

▶ Exact time of transition between states for all persons

Multistate models (MSintro)

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Multistate models (MSintro

Lung cancer survival	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))
computations	63.3405 ⁸ 61.0777 ⁸
	Formally there is a significant difference in survival between men and
Multistate models: Occurrence rates, cumulative risks, competing risks,	<pre>WOMEN > survdiff(Surv(time, status==2) ~ sex, data = lung)</pre>
state probabilities with multiple states and time scales with ${f R}$ and Epi::Lexis	Call: survdiff(formula = Surv(time, status == 2) ~ sex, data = lung)
Baker HDI, 22-23 February 2023	N Observed Expected (O-E)^2/E (O-E)^2/V
http://bendixcarstensen.com/AdvCoh/courses/Melb-2023 surv	sex=W 90 53 73.4 5.68 10.3
	Chisq= 10.3 on 1 degrees of freedom, p= 0.001 Lung cancer survival (serv) 23/130
	Rates and rate-ratios
Prerequisites	
<pre>> library(Epi) > library(popEpi)</pre>	Occurrence rate:
<pre>> # popEpi::splitMulti returns a data.frame rather than a data.table > options("popEpi.datatable" = FALSE)</pre>	$\lambda(t) = \lim_{h o 0} \mathrm{P} \left\{ event \; in \; (t,t+h] \; \; alive at \; t ight\} / h$
	 —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 (x1, x2):
	$\lambda(t, x) = \lambda_0(t) \exp(\beta_1 x_1 + \beta_2 x_2)$
	—mortality depends on the person's sex and age, say.
	Data looks like data for a K-M analysis plus covariate values
Lung cancer survival (ssrv) 19/130	Lung cancer survival (ssrv) 24/130
The lung data set	Rates and rate-ratios: Simple Cox model
<pre>> library(survival) > data(lung) > lung\$sex <- factor(lung\$sex,</pre>	Now explore how sex and age (at diagnosis) influence the mortality—note that in a Cox-model we are addressing the mortality
+ levels = 1:2, + labels = c("M", "W"))	rate and not the survival:
<pre>> lung%time <- lung%time / (365.25/12) > head(lung)</pre>	<pre>> c0 <- coxph(Surv(time, status == 2) ~ sex , data = lung) > c1 <- coxph(Surv(time, status == 2) ~ sex + age, data = lung)</pre>
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	> summary(c1) > ci.exp(c0) > ci.exp(c1)
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	What variables from lung are we using?
Lung cancer survival (szrv) 20 / 130	Lung cancer survival (serv) 25/130
Survival function	> c0 <- comp(Surv(time status == 2) ~ sex data = lung)
 Use survfit to construct the Kaplan-Meier estimator of overall 	<pre>> c0 <- coxph(Surv(time, status == 2) ~ sex , data = lung) > c1 <- coxph(Surv(time, status == 2) ~ sex + age, data = lung) > summary(c1)</pre>
survival:	Call: coxph(formula = Surv(time, status == 2) ~ sex + age, data = lung)
> ?Surv > ?survfit	n= 228, number of events= 165
<pre>> km <- survfit(Surv(time, status == 2) ~ 1, data = lung) > km</pre>	coef exp(coef) se(coef) z Pr(> z) sexW -0.513219 0.598566 0.167458 -3.065 0.00218 **
Call: survfit(formula = Surv(time, status == 2) ~ 1, data = lung) n events median 0.95LCL 0.95UCL	age 0.017045 1.017191 0.009223 1.848 0.06459 . Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
[1,] 228 165 10.2 9.36 11.9 > # summary(km) # very long output	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
Lung cancer survival (1017) 21/130	Wald test = 13.47 on 2 df, p=0.001 Score (logrank) test = 13.72 on 2 df, p=0.001 Lung cancer survised (surv) 26/130
We can plot the survival curve—this is the default plot for a survfit object:	> ci.exp(c0)
> plot(km)	exp(Est.) 2.5% 97.5% sexW 0.5880028 0.4237178 0.8159848
	<pre>> ci.exp(c1) exp(Est.) 2.5% 97.5% sexW 0.598566 0.4310936 0.8310985</pre>
What is the median survival? What does it mean? Explore if survival patterns between men and women are different:	age 1.017191 0.9989686 1.0357467
> kms <- survfit(Surv(time, status == 2) ~ sex, data = lung) > kms	What do these estimates mean?
