(15, 65, 10))4  2.596  0.097 26.631 0.000
Ns(A, knots = seq(15, 65, 10))5  -0.780  0.042 -18.748 0.000
P          0.024  0.001 32.761 0.000
> pp <- seq(1945,1995,0.2)
> Cp <- cbind( pp ) - 1970
> head( Cp )
      pp
[1,] -25.0
[2,] -24.8
[3,] -24.6
[4,] -24.4
[5,] -24.2
[6,] -24.0
> ci.exp( msp, subset="P" )
exp(Est.)  2.5%  97.5%
P  1.024235 1.022769 1.025704
> matplot( pp, ci.exp( msp, subset="P", ctr.mat=Cp ),
```

Multiple time scales and continuous rates (crv-mod) 150 / 220

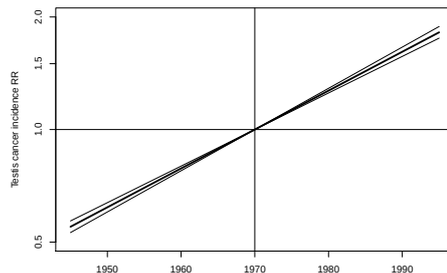
## A spline period effect

Because we have the age-effect with the rate dimension, the period effect is a RR

```
> msp <- glm( D ~ Ns(A,knots=seq(15,65,10)) +
+           Ns(P,knots=seq(1950,1990,10),ref=1970),
+           offset=log(Y), family=poisson, data=testisDK
> round( ci.exp( msp ), 3 )
      exp(Est.)  2.5%
Ns(A, knots = seq(15, 65, 10))1  0.000  0.000
Ns(A, knots = seq(15, 65, 10))2  8.327  7.452
Ns(A, knots = seq(15, 65, 10))3  5.528  4.842
Ns(A, knots = seq(15, 65, 10))4  1.007  0.894
Ns(A, knots = seq(15, 65, 10))5 13.447 11.107
Ns(A, knots = seq(15, 65, 10))6  0.458  0.422
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)1  1.711  1.526
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)2  2.190  2.028
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)3  3.222  2.835
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)4  2.299  2.149
```

Multiple time scales and continuous rates (crv-mod) 154 / 220

## Period effect



```
> matplot( pp, ci.exp( msp, subset="P", ctr.mat=Cp ),
+         log="y", ylim=c(0.5,2), xlab="Date",
+         ylab="Testis cancer incidence RR",
+         type="l", lty=1, lwd=c(3,1,1), col="black" )
> abline( h=1, v=1970 )
```

Multiple time scales and continuous rates (crv-mod) 151 / 220

## A spline period effect

```
> Cp <- Ns( pp, knots=seq(1950,1990,10),ref=1970)
> head( Cp, 4 )
      1      2      3      4
[1,] -0.6666667  0.0142689462 -0.5428068  0.3618712
[2,] -0.6666667  0.0091980207 -0.5275941  0.3517294
[3,] -0.6666667  0.0041270951 -0.5123813  0.3415875
[4,] -0.6666667 -0.0009438304 -0.4971685  0.3314457
> ci.exp( msp, subset="P" )
      exp(Est.)  2.
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)1  1.710808 1.5259
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)2  2.189650 2.0278
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)3  3.221563 2.8351
Ns(P, knots = seq(1950, 1990, 10), ref = 1970)4  2.298946 2.1491
> matplot( pp, ci.exp( msp, subset="P", ctr.mat=Cp ),
+         log="y", ylim=c(0.5,2), xlab="Date",
+         ylab="Testis cancer incidence RR",
+         type="l", lty=1, lwd=c(3,1,1), col="black" )
```

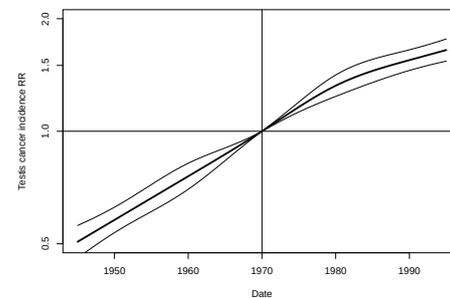
Multiple time scales and continuous rates (crv-mod) 155 / 220

## A quadratic period effect

```
> mspq <- glm( D ~ Ns(A,knots=seq(15,65,10)) + P + I(P^2),
+             offset=log(Y), family=poisson, data=testisDK
> round( ci.exp( mspq ), 3 )
      exp(Est.)  2.5%  97.5%
(Intercept)  0.000  0.000  0.000
Ns(A, knots = seq(15, 65, 10))1  8.356  7.478  9.337
Ns(A, knots = seq(15, 65, 10))2  5.513  4.829  6.295
Ns(A, knots = seq(15, 65, 10))3  1.006  0.894  1.133
Ns(A, knots = seq(15, 65, 10))4 13.439 11.101 16.269
Ns(A, knots = seq(15, 65, 10))5  0.458  0.422  0.497
P          2.189  1.457  3.291
I(P^2)     1.000  1.000  1.000
> Cq <- cbind( pp-1970, pp^2-1970^2 )
> head( Cq )
      [,1]      [,2]
[1,] -25.0 -97875.00
[2,] -24.8 -97096.96
[3,] -24.6 -96318.84
[4,] -24.4 -95540.64
[5,] -24.2 -94762.36
[6,] -24.0 -93984.00
> ci.exp( mspq, subset="P" )
exp(Est.)  2.5%  97.5%
```

Multiple time scales and continuous rates (crv-mod) 152 / 220

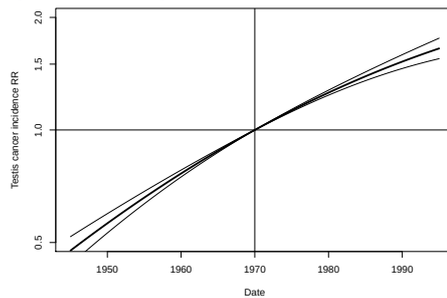
## Period effect



```
> matplot( pp, ci.exp( mspq, subset="P", ctr.mat=Cq ),
+         log="y", ylim=c(0.5,2), xlab="Date",
+         ylab="Testis cancer incidence RR",
+         type="l", lty=1, lwd=c(3,1,1), col="black" )
> abline( h=1, v=1970 )
```

Multiple time scales and continuous rates (crv-mod) 156 / 220

## A quadratic period effect



