## **Biostatistics II: Survival Analysis**

#### **Dimitris Rizopoulos**

Department of Biostatistics, Erasmus University Medical Center



d.rizopoulos@erasmusmc.nl



@drizopoulos

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- We are interested in the Time until a prespecified event of interest occurs
  - $\triangleright$  time until a patient dies from a serious disease
  - ▷ time until metastasis
  - $\triangleright$  time until a machine breaks down

 $\triangleright \dots$ 

- Statistical analysis of time-to-event outcomes (aka Survival Analysis)
  - > Describe the distribution of the survival times
    - \* shape of the survival distribution
    - \* location measures, e.g., median survival time



▷ Inference, i.e., understand prognostic factors (strength and shape)

- \* is the new treatment prolonging survival time of patients?
- \* are non-smokers surviving longer than smokers?
- \* is the new machine lasting longer than the old one?
- \* statistical modelling



- Throughout this part we will use several equivalent names for the Time until the event of interest occurs, namely
  - ▷ time-to-event data
  - $\triangleright$  event time data
  - $\triangleright$  event times
  - $\triangleright$  survival times
  - $\triangleright$  survival data
  - $\triangleright$  failure times
  - $\triangleright$  failure time data



- We will learn which are the special characteristics of event time data and why they require special treatment (from a statistical point of view)
- From this part it will become clear
  - ▷ which statistical tools are applicable for this kind of data,
  - > which are their advantages and disadvantages, and
  - ▷ which are the optimal inferential strategies
- What is there further in survival analysis than what we will cover in this part



# • Part I: Introduction

- $\triangleright$  Data sets that we will use throughout this part
- $\triangleright$  Features of time-to-event data
- ▷ Censoring
- $\triangleright$  Truncation



## • Part II: Basic Tools in Survival Analysis

- ▷ Basic tools in survival analysis
  - \* Survival function
  - \* Cumulative distribution function
  - \* Density function
  - \* Hazard function
  - \* Cumulative hazard function
- $\triangleright$  Relationships between them



- Part III: Estimation & Statistical Inference
  - Basic notation for censored event time data
  - $\triangleright$  Estimating the survival function
    - \* the Kaplan-Meier estimator
    - \* the Breslow estimator
  - ▷ Comparing survival functions
    - \* the log-rank test
    - \* the Peto & Peto modified Gehan-Wilcoxon test



- **Part IV:** Regression Models for Time-to-Event Data
  - > Accelerated failure time models
  - Cox proportional hazards model
  - Parametric proportional hazards models
  - $\triangleright$  For each of the above
    - \* Estimation
    - \* Interpretation of parameters
    - \* Hypothesis testing
    - \* Effect plots
    - \* Checking the model's assumptions
    - \* General statistical modeling strategies



- **Part V:** Extensions of the Cox Model
  - ▷ Expected survival
  - $\triangleright$  Stratified Cox model
  - $\triangleright$  Time-dependent covariates
  - Clustered Event Time Data
  - ▷ Competing risks
  - $\triangleright$  Discrimination



- Lectures & Practice Sessions:
  - $\triangleright$  theory sessions
  - ▷ software practicals (build up approach)
- Software
  - $\triangleright$  practice sessions will be in R
  - b we will use online tutorials that provide you with hints on how to solve the exercises



- Material:
  - $\triangleright$  Course Notes
  - ▷ Survival Analysis in R Companion
- Within the course notes there are several examples of R code which are denoted by the symbol 'R> '
  - ▷ more examples in the Survival Analysis in R Companion & during the practicals

More than what we are going to cover



- Standard texts
  - Kalbfleisch, J. and Prentice, R. (2002). The Statistical Analysis of Failure Time Data, 2nd Ed.. New York: Wiley.
  - ▷ Cox, D. and Oakes, D. (1984). *Analysis of Survival Data*. London: Chapman & Hall.
  - Parmar, M. and Machin, D. (1996). Survival Analysis: A Practical Approach. New York: Wiley.
  - Description Freedom Freedom Freedom Press, P. (2000). Modeling Survival Data: Extending the Cox Model. New York: Springer-Verlag.
  - Harrell, F. (2001). Regression Modeling Strategies: With Applications to Linear Models, Logistic Regression, and Survival Analysis. New York: Springer-Verlag.



- Klein, J. and Moeschberger, M. (2003). Survival Analysis Techniques for Censored and Truncated Data. New York: Springer-Verlag.
- Kleinbaum, D. and Klein, M. (2005). Survival Analysis A Self-Learning Text. New York: Springer-Verlag.
- More theoretical texts
  - Fleming, T. and Harrington, D. (1991). Counting Processes and Survival Analysis. New York: Wiley.
  - Andersen, P., Borgan, O., Gill, R. and Keiding, N. (1993). Statistical Models Based on Counting Processes. New York: Springer-Verlag.



- Some of the books referenced above also contain software examples in R, SAS and other statistical software programs
- Intro to R with some code for survival analysis
  - ▷ Dalgaard, P. (2008) Introductory Statistics with R, 2nd Ed. New York: Springer-Verlag.
  - ▷ Venables, W. and Ripley, B. (2002) *Modern Applied Statistics with S*. New York: Springer-Verlag.



- Interaction will be important for the comprehension of all the material that we will cover
- Therefore, you are welcome to interrupt and ask questions

Part I

# Introduction



- Survival of 184 patients on the waiting list for the Stanford heart transplant program
- Outcomes of interest:
  - $\triangleright$  time to death
  - ⊳ age
  - $\triangleright$  T5 tissue mismatch score



- A study of prognostic variables in 228 lung cancer patients conducted by the North Central Cancer Treatment Group
- Outcomes of interest:
  - $\triangleright$  time to death
  - ⊳ age
  - $\triangleright$  sex
  - $\triangleright$  ECOG performance score (physician's estimate); values: 0 4
  - $\triangleright$  Karnofsky performance score (physician's estimate); values: 20, 30, ..., 100



- 467 HIV infected patients who had failed or were intolerant to zidovudine therapy (AZT)
- The aim of this study was to compare the efficacy and safety of two alternative antiretroviral drugs
- Outcomes of interest:
  - $\triangleright$  time to death
  - ▷ randomized treatment: 230 patients didanosine (ddl) and 237 zalcitabine (ddC)
  - $\triangleright$  gender
  - $\triangleright$  AZT: failure or intolerance



- Outcomes of interest:
  - ▷ prevOI: previous opportunistic infections
  - $\triangleright$  CD4 cell count measurements

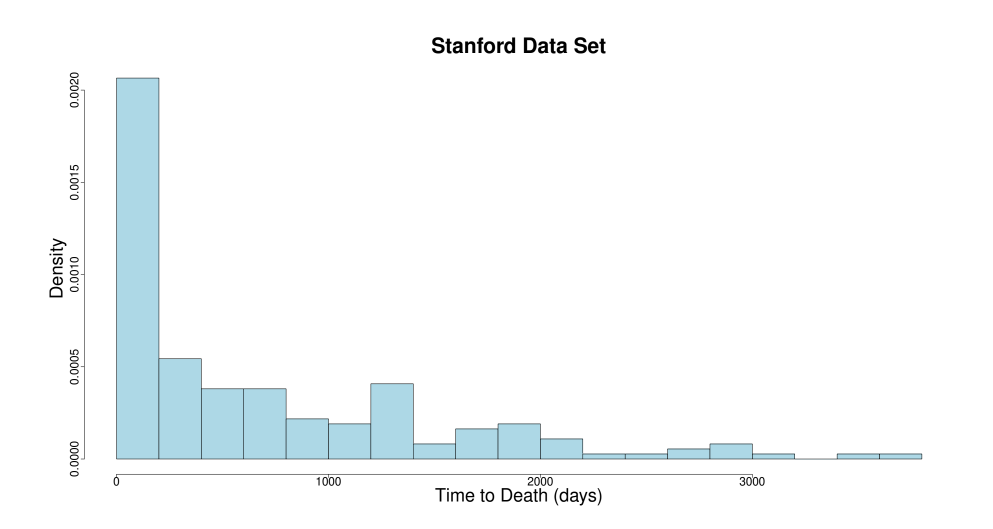


- Primary Biliary Cirrhosis (PBC):
  - ▷ a chronic, fatal but rare liver disease
  - > characterized by inflammatory destruction of the small bile ducts within the liver
- Data collected by Mayo Clinic from 1974 to 1984 (Murtaugh et al, Hepatology, 1994)
- Outcomes of interest:
  - ▷ time to death and/or time to liver transplantation
  - > randomized treatment: 158 patients received D-penicillamine and 154 placebo
  - $\triangleright$  age at baseline
  - Iongitudinal bilirubin levels

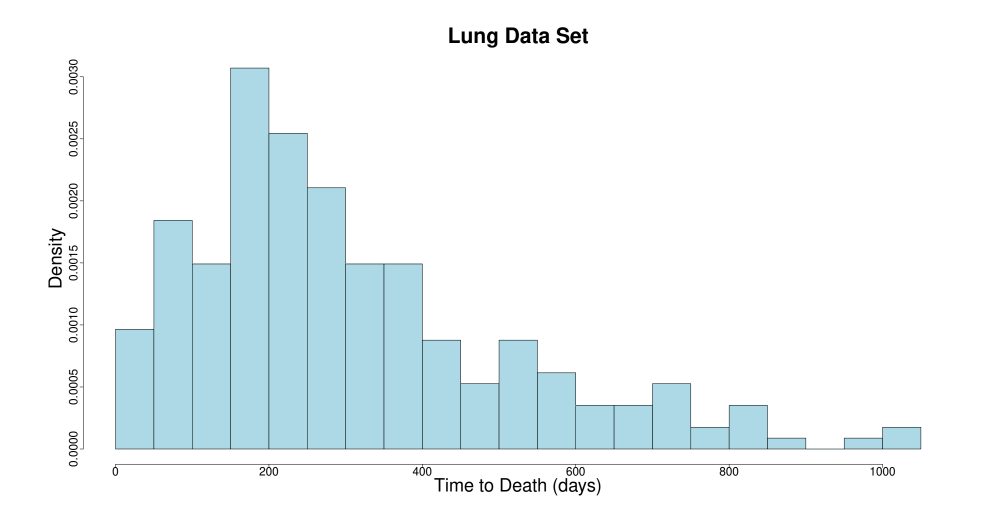


- 407 patients who underwent primary renal transplantation from deceased or living donor
- Outcomes of interest:
  - $\triangleright$  time to graft failure
  - ▷ smoking status
  - $\triangleright$  history of dialysis

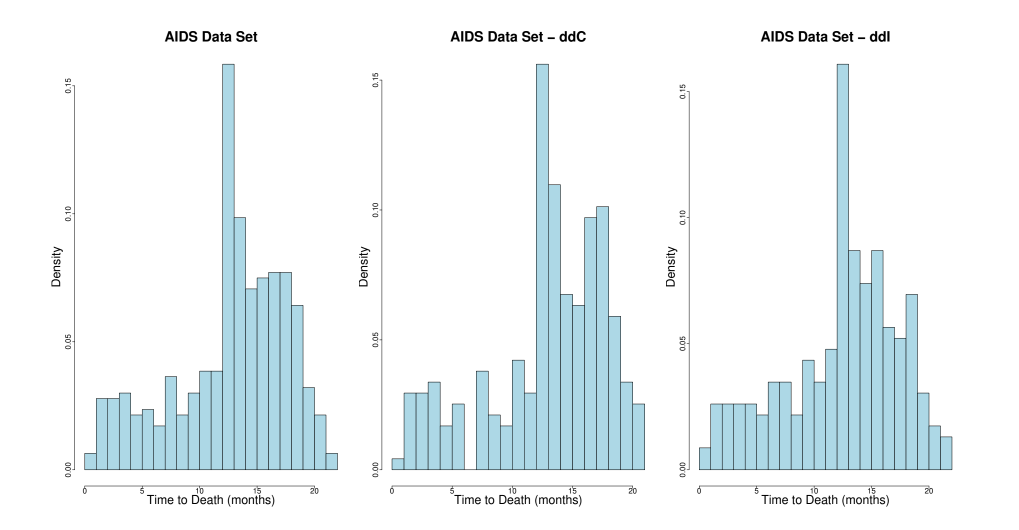




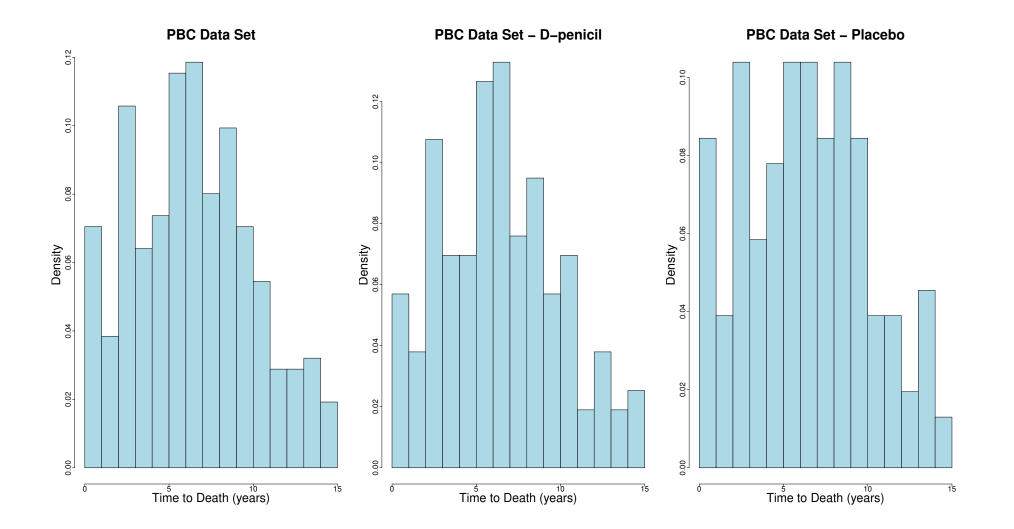




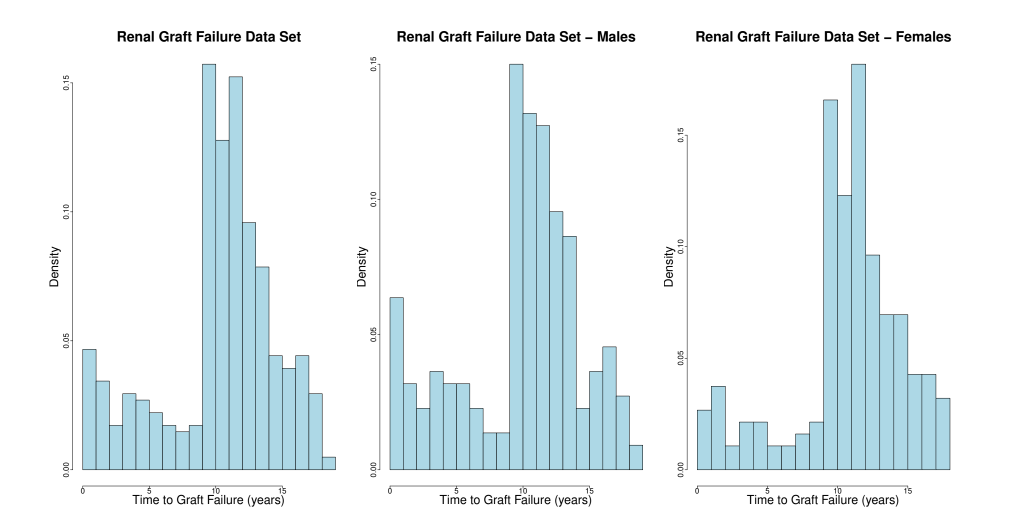












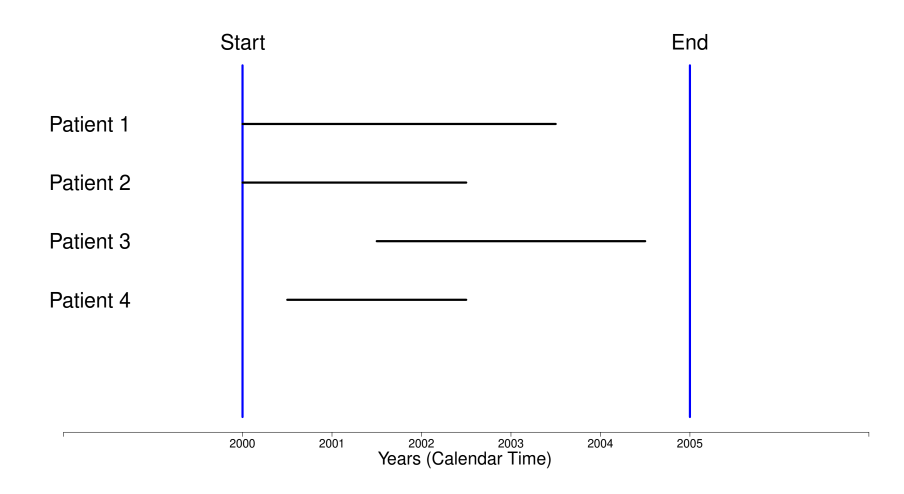


- Survival times are non-negative
  - ▷ in many cases the time to failure can have unusual distribution, i.e., does <u>not</u> look like a Normal
  - $\triangleright$  skewed to the right or to the left
- Naive analysis of untransformed times may produce invalid results

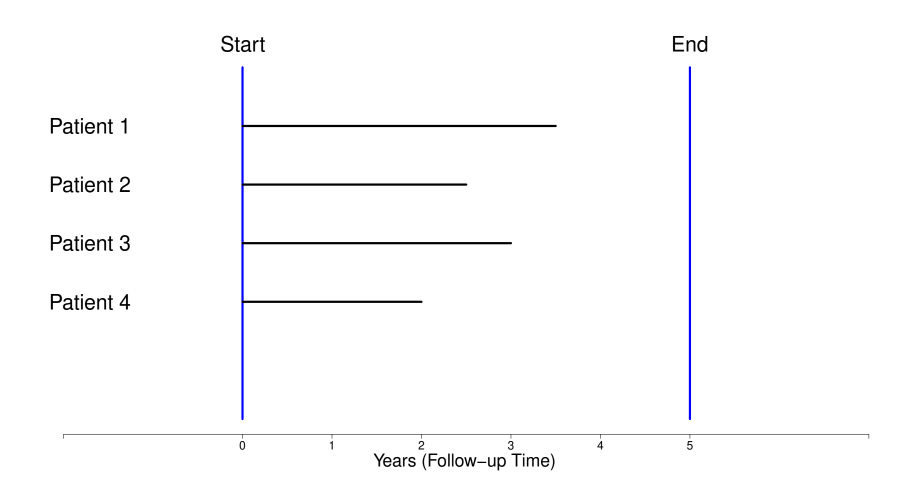


- Some patients enter the study at some point later than its start that is, at different calendar times
- In the analysis of failure time data we are only interested in the survival time that is, how long did the patient survive, i.e., how long was she *at risk* of the event
- **Crucial Assumption**: the distribution of survival times of those who enter early is the same as the distribution of the ones who enter late
  - ▷ this is violated if patients who enter later are expected to live longer (or shorter)







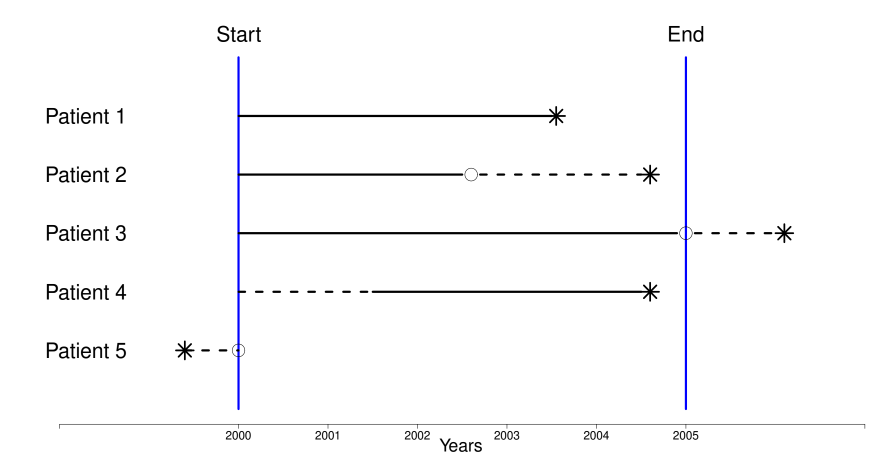


## 1.4 Censoring



- The time-to-event is only partially known for some patients in the study
- Types of censoring
  - ▷ right censoring
  - ▷ left censoring
  - ▷ interval censoring
- **Caution:** failure to take censoring into account can produce serious bias in estimates of the distribution of event times and related quantities







- Before talking in more detail about censoring ...
- Patients who had the event within the study period
  - $\triangleright$  Patient 1 was under observation from the start of the study until 3.5 years when he had the event  $\Rightarrow$  the time-to-event equals 3.5 years
  - $\triangleright$  Patient 4 enter the study after 1.5 years from the start (late entry), and she had the event at 4.6 years  $\Rightarrow$  the time-to-event equals 4.6 1.5 = 3.1 years
    - \* why can't we treat Patient 4 as observed for the full 5-year period since we know that she has survived 1.5 years?
    - \* had this patient died before 1.5 years, she would not have had the opportunity to enroll the study, and the event would have never been observed  $\Rightarrow$  <u>biases</u> survival time upwards

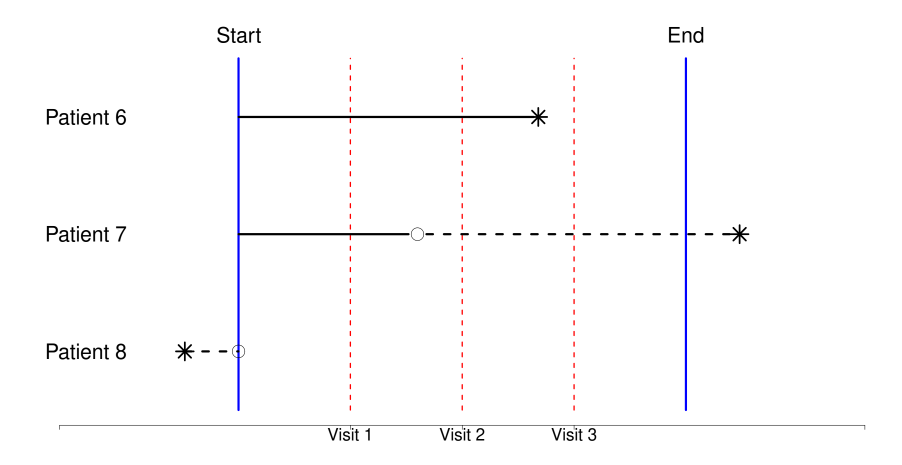


- Right censoring  $\Rightarrow$  the survival time is above a certain value
- Types of right censoring Examples:
  - $\triangleright$  Fixed type I: Patient 3 reached the end of the study  $\Rightarrow$  we know this patient had the event after 5 years
  - ▷ Fixed type II: a study ends when there is a prespecified number of events
  - $ightarrow \frac{Random}{Random}$ : Patient 2 moved to a new location at 2.6 years  $\Rightarrow$  we know this patient had the event after 2.6 years



- $\bullet$  Left censoring  $\Rightarrow$  the survival time is below a certain value
- Example:
  - ▷ Patient 5 had the event before the start of the study







- Interval censoring:  $\Rightarrow$  the survival time is between two values
- Example:
  - b during the study period there are 3 planned visits at which it is checked whether the event has occurred
  - $\triangleright$  Patient 6 did not yet have the event at Visit 2 but she had it at Visit 3  $\Rightarrow$  we know that she had the event in between Visits 2 and 3
  - $\triangleright$  Patient 7 did not yet have the event at Visit 1 and she left the study before Visit 2  $\Rightarrow$  we know that she had the event at some point after Visit 1
  - Patient 8 had the event before the stat of the study

Interval censoring includes left and right censoring as special cases



- Non-informative versus Informative Censoring
  - > a patient is excluded from the study because he decided to move to a new location from which he cannot easily reach the study center
  - ▷ a patient is excluded from the study because his condition deteriorates (e.g., adverse event) and his physician decides to give him a rescue medication
- What is the substantiative difference in the above two situations?



- Non-informative versus Informative Censoring
  - > a patient is excluded from the study because he decided to move to a new location from which he cannot easily reach the study center
  - ▷ a patient is excluded from the study because his condition deteriorates (e.g., adverse event) and his physician decides to give him a rescue medication
- What is the substantiative difference in the above two situations?
  - $\triangleright$  in the second case withdrawal at time c may indicate death is likely to happen sooner than might have been expected otherwise

Informative Censoring: lost to follow-up for reasons related to the event time



- Problems with informative censoring
  - $\triangleright$  biased estimates
  - ▷ inaccurate statistical inference
- <u>Note</u>: histograms revisited interpretation should be done with caution in the presence of censoring



- Truncation has a similar flavor to censoring (both are handled in a a similar manner analytically) but we should distinguish between the two terms
- Censoring period:
  - b during this period the subject is no longer under observation, but she may experience the event of interest
- Truncation period
  - b during this period the subject is no longer under observation, but she cannot experience the event of interest



- Similarly to censoring, there are 2 types of truncation
- <u>Left truncation</u>: a subject enters the population at risk at some stage after the start of the study, and we know that there is no way that the event of interest could have occurred <u>before</u> this date
- Right truncation: a subject leaves the population at risk at some stage after the start of the study, and we know that there is no way that the event of interest could have occurred <u>after</u> this date



- Leukemia patients are given a drug or placebo. Survival time is the duration from remission to relapse. The study ends at 52 weeks with some patients yet to relapse
  - A left censoring B right censoring
  - C left truncation
  - D right truncation



- Leukemia patients are given a drug or placebo. Survival time is the duration from remission to relapse. The study ends at 52 weeks with some patients yet to relapse
  - A left censoring B **right censoring** C left truncation
  - D right truncation



- College students are asked the age at which they first tried marijuana. Some answer never, and some report using it but forget when
  - A left censoring B right censoring
  - C left truncation
  - D right truncation



• College students are asked the age at which they first tried marijuana. Some answer never, and some report using it but forget when

A left censoringB right censoringC left truncationD right truncation



- The age at which children are able to count from 1–10 at school. Some children are already able to count before joining school
  - A true event
  - B interval censoring
  - C left truncation
  - D left censoring



- The age at which children are able to count from 1–10 at school. Some children are already able to count before joining school
  - A true event
  - B interval censoring
  - ${\sf C}$  left truncation
  - D left censoring



• For patients who have been hospitalized for a heart attack, we are interested in testing whether a new treatment that they take after they have been discharged prolongs survival. A patient died in the hospital

A left censoring B left truncation C true event

D interval censoring



• For patients who have been hospitalized for a heart attack, we are interested in testing whether a new treatment that they take after they have been discharged prolongs survival. A patient died in the hospital

A left censoring

## **B** left truncation

- C true event
- D interval censoring



- We are interested in identifying prognostic factors for the survival of ovarian cancer patients. Only patients who have survived at least 5 years after diagnosis are included in the study
  - A informative left truncation
  - B left truncation
  - C left censoring
  - D informative left censoring



- We are interested in identifying prognostic factors for the survival of ovarian cancer patients. Only patients who have survived at least 5 years after diagnosis are included in the study
  - A informative left truncation
  - **B** left truncation
  - ${\sf C}$  left censoring
  - D informative left censoring



- For patients who start feeling better, the physicians decide to exclude them from the study
  - A right truncation
  - B right censoring
  - C informative right truncation
  - D informative right censoring



- For patients who start feeling better, the physicians decide to exclude them from the study
  - A right truncation
  - B right censoring
  - ${\sf C}$  informative right truncation
  - D informative right censoring



- We are interested in the years spent in retirement. However, some died before getting retired
  - A left censoring
  - B left truncation
  - C right censoring
  - D right truncation



- We are interested in the years spent in retirement. However, some died before getting retired
  - A left censoring B **left truncation**
  - ${\sf C}$  right censoring
  - D right truncation



- Time-to-event data exhibit special characteristics:
  - $\triangleright$  skewed distributions
  - ▷ censoring and/or truncation
- Standard statistical tools do not work optimally for survival data  $\Rightarrow$  specialized statistical techniques are required

Part II

## **Basic Tools in Survival Analysis**



- We define T to be a positive random variable denoting the time-to-event
- $\bullet$  There are many ways to represent and describe the distribution of T
- The most useful in survival analysis is the *Survival Function*

