

Multistate models:

Occurrence rates, cumulative risks, competing risks,
state probabilities with multiple states and time scales in
Register Research with R and Epi::Lexis

Bendix Carstensen Steno Diabetes Center Copenhagen
Herlev, Denmark

<http://BendixCarstensen.com>

Dorte Vistisen Steno Diabetes Center Copenhagen

SDCG, Nuuk, 1-3 March 2022

From /home/bendix/teach/AdvCoh/courses/Nuuk.2022/slides/slides.tex

Monday 28th February, 2022, 21:00

1/ 141

R installed

We will be using the Epi package

```
> library(Epi)
> library(tidyverse)
```

and sometimes also functions from the tidyverse package — but
beware some conflicts exist

Rintro

2/ 141

R is a calculator—use the console

```
> 3 + 2
[1] 5
> x <- 3 * 7
> x
[1] 21
> 5 / 12 -> z
> z
[1] 0.4166667
```

R operates on **objects** — vectors, data frames, models etc.

Rintro

3/ 141

Everything is a function

Fit a regression model

```
> m1 <- lm(y ~ x, data = dd)
```

uses the variables `y` and `x` from the data frame `dd` and saves the result in `m1`, `m1` is an object of class `lm`

But it prints nothing.

You get the result printed by typing the name of the object:

```
> m1  
Call:  
lm(formula = y ~ x, data = dd)  
  
Coefficients:  
(Intercept) x  
-2.6360 0.3888
```

Rintro

4 / 141

A summary method exists for some objects

```
> summary(m1)  
Call:  
lm(formula = y ~ x, data = dd)  
  
Residuals:  
    Min      1Q  Median      3Q     Max  
-0.83379 -0.31168 -0.06909  0.30041  0.87754  
  
Coefficients:  
            Estimate Std. Error t value Pr(>|t|)  
(Intercept) -2.63598   0.21710 -12.14 3.05e-16  
x             0.38882   0.02847  13.66 < 2e-16  
  
Residual standard error: 0.4394 on 48 degrees of freedom  
Multiple R-squared:  0.7954, Adjusted R-squared:  0.7911  
F-statistic: 186.6 on 1 and 48 DF, p-value: < 2.2e-16
```

Rintro

5 / 141

A str method exists for all objects

```
> str(m1)  
List of 12  
 $ coefficients : Named num [1:2] -2.636 0.389  
   ..- attr(*, "names")= chr [1:2] "(Intercept)" "x"  
 $ residuals    : Named num [1:50] 0.597 0.878 0.499 0.112 0.142 ...  
   ..- attr(*, "names")= chr [1:50] "1" "2" "3" "4" ...  
 $ effects      : Named num [1:50] -1.4519 6.0016 0.49 -0.0472 0.1142 ...  
   ..- attr(*, "names")= chr [1:50] "(Intercept)" "x" "" "" ...  
 $ rank         : int 2  
 $ fitted.values: Named num [1:50] -0.5893 0.0796 0.5955 -0.5328 0.4572 ...  
   ..- attr(*, "names")= chr [1:50] "1" "2" "3" "4" ...  
 $ assign        : int [1:2] 0 1  
 $ qr           :List of 5  
   ..$ qr    : num [1:50, 1:2] -7.071 0.141 0.141 0.141 0.141 ...  
   ...- attr(*, "dimnames")=List of 2  
   ... .$. : chr [1:50] "1" "2" "3" "4" ...  
   ... .$. : chr [1:2] "(Intercept)" "x"  
   ...- attr(*, "assign")= int [1:2] 0 1  
   ..$ qraux: num [1:2] 1.14 1  
   ..$ pivot: int [1:2] 1 2  
   #> #> #> #> #>
```

Rintro

6 / 141

When something goes wrong

... you want to see how your data looks — is it as you expected?

Use these functions to see what you have:

```
> summary(dd)
> str(dd)
> dim(dd)
> length(dd)
> names(dd)
> class(dd)
> mode(dd)
```

You can create your own functions

Calculate logit (log-odds) from probability:

```
> p21 <- function(pr)
+   {
+     odds <- log(pr / (1 - pr))
+     odds
+ }
```

function name is `p21` (probability `2 1`ogit)

argument name is `pr`

function value is the value of the last expression in the

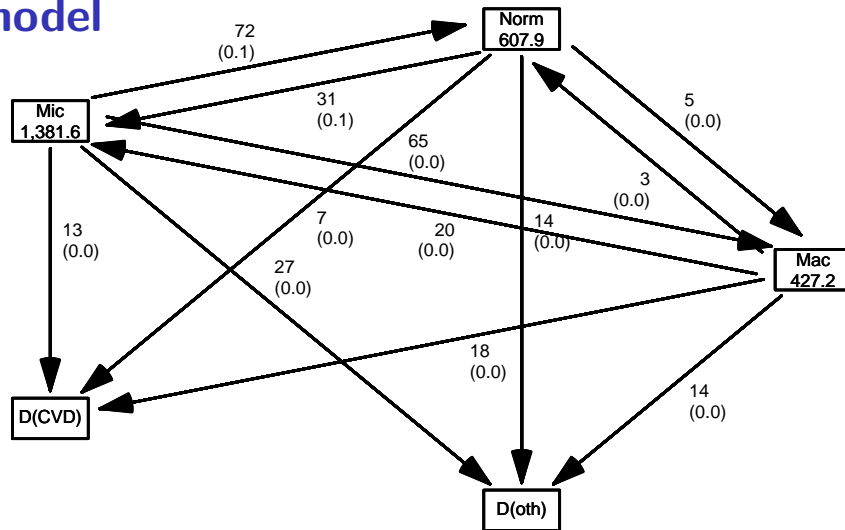
function body (which is what is between the `{}`s):

```
> p21(p = 0.37)
[1] -0.5322168
```

argument value is `0.37`

Your turn — practicals chapter 1!

A multistate model



A multistate model

- ▶ Not really a model
- ▶ What is the data:
 - ▶ Sequence of transitions: (when, from, to)
... same as:
 - ▶ sequence of: (state time, next state)
- ▶ What are the target parameters:
 - ▶ Rates (the arrows)
 - ▶ State probabilities (of being in a state at a given time)
 - ▶ Survival probability
 - ▶ Sojourn times (how long time do you spend in a state)
 - ▶ Probability of ever visiting a state

What is a statistical model

- ▶ Specification of a statistical machinery that could have generated data
- ▶ ... so when we have a statistical model we can simulate a data set
- ▶ The basis for the likelihood of data is the statistical model
⇒ Estimation of parameters in the model
- ▶ Parameter estimates needed for prediction of rates (hazards)

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")

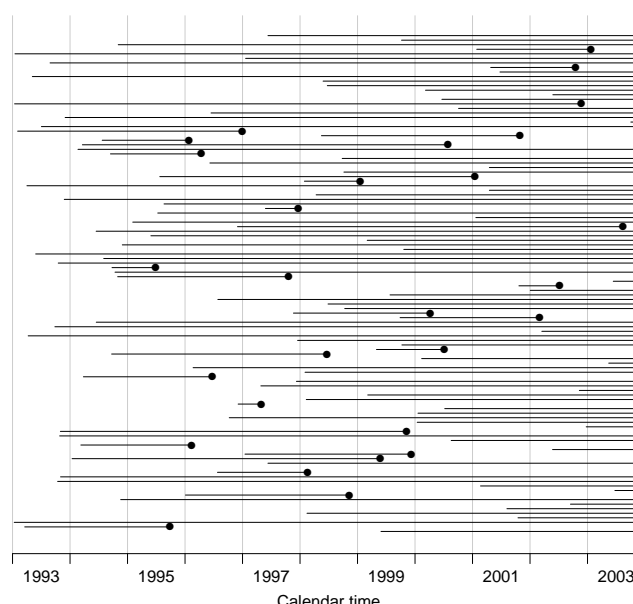
Examples of time-to-event measurements

- ▶ 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

Each line a person

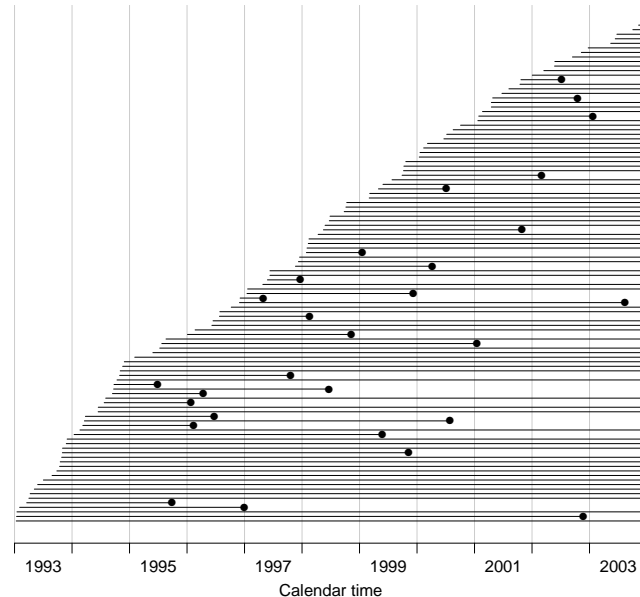
Each blob a death

Study ended at 31
Dec. 2003



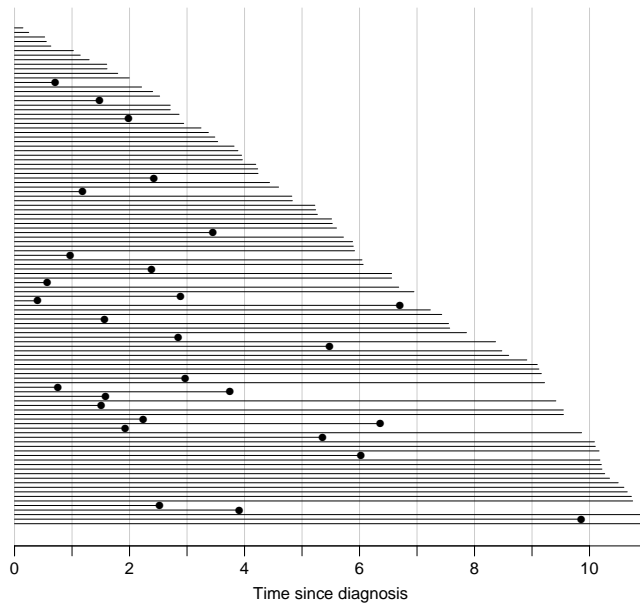
Ordered by date of entry

Most likely the order in your database.



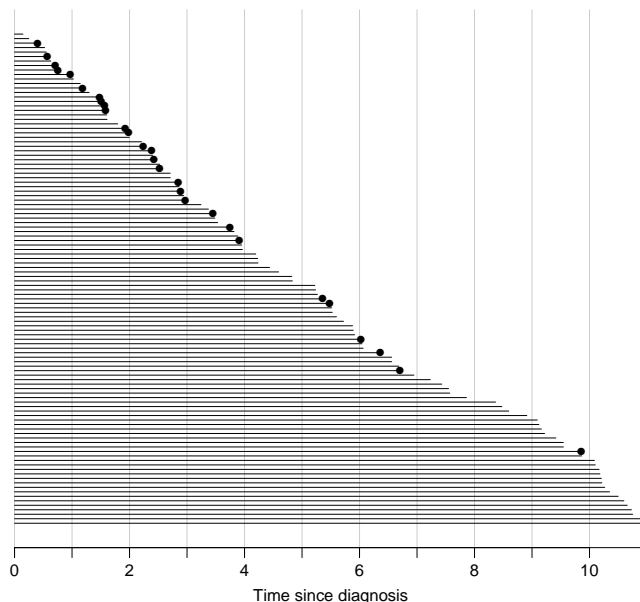
16 / 141

Timescale changed to
“Time since diagnosis”.



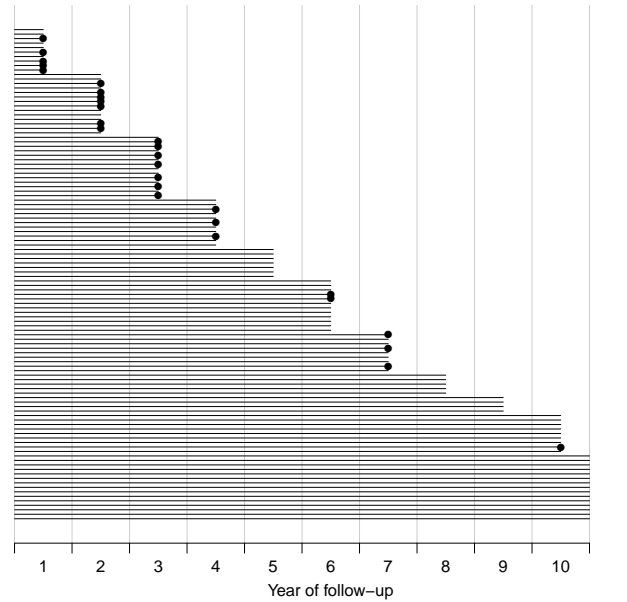
17 / 141

Patients ordered by survival time.