```
> matplot( pp, ci.exp( mspq, subset="P", ctr.mat=Cq ),
+         log="y", ylim=c(0.5,2), xlab="Date",
+         ylab="Testis cancer incidence RR",
+         type="l", lty=1, lwd=c(3,1,1), col="black" )
> abline( h=1, v=1970 )
```

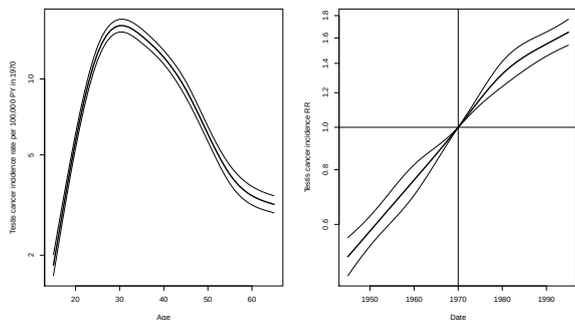
Multiple time scales and continuous rates (crv-mod) 153 / 220

## Period effect

```
> par( mfrow=c(1,2) )
> matplot( aa, ci.pred( msp, newdata=data.frame(A=aa,P=1970,Y=1)
+             log="y", xlab="Age",
+             ylab="Testis cancer incidence rate per 100,000 PY in
+             type="l", lty=1, lwd=c(3,1,1), col="black" )
> matplot( pp, ci.exp( msp, subset="P", ctr.mat=Cp ),
+         log="y", xlab="Date", ylab="Testis cancer incidence R
+         type="l", lty=1, lwd=c(3,1,1), col="black" )
> abline( h=1, v=1970 )
```

Multiple time scales and continuous rates (crv-mod) 157 / 220

## Age and period effect



Multiple time scales and continuous rates (crv-mod)

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## Likelihood for multistate follow-up

Modern Demographic  
Methods in Epidemiology  
with R

23 November 2015

University of Melbourne

<http://BendixCarstensen/AdvCoh/Melb-2015>

ms-lik

## Period effect