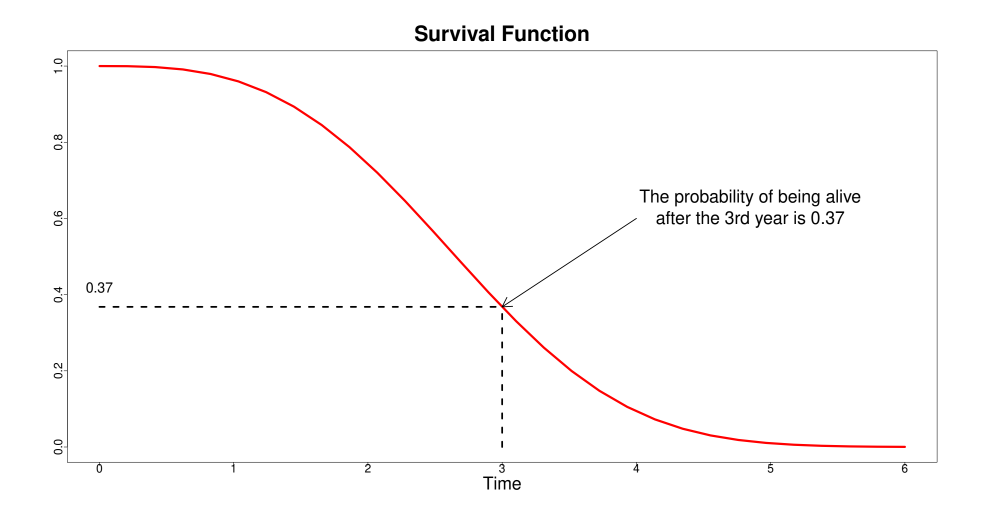
$$S(t) = \Pr(T > t)$$

• It denotes the probability of being alive up to time t (i.e., dying after t)



- Properties of the survival function
  - $\triangleright$  it is constrained between 0 and 1
  - $\triangleright$  it is a decreasing function of time, i.e.,
    - \* at time t = 0 all patients are alive
    - \* at time  $t=\infty$  all patients have died
- Note: in some settings patients can be cured and thus, it may not be reasonable to assume that all patients would die from the disease under study
  - ▷ a class of statistical models (aka Cure rate models) has been developed to deal with such phenomena (outside the scope of this course)







- The survival function is related to the cumulative distribution and the probability density function
- The *Cumulative Distribution Function* (CDF)

 $F(t) = \Pr(T \le t) = 1 - S(t)$ 

denotes the probability of dying until time  $\boldsymbol{t}$ 

- Properties of the CDF
  - $\triangleright$  it is constrained between 0 and 1
  - $\triangleright$  it is an increasing function of time

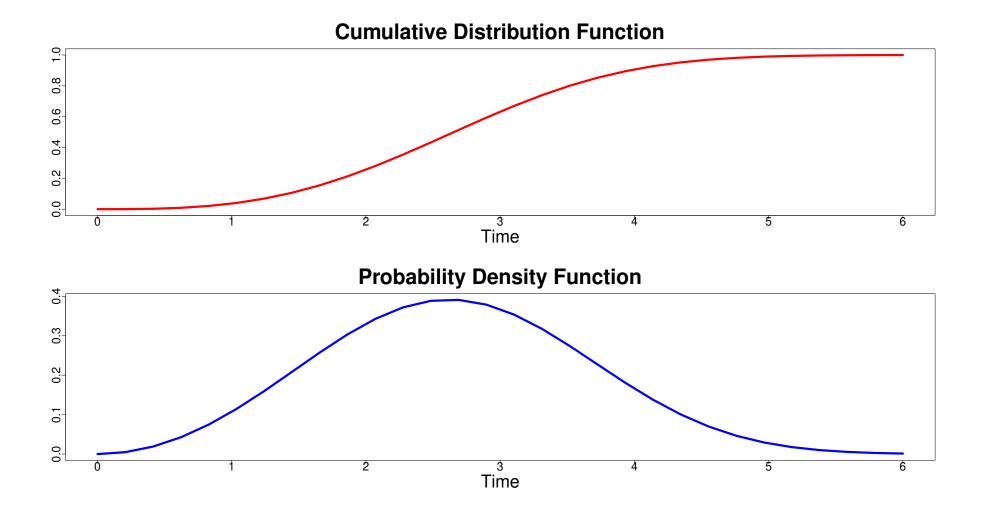


• The *Probability Density Function* (pdf)

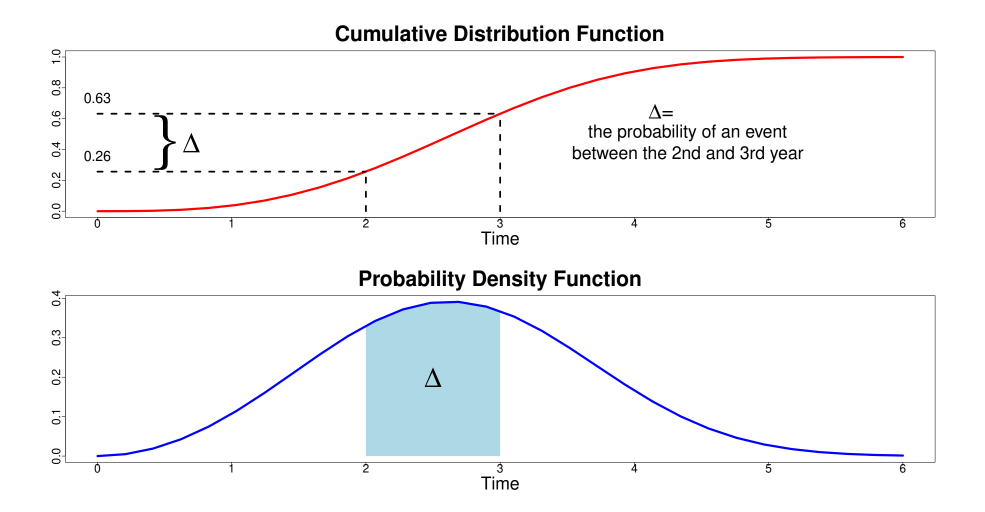
$$f(t) = \frac{dF(t)}{dt} \quad \text{ or } \quad F(t) = \int_0^t f(s) \; ds$$

denotes how dense is the probability of dying in a specific time interval











- Another useful notion is the risk of an event
- The Hazard Function

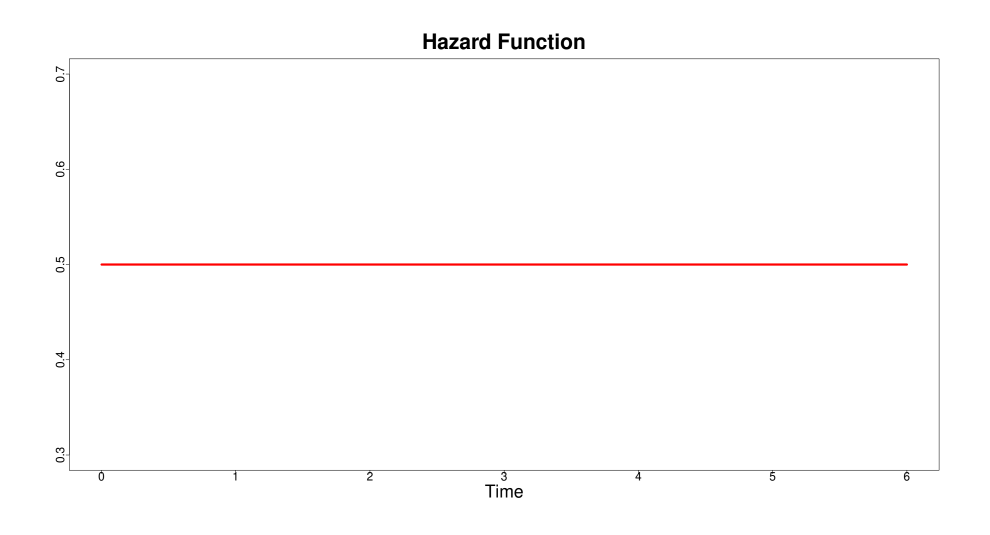
$$h(t) = \lim_{s \to 0} \frac{\Pr(t \le T < t + s \mid T \ge t)}{s}$$

is the instantaneous risk of an event at time t, given that the event has not occurred until time t

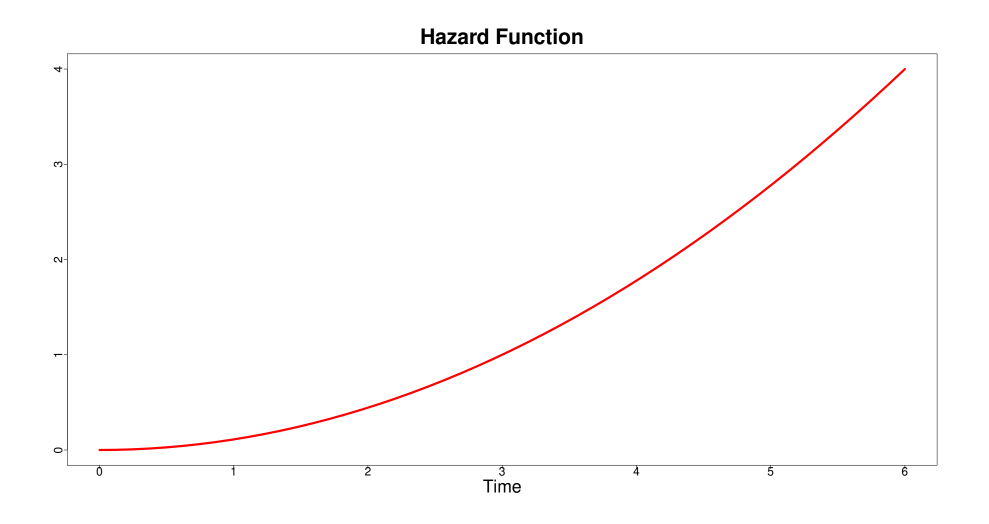


- Note: the hazard is <u>not</u> a probability ⇒ can be interpretable as the expected number of events per individual per unit of time
  - $\triangleright$  it has to be positive
  - ▷ but it can be (much) greater than 1

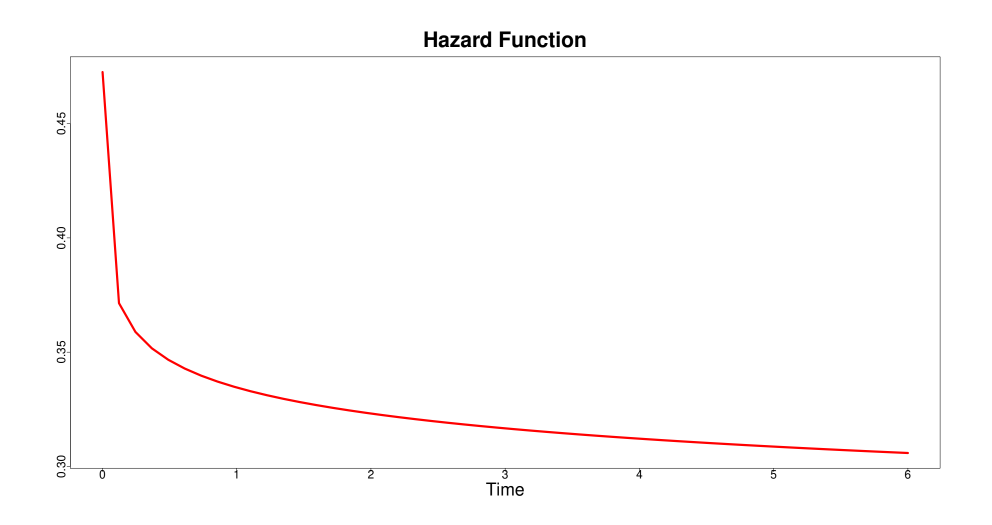




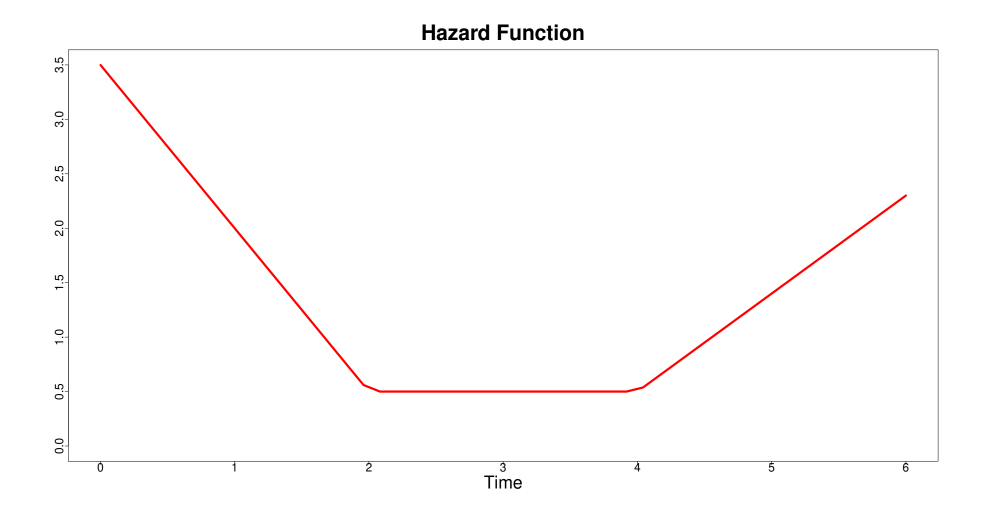




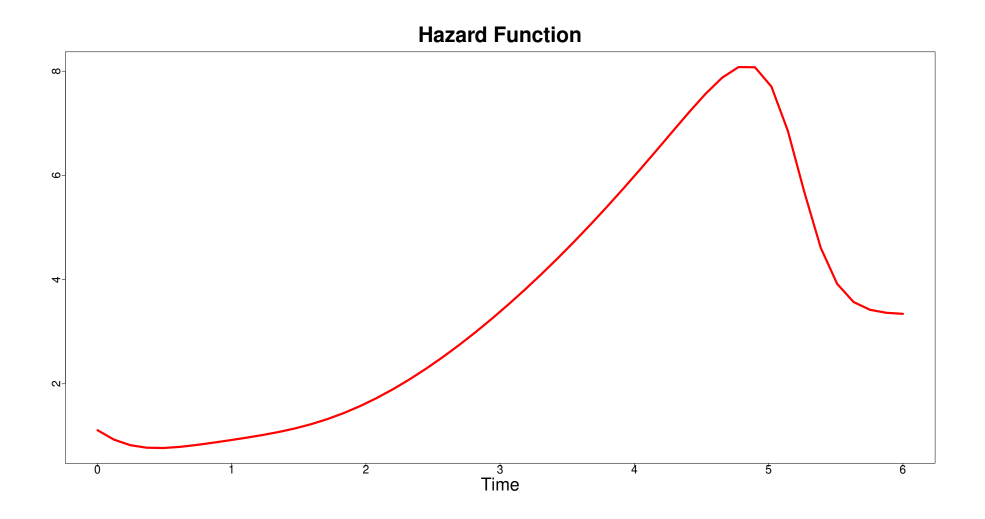














• The *Cumulative Hazard Function* is the integrated hazard function:

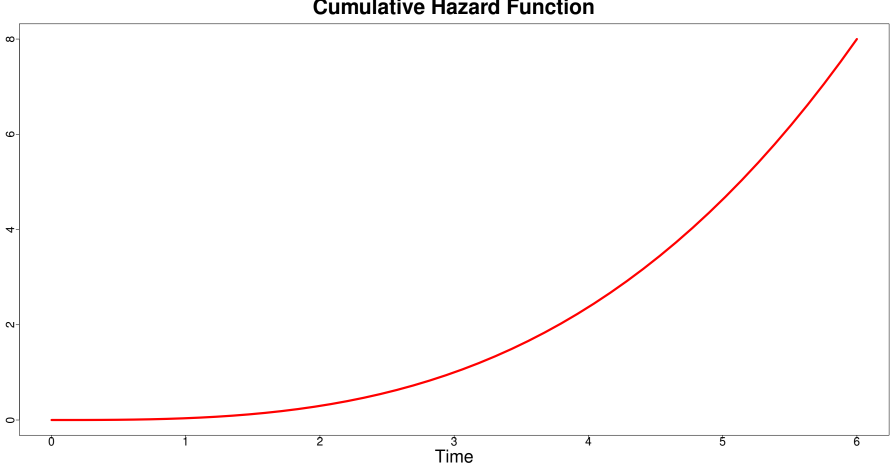
$$H(t) = \int_0^t h(s) \ ds$$

denotes the cumulative risk up to time t, i.e., the expected number of events that have occurred by time t

- Again this is <u>not</u> a (cumulative) probability
  - $\triangleright$  has to be positive
  - > increasing function of t (as the time progresses we expect more events to have occurred)

# 2.4 The Cumulative Hazard Function (cont'd)

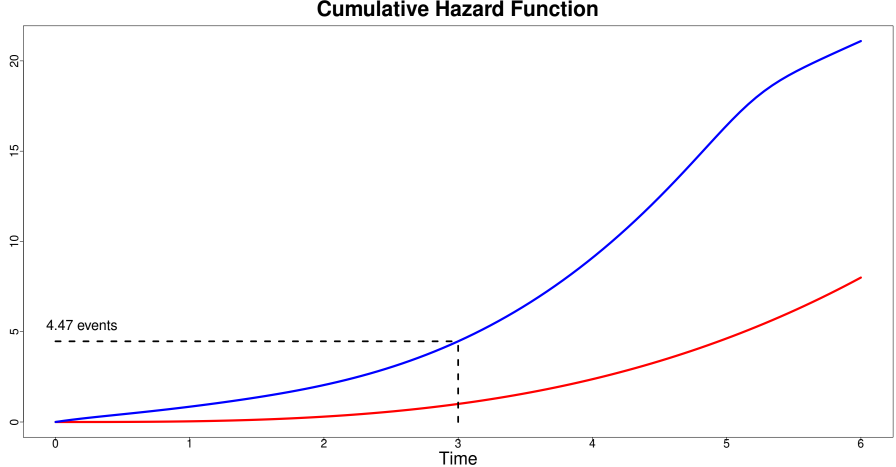




#### **Cumulative Hazard Function**

# 2.4 The Cumulative Hazard Function (cont'd)





**Cumulative Hazard Function** 

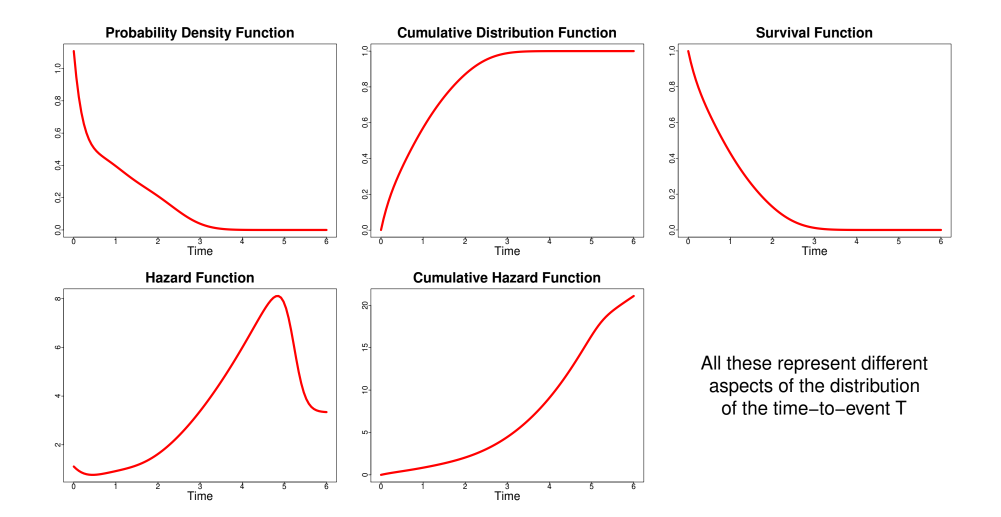


• All functions we have seen so far are related

▷ if you know one you know all!

$$f(t) = \frac{dF(t)}{dt} \qquad F(t) = 1 - S(t)$$
$$h(t) = \frac{f(t)}{S(t)} \qquad H(t) = \int_0^t h(s) \, ds$$
$$H(t) = -\log S(t)$$
$$S(t) = \exp\{-H(t)\} = \exp\{-\int_0^t h(s) \, ds$$







• *Median Life Length* or *Median Survival* is the time by which half of the subjects will experience the event – it is defined as

$$T_{0.5} = S^{-1}(0.5)$$

$$= H^{-1}(\log 2)$$

where  $S^{-1}(\cdot)$  and  $H^{-1}(\cdot)$  are the inverse survival and cumulative hazard functions, respectively



• *Mean Survival* or *Average Survival* is the expected failure time – is defined as

$$\mu = \int_0^\infty t f(t) \, dt$$

$$= \int_0^\infty S(t) \ dt$$

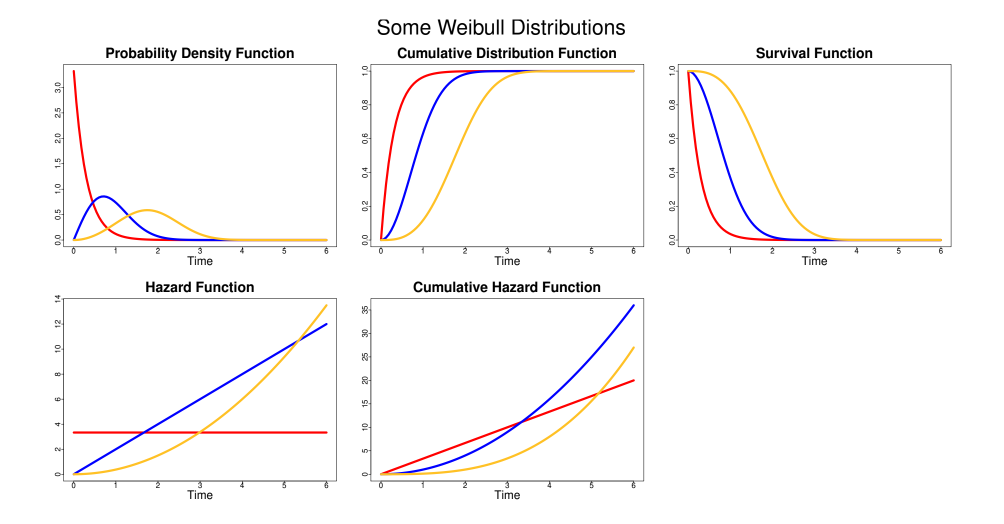
• *Expected Future Lifetime* is the expected value of future lifetime given survival up to time point  $t_0$ 

$$\tilde{\mu} = \frac{1}{S(t_0)} \int_0^\infty t f(t+t_0) dt$$

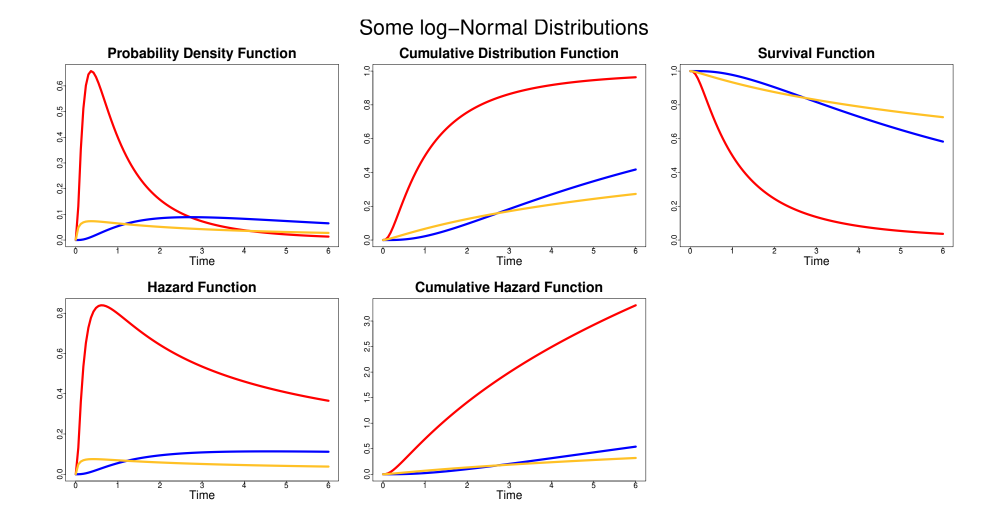


- In the literature there have been proposed many distributions for time-to-event random variables
- Some of the most popular are
  - ▷ Weibull (it has as special case the Exponential)
  - ▷ Gamma (it has as special case the Exponential)
  - ⊳ log-Normal
  - $\triangleright$  log-Student's-t
  - $\triangleright$  log-Logistic











- The basic functions to describe the distribution of time-to-event data
  - $\triangleright$  survival function
  - $\triangleright$  cumulative distribution and probability density function
  - $\triangleright$  hazard and cumulative hazard function
- All these functions are related



- Statistical measures
  - ▷ median survival
  - ▷ mean survival
  - ▷ expected future lifetime

Part III

### **Estimation & Statistical Inference**



- We have a sample of failure times  $\Rightarrow$  What is the available information?
- Notation (*i* denotes the patient)
  - $\triangleright T_i^*$  'true' time-to-event
  - $\triangleright$  because of censoring we do <u>not</u> always observe  $T_i^*$
  - $\triangleright C_i$  the censoring time (e.g., the end of the study or a random censoring time)
- Available data for each patient

 $\triangleright$  observed event time:  $T_i = \min(T_i^*, C_i)$ 

 $\triangleright$  event indicator:  $\delta_i = 1$  if event;  $\delta_i = 0$  if censored



Patient	$T_i^*$	$C_i$	$T_i$	$\delta_i$
1	3.5		3.5	1
2	3.4	2.2	2.2	0
3	5.7	5	5	0
:	:	:	:	:

The end of the study is at 5 years



- Based on the available information  $\{T_i, \delta_i\}$  we wish to estimate various quantities of interest, e.g.,
  - $\triangleright$  the Survival function
  - ▷ mean survival time
  - $\triangleright$  median survival time
  - $\triangleright$  specific quantiles
    - \* by which follow-up time 25% of the patients is still alive

 $\triangleright \dots$ 



- Aim: estimate the Survival Function S(t) based on a sample of failure times  $T_1, \ldots, T_n$ 
  - $\triangleright$  <u>Remember</u>: S(t) is the probability of being alive at time t (see Section 2.1)
- If there was no censoring, we could simply

$$\hat{S}(t) = \frac{\text{number of patients alive at time } t}{n} = \frac{1}{n} \sum_{i=1}^{n} I(T_i > t)$$

where  $I(T_i > t)$  equals 1 if  $T_i > t$ , and 0 otherwise

• However, we <u>do</u> have censored observations



- To take into account censoring in the estimation of S(t) we will use the law of total probability
- For instance, the probability of surviving 2 years can be computed as:

$$S(2) = \Pr(T_i > 2) = \Pr(T_i > 1) \times \Pr(T_i > 2 \mid T_i > 1)$$

• In words, the probability of surviving year 2 is the product of

 $\triangleright$  the probability of surviving year 1 and

 $\triangleright$  the conditional probability of surviving up to year 2 given still being alive at year 1



 $\bullet$  So S(2) can be estimated by

$$\hat{S}(2) = \frac{\# \text{ patients alive at year 1}}{\# \text{ patients at risk up to year 1}} \times \frac{\# \text{ patients alive at year 2}}{\# \text{ patients at risk up to year 2}}$$

 $\bullet$  If we apply this idea repeatedly, we can obtain survival probabilities for every time point t



- Let  $t_1$ ,  $t_2$ , ...,  $t_k$  denote the unique event times in the sample at hand
- We account for censoring by suitably adjusting the risk set ⇒ the Kaplan-Meier Estimator

$$\hat{S}_{KM}(t) = \prod_{i: t_i \le t} \frac{r_i - d_i}{r_i}$$

where  $d_i$  is the number of events at time  $t_i$ , and  $r_i$  the number of patients still at risk at time  $t_i$ 

 $\triangleright$  still at risk means alive and not censored



• A small example

 $1 5^+ 6 6 8 8^+ 9 11^+$ 

+ denotes a censored time

i	$t_i$	$r_i$	$d_i$	$(r_i - d_i)/r_i$
1	1	8	1	7/8
2	6	6	2	4/6
3	8	4	1	3/4
4	9	2	1	1/2



$$\hat{S}_{KM}(t) = 1, \quad 0 \le t < 1$$

$$= 7/8 = 0.875, \quad 1 \le t < 6$$

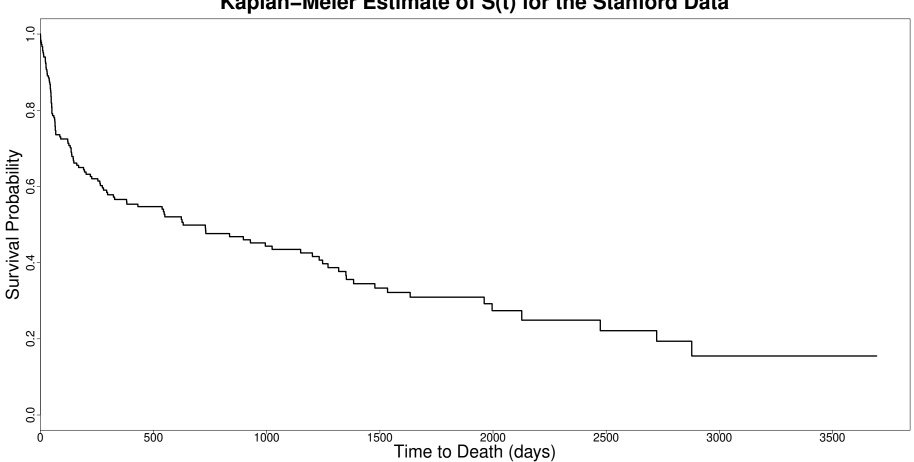
$$= (7/8)(4/6) = 0.583, \quad 6 \le t < 8$$

$$= (7/8)(4/6)(3/4) = 0.438, \quad 8 \le t < 9$$

$$= (7/8)(4/6)(3/4)(1/2) = 0.219, \quad 9 \le t < 11$$

 $\underline{\text{Note:}}$  the estimate of S(t) is undefined for t>11 since not all subjects have died by t=11