18 / 141

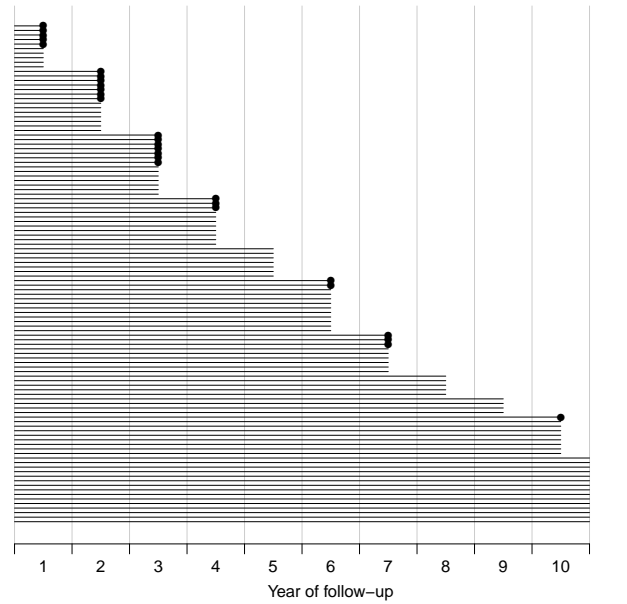
Survival times
grouped into
bands of survival.



surv-rate

19 / 141

Patients ordered
by survival status
within each band.



surv-rate

20 / 141

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 |

Life-table estimator of death probability: $D/(N - L/2)$

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

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

21 / 141

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 |

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

Estimated risk in year 2 for Stage I women is $7/96.5 = 0.0725$

Estimated risk in year 3 for Stage I women is $7/82.5 = 0.0848$

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

Estimated 2 year survival is $0.9535 \times (1 - 0.0725) = 0.8843$

Estimated 3 year survival is $0.8843 \times (1 - 0.0848) = 0.8093$

This is the life-table estimator.

surv-rate

22 / 141

- ▶ No need to use 1 year intervals
- ▶ Very small intervals will leave at most 1 censoring or 1 death in each
- ▶ Interval with 1 death and n_t persons at risk:
 $P\{\text{Death}\} = 1/n_t$
- ▶ corresponding death probability $(n_t - 1)/n_t$
- ▶ if you multiply these over times with event:

$$S(t) = \prod_{t \text{ with event}} (n_t - 1)/n_t$$

... you have the **Kaplan-Meier estimator**

- ▶ looks complicated but just a question of book keeping

surv-rate

23 / 141

Prerequisites

```
> library(Epi)
> library(popEpi)
> # popEpi::splitMulti returns a data.frame rather than a data.table
> options("popEpi.datatable" = FALSE)
```

surv

24 / 141

The lung data set

```
> library(survival)
> data(lung)
> lung$sex <- factor(lung$sex,
+                       levels = 1:2,
+                       labels = c("M", "W"))
> lung$time <- lung$time / (365.25/12)
> head(lung)

  inst      time status age sex ph.ecog ph.karno pat.karno meal.cal wt.loss
1    3 10.053388     2   74   M       1        90       100     1175      NA
2    3 14.948665     2   68   M       0        90       90      1225      15
3    3 33.182752     1   56   M       0        90       90      NA      15
4    5  6.899384     2   57   M       1        90       60      1150      11
5    1 29.010267     2   60   M       0       100       90      NA      0
6   12 33.577002     1   74   M       1        50       80      513      0
```

surv

25/ 141

Survival function

- ▶ Use `survfit` to construct the Kaplan-Meier estimator of overall survival:

```
> ?Surv
> ?survfit

> km <- survfit(Surv(time, status == 2) ~ 1, data = lung)
> km
Call: survfit(formula = Surv(time, status == 2) ~ 1, data = lung)

      n  events  median 0.95LCL 0.95UCL
228.00 165.00   10.18    9.36   11.93
> # summary(km) # very long output
```

surv

26/ 141

We can plot the survival curve—this is the default plot for a `survfit` object:

```
> plot(km)
```

What is the median survival? What does it mean? Explore if survival patterns between men and women are different:

```
> kms <- survfit(Surv(time, status == 2) ~ sex, data = lung)
> kms
Call: survfit(formula = Surv(time, status == 2) ~ sex, data = lung)

      n  events median 0.95LCL 0.95UCL
sex=M 138     112    8.87    6.97   10.2
sex=W  90      53   14.00   11.43   18.1
```

surv

27/ 141

We see that men have worse survival than women, but they are also a bit older (`age` is age at diagnosis of lung cancer):

```
> with(lung, tapply(age, sex, mean))
```

| M | W |
|----------|----------|
| 63.34058 | 61.07778 |

Formally there is a significant difference in survival between men and women

```
> survdiff(Surv(time, status==2) ~ sex, data = lung)
```

Call:

```
survdiff(formula = Surv(time, status == 2) ~ sex, data = lung)
```

| | N | Observed | Expected | $(O-E)^2/E$ | $(O-E)^2/V$ |
|-------|-----|----------|----------|-------------|-------------|
| sex=M | 138 | 112 | 91.6 | 4.55 | 10.3 |
| sex=W | 90 | 53 | 73.4 | 5.68 | 10.3 |

Chisq= 10.3 on 1 degrees of freedom, p= 0.001

surv

28 / 141

Rates and rate-ratios

- ▶ Occurrence **rate**:

$$\lambda(t) = \lim_{h \rightarrow 0} P\{\text{event in } (t, t+h] \mid \text{alive at } t\} / h$$

—measured in probability per time: time⁻¹

- ▶ observation in a survival study: (exit status, time alive)
- ▶ empirical rate $(d, y) = (\text{deaths}, \text{time})$
- ▶ the Cox model is a model for rates as function of time (t) and covariates (x_1, x_2) :

$$\lambda(t, x) = \lambda_0(t) \exp(\beta_1 x_1 + \beta_2 x_2)$$

—mortality depends on the person's sex and age, say.

- ▶ Data looks like data for a K-M analysis **plus** covariate values

surv

29 / 141

Rates and rate-ratios: Simple Cox model

Now explore how sex and age (at diagnosis) influence the mortality—note that in a Cox-model we are addressing the mortality rate and not the survival:

```
> c0 <- coxph(Surv(time, status == 2) ~ sex, data = lung)
> c1 <- coxph(Surv(time, status == 2) ~ sex + age, data = lung)
> summary(c1)
> ci.exp(c0)
> ci.exp(c1)
```

What variables from `lung` are we using?

surv

30 / 141

```

> c0 <- coxph(Surv(time, status == 2) ~ sex      , data = lung)
> c1 <- coxph(Surv(time, status == 2) ~ sex + age, data = lung)
> summary(c1)

Call:
coxph(formula = Surv(time, status == 2) ~ sex + age, data = lung)

n= 228, number of events= 165

      coef exp(coef)   se(coef)      z Pr(>|z|)
sexW -0.513219  0.598566  0.167458 -3.065  0.00218
age   0.017045  1.017191  0.009223  1.848  0.06459

      exp(coef) exp(-coef) lower .95 upper .95
sexW    0.5986   1.6707    0.4311   0.8311
age     1.0172    0.9831    0.9990   1.0357

Concordance= 0.603  (se = 0.025 )
Likelihood ratio test= 14.12  on 2 df,  p=9e-04
Wald test           = 13.47  on 2 df,  p=0.001
Score (logrank) test = 13.72  on 2 df,  p=0.001

```

surv

31/ 141

```

> ci.exp(c0)

      exp(Est.)    2.5%    97.5%
sexW 0.5880028 0.4237178 0.8159848

> ci.exp(c1)

      exp(Est.)    2.5%    97.5%
sexW 0.598566 0.4310936 0.8310985
age   1.017191 0.9989686 1.0357467

```

What do these estimates mean?

$$\lambda(t, x) = \lambda_0(t)\exp(\beta_1 x_1 + \beta_2 x_2)$$

Where is β_1 ? Where is β_2 ? Where is $\lambda_0(t)$?

What is the mortality RR for a 10 year age difference?

surv

32/ 141

If mortality is assumed constant ($\lambda(t) = \lambda$), then the likelihood for the Cox-model is equivalent to a Poisson likelihood, which can be fitted using the `poisreg` family from the `Epi` package:

```

> ?poisreg

> p1 <- glm(cbind(status == 2, time) ~ sex + age,
+             family = poisreg,
+             data = lung)
> ci.exp(p1) # Poisson

      exp(Est.)    2.5%    97.5%
(Intercept) 0.03255152 0.01029228 0.1029511
sexW        0.61820515 0.44555636 0.85777537
age         1.01574132 0.99777446 1.0340317

> ci.exp(c1) # Cox

      exp(Est.)    2.5%    97.5%
sexW    0.598566 0.4310936 0.8310985
age     1.017191 0.9989686 1.0357467

```

surv

33/ 141

Sex and age effects are quite close between the Poisson and the Cox models.

Poisson model has an intercept term, the estimate of the (assumed) constant underlying mortality.

The risk time part of the response (second argument in the `cbind`) was entered in units of months (remember we rescaled in the beginning?), the (`Intercept`) (taken from the `ci.exp`) is a rate per 1 person-month.

What age and sex does the (`Intercept`) refer to?

```
> ci.exp(p1) # Poisson
            exp(Est.)    2.5%    97.5%
(Intercept) 0.03255152 0.01029228 0.1029511
sexW        0.61820515 0.44555636 0.8577537
age         1.01574132 0.99777446 1.0340317
```

surv

34 / 141

poisreg and poisson

poisreg: `cbind(d,y) ~ ...`

```
> p1 <- glm(cbind(status == 2, time) ~ sex + age,
+             family = poisreg,
+             data = lung)
```

poisson: `d ~ ... + offset(log(y))`

```
> px <- glm(status == 2 ~ sex + age + offset(log(time)),
+             family = poisson,
+             data = lung)
> ## or:
> px <- glm(status == 2 ~ sex + age,
+             offset = log(time),
+             family = poisson,
+             data = lung)
```

surv

35 / 141

Representation of follow-up: Lexis object

```
> L1 <- Lexis(exit = list(tfl = time),
+               exit.status = factor(status,
+                                     levels = 1:2,
+                                     labels = c("Alive", "Dead")),
+               data = lung)
```

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

NOTE: entry is assumed to be 0 on the tfl timescale.

```
> head(L1)
```

| | tfl | lex.dur | lex.Cst | lex.Xst | lex.id | inst | time | status | age | sex | ph.ecog | ph.karnofsky |
|---|-----|-----------|---------|---------|--------|------|-----------|--------|-----|-----|---------|--------------|
| 1 | 0 | 10.053388 | Alive | Dead | 1 | 3 | 10.053388 | 2 | 74 | M | 1 | |
| 2 | 0 | 14.948665 | Alive | Dead | 2 | 3 | 14.948665 | 2 | 68 | M | 0 | |
| 3 | 0 | 33.182752 | Alive | Alive | 3 | 3 | 33.182752 | 1 | 56 | M | 0 | |
| 4 | 0 | 6.899384 | Alive | Dead | 4 | 5 | 6.899384 | 2 | 57 | M | 1 | |
| 5 | 0 | 29.010267 | Alive | Dead | 5 | 1 | 29.010267 | 2 | 60 | M | 0 | |
| 6 | 0 | 33.577002 | Alive | Alive | 6 | 12 | 33.577002 | 1 | 74 | M | 1 | |

meal.cal wt.loss

| | meal.cal | wt.loss |
|---|----------|---------|
| 1 | 1175 | NA |
| 2 | 1225 | 15 |
| 3 | NA | 15 |

surv

36 / 141

New variables in a Lexis object

- `tfl`: time from lung cancer **at the time of entry**, therefore it is 0 for all persons; the entry time is 0 from the entry time. But it defines a **timescale**.
- `lex.dur`: the **length** of time a person is in state `lex.Cst`, here measured in months, because `time` is.
- `lex.Cst`: Current **state**, the state in which the `lex.dur` time is spent.
- `lex.Xst`: eXit **state**, the state to which the person moves after the `lex.dur` time in `lex.Cst`.
- `lex.id`: an id of each record in the source dataset. Can be explicitly set by `id=`.

surv

37 / 141

Lexis object: Overview of follow-up

Overkill?

The point is that the machinery generalizes to multistate data.

```
> summary(L1)
```

Transitions:

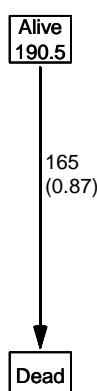
| To | From | Alive | Dead | Records: | Events: | Risk time: | Persons: |
|-------|-------|-------|------|----------|---------|------------|----------|
| Alive | Alive | 63 | 165 | 228 | 165 | 2286.42 | 228 |

What is the average follow-up time for persons?

surv

38 / 141

```
> boxes(L1, boxpos = TRUE, scale.Y = 12, digits.R = 2)
```



Explain the numbers in the graph.

surv

39 / 141

Cox model using the Lexis-specific variables:

```
> cl <- coxph(Surv(tfl,
+                     tfl + lex.dur,
+                     lex.Xst == "Dead") ~ sex + age,
+                     data = Ll)

Surv(from-time, to-time, event indicator)
```

Using the Lexis features:

```
> cL <- coxph.Lexis(Ll, tfl ~ sex + age)

survival::coxph analysis of Lexis object Ll:
Rates for the transition Alive->Dead
Baseline timescale: tfl

> round(cbind(ci.exp(cL),
+             ci.exp(cL)), 3)

exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5%
sexW      0.599 0.431 0.831      0.599 0.431 0.831
age       1.017 0.999 1.036      1.017 0.999 1.036
```

surv 40 / 141

The crude Poisson model:

```
> pc <- glm(cbind(lex.Xst == "Dead", lex.dur) ~ sex + age,
+             family = poisreg,
+             data = Ll)
```

or even simpler, by using the Lexis features:

```
> pL <- glm.Lexis(Ll, ~ sex + age)

stats::glm Poisson analysis of Lexis object Ll with log link:
Rates for the transition: Alive->Dead

> round(cbind(ci.exp(pL),
+             ci.exp(pc)), 3)

exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5%
(Intercept) 0.033 0.010 0.103      0.033 0.010 0.103
sexW        0.618 0.446 0.858      0.618 0.446 0.858
age         1.016 0.998 1.034      1.016 0.998 1.034
```

surv

41 / 141

Poisson and Cox model

The crude Poisson model is a Cox-model with the (quite brutal) assumption that baseline rate is constant over time.

But results are similar:

```
> round(cbind(ci.exp(cL),
+             ci.exp(pL)[-1,]), 3)

exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5%
sexW      0.599 0.431 0.831      0.618 0.446 0.858
age       1.017 0.999 1.036      1.016 0.998 1.034
```

surv

42 / 141

Likelihood and records

Suppose a person is alive from t_e (entry) to t_x (exit) and that the person's status at t_x is d , where $d = 0$ means alive and $d = 1$ means dead. If we choose, say, two time points, t_1, t_2 between t_e and t_x , standard use of conditional probability (formally, repeated use of Bayes' formula) gives

$$\begin{aligned} P\{d \text{ at } t_x \mid \text{entry at } t_e\} &= P\{\text{survive } (t_e, t_1] \mid \text{alive at } t_e\} \times \\ &\quad P\{\text{survive } (t_1, t_2] \mid \text{alive at } t_1\} \times \\ &\quad P\{\text{survive } (t_2, t_x] \mid \text{alive at } t_2\} \times \\ &\quad P\{d \text{ at } t_x \mid \text{alive at } t_3\} \end{aligned}$$

Rates and likelihood

For a start assume that the mortality is constant over time $\lambda(t) = \lambda$:

$$\begin{aligned} P\{\text{death during } (t, t+h]\} &\approx \lambda h & (1) \\ \Rightarrow P\{\text{survive } (t, t+h]\} &\approx 1 - \lambda h \end{aligned}$$

where the approximation gets better the smaller h is.

Dividing follow-up time

- ▶ Survival for a time span: $y = t_x - t_e$
- ▶ Subdivided in N intervals, each of length $h = y/N$
- ▶ Survival probability for the entire span from t_e to t_x is the **product** of probabilities of surviving each of the small intervals, conditional on being alive at the beginning each interval:

$$P\{\text{survive } t_e \text{ to } t_x\} \approx (1 - \lambda h)^N = \left(1 - \frac{\lambda y}{N}\right)^N$$

Dividing follow-up time

- ▶ From mathematics it is known that $(1 + x/n)^n \rightarrow \exp(x)$ as $n \rightarrow \infty$ (some define $\exp(x)$ this way).
- ▶ So if we divide the time span y in small pieces we will have that $N \rightarrow \infty$:

$$P\{\text{survive } t_e \text{ to } t_x\} \approx \left(1 - \frac{\lambda y}{N}\right)^N \rightarrow \exp(-\lambda y), \quad N \rightarrow \infty \quad (2)$$

- ▶ The contribution to the likelihood from a person observed for a time span of length y is $\exp(-\lambda y)$, and the contribution to the log-likelihood is therefore $-\lambda y$.

Dividing follow-up time

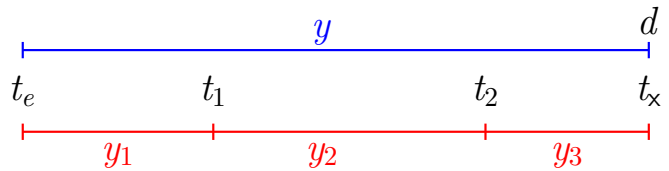
- ▶ A person dying at the end of the last interval, the contribution to the likelihood from the last interval will be
- ▶ the probability surviving till just before the end of the interval,
- ▶ **multiplied by**
- ▶ the probability of dying in the last tiny instant (of length ϵ) of the interval
- ▶ The probability of dying in this tiny instant is $\lambda\epsilon$
- ▶ log-likelihood contribution from this last instant is $\log(\lambda\epsilon) = \log(\lambda) + \log(\epsilon)$.