Call: survfit(formula = Surv(time, status == 2) ~ sex, data = lung)	$\lambda(t, x) = \lambda_0(t) \exp(\beta_1 x_1 + \beta_2 x_2)$
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	Where is eta_1 ? Where is eta_2 ? Where is $\lambda_0(t)$? What is the mortality RR for a 10 year age difference?
Lung cancer survival (surv) 22 / 130	Lung cancer survival (serv) 27 / 130

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:

Lung cancer survival (sur-

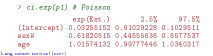
> ?poisreg

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?



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)
```

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

$$\begin{split} \mathrm{P}\left\{d \text{ at } t_x \mid \text{ entry at } t_e\right\} &= \mathrm{P}\left\{\text{survive } (t_e, t_1] \mid \text{alive at } t_e\right\} \times \\ & \mathrm{P}\left\{\text{survive } (t_1, t_2] \mid \text{alive at } t_1\right\} \times \\ & \mathrm{P}\left\{\text{survive } (t_2, t_x] \mid \text{alive at } t_2\right\} \times \\ & \mathrm{P}\left\{d \text{ at } t_x \mid \text{alive just before } t_x\right\} \end{split}$$

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Rates and likelihood

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

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

where the approximation gets better the smaller h is.

Dividing follow-up time

- \blacktriangleright Survival for a time span: $y=t_x-t_e$