#### Kaplan–Meier Estimate of S(t) for the Stanford Data



- The variance of  $\hat{S}_{KM}(t)$  can be estimated using Greenwood's formula
- Using the formula and asymptotic normality of  $\hat{S}_{KM}(t)$ , we can derive a 95% confidence interval
- <u>Problem</u>: this can exceed 1 or fall below 0!
- A better asymmetric 95% confidence interval for  $\hat{S}_{KM}(t)$  that respects the boundaries is derived from a symmetric 95% confidence interval for either

$$\hat{H}_{KM}(t) = -\log \hat{S}_{KM}(t) \quad \text{or} \quad \log \hat{H}_{KM}(t) = \log\{-\log \hat{S}_{KM}(t)\}$$



• An estimate for the variance of  $\log \hat{H}_{KM}(t)$  is obtained by

$$\hat{\mathsf{var}}\{\log \hat{H}_{KM}(t)\} = \frac{\sum_{i:t_i \le t} d_i / \{r_i(r_i - d_i)\}}{\left[\sum_{i:t_i \le t} \log\{(r_i - d_i)/r_i\}\right]^2}$$

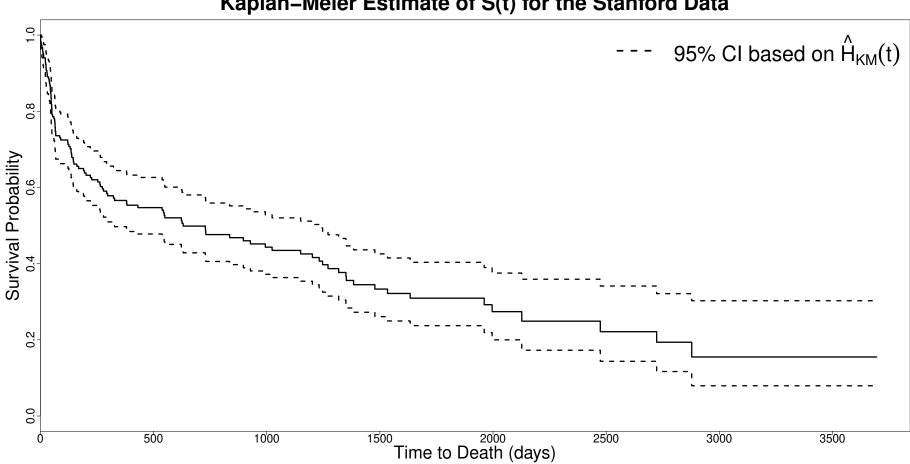
• Based on the estimated variance for  $\log \hat{H}_{KM}(t)$  we calculate the confidence interval

$$[a,b] = \log \hat{H}_{KM}(t) \mp 1.96 \times \sqrt{\mathsf{var}\{\log \hat{H}_{KM}(t)\}}$$

the confidence intervals for  $\hat{S}_{KM}(t)$  is then obtained as

$$\left[\exp\{-\exp(b)\}, \exp\{-\exp(a)\}\right]$$









**R>** Survival analysis in R

- > the basic package for survival analysis in R is the **survival** package
- ▷ this is a recommended package, i.e., you do not have to separately install it; you automatically install it whenever you install R
- $\triangleright$  however, in order to use it, you will need to load it using the command

library("survival")

R> We will also need data from the JM package
> you either need to install this package using install.packages("JM")
> or directly load the R workspace from CANVAS



- R> A key function in R that is used to specify the available event time information in a sample at hand is Surv()
- R> For right censored failure times (i.e., what we will see in this course) we need to provide the observed event times time, and the event indicator status, which equals 1 for true failure times and 0 for right censored times

Surv(time, status)



R> The function that is used to produce the Kaplan-Meier estimate of a survival function is survfit() – for the Stanford data we have

```
KM <- survfit(Surv(time, status) ~ 1, data = stanford2)</pre>
```

plot(KM)

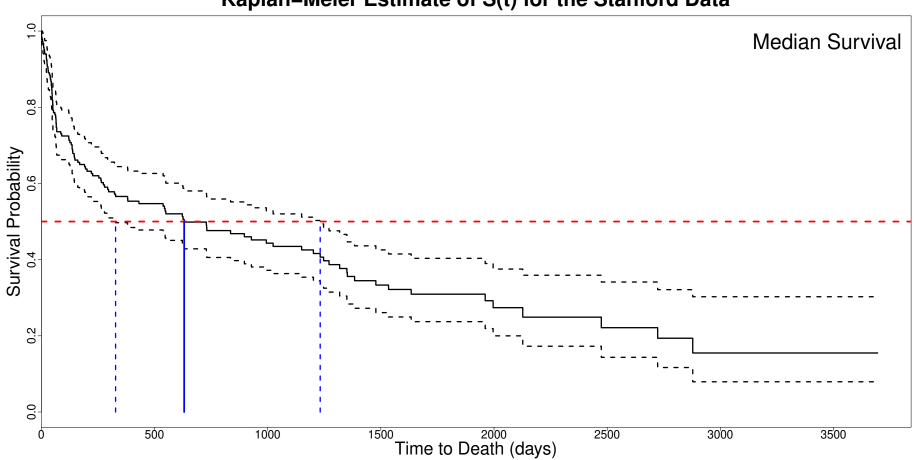


- The estimated survival function can be used to extract estimates of specific percentiles of interest, such as the median survival time (see Section 2.6)
- The following procedure is followed:
  - ▷ draw a horizontal line at the specific probability level of interest (e.g., 0.5 for the median survival time, 0.25 for the 1st quantile, etc.)
  - b the time point at which this horizontal line intersects with the survival curve is the estimated survival time of interest
  - ▷ a 95% confidence interval for this survival time is obtained by extracting the times at which the horizontal line intersects with the 95% confidence interval of the survival function



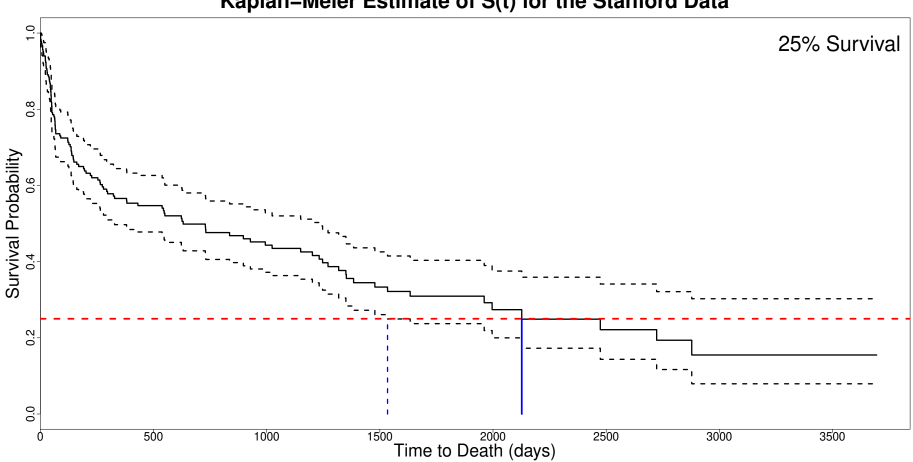
- If the horizontal line does not cross either the survival curve or its confidence interval, then the desired percentile and/or its confidence limits cannot be specified from the nonparametric estimate of S(t)
- Example: for the Stanford data and based on the Kaplan-Meier estimate of the survival function we compute
  - ▷ the median survival time (i.e., the time at which 50% of the patients is still alive)
  - ▷ 1st quantile survival time (i.e., the time at which 25% of the patients is still alive)





### Kaplan–Meier Estimate of S(t) for the Stanford Data





Kaplan–Meier Estimate of S(t) for the Stanford Data



# • We obtain

% Alive	Time	95% Lower	95% Upper
	days	Limit (days)	Limit (days)
50	631	328	1232
25	2127	1534	NA



R> We use the build-in function quantile(); for example,

KM <- survfit(Surv(time, status) ~ 1, data = stanford2)
quantile(KM, 1 - c(0.25, 0.50))</pre>



- $\bullet$  Using a similar approach to the Kaplan-Meier, we can also estimate the cumulative hazard function H(t)
- Remember: H(t) denotes the expected number of events up to and including time point t (see Section 2.4)
- $\bullet$  So a natural estimator of H(t) is

$$\hat{H}_{NA}(t) = \sum_{i: t_i \le t} \frac{d_i}{r_i}$$

which is called the Nelson-Aalen estimator



- <u>Remember</u>: the cumulative hazard function is related to the survival function (see Section 2.5)
- Therefore, an estimator for the survival function based on the Nelson-Aalen estimator is

$$\hat{S}_B(t) = \exp\{-\hat{H}_{NA}(t)\} = \prod_{i: t_i \le t} \exp(-d_i/r_i)$$

which has been suggested by Breslow and therefore it is known as the *Breslow Estimator* 



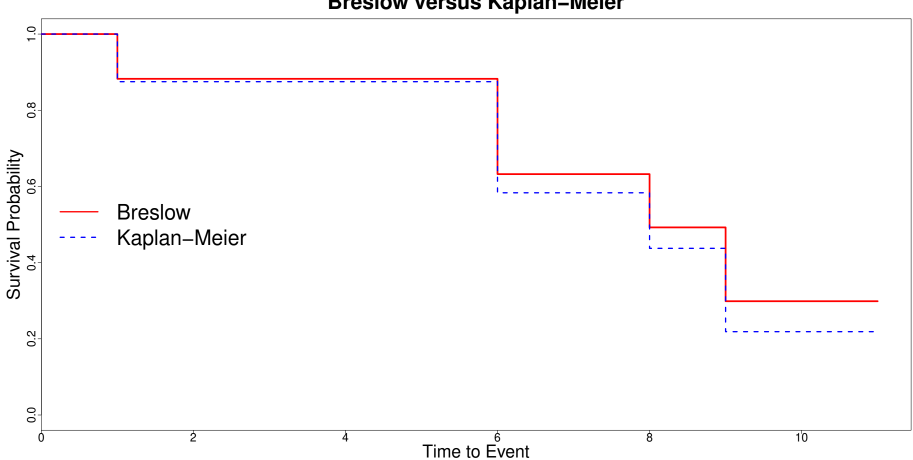
• Using the same example

$$1 5^{+} 6 6 8 8^{+} 9 11^{+}$$

 $+\ {\rm denotes}\ {\rm a}\ {\rm censored}\ {\rm time}$ 

i	$t_i$	$r_i$	$d_i$	$\frac{r_i - d_i}{r_i}$	$d_i/r_i$	$\hat{S}_B(t)$
1	1	8	1	7/8	1/8	$\exp(-1/8) = 0.882$
2	6	6	2	4/6	2/6	$\exp\{-(1/8 + 2/6)\} = 0.632$
3	8	4	1	3/4	1/4	$\exp\{-(1/8 + 2/6 + 1/4)\} = 0.492$
4	9	2	1	1/2	1/2	$\exp\{-(1/8 + 2/6 + 1/4 + 1/2)\} = 0.297$





# **Breslow versus Kaplan–Meier**



- The difference between the Kaplan-Meier and the Breslow estimators is always very small
  - $\triangleright$  as  $n \to \infty$  the two estimates are equivalent
- The Breslow estimator is biased upwards, especially close to zero, but it has lower variance
  - $\triangleright$  if the largest observed time T in the data set is an event, then  $\hat{S}_{KM}(t) = 0$ whereas  $\hat{S}_B(t)$  is positive



R> The Breslow estimator of the survival function is again computed using function survfit(), however now the type argument needs to be specified – for the Stanford data we have

```
Brs <- survfit(Surv(time, status) ~ 1, data = stanford2,
    type = "fleming-harrington")
```

plot(Brs)

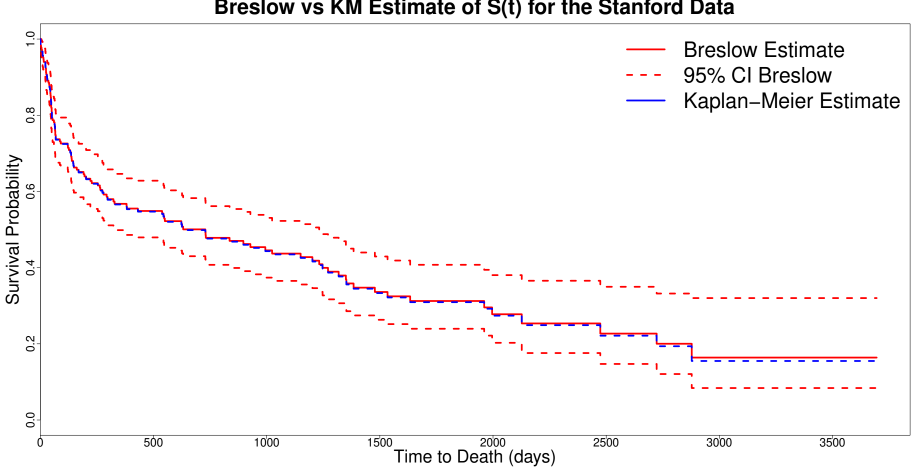


- The variance for the Breslow estimator is based on a similar approach as for the Kaplan-Meier estimator
- The same also holds for the calculation of the 95% confidence intervals
  - $\triangleright$  that is, confidence intervals are computed for  $\log H(t)$  and then back-transformed using the relation

$$\exp\left[-\exp\left\{95\%\ \mathsf{CI}\ \mathsf{for}\ \log H(t)
ight\}
ight]$$

to obtain confidence intervals for  $\boldsymbol{S}(t)$ 





### Breslow vs KM Estimate of S(t) for the Stanford Data

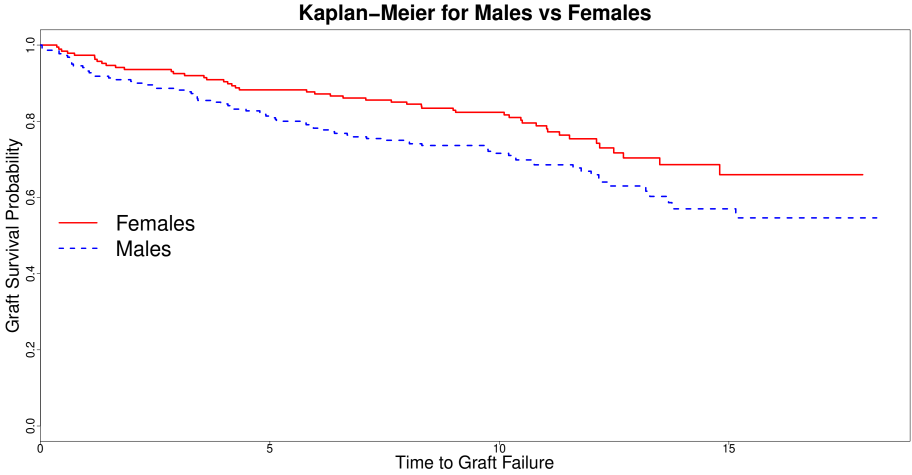


- As before, we observe that the two estimators are indistinguishable
- The Kaplan-Meier is more popular
- However, a lot of theoretical developments in statistics have been based on the Breslow estimator



- We have 2 groups of patients
  - ▷ treatment vs placebo
  - $\triangleright$  females vs males
  - ▷ history of diabetes, Yes vs No
  - $\triangleright \dots$
- Question of Interest: how can we compare these groups with respect to survival
- We can estimate separate survival curves for the 2 groups,

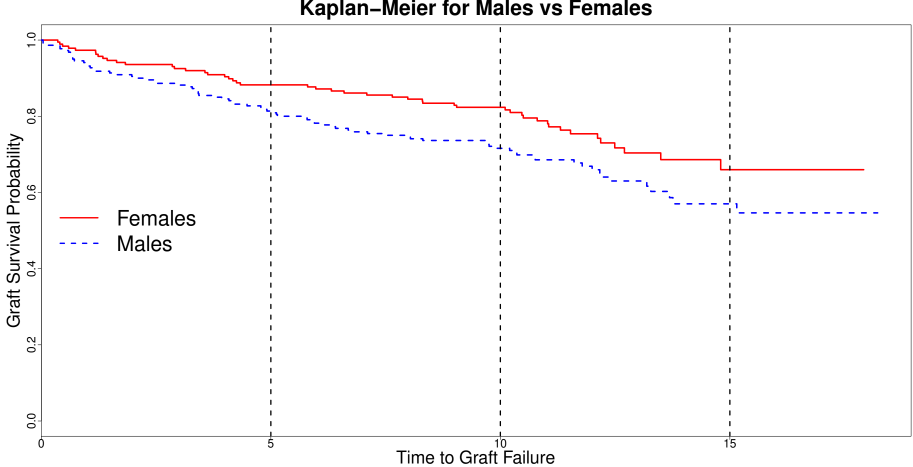






- but how to compare these survival curves?
- We could compare at a specific time point
- At which time point?
  - $\triangleright$  start of follow-up
  - $\triangleright$  end of follow-up
  - ▷ intermediate points
  - $\triangleright \ldots$





# Kaplan–Meier for Males vs Females



- Not very informative because the difference between the survival curves can be greater at some time points than others
- Alternatively, it seems more appropriate to compare the 2 survival curves over the whole follow-up period
- Formally, we are interested in testing the following set of hypotheses

 ${\it H}_0\,$  : the distribution of survival times is the same for the 2 groups

 $H_a$  : it is not the same



- The most famous statistical test to test this hypothesis is the *Mantel-Haenszel Test* (aka Log-Rank Test)
- This is a nonparametric test
  - ▷ no distributional assumption is made for the survival times of the 2 groups
- The philosophy behind it is to construct  $2 \times 2$  contingency tables for each unique event time, and compare observed with expected numbers of events.



• In particular, let  $t_{(i)}$  denote the *i*th ordered event time in the 2 groups combined

	Group 1	Group 2	Total
Event	$d_{1i}$	$d_{2i}$	$d_i$
No Event	$r_{1i} - d_{1i}$	$r_{2i} - d_{2i}$	$r_i - d_i$
At risk	$r_{1i}$	$r_{2i}$	$r_i$

▷  $d_{ji}$  is the number of subjects experiencing the event at time  $t_{(i)}$  in group j▷  $r_{ji}$  is the number of subjects at risk at time  $t_{(i)}$  in group j▷  $d_i$  is the total number of subjects experiencing the event

 $\triangleright r_i$  is the total number of subjects at risk



- In the case of no ties, one of  $d_{1i}$  and  $d_{2i}$  will be 1 and the other 0
- Under the null hypothesis (i.e., the population survival curves are the same in the 2 groups), we can estimate the expected number of subjects experiencing the event at time  $t_{(i)}$

$$\hat{E}_{ji} = \frac{d_i r_{ji}}{r_i}$$

• The variance of  $\hat{E}_{ji}$  can be estimated by

$$\operatorname{var}(\hat{E}_{ji}) = \frac{r_{1i}r_{2i}d_i(r_i - d_i)}{r_i^2(r_i - 1)}$$



- We construct this  $2 \times 2$  contingency table for every observed event time  $t_{(1)}, \ldots, t_{(m)}$
- Then, the log-rank test has the form of a standard  $X^2$  statistic, i.e.,

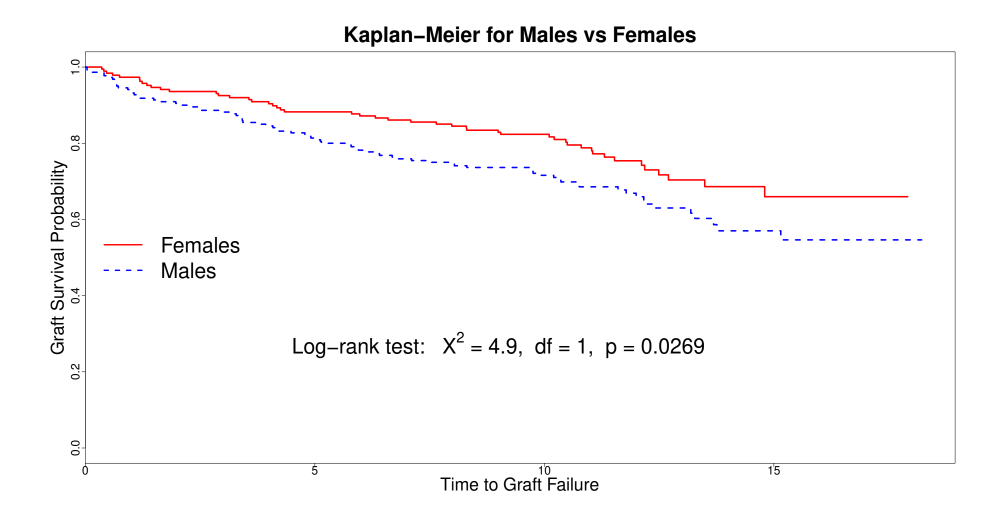
$$X^{2} = \frac{\left(\sum_{i=1}^{m} d_{1i} - \hat{E}_{1i}\right)^{2}}{\sum_{i=1}^{m} \operatorname{var}(\hat{E}_{1i})}$$

• Under the null hypothesis, this statistic is asymptotically distributed as  $\chi^2_1$ 



• Example: for the Renal Graft failure data we are interested in testing whether the survival curve of males is different from the one of females







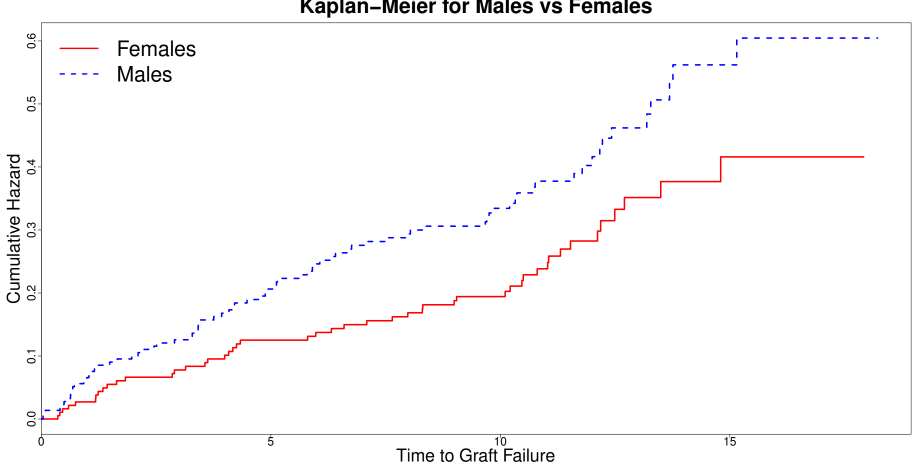
R> The Log-Rank test is computed using function survdiff() – for the Renal data we used

```
survdiff(Surv(Time, failure) ~ gender, data = rgf)
```



- The performance of the Log-Rank Test is compromised when
  - ▷ censoring is informative
  - ▷ the hazard of an event in the one group is not proportional to the hazard of an event in the other group (*proportional hazards assumption*)
- Feature: it places the same weight on all follow-up times





# Kaplan–Meier for Males vs Females



- An alternative test to compare the survival curves, is the modified by *Peto and Peto Gehan-Wilcoxon test*
- Compared to the log-rank test, this test
  - ▷ is more powerful when the hazard functions of the 2 groups are not proportional
  - ▷ puts more emphasis on earlier event times



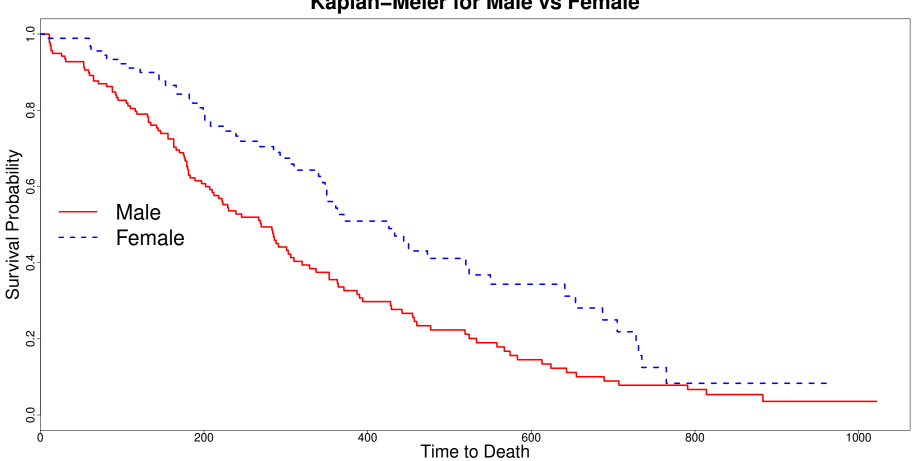
• The Peto-Wilcoxon statistic has a similar form as the  $X^2$  statistic of the log-rank test

$$W = \frac{\left\{\sum_{i=1}^{m} r_i (d_{1i} - \hat{E}_{1i})\right\}^2}{\sum_{i=1}^{m} r_i^2 \operatorname{var}(\hat{E}_{1i})}$$

which weighs the differences between observed and expected deaths in group 1 by the factor  $r_i$  – asymptotically (i.e., in large samples) W has a  $\chi_1^2$  distribution

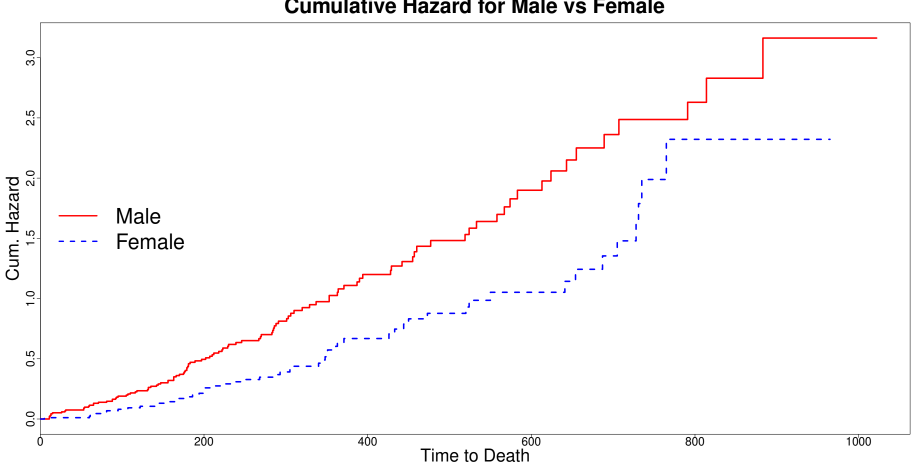
• Example: to illustrate the difference between the two tests, we compare with both the survival curves of males versus females for the Lung data set





### Kaplan–Meier for Male vs Female





### **Cumulative Hazard for Male vs Female**



• Log-rank test

 $\triangleright X^2 = 10.3, df = 1, p = 0.00131$ 

• Peto & Peto modification of the Gehan-Wilcoxon test

 $\triangleright X^2 = 12.7, df = 1, p = 0.00036$ 

- In both cases the result is significant, but the p-value from the Log-rank test is almost 4 times the p-value of the Peto & Peto modified Gehan-Wilcoxon test
  - b the survival curves are much closer at the end of the follow-up than in the beginning



R> The Peto & Peto modified Gehan-Wilcoxon test is again computed using function survdiff(); however, we need to set argument rho to 1 – for the Lung data we used

survdiff(Surv(time, status == 2) ~ sex, data = lung, rho = 1)



- Which of the 2 tests should be preferred?
  - $\triangleright$  if the survival curves cross, then both tests are not optimal
  - check if the proportional hazards assumption is (seriously) violated using the cumulative Hazards plot
  - b the log-rank test will be more powerful if the proportional hazards assumption is valid
  - $\triangleright$  otherwise use the Peto-Wilcoxon test
- However, it would not be fair to decide which test to use based on where the survival curves are further apart, i.e.,
  - > differences in earlier vs later survival times



- We need special notation to take into account censored data
  - $\triangleright T_i$  observed event time
  - $\triangleright \delta_i$  equals 1 if  $T_i$  is a true event, and 0 if it is a censoring time
- Estimators of the survival function
  - ▷ Kaplan-Meier
  - $\triangleright$  Breslow



- Statistical tests to compare survival functions
  - ▷ log-rank test
  - $\triangleright$  Gehan-Wilcoxon test

 $\mathbf{Part}~\mathbf{IV}$ 

## **Regression Models for Time-to-Event Data**



- We have seen how we can compare the survival curves of groups of patients
  - ▷ log-rank test
  - ▷ Peto & Peto modified Gehan-Wilcoxon test
- However, in many cases we may have more complex research questions for example,
  - what is the effect of weight on survival (continuous covariate which we do not want to categorize)
  - b what is the effect of treatment if we control for other variables (e.g., age at baseline, history of other diseases, etc.)