Total likelihood

The total likelihood for one person is the product of all these terms from the follow-up intervals (i) for the person; and the log-likelihood (ℓ) is therefore:

$$\begin{aligned}\ell(\lambda) &= -\lambda \sum_i y_i + \sum_i d_i \log(\lambda) + \sum_i d_i \log(\epsilon) \\ &= \sum_i (d_i \log(\lambda) - \lambda y_i) + \sum_i d_i \log(\epsilon)\end{aligned}$$

The last term does not depend on λ , so can be ignored



Probability

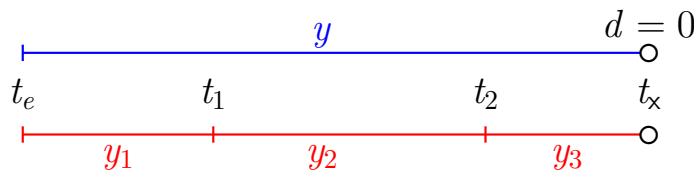
$$P(d \text{ at } t_x | \text{entry } t_e)$$

$$\begin{aligned} &= P(\text{surv } t_e \rightarrow t_1 | \text{entry } t_e) \\ &\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1) \\ &\times P(d \text{ at } t_x | \text{entry } t_2) \end{aligned}$$

log-Likelihood

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

$$\begin{aligned} &= 0 \log(\lambda) - \lambda y_1 \\ &+ 0 \log(\lambda) - \lambda y_2 \\ &+ d \log(\lambda) - \lambda y_3 \end{aligned}$$



Probability

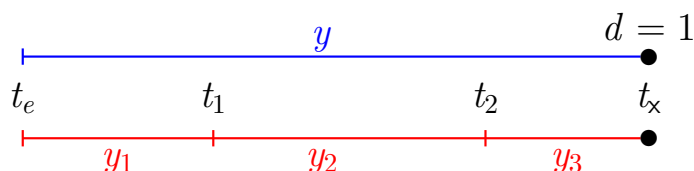
$$P(\text{surv } t_e \rightarrow t_x | \text{entry } t_e)$$

$$\begin{aligned} &= P(\text{surv } t_e \rightarrow t_1 | \text{entry } t_e) \\ &\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1) \\ &\times P(\text{surv } t_2 \rightarrow t_x | \text{entry } t_2) \end{aligned}$$

log-Likelihood

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

$$\begin{aligned} &= 0 \log(\lambda) - \lambda y_1 \\ &+ 0 \log(\lambda) - \lambda y_2 \\ &+ 0 \log(\lambda) - \lambda y_3 \end{aligned}$$



Probability

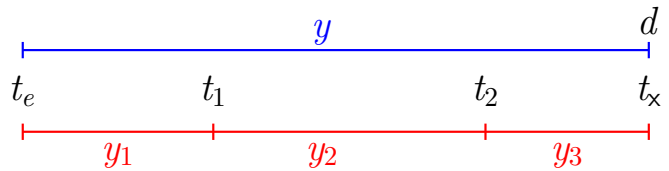
$$P(\text{event at } t_x | \text{entry } t_e)$$

$$\begin{aligned} &= P(\text{surv } t_e \rightarrow t_1 | \text{entry } t_e) \\ &\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1) \\ &\times P(\text{event at } t_x | \text{entry } t_2) \end{aligned}$$

log-Likelihood

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

$$\begin{aligned} &= 0 \log(\lambda) - \lambda y_1 \\ &+ 0 \log(\lambda) - \lambda y_2 \\ &+ 1 \log(\lambda) - \lambda y_3 \end{aligned}$$



Probability

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

$$\begin{aligned} &= P(\text{surv } t_e \rightarrow t_1 | \text{entry } t_e) \\ &\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1) \\ &\times P(d \text{ at } t_x | \text{entry } t_2) \end{aligned}$$

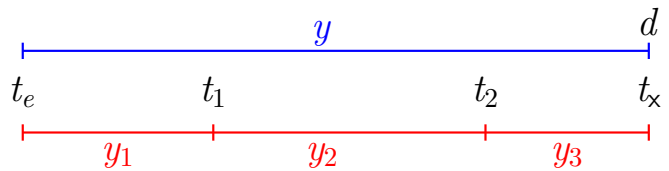
log-Likelihood

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

$$\begin{aligned} &= 0 \log(\lambda) - \lambda y_1 \\ &+ 0 \log(\lambda) - \lambda y_2 \\ &+ d \log(\lambda) - \lambda y_3 \end{aligned}$$

xsurv

52 / 141



Probability

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

$$\begin{aligned} &= P(\text{surv } t_e \rightarrow t_1 | \text{entry } t_e) \\ &\times P(\text{surv } t_1 \rightarrow t_2 | \text{entry } t_1) \\ &\times P(d \text{ at } t_x | \text{entry } t_2) \end{aligned}$$

log-Likelihood

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

$$\begin{aligned} &= 0 \log(\lambda_1) - \lambda_1 y_1 \\ &+ 0 \log(\lambda_2) - \lambda_2 y_2 \\ &+ d \log(\lambda_3) - \lambda_3 y_3 \end{aligned}$$

— allows different rates (λ_i) in each interval

xsurv

53 / 141

Baseline hazard: splitting time

```
> S1 <- splitMulti(L1, tf1 = 0:36)
> summary(L1)

Transitions:
  To
From   Alive Dead  Records:  Events: Risk time: Persons:
    Alive    63   165      228      165    2286.42     228

> summary(S1)

Transitions:
  To
From   Alive Dead  Records:  Events: Risk time: Persons:
    Alive   2234   165     2399      165    2286.42     228
```

What happened to no. records?

What happened to amount of risk time?

What happened to no. events?

surv

54 / 141

```

> wh <- names(L1)[1:10] # names of variables in some order
> subset(L1, lex.id == 10) [,wh]

    tfl  lex.dur lex.Cst lex.Xst lex.id inst      time status age sex
10   0 5.453799   Alive     Dead     10      7 5.453799      2 61   M

> subset(S1, lex.id == 10) [,wh]

    tfl  lex.dur lex.Cst lex.Xst lex.id inst      time status age sex
163  0 1.0000000   Alive     Alive     10      7 5.453799      2 61   M
164  1 1.0000000   Alive     Alive     10      7 5.453799      2 61   M
165  2 1.0000000   Alive     Alive     10      7 5.453799      2 61   M
166  3 1.0000000   Alive     Alive     10      7 5.453799      2 61   M
167  4 1.0000000   Alive     Alive     10      7 5.453799      2 61   M
168  5 0.4537988   Alive     Dead     10      7 5.453799      2 61   M

```

In `S1` each record now represents a small interval of follow-up for a person, so each person has many records.

surv

55 / 141

Natural splines for baseline hazard

```

> ps <- glm(cbind(lex.Xst == "Dead", lex.dur)
+             ~ Ns(tfl, knots = seq(0, 36, 12)) + sex + age,
+             family = poisreg,
+             data = S1)

```

or even simpler:

```

> ps <- glm.Lexis(S1, ~ Ns(tfl, knots = seq(0, 36, 12)) + sex + age)

stats::glm Poisson analysis of Lexis object S1 with log link:
Rates for the transition: Alive->Dead

> ci.exp(ps)

            exp(Est.)      2.5%      97.5%
(Intercept) 0.0189837 0.005700814 0.06321569
Ns(tfl, knots = seq(0, 36, 12))1 2.4038681 0.809442081 7.13896863
Ns(tfl, knots = seq(0, 36, 12))2 4.1500822 0.436273089 39.47798357
Ns(tfl, knots = seq(0, 36, 12))3 0.8398973 0.043928614 16.05849662
sexW        0.5987171 0.431232662 0.83124998
age         1.0165872 0.998377104 1.03512945

```

surv

56 / 141

Comparing with estimates from the Cox-model and from the model with constant baseline:

```

> round(cbind(ci.exp(cl),
+             ci.exp(ps, subset = c("sex", "age")),
+             ci.exp(pc, subset = c("sex", "age"))), 3)

            exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5%
sexW      0.599 0.431 0.831      0.599 0.431 0.831      0.618 0.446 0.858
age       1.017 0.999 1.036      1.017 0.998 1.035      1.016 0.998 1.034

```

surv

57 / 141

But where is the baseline hazard?

`ps` is a model for the hazard so we can predict the value of it at defined values for the covariates in the model:

```
> prf <- data.frame(tfl = seq(0, 30, 0.2),
+                      sex = "W",
+                      age = 60)
```

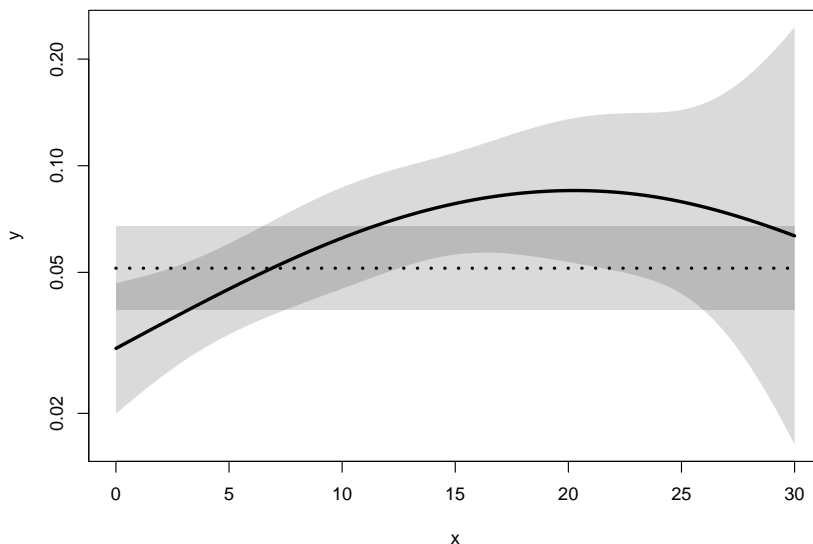
We can over-plot with the predicted rates from the model where mortality rates are constant, the only change is the model (`pc` instead of `ps`):

```
> matshade(prf$tfl, ci.pred(ps, prf),
+            plot = TRUE, log = "y", lwd = 3)
> matshade(prf$tfl, ci.pred(pc, prf), lty = 3, lwd = 3)
```

surv

58 / 141

Here is the baseline hazard!



surv What are the units on the y -axis? Describe the mortality rates

59 / 141

Survival function and hazard function

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

Simple, but the CI for $S(t)$ not so simple...

Implemented in the `ci.surv` function

Arguments: 1:model, 2:prediction data frame, 3:equidistance

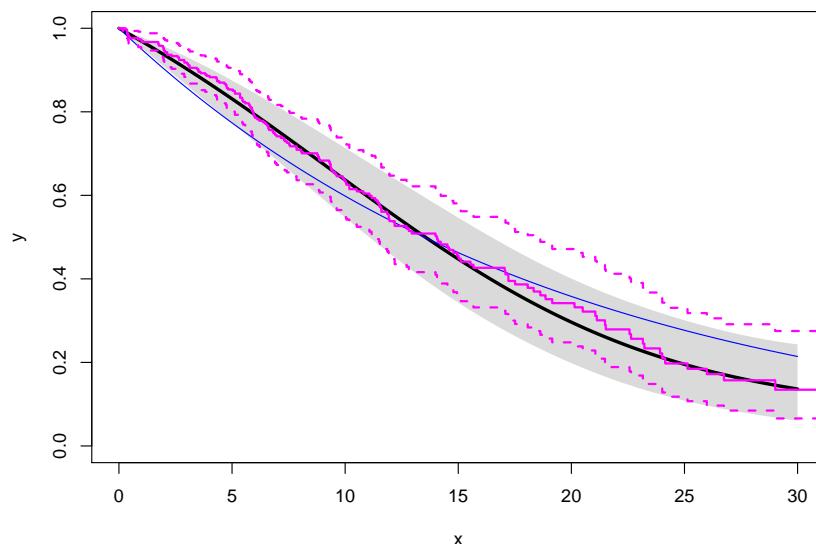
Prediction data frame must correspond to a sequence of equidistant time points:

```
> matshade(prf$tfl, ci.surv(ps, prf, intl = 0.2),
+            plot = TRUE, ylim = 0:1, lwd = 3)
> lines(prf$tfl, ci.surv(pc, prf, intl = 0.2)[,1], col="blue")
> lines(survfit(c1, newdata = data.frame(sex = "W", age = 60)),
+        lwd = 2, lty = 1, col="magenta")
```

surv

60 / 141

Survival functions



surv

61 / 141

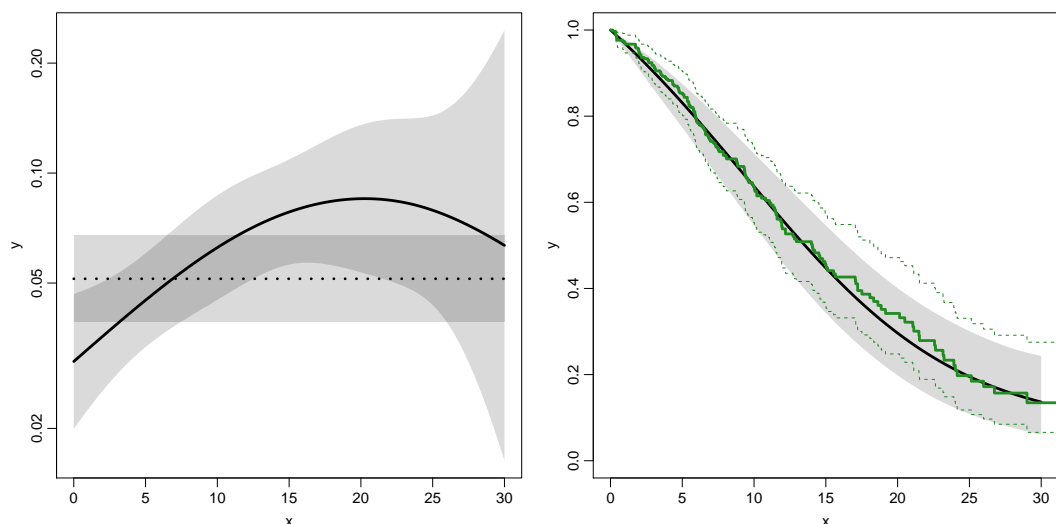
Hazard and survival functions

```
> par(mfrow = c(1,2), mar=c(3,3,1,1), mgp=c(3,1,0)/1.6)
> #
> # hazard scale
> matshade(prf$tfl, ci.pred(ps, prf),
+           plot = TRUE, log = "y", lwd = 3)
> matshade(prf$tfl, ci.pred(pc, prf), lty = 3, lwd = 3)
> #
> # survival
> matshade(prf$tfl, ci.surv(ps, prf, intl = 0.2),
+           plot = TRUE, ylim = 0:1, lwd = 3)
> lines(survfit(c1, newdata = data.frame(sex = "W", age = 60)),
+        col = "forestgreen", lwd = 3, conf.int = FALSE)
> lines(survfit(c1, newdata = data.frame(sex = "W", age = 60)),
+        col = "forestgreen", lwd = 1, lty = 1)
```