- $\blacktriangleright\,$ 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:

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

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Dividing follow-up time

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

 $P\left\{\text{survive } t_e \text{ to } t_x\right\} \approx \left(1 - \frac{\lambda y}{N}\right)^N \to \exp(-\lambda y), \quad N \to \infty$ (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
- \blacktriangleright the probability of dying in the last tiny instant (of length $\epsilon)$ of the interval
- \blacktriangleright The probability of dying in this tiny instant is $\lambda\epsilon$
- ► log-likelihood contribution from this last instant is $\log(\lambda \epsilon) = \log(\lambda) + \log(\epsilon)$.

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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 the sum of the log-likelihood terms:

$$\ell(\lambda) = \sum_{i} (-\lambda y_i + d_i \log(\lambda) + d_i \log(\epsilon))$$
$$= \sum_{i} (d_i \log(\lambda) - \lambda y_i) + \sum_{i} d_i \log(\epsilon)$$

The last term does not depend on λ_i so it can be ignored

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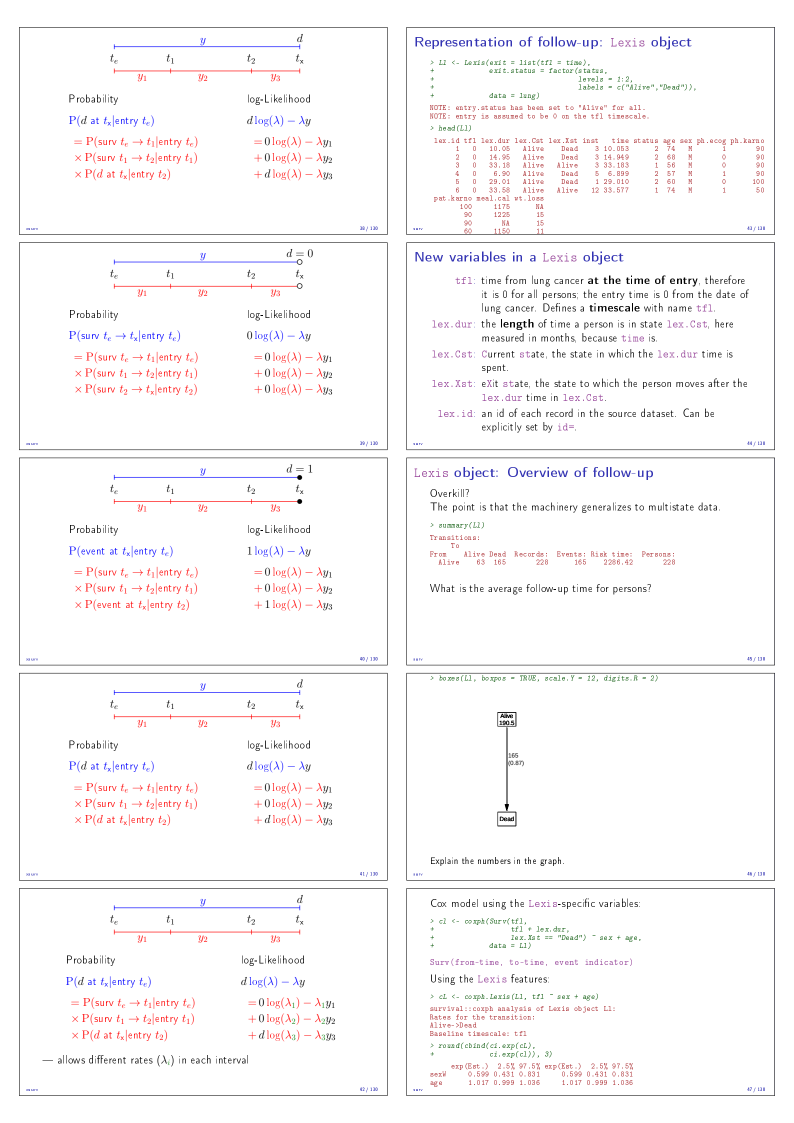
Total log-likelihood

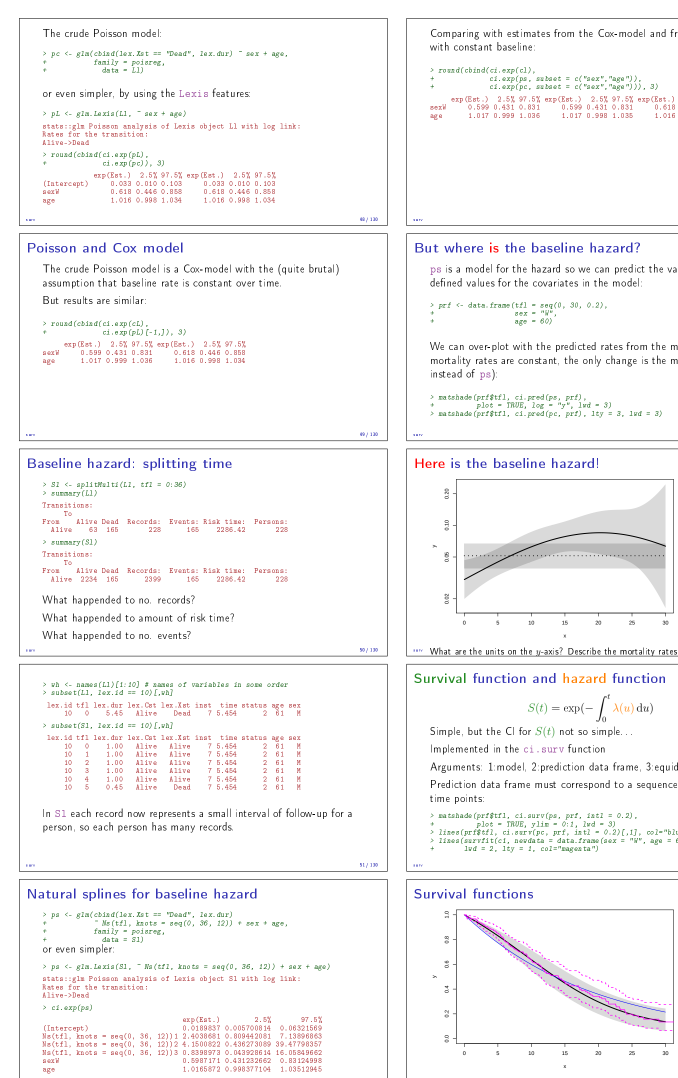
▶ ... for the follow up of 1 person is (the **rate** likelihood):

$$\sum_{i} \left(d_i \log(\lambda) - \lambda y_i \right)$$

- this is also the likelihood for independent Poisson variates d_i with means λy_i .