- To handle such type of questions we will use statistical models
- Statistical models are usually developed for one of the following reasons
  - $\triangleright$  effect estimation
  - ▷ hypothesis testing
  - $\triangleright$  prediction
- Different modelling strategies apply depending on the reason for which we develop a statistical model



• In standard statistics we have the *Simple Linear Regression Model*:

$$Y_i = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip} + \varepsilon_i$$

where

 $\triangleright Y_i$  denotes the value of the response variable for the *i*th subject

 $\triangleright X_{i1}, \ldots, X_{ip}$  denote the value of the *p* explanatory variables (aka covariates)  $\triangleright \beta_0, \ldots, \beta_p$  regression coefficients

 $\triangleright \varepsilon_i$  random error terms – usually

$$\varepsilon_i \sim \mathcal{N}(0, \sigma^2)$$



- For survival data we have two complications:
  - ▷ the response variable is Time, which is always positive
  - ▷ censoring
- The solution to the first problem is simple, namely

Use  $\log T_i^*$  instead of  $T_i^*$  as a response variable

• Therefore, we obtain the model

$$\log T_i^* = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip} + \varepsilon_i$$



• This model is known as the:

Accelerated Failure Time Model



- The implications of censoring are twofold
  - estimation is more difficult from a theoretical point of view (but nowadays straightforward with modern computer software)
  - > sensitivity to distributional assumptions for the error terms
    - \* contrary to the linear regression model which is relatively robust against misspecification of the errors' distribution



- Therefore, AFT models are not only based on the normal distribution but other distributions as well
  - $\triangleright$  Student's-*t* distribution heavier tails than the normal
  - ▷ Logistic distribution
  - $\triangleright$  Extreme value distribution this corresponds to the Weibull distribution for  $T_i$ , and it also has as a special case the Exponential distribution

 $\triangleright \dots$ 



- The estimation of the parameters in the AFT model is typically based on the Maximum Likelihood (ML) method
- A (brief) review of ML: we want to find the values of the parameters that are more 'likely' in the light of the data
- As measure of likelihood we use the density function but we treat it as a function of the parameters given the sample at hand

$$L(\theta) = \prod_{i=1}^{n} f(y_i; \theta)$$

where  $y_i$  are the data, and  $\theta$  the parameters



- The most 'likely' parameter values in the light of the data are the values that maximize the likelihood function
- For numerical reasons, it is more convenient to work with the log-likelihood function

$$\ell(\theta) = \sum_{i=1}^{n} \log f(y_i; \theta)$$

• The value of  $\theta$  that maximizes  $L(\theta)$  also maximizes  $\ell(\theta) \Rightarrow$  sufficient to maximize  $\ell(\theta)$ 



- ML for censored data requires special treatment because not all subject provide the same kind of information
- <u>Remember</u>: the observed event time  $T_i$  is the true failure times  $T_i^*$  if subject i had the event or the last time point at which we know this subject was still alive (i.e., in this case  $T_i^* > T_i$ ) (see Section 3.1)
- Therefore, we have two sources of information in the log-likelihood function
  - $\triangleright$  subjects who experience the event  $\Rightarrow$  we use  $f(T_i;\theta)$ , the density function of the assumed distribution for  $T_i$
  - $\triangleright$  subjects who did **not** experience the event  $\Rightarrow$  we use  $\Pr(T_i^* > T_i) = S(T_i; \theta)$ , the survival function of the assumed distribution for  $T_i$



• Thus, the likelihood and log-likelihood functions take the form

$$L(\theta) = \prod_{i=1}^{n} f(T_i; \theta)^{\delta_i} \times S(T_i; \theta)^{1-\delta_i} \Rightarrow$$

$$\ell(\theta) = \sum_{i=1}^{n} \delta_i \log f(T_i; \theta) + (1 - \delta_i) \log S(T_i; \theta)$$

## where

- $\triangleright T_i$  denotes the observed event times, and  $\delta_i$  is the event indicator
- $\triangleright$   $\theta$  denotes all model parameters, i.e., the  $\beta$  's and the variance parameter of the error terms  $\sigma^2$



• An alternative formulation of the log-likelihood (especially useful for proportional hazards models that we will see later) is in terms of the hazard function

$$\ell(\theta) = \sum_{i=1}^{n} \delta_i \log h(T_i; \theta) - H(T_i; \theta)$$

where we use the relations (see Section 2.5)

$$h(t) = \frac{f(t)}{S(t)}$$
$$S(t) = \exp\{-H(t)\}$$



- The Maximum Likelihood Estimates (MLEs) cannot be obtained analytically
- Therefore, to find the MLEs we use optimization algorithms that maximize the log-likelihood with respect to  $\theta$  numerically
  - Newton-Raphson algorithm
  - ▷ quasi-Newton algorithm
- These algorithms are implemented in standard software, such as R and SAS



• The obtained MLEs, which are usually denoted as  $\hat{\theta}$ , are asymptotically (i.e., when the sample size is large) normally distributed

$$\hat{\theta} \sim \mathcal{N}(\theta_0, \{\mathcal{I}(\theta_0)\}^{-1})$$

where

 $\triangleright \theta_0$  denotes the true parameter values

 $\triangleright \mathcal{I}(\theta_0)$  Fisher Information matrix



• The model is

$$\log T_i^* = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip} + \varepsilon_i$$

• One-unit change in variable  $X_1$  corresponds to

$$\log T_i^* = \beta_0 + \beta_1 x + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$$

$$\log T_i^* = \beta_0 + \beta_1(x+1) + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$$

• Therefore,

$$\beta_1 = \log T_i^* - \log T_i^*$$



- In general, one-unit change in variable  $X_j$ , (j = 1, ..., p) corresponds to
  - $\triangleright$  a  $\beta_j$  change in the average  $\log T_i^*$

 $\triangleright$  multiplies average  $T_i^*$  by a factor of  $\exp(\beta_j)$ 

- Now it is more clear where the name Accelerated Failure Time models stems from
  - $\triangleright$  the regression coefficients  $\beta$  directly quantify whether the survival time accelerates or decelerates for a one-unit change in the covariate values



- Example: for the PBC data, we are interested in the treatment effect on survival times after correcting for the effects of Gender and Age at baseline
- To put it in a regression model notation

$$\log T_i^* = \beta_0 + \beta_1 \operatorname{Treat}_i + \beta_2 \operatorname{Sex}_i + \beta_3 \operatorname{Age}_i + \varepsilon_i$$

we are interested in  $\beta_1$ 

• We fit the model assuming normal error terms



• The results are

	est.	(s.e.)
$\beta_0$ – Intercept	4.42	(0.60)
$\beta_1$ – D-penicil	0.21	(0.19)
$\beta_2$ – Female	0.30	(0.28)
$\beta_3$ – Age	-0.05	(0.06)



- The coefficient for the active treatment is  $\beta_1=0.21$
- This means that for patients of the same gender and of the same age at baseline,
  - b the log survival time is 0.21 larger on average for patients receiving D-penicil compared to patients receiving placebo
  - $\triangleright$  the average survival time for the D-penicil patients is  $\exp(0.21)=1.23$  times the average survival time of the placebo patients



**R>** AFT models are fitted using function surveg(). The dist argument specifies the assumed distribution for  $T_i^*$  (not the error terms  $\varepsilon_i$ ) – for the PBC data the following code produces the fit under the Weibull and log-normal distributions:

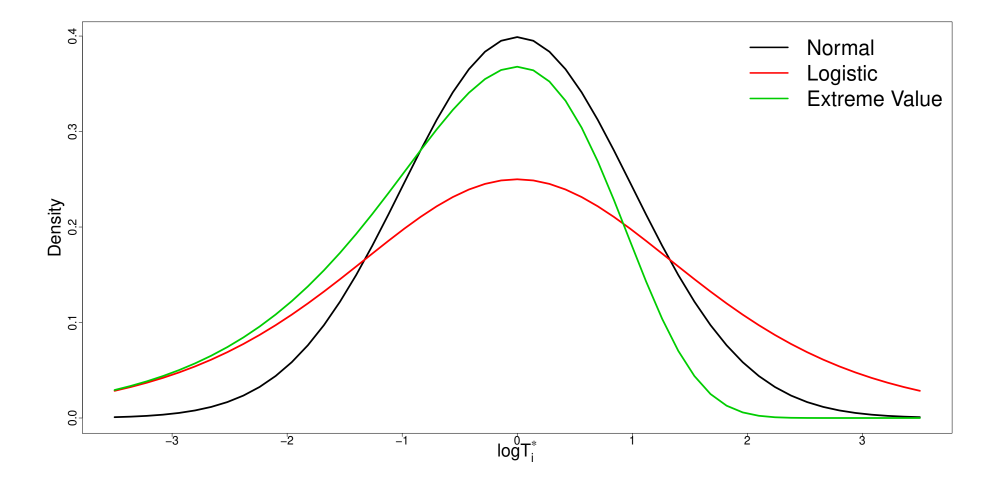
```
dist = "lognormal")
```



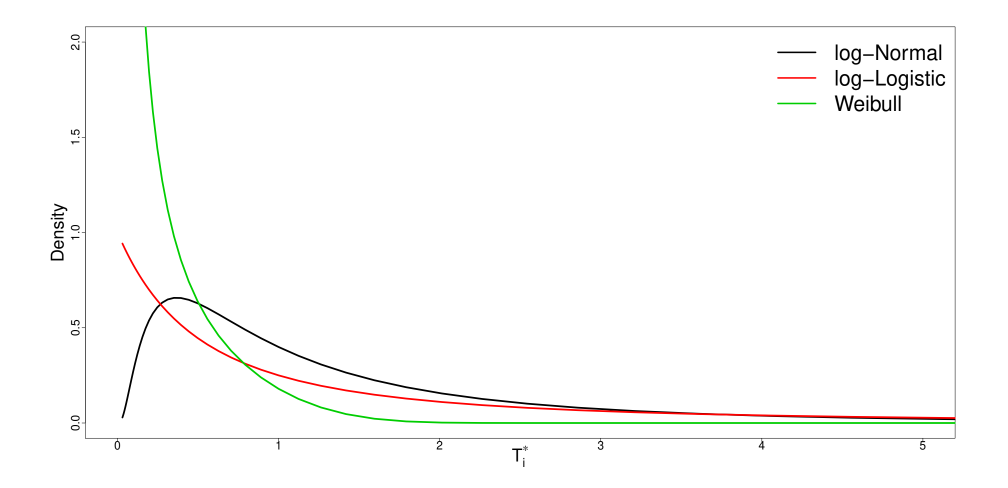
## **R>** The available distributions for $T_i^*$ in R are:

$T_i^*$	$\log T_i^*$	
Weibull / Exponential	Extreme Value	
log-Normal	Normal	
log-Logistic	Logistic	
Normal	—	
Logistic		











• Sensitivity analysis under different error distributions

	Expontl	Weibull	log-Normal	log-Logistic
	est. $(s.e.)$	est. $(s.e.)$	est. $(s.e.)$	est. $(s.e.)$
Intercept	4.33(0.52)	4.14 (0.48)	4.42 (0.60)	4.00(0.55)
D-penicil	0.14(0.17)	0.13(0.16)	$0.21 \ (0.19)$	0.15(0.17)
Female	0.48(0.22)	0.44 (0.20)	0.30(0.28)	0.38(0.24)
Age	-0.04(0.01)	-0.04(0.07)	-0.05(0.06)	-0.04(0.01)



- We observe that the distributional assumptions for the error terms have an effect on the derived parameter estimates and standard errors
- For instance, the gender effect ranges

 $\triangleright$  from 0.30 (s.e. = 0.28) for the log-normal model

- $\triangleright$  to 0.48 (s.e. = 0.22) for the exponential model
- We will come back to this later!



- Regression models often contain several explanatory variables, possibly interacting with each other  $\Rightarrow$  it is difficult to understand how the relationships between these explanatory variables and the response variable interplay
- <u>Solution</u>: communicate the results of a statistical model using Effect Plots
   > a picture is worth a 1000 words
- What are Effect Plots
  - $\triangleright$  choose a specific effect, i.e., a specific combination of the levels of the covariates
    - \* for categorical covariates, specific categories
    - \* for continuous covariates, specific quantiles or range of values
  - ▷ calculate the fitted average, including also the standard error of the fit



b display this fitted average with the associated 95% pointwise Confidence Intervals, preferably using trellis plots

- Example: for the AFT model fitted to the PBC data, we display how the average survival time of female patients changes with respect to the age at baseline, separately for the 2 treatment groups
- In model terms
  - > Treat: both groups are considered, i.e., 'placebo' and 'D-penicil'
  - ▷ Sex: set to 'female'
  - ▷ Age: varies from the minimum observed age to the maximum



• Based on the AFT model on p.128 and the results on p.129, we can calculate the average log failure time as estimated by the model

Placebo, Age = 30  $\Rightarrow \log \hat{T}_i^* = 4.42 + 0.30 - 0.05 \times 30 = 3.27$ D-penicil, Age = 30  $\Rightarrow \log \hat{T}_i^* = 4.42 + 0.21 + 0.30 - 0.05 \times 30 = 3.48$ 

Placebo, Age = 40  $\Rightarrow \log \hat{T}_i^* = 4.42 + 0.30 - 0.05 \times 40 = 2.78$ D-penicil, Age = 40  $\Rightarrow \log \hat{T}_i^* = 4.42 + 0.21 + 0.30 - 0.05 \times 40 = 2.99$ 

Placebo, Age = 50  $\Rightarrow \log \hat{T}_i^* = 4.42 + 0.30 - 0.05 \times 50 = 2.30$ D-penicil, Age = 50  $\Rightarrow \log \hat{T}_i^* = 4.42 + 0.21 + 0.30 - 0.05 \times 50 = 2.51$ 



• And then we can transform back to the original time scale using exp(), i.e.,

Placebo, Age = 30 
$$\Rightarrow \hat{T}_i^* = \exp(4.42 + 0.30 - 0.05 \times 30) = 26.19$$
  
D-penicil, Age = 30  $\Rightarrow \hat{T}_i^* = \exp(4.42 + 0.21 + 0.30 - 0.05 \times 30) = 32.38$ 

Placebo, Age = 40 
$$\Rightarrow \hat{T}_i^* = \exp(4.42 + 0.30 - 0.05 \times 40) = 16.14$$
  
D-penicil, Age = 40  $\Rightarrow \hat{T}_i^* = \exp(4.42 + 0.21 + 0.30 - 0.05 \times 40) = 19.95$ 

Placebo, Age = 50  $\Rightarrow \hat{T}_i^* = \exp(4.42 + 0.30 - 0.05 \times 50) = 9.95$ D-penicil, Age = 50  $\Rightarrow \hat{T}_i^* = \exp(4.42 + 0.21 + 0.30 - 0.05 \times 50) = 12.30$ 



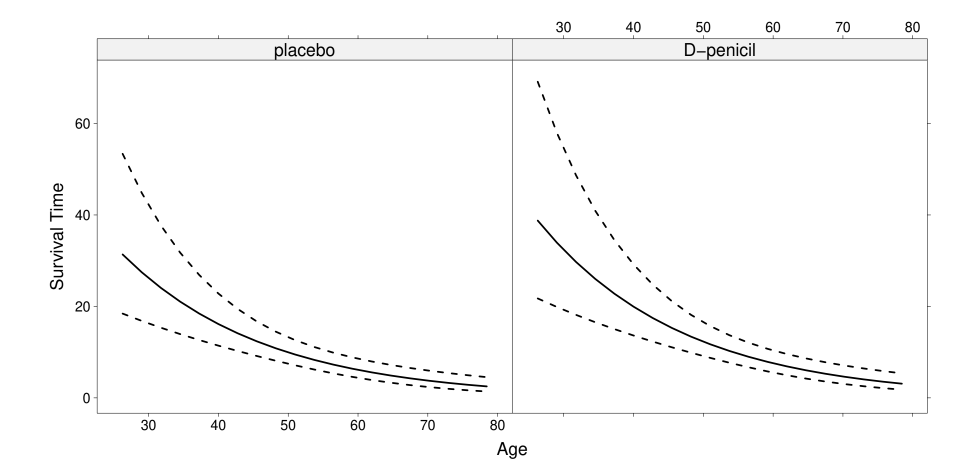
• In addition, for the estimated  $\log \widehat{T}_i^*$  we also obtain a standard error – using this standard error we can calculate 95% pointwise confidence intervals

$$\log \widehat{T}_i^* \mp 1.96 \times s.e.(\log \widehat{T}_i^*)$$

which can also be transformed to the original scale

$$\exp\left\{\log\widehat{T}_i^* \mp 1.96 \times s.e.(\log\widehat{T}_i^*)\right\}$$







**R>** To produce effects, first you fit the desired AFT model using survreg()



R> At the second stage you need to construct a data frame that contains the combination of covariates for which you would like to compute effects

```
ND <- expand.grid(
    age = seq(27, 78, length.out = 20),
    sex = factor("female", levels = c("male", "female")),
    drug = c("placebo", "D-penicil")
)</pre>
```



R> This data frame is then used in the predict(), which provides estimates for the desired effects and their standard errors

```
prs <- predict(fit, ND, se.fit = TRUE, type = "lp")
ND$pred <- prs[[1]]
ND$se <- prs[[2]]
ND$lo <- exp(ND$pred - 1.96 * ND$se)
ND$up <- exp(ND$pred + 1.96 * ND$se)
ND$up <- exp(ND$pred + 1.96 * ND$se)</pre>
```



**R>** Finally we plot the result

```
library("lattice")
xyplot(pred + lo + up ~ age | drug, data = ND, type = "l",
    lty = c(1, 2, 2), col = "black", lwd = 2, xlab = "Age",
    ylab = "Survival Time")
```



• As we have seen, one of the purposes of modelling is to test complex hypothesis of scientific interest – in general, we may be interested in

$$H_0: \theta = \theta_0$$
$$H_a: \theta \neq \theta_0$$

- Since we have fitted the AFT model under maximum likelihood, the standard statistical tests are available
- That is we can choose from
  - > Wald test
    > Likelihood Ratio test
  - $\triangleright$  Score test



• The Wald test is defined as

$$(\hat{\theta}_a - \theta_0)^{\top} \{ \mathsf{var}(\hat{\theta}_a) \}^{-1} (\hat{\theta}_a - \theta_0) \sim \chi_p^2$$

where

 $\triangleright \hat{\theta}_a \text{ the maximum likelihood estimate under the alternative hypothesis}$  $\triangleright v\hat{a}r(\hat{\theta}_a) = \left\{-\partial^2 \ell(\theta)/\partial \theta^\top \partial \theta \Big|_{\theta=\hat{\theta}_a}\right\}^{-1} \text{ denotes the covariance matrix of the MLEs}$  $\triangleright p \text{ denotes the number of parameters being tested}$ 

• The Wald test requires fitting the model only under the alternative hypothesis



• The Score test is defined as

$$\mathcal{S}(\hat{\theta}_0)^\top \hat{\mathrm{var}}(\hat{\theta}_0) \mathcal{S}(\hat{\theta}_0) \sim \chi_p^2$$

where

▷  $\hat{\theta}_0$  the maximum likelihood estimate under the null hypothesis ▷  $S(\theta) = \partial \ell(\theta) / \partial \theta$  denotes the score vector ▷  $v\hat{a}r(\hat{\theta}_0)$  denotes the covariance matrix of the MLEs ▷ v denotes the number of parameters being tested

- $\triangleright p$  denotes the number of parameters being tested
- The Score test requires fitting the model only under the null hypothesis



• The likelihood ratio test (LRT) is defined as

$$-2 \times \{\ell(\hat{\theta}_0) - \ell(\hat{\theta}_a)\} \sim \chi_p^2$$

where

 $\triangleright \ell(\cdot)$  the value of the log-likelihood function  $\triangleright \hat{\theta}_0$  the maximum likelihood estimate under the null hypothesis

- $\triangleright \hat{\theta}_a$  the maximum likelihood estimate under the alternative hypothesis
- $\triangleright p$  denotes the number of parameters being tested
- The LRT requires fitting the model under both the null & alternative hypotheses



- Asymptotically (i.e., for large samples) these three tests are equivalent
- Advice: prefer to use the likelihood ratio test over the other two
- Why:
  - $\triangleright$  it has better theoretical properties
  - $\triangleright$  it makes you carefully think about the hypotheses being tested
- <u>Caveat</u>: if you have missing data in the variable(s) being tested then you have to work with complete cases ⇒ decreased efficiency



- Example: in the AIDS data set we assume that the average failure time may be different
  - ▷ for males and females (main effect of Sex), and
  - ▷ for patients with AZT intolerance (main effect of AZT), but also that
  - b the effect of AZT intolerance on the average failure time is not expected to be the same in males and females (interaction effect)
- We are interested in testing whether AZT intolerance has any effect at all in the average failure time (overall effect of AZT)



• The full model (i.e., the model under the alternative hypothesis)

$$\log T_i^* = \beta_0 + \beta_1 \mathbf{Sex}_i + \beta_2 \mathbf{AZT}_i + \beta_3 \mathbf{Sex} : \mathbf{AZT}_i + \varepsilon_i$$

we fit the model assuming the extreme value distribution for the error terms (Weibull distribution for  $T_i^*$ )

• The reduced model (i.e., the model under the null hypothesis) **must be** a special case of the full model

$$\log T_i^* = \beta_0 + \beta_1 \mathbf{Sex}_i + \varepsilon_i$$



• The models are

$$H_0: \log T_i^* = \beta_0 + \beta_1 \mathbf{Sex}_i + \varepsilon_i$$

$$H_a: \log T_i^* = \beta_0 + \beta_1 \mathbf{Sex}_i + \beta_2 \mathbf{AZT}_i + \beta_3 \mathbf{Sex}: \mathbf{AZT}_i + \varepsilon_i$$

• Therefore, the hypothesis that we are interested in is

$$H_0 : \beta_2 = \beta_3 = 0$$

 $H_a$  : at least one of  $\beta_2$  and  $\beta_3$  is different from 0



• Understanding the models:

▷ reference levels: for Sex is 'female' and for AZT is 'failure'

Female, AZT Fail  $\Rightarrow \log T_i^* = \beta_0$   $+ \varepsilon_i$ Male, AZT Fail  $\Rightarrow \log T_i^* = \beta_0 + \beta_1$   $+ \varepsilon_i$ Female, AZT Intol  $\Rightarrow \log T_i^* = \beta_0$   $+ \beta_2$   $+ \varepsilon_i$ Male, AZT Intol  $\Rightarrow \log T_i^* = \beta_0 + \beta_1 + \beta_2 + \beta_3 + \varepsilon_i$ 



• Understanding the models:

▷ reference level for Sex is 'female'

Female 
$$\Rightarrow \log T_i^* = \beta_0 + \varepsilon_i$$
  
Male  $\Rightarrow \log T_i^* = \beta_0 + \beta_1 + \varepsilon_i$ 



- Likelihood ratio test
  - $\triangleright$  log-likelihood under the reduced model  $\ell(\hat{\theta}_0) = -826.18$
  - $\triangleright$  log-likelihood under the alternative model  $\ell(\hat{\theta}_a) = -813.02$
  - $\triangleright$  parameters being tested p=2
  - $\triangleright LRT = -2 \times \{-826.18 (-813.02)\} = 26.32, df = 2, p-value < 0.001$



R> The LRT and the associated p-value is computed by the anova() method. This function calculates the LRT based on two fitted AFT models – the user is responsible to supply nested models for the LRT to be valid

```
anova(fit.null, fit.alt)
```



- All statistical models are based on assumptions
- In order the derived results from a model to be valid, these assumptions need to hold or at least not to be severely violated
- For the AFT model the assumptions are

 $\triangleright$  linearity (to be discussed later – see Section 4.5.1)

- ▷ additivity (to be discussed later see Section 4.5.2)
- > error terms distribution



• We rewrite the AFT model

$$\log T_i^* = \beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip} + \sigma \varepsilon_i \implies$$

$$\varepsilon_i = \{\log T_i^* - (\beta_0 + \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip})\} / \sigma$$

• Thus, under the AFT model we can check the assumptions for the error terms using the residuals

$$r_{i} = \{\log T_{i} - (\hat{\beta}_{0} + \hat{\beta}_{1}X_{i1} + \hat{\beta}_{2}X_{i2} + \ldots + \hat{\beta}_{p}X_{ip})\}/\hat{\sigma}$$

where

$$\triangleright \hat{\beta}$$
 denotes the estimated regression coefficients

 $\triangleright \, \hat{\sigma}$  the estimated scale parameter of the error terms

Survival Analysis



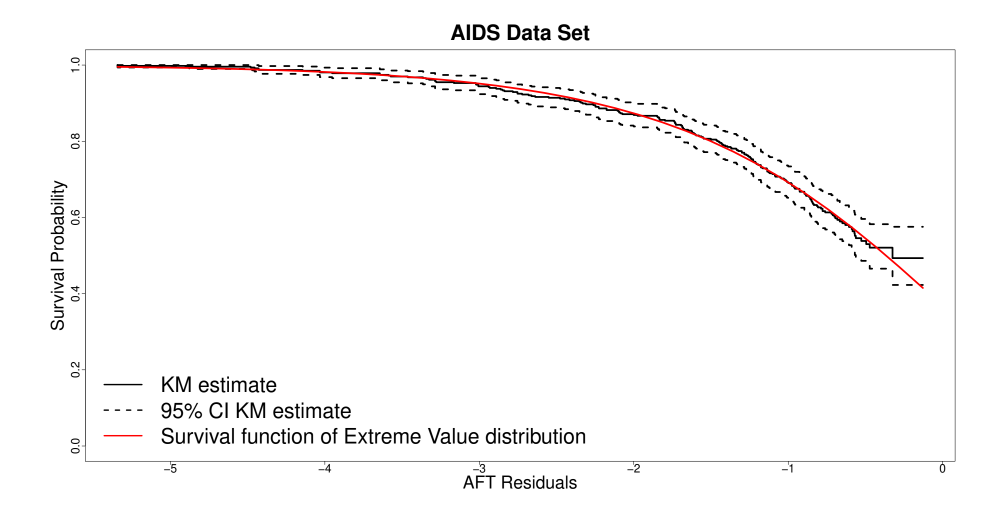
- The error terms are defined with respect to the true failure times  $T_i^*$ . However, the residuals are defined based on the observed failure times  $T_i$
- Thus, when  $T_i$  is censored  $r_i$  will be censored as well
- Censoring must be taken into account when using the residuals to check the AFT model assumptions
- For instance, we can use the Kaplan-Meier estimate of  $r_i$  and compare it with the assumed survival function for the error terms



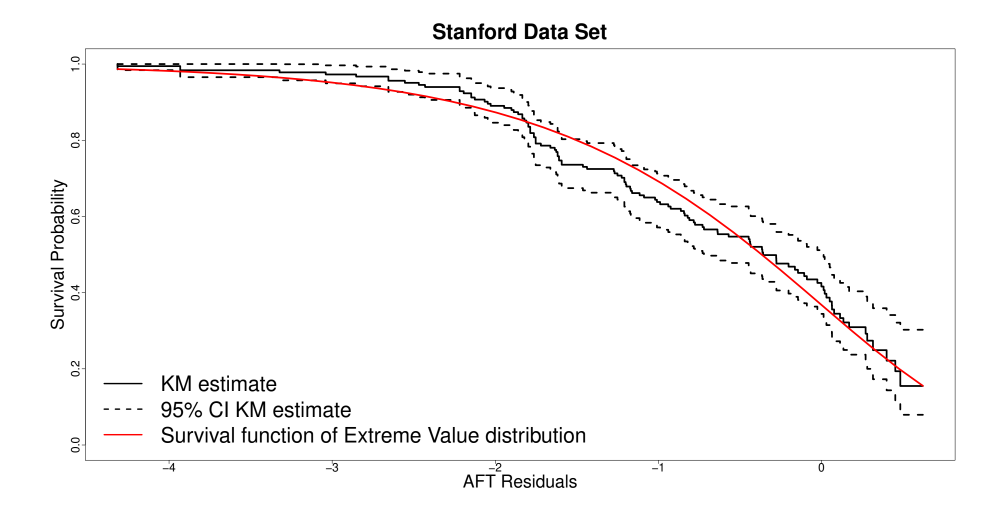
- Example: we display the AFT residuals for the Weibull model fitted to
   the AIDS data set using as only covariate the randomized treatment
   the Stanford data set with only an intercept term
- In particular, we compare
  - $\triangleright$  the Kaplan-Meier estimate of the survival function of  $r_i$ , with
  - ▷ the survival function of the Extreme Value distribution (this is the distribution which we have assumed for the error terms)
- If the model fits the data well, the two survival functions should be close to each other

## 4.2.6 AFT Models – Checking Assumptions (cont'd)











- We observe that the Weibull model provides a much better fit to the AIDS data set than to the Stanford data set
- Therefore, conclusions extracted from the fit of the Weibull model to the Stanford data set should be treated with some caution!