surv

62 / 141

Hazard and survival functions



surv

63 / 141

K-M estimator and smooth Poisson model

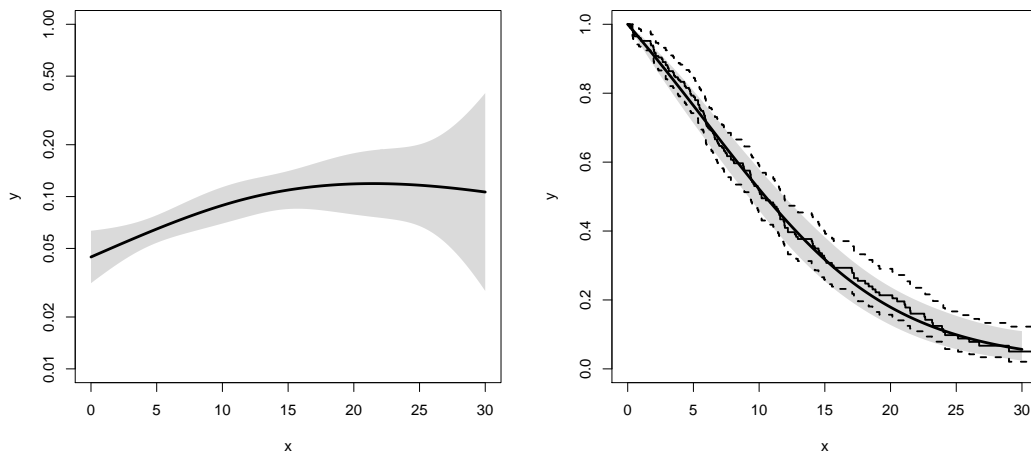
Kaplan-Meier estimator and compared to survival from corresponding Poisson-model, which is one with time (`tfl`) as the only covariate:

```
> par(mfrow=c(1,2))
> pk <- glm(cbind(lex.Xst == "Dead",
+                   lex.dur) ~ Ns(tfl, knots = seq(0, 36, 12)),
+                   family = poisreg,
+                   data = S1)
> # hazard
> matshade(prf$tfl, ci.pred(pk, prf),
+            plot = TRUE, log = "y", lwd = 3, ylim = c(0.01,1))
> # survival from smooth model
> matshade(prf$tfl, ci.surv(pk, prf, intl = 0.2) ,
+            plot = TRUE, lwd = 3, ylim = 0:1)
> # K-M estimator
> lines(km, lwd = 2)
```

surv

64 / 141

K-M estimator and smooth Poisson model



surv

65 / 141

K-M estimator and smooth Poisson model

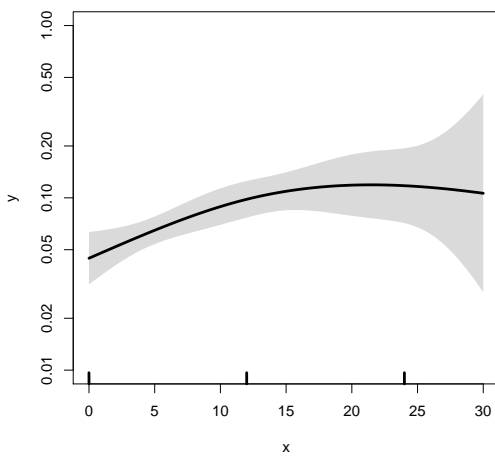
We can explore how the tightness of the knots in the smooth model influence the underlying hazard and the resulting survival function:

```
> zz <- function(dk) # distance between knots
+ {
+   par(mfrow=c(1,2))
+   kn <- seq(0, 36, dk)
+   pk <- glm(cbind(lex.Xst == "Dead",
+                   lex.dur) ~ Ns(tfl, knots = kn),
+                   family = poisreg,
+                   data = S1)
+   matshade(prf$tfl, ci.pred(pk, prf),
+             plot = TRUE, log = "y", lwd = 3, ylim = c(0.01,1))
+   rug(kn, lwd=3)
+
+   matshade(prf$tfl, ci.surv(pk, prf, intl = 0.2) ,
+             plot = TRUE, lwd = 3, ylim = 0:1)
+   lines(km, lwd = 2, col = "forestgreen")
+ }
> zz(12)
```

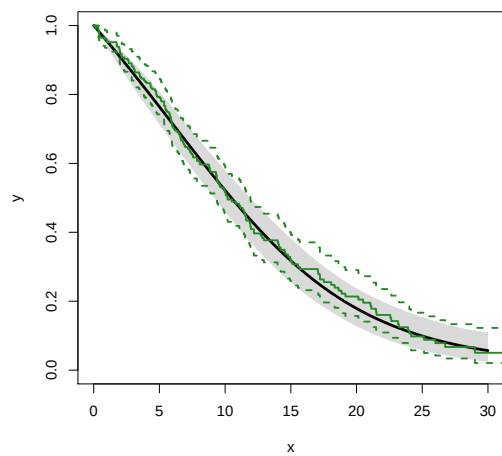
surv

66 / 141

K-M estimator and smooth Poisson model

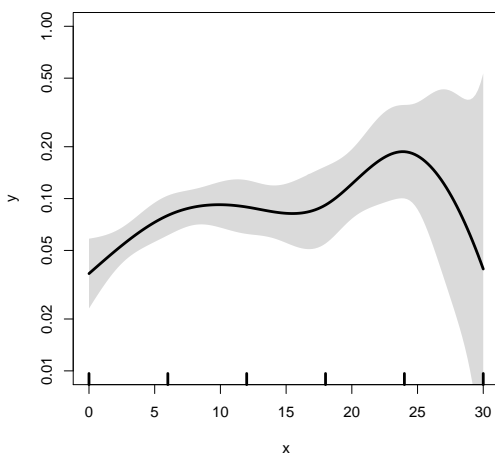


surv

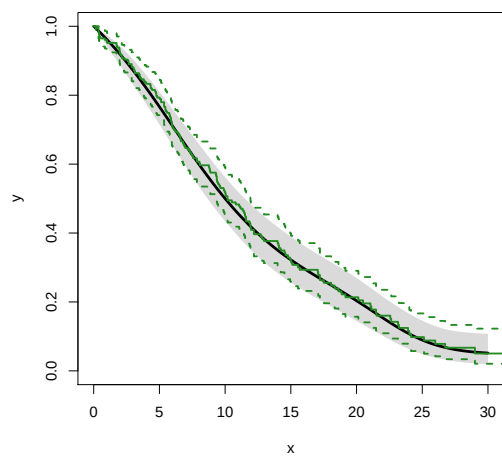


67 / 141

K-M estimator and smooth Poisson model

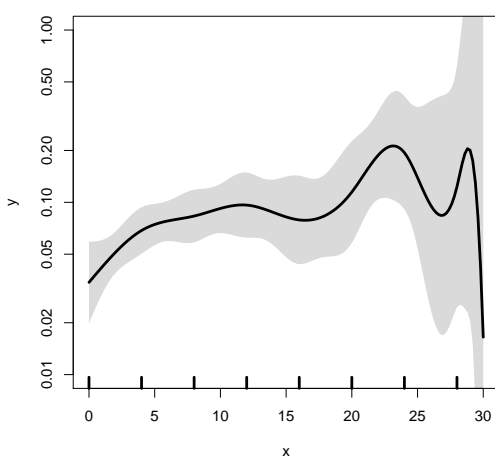


surv

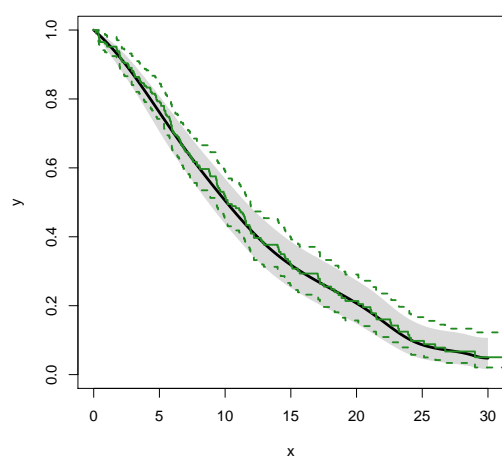


68 / 141

K-M estimator and smooth Poisson model

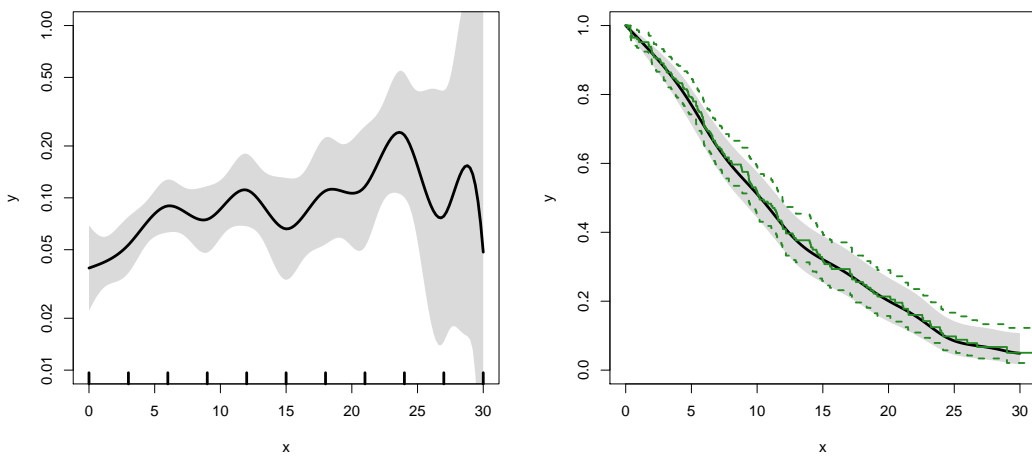


surv



69 / 141

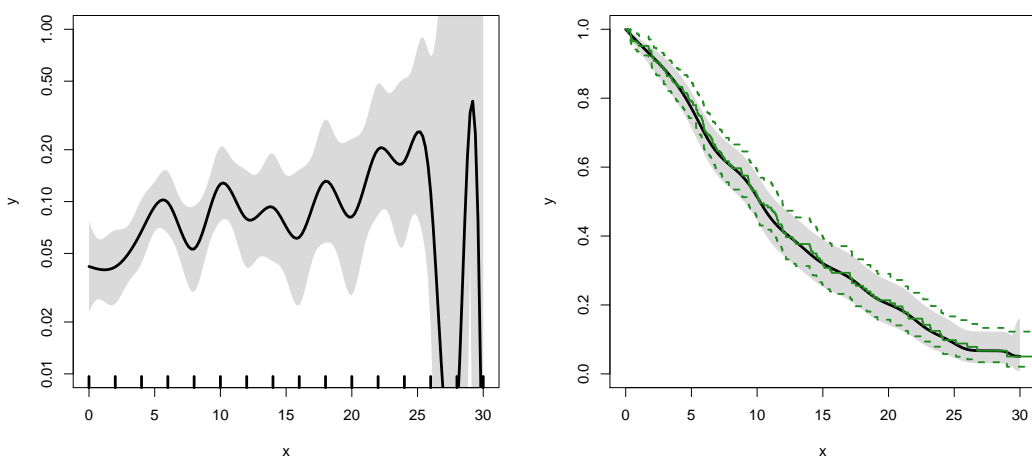
K-M estimator and smooth Poisson model



surv

70 / 141

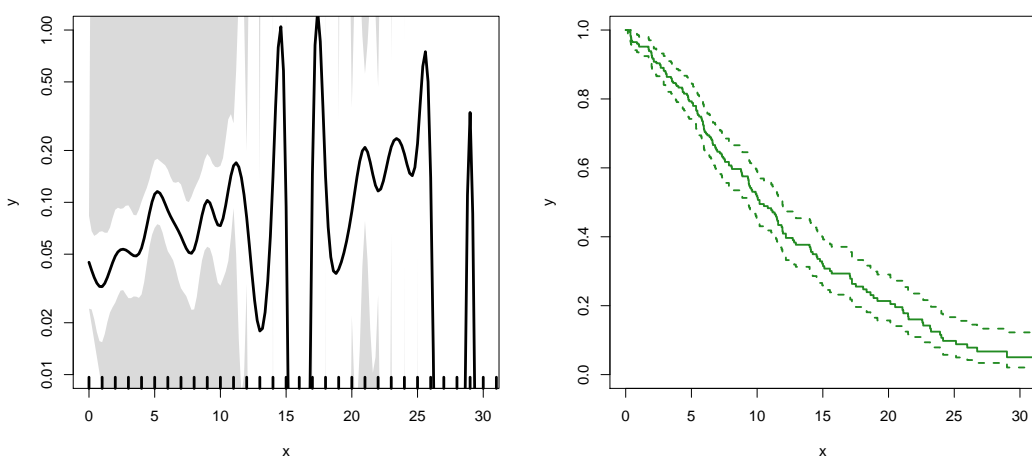
K-M estimator and smooth Poisson model



surv

71 / 141

K-M estimator and smooth Poisson model



surv

72 / 141

Survival analysis summary

- ▶ 1 to 1 correspondence between hazard function and survival function
- ▶ K-M and Cox use a very detailed baseline hazard (omits it)
- ▶ Smooth parametric hazard function more credible:
 - ▶ Define `Lexis` object
 - ▶ Split along time
 - ▶ Fit Poisson model
 - ▶ Prediction data frame
 - ▶ `ci.pred` to get baseline rates
 - ▶ `ci.surv` to get baseline survival

surv

73 / 141

```
> data(lung)
> lung$sex <- factor(lung$sex, labels=c("M", "F"))
> Lx <- Lexis(exit = list(tfe=time),
+   exit.status = factor(status, labels = c("Alive", "Dead")),
+   data = lung)
> sL <- splitMulti(Lx, tfe=seq(0, 1200, 10))
```

Smooth parametric hazard function

```
> m0 <- glm.Lexis(sL, ~ Ns(tfe, knots = seq(0, 1000, 200)) + sex + age)
```

Prediction data frame

```
> nd <- data.frame(tfe = seq(0, 900, 20) + 10, sex = "M", age = 65)
```

Predictions

```
> rate <- ci.pred(m0, nd) * 365.25 # per year, not per day
> surv <- ci.surv(m0, nd, int = 20)
```