- even though the d_i s are neither Poisson nor independent
- Different models can have the same (log)likelihood:
 - model for follow-up of a person (d_i, y_i), constant rate λ
 model for independent Poisson variates (d_i), mean λy_i

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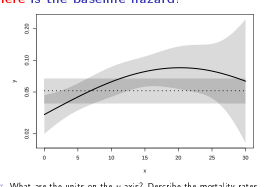
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Comparing with estimates from the Cox-model and from the model ci.exp(ps, subset = c("sex", "age")),
ci.exp(pc, subset = c("sex", "age"))), 3) exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5% exp(Est.) 2.5% 97.5% 0.599 0.431 0.831 0.599 0.431 0.831 0.618 0.446 0.858 1.017 0.999 1.036 1.017 0.998 1.035 1.016 0.998 1.034 53 / 130 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:

We can over-plot with the predicted rates from the model where mortality rates are constant, the only change is the model (pc

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



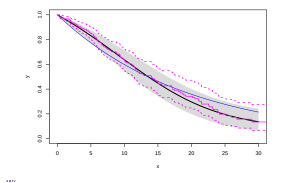
Survival function and hazard function

$$S(t) = \exp(-\int_0^t \lambda(u) \,\mathrm{d}u)$$

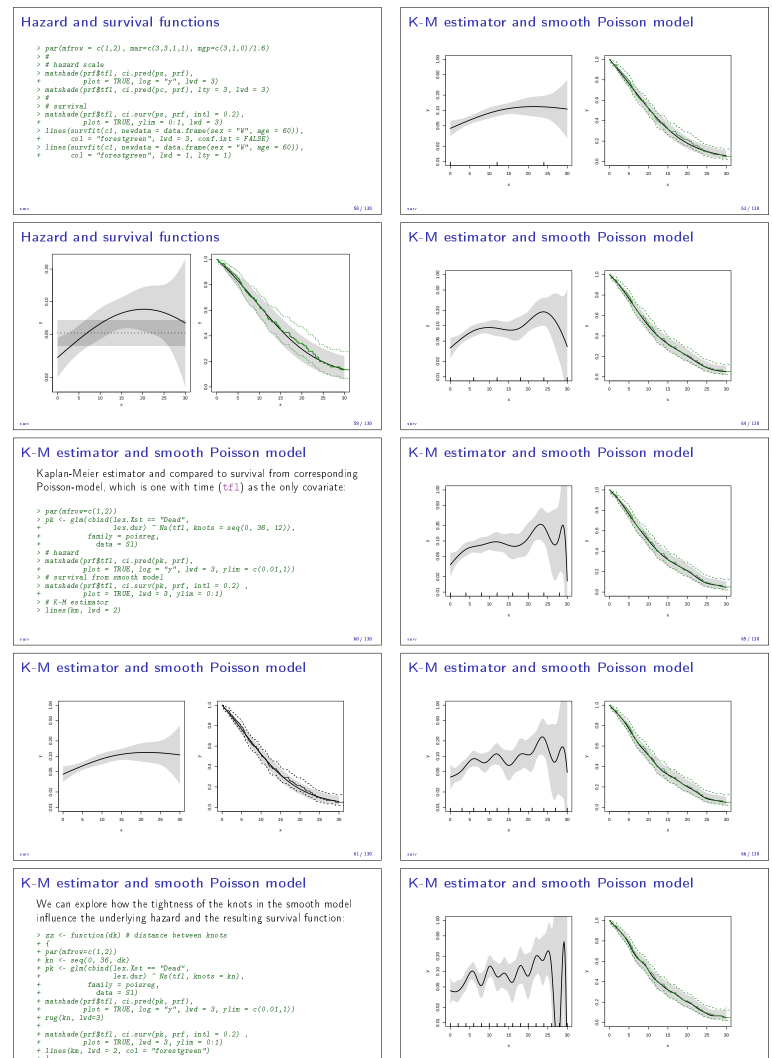
Simple, but the Cl for S(t) not so simple...