**R>** The AFT residuals can be easily calculated using their definition

# extract fitted values
fits <- fit.weib\$linear.predictors</pre>

```
# compute the AFT residuals
resids <- (log(fit.weib$y[, 1]) - fits) / fit.weib$scale</pre>
```



R> Following we need to compute the Kaplan-Meier estimate of the survival function of these residuals and compare it with the survival function of the distribution of the error terms

resKM <- survfit(Surv(resids, death) ~ drug, data = aids.id)</pre>

```
# plot the KM estimate
plot(resKM, mark.time = FALSE)
```

```
# superimpose the survival function of the assumed
# extreme value distribution
xx <- seq(min(resids), max(resids), length.out = 35)
yy <- exp(- exp(xx))
lines(xx, yy, col = "red", lwd = 2)
```



- So far we have seen that parametric models (i.e., models which make specific distributional assumptions (e.g., log-Normal, Weibull, log-Logistic, etc.) for the event time data) are sensitive, mainly due to censoring
- That is, changing the distribution of the error terms in the AFT model can have a profound effect on parameter estimates and standard errors, and as a result on inference
- This problem has lead to the development of **Semiparametric Regression Models** 
  - ▷ i.e., regression models (parametric component) that make no assumptions for the distribution of the failure times (nonparametric component)



• The most well known semiparametric regression model for survival data is the

Cox Proportional Hazards Model

proposed by sir D.R. Cox in 1972 (Journal of the Royal Statistical Society, Series B)



- Before going into more details about the Cox model,
- Let's review some common ratios in statistics
  - $\triangleright$  odds ratio
  - ▷ risk ratio (aka relative risk)
  - $\triangleright$  hazards ratio



• Odds ratio

$$OR = \frac{p_A/(1-p_A)}{p_B/(1-p_B)} = \frac{p_A(1-p_B)}{p_B(1-p_A)}$$

is the ratio of odds of the event occurring in group A to the odds of it occurring in group B  $% \left( A_{1}^{2}\right) =0$ 

 $\triangleright p_A$  the probability of event in group A  $\triangleright p_B$  the probability of event in group B



• Risk ratio (Relative Risk)

$$RR = \frac{p_A}{p_B}$$

is the ratio of the probability of event in group A to the probability of event in group  ${\sf B}$ 



• Hazards ratio

$$HR = \frac{h_A(t)}{h_B(t)}$$

is the ratio of the hazard of an event at time t for group A to the hazard of an event at t for group B

- The Hazard Ratio is related to the Relative Risk but they are not exactly equivalent
- The difference between the two can be illustrated in the following hypothetical example



- Say that we have 3 patients with increasingly worse prognostic factors
  - ▷ e.g., higher age corresponds to higher risk, and Patient 1 is 20 years old, Patient 2 is 40 years old and Patient 3 is 60 years old
- We assume proportional hazards, that is

 $h_A(t) = 0.5h_B(t)$ 

where  $h_A(t)$  is the hazard of the treated group, and  $h_B(t)$  the hazard of the control

• <u>Remember</u>: the survival and hazard functions are related (see Section 2.5)

$$S(t) = \exp\left(-\int_0^t h(s) \, ds\right)$$



• Therefore we have

$$S_A(t) = \exp\left(-\int_0^t h_A(s) \, ds\right)$$

$$= \exp\left(-\int_0^t 0.5h_B(s) \ ds\right)$$

$$= \left\{ \exp\left(-\int_0^t h_B(s) \ ds\right) \right\}^{0.5}$$

 $= \{S_B(t)\}^{0.5}$ 



Patient	5 Y. Survival		Risk Ratio
	В	А	(A/B)
1	0.90	0.95	0.05/0.10 = 0.50
2	0.50	0.71	0.29/0.50 = 0.58
3	0.20	0.45	0.55/0.80 = 0.69

• The risk ratio depends on the survival rate of the B patients



• The Cox model assumes that the effect of covariates is multiplicative in the hazard scale, i.e.,

$$h_i(t) = h_0(t) \exp(\beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}) \Rightarrow$$

$$\log h_i(t) = \log h_0(t) + \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$$

where

- $\triangleright X_{i1}, \ldots, X_{ip}$  denote p explanatory variables (aka covariates) <u>note</u>: there is no intercept term!
- $> h_i(t)$  denotes the hazard of an event for patient i at time t
- $\triangleright h_0(t)$  denotes the baseline hazard



- The baseline hazard  $h_0(t)$  represents the hazard of an event when all the covariates or all the  $\beta {\rm s}$  are 0
- That is,  $h_0(t)$  represents the instantaneous risk of experiencing the event at time t, without the influence of any covariate
- Therefore
  - $\triangleright$  if a covariate has a beneficial effect, it will decrease this baseline risk
  - $\triangleright$  if, on the other hand, it has a harmful effect, it will increase  $h_0(t)$



- The Cox model makes no assumptions for the baseline hazard function (nonparametric component)
- The model parameters  $\beta_1, \ldots, \beta_p$  are estimated using a Semi-Parametric Maximum Likelihood (SPML) estimation approach
- In particular,  $\beta_1, \ldots, \beta_p$  are obtained as the values that maximize the log partial likelihood function

$$p\ell(\beta) = \sum_{i=1}^{n} \delta_i \Big[ (X_i^{\top}\beta) - \log \Big\{ \sum_{T_j \ge T_i} \exp(X_j^{\top}\beta) \Big\} \Big]$$

where  $X_i^\top \beta = \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$ 



• The obtained Maximum Partial Likelihood Estimates, which are usually denoted as  $\hat{\beta}$ , are asymptotically (i.e., when the number of events is large) normally distributed

$$\hat{\beta} \sim \mathcal{N}(\beta_0, \{\mathcal{I}_p(\beta_0)\}^{-1})$$

where

 $\rhd \beta_0$  denotes the true values of parameters  $\beta$ 

 $\triangleright$  { $\mathcal{I}_p(\beta_0)$ } expected information matrix based on the partial likelihood



- Partial likelihood can be considered as measure of how well the model can order the patients with respect to their survival time.
- **Problem:** some times the hazard ratio for one covariate can be 0 for example, consider the following data set

	Alive	Dead
Treatment	40	0
Control	30	10

Because the treatment group has no deaths its hazard rate is 0



- $\bullet$  If a Cox model is fitted to such data, then the estimated regression coefficient for treatment is  $\infty$
- $\bullet$  Software packages of course cannot detect this problematic case and will usually produce a large in magnitude estimate for  $\beta$
- Therefore, if in the software output you observe a relatively large value for a  $\beta$ , be alarmed and check your data to see why is this happening (e.g., make tables)



• The model is

$$\log h_{i}(t) = \log h_{0}(t) + \beta_{1}X_{i1} + \beta_{2}X_{i2} + \ldots + \beta_{p}X_{ip}$$

• One-unit change in variable  $X_1$ , (j = 1, ..., p) corresponds to

$$\log h_i(t) = \log h_0(t) + \beta_1 x + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$$

$$\log h_i(t) = \log h_0(t) + \beta_1(x+1) + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$$

• Therefore,

$$\beta_1 = \log h_i(t) - \log h_i(t) \implies \exp(\beta_1) = \frac{h_i(t)}{h_i(t)}$$



• In general, one-unit change in variable  $X_j$ , (j = 1, ..., p) corresponds to

 $\triangleright$  a  $\beta_j$  change of  $\log\{h_i(t)/h_0(t)\}$ 

- $\triangleright$  increases  $h_i(t)/h_0(t)$  by a factor of  $\exp(\beta_j)$  (if  $\beta_j < 0$ , then  $\exp(\beta_j) < 1$  and therefore the risk is decreased)
- <u>Note</u>: the interpretation of  $\beta_j$  (j = 1, ..., p) is different from the one in the AFT model



- Example: for the PBC data, we are interested in the treatment effect on the hazard of an event after correcting for the effects of Gender and Age at baseline
- To put in a regression model notation

$$\log h_i(t) = \log h_0(t) + \beta_1 \operatorname{Treat}_i + \beta_2 \operatorname{Sex}_i + \beta_3 \operatorname{Age}_i$$

$$h_i(t) = h_0(t) \exp(\beta_1 \operatorname{Treat}_i + \beta_2 \operatorname{Sex}_i + \beta_3 \operatorname{Age}_i)$$

which means that we are interested in  $\beta_1$ 



• The results are

	est.	$\exp(\text{est.})$	(s.e.)
$\beta_1$ – D-penicil	-0.15	0.86	(0.17)
$\beta_2$ – Female	-0.47	0.62	(0.22)
$\beta_3 - Age$	0.42	1.04	(0.01)



- The estimate for the treatment effect is  $\beta_1=-0.15$
- This means that for patients of the same gender and of the same age at baseline,
  - ▷ the log hazard of the D-penicillamine group is at any fixed point in time 0.15 lower than the log hazard of the placebo group
  - $\triangleright$  the hazard ratio of the D-penicillamine group to the placebo group is  $\exp(-0.15)=0.86$



- $\bullet$  For quantitative covariates  $\beta$  quantifies the effect of one-unit change in the hazard
- For instance, for age  $\beta_3 = 0.42$ . Therefore, for patients of the same sex and who receive the same treatment,
  - > the log hazard is increased by 0.42 for each one year increase in the baseline age
  - $\triangleright$  the hazard of a patient at d+1 years old to the hazard of a patient at d years old is  $\exp(0.42)=1.04$



R> Cox models are fitted using function coxph(). This has the same syntax as survreg() – for instance, for the PBC data the following code fits the Cox model that contains the main effects of 'drug', 'sex' and 'age':

coxph(Surv(years, status2) ~ drug + sex + age, data = pbc2.id)



- As also mentioned in the case of AFT models, Effect Plots are the optimal way for communicating the information in a model
- Example: we fit a Cox model for the Lung data that contains
  - $\triangleright$  main effect of gender
  - ▷ main effect of age at baseline
  - ▷ main effect of Karnofsky performance score
  - $\triangleright$  interaction effect of gender with age



• In model notation we have

$$h_i(t) = h_0(t) \exp(\beta_1 \operatorname{Sex}_i + \beta_2 \operatorname{Age}_i + \beta_3 \operatorname{Karno}_i + \beta_4 \operatorname{Sex}: \operatorname{Age}_i)$$

- We are interested in seeing how the risk of death (at any time point) changes with increasing age at baseline and separately for males and females
- Since in the Cox model we also corrected for the effect of the Karnofsky performance score, we need to specify a value for it
  - $\triangleright$  a reasonable choice is the median Karnofsky score, which in our sample equals 80



• The results are

	est.	$\exp(\text{est.})$	(s.e.)
Female	0.26	1.30	(1.22)
Age	0.02	1.02	(0.01)
Karn Score	-0.01	0.99	(0.01)
Age:Female	-0.01	0.99	(0.02)



• Based on these results, we can calculate the average log hazard rate as estimated by the model ('M' denotes males and 'F' females)

M, Age = 45  $\Rightarrow \log\{h_i(t)/h_0(t)\} = 0.02 \times 45 - 0.01 \times 80 = -0.35$ F, Age = 45  $\Rightarrow \log\{h_i(t)/h_0(t)\} = 0.26 + (0.02 - 0.01) \times 45 - 0.01 \times 80 = -0.63$ 

M, Age = 55  $\Rightarrow \log\{h_i(t)/h_0(t)\} = 0.02 \times 55 - 0.01 \times 80 = -0.18$ F, Age = 55  $\Rightarrow \log\{h_i(t)/h_0(t)\} = 0.26 + (0.02 - 0.01) \times 55 - 0.01 \times 80 = -0.58$ 

M, Age = 65  $\Rightarrow \log\{h_i(t)/h_0(t)\} = 0.02 \times 65 - 0.01 \times 80 = -0.02$ F, Age = 65  $\Rightarrow \log\{h_i(t)/h_0(t)\} = 0.26 + (0.02 - 0.01) \times 65 - 0.01 \times 80 = -0.54$ 



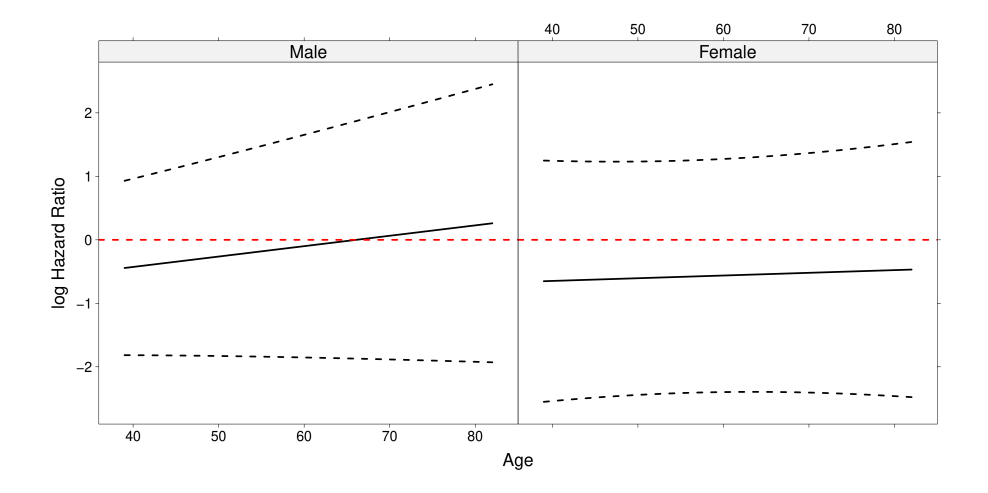
• In addition, for the estimated  $\log \widehat{HR} = \log\{h_i(t)/h_0(t)\}\$  we also obtain a standard error – using this standard error we can calculate 95% pointwise confidence intervals

$$\log \widehat{HR} \mp 1.96 \times s.e.(\log \widehat{HR})$$

which can also be transformed to the hazard scale by

$$\exp\left\{\log\widehat{HR}\mp 1.96\times s.e.(\log\widehat{HR})\right\}$$







- R> The construction of effect plots for Cox models proceeds in the same manner as for AFT models – we start by fitting the desired Cox model
- fit <- coxph(Surv(time, status) ~ age \* sex + ph.karno, data = lung)</pre>



R> Following we construct a data frame that contains the combination of covariates for which we would like to compute effects

```
ND <- expand.grid(
    age = seq(39, 82, length.out = 20),
    sex = c("male", "female"),
    ph.karno = 80
)
```



R> This data frame is then used in the predict(), which provides estimates for the desired effects and their standard errors

```
prs <- predict(fit, newdata = ND, type = "lp", se.fit = TRUE)
ND$pred <- prs[[1]]
ND$se <- prs[[2]]
ND$lo <- ND$pred - 1.96 * ND$se
ND$up <- ND$pred + 1.96 * ND$se</pre>
```



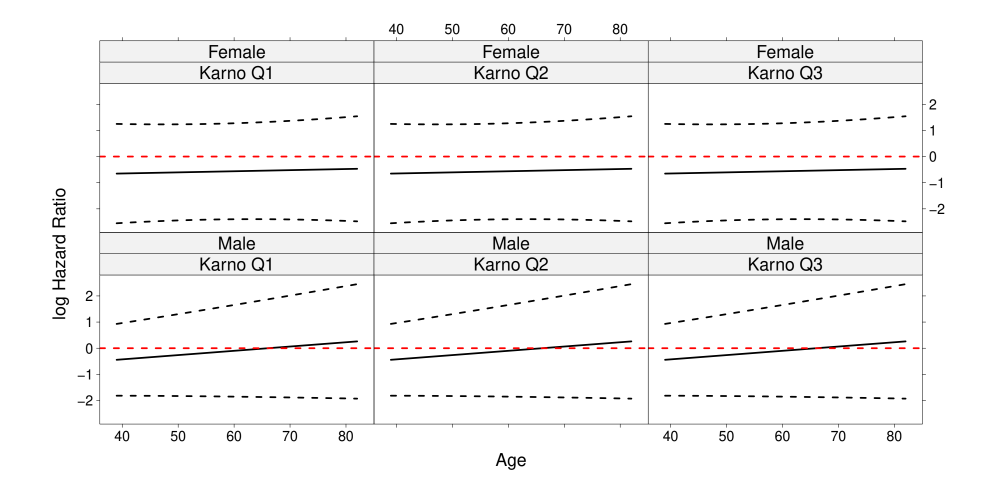
**R>** Finally, we plot the result

```
library("lattice")
xyplot(pred + lo + up ~ age | sex, data = ND,
    type = "l", lty = c(1, 2, 2), lwd = 2, col = "black",
    abline = list(h = 0, lty = 2),
    xlab = "Age", ylab = "log Hazard Ratio")
```



- Note that effect plots can be even more flexible
- For example, we may be interested in how the risk of death changes
  - $\triangleright$  as baseline age increases
  - $\triangleright$  separately for males and females
  - ▷ as the Karnofsky score increases from its 1st quantile value (75), to the median value (80) to the 3rd quantile (90)

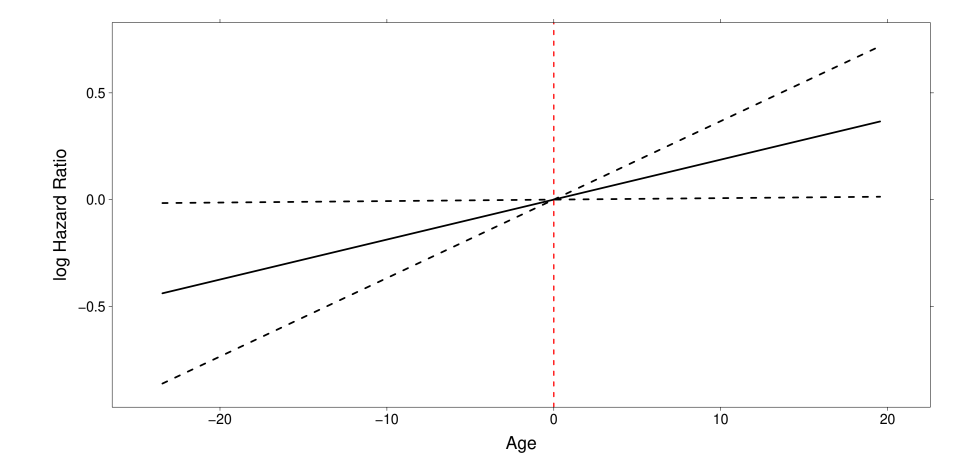






- A peculiarity in the Cox model arises when we compute the log hazard ratio for the reference level of continuous covariate
  - $\triangleright$  the standard error for the predicted by the Cox model value is effectively 0
- Example: for the Lung data we fit a Cox model in which we correct for centered age (i.e., age mean(age)), and we produce the corresponding effect plot







- Hypothesis testing for the Cox model follows the same principles as in AFT models
- Namely for testing the general hypothesis,

$$H_0: \beta = \beta_0$$
$$H_a: \beta \neq \beta_0$$

• <u>Remember</u>:  $\beta$  have the interpretation of log hazard ratio (see Section 4.3.4)



- We can use
  - ▷ Wald test
    ▷ Likelihood Ratio test (LRT)
  - $\triangleright$  Score test
- Important (technical) difference: the Score and LRT statistics are now based on the partial likelihood and not on the full likelihood, as in AFT models
- **Special Case:** when we are interested in testing whether the hazard ratio for a categorical covariate (e.g., treatment) is 1, then

▷ Score test of the Cox model exactly equivalent to the log-rank test



- Asymptotically (i.e., when the number of events is large) the three test statistics are equivalent and converge to the same value (and therefore *p*-value)
- In some cases we may observe differences
- Example: in the PBC data set we are interested to know whether gender has a significant impact on survival



	Statistic	df	<i>p</i> -value
Wald	8.87	1	0.0029
Score (log-rank)	9.18	1	0.0024
LRT	7.73	1	0.0054

- In all cases the result is significant; however, we observe some differences in the value of the statistics and therefore in the level of the *p*-value
- Advice: (as in AFT models) prefer to use the likelihood ratio test over the other two because it has better statistical properties



- Example: in the PBC data set we account for
  - $\triangleright$  main effect of treatment
  - ▷ main effect of age at baseline
  - ▷ interaction effect between treatment and age

and we are interested in the overall treatment effect

• In model terms

$$H_0: h_i(t) = h_0(t) \exp(\beta_2 \operatorname{Age}_i)$$

$$H_a: h_i(t) = h_0(t) \exp \left( eta_1 \texttt{Treat}_i + eta_2 \texttt{Age}_i + eta_3 \texttt{Treat}: \texttt{Age}_i 
ight)$$



• Understanding the models (reference level for Treat is 'placebo'):

 $\triangleright$  Under  $H_0$ 

$$h_i(t) = h_0(t) \exp(\beta_2 \operatorname{Age}_i)$$

 $\triangleright$  Under  $H_a$ 

placebo  $\Rightarrow h_i(t) = h_0(t) \exp(\beta_2 \operatorname{Age}_i)$ D-penicillamine  $\Rightarrow h_i(t) = h_0(t) \exp(\beta_1 + (\beta_2 + \beta_3) \operatorname{Age}_i)$ 



• In this example

$$H_0: \quad \beta_1 = \beta_3 = 0$$
  
 $H_a: \quad \text{at least one is different from } 0$ 

• Likelihood ratio test

 $\triangleright$  log partial-likelihood under the reduced model  $p\ell(\beta_1 = 0, \hat{\beta}_2, \beta_3 = 0) = -712.41$ 

- $\triangleright$  log partial-likelihood under the alternative model  $p\ell(\hat{\beta}_1, \hat{\beta}_2, \hat{\beta}_3) = -711.29$
- $\triangleright$  parameters being tested p=2

$$\triangleright$$
 LRT =  $-2 \times \{-712.41 - (-711.29)\} = 2.24$ ,  $df = 2$ , p-value = 0.326



R> As for AFT models, the LRT and the associated *p*-value for Cox models is computed by the anova() method. Again, this function accepts two arguments, namely the fitted Cox models under the null and alternative hypothesis

fit.null <- coxph(Surv(years, status2) ~ age, data = pbc2.id)
fit.alt <- coxph(Surv(years, status2) ~ drug\*age, data = pbc2.id)</pre>

anova(fit.null, fit.alt)



- Usually, before starting a study we need to know how many subjects we should enroll such that we have a high chance to find a statistical significant difference ⇒
   Sample size determination
- For instance, say we have two groups A and B and we want to detect a hazard ratio of magnitude  $\exp(\beta)$  for a prespecified power level

 $h_A(t) = h_B(t) \exp(\beta)$ 

• <u>Remember</u>: power is the probability that we will find a statistically significant difference between the two groups, **given** that this difference truly exists



• Under the Score test for the Cox model (which is equivalent to the log-rank test) we have the formula

$$\# \text{ events} = rac{(c_{lpha}+z_{power})^2}{p(1-p)\beta^2}$$

where

 $\rhd \beta$  is the log hazard ratio

 $\triangleright p$  is the proportion of patients in the active treatment group

 $\rhd \alpha$  is the Type I error, and  $c_\alpha$  the critical value for the test

- $\rhd z_{\nu}$  is the upper  $\nu$  quantile of the standard normal distribution
- <u>Note</u>: the power depends on the number of events and **not** on the sample size



 $\bullet$  Number of events required under equal allocation in the two groups, with two-sided  $\alpha=0.05$ 

	$\exp(eta)$					
Power	1.10	1.20	1.40	1.60	1.80	2
	1692					
70%	2718	743	218	112	71	51
80%	3456	944	277	142	91	65
90%	4627	1264	371	190	190	87



**R>** The following function computes the number of deaths based on the formula presented in p.213

```
deaths <- function (beta, p, power, alpha = 0.05) {
    # 'beta' the log hazard ratio
    # 'p' the proportion of subjects in the treatment group
    # 'power' the desired level of power
    # 'alpha' the Type I error
    q <- 1 - p
    z.power <- qnorm(power)
    ca <- abs(qnorm(alpha / 2))
    round((ca + z.power)^2 / (p * q * beta^2))
}</pre>
```



R> For instance, to compute the number of events required for a hazard ratio of 1.15, for two groups with the same number of patients on average and power of 90% we use:

deaths(log(1.15), 0.5, 0.9)



- <u>Challenge</u>: how many patients to enroll in order to obtain the required number of events
  - ▷ enrollment period & enrollment rate

 $\triangleright$  study closer

 $\triangleright$  survival probabilities for the control group

- How can we have a rough idea about the log hazard ratio  $\beta$ , in advance
  - ▷ for rare events we have  $\exp(\beta) \approx F_A(t)/F_B(t)$  (F(t) denotes the CDF; check Section 2.2)
  - b that is, a treatment that halves the hazard ratio will approximately halve the proportion of events



- The main motivation to introduce the semiparametric Cox model was to avoid the impact of a possibly wrong assumption for the distribution of the event times
- However, all statistical models make assumptions in the Cox model we make no assumption for the distribution of  $T_i^*$  but we do make other assumptions:
  - $\triangleright$  linearity (to be discussed later see Section 4.5.1)
  - ▷ additivity (to be discussed later see Section 4.5.2)
  - > proportional hazards (PH)
- If one or more of these assumptions are seriously violated, then the results we obtain from the Cox model may not be trustworthy!



- In practice, PH means that the effect of a covariate in the risk of an event is **constant over time**
- Some times the PH assumption may not be reasonable, e.g.,
  - ▷ the new treatment requires a time period to start working ⇒ at the beginning of follow-up the risk of the treatment group is the same as in the control group, however we expect that later the risk of the treatment group will decrease

▷...