Plot the rates

```
> matshade(nd$tfe, rate, log = "y", plot = TRUE)
```

Plot the survival function

```
> matshade(nd$tfe - 10, surv, ylim = c(0, 1), plot = TRUE)
```

surv

74 / 141

```
> library(survival)
> library(Epi)
> library(popEpi)
> # popEpi::splitMulti returns a data.frame rather than a data.table
> options("popEpi.datatable" = FALSE)
> library(tidyverse)
> clear()
```

```
> data(DMlate)
> # str(DMlate)
> set.seed(1952)
> DMlate <- DMlate[sample(1:nrow(DMlate), 2000),]
> str(DMlate)

'data.frame': 2000 obs. of  7 variables:
 $ sex : Factor w/ 2 levels "M","F": 2 1 2 1 1 1 1 1 1 ...
 $ dobth: num  1964 1944 1957 1952 1952 ...
 $ dodm : num  2003 2006 2008 2007 2003 ...
 $ dodth: num  NA NA NA NA NA NA NA NA NA ...
 $ dooad: num  NA 2006 NA 2007 2006 ...
 $ doins: num  NA NA NA 2008 NA ...
 $ dox  : num  2010 2010 2010 2010 2010 ...
```

cmp

75 / 141

Lexis object from DM to Death

```
> Ldm <- Lexis(entry = list(per = dodm,
+                               age = dodm - dobth,
+                               tfd = 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.
NOTE: Dropping 1 rows with duration of follow up < tol

> summary(Ldm)

Transitions:
  To
From   DM Dead  Records:  Events: Risk time: Persons:
  DM 1521  478      1999      478    10742.34      1999
```

cmpr

76 / 141

Cut follow-up at the date of OAD

```
> Cdm <- cutLexis(Ldm,
+                     cut = Ldm$dooad,
+                     timescale = "per",
+                     new.state = "OAD")
> summary(Cdm)

Transitions:
  To
From   DM  OAD Dead  Records:  Events: Risk time: Persons:
  DM   685   634   226      1545     860    5414.3     1545
  OAD   0   836   252      1088     252    5328.1     1088
  Sum  685  1470   478      2633    1112    10742.3     1999
```

cmpr

77 / 141

Cut follow-up at the date of OAD, dooad

```
> subset(Ldm, lex.id %in% c(2:3,20))[,c(1:7,12)]

  per    age tfd lex.dur lex.Cst lex.Xst lex.id  dooad
235221 2005.6 61.517    0  4.3532      DM      DM      2 2005.8
230872 2007.9 51.097    0  2.1109      DM      DM      3    NA
114618 2006.0 73.183    0  3.7919      DM    Dead     20 2007.0

> subset(Cdm, lex.id %in% c(2:3,20))[,c(1:7,12)]

  per    age    tfd lex.dur lex.Cst lex.Xst lex.id  dooad
2  2005.6 61.517 0.00000 0.13415      DM      OAD      2 2005.8
2001 2005.8 61.651 0.13415 4.21903      OAD      OAD      2 2005.8
3   2007.9 51.097 0.00000 2.11088      DM      DM      3    NA
20  2006.0 73.183 0.00000 1.01848      DM      OAD     20 2007.0
2019 2007.0 74.201 1.01848 2.77344      OAD    Dead     20 2007.0
```

cmpr

78 / 141

Restrict to those alive in DM

```
> Adm <- subset(Cdm, lex.Cst == "DM")
> summary(Adm)

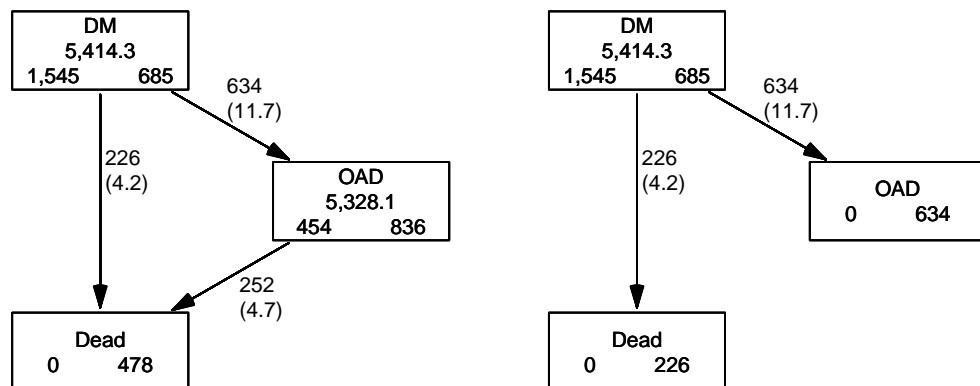
Transitions:
  To
From  DM OAD Dead  Records:  Events: Risk time: Persons:
      DM 685 634 226      1545      860      5414.3      1545

> par(mfrow=c(1,2))
> boxes(Cdm, boxpos = TRUE, scale.R = 100, show.BE = TRUE)
> boxes(Adm, boxpos = TRUE, scale.R = 100, show.BE = TRUE)
```

cmpr

79 / 141

Transitions in Cdm and Adm



cmpr

80 / 141

Survival function?

$$\begin{aligned} S(t) &= \exp \left(- \int_0^t \lambda(u) + \mu(u) \, du \right) \\ S(t) &= \exp \left(- \int_0^t \lambda(u) \, du \right) \\ S(t) &= \exp \left(- \int_0^t \mu(u) \, du \right) \end{aligned}$$

cmpr

81 / 141

Survival function?

- ▶ Regarding either Dead or OAD as censorings — or neither?
- ▶ **Simple survival:** what is the probability of being in each of the states Alive and Dead
 - depends on **one** rate, Alive → Dead
- ▶ **Competing risks:** what is the probability of being in each of the states DM, OAD and Dead
 - depends on **two** rates, DM → OAD and DM → Dead

cmpr

82 / 141

Survival function and Cumulative risk function

`survfit` does the trick; the requirements are:

1. (start, stop, event) arguments to `Surv`
2. the third argument to the `Surv` function is a factor
3. an `id` argument is given, pointing to an id variable that links together records belonging to the same person.
4. the initial state (DM) must be the first level of the factor `lex.Xst`

cmpr

83 / 141

Survival function and Cumulative risk function

```
> levels(Adm$lex.Xst)
[1] "DM"    "OAD"   "Dead"
> m3 <- survfit(Surv(tfd, tfd + lex.dur, lex.Xst) ~ 1,
+                  id = lex.id,
+                  data = Adm)
> # names(m3)
> m3$states
[1] "(s0)"  "OAD"   "Dead"
> head(cbind(time = m3$time, m3$pstate))
      time
[1,] 0.0027379 0.99871 0.0012945 0.00000000
[2,] 0.0054757 0.99288 0.0064725 0.00064725
[3,] 0.0082136 0.98900 0.0090615 0.00194175
[4,] 0.0109514 0.98770 0.0097087 0.00258900
[5,] 0.0136893 0.98382 0.0135922 0.00258900
[6,] 0.0164271 0.98058 0.0168285 0.00258900
```

cmpr

84 / 141

Survival function and cumulative risks—formulae

$$\begin{aligned} S(t) &= \exp\left(-\int_0^t \lambda(u) + \mu(u) du\right) \\ R_{\text{Dead}}(t) &= \int_0^t \mu(u) S(u) du \\ R_{\text{OAD}}(t) &= \int_0^t \lambda(u) S(u) du \\ &= \int_0^t \lambda(u) \exp\left(-\int_0^u \lambda(s) + \mu(s) ds\right) du \end{aligned}$$

$$S(t) + R_{\text{OAD}}(t) + R_{\text{Dead}}(t) = 1, \quad \forall t$$

cmpr

85 / 141

Survival function and cumulative risks

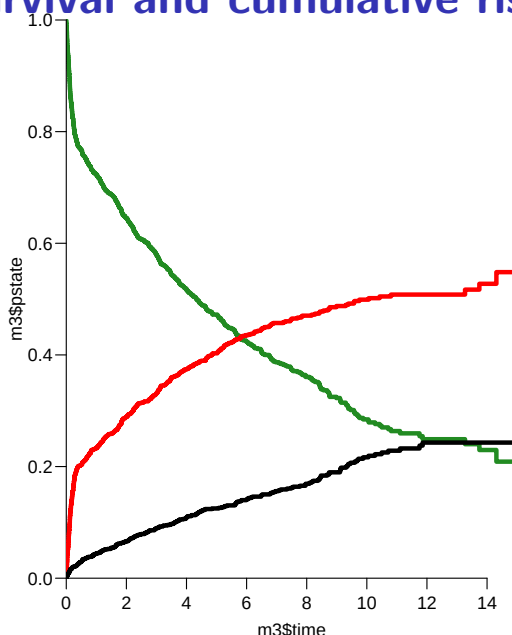
```
> par( mfrow=c(1,2) )
> matplot(m3$time, m3$pstate,
+           type="s", lty=1, lwd=4,
+           col=c("ForestGreen","red","black"),
+           xlim=c(0,15), xaxs="i",
+           ylim=c(0,1), yaxs="i" )
> stackedCIF(m3, lwd=3, xlim=c(0,15), xaxs="i", yaxs="i" )
> text(rep(12,3), c(0.9,0.3,0.6), levels(Cdm))
> box(bty="o")

> par( mfrow = c(1,2) )
> matshade(m3$time, cbind(m3$pstate,
+                         m3$lower,
+                         m3$upper)[,c(1,4,7,2,5,8,3,6,9)],
+           plot = TRUE, lty = 1, lwd = 4,
+           col = c("ForestGreen","red","black"),
+           xlim=c(0,15), xaxs="i",
+           ylim = c(0,1), yaxs = "i" )
> stackedCIF(m3, lwd=3, xlim=c(0,15), xaxs="i", yaxs="i" )
> text(rep(12,3), c(0.9,0.3,0.6), levels(Cdm))
> box(bty="o")
```

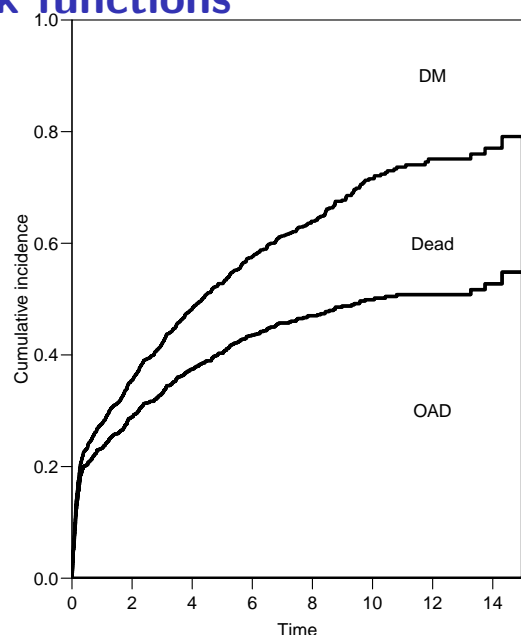
cmpr

86 / 141

Survival and cumulative risk functions

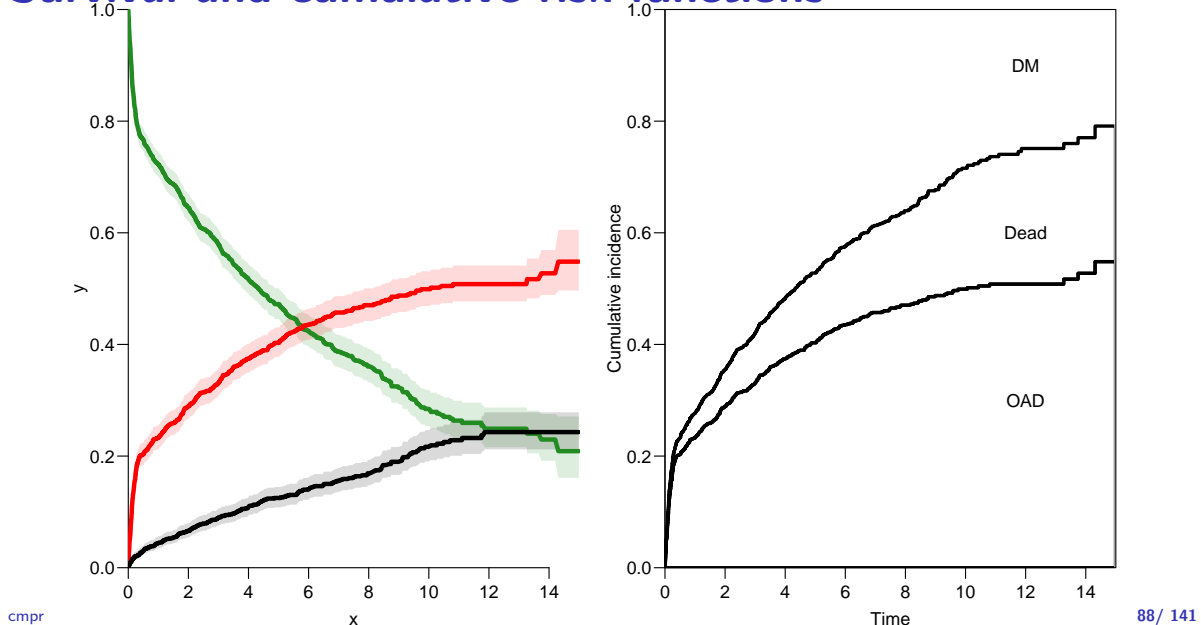


cmpr



87 / 141

Survival and cumulative risk functions



88 / 141

Survival function and cumulative risks—don't

$$\begin{aligned}
 S(t) &= \exp\left(-\int_0^t \lambda(u) + \mu(u) du\right) \\
 R_{\text{Dead}}(t) &= \int_0^t \mu(u) S(u) du \\
 R_{\text{OAD}}(t) &= \int_0^t \lambda(u) S(u) du \\
 &= \int_0^t \lambda(u) \exp\left(-\int_0^u \lambda(s) + \mu(s) ds\right) du \\
 &\neq \int_0^t \lambda(u) \exp\left(-\int_0^u \lambda(s) ds\right) du \\
 &= 1 - \exp\left(-\int_0^t \lambda(s) ds\right) \text{ — nice formula, but wrong!}
 \end{aligned}$$

Probability of OAD **assuming** Dead does not exist **and** rate of OAD unchanged!

cmpr

89 / 141

Survival function and cumulative risks—don't

```

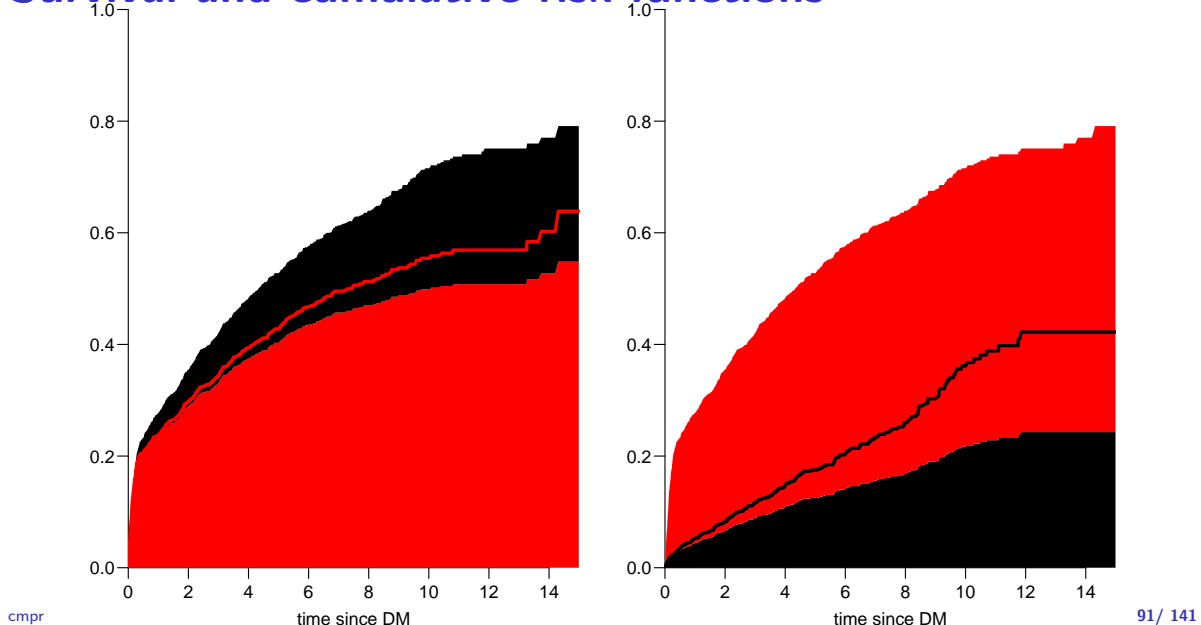
> m2 <- survfit(Surv(tfd,
+                     tfd + lex.dur,
+                     lex.Xst == "OAD" ) ~ 1,
+                     data = Adm)
> M2 <- survfit(Surv(tfd,
+                     tfd + lex.dur,
+                     lex.Xst == "Dead") ~ 1,
+                     data = Adm)
> par(mfrow = c(1,2))
> mat2pol(m3$pstate, c(2,3,1), x = m3$time,
+           col = c("red", "black", "transparent"),
+           xlim=c(0,15), xaxis="i",
+           yaxis = "i", xlab = "time since DM", ylab = "" )
> lines(m2$time, 1 - m2$surv, lwd = 3, col = "red" )
> mat2pol(m3$pstate, c(3,2,1), x = m3$time, yaxis = "i",
+           col = c("black","red","transparent"),
+           xlim=c(0,15), xaxis="i",
+           yaxis = "i", xlab = "time since DM", ylab = "" )
> lines(M2$time, 1 - M2$surv, lwd = 3, col = "black" )