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

Prediction data frame must correspond to a sequence of equidistant



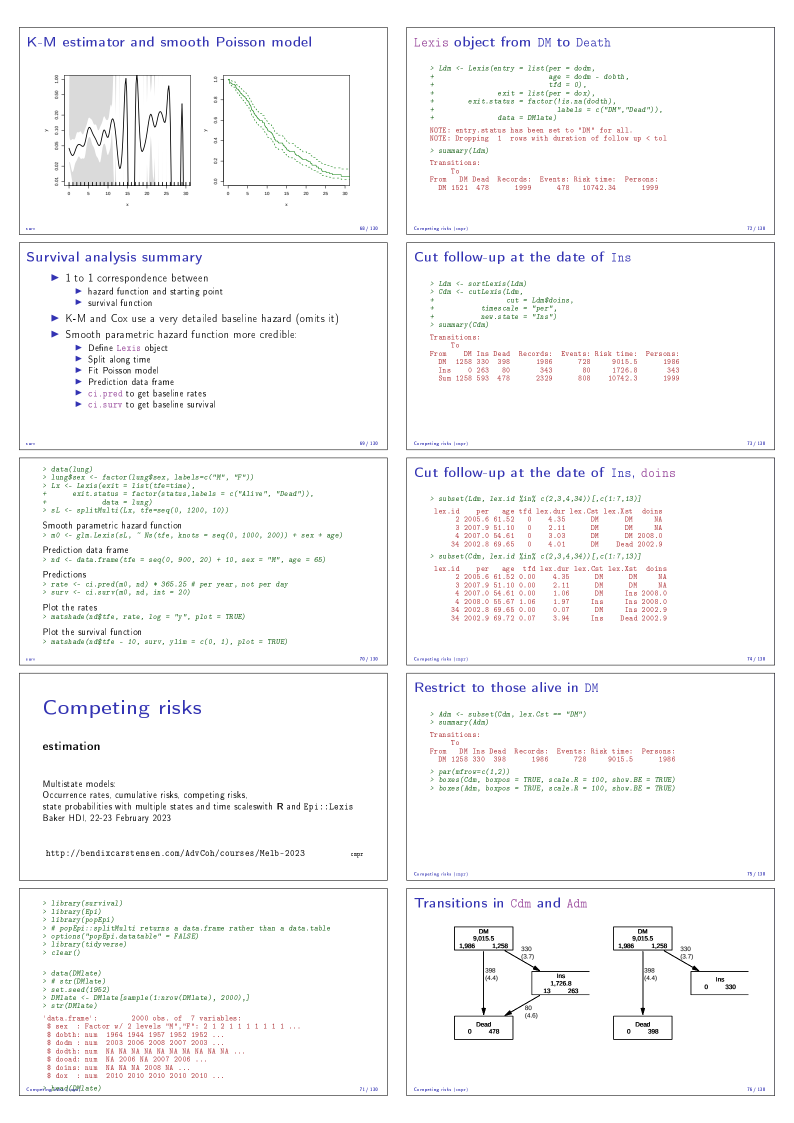
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> zz(12)



Survival function?

$$S(t) = \exp\left(-\int_0^t \lambda(u) + \mu(u) \, \mathrm{d}u\right)$$

$$S(t) = \exp\left(-\int_0^t \lambda(u) \, \mathrm{d}u\right)$$

$$S(t) = \exp\left(-\int_0^t \mu(u) \, \mathrm{d}u\right)$$

Competing risks (cmpr)

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Survival function?

- Regarding either Dead or Ins 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
 ightarrow DeadCompeting risks: the probability of being in each of the states DM, Ins and Dead —depends on two rates, $\texttt{DM} \rightarrow \texttt{Ins}$ and $\texttt{DM} \rightarrow \texttt{Dead}$

Competing risks (cmpr)

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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

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Survival function and Cumulative risk function

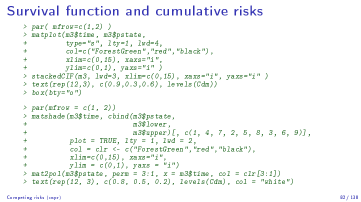
ex.Xst)		
s" "Dead"		
<pre>> m3 <- survfit(Surv(tfd, tfd + lex.dur, lex.Xst) ~ 1, + id = lex.id, + data = Adm) > # names(m3) > m3\$states</pre>		
s" "Dead"		
ime = m3\$time, m3\$pstate))		
0.99950 0.0000000 0.00050352 0.99748 0.001070 0.00151057 0.99547 0.0025184 0.00201435 0.99396 0.0040297 0.00201435 0.99295 0.0050373 0.00201435 0.98942 0.0085637 0.00201435		
	<pre>" "Dead" (Surv(tfd, tfd + lex.dur, lex.Xst) ' id = lex.id, ata = Adm) " "Dead" me = m3\$time, m3\$pstate)) 0.99950 0.0000000 0.00050352 0.99748 0.0010070 0.00151057 0.99547 0.0025184 0.00201435 0.99295 0.0050373 0.00201435</pre>	

____this is called the Aalen-Johansen estimator of state probabilities 80 / 130

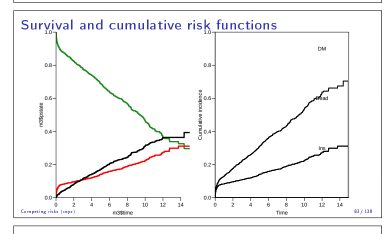