- How can we test for PH? We distinguish between
  - categorical covariates with a small number of levels (e.g., treatment, gender, etc.)
     continuous covariates (e.g., age, weight, etc.)
- For categorical covariates we can compare appropriately transformed Kaplan-Meier estimates of the survival functions of the different groups



• In particular, under the PH assumption we have (see pp. 173-175)

$$S_A(t) = \{S_B(t)\}^{\exp(\beta X_i)} \Rightarrow$$

 $\log S_A(t) = \exp(\beta X_i) \log S_B(t) \Rightarrow$ 

 $-\log S_A(t) = \exp(\beta X_i) \{-\log S_B(t)\} \Rightarrow$ 

$$\log\{-\log S_A(t)\} = \beta X_i + \log\{-\log S_B(t)\}$$

• Therefore, if PH holds and we plot the Kaplan-Meier estimates of  $\log\{-\log S_A(t)\}\$ and  $\log\{-\log S_B(t)\}\$ , we expect two approximately parallel lines



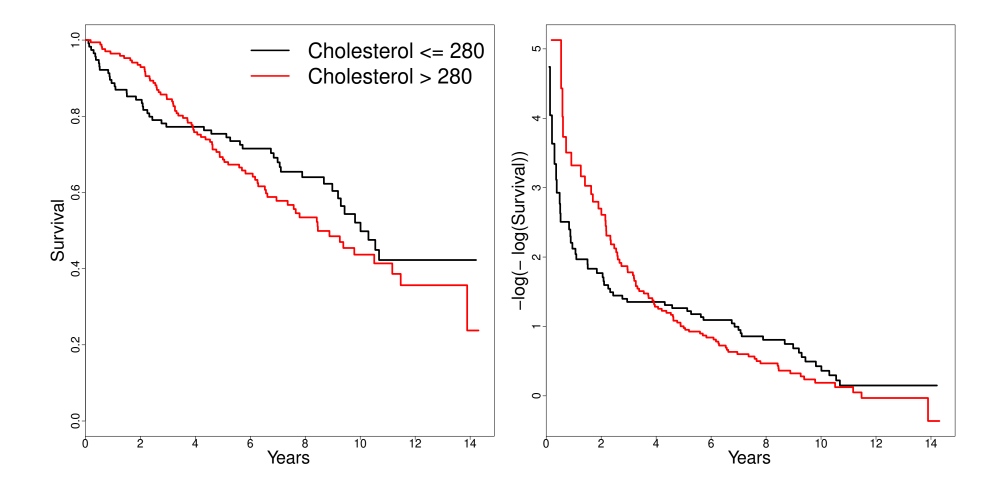
• Example: for the PBC data set we want to test for a impact of cholesterol on the hazard of event – we fit the Cox model

$$h_i(t) = h_0(t) \exp(\beta X_i)$$

where  $X_i = 1$  for patients who had serum cholesterol greater that 280, and  $X_i = 0$  otherwise

- We obtain HR = 1.226, p-value = 0.267
- We test whether the assumption holds graphically by comparing the Kaplan-Meier estimates of  $\log\{-\log S(t)\}$  for the two groups

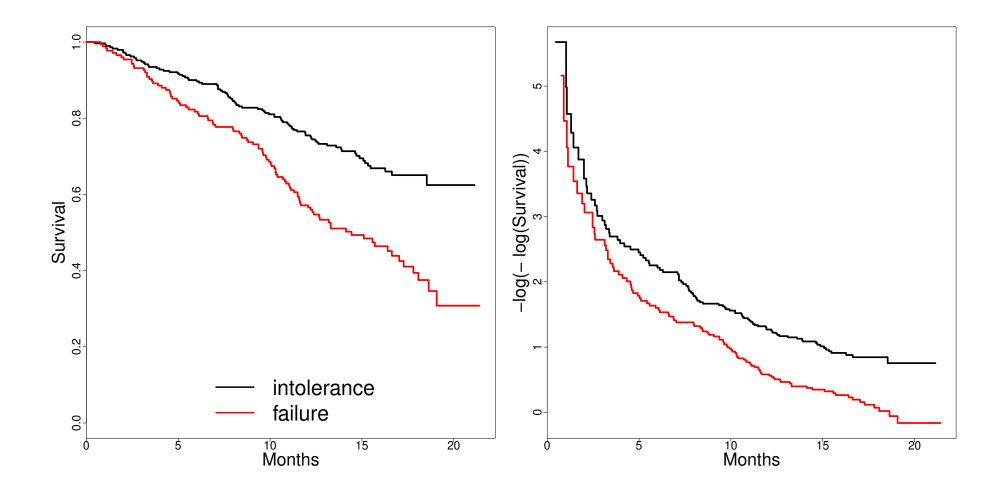






- We observe that the PH assumption seems questionable
- <u>Note</u>: PH may not hold even if the survival curves do not cross over
- Example: for the AIDS data set we want to test for differences in the risk of death between the different AZT groups
  - ▷ AZT intolerance
  - ▷ AZT failure







• The survival curves do not cross but the  $\log\{-\log S(t)\}$  lines are not parallel



**R>** To transform the Kaplan-Meier estimate of S(t) to  $-\log\{-\log S(t)\}$  we use the fun argument of the plot() method



- When the covariate has a few levels, the Kaplan-Meier can be easily used to test the PH assumption
- However, when the covariate has many levels or is continuous, the Kaplan-Meier plot is not useful for discerning either the fact or the pattern of non-proportional hazards
  - b we could collapse some levels of the categorical covariate or discritize the continuous covariate but there is no objective way to do this
- To deal with this we will (hypothetically for now) consider an extension of the Cox model



• Cox model with a *time-dependent coefficient* 

 $h_i(t) = h_0(t) \exp\{X_i\beta(t)\}$ 

- $\bullet$  The impact of covariate X on the hazard varies with time
- Grambsch and Therneau (Biometrika, 1994) have shown that, if  $\hat{\beta}$  is the estimated coefficient from the ordinary (time-independent) Cox model, then

 $\beta(t) \approx \hat{\beta} + E\{s^*(t)\}$ 

where  $s^{\ast}(t)$  is the scaled Schoenfeld residual

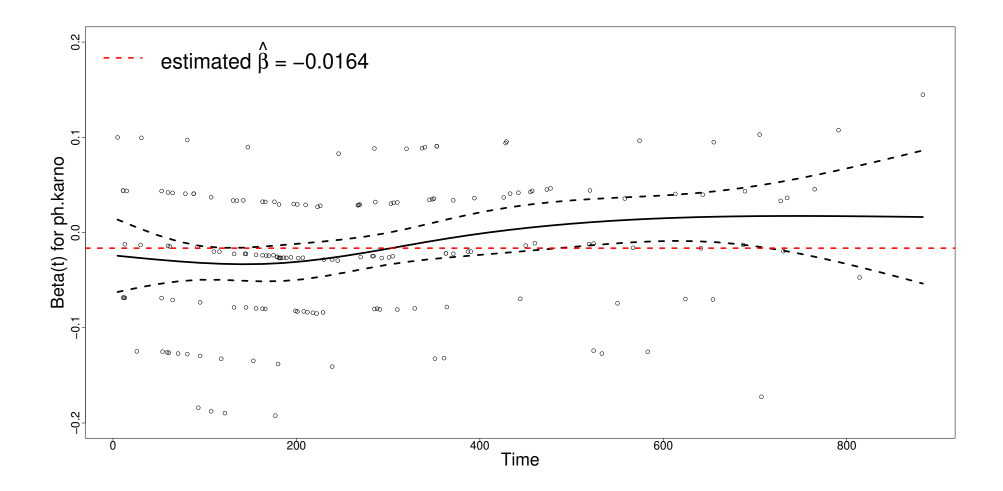


- The formula and rationale behind the scaled Schoenfeld residuals is rather technical > we will not give them here (see Therneau & Grambsch (2000) for more info)
- Plotting scaled Schoenfeld residuals against time or suitable transformation of time, reveals violations of the PH assumption
- An additional advantage of the scaled Schoenfeld residuals is that they can be used to test PH both graphically and via a formal statistical test



- Example: for the Lung data we are interested in the relationship between the Karnofsky performance score and the risk of death
- The ordinary Cox model gives a significant result:
  - ightarrow HR = 0.984, *p*-value = 0.005
- We check the PH assumption using the scaled Schoenfeld residuals
   ▷ we plot the approximated β(t) for Karnofsky score against time







- We have some indications that the effect is not constant in time
- We can formally test for non-proportionality
  - ▷ correlation between scaled Schoenfeld residuals and time

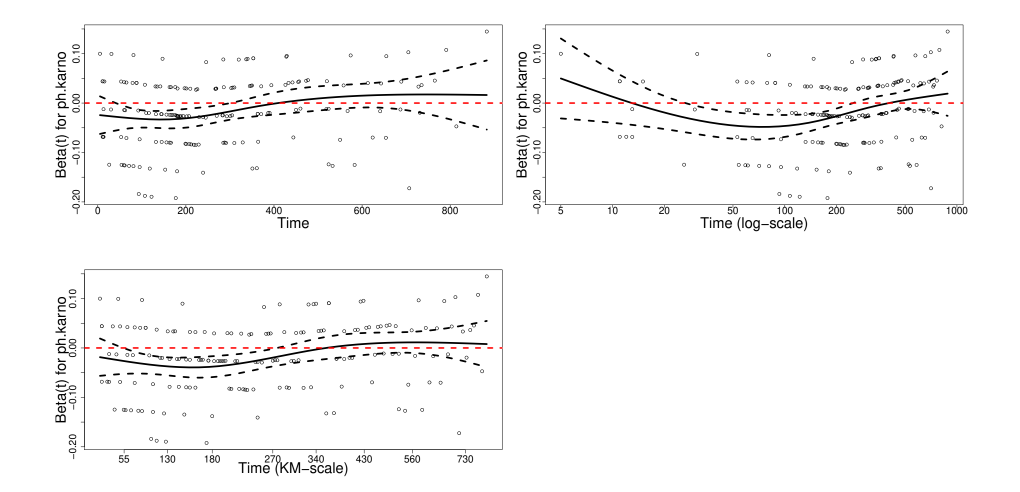
 $\begin{array}{ll} H_0: & \rho=0 \mbox{ (correlation 0 means PH holds)} \\ H_a: & \rho\neq 0 \end{array}$ 

 $\triangleright$  for the Karnofsky score:  $\hat{\rho} = 0.234$ ,  $\chi^2 = 8.08$ , df = 1, p-value = 0.0045



- So far we have used  $\beta(t)$  vs t, but we can also try other time transformations  $\rhd \beta(t)$  vs  $\log(t)$ 
  - $\rhd \beta(t)$  vs Kaplan-Meier transform of t
- The last option is more difficult to interpret in practice,
  - $\triangleright$  (it is calculated as the inverse CDF, with  $CDF(t) = 1 \hat{S}_{KM}(t)$ )
  - $\triangleright$  and it aims to spread the residuals evenly in time, avoiding problems with outliers
- We illustrate for the previous example







- The spread of the residuals is not the same for all time scales
  - $\triangleright$  for t more residuals to the left for  $\log(t)$  more to the right for KM(t) more evenly spaced
- This can have a (substantial) effect on the resulting inferences for non-proportionality

	ho	Statistic	<i>p</i> -value
t	0.234	8.08	0.0045
$\log(t)$	0.139	2.85	0.0913
KM(t)	0.232	7.95	0.0048



- The optimal transformation depends on the specific data set at hand
- General guideline: a transformation that spreads the residuals evenly over time such that the plot and the test are not affected by outliers



R> The test for non-proportional hazards based on the scaled Schoenfeld residuals is calculated by function cox.zph() – argument transform specifies the time scale

```
fit.ph <- coxph(Surv(time, status) ~ ph.karno, data = lung)</pre>
```

```
zph1 <- cox.zph(fit.ph, transform = "identity")
zph2 <- cox.zph(fit.ph, transform = "log")
zph3 <- cox.zph(fit.ph, transform = "km")</pre>
```

zph1 zph2 zph3



**R>** The plot() method is used to produce the plots of  $\beta(t)$  versus the selected time scales

```
par(mfrow = c(2, 2))
```

```
plot(zph1)
abline(h = 0, lty = 2, lwd = 2, col = "red")
plot(zph2)
abline(h = 0, lty = 2, lwd = 2, col = "red")
plot(zph3)
abline(h = 0, lty = 2, lwd = 2, col = "red")
```



- What if the Proportional Hazards assumption is violated?
  - ▷ does it really matter? How serious is the non-proportionality (see p. 232)
- If proportionality is not seriously affected, then what we will obtain is the average Hazard Ratio, averaged over the event times



- If non-proportionality is large, then
  - b incorporate covariates with nonproportional effects as stratification factors into the model (see Section 5.2)
  - $\triangleright$  partition the time axis: the PH assumption may hold over short time periods

\* caveat: less efficient

- ▷ model non-proportionality using time-dependent covariates (see Section 5.3)
- ▷ use nonproportional hazards models, such as AFT models (see Section 4.2)
  - \* <u>caveat</u>: if PH does hold for some covariates, the AFT model will assume that it does not



- The Cox model is a semiparametric proportional hazards model
   we have made no assumptions for the baseline hazard
- However, we also have parametric proportional hazard models
   we assume that baseline hazard has a specific parametric form
- $\bullet$  If we assume that time-to-event  $T^{\ast}_i$  follows a Weibull distribution then we obtain the model



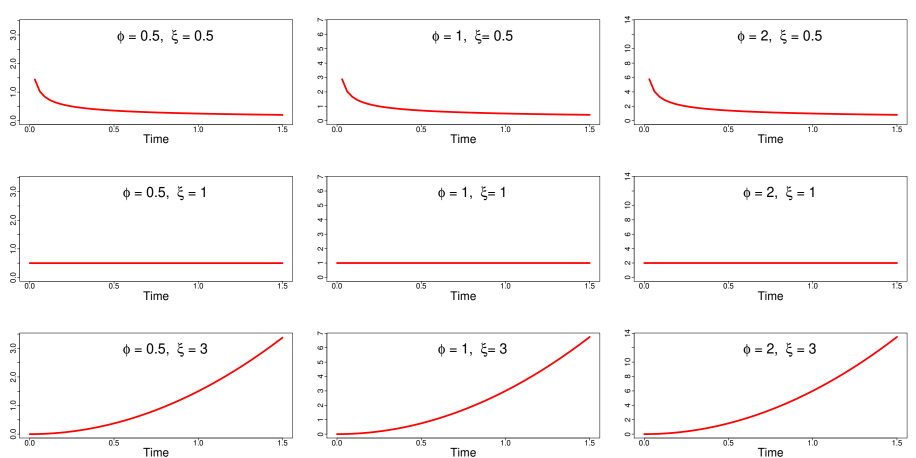
$$h_i(t) = h_0(t) \exp(\beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip})$$

where

$$h_0(t) = \phi \xi t^{\xi - 1}$$

- The interpretation of the  $\beta$  parameters is exactly the same as in the Cox model
- The other parameters control the shape and scale of the distribution of  $T^{\ast}_i$





## Some Weibull Hazard Functions

Survival Analysis



• Example: we compare the Cox and Weibull models for the Stanford data where we control for the effects of age and t5 mismatch score

CoxWeibullvalue (std. err.)value (std. err.)age0.0296 (0.0114)0.0315 (0.0117)t50.1704 (0.1833)0.1738 (0.1823)

- We observe small differences between the two models
  - ▷ for this data set the Weibull assumption seems reasonable this however is not always the case



• The Weibull model (and its special case the Exponential model) is the only model that has both an AFT and PH formulation

$$\log T_i^* = \beta_0 + \beta_1^{aft} X_i + \sigma \varepsilon_i \iff$$

$$h_i(t) = \phi \xi t^{\xi - 1} \exp(\beta_1^{ph} X_i)$$

with the following correspondence

$$\phi = \exp(-\beta_0/\sigma), \qquad \beta_1^{ph} = -\beta_1^{aft}/\sigma, \qquad \xi = 1/\sigma$$



R> Using the previously defined relations we can transform the AFT estimated regression coefficients to PH coefficients for the Weibull model

betasPH <- - coef(fit.weib) / fit.weib\$scale</pre>



**R>** For the calculation of standard errors of the PH estimated coefficients, we need to apply the Delta method

```
# we obtain covariances for the logarithm of the scale parameter
v.betas <- vcov(fit.weib)
transf <- list(~ -x1/exp(x4), ~ -x2/exp(x4), ~ -x3/exp(x4))
library(msm) # load package 'msm'
ses <- deltamethod(transf, c(coef(fit.weib), log(fit.weib$scale)),
v.betas)
```

```
# We compare with the Cox model
fit.cph <- coxph(Surv(time, status) ~ age + t5, data = stanford2)
summary(fit.cph)
cbind("coef" = betasPH, "se(coef)" = ses)
```



- When modelling continuous covariates it is customary to assume that such covariates affect linearly the log hazard ratio (in PH models) or the average log failure time (in AFT models)
- However, this assumption is very restrictive and in many real applications it may not hold
  - ▷ increasing age from 20y to 25y does not increase the risk in the same amount as increasing age from 60y to 65y
- Wrongly assuming linearity may affect the resulting inference for such covariates as well as the predictive ability of the model



- Therefore, it is highly advisable not to restrict a priori the effects of continuous predictors to be linear and let the data tell you the true story
- The easiest way to relax linearity is to assume polynomial effects

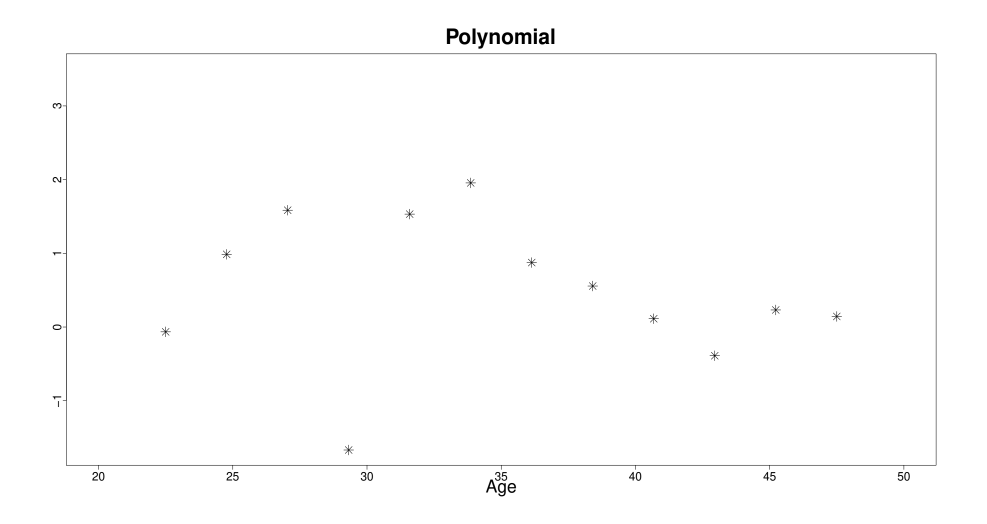
$$\beta_0 + \beta_1 X_i + \beta_2 X_i^2 + \beta_3 X_i^3 + \dots$$

• However, polynomials have some disadvantages

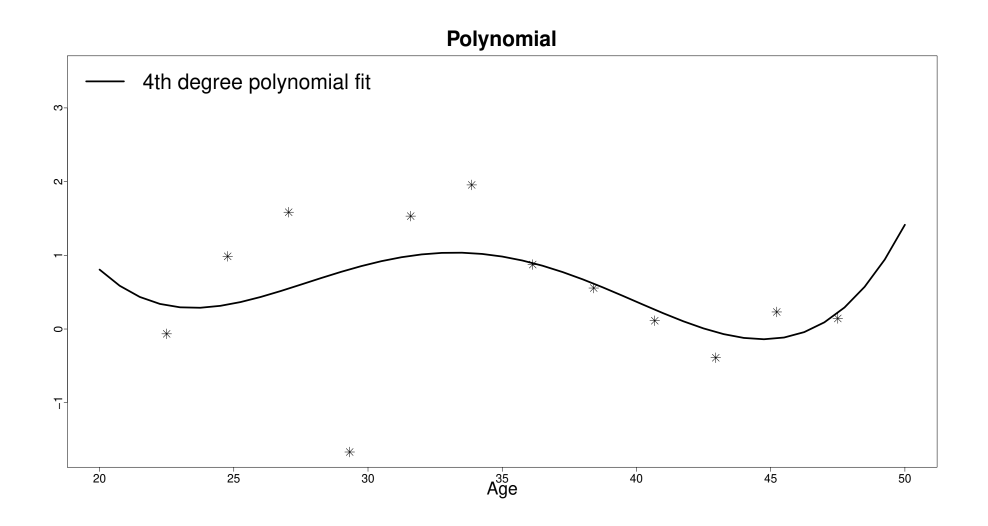
 $\triangleright$  they are not local  $\Rightarrow$  changing one data point will affect the overall fit

> numerically ill-conditioned (however, not too worrisome with modern software)

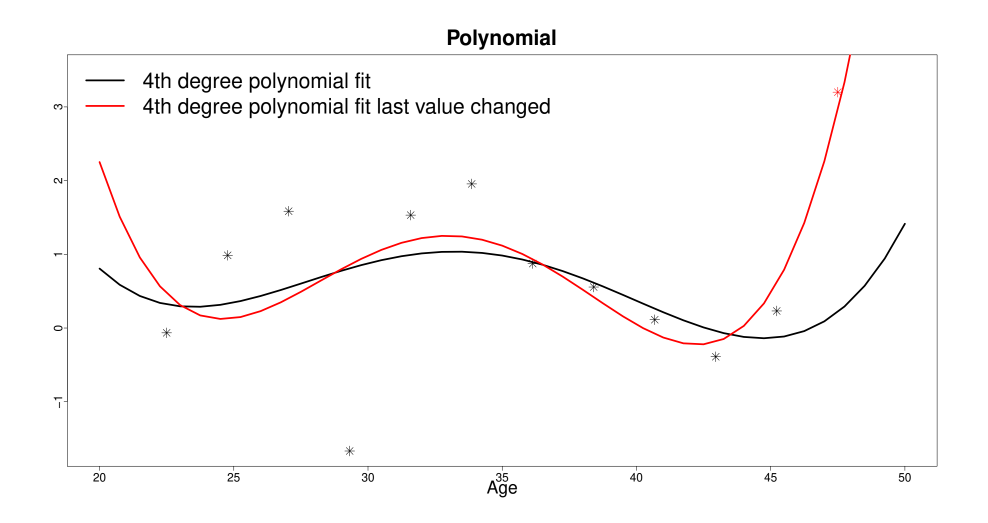








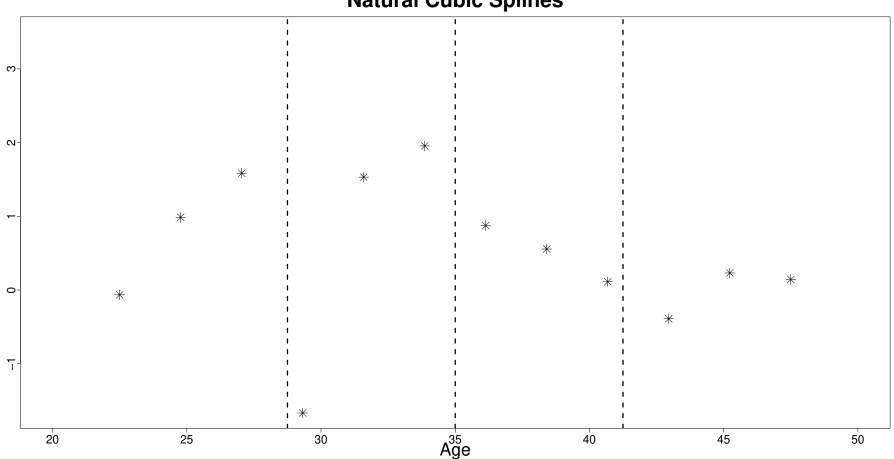






- An alternative approach to relax the linearity assumption of continuous predictors is to use regression splines
- Idea behind regression splines: use polynomials but locally
  - Split the range of values of the continuous predictor into subintervals using a series of knots
  - b within each subinterval assume that the effect of the predictor is nonlinear and can be approximated by a cubic polynomial
  - > put extra smoothness assumptions, i.e., the cubic polynomial fits between neighboring subintervals must be connected



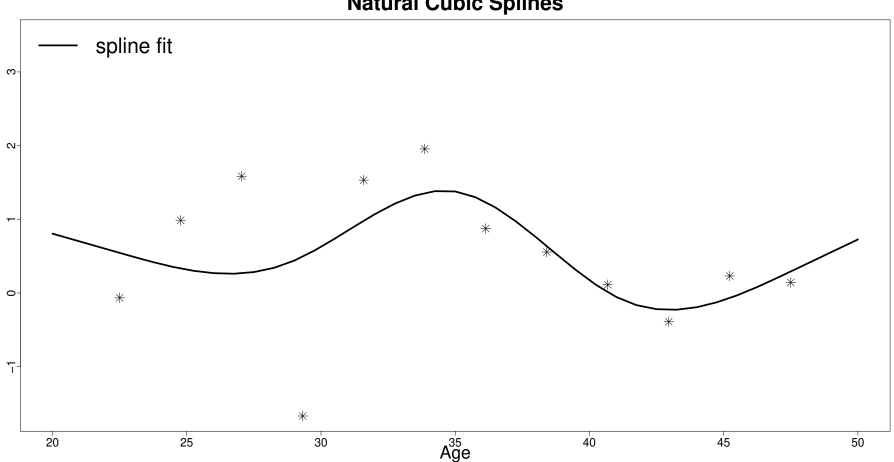


## **Natural Cubic Splines**



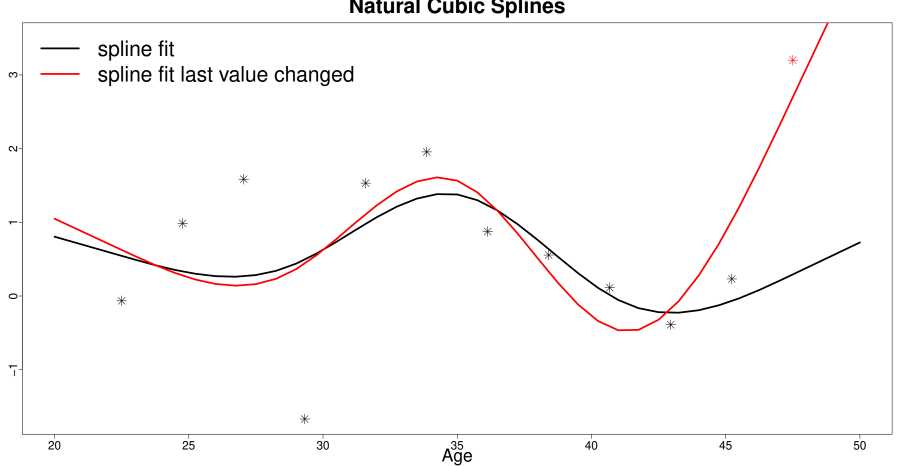
- There are several types of regression splines available
  - b advisable to use natural cubic splines, which assume linearity outside the boundary knots – better statistical properties
- Other approaches (we are not going to discuss them here)
  - $\triangleright$  penalized splines
  - ▷ local regression
  - $\triangleright$  wavelets
  - $\triangleright \dots$





**Natural Cubic Splines** 



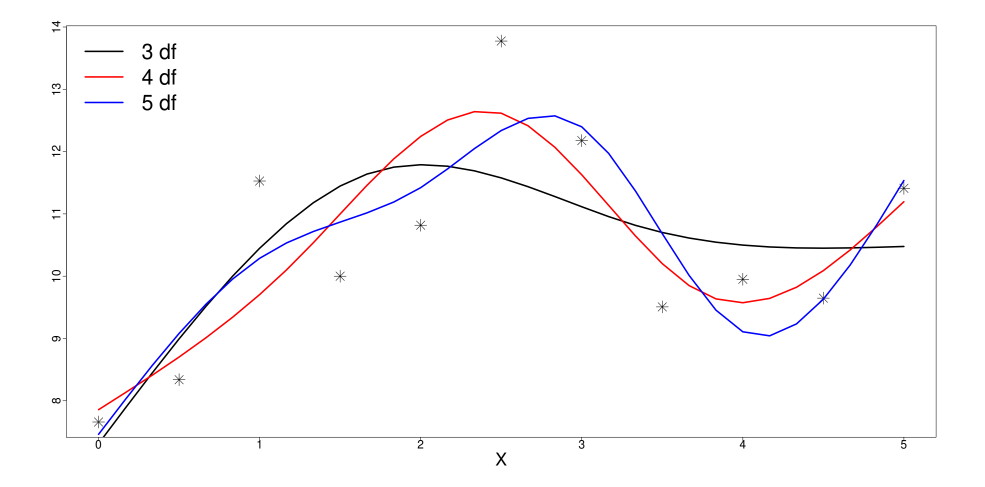


**Natural Cubic Splines** 



- As also in the case of the polynomials, we can tune the degree of nonlinearity by specifying the degrees of freedom for the spline
  - ▷ increasing the degrees of freedom results in more flexible modelling
  - ▷ bias-variance tradeoff

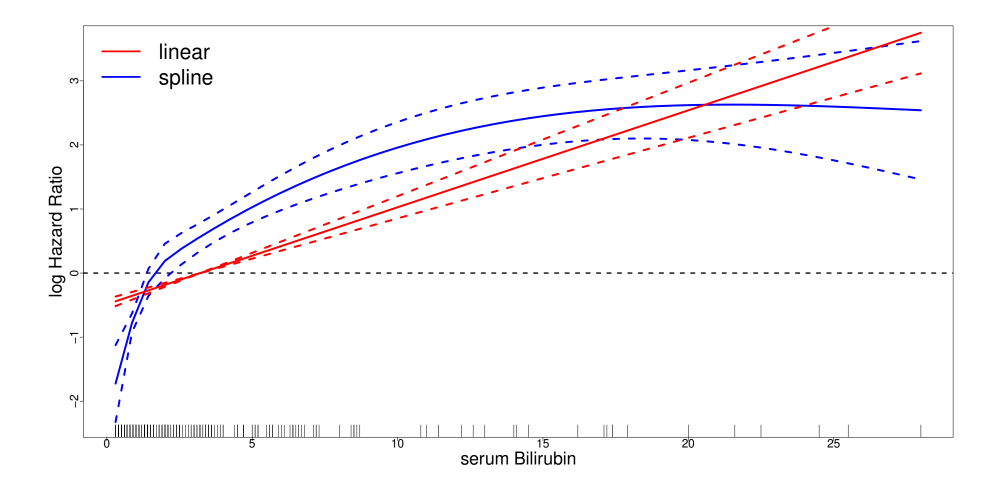






- Example: for the PBC data we are interested in the effect of serum Bilirubin levels in the hazard of death
  - $\triangleright$  we will fit two Cox models,
  - $\triangleright$  one where the effect of serum Bilirubin is restricted to be linear, and
  - $\triangleright$  one where the effect of serum Bilirubin is allowed to be nonlinear
- <u>Note</u>: these are still PH models relaxing linearity it does not mean that we allow the effect to change in time







- R> Polynomial and spline nonlinear effects can be easily specified in both AFT and Cox models, within the formula argument
- R> Orthogonal polynomials are defined using function poly()
- # 3rd degree polynomial for serum Bilirubin coxph(Surv(years, status2) ~ poly(serBilir, 3), data = pbc2.id)

(Orthogonal polynomials provide the same fit as the standard polynomials but with better numerical properties)