```
> par( mfrow=c(1,2) )
> matplot( aa, ci.pred( msp, newdata=data.frame(A=aa,P=1970,Y=1
+ log="y", xlab="Age",
+ ylim=c(2,20), xlim=c(15,65),
+ ylab="Testis cancer incidence rate per 100,000 PY in
+ type="l", lty=1, lwd=c(3,1,1), col="black" )
> matplot( pp, ci.exp( msp, subset="P", ctr.mat=Cp ),
+ log="y", xlab="Date",
+ ylim=c(2,20)/sqrt(2*20), xlim=c(15,65)+1930,
+ ylab="Testis cancer incidence RR",
+ type="l", lty=1, lwd=c(3,1,1), col="black" )
> abline( h=1, v=1970 )
```

Multiple time scales and continuous rates (crv-mod)

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## Likelihood for transition through states

$A \rightarrow B \rightarrow C \rightarrow$

- ▶ given start of observation in **A** at time  $t_0$
- ▶ transitions at times  $t_B$  and  $t_C$
- ▶ survival in **C** till (at least) time  $t_x$ :

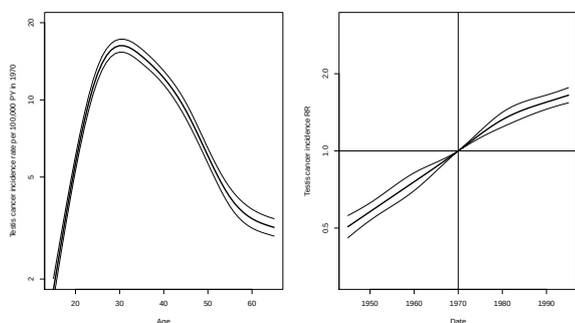
$$L = P\{\text{survive } t_0 \rightarrow t_B \text{ in } \mathbf{A}\} \\ \times P\{\text{transition } \mathbf{A} \rightarrow \mathbf{B} \text{ at } t_B \mid \text{alive in } \mathbf{A}\} \\ \times P\{\text{survive } t_B \rightarrow t_C \text{ in } \mathbf{B} \mid \text{entered } \mathbf{B} \text{ at } t_B\} \\ \times P\{\text{transition } \mathbf{B} \rightarrow \mathbf{C} \text{ at } t_C \mid \text{alive in } \mathbf{B}\} \\ \times P\{\text{survive } t_C \rightarrow t_x \text{ in } \mathbf{C} \mid \text{entered } \mathbf{C} \text{ at } t_C\}$$

- ▶ Product of likelihood contributions for each transition
- each one as for a survival model

Likelihood for multistate follow-up (ms-lik)

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## Age and period effect



Multiple time scales and continuous rates (crv-mod)

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## Likelihood contributions reflected in Lexis object

$$L = P\{\text{survive } t_0 \rightarrow t_B \text{ in } \mathbf{A}\} \\ \times P\{\text{transition } \mathbf{A} \rightarrow \mathbf{B} \text{ at } t_B \mid \text{alive in } \mathbf{A}\} \\ \times P\{\text{survive } t_B \rightarrow t_C \text{ in } \mathbf{B} \mid \text{entered } \mathbf{B} \text{ at } t_B\} \\ \times P\{\text{transition } \mathbf{B} \rightarrow \mathbf{C} \text{ at } t_C \mid \text{alive in } \mathbf{B}\} \\ \times P\{\text{survive } t_C \rightarrow t_x \text{ in } \mathbf{C} \mid \text{entered } \mathbf{C} \text{ at } t_C\}$$

| lex.id | time | lex.dur   | lex.Cst | lex.Xst |
|--------|------|-----------|---------|---------|
| 1      | t_0  | t_B - t_0 | A       | B       |
| 1      | t_B  | t_C - t_B | B       | C       |
| 1      | t_C  | t_x - t_C | C       | C       |

Likelihood for multistate follow-up (ms-lik)

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## Age and period effect with ci.exp

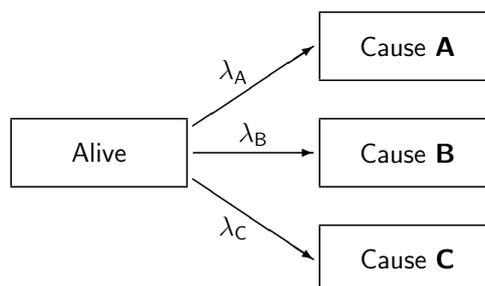
- ▶ In rate models there is always one term with the **rate** dimension — usually **age**
- ▶ But it must refer to a specific **reference** value for **all other** variables (P).
- ▶ **All** parameters must be used in computing rates, at some reference value(s).
- ▶ For the “other” variables, report the RR **relative** to the reference point.
- ▶ Only parameters relevant for the variable (P) used.
- ▶ Contrast matrix is a **difference** between (splines at) the prediction points and the reference point.

Multiple time scales and continuous rates (crv-mod)

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## Competing risks

But you may die from more than one cause  
(or move to more than one state):



Likelihood for multistate follow-up (ms-lik)

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## Cause-specific intensities

$$\lambda_A(t) = \lim_{h \rightarrow 0} \frac{P \{\text{death from cause A in } (t, t+h] \mid \text{alive at } t\}}{h}$$

$$\lambda_B(t) = \lim_{h \rightarrow 0} \frac{P \{\text{death from cause B in } (t, t+h] \mid \text{alive at } t\}}{h}$$

$$\lambda_C(t) = \lim_{h \rightarrow 0} \frac{P \{\text{death from cause C in } (t, t+h] \mid \text{alive at } t\}}{h}$$

Total mortality rate:

$$\lambda_{\text{Total}}(t) = \lim_{h \rightarrow 0} \frac{P \{\text{death from any cause in } (t, t+h] \mid \text{alive at } t\}}{h}$$

Likelihood for multistate follow-up (ms-lik)

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## Components of the likelihood

The log-likelihood is made up of three contributions:

- ▶ one for cause A,
- ▶ one for cause B and
- ▶ one for cause C

**Deaths** are the cause-specific deaths,

but the **person-years** are the **same** in all contributions.

The person-years appear once for each transition **out** of a state.

Likelihood for multistate follow-up (ms-lik)

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## Cause-specific intensities

For small  $h$ ,  $P \{2 \text{ events in } (t, t+h]\} \approx 0$ , so:

$$P \{\text{death from any cause in } (t, t+h] \mid \text{alive at } t\}$$

$$= P \{\text{death from cause A in } (t, t+h] \mid \text{alive at } t\} + P \{\text{death from cause B in } (t, t+h] \mid \text{alive at } t\} + P \{\text{death from cause C in } (t, t+h] \mid \text{alive at } t\}$$

$$\implies \lambda_{\text{Total}}(t) = \lambda_A(t) + \lambda_B(t) + \lambda_C(t)$$

Intensities are additive,

**if** they all refer to the

**same risk set**, in this case "Alive".

Likelihood for multistate follow-up (ms-lik)

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## Likelihood for multiple states

- ▶ **Product** of likelihoods for each transition — each one as for a survival model
- ▶ **conditional** on being alive at (observed) entry to current state
- ▶ **Risk time** is the risk time in the current ("From", `lex.Cst`) state
- ▶ **Events** are transitions to the "To" state (`lex.Xst`)
- ▶ All other transitions out of "From" are treated as **censorings** (but they are not)
- ▶ Fit models separately for each transition or jointly for all

Likelihood for multistate follow-up (ms-lik)

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## Likelihood for competing risks

Data:

$Y$  - person years in "Alive"

$D_A$  - deaths from cause A

$D_B$  - deaths from cause B

$D_C$  - deaths from cause C

Now, assume for a start that transition rates between states are constant.

Likelihood for multistate follow-up (ms-lik)

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## Time varying rates:

- ▶ The same type of analysis as with a constant rates, but data must be
- ▶ split in intervals sufficiently small to justify an assumption of constant rate (intensity),
- ▶ the model should allow for a separate rate for each interval,
- ▶ but constrained to follow model with a smooth effect of the time-scale values allocated to each interval.

Likelihood for multistate follow-up (ms-lik)

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## Likelihood for competing risks

A survivor contributes to the log-likelihood:

$$\log(P \{\text{Survival for a time of } y\}) = -(\lambda_A + \lambda_B + \lambda_C)y$$

A death from cause **A** contributes an additional  $\log(\lambda_A)$ , from cause **B** an additional  $\log(\lambda_B)$  etc.

The total log-likelihood is then:

$$\begin{aligned} \ell(\lambda_A, \lambda_B, \lambda_C) &= D_A \log(\lambda_A) + D_B \log(\lambda_B) + D_C \log(\lambda_C) \\ &\quad - (\lambda_A + \lambda_B + \lambda_C)Y \\ &= [D_A \log(\lambda_A) - \lambda_A Y] + \\ &\quad [D_B \log(\lambda_B) - \lambda_B Y] + \\ &\quad [D_C \log(\lambda_C) - \lambda_C Y] \end{aligned}$$

Likelihood for multistate follow-up (ms-lik)

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## Practical implications

- ▶ Empirical rates  $((d, y)$  from each individual) will be the same for all analyses except for those where deaths occur.
- ▶ Analysis of cause **A**:
  - ▶ Contributions  $(1, y)$  only for those intervals where a cause **A** death occurs.
  - ▶ Intervals with cause **B** or **C** deaths (or no deaths) contribute only  $(0, y)$  treated as censorings.

Likelihood for multistate follow-up (ms-lik)

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| original |      |       |       |     |     |     | expanded |      |    |       |    |
|----------|------|-------|-------|-----|-----|-----|----------|------|----|-------|----|
| id       | time | cause | xx    | d.A | d.B | d.C | id       | time | dd | xx    | Tr |
| 1        | 1    | B     | 0.50  | 0   | 1   | 0   | 1        | 1    | 0  | 0.50  | A  |
| 2        | 1    | NA    | 1.00  | 0   | 0   | 0   | 2        | 1    | 0  | 1.00  | A  |
| 3        | 8    | B     | -1.74 | 0   | 1   | 0   | 3        | 8    | 0  | -1.74 | A  |
| 4        | 3    | A     | -0.55 | 1   | 0   | 0   | 4        | 3    | 1  | -0.55 | A  |
| 5        | 7    | NA    | -0.58 | 0   | 0   | 0   | 5        | 7    | 0  | -0.58 | A  |
| 6        | 7    | C     | -0.04 | 0   | 0   | 1   | 6        | 7    | 0  | -0.04 | A  |
|          |      |       |       |     |     |     | 1        | 1    | 1  | 0.50  | B  |
|          |      |       |       |     |     |     | 2        | 1    | 0  | 1.00  | B  |
|          |      |       |       |     |     |     | 3        | 8    | 1  | -1.74 | B  |
|          |      |       |       |     |     |     | 4        | 3    | 0  | -0.55 | B  |
|          |      |       |       |     |     |     | 5        | 7    | 0  | -0.58 | B  |
|          |      |       |       |     |     |     | 6        | 7    | 0  | -0.04 | B  |
|          |      |       |       |     |     |     | 1        | 1    | 0  | 0.50  | C  |
|          |      |       |       |     |     |     | 2        | 1    | 0  | 1.00  | C  |
|          |      |       |       |     |     |     | 3        | 8    | 0  | -1.74 | C  |
|          |      |       |       |     |     |     | 4        | 3    | 0  | -0.55 | C  |
|          |      |       |       |     |     |     | 5        | 7    | 0  | -0.58 | C  |
|          |      |       |       |     |     |     | 6        | 7    | 1  | -0.04 | C  |

... accomplished by stack.Lexis

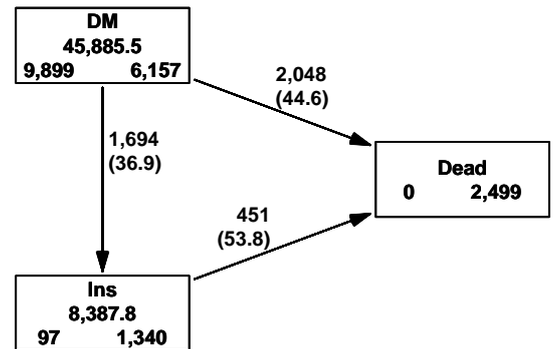
Implemented in the stack.Lexis function:

```
> dmi <- cutLexis( dml, cut = dml$doins,
+                 new.state = "Ins",
+                 precursor = "DM" )
> summary( dmi )
Transitions:
To
From  DM  Ins  Dead  Records:  Events:  Risk time:  Persons:
DM  6157 1694 2048    9899    3742  45885.49  9899
Ins    0 1340  451    1791    451    8387.77  1791
Sum  6157 3034 2499   11690   4193  54273.27  9996
> boxes( dmi, boxpos = list(x=c(20,20,80),
+                           y=c(80,20,50)),
+        scale.R=1000, show.BE=TRUE, hmult=1.2, wmult=1.1 )
```

### Lexis objects (data frame)

- Represents the **follow-up**
- `lex.dur` contains the total time at risk for (any) event
- `lex.Cst` is the state in which this time is spent
- `lex.Xst` is the state to which a transition occurs
  - if no transition, the same as `lex.Cst`.

This is used for modelling of single transitions between states — and multiple transitions with no two originating in the same state.



### stacked.Lexis objects (data frame)

- Represents the **likelihood** contributions
- `lex.dur` contains the total time at risk for (any) event
- `lex.Tr` is the transition to which the record contributes
- `lex.Fail` is the event (failure) indicator for the transition in question.

This is used for joint modelling of **all** transition in a multistate set-up.

Particularly with several rates originating in the **same** state (competing risks).

Implemented in the stack.Lexis function:

```
> options( digits=3, width=200 )
> st.dmi <- stack( dmi )
> print( st.dmi[1:6,], row.names=F )
  Per Age DMdur lex.dur lex.Cst lex.Xst lex.Tr lex.Fail lex.id
1999 58.7 0 11.080 DM DM->Ins FALSE 1
2003 64.1 0 6.689 DM DM->Ins FALSE 2
2005 86.3 0 5.446 DM DM->Ins FALSE 3
2009 44.0 0 0.736 DM DM->Ins FALSE 4
2009 75.8 0 1.344 DM DM->Ins FALSE 5
2008 80.0 0 2.037 DM Dead->Ins FALSE 6
> str( st.dmi )
Classes 'stacked.Lexis' and 'data.frame': 21589 obs. of 16 vari
 $ Per : num 1999 2003 2005 2009 2009 ...
 $ Age : num 58.7 64.1 86.3 44 75.8 ...
 $ DMdur : num 0 0 0 0 0 0 0 ...
 $ lex.dur : num 11.08 6.689 5.446 0.736 1.344 ...
 $ lex.Cst : Factor w/ 3 levels "DM","Ins","Dead": 1 1 1 1 1 1
 $ lex.Xst : Factor w/ 3 levels "DM","Ins","Dead": 1 1 1 1 3 1
 $ lex.Tr : Factor w/ 3 levels "DM->Ins","DM->Dead",...: 1 1 1 1
 $ lex.Fail: logi FALSE FALSE FALSE FALSE FALSE ...
 $ lex.id : int 1 2 3 4 5 6 7 8 9 10 ...
 $ sex : Factor w/ 2 levels "M","F": 2 1 2 2 1 2 1 2 1 ...
 $ dobth : num 1940 1939 1918 1965 1933 ...
 $ dodm : num 1999 2003 2005 2009 2009 ...
```

Implemented in the stack.Lexis function:

```
> library( Epi )
> data(DMLate)
> head(DMLate)
  sex  dobth  dodm  dodth  dooad doins  dox
50185 F 1940.256 1998.917 NA NA NA 2009.997
307563 M 1939.218 2003.309 NA 2007.446 NA 2009.997
294104 F 1918.301 2004.552 NA NA NA 2009.997
336439 F 1965.225 2009.261 NA NA NA 2009.997
245651 M 1932.877 2008.653 NA NA NA 2009.997
216824 F 1927.870 2007.886 2009.923 NA NA 2009.923
> dml <- Lexis( entry = list(Per = dodm,
+                           Age = dodm-dobth,
+                           DMdur = 0 ),
+              exit = list(Per = dox ),
+              exit.status = factor(!is.na(dodth),
+                                 labels=c("DM","Dead")),
+              data = DMLate )
NOTE: entry.status has been set to "DM" for all.
```

Implemented in the stack.Lexis function:

```
> print( subset( dmi, lex.id %in% c(13,15,28) ), row.names=FA
  Per Age DMdur lex.dur lex.Cst lex.Xst lex.Tr lex.Fail lex.
2003 58.1 0.0 2.804 DM Dead 13 M 1938 1997
2005 60.9 2.8 4.643 Ins Ins 15 M 1944 2003
1999 73.7 0.0 8.701 DM Ins 28 F 1925 1999
2007 82.4 8.7 0.977 Ins Dead 28 F 1925 1999
> print( subset( st.dmi, lex.id %in% c(13,15,28) ), row.names=FA
  Per Age DMdur lex.dur lex.Cst lex.Xst lex.Tr lex.Fail lex.
1997 59.4 0.0 0.890 DM Dead DM->Ins FALSE
2003 58.1 0.0 2.804 DM Ins DM->Ins TRUE
1999 73.7 0.0 8.701 DM Ins DM->Ins TRUE
1997 59.4 0.0 0.890 DM Dead DM->Dead TRUE
2003 58.1 0.0 2.804 DM Ins DM->Dead FALSE
1999 73.7 0.0 8.701 DM Ins DM->Dead FALSE
2005 60.9 2.8 4.643 Ins Ins Ins->Dead FALSE
2007 82.4 8.7 0.977 Ins Dead Ins->Dead TRUE
```

## Analysis of rates in multistate models

- ▶ Interactions between all covariates (including time) and state (lex.Cst):  
⇔ separate analyses of all transition rates.
- ▶ Only interaction between state (lex.Cst) and time(scales):  
⇔ same covariate effects for all causes transitions, but separate baseline hazards — “stratified model”.
- ▶ Main effect of state only (lex.Cst):  
⇔ proportional hazards
- ▶ No effect of state:  
⇔ identical baseline hazards — hardly ever relevant.

Likelihood for multistate follow-up (ms-11k)

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## Lifetime risk

Modern Demographic  
Methods in Epidemiology  
with R  
23 November 2015  
University of Melbourne  
<http://BendixCarstensen/AdvCoh/Melb-2015>

DK-lung

## Analysis approaches and data representation

- ▶ Lexis objects represents the precise follow-up in the cohort, in states and along timescales
- ▶ — used for analysis of single transition rates.
- ▶ stacked.Lexis objects represents contributions to the total likelihood
- ▶ — used for joint analysis of (all) rates in a multistate setup
- ▶ . . . which is the case if you want to specify common effects between different transitions.

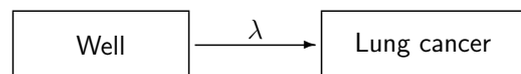
Likelihood for multistate follow-up (ms-11k)

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## Competing risk interpretation

The problems with competing risk models **only** comes when estimated intensities (rates) are used to produce probability statements.

Classical set-up in cancer-registries:



Common statement:

$$P \{ \text{Lung cancer before age 75} \} = 1 - e^{-\Lambda(75)}$$

This is not quite right.

Lifetime risk (DK-lung)

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## Assumptions in competing risks

“Classical” way of looking at survival data: description of the distribution of time to death.

For competing risks that would require three variables:

$T_A$ ,  $T_B$  and  $T_C$ , representing times to death from each of the three causes.

But at most one of these is observed.

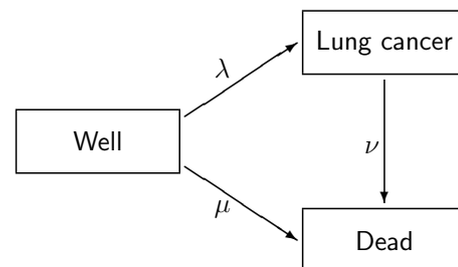
Often it is stated that these must be assumed independent in order to make the likelihood machinery work

1. It is not necessary.
2. Independence can never be assessed from data.

Likelihood for multistate follow-up (ms-11k)

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## How the world really looks



Illness-death model, mortality of lung cancer patients ( $\nu$ ) not relevant here, we only want to find out how many pass through “Lung cancer”

Lifetime risk (DK-lung)

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An account of these problems is given in:

PK Andersen, SZ Abildstrøm & S Rosthøj:

**Competing risks as a multistate model**,  
*Statistical Methods in Medical Research*; **11**, 2002: pp. 203–215

Per Kragh Andersen, Ronald B Geskus, Theo de Witte & Hein Putter:

**Competing risks in epidemiology: possibilities and pitfalls**,

*International Journal of Epidemiology*; 2012: pp. 1–10

Contains examples where both dependent and independent “cause specific survival times” gives rise to the same set of cause specific rates.

Likelihood for multistate follow-up (ms-11k)

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## How many get lung cancer before age $a$ ?

$$P \{ \text{Lung cancer before age 75} \} \neq 1 - e^{-\Lambda(75)}$$

the r.h.s. does not take the possibility of death prior to lung cancer into account.

- ▶  $1 - e^{-\Lambda(75)}$  often stated as the probability of lung cancer before age 75, assuming all other causes of death absent.
- ▶ Lung cancer rates are however observed in a mortal population.
- ▶ If all other causes of death were absent, this would assume that lung “cancer rates” remained the same.

Lifetime risk (DK-lung)

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How it really is:

$P\{\text{Lung cancer diagnosis before age } a\}$

$$\begin{aligned}
 &= \int_0^a P\{\text{Lung cancer at age } u\} du \\
 &= \int_0^a P\{\text{Lung cancer in age } (u, u + du) \mid \text{alive at } u\} \\
 &\quad \times P\{\text{alive at } u \text{ without lung cancer}\} du \\
 &= \int_0^a \lambda(u) \exp\left(-\int_0^u \mu(s) + \lambda(s) ds\right) du
 \end{aligned}$$

R-commands needed to do the calculations:

```

cr.death <- cumsum( D/Y )
cr.lung <- cumsum( lung/Y )
p.simple <- 1 - exp( -cr.lung )
p.lung <- cumsum( lung/Y *
                 exp( -(cr.death+cr.lung) ) )
matlines( age, 100*cbind( cr.lung, p.simple, p.lung ),
          type="l", lty=1, lwd=2*c(2,2,3),
          col=c("black","blue","red") )

```

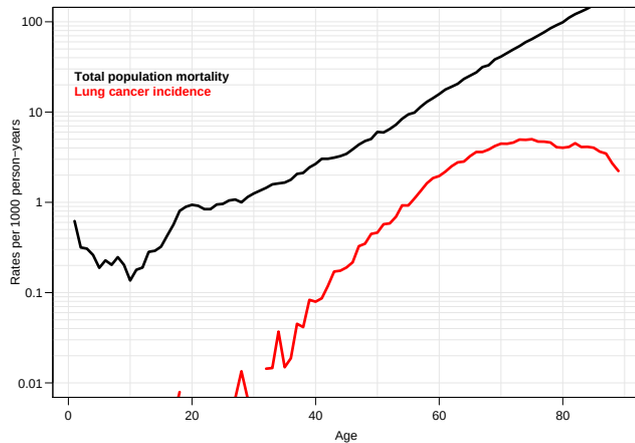
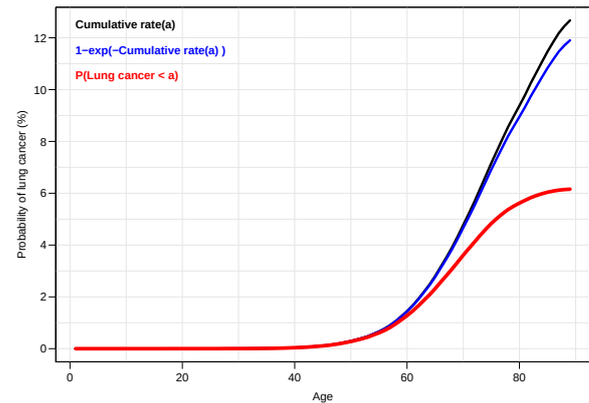
## Probability of lungcancer

The rates are easily plotted for inspection in R:

```

matplot( age, 1000*cbind( D/Y, lung/Y ),
        log="y", type="l", lty=1, lwd=3,
        ylim=c(0.01,100), xlab="Age",
        ylab="Rates per 1000 person-years" )

```



## Assumptions

- ▶ The calculation and the statement “6% of Danish males will get lung cancer” assumes that the lung cancer rates and the mortality rates in the file apply to a cohort of men.
- ▶ But they are cross-sectional rates, so the assumption is one of steady state of:
  1. mortality rates (which is dubious)
  2. lung cancer incidence rates (which is appalling).
- ▶ However, the machinery can be applied to any set of rates for competing risks, regardless of how they were estimated.

The probability that a person contracts lung cancer before age  $a$  is:

$$\begin{aligned}
 &\int_0^a \lambda(u) \exp\left(-\int_0^u \mu(s) + \lambda(s) ds\right) du \\
 &= \int_0^a \lambda(u) \exp\left(-\left(M(u) + \Lambda(u)\right)\right) du
 \end{aligned}$$

$M(u)$  is the cumulative mortality rate.

$\Lambda(u)$  is the cumulative lung cancer incidence rate.

## Life expectancy and life lost

Modern Demographic  
 Methods in Epidemiology  
 with R  
 23 November 2015  
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## Life expectancy

The expected lifetime (at birth) is the variable age ( $a$ ) integrated with respect to the distribution of age at death:

$$EL = \int_0^{\infty} af(a) da$$

where  $f$  is the density of the distribution of lifetimes.

Simplest computed as the area under the survival curve:

$$EL = \int_0^{\infty} S(a) da$$

## Lifetime lost in practice

- ▶ Compute mortality rates at age midpoints of small intervals (1/10 year long, say): 0.05, 0.15, 0.25, ... —  $\lambda(a)$ , lambda
- ▶ Compute the integral by summing  $\lambda(a) \times 0.1$  cumsum(lambda\*0.1) —  $\Lambda(a)$
- ▶ Compute survival function as exp of minus this  $S \leftarrow \exp(-\text{cumsum}(\text{lambda} * 0.1))$
- ▶ Expected life time at age 40, say, is then the integral of the conditional survival:  $\text{sum}(S[400:1000]/S[400]) * 0.1$
- ▶ Compute both for well and dis, and subtract.
- ▶ — now you do the practical. . .

## Life expectancy at age $a$

Use the **conditional** survival function, given alive at age  $a$

$$P(\text{Survive till } t | \text{alive at } a) = S(t)/S(a)$$

Life expectancy at age  $a$ :

$$EL(a) = \int_a^{\infty} S(t)/S(a) dt$$

— the area under the conditional survival function.

## Reporting a multistate model

Modern Demographic  
Methods in Epidemiology  
with R  
23 November 2015  
University of Melbourne  
<http://BendixCarstensen/AdvCoh/Melb-2015>

ms-rep

## Lifetime lost

— due to a disease is the **difference** between the expected residual lifetime for a diseased person and a non-diseased (well) person at the same age:

$$LL(a) = \int_a^{\infty} S_{\text{Well}}(u)/S_{\text{Well}}(a) - S_{\text{Diseased}}(u)/S_{\text{Diseased}}(a) du$$

Note that the survival for a “well” person,  $S_{\text{Well}}(a)$  must be defined:

- ▶ includes the possibility to become diseased (increase mortality)
- ▶ **or** assumes immunity to the disease

## Multistate models

- ▶ Outcomes are transitions between states, with times
- ▶ Covariates are measurements and timescales
- ▶ Models describe the single transition rates
- ▶ Results are:
  - ▶ Description of rates — how do they depend time etc.
  - ▶ Prediction of state occupancy: What is the probability that a person is in a given state at a given time?
- ▶ This illustrates the latter.

## Lifetime lost using rates

- ▶ age-specific mortality rates  $\lambda(a)$
- ▶ survival function  $S(a) = \exp(-\int_0^a \lambda(u) du)$
- ▶ residual lifetime  $EL(a) = \int_a^{\infty} S(u) du$
- ▶ do for “well” and “dis”
- ▶ life lost at age  $a$ :  $LL(a) = EL_{\text{Well}}(a) - EL_{\text{Dis}}(a)$

## Diabetes patient mortality

```
> library(Epi)
> data(DMlate)
> dml <- Lexis( entry = list(Per=dodm, Age=dodm-dobth, DMdur=0 )
+             exit = list(Per=dox),
+             exit.status = factor(!is.na(dodth), labels=c("DM", "Dead"
+             data = DMlate )
```

NOTE: entry.status has been set to "DM" for all.

```
> summary(dml)
```

Transitions:

```
To
From DM Dead Records: Events: Risk time: Persons:
DM 7497 2499 9996 2499 54273.27 9996
```

## ... subdivided by insulin status

Split follow-up at insulin,  
introduce a new timescale and  
split non-precursor states:

```
> dmi <- cutLexis( dml, cut = dml$doins,
+                 pre = "DM",
+                 new.state = "Ins",
+                 new.scale = "t.Ins",
+                 split.states = TRUE )
> summary( dmi )
Transitions:
  To
From DM Ins Dead Dead(Ins) Records: Events: Risk time: Pe
DM 6157 1694 2048 0 9899 3742 45885.49
Ins 0 1340 0 451 1791 451 8387.77
Sum 6157 3034 2048 451 11690 4193 54273.27
> boxes( dmi, boxpos=list(x=c(20,20,80,80),y=c(80,20,80,20)),
+       scale.R=1000, show.BE=TRUE, hmult=1.2, wmult=1.2 )
```

Reporting a multistate model (ms-rep)

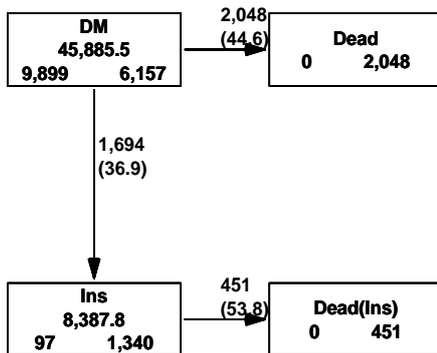
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## Fit Poisson models to transition rates