Survival function and cumulative risks-formulae $S(t) = \exp\left(-\int_0^t \lambda(u) + \mu(u) \,\mathrm{d}u\right)$ $R_{\text{Dead}}(t) = \int_0^t \mu(u) S(u) \, \mathrm{d}u$ $R_{\text{Ins}}(t) = \int_0^t \lambda(u) S(u) \, \mathrm{d}u)$ $= \int_0^{J_0} \lambda(u) \exp\left(-\int_0^u \lambda(s) + \mu(s) \,\mathrm{d}s\right) \mathrm{d}u$

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

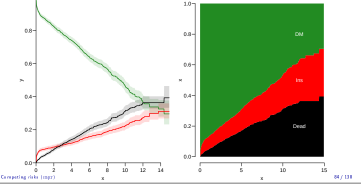












Survival function and cumulative risks-don't

$$\begin{split} S(t) &= \exp\left(-\int_{0}^{t} \lambda(u) + \mu(u) \,\mathrm{d}u\right) \\ R_{\text{Dead}}(t) &= \int_{0}^{t} \mu(u)S(u) \,\mathrm{d}u \\ R_{\text{Ins}}(t) &= \int_{0}^{t} \lambda(u)S(u) \,\mathrm{d}u \\ &= \int_{0}^{t} \lambda(u) \exp\left(-\int_{0}^{u} \lambda(s) + \mu(s) \,\mathrm{d}s\right) \,\mathrm{d}u \\ &\neq \int_{0}^{t} \lambda(u) \exp\left(-\int_{0}^{u} \lambda(s) \,\mathrm{d}s\right) \,\mathrm{d}u \\ &= 1 - \exp\left(-\int_{0}^{t} \lambda(s) \,\mathrm{d}s\right) - \text{nice formula, but wrong!} \\ \underset{\text{Completion bility of Ins assuming Dead does not exist and rate of Ins unchanged!}{} \end{split}$$

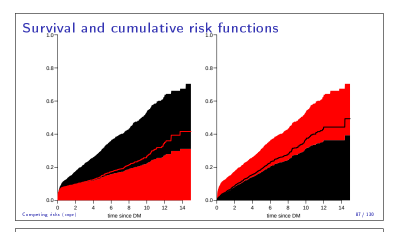
Survival function and cumulative risks-don't

```
m2 <- survfit(Surv(tfd,
tfd + lex.dur)
              lex.uur,
lex.Xst == "Ins" ) ~ 1,
data = Adm)
(Comp(15)
M2 <- survfit(Surv(tfd,
tfd + lex.dur,
lex.Xst == "De
"Dead") ~ 1,
```

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Competing risks (cmpr)



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.
- These models have no common parameters (effects of time or other covariates are not constrained to be the same).
- ... not for technical or statistical reasons, but for substantial reasons

it is unlikely that rates of different types of event (Insulin initiation and death, say) depend on time in the same way.