R> Natural cubic splines are defined using function ns() of package splines

# natural cubic splines for serum Bilirubin with 3 df
coxph(Surv(years, status2) ~ ns(serBilir, 3), data = pbc2.id)



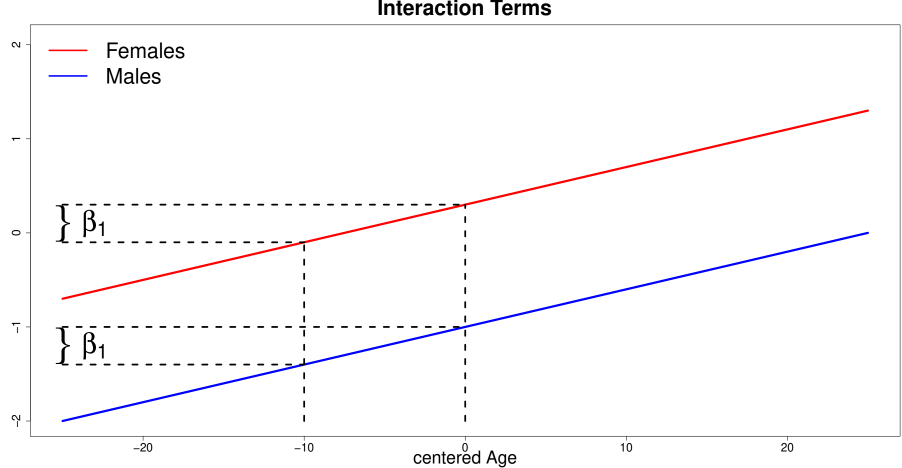
• The additivity assumption can be relaxed by considering meaningful interaction effects

Model I  $h_i(t) = h_0(t) \exp(\beta_1 \operatorname{Age}_i + \beta_2 \operatorname{Sex}_i)$ 

Model II  $h_i(t) = h_0(t) \exp(\beta_1 \operatorname{Age}_i + \beta_2 \operatorname{Sex}_i + \beta_3 \operatorname{Sex} \cdot \operatorname{Age}_i)$ 

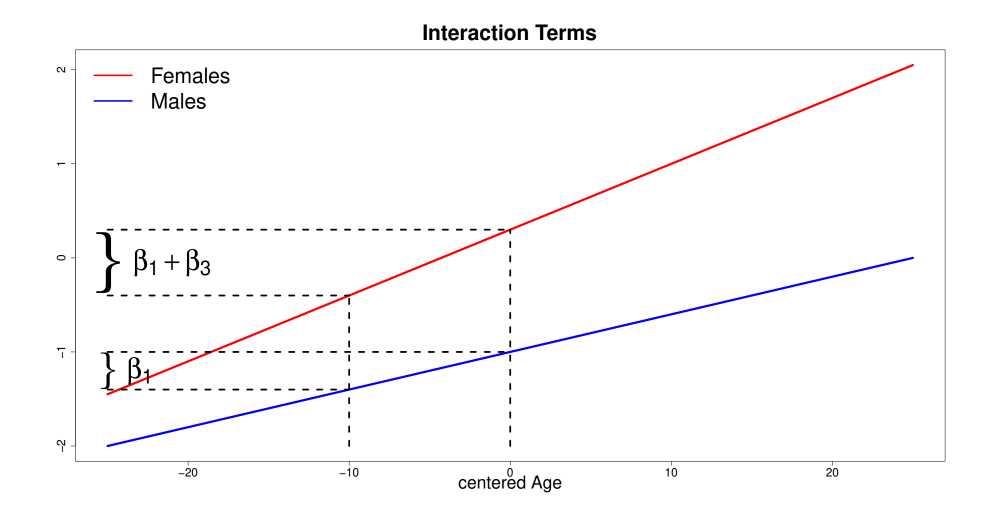
- Model I contains only additive effects whereas Model II contains also the interaction of age with gender
  - $\triangleright$  Model I assumes that the age effect is the same for males and females
  - $\triangleright$  Model II relaxes this assumption





**Interaction Terms** 







- More care is required in extracting conclusions from models with interaction terms
- For instance, in Model II let the reference level for gender being 'Male', then
  - $\triangleright \beta_1$  is the effect of Age for males
  - $\rhd \beta_2$  is the effect of females but for 0 years old!
  - $\triangleright$  (we could center Age, in which case  $\beta_2$  is the effect of females for the mean age)
- However, it is clear that it is difficult to interpret  $\beta_1$  and  $\beta_2$  in isolation



## • For Hypothesis testing:

Null or Alternative Hypothesis	In terms of Parameters
Effect of age is independent of sex	$H_0:\beta_3=0$
Age effects are parallel	
Age and sex are additive	
Age interacts with sex	$H_a:\beta_3\neq 0$
Sex modifies effect of age	
Age and sex are nonadditive	

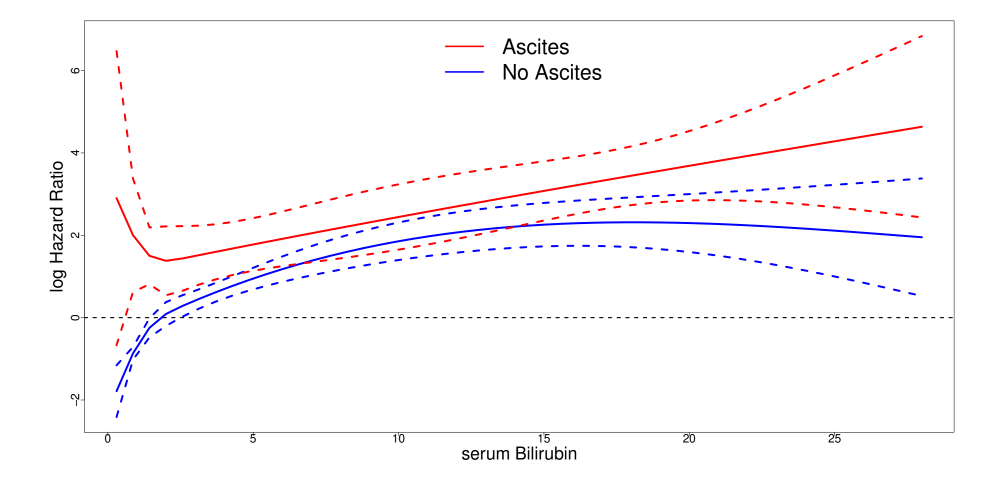


Null or Alternative Hypothesis	In terms of Parameters
Age is not associated with risk of an event	$H_0: \beta_1 = \beta_3 = 0$
Age is associated with risk of an event	$H_a: \beta_1 \neq 0 \text{ or } \beta_3 \neq 0$
Sex is not associated with risk of an event	$H_0: \beta_2 = \beta_3 = 0$
Sex is associated with risk of an event	$H_a: \beta_2 \neq 0 \text{ or } \beta_3 \neq 0$
Neither Age nor Sex are associated	$H_0: \beta_1 = \beta_2 = \beta_3 = 0$
with risk of an event	
Either Age or Sex are associated	$H_a: \beta_1 \neq 0 \text{ or } \beta_2 \neq 0 \text{ or } \beta_3 \neq 0$
with risk of an event	



- Interactions can be easily combined with splines
- Example: for the PBC data set we want to investigate the effects of serum Bilirubin and presence of ascites in the risk of death we relax
  - ▷ the linearity assumption using splines for serum Bilirubin, and
  - b the additivity assumption by taking the interaction of the nonlinear terms with ascites







- Fitting too complex models (i.e., models with too many parameters) may result in *Overfitting*
- Overfitting has two important consequences for the fitted models
  - $\triangleright$  estimated effects have increased variance  $\Rightarrow$  influences confidence intervals and p-values
  - ▷ predicted values from the model do not agree with observed values from future data sets (from the same population) ⇒ the model does not validate well



- To avoid overfitting and depending on the amount of information we have available in the data, there is only a limited number of parameters that we can reliably estimate
- <u>Note</u>: the number of parameters is not, in general, equal to the number of covariates
  - $\triangleright$  categorical covariates with k levels are represented by k-1 dummy variables  $\Rightarrow$  they require k-1 parameters
  - $\triangleright$  allowing for nonlinearity with splines  $\Rightarrow$  number of parameters equals the number of degrees of freedom



• For survival data the number of parameters we can reliably estimate is (rule of thumb)

$$\frac{\# \text{ number of events}}{10} \quad \text{or} \quad \frac{\# \text{ number of events}}{15}$$

• This number can be boosted a bit by censored observations in the case of non-proportional hazards models



- General regression modelling strategies
- How to develop statistical models, with aim to
  - $\triangleright$  effect estimation
  - ▷ hypothesis testing
  - $\triangleright$  prediction
- Most of these guidelines are not only applicable to regression models for time-to-event data but also to other types of models (e.g., linear regression, logistic regression, etc.)



- How to spend your available number of parameters
  - ▷ decide how important each one of the predictors is
    - \* using prior subject-matter expert knowledge
    - \* <u>not</u> using plots of each predictor with the outcome (this will lead to overoptimistic p-values)
  - b if some levels of categorical predictors have very low frequencies consider collapsing these levels with other levels
  - > for important continuous predictors relax the linearity assumption using splines
  - ▷ consider meaningful interactions



▷ fit the model and check assumptions using residuals plots

- $\triangleright$  test the hypothesis of interest
- > present the model using effect plots

> calculate predictions

- Test for complex terms
  - ▷ try to simplify the model using global tests, i.e.,
  - $\triangleright$  test all interaction terms simultaneously; if p>0.15 all interaction terms can be omitted
  - $\triangleright$  apply an analogue global test for the nonlinear terms



- If the model is built with purpose to estimate a specific effect of interest and to test a specific hypothesis, then
  - ▷ do not conserve degrees of freedom for the predictor(s) of interest
  - $\triangleright$  do not try to simplify the model by excluding insignificant predictors in fact p-values obtained by the full model fit will be more accurate



- When we want to measure the effect of more than one predictors we use statistical models for time-to-event data
- We have two main options
  - ▷ Accelerated Failure Time (AFT) models
  - ▷ Proportional Hazards (PH) models
- AFT models measure the effect of predictors on the average failure time > they are the analogue of linear regression for event time data



- PH models measure the effect of predictors on the risk of an event
- AFT models are mainly parametric
  - $\triangleright$  they make specific assumptions for the distribution of the event times
  - ▷ due to censoring they are sensitive to misspecification of this distribution
- This lead to the development of the semiparametric Cox PH model
  - ▷ we make the PH assumption, i.e., the effect of covariate is multiplicative in the hazard scale
  - $\triangleright$  no assumption for the distribution of the event times



- For both modelling frameworks we have seen
  - ▷ how to estimate the model parameters
  - > interpretation of model parameters
  - communicating the results of the model using effect plots
  - ▷ hypothesis testing
  - $\triangleright$  how to check the model assumptions
- In addition for the Cox model
  - $\triangleright$  sample size calculations based on the score test



- Regression modelling strategies
  - > develop models for effect estimation, hypotheses testing or prediction
  - ▷ relax linearity of continuous predictors using splines
  - > decide how many parameters to include depending on the number of events
  - ▷ decide how many degrees of freedom to spend for each covariate in advance

 $\mathbf{Part}~\mathbf{V}$ 

## Extensions of the Cox Model



- The Cox model describes the relationship between the survival curves of the different levels of a covariate using a single number  $\Rightarrow$  the Hazard Ratio
- However, unfortunately, the interpretation of the HR is not that straightforward
   in many cases it is desirable to compare differences between groups in a more easily interpretable scale
- What is often medically relevant is to compare survival probabilities
  - ▷ e.g., how much greater is the probability of surviving at least 5 years in the treatment group than the probability of surviving at least 5 years in the placebo group



- Our aim is to obtain survival probabilities based on the output of the Cox model
- <u>Remember:</u>
  - ▷ in the Cox model we do not need to estimate the baseline hazard we leave it completely unspecified; (see Section 4.3.2)
  - ▷ however, the hazard is directly related to the survival function; (see Section 2.5)
  - b therefore in order to estimate survival probabilities we first need an estimator for the baseline hazard function



• A semiparametric estimator of the survival function based on the output of the Cox model is given by

$$\hat{S}_B(t) = \exp\left\{-\hat{H}_0(t)\exp(\hat{\beta}_1 X_{i1} + \hat{\beta}_2 X_{i2} + \ldots + \hat{\beta}_p X_{ip})\right\}$$

where the baseline cumulative hazard is estimated by

$$\hat{H}_0(t) = \sum_{i=1}^n \frac{I(T_i \le t)\delta_i}{\sum_{j \in \mathcal{R}_i} \exp(\hat{\beta}_1 X_{j1} + \hat{\beta}_2 X_{j2} + \dots + \hat{\beta}_p X_{jp})}$$

with  $\mathcal{R}_i = \{j \text{ for which } T_j \geq T_i\}$  denoting the risk set, i.e., the subjects which did not have the event yet and are not censored



- This is, in fact, an extension of the Breslow estimator we have used for the estimation of the survival function when we had no covariates (see Section 3.3)
- Its variance can be computed using similar arguments as in the no-covariates case
- Example: for the renal graft failure data we are interested in the 5 and 10 year survival rates separately for male and female patients, controlling also for age and weight



• The Cox model has the form

$$h_i(t) = h_0(t) \exp(\beta_1 \mathbf{Sex}_i + \beta_2 \mathbf{Age}_i + \beta_3 \mathbf{Weight}_i)$$

$$\triangleright$$
 HR Sex = 1.01,  $p = 0.130$ 

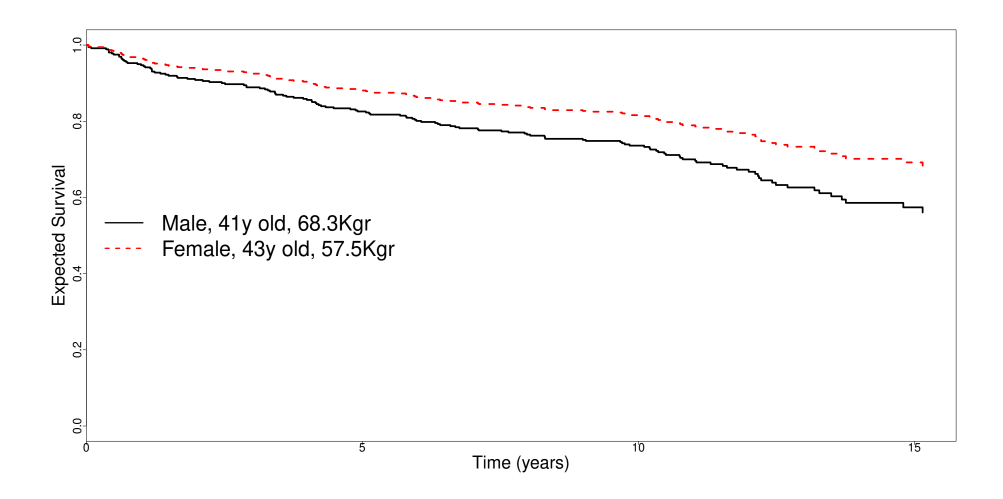
$$\triangleright$$
 HR Age = 0.98,  $p = 0.011$ 

- $\triangleright$  HR Weight = 1.25, p = 0.290
- The following figure depicts estimated survival curves separately for

▷ the median male (41y old, 68.3Kgr)

▷ the median female (43y old, 57.5Kgr)

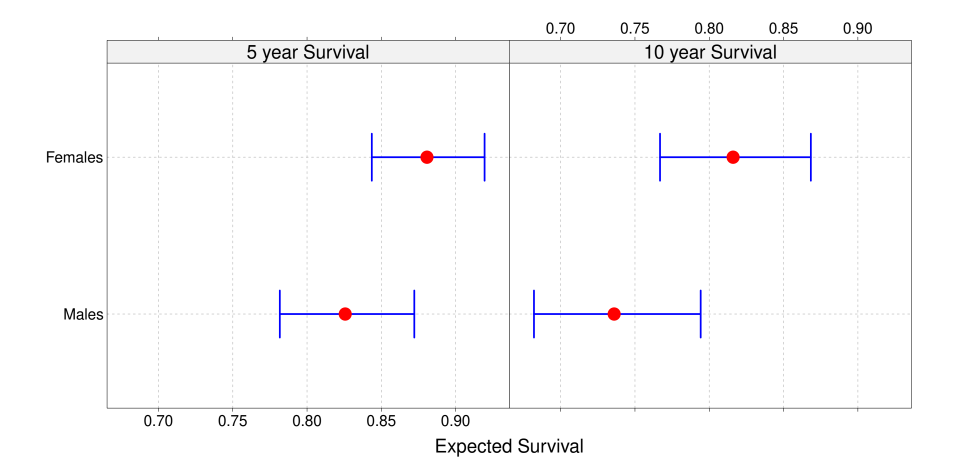




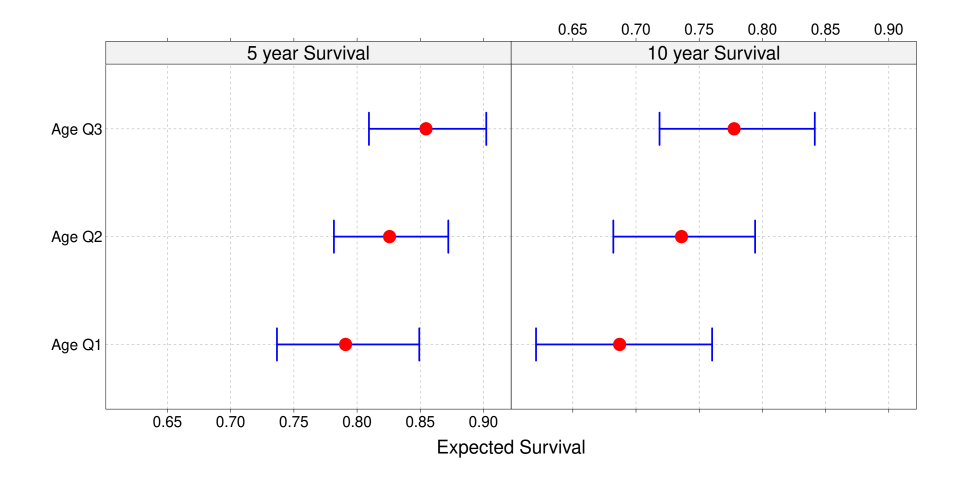


- The next two figures illustrate, estimates of
  - $\triangleright$  5y and 10y survival for the median male, and median female
  - $\triangleright$  5y and 10y survival, for males of 68.3Kgr and
    - \* 30.75 years old (Q1)
    - \* 41 years old (Q2 median)
    - \* 51 years old (Q3)











- A strange result:
  - > the HR for Age is statistically significant different than 1 (HR Age = 0.98, p = 0.011);
  - $\triangleright$  however, the confidence intervals for 5y and 10y survival overlap for increasing Age
  - $\triangleright$  how is this possible?
- <u>Remember</u>: we compare survival curves over the whole follow-up period (see Section 3.4)
   > a significant HR does not mean that the survival curves differ everywhere
  - $\triangleright$  for instance, the 95% confidence intervals at t=0 will almost certainly always overlap



R> Estimates of survival probabilities from Cox models are produced in a similar manner as effect plots – we start with a fitted Cox model and a data frame that contains the specific combinations of covariate values for which we wish to estimate survival



- R> The baseline survival function is estimated by survfit() note that now the first argument is not a formula (as when we used this function compute the Kaplan-Meier estimate) but a fitted Cox model
- R> The data frame that contains the specific combinations of covariate values is supplied in argument newdata – the summary() method can be used to provide survival probabilities estimates for specific follow-up times

```
sfit <- survfit(fit, newdata = ND)</pre>
```

```
sum.sfit <- summary(sfit, times = c(5, 10))</pre>
```



R> The survival estimates with the associated 95% confidence intervals can be extracted using

```
out <- rbind(ND, ND)
out$times <- gl(2, 2, labels = c("5 year", "10 year"))
out$surv <- c(t(sum.sfit$surv))
out$lower <- c(t(sum.sfit$lower))
out$upper <- c(t(sum.sfit$upper))
out</pre>
```



- In many cases it may be unreasonable to assume that the baseline hazard is the same for groups of patients
  - $\triangleright$  multi-center clinical trials  $\Rightarrow$  varying patient populations are likely to have different baseline survival curves

 $\triangleright \dots$ 

- Violation of the proportional hazards assumption
  - $\triangleright$  for categorical covariates  $\Rightarrow$  the log hazard functions of the different levels are not parallel



• The *Stratified Cox Model* allows for multiple strata that divide subject into disjoint groups

$$h_i(t) = h_{0k}(t) \exp(\beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip})$$

where

- $\triangleright$  each of the  $k = 1, \ldots, K$  strata has a distinct baseline hazard  $h_{0k}(t)$
- $\triangleright$  the effect of covariates is the same for all strata (i.e., the coefficients  $\beta$  do not depend on k)



- Example: for the PBC data set we are interested in the main effects of age and logarithm of serum Bilirubin we present three analyses:
  - $\triangleright$  with no stratification
  - ▷ stratify for the different edema categories (3 categories)
  - $\triangleright$  stratify for the different edema and gender categories ( $3 \times 2 = 6$  categories)



	No Strata		Edema		Edema × Sex	
	Value	Std. Err	Value	Std. Err	Value	Std. Err
Age	0.045	0.007	0.045	0.008	0.044	0.009
log ser Bilir	1.091	0.092	0.982	0.097	1.066	0.105

• We observe some differences, especially for the effect of log serum Bilirubin



R> To fit the stratified Cox model we need to specify the stratification variables – in coxph() we use function strata() within the formula argument

R> More than one stratification variables are specified as multiple arguments in strata()

```
coxph(Surv(years, status2) ~ age + log(serBilir) +
    strata(edema, sex), data = pbc2.id)
```



- The standard stratified Cox model assumes that the covariate effects are equal across strata
  - ▷ not always a reasonable assumption
- Interactions between strata and covariates can be easily included
  - ▷ if all of the covariate by strata interaction terms are added, then the results are identical to doing separate fits per stratum
- Example: in the stratified Cox model fitted in the PBC data set we are interested in testing whether the effect of serum Bilirubin is equal among the different edema categories



	Value	Std. Err	<i>p</i> -value
Age	0.043	0.008	< 0.001
log ser Bilir	1.082	0.118	< 0.001
log ser Bilir – edema no diuretics	-0.022	0.241	0.926
log ser Bilir – edema despite diuretics	-0.753	0.259	0.004

 The omnibus *p*-value for the interaction parameters is 0.024 ⇒ the effect of the log serum Bilirubin is not equal among the edema categories

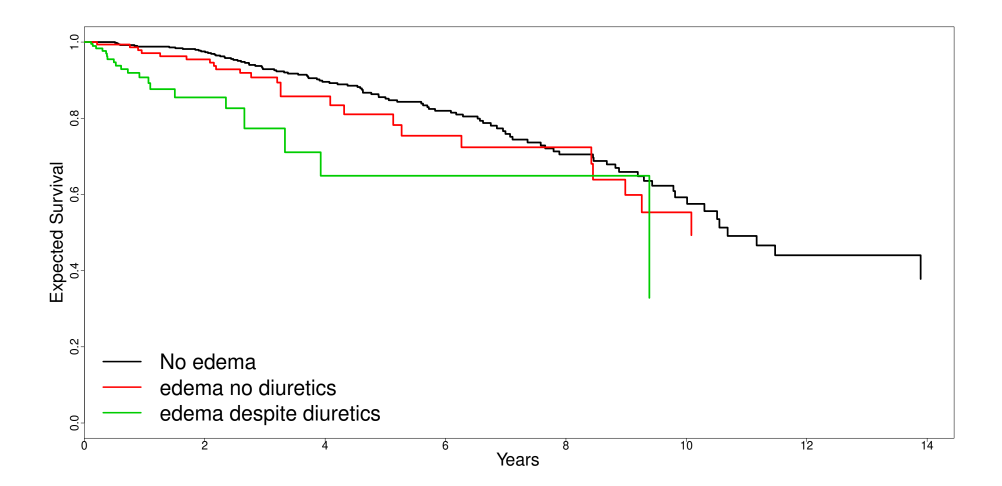


**R>** Covariate by strata interactions are inlcuded in the standard way



- Estimating survival probabilities in stratified Cox models,
  - $\triangleright$  a separate baseline hazard for each stratum, thus
  - $\triangleright$  we obtain a different survival curve for each stratum
- Example: for the stratified Cox model fitted in the PBC data set we are interested to estimate the survival function of 40 year old patient with serum Bilirubin equal to 2







- Advantages of stratification
  - $\triangleright$  it gives the more general adjustment for a confounding variable
  - $\triangleright$  available in standard software
- Disadvantages of stratification
  - $\triangleright$  no direct estimate of the importance of the stratification factor is produced (no p-value)
  - ▷ large number of strata may result in decreased efficiency



- In many cases, we are interested in the effect of covariates whose value changes in time
  - ▷ treatment (e.g., dose) changes with time
  - ▷ time-dependent exposure (e.g., smoking)

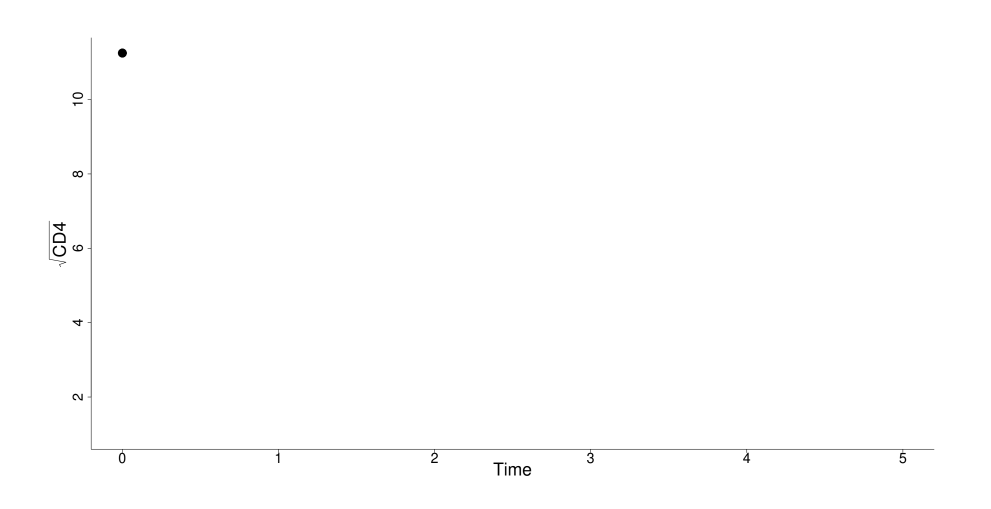
 $\triangleright$  longitudinal measurements on the patient level (e.g., blood values)

 $\triangleright \dots$ 

• Example: in the AIDS data set we have repeated measurements of the CD4 cell count, which is a marker for the condition of the immune system

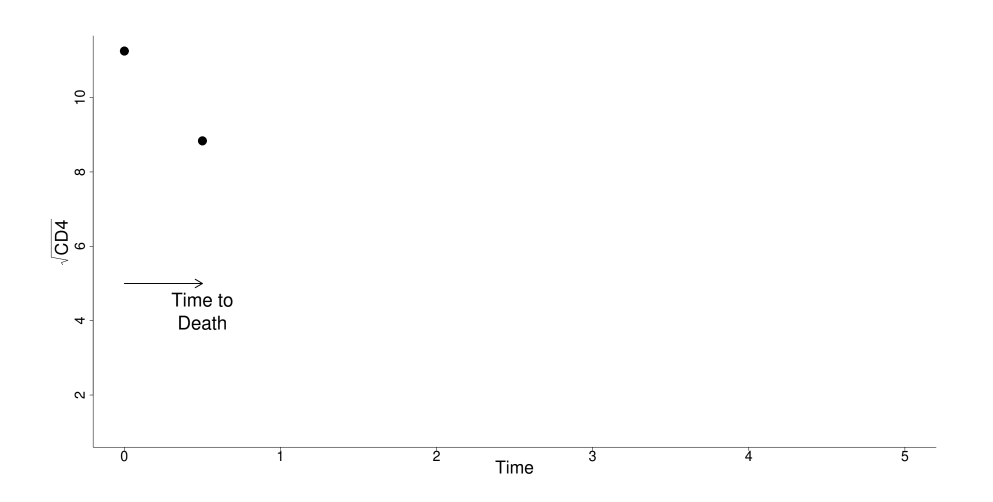
## 5.3 Time-Dependent Covariates (cont'd)



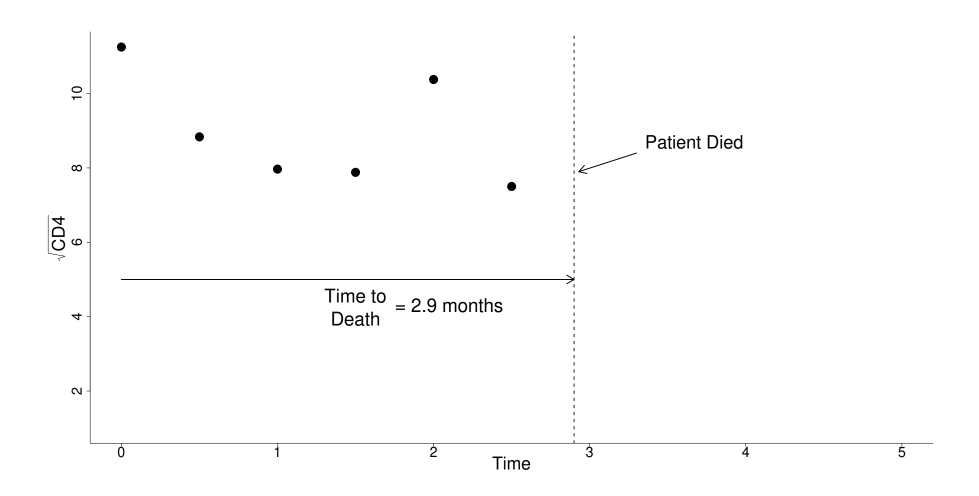


## 5.3 Time-Dependent Covariates (cont'd)











- There are two types of time-dependent covariates (Kalbfleisch & Prentice, *The Stat. Anal. of Failure Time Data*, 2002)
  - $\triangleright$  External (aka exogenous): the value of the covariate at time point t is not affected by the occurrence of an event at time point u, with t > u
  - ▷ Internal (aka endogenous): not External
- This is a difficult concept and we will try to explain it with an example...