```

cmpr

90 / 141

Survival and cumulative risk functions



Cause-specific rates

- ▶ There is nothing wrong with modeling the cause-specific event-rates, the problem lies in how you transform them into probabilities.
- ▶ The relevant model for a competing risks situation normally consists of separate models for each of the cause-specific rates.
- ▶ . . . not for technical or statistical reasons, but for **substantial** reasons:
it is unlikely that rates of different types of event (OAD initiation and death, say) depend on time in the same way.

cmpr

92 / 141

Cause-specific rates

```
> Sdm <- splitMulti(Adm, tfd = seq(0, 20, 0.1))
> summary(Adm)

Transitions:
  To
From   DM OAD Dead  Records:  Events: Risk time: Persons:
      DM 685 634 226      1545      860     5414.3      1545

> summary(Sdm)

Transitions:
  To
From   DM OAD Dead  Records:  Events: Risk time: Persons:
      DM 54064 634 226     54924      860     5414.3      1545
```

cmpr

93 / 141

Cause-specific rates

```
> round(cbind(
+ with(subset(Sdm, lex.Xst == "OAD" ), quantile(tfd + lex.dur, 0:5/5)),
+ with(subset(Sdm, lex.Xst == "Dead"), quantile(tfd + lex.dur, 0:5/5))), 2)
 [,1]  [,2]
0%    0.00  0.01
20%   0.09  0.51
40%   0.24  1.73
60%   1.27  3.58
80%   3.37  6.20
100%  14.31 11.86

> okn <- c(0, 0.5, 3, 10)
> dkn <- c(0, 2.0, 5, 9)
> OAD.glm <- glm.Lexis(Sdm, ~ Ns(tfd, knots = okn), to = "OAD" )

stats::glm Poisson analysis of Lexis object Sdm with log link:
Rates for the transition: DM->OAD

> Dead.glm <- glm.Lexis(Sdm, ~ Ns(tfd, knots = dkn), to = "Dead" )

stats::glm Poisson analysis of Lexis object Sdm with log link:
Rates for the transition: DM->Dead
```

cmpr

94 / 141

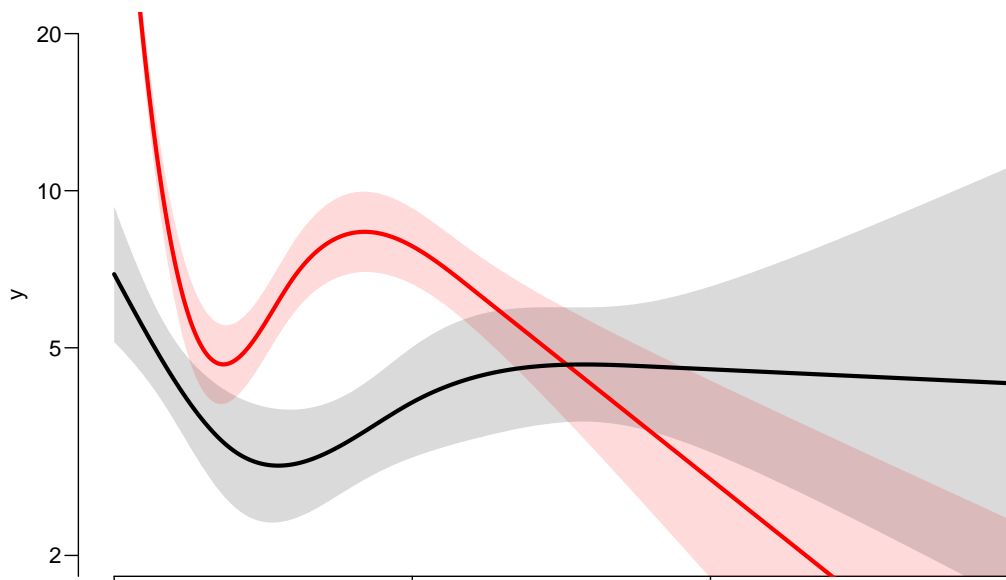
Cause-specific rates

```
> int <- 0.01
> nd <- data.frame(tfd = seq(0, 15, int))
> l.glm <- ci.pred(OAD.glm, nd)
> m.glm <- ci.pred(Dead.glm, nd)
> matshade(nd$tfid,
+            cbind(l.glm, m.glm) * 100,
+            plot = TRUE,
+            yaxs="i", ylim = c(0, 20),
+            # log = "y", ylim = c(2, 20),
+            col = rep(c("red","black"), 2), lwd = 3)
```

cmpr

95 / 141

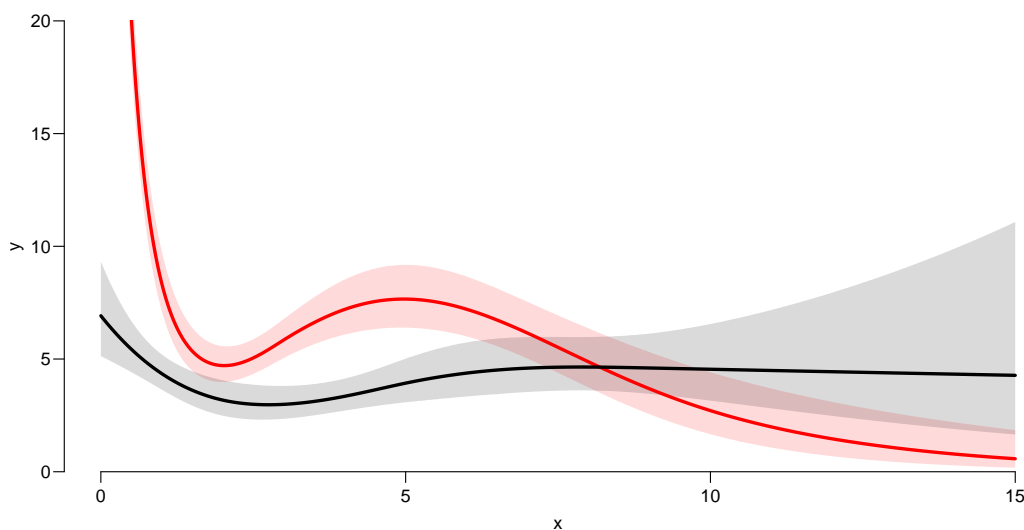
Survival and cumulative risk functions



cmpr

96 / 141

Survival and cumulative risk functions



cmpr

97 / 141

Integrals with R

- ▶ Integrals look scary to many people, but they are really just areas under curves.
- ▶ The key is to understand how a curve is represented in R.
- ▶ A curve of the function $\mu(t)$ is a set of two vectors: one vector of ts and one vector $y = \mu(t)s$.
- ▶ When we have a model such as the `glm` above that estimates the mortality as a function of time (`tfid`), we can get the mortality as a function of time by first choosing the timepoints, say from 0 to 15 years in steps of 0.01 year (≈ 4 days), using `ci.pred`
- ▶ Then use the formulae with all the integrals to get the state probabilities.

cmpr

98 / 141

Integrals with R

```
> t <- seq(0, 15, 0.01)
> nd <- data.frame(tfid = t)
> mu <- ci.pred(Dead.glm, nd)[,1]
> head(cbind(t, mu))

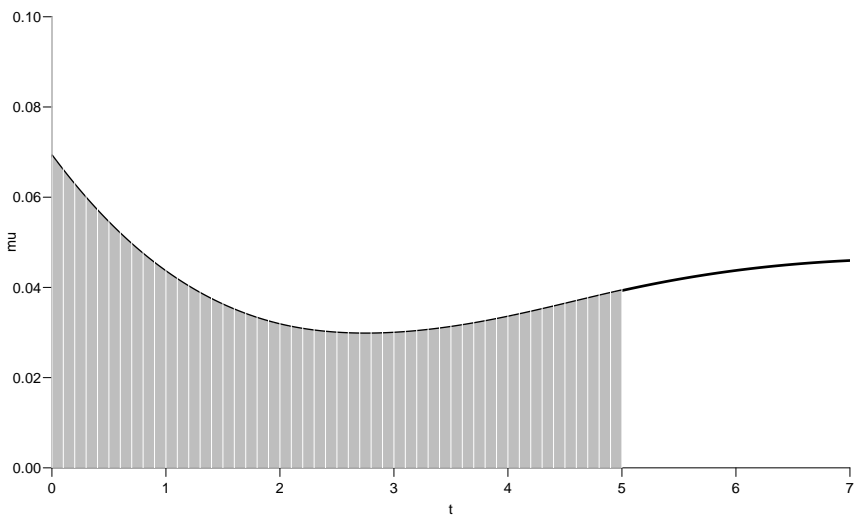
      t        mu
1 0.00 0.069190
2 0.01 0.068853
3 0.02 0.068517
4 0.03 0.068183
5 0.04 0.067851
6 0.05 0.067520

> plot(t, mu, type="l", lwd = 3,
+       xlim = c(0, 7), xaxs = "i",
+       ylim = c(0, 0.1), yaxs = "i")
> polygon(t[c(1:501, 501:1)], c(mu[1:501], rep(0, 501)),
+           col = "gray", border = "transparent")
> abline(v=0:50/10, col="white")
```

cmpr

99 / 141

Integrals with R



cmpr

100 / 141

Numerical integration with R

```
> mid <- function(x) x[-1] - diff(x) / 2  
> (x <- c(1:5, 7, 10))  
[1] 1 2 3 4 5 7 10  
> mid(x)  
[1] 1.5 2.5 3.5 4.5 6.0 8.5
```

`mid(x)` is a vector that is 1 shorter than the vector `x`, just as `diff(x)` is.

So if we want the integral over the period 0 to 5 years, we want the sum over the first 500 intervals, corresponding to the first 501 interval endpoints:

```
> cbind(diff(t), mid(mu))[1:5,]  
[,1] [,2]  
2 0.01 0.069022  
3 0.01 0.068685  
4 0.01 0.068350  
5 0.01 0.068017  
6 0.01 0.067686
```

101 / 141

Numerical integration with R

In practice we will want the integral **function** of μ , so for every t we want $M(t) = \int_0^t \mu(s) d(s)$. This is easily accomplished by the function `cumsum`:

```
> Mu <- c(0, cumsum(diff(t) * mid(mu)))  
> head(cbind(t, Mu))  
  
t Mu  
0.00 0.00000000  
2 0.01 0.00069022  
3 0.02 0.00137707  
4 0.03 0.00206057  
5 0.04 0.00274074  
6 0.05 0.00341760
```

Note the first value which is the integral from 0 to 0, so by definition 0.

cmpr

102 / 141

Cumulative risks from parametric models

If we have estimates of λ and μ as functions of time, we can derive the cumulative risks.

In practice this will be by numerical integration; compute the rates at closely spaced intervals and evaluate the integrals as sums. This is easy.

but what is not so easy is to come up with confidence intervals for the cumulative risks.

cmpr

103 / 141

Simulation of cumulative risks: `ci.Crisk`

1. generate a random vector from the multivariate normal distribution with mean equal to the parameters of the model, and variance-covariance equal to the estimated variance-covariance of the parameter estimates
2. use this to generate a simulated set of rates $(\lambda(t), \mu(t))$, evaluated a closely spaced times
3. use these in numerical integration to derive state probabilities at these times
4. repeat 1000 times, say, to obtain 1000 sets of state probabilities at these times
5. use these to derive confidence intervals for the state probabilities as the 2.5 and 97.5 percentiles of the state probabilities at each time

cmpr

104 / 141

Cumulative risks from parametric models

```
> cR <- ci.Crisk(mods = list(OAD = OAD.glm,
+                               Dead = Dead.glm),
+                               nd = nd)

NOTE: Times are assumed to be in the column tfd at equal distances of 0.01

> str(cR)

List of 4
 $ Crisk: num [1:1501, 1:3, 1:3] 1 0.992 0.984 0.976 0.969 ...
   ..- attr(*, "dimnames")=List of 3
   ...$ tfd : chr [1:1501] "0" "0.01" "0.02" "0.03" ...
   ...$ cause: chr [1:3] "Surv" "OAD" "Dead"
   ...$ : chr [1:3] "50%" "2.5%" "97.5%"
 $ Srisk: num [1:1501, 1:2, 1:3] 0 0.000692 0.001375 0.002049 0.002715 ...
   ..- attr(*, "dimnames")=List of 3
   ...$ tfd : chr [1:1501] "0" "0.01" "0.02" "0.03" ...
   ...$ cause: chr [1:2] "Dead" "Dead+OAD"
   ...$ : chr [1:3] "50%" "2.5%" "97.5%"
 $ Stime: num [1:1501, 1:3, 1:3] 0 0.00996 0.01984 0.02964 0.03937 ...
   ..- attr(*, "dimnames")=List of 3
   ...$ tfd : chr [1:1501] "0" "0.01" "0.02" "0.03" ...
```

cmpr

105 / 141

Cumulative risks from parametric models

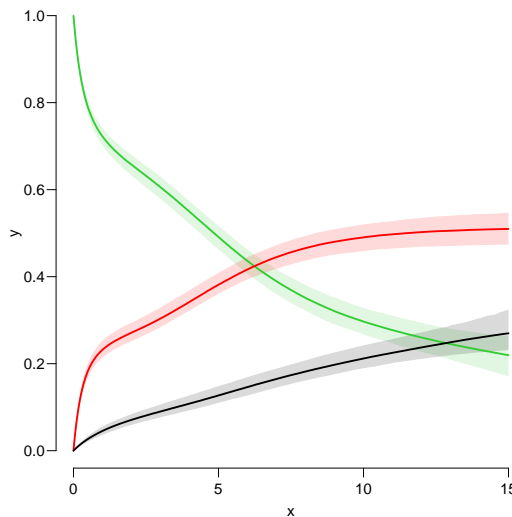
So now plot the cumulative *risk*s of being in each of the states (the **Crisk** component):

```
> matshade(cR$time, cbind(cR$Crisk[,1,],
+                           cR$Crisk[,2,],
+                           cR$Crisk[,3,]), plot = TRUE,
+                           lwd = 2, col = c("limegreen", "red", "black"))
```

cmp

106 / 141

Survival and cumulative risk functions



cmp

107 / 141

Stacked probabilities: (matrix 2 polygons)

```
> mat2pol(cR$Crisk[,3:1,1], col = c("forestgreen", "red", "black")[3:1])
```

1st argument to **mat2pol** must be a 2-dimensional matrix, with rows representing the *x*-axis of the plot, and columns states.