Competing risks (cmpr)

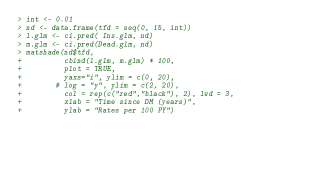
Cause-specific rates > Sdm <- splitMulti(Adm, tfd = seq(0, 20, 0.1)) > summary(Adm) Transitions: To DM Ins Dead Records: Events: Risk time: Persons 1258 330 398 1986 728 9015.5 1988 From DM 1258 330 398 1986 > summary(Sdm) Transitions: To From DM Ins Dead Records: Events: Risk time: Persons: DM 90419 330 398 91147 728 9015.5 1986

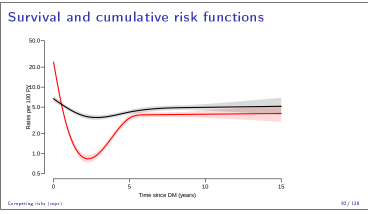
Competing risks (cmpr)

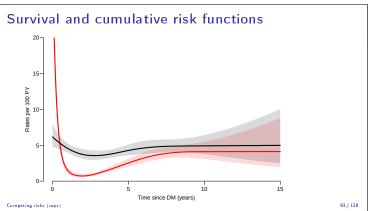
Cause-specific rates > round(cbind(
+ with(ent) + vinh(volma(volma(), lex.Xst == "Ins"), quantile(tfd + lex.dur, 0:4/4)),
+ with(subset(Sdm, lex.Xst == "Dead"), quantile(tfd + lex.dur, 0:4/4))), 2) [,1] [,2] 0% 0.01 0.01 25% 0.07 1.15 50% 1.07 3.01 75% 5.19 5.69 100% 13.74 14.38 > ikn <- c(0, 0.5, 3, 10) > dkn <- c(0, 2.0, 5, 9) > Ins.glm <- glm.Lexis(Sdm, ~ Ns(tfd, knots = ikn), to = "Ins") stats::glm Poisson analysis of Lexis object Sdm with log link: Rates for the transition: DM->Ins > 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 90 / 130 eting risks (cmpr

Cause-specific rates

Competing risks (cmpr)







Integrals with R

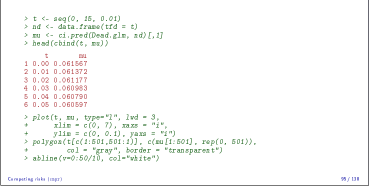
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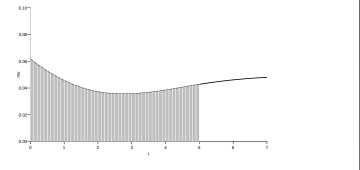
- Integrals look scary to many people, but they are really just areas under curves.
- ▶ In R, a curve of the function $\mu(t)$ is a set of two vectors: one vector of ts and one vector $y = \mu(t)s$.
- \blacktriangleright When we have a model such as the glm above that estimates the mortality as a function of time (tfd), we can get the mortality as a funtion of time by first choosing the timepoints, say from 0 to 15 years in steps of 0.01 year (≈ 4 days)
- ▶ Using ci.pred on this gives the predicted rates
- Then use the formuale with all the integrals to get the state probabilities.

Competing risks (cmpr)

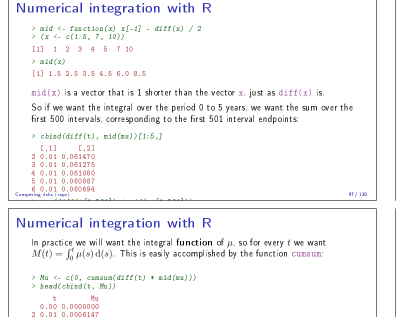
Integrals with R



Integrals with R



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0.00 0.0000000 0.01 0.0006147 0.02 0.0012274 4 0.03 0.0018383 5 0.04 0.0024471

6 0.05 0.0030541

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

Competing risks (cmpr)

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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.