```
> DM.Ins <- glm( (lex.Xst=="Ins") ~ Ns( Age , knots=ai.kn ) +
+               Ns( DMdur, knots=di.kn ) +
+               I(Per-2000) + sex,
+               family=poisson, offset=log(lex.dur),
+               data = subset(Si,lex.Cst=="DM") )
> DM.Dead <- glm( (lex.Xst=="Dead") ~ Ns( Age , knots=ad.kn ) +
+               Ns( DMdur, knots=dd.kn ) +
+               I(Per-2000) + sex,
+               family=poisson, offset=log(lex.dur),
+               data = subset(Si,lex.Cst=="DM") )
> Ins.Dead <- glm( (lex.Xst=="Dead(Ins)") ~ Ns( Age , knots=ad.
+               Ns( DMdur, knots=dd.
+               Ns( t.Ins, knots=td.
+               I(Per-2000) + sex,
+               family=poisson, offset=log(lex.dur),
+               data = subset(Si,lex.Cst=="Ins") )
```

Reporting a multistate model (ms-rep)

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Reporting a multistate model (ms-rep)

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Put the fitted models into an object representing  
the transitions

```
> Tr <- list( "DM" = list( "Ins" = DM.Ins,
+                         "Dead" = DM.Dead ),
+            "Ins" = list( "Dead(Ins)" = Ins.Dead ) )
> lapply( Tr, names )
$DM
[1] "Ins" "Dead"

$Ins
[1] "Dead(Ins)"
```

Reporting a multistate model (ms-rep)

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Split the follow in 3-month intervals for modelling

```
> Si <- splitLexis( dmi, 0:60/4, "DMdur" )
> summary( Si )
Transitions:
  To
From DM Ins Dead Dead(Ins) Records: Events: Risk time: Pe
DM 184986 1694 2048 0 188728 3742 45885.49
Ins 0 34707 0 451 35158 451 8387.77
Sum 184986 36401 2048 451 223886 4193 54273.27
> summary( dmi )
Transitions:
  To
From DM Ins Dead Dead(Ins) Records: Events: Risk time: Pe
DM 6157 1694 2048 0 9899 3742 45885.49
Ins 0 1340 0 451 1791 451 8387.77
Sum 6157 3034 2048 451 11690 4193 54273.27
```

Reporting a multistate model (ms-rep)

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Define an initial object

— note the combination of `select=` and `NULL`  
which ensures that the relevant attributes from the  
Lexis object `Si` are carried over to `ini` (using  
`Si[NULL, 1:9]` will lose essential attributes)

```
> ini <- subset(Si,select=1:9)[NULL,]
> ini[1:2,"lex.Cst"] <- "DM"
> ini[1:2,"Per"] <- 1995
> ini[1:2,"Age"] <- 60
> ini[1:2,"DMdur"] <- 5
> ini[1:2,"sex"] <- c("M","F")
> ini
lex.id Per Age DMdur t.Ins lex.Cst lex.Xst sex
1 NA 1995 60 5 NA NA DM <NA> M
2 NA 1995 60 5 NA NA DM <NA> F
```

Reporting a multistate model (ms-rep)

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Define knots for spline modelling of the rates:

```
> nk <- 4
> ( ai.kn <- with( subset(Si,lex.Xst=="Ins"),
+                 quantile( Age+lex.dur, probs=(1:nk-0.5)/nk )
+                 12.5% 37.5% 62.5% 87.5%
27.68241 49.61893 61.88364 75.56211
+ ( ad.kn <- with( subset(Si,lex.Xst=="Dead"),
+                 quantile( Age+lex.dur, probs=(1:nk-0.5)/nk )
+                 12.5% 37.5% 62.5% 87.5%
63.61875 74.98700 81.38501 89.26831
+ ( di.kn <- with( subset(Si,lex.Xst=="Ins"),
+                 quantile( DMdur+lex.dur, probs=(1:nk-0.5)/nk
+                 1.50 4.25 7.00 10.50
+ ( dd.kn <- with( subset(Si,lex.Xst=="Dead"),
+                 quantile( DMdur+lex.dur, probs=(1:nk-0.5)/nk
+                 12.5% 37.5% 62.5% 87.5%
0.3778234 1.9582478 4.3370979 8.0232717
+ ( td.kn <- with( subset(Si,lex.Xst=="Dead(Ins)",
+                 quantile( t.Ins+lex.dur, probs=(1:nk-0.5)/nk
+                 12.5% 37.5% 62.5% 87.5%
0.1759069 1.0095825 2.7939767 6.3579740
> library( splines )
```

Reporting a multistate model (ms-rep)

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Simulate 10,000 of each sex using the estimated  
models in `Tr`:

```
> system.time(
+ simL <- simLexis( Tr, ini, time.pts=seq(0,11,0.5), N=10000 ) )
user system elapsed
28.347 0.096 28.441
> summary( simL )
Transitions:
  To
From DM Ins Dead Dead(Ins) Records: Events: Risk time: P
DM 8919 6071 5010 0 20000 11081 150535.86
Ins 0 4328 0 1743 6071 1743 33223.09
Sum 8919 10399 5010 1743 26071 12824 183758.95
> subset( simL, lex.id < 3 )
lex.id Per Age DMdur t.Ins lex.dur lex.Cst lex.
1 1 1995.000 60.00000 5.000000 NA 11.000000 DM
2 2 1995.000 60.00000 5.000000 NA 4.303086 DM
3 2 1999.303 64.30309 9.303086 0 6.696914 Ins
```

Reporting a multistate model (ms-rep)

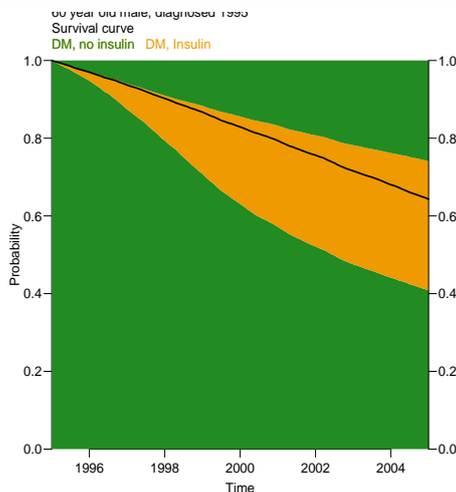
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We now have a dataframe (Lexis object) with simulated follow-up of 10,000 men and 10,000 women.

We then find the number of persons in each state at a specified set of times.