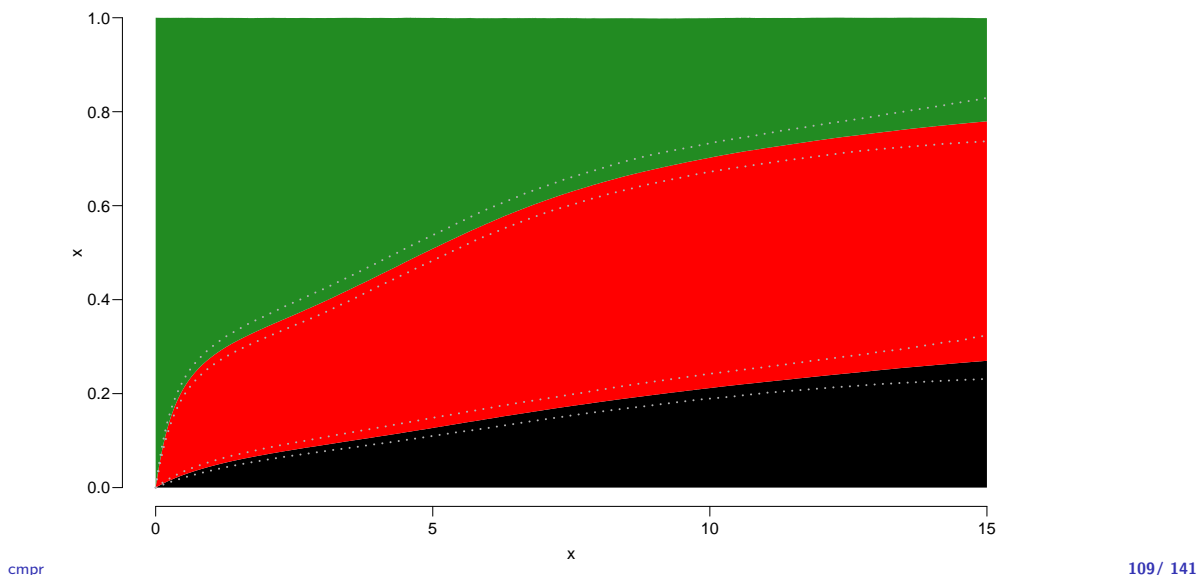
The component **Srisk** has the confidence limits of the stacked probabilities:

```
> mat2pol(cR$Crisk[,3:1,1], col = c("forestgreen", "red", "black")[3:1])
> matlines(as.numeric(dimnames(cR$Srisk)[["tfd"]]),
+            cbind(cR$Srisk[,"Dead"      ,2:3],
+                  cR$Srisk[,"Dead+OAD",2:3]),
+            lty = 3, lwd = 2, col = gray(0.7))
```

cmp

108 / 141

Survival and cumulative risk functions



Expected life time: using simulated objects

The areas between the lines (up to say 10 years) are **expected sojourn times**, that is:

- ▶ expected years alive without OAD
- ▶ expected years lost to death without OAD
- ▶ expected years after OAD, including years dead after OAD

Not all of these are of direct relevance; actually only the first may be so.

They are available (with simulation-based confidence intervals) in the component of `cR$Stime` (*Sojourn time*).

cmpr

110 / 141

Expected life time: using simulated objects

A relevant quantity would be the expected time alive without OAD during the first 5, 10 and 15 years:

```
> str(cR$Stime)
num [1:1501, 1:3, 1:3] 0 0.00996 0.01984 0.02964 0.03937 ...
- attr(*, "dimnames")=List of 3
..$ tfd : chr [1:1501] "0" "0.01" "0.02" "0.03" ...
..$ cause: chr [1:3] "Surv" "OAD" "Dead"
..$      : chr [1:3] "50%" "2.5%" "97.5%"

> round(cR$Stime[c("5", "10", "15"), "Surv", ], 1)
tfd 50% 2.5% 97.5%
5   3.2  3.1  3.3
10  5.1  4.9  5.3
15  6.4  6.0  6.7
```

cmpr

111 / 141

BBackground: Steno 2 trial

- ▶ Clinical trial for diabetes ptt. with kidney disease (micro-albuminuria)
- ▶ 80 ptt. randomised to either of
 - ▶ Conventional treatment
 - ▶ Intensified multifactorial treatment
- ▶ 1993–2001
- ▶ follow-up till 2018

msmt

112 / 141

Steno 2 trial: goal

- ▶ Is there a treatment effect on:
 - ▶ CVD mortality
 - ▶ non-CVD mortality
 - ▶ Albuminuria state
- ▶ Rate-ratios
- ▶ Life times
- ▶ Changes in clinical parameters

msmt

113 / 141

```
> data(steno2)
> steno2 <- cal.yr(steno2)
> steno2 <- transform(steno2,
+                      doEnd = pmin(doDth, doEnd, na.rm = TRUE))
> str(steno2)

'data.frame': 160 obs. of 14 variables:
 $ id      : num  1 2 3 4 5 6 7 8 9 10 ...
 $ allo    : Factor w/ 2 levels "Int","Conv": 1 1 2 2 2 2 2 1 1 1 ...
 $ sex     : Factor w/ 2 levels "F","M": 2 2 2 2 2 2 1 2 2 2 ...
 $ baseCVD : num  0 0 0 0 0 1 0 0 0 0 ...
 $ deathCVD: num  0 0 0 0 1 0 0 0 0 1 0 ...
 $ doBth   : 'cal.yr' num  1932 1947 1943 1945 1936 ...
 $ doDM    : 'cal.yr' num  1991 1982 1983 1977 1986 ...
 $ doBase  : 'cal.yr' num  1993 1993 1993 1993 1993 ...
 $ doCVD1  : 'cal.yr' num  2014 2009 2002 1995 1994 ...
 $ doCVD2  : 'cal.yr' num  NA 2009 NA 1997 1995 ...
 $ doCVD3  : 'cal.yr' num  NA 2010 NA 2003 1998 ...
 $ doESRD  : 'cal.yr' num  NaN NaN NaN NaN 1998 ...
 $ doEnd   : 'cal.yr' num  2015 2015 2002 2003 1998 ...
 $ doDth   : 'cal.yr' num  NA NA 2002 2003 1998 ...
```

msmt

114 / 141

A Lexis object

```
> L2 <- Lexis(entry = list(per = doBase,
+                               age = doBase - doBth,
+                               tfi = 0),
+                 exit = list(per = doEnd),
+                 exit.status = factor(deathCVD + !is.na(doDth),
+                                       labels=c("Mic", "D(oth)", "D(CVD)")),
+                 id = id,
+                 data = steno2)
```

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

Explain the coding of `exit.status`.

msmt

115 / 141

A Lexis object

```
> summary(L2, t = TRUE)

Transitions:
  To
From  Mic D(oth) D(CVD)  Records:  Events: Risk time: Persons:
      Mic   67     55     38       160       93    2416.59       160

Timescales:
per age tfi
  ""  ""   ""
```

How many persons are there in the cohort?

How many deaths are there in the cohort?

How much follow-up time is there in the cohort?

How many states are there in the model (so far)?

msmt

116 / 141

Albuminuria status

```
> data(st2alb) ; head(st2alb, 3)

  id      doTr state
1  1 1993-06-12   Mic
2  1 1995-05-13  Norm
3  1 2000-01-26   Mic

> cut2 <- rename(cal.yr(st2alb),
+                   lex.id = id,
+                   cut = doTr,
+                   new.state = state)
> with(cut2, addmargins(table(table(lex.id))))
```

| | 1 | 2 | 3 | 4 | 5 | Sum |
|---|----|----|----|----|-----|-----|
| 4 | 25 | 40 | 46 | 41 | 156 | |

What does this table mean?

msmt

117 / 141

Albuminuria status as states

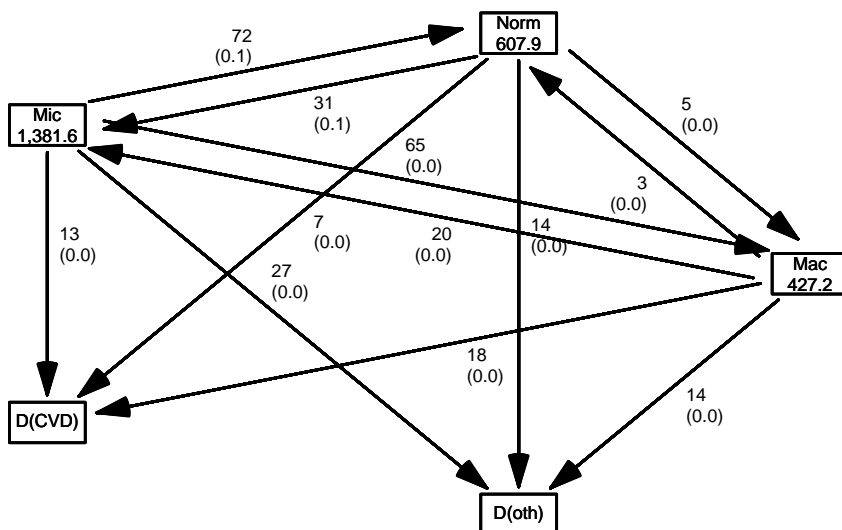
```
> L2$per <- as.numeric(L2$per)
> cut2$cut <- as.numeric(cut2$cut)
> L3 <- rcutLexis(L2, cut2, time = "per")
> summary(L3)
```

Transitions:

| From | Mic | Norm | Mac | D(oth) | D(CVD) | Records: | Events: | Risk | time: | Persons: |
|------|-----|------|-----|--------|--------|----------|---------|---------|-------|----------|
| Mic | 299 | 72 | 65 | 27 | 13 | 476 | 177 | 1381.57 | | 160 |
| Norm | 31 | 90 | 5 | 14 | 7 | 147 | 57 | 607.86 | | 69 |
| Mac | 20 | 3 | 44 | 14 | 18 | 99 | 55 | 427.16 | | 64 |
| Sum | 350 | 165 | 114 | 55 | 38 | 722 | 289 | 2416.59 | | 160 |

```
> boxes(L3, boxpos = TRUE, cex = 0.8)
```

What's wrong with this



What's in jump

```
> (jump <-
+ subset(L3, (lex.Cst == "Norm" & lex.Xst == "Mac") |
+         (lex.Xst == "Norm" & lex.Cst == "Mac"))[,,
+         c("lex.id", "per", "lex.dur", "lex.Cst", "lex.Xst")])
```

| | lex.id | per | lex.dur | lex.Cst | lex.Xst |
|-----|--------|----------|------------|---------|---------|
| 291 | 70 | 1999.487 | 2.6748802 | Mac | Norm |
| 353 | 86 | 2001.759 | 12.8158795 | Norm | Mac |
| 506 | 130 | 2000.910 | 1.8781656 | Mac | Norm |
| 511 | 131 | 1997.756 | 4.2354552 | Norm | Mac |
| 525 | 136 | 1997.214 | 0.4709103 | Mac | Norm |
| 526 | 136 | 1997.685 | 4.2436687 | Norm | Mac |
| 654 | 171 | 1996.390 | 5.3388090 | Norm | Mac |
| 676 | 175 | 2004.585 | 9.8836413 | Norm | Mac |

—and what will you do about it?

How to fix things

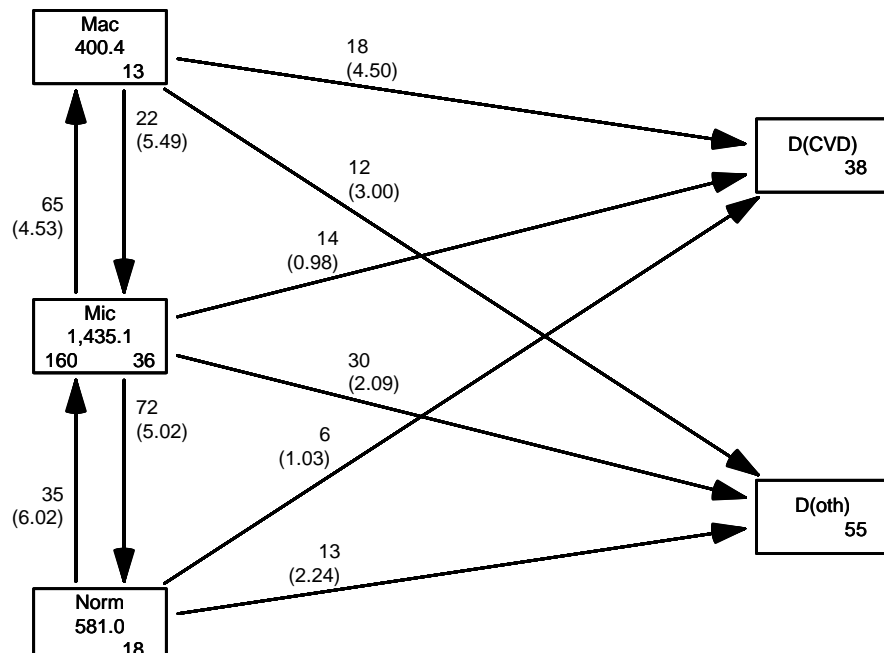
```
> set.seed(1952)
> xcut <- transform(jump,
+                      cut = per + lex.dur * runif(per, 0.1, 0.9),
+                      new.state = "Mic")
> xcut <- select(xcut, c(lex.id, cut, new.state))
> L4 <- rcutLexis(L3, xcut)
> L4 <- Relevel(L4, c("Norm", "Mic", "Mac", "D(CVD)", "D(oth)"))
> summary(L4)
```

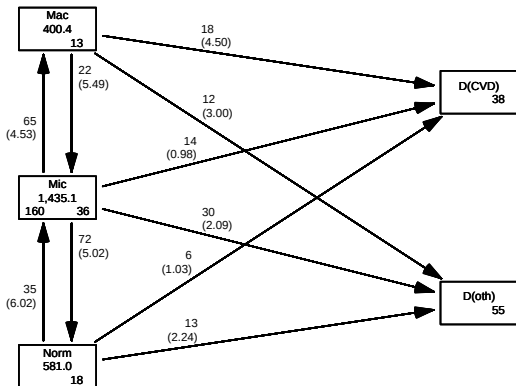
Transitions:

| From | Norm | Mic | Mac | D(CVD) | D(oth) | Records: | Events: | Risk time: | Persons: |
|------|------|-----|-----|--------|--------|----------|---------|------------|----------|
| Norm | 90 | 35 | 0 | 6 | 13 | 144 | 54 | 581.04 | 66 |
| Mic | 72 | 312 | 65 | 14 | 30 | 493 | 181 | 1435.14 | 160 |
| Mac | 0 | 22 | 41 | 18 | 12 | 93 | 52 | 400.41 | 60 |
| Sum | 162 | 369 | 106 | 38 | 55 | 730 | 287 | 2416.59 | 160 |

Plot the boxes

```
> boxes(L4, boxpos = list(x = c(20, 20, 20, 80, 80),
+                           y = c(10, 50, 90, 75, 25)),
+        show.BE = "nz",
+        scale.R = 100, digits.R = 2,
+        cex = 0.9, pos.arr = 0.3)
```





Explain all the numbers in the graph.