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

Competing risks (cmpr

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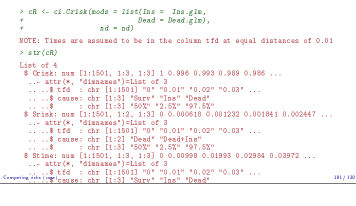
Simulation of cumulative risks: ci.Crisk

 1_{\cdot} a random vector from the multivariate normal distribution with

- mean equal to the parameters of the model,
 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
- $\ensuremath{\mathsf{3.}}$ derive state probabilities at these times by numerical integration
- 4. repeat to obtain, say, 1000 sets of state probabilities at these times
- 5. derive confidence intervals for the state probabilities as the 2.5 and 97.5 percentiles of the state probabilities at each time

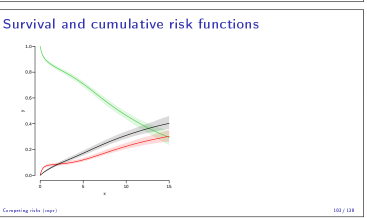
This machinery is implemented in the function ci.Crisk in Epi Competing disks (copr)

Cumulative risks from parametric models



Cumulative risks from parametric models So now plot the cumulative risks of being in each of the states (the Crisk component): > matshade(as.numeric(dimnames(cR\$Crisk)[[1]]), + cbind(cR\$Crisk[,1,], + cR\$Crisk[,2,], + cR\$Crisk[,2,], plot = TRUE, + lwd = 2, col = c("limegreen", "red", "black"))

Competing risks (cmpr)



Stacked probabilities: (matrix 2 polygons)

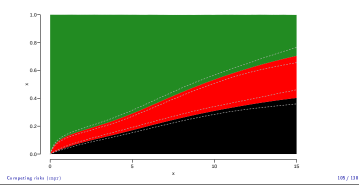
> mat2pol(cR\$Crisk[,3:1,1], col = c("forestgreen","red","black")[3:1])

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

The component ${\tt Srisk}$ has the confidence limits of the stacked probabilities:

Competing risks (cmpr)

Survival and cumulative risk functions



Expected life time: using simulated objects

The areas between the lines (up to say 10 years) are $\mbox{expected}$ sojourn times, that is:

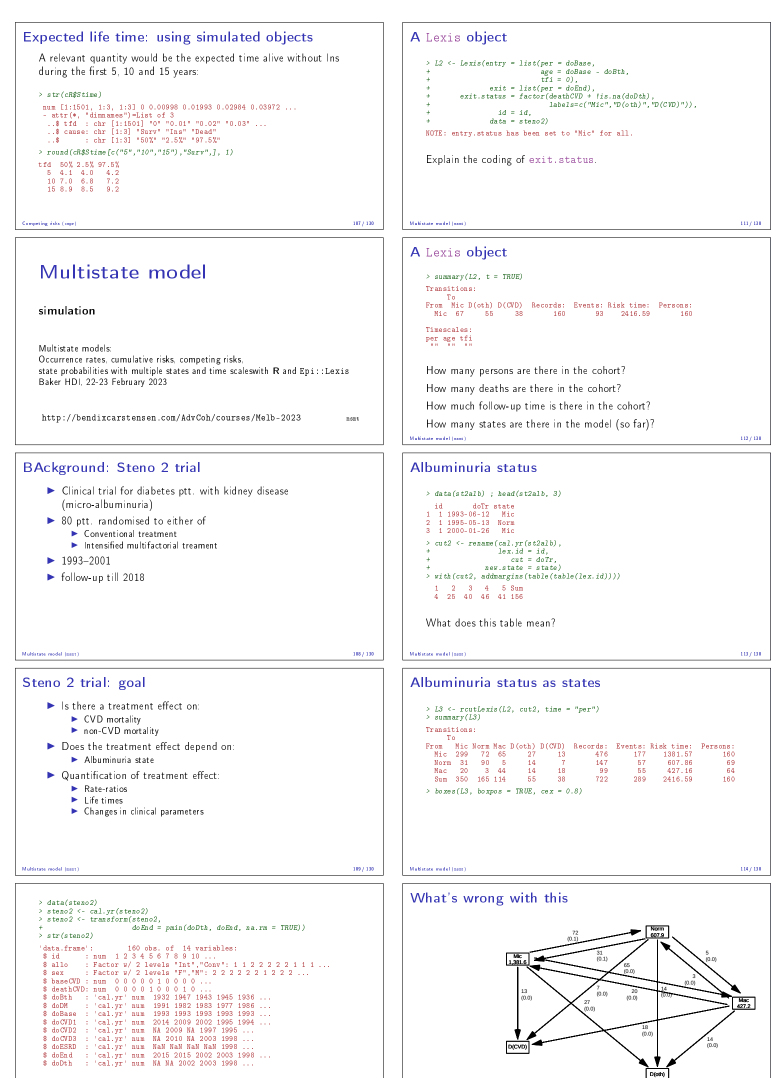
- expected years alive without Ins
- expected years lost to death without Ins
- expected years after Ins, including years dead after Ins

Not all 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).

Competing risks (cmpr)

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Multistate model (msmt)

Multistate model (msmt)

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