```
> nSt <- nState( subset(simL,sex=="M"),
+               at=seq(0,10,0.1), from=1995, time.scale="Per" )
> nSt
```

| when   | State | DM    | Ins | Dead | Dead(Ins) |
|--------|-------|-------|-----|------|-----------|
| 1995   |       | 10000 | 0   | 0    | 0         |
| 1995.1 |       | 9950  | 18  | 32   | 0         |
| 1995.2 |       | 9900  | 41  | 59   | 0         |
| 1995.3 |       | 9843  | 69  | 86   | 2         |
| 1995.4 |       | 9802  | 80  | 116  | 2         |
| 1995.5 |       | 9757  | 93  | 147  | 3         |
| 1995.6 |       | 9694  | 115 | 187  | 4         |
| 1995.7 |       | 9644  | 137 | 215  | 4         |
| 1995.8 |       | 9589  | 165 | 242  | 4         |
| 1995.9 |       | 9535  | 191 | 269  | 5         |
| 1996   |       | 9479  | 220 | 293  | 8         |
| 1996.1 |       | 9411  | 252 | 323  | 14        |



Show the cumulative prevalences in a different order than that of the state-level ordering and plot them using all defaults:

```
> pp <- pState( nSt, perm=c(1,2,4,3) )
> head( pp )
```

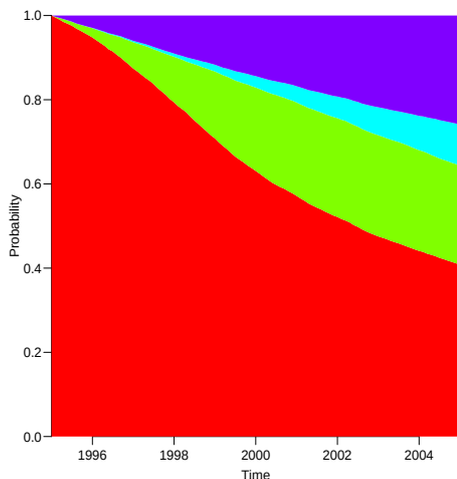
| when   | State | DM     | Ins    | Dead(Ins) | Dead |
|--------|-------|--------|--------|-----------|------|
| 1995   |       | 1.0000 | 1.0000 | 1.0000    | 1    |
| 1995.1 |       | 0.9950 | 0.9968 | 0.9968    | 1    |
| 1995.2 |       | 0.9900 | 0.9941 | 0.9941    | 1    |
| 1995.3 |       | 0.9843 | 0.9912 | 0.9914    | 1    |
| 1995.4 |       | 0.9802 | 0.9882 | 0.9884    | 1    |
| 1995.5 |       | 0.9757 | 0.9850 | 0.9853    | 1    |

```
> plot( pp )
```

We could also use a Cox-model for the mortality rates assuming the two mortality rates to be proportional:

When we fit a Cox-model, `lex.dur` must be used in the `Surv()` function, and the `I()` construction must be used when specifying intermediate states as covariates, since factors with levels not present in the data will create NAs in the parameter vector returned by `coxph`, which in return will crash the simulation machinery.

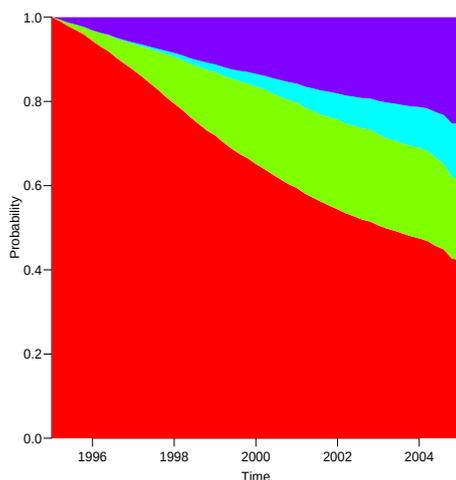
```
> library( survival )
> Cox.Dead <- coxph( Surv( DMdur, DMdur+lex.dur,
+                         lex.Xst %in% c("Dead(Ins)", "Dead") ) ~
+                         Ns( Age-DMdur, knots=ad.kn ) +
+                         I(lex.Cst=="Ins") +
+                         I(Per-2000) + sex,
+                         data = Si )
```

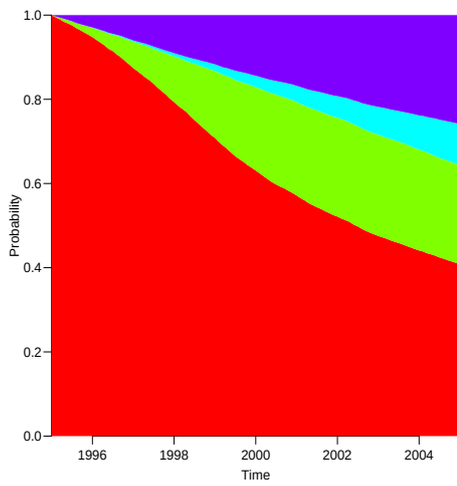


```
> Cr <- list( "DM" = list( "Ins" = DM.Ins,
+                         "Dead" = Cox.Dead ),
+            "Ins" = list( "Dead(Ins)" = Cox.Dead ) )
> simL <- simLexis( Cr, ini, time.pts=seq(0,11,0.2), N=10000 )
> nSt <- nState( subset(simL,sex=="M"),
+               at=seq(0,10,0.2), from=1995, time.scale="Per" )
> pp <- pState( nSt, perm=c(1,2,4,3) )
> plot( pp )
```

We can show the results in an clearer way, buy choosing colors wiser:

```
> clr <- c("orange2","forestgreen")
> par( las=1, mar=c(3,3,3,3) )
> plot( pp, col=clr[c(2,1,1,2)] )
> lines( as.numeric(rownames(pp)), pp[,2], lwd=2 )
> mtext( "60 year old male, diagnosed 1995", side=3, line=2.5, a
> mtext( "Survival curve", side=3, line=1.5, adj=0 )
> mtext( "DM, no insulin DM, Insulin", side=3, line=0.5, adj=0
> mtext( "DM, no insulin", side=3, line=0.5, adj=0, col=clr[2] )
> axis( side=4 )
```





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## References

Now your turn...

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