Describe the overall effect of albuminuria on the two mortality rates.

Modeling transition rates

- ▶ A model with a smooth effect of timescales on the rates require follow-up in small bits
- ▶ Achieved by `splitLexis` (or `splitMulti` from `popEpi`)
- ▶ Compare the `Lexis` objects

```
> S4 <- splitMulti(L4, tfi = seq(0, 25, 1/2))
> summary(L4)
```

Transitions:

To

| From | Norm | Mic | Mac | DCVD | D(oth) | Records: | Events: | Risk time: | Persons: |
|------|------|-----|-----|------|--------|----------|---------|------------|----------|
| Norm | 90 | 35 | 0 | 6 | 13 | 144 | 54 | 581.04 | 66 |
| Mic | 72 | 312 | 65 | 14 | 30 | 493 | 181 | 1435.14 | 160 |
| Mac | 0 | 22 | 41 | 18 | 12 | 93 | 52 | 400.41 | 60 |
| Sum | 162 | 369 | 106 | 38 | 55 | 730 | 287 | 2416.59 | 160 |

```
> summary(S4)
```

Transitions:

To

| From | Norm | Mic | Mac | DCVD | D(oth) | Records: | Events: | Risk time: | Persons: |
|------|------|------|-----|------|--------|----------|---------|------------|----------|
| Norm | 1252 | 35 | 0 | 6 | 13 | 1306 | 54 | 581.04 | 66 |
| Mic | 72 | 3101 | 65 | 14 | 30 | 3282 | 181 | 1435.14 | 160 |
| Mac | 0 | 22 | 844 | 18 | 12 | 896 | 52 | 400.41 | 60 |
| Sum | 1324 | 3158 | 909 | 38 | 55 | 5484 | 287 | 2416.59 | 160 |

How the split works:

```
> subset(L4, lex.id == 96)[,1:7]

  per      age      tfi      lex.dur lex.Cst lex.Xst lex.id
417 1993.650 51.53183 0.0000000 0.4544832      Mic     Norm    96
418 1994.104 51.98631 0.4544832 2.5790554      Norm     Norm    96
419 1996.683 54.56537 3.0335387 1.9028063      Norm     Norm    96
420 1998.586 56.46817 4.9363450 2.8966461      Norm D(CVD)    96

> subset(S4, lex.id == 96)[c(1:5,NA,17:19),1:7]

  lex.id      per      age      tfi      lex.dur lex.Cst lex.Xst
3138      96 1993.650 51.53183 0.0000000 0.45448323      Mic     Norm
3139      96 1994.104 51.98631 0.4544832 0.04551677      Norm     Norm
3140      96 1994.150 52.03183 0.5000000 0.50000000      Norm     Norm
3141      96 1994.650 52.53183 1.0000000 0.50000000      Norm     Norm
3142      96 1995.150 53.03183 1.5000000 0.50000000      Norm     Norm
NA        NA       NA       NA          NA          NA <NA>   <NA>
3154      96 2000.150 58.03183 6.5000000 0.50000000      Norm     Norm
3155      96 2000.650 58.53183 7.0000000 0.50000000      Norm     Norm
3156      96 2001.150 59.03183 7.5000000 0.33299110      Norm D(CVD)
```

msmt

127 / 141

```
> subset(L4, lex.id == 159)[,1:7]
```

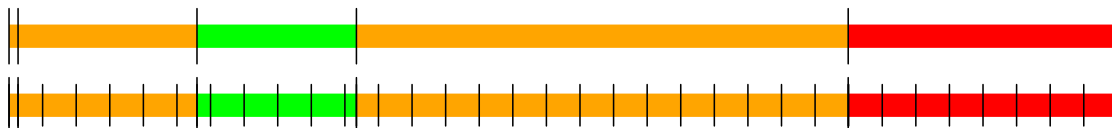
```
  per      age      tfi      lex.dur lex.Cst lex.Xst lex.id
646 1994.025 67.49624 0.0000000 0.1341547      Mic     Mic    159
647 1994.159 67.63039 0.1341547 2.6639288      Mic     Norm    159
648 1996.823 70.29432 2.7980835 2.3737166      Norm     Mic    159
649 1999.196 72.66804 5.1718001 7.3210130      Mic     Mac    159
650 2006.517 79.98905 12.4928131 3.9479808      Mac D(CVD)    159
```

```
> subset(S4, lex.id == 159)[c(1:2,NA,6:7,NA,12:13,NA,27:28,NA,36:37),1:7]
```

```
  lex.id      per      age      tfi      lex.dur lex.Cst lex.Xst
4853      159 1994.025 67.49624 0.0000000 0.1341547      Mic     Mic
4854      159 1994.159 67.63039 0.1341547 0.3658453      Mic     Mic
NA        NA       NA       NA          NA          NA <NA>   <NA>
4858      159 1996.025 69.49624 2.0000000 0.5000000      Mic     Mic
4859      159 1996.525 69.99624 2.5000000 0.2980835      Mic     Norm
NA.1       NA       NA       NA          NA          NA <NA>   <NA>
4864      159 1998.525 71.99624 4.5000000 0.5000000      Norm     Norm
4865      159 1999.025 72.49624 5.0000000 0.1718001      Norm     Mic
NA.2       NA       NA       NA          NA          NA <NA>   <NA>
4879      159 2005.525 78.99624 11.5000000 0.5000000      Mic     Mic
4880      159 2006.025 79.49624 12.0000000 0.4928131      Mic     Mac
NA.3       NA       NA       NA          NA          NA <NA>   <NA>
4888      159 2009.525 82.99624 15.5000000 0.5000000      Mac     Mac
msmt 4889      159 2010.025 83.49624 16.0000000 0.4407940      Mac D(CVD)
```

128 / 141

How the split works



Same amount of follow-up

Same transitions

More intervals (5, resp. 37)

Different value of time scales between intervals

msmt

129 / 141

Purpose of the split

- ▶ Assumption of constant rate in each interval
- ▶ All intervals are (shorter than) 0.5 years
- ▶ Magnitude of the rates depend on covariates:
 - ▶ fixed covariates
 - ▶ time scales
 - ▶ randomly varying covariates (not now)
- ▶ value of covariates differ between intervals
- ▶ each record contributes one term to the (log-)likelihood for a specific rate
 - from a given origin state (`lex.Cst`)
 - to a given destination state (`lex.Cst`).
- ▶ —looks as the likelihood for a single Poisson observation

msmt

130 / 141

Modeling the rate: Mic → D(CVD)

```
> mr <- glm(cbind(lex.Xst == "D(CVD)" & lex.Cst != lex.Xst,  
+                 lex.dur)  
+             ~ Ns(tfi, knots = seq(0, 20, 5)) +  
+             Ns(age, knots = seq(50, 80, 10)),  
+             family = poisreg,  
+             data = subset(S4, lex.Cst == "Mic"))
```

... the same as:

```
> mp <- glm((lex.Xst == "D(CVD)" & lex.Cst != lex.Xst)  
+             ~ Ns(tfi, knots = seq(0, 20, 5)) +  
+             Ns(age, knots = seq(50, 80, 10)),  
+             offset = log(lex.dur),  
+             family = poisson,  
+             data = subset(S4, lex.Cst == "Mic"))  
> summary(coef(mr) - coef(mp))
```

| | Min. | 1st Qu. | Median | Mean | 3rd Qu. | Max. |
|--|------------|------------|------------|------------|-----------|-----------|
| | -1.315e-12 | -2.389e-13 | -2.343e-14 | -1.540e-13 | 7.050e-15 | 6.466e-13 |

msmt

131 / 141

Modeling the rate: Mic → D(CVD)

A convenient wrapper for Lexis objects:

```
> mL <- glm.Lexis(S4,  
+                     ~ Ns(tfi, knots = seq(0, 20, 5)) +  
+                     Ns(age, knots = seq(50, 80, 10)),  
+                     from = "Mic",  
+                     to = "D(CVD)")  
  
stats::glm Poisson analysis of Lexis object S4 with log link:  
Rates for the transition: Mic->D(CVD)  
  
> summary(coef(mr) - coef(mL))  
  
Min. 1st Qu. Median Mean 3rd Qu. Max.  
0 0 0 0 0 0
```

msmt

132 / 141

`glm.Lexis` by default models all transitions **to** absorbing states, from states preceding these

```
> mX <- glm.Lexis(S4,
+                     ~ Ns(tfi, knots = seq( 0, 20, 5)) +
+                     Ns(age, knots = seq(50, 80, 10)) +
+                     lex.Cst)

stats::glm Poisson analysis of Lexis object S4 with log link:
Rates for transitions: Norm->D(CVD), Mic->D(CVD), Mac->D(CVD), Norm->D(oth), Mic->
```

Describe the model(s) in `mX`:

- ▶ What rates are modeled ?
- ▶ How are they modeled (assumptions about shapes) ?
- ▶ What are the differences between the rates modeled?
- ▶ What would you rather do?

```
> mox <- glm.Lexis(S4, ~ Ns(tfi, knots = seq( 0, 20, 5)) +
+                     Ns(age, knots = seq(50, 80, 10)) +
+                     lex.Cst / allo,
+                     to = "D(oth)")

stats::glm Poisson analysis of Lexis object S4 with log link:
Rates for transitions: Norm->D(oth), Mic->D(oth), Mac->D(oth)

> mCx <- glm.Lexis(S4, ~ Ns(tfi, knots = seq( 0, 20, 5)) +
+                     Ns(age, knots = seq(50, 80, 10)) +
+                     lex.Cst / allo,
+                     to = "D(CVD)")

stats::glm Poisson analysis of Lexis object S4 with log link:
Rates for transitions: Norm->D(CVD), Mic->D(CVD), Mac->D(CVD)
```

```
> det <- glm.Lexis(S4, ~ Ns(tfi, knots = seq( 0, 20, 5)) +
+                     Ns(age, knots = seq(50, 80, 10)) +
+                     lex.Cst / allo,
+                     from = c("Norm", "Mic"),
+                     to = c("Mic", "Mac"))

stats::glm Poisson analysis of Lexis object S4 with log link:
Rates for transitions: Norm->Mic, Mic->Mac

> imp <- glm.Lexis(S4, ~ Ns(tfi, knots = seq( 0, 20, 5)) +
+                     Ns(age, knots = seq(50, 80, 10)) +
+                     lex.Cst / allo,
+                     from = c("Mac", "Mic"),
+                     to = c("Mic", "Norm"))

stats::glm Poisson analysis of Lexis object S4 with log link:
Rates for transitions: Mac->Mic, Mic->Norm
```

Specification of the model

```
> Tr <- list(Norm = list("Mic" = det,
+                         "D(oth)" = mox,
+                         "D(CVD)" = mCx),
+             Mic = list("Mac" = det,
+                        "Norm" = imp,
+                        "D(oth)" = mox,
+                        "D(CVD)" = mCx),
+             Mac = list("Mic" = imp,
+                        "D(oth)" = mox,
+                        "D(CVD)" = mCx))
> lapply(Tr, names)

$Norm
[1] "Mic"      "D(oth)"   "D(CVD)"

$Mic
[1] "Mac"      "Norm"     "D(oth)"   "D(CVD)"

$Mac
[1] "Mic"      "D(oth)"   "D(CVD)"

msmt
```

136 / 141

Specification of the prediction population

```
> ini <- L2[,c("per", "age", "tfi")]
> ini <- rbind(transform(ini, lex.Cst = factor("Mic"), allo = factor("Int")),
+               transform(ini, lex.Cst = factor("Mic"), allo = factor("Conv")))
> ini$lex.Cst <- factor(ini$lex.Cst, levels = levels(L4))
> str(ini)

Classes 'Lexis' and 'data.frame': 320 obs. of 5 variables:
 $ per    : num  1993 1993 1993 1993 1993 ...
 $ age    : 'cal.yr' num  61.1 46.6 49.9 48.5 57.3 ...
 $ tfi    : num  0 0 0 0 0 0 0 0 0 0 ...
 $ lex.Cst: Factor w/ 5 levels "Norm","Mic","Mac",...: 2 2 2 2 2 2 2 2 2 2 ...
 $ allo   : Factor w/ 2 levels "Int","Conv": 1 1 1 1 1 1 1 1 1 1 ...
 - attr(*, "breaks")=List of 3
   ..$ per: NULL
   ..$ age: NULL
   ..$ tfi: NULL
 - attr(*, "time.scales")= chr  "per" "age" "tfi"
 - attr(*, "time.since")= chr  "" "" ""

msmt
```

137 / 141

Simulating a cohort

```
> set.seed(1952)
> system.time(
+   Sorg <- simLexis(Tr = Tr, # models for each transition
+                     init = ini, # cohort of straters
+                     N = 10, # how many copies of each person in ini
+                     t.range = 20, # how long should we simulate before censoring
+                     n.int = 100))# how many intervals for evaluating rates

  user  system elapsed
19.460   9.104  17.566

msmt
```

138 / 141

Simulated cohort

```
> summary(Sorg)

Transitions:
  To
From   Norm  Mic  Mac D(CVD) D(oth)  Records:  Events: Risk time: Persons:
  Norm  387  655    0    114    266      1422     1035  11582.90    1300
  Mic   1422 646 1302    281    574      4225     3579  26869.84    3200
  Mac     0 370 308    383    241      1302     994   7831.63    1206
  Sum  1809 1671 1610    778   1081      6949     5608  46284.38    3200
```

msmt

139 / 141

Who is where when?

```
> Nst <- nState(Sorg,
+                   at = seq(0, 20, 0.2),
+                   from = 0,
+                   time.scale = "tfi")
> str(Nst)

'table' int [1:101, 1:5] 0 88 167 233 295 ...
- attr(*, "dimnames")=List of 2
..$ when : chr [1:101] "0" "0.2" "0.4" "0.6" ...
..$ State: chr [1:5] "Norm" "Mic" "Mac" "D(CVD)" ...

> head(Nst)

  State
when Norm  Mic  Mac D(CVD) D(oth)
  0     0 3200    0    0    0
  0.2   88 3077   33    2    0
  0.4   167 2966   62    5    0
  0.6   233 2863   98    6    0
  0.8   295 2772  120   13    0
  1     341 2693  148   17    1
```

msmt

140 / 141

Who is where when?

```
> Nint <- nState(subset(Sorg, allo == "Int"),
+                   at = seq(0, 20, 0.1),
+                   from = 0,
+                   time.scale = "tfi")
> Nconv<- nState(subset(Sorg, allo == "Conv"),
+                   at = seq(0, 20, 0.1),
+                   from = 0,
+                   time.scale = "tfi")
> head(Nint , 4)

  State
when Norm  Mic  Mac D(CVD) D(oth)
  0     0 1600    0    0    0
  0.1   24 1569    6    1    0
  0.2   55 1533   11    1    0
  0.3   76 1507   15    2    0

> head(Nconv, 4)

  State
when Norm  Mic  Mac D(CVD) D(oth)
  0     0 1600    0    0    0
```

msmt

141 / 141