- Example: Consider a study on asthma, in particular on the time until an asthma attack for a group of patients
- We have two time-varying covariates: Pollution levels & a biomarker for asthma
- $\bullet$  Say a patient had an asthma attack at a particular time point u
  - $\triangleright$  Pollution levels
    - \* will the pollution levels at time t > u be affected by the fact that the patient had an attack at  $u? \Rightarrow No$
  - $\triangleright$  Biomarker
    - \* will the biomarker level at time t > u be affected by the fact that the patient had an attack at  $u? \Rightarrow Yes$



- It is **very important** to distinguish between these two types of time-dependent covariates, because it determines the type of analysis that it should be followed
- If we treat internal covariates as external, we may produce spurious results



• The Cox model can be extended to handle <u>external</u> time-dependent covariates

$$h_i(t) = h_0(t) \exp\{\beta^\top X_i + \alpha \ m_i(t)\}$$

where

- $\triangleright \beta^{\top} X_i = \beta_1 X_{i1} + \beta_2 X_{i2} + \ldots + \beta_p X_{ip}$  denotes the baseline covariates as we had so far
- $> m_i(t)$  denotes the value of the time-dependent covariate at time t
- $\triangleright \alpha$  quantifies the effect of this covariate at time t to the hazard of an event at the same time point



- When we want to fit the Cox model taking into account the effect of external time-dependent covariates we need to use the counting process formulation
  - > this is a rather technical subject which we will not describe in detail here



• To use this formulation the data must be organized in a long format

Patient	Start	Stop	Event	$m_i(t)$	Age
1	0	135	1	5.5	45
2	0	65	0	2.2	38
2	65	120	0	3.1	38
2	120	155	1	4.1	38
3	0	115	0	2.5	29
3	115	202	0	2.9	29
:	:	:	:	:	



- Example: in the Stanford data
  - ▷ we measure the time until patients die
  - ▷ some patients received a heart transplantation
  - b the dichotomous covariate yes/no transplantation can be considered as an external time-dependent covariate
  - b we are interested in testing whether transplantation has a beneficial effect in survival



• The model that we fit has the form

$$h_i(t) = h_0(t) \exp\{\alpha m_i(t) + \beta A g e_i\}$$

## where

 $\triangleright m_i(t) = 1$  if patient *i* had a transplantation at some time  $u \le t$  $\triangleright m_i(t) = 0$  if patient *i* did not have a transplantation by time *t* 

• We obtain the results

 $ightarrow \exp(\hat{\alpha}) = 0.995, \ p = 0.989$ 

b transplantation does not improve survival



R> Cox models with external time-dependent covariates are fitted using the counting process notation – the data need to be arranged in the long format as in p.311

coxph(Surv(start, stop, event) ~ transplant, data = heart)



- <u>Note</u>: even though time-dependent covariates may produce better insights for the phenomenon under study, some times you may encounter surprising results
- Example: Cavender et al. (1992, J. Am. Coll. Cardiol.) conducted an analysis to test the effect of cigarette smoking on survival of patients who underwent coronary artery surgery
  - b the estimated effect of current cigarette smoking was positive on survival although not significant (i.e., patient who smoked had higher probability of survival)
  - b most of those who had died were smokers but many stopped smoking at the last follow-up before their death



- The approaches these authors used to take this feature into account were
  - instead of the last they used the pre-last assessment as time-dependent covariate (lagged covariates)
  - $\triangleright$  to use the percentage of the follow-up period that the patient smoked
  - > both resulted in a statistically significant increased risk from cigarette smoking
- The choice of the functional form of a time-dependent covariate can have a substantial impact on the derived results

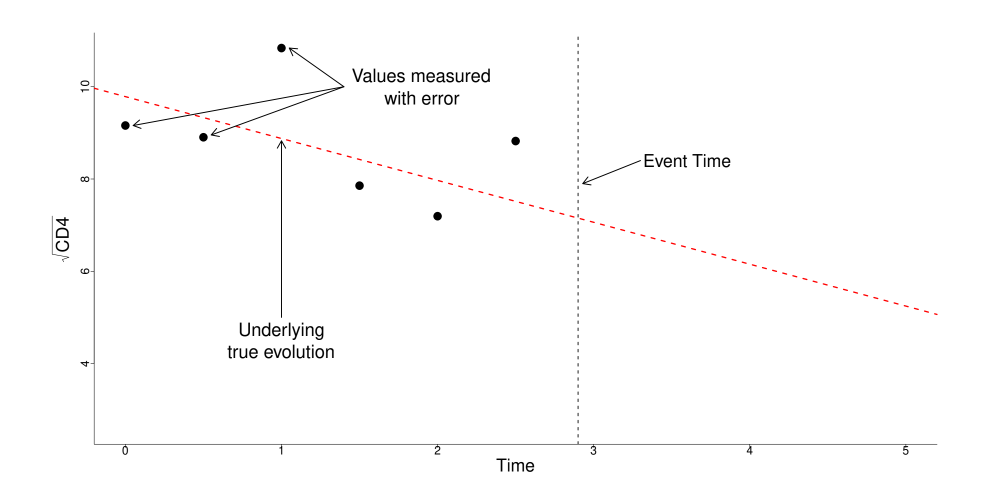


• Let's turn our attention to internal covariates

b what are the challenges with such covariates and why the time-dependent Cox model is inappropriate for them

- They have a stochastic nature
  - $\triangleright$  they contain measurement error
  - we do not have the complete history available(by 'history' we mean the values at any time point)







- To solve this problem, a new class of statistical models has been developed that jointly models the time-to-event outcome with the longitudinal responses
- Intuitively, these models reconstruct the history of the time-dependent covariate and then this estimated history is included as a covariate in the survival model

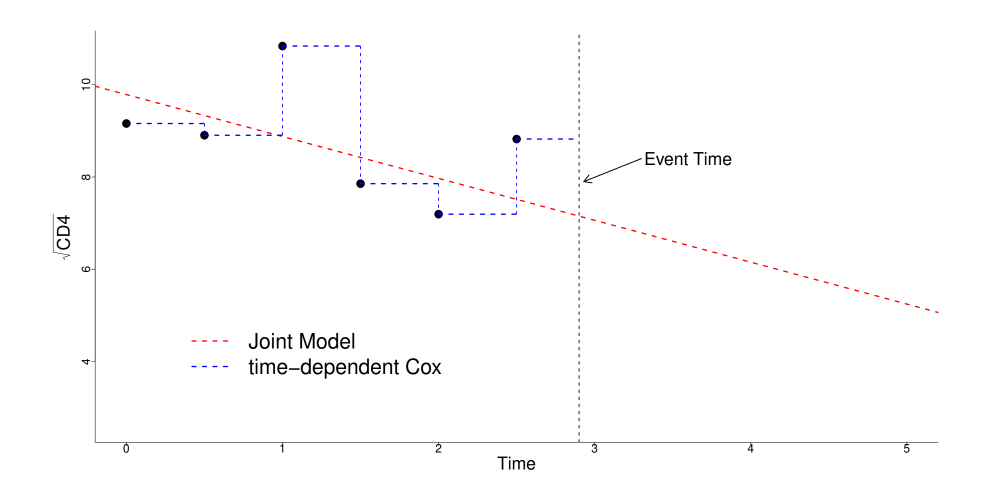


- To illustrate the virtues of joint modelling, we compare with the standard time-dependent Cox model
  - $\triangleright$  i.e., we ignore the measurement error in the CD4 cell count

	Joint Model	Naive TD Cox
	value (std.err)	value (std.err)
Treat	0.35 (0.15)	0.33(0.15)
CD4	-1.10(0.12)	-0.72(0.08)

• Clearly, there is a considerable effect of ignoring the measurement error, especially for the effect of CD4!







- **R>** Such joint models are fitted using functions from **JM** package
- **R>** They require fitting separately a linear mixed effects model and a Cox models
- **R>** A detailed example can be found in the **Survival Analysis in R Companion**



- In many case studies sample units are clustered
  - ▷ patients are clustered within hospitals
  - ▷ children are clustered with schools or families
  - ▷ recurrent asthma attacks (cluster is the patient)

 $\triangleright \dots$ 

- Subjects from the same cluster are expected to be (positively) correlated
  - ▷ if a farmer treats his herd better than an other one, then his cows are expected to live longer



- So far we have assumed that the time-to-event for one subject is completely independent from the time-to-event of another subject
- However, subjects from the same cluster cannot be considered as independent
- Therefore, clustering **must** be taken into account in our analysis



- What are the implications of clustering
  - ▷ it does not affect consistency
  - $\triangleright$  it does affect efficiency
- In practice, this implies that
  - ▷ the estimated effects (i.e., parameter estimates) from the Cox model are valid
  - $\triangleright$  the estimated standard errors are not  $\Rightarrow$  we need to adjust them



• To estimate the standard errors taking into account clustering the **grouped** Jackknife method works satisfactorily

$$\mathbf{v}\hat{\mathbf{a}}\mathbf{r}(\hat{\beta}) = \left(\frac{C-p}{C}\right)\sum_{c=1}^{C}(\hat{\beta}_{-c} - \hat{\beta})(\hat{\beta}_{-c} - \hat{\beta})^{\top}$$

where

- $\triangleright \ C$  is the number of clusters
- $\triangleright p$  the number of parameters
- $\triangleright \hat{\beta}$  the parameter estimates using all the clusters
- $\triangleright \, \hat{\beta}_{-c}$  the parameter estimates excluding cluster c



- Example: the patients in the Lung data set are clustered in institution (we have ignored this feature in the previous analyses of this data set)
  - ▷ we fit a Cox model in which we correct for age, gender, and the Karnofsky performance score



	Value	Std. Err.	Std. Err.	Ratio Std. Err.
		Naive	Jackknife	Naive / Jackknife
Age	0.012	0.0094	0.0062	$1.52~(52\%\uparrow)$
Sex	-0.497	0.1679	0.1252	$1.34~(34\%\uparrow)$
Karno	-0.013	0.0059	0.0086	$0.69~(31\%\downarrow)$

• We observe some considerable differences between the naive and jackknife standard errors



- R> To fit a marginal Cox model for clustered data we need to specify which observations belong to the same cluster – this is achieved using function cluster() within the formula argument of coxph()
- R> By default both the naive and Jackknife (termed 'robust') standard errors are included in the output

coxph(Surv(time, status) ~ age + cluster(inst), data = lung)



• An alternative approach to handle correlated event time data is to use frailty terms

$$h_{ij}(t) = h_0(t) \ \omega_i \ \exp(\beta_1 X_{ij1} + \beta_2 X_{ij2} + \ldots + \beta_p X_{ijp})$$

where

- $\triangleright h_{ij}(t)$  is the hazard for subject j in cluster i
- $\triangleright \omega_i$  is the frailty term, which is an unobserved random variable that
  - \* is shared in all subjects in the same cluster  $\Rightarrow$  induces positive correlation between the subjects of each cluster
  - \* explains heterogeneity between clusters



• Basic Assumption: given the frailty term subjects from the same cluster have independent hazard functions – the log-likelihood can be written as (see p.123)

$$\ell(\theta) = \sum_{i=1}^{n} \log \left[ \int \prod_{i=1}^{n_i} \left\{ h_0(T_{ij}) \omega_i \exp(\beta^\top X_{ij}) \right\}^{\delta_{ij}} \exp\left\{ -H_0(T_{ij}) \omega_i \exp(\beta^\top X_{ij}) \right\} \right]$$
$$f(\omega_i; \sigma) \ d\omega_i$$

where

- $\triangleright$  we assume that  $\omega_i$  follows a distribution (e.g., Gamma, log-Normal, etc.), with density  $f(\omega_i; \sigma)$
- $\rhd \sigma$  is a scale parameter that quantifies correlation within cluster  $\Rightarrow$  heterogeneity between clusters



b maximizing the log-likelihood is a bit more difficult than for the Cox model, but it is available in current software

- Important: the interpretation of parameters is different between
  - ▷ Cox model with grouped jackknife variance estimator (marginal model)
  - ▷ frailty models (conditional model)



• Example: for the Lung data set we fit a marginal Cox model and a frailty Cox model, where in both we correct for the age effect

 Marginal
 Frailty

 Value
 Std. Err.
 Value
 Std. Err.

 Age
 0.0186
 0.0071
 0.0194
 0.0093

  $\sigma$  —
 0.19
 0.0093

 $\rhd \exp(0.0194) = 1.020$  is the hazard ratio for 1 year increase in age for patients in the same institution

 $ightarrow \exp(0.0186) = 1.018$  is the hazard ratio for 1 year increase in age independently from the institution (pooled effect)



R> Cox models with frailty terms are fitted similarly to marginal models – now function frailty() is used within the formula argument of coxph() to identify subjects belonging to the same cluster

coxph(Surv(time, status) ~ age + frailty(inst, df = 4), data = lung)



- Features of frailty models
  - > they directly provide a measure of the correlation between clusters
  - ▷ they can be more efficient
  - > sensitivity in the assumed distribution for the frailty terms
  - > limited availability of model checking tools
- The choice between marginal and frailty models should be dictated by the focus of inference, i.e., conditional versus marginal



- Often in clinical studies we are faced with multiple endpoints, e.g.,
  - $\triangleright$  cancer studies: recurrence of the disease and death
  - $\triangleright$  cancer studies: death from cancer and death due to other causes
  - > cardiovascular studies: nonfatal myocardial infraction and death
- In some cases it makes sense to combine the endpoints
  - b e.g., time until either recurrence of the disease or death (aka Disease Free Survival)
- We end up with a single composite event of interest and therefore, all tools that we have seen so far for the analysis of survival times can be used



- However, in some cases we may be interested in one of the possible events
  - ▷ e.g., we are interested specifically in the time until recurrence of cancer but not in the time until death
  - $\triangleright$  or we are interested separately in the time until recurrence and the time until death
- <u>Problem</u>: occurrence of another event prevents occurrence of the event of main interest
  - $\triangleright$  e.g., for patients who died we can never observe the recurrence of cancer
- The simultaneous consideration of more than one events, which are exclusive with each other is known as a **Competing Risks** problem



- **Crucial distinction:** are the competing risks independent?
  - ▷ if yes, then treating all events from all other causes (except from the one of interest) as censored will produce valid results
  - ▷ if not, then treating all other events as censored will produce **biased** results
- Examples:
  - $\triangleright$  in a study on the survival of cancer patients a patient dies in a car accident  $\Rightarrow$  independent endpoints
  - $\triangleright$  in a study on the survival of cancer patients a patient dies from a heart attack  $\Rightarrow$  probably dependent endpoints
  - $\triangleright$  in a study on patients with osteoporosis were are interested in the time-to-fracture; however, some patients die  $\Rightarrow$  dependent or independent endpoints?



- <u>Caveat:</u> you cannot test for independence
  - ▷ unless unverifiable assumptions are made
- Thus, if you are not sure if the independence assumption is satisfied, you should always do an analysis that allows for dependent competing risks



• Notation for competing risks

 $\triangleright T_1^*, T_2^*, \ldots, T_K^*$  time-to-failure from each one of the K causes

 $\triangleright C$  censoring time (censoring independent of all  $T_k^*$ ,  $k = 1, \ldots, K$ )

what we observe is

 $\triangleright T = \min(T_1^*, T_2^*, \dots, T_K^*, C)$  $\triangleright D = 0, 1, \dots, K \text{ with } 0 \text{ denoting censored time, and } D = k \text{ failure from cause } k$ 



• What is estimable from the available data is the *cause-specific hazard function* 

$$h_k(t) = \lim_{s \to 0} \frac{\Pr(t \le T < t + s, D = k \mid T \ge t)}{s}$$

which is the hazard of failing from cause k at time t

• Anything that can be derived (uniquely) from the cause-specific hazard is also estimable



• <u>Remember</u>: based on the hazard function we can derive the survival function using the relation (see Section 2.5)

$$S_k(t) = \exp\left\{-\int_0^t h_k(u) \ du\right\}$$

which denotes the probability of failure from cause k after time t if

$$h_j(t) = 0$$
, for  $j = 1, \ldots, K$  excluding  $k$ ,

that is, in the **absence** of competing risks



- Therefore,  $S_k(t)$  denotes the survival function for cause k in a hypothetical population where failure from other causes has been eliminated
  - ▷ not very relevant for practical use (even though frequently used in medical papers)
- <u>Note</u>: if we use the Kaplan-Meier estimator to estimate the survival function treating events from other causes as censored, then we actually estimate  $S_k(t)$

 $\triangleright$  bias  $\Rightarrow$  the probability of failing is overestimated

• Need for more easily interpretable functions



• The Overall survival function

$$S(t) = \exp\left\{-\sum_{k=1}^{K}\int_{0}^{t}h_{k}(u) \ du\right\}$$

describes the probability of not having failed from any cause by time  $t \$ 

• The *Cumulative incidence function* of cause k

$$F_k(t) = \int_0^t h_k(u) S(u) \ du$$

describes the probability of failing from cause k before time t



- The cumulative incidence function can be estimated in a similar manner as the Kaplan-Meier estimate of the survival function
- <u>Remember</u>: the Kaplan-Meier estimator was based on the law of total probability (see Section 3.2)
- $\bullet$  We follow the notation of Section 3.2
  - $\triangleright t_1, t_2, \ldots, t_m$  denote the unique event times in the sample at hand
  - $\triangleright d_i$  is the number of events at time  $t_i$ , from all causes
  - $\triangleright d_{ki}$  is the number of events at time  $t_i$ , from cause k
  - $\triangleright r_i$  the number of patients still at risk at time  $t_i$



• The overall survival function S(t) can be estimated using the Kaplan-Meier estimator, without considering the cause of failure (see Section 3.2)

$$\hat{S}(t) = \prod_{i:t_i \le t} \frac{r_i - d_i}{r_i}$$

• The cause-specific hazard function can be estimated by

 $\hat{h}_k(t) =$  Prob failing from cause k at time  $t_i$  given survival up to time  $t_{i-1}$ 

$$= \frac{d_{ki}}{r_i}$$



• Using the definition of the cumulative incidence function

$$F_k(t) = \int_0^t h_k(u) S(u) \ du$$

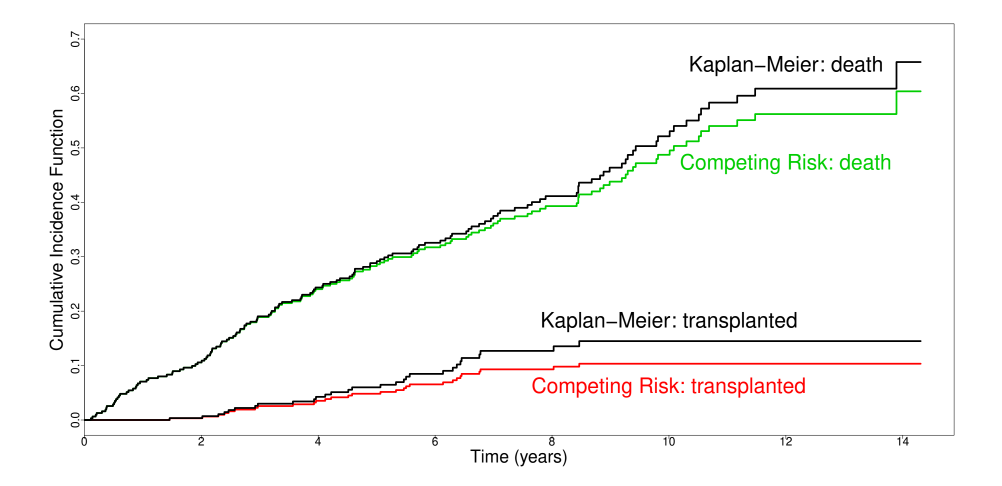
we obtain the estimator

$$\hat{F}_k(t) = \sum_{i:t_i \le t} \hat{h}_k(t_i) \,\hat{S}(t_{i-1})$$



- Example: in the PBC data and when interest is in the time until transplantation, death is a competing risk (i.e., if a patient dies he can never have a transplantation)
- We compare the estimates of the cumulative incidence function for both events (death and transplantation) using
  - ▷ the naive Kaplan-Meier estimator (that treats events from other causes as censored), and
  - ▷ the estimator we derived previously







• As expected, we observe that the naive Kaplan-Meier overestimates the cumulative incidence function



- R> The competing risks estimate of the cumulative incidence function is produced by the survfit() function
- R> Two differences: (i) the event indicator status3 is 1 whenever either of the two events occurred; (ii) argument etype is used to distinguish between the events of interest

survfit(Surv(years, status3) ~ 1, data = pbc2.id, etype = status)



- Inclusion of covariates: two different approaches
  - ▷ Cox model for the cause-specific hazards
    - \* straightforward to implement in standard software (R, SAS)
    - \* more difficult to get estimates of the cumulative incidence function
    - \* proportional hazards assumption for the cause-specific hazards
  - ▷ Fine & Gray model (JASA, 1999)
    - \* hazard ratios interpretable on the cumulative incidence scale
    - \* only available in R (package cmprsk)



• We will only illustrate the Cox model for the cause-specific hazards

$$h_{ki}(t) = h_{k0}(t) \exp(\beta_{k1}X_{ki1} + \beta_{k2}X_{ki2} + \ldots + \beta_{kp}X_{kip})$$

where

 $\triangleright h_{ki}(t)$  the hazard for patient *i* for cause *k* 

 $\triangleright h_{k0}(t)$  the baseline hazard for cause k

 $\triangleright \beta_{k1}, \ldots, \beta_{kp}$  log hazard ratios for the covariates  $X_{k1}, \ldots, X_{kp}$  for cause k

• For its estimation it is correct to treat each failure from other causes as censored



- Example: in the PBC data we are interested in the effects of treatment and age in the hazards for transplantation and death
- We fit the cause-specific hazard models

$$h_{1i}(t) = h_{10}(t) \exp(eta_{11} \texttt{Treat}_i + eta_{12} \texttt{Age}_i)$$

$$h_{2i}(t) = h_{20}(t) \exp(\beta_{21} \operatorname{Treat}_i + \beta_{22} \operatorname{Age}_i)$$

where

 $\triangleright k = 1$  for transplantation, and  $\triangleright k = 2$  for death



Transplantation	est.	$\exp(\text{est.})$	s.e.	<i>p</i> -value
D-penicil – $\beta_{11}$	-0.24	0.79	0.38	0.530
$Age - \beta_{12}$	-0.10	0.91	0.02	< 0.001

Death	est.	$\exp(\text{est.})$	s.e.	<i>p</i> -value
D-penicil – $\beta_{21}$	-0.16	0.85	0.17	0.347
$Age - \beta_{22}$	0.05	1.05	0.01	< 0.001



- We observe that after correcting for treatment
  - ▷ younger patients will receive a transplant sooner, whereas
  - ▷ older patient have greater risk of dying
- These risk estimates **cannot** be converted to relative survival probabilities using the known formula (see pp. 173–175)

$$S(t) = \left\{ S_0(t) \right\}^{\exp(\beta^\top X)}$$

due to the competition between the different causes



• Moreover, the obtained parameter estimates from the cause-specific hazards models do **not** satisfy the relation

$$F_k(t) = \left\{ F_{k0}(t) \right\}^{\exp(\beta_k^\top X_k)}$$

that is, the cumulative incidence functions of different groups (e.g., treated vs untreated) are allowed to cross over

 $\triangleright$  this may or may not be reasonable



• The Fine & Gray model was developed to provide parameters, which satisfy this relation for the cumulative incidence function, i.e.,

$$F_k(t) = \left\{ F_{k0}(t) \right\}^{\exp(\beta_k^\top X_k)}$$

- We will not cover this model further here
  - ▷ more information can be found in Putter et al. (Stat. in Med., 2007)



R> Cox regression for the cause-specific hazards is straightforward to implement using coxph() – for the PBC data set we used



- Often we are interested in assessing the discriminative capability of a covariate
  - can we use LDL cholesterol levels to discriminate between patients with low and high risk of heart disease
  - can we use PSA levels to discriminate between patients with low and high risk of prostate cancer

▷...

• Example: in the AIDS data set we have seen that the baseline CD4 cell count is highly associated with the risk of the death

▷ but how good is CD4 cell count in discriminating between patients?



- We denote the marker (e.g., CD4 cell count) by  $M_i \Rightarrow$  for any threshold c we can define a prediction rule
  - $\triangleright$  if  $M_i > c$ , we classify patient i as a case (she had the event)
  - $\triangleright$  if  $M_i \leq c$ , we classify patient *i* as a non-case (she didn't have the event)
- We borrow ideas from standard ROC analysis let  $D_i$  the case indicator, then

 $\triangleright$  Sensitivity (true positive rate):  $\Pr(M_i > c \mid D_i = 1)$ 

 $\triangleright$  Specificity (true negative rate):  $\Pr(M_i \leq c \mid D_i = 0)$ 



• To depict the discriminative capability of the marker for all possible threshold values *c*, we construct the Receiver Operating Characteristic (ROC) curve

$$TP(c) \ = \ \mathsf{Pr}(M > c \mid D = 1)$$

$$FP(c) = \Pr(M > c \mid D = 0)$$

 $\mathsf{ROC} \hspace{0.1 in}:\hspace{0.1 in} \{FP(c),TP(c)\}, \hspace{0.1 in} \text{for every} \hspace{0.1 in} c$ 



• To summarize the discriminative capability of the marker, we use the area under the ROC curve (AUC)

$$\mathsf{AUC} = \int_0^1 \mathsf{ROC}(p) \, dp$$

- Intuitive interpretation:
  - $\triangleright$  for a randomly chosen pair of patients  $\{i, j\}$  where i is a case and j a control, the AUC is the probability that the marker value for the case is greater than the marker value for the control:

$$AUC = Pr(M_i > M_j | D_i = 1, D_j = 0)$$



- In survival analysis we have to account for time
  - $\triangleright$  the status of some patients changes at some time point
- Possible solution: consider the status of the patients at the end of the study
- Problems:
  - random right censoring, e.g., if a patient is lost to follow-up before the end of the study, her status is unknown at the end of the study
  - b we lose the dynamic nature, e.g., considering discrimination at early stages may be more informative than at the end of the study



- Due to the time dimension, we can have more than one definitions for **Cases** and **Controls**
- For any time t, we can define Cases as
   event (disease, death) before time t
   event (disease, death) at time t
- For any time t, we can define **Controls** as
  - $\triangleright$  event-free through time t
  - $\triangleright$  event-free through a fixed follow-up time  $t^*$



- In the following we follow the work of Heagerty, Lumley & Pepe (Biometrics, 2000)
- At time t, we define
  - $\triangleright$  **Case**: if a patient had the event at any time before  $t \Rightarrow T_i^* \leq t$

 $\triangleright$  **Control**: if a patient did not have the event by time  $t \Rightarrow T_i^* > t$ 

- Features
  - $\triangleright$  at any time t, the entire population is classified as either case or a control
  - $\triangleright$  a patient plays the role of a control for all  $t < T^*_i$  , but she then contributes as a case for  $t \geq T^*_i$



• We can now define the True Positive (sensitivity) and False Positive (1 – specificity) rates

$$TP(c,t) = \Pr(M_i > c \mid T_i^* \le t)$$

$$FP(c,t) \ = \ \mathsf{Pr}(M > c \mid T_i^* > t)$$

- The corresponding ROC and AUC are calculated in exactly the same way as in p.362 and p.363, respectively
- <u>Note:</u> now we have *time-dependent* sensitivity, specificity, ROCs and AUCs



- $\bullet$  We have defined the accuracy measures in the survival context using  $T^{\ast}_i$ , i.e., the true event time
- However, due to censoring we actually only observe  $\{T_i, \delta_i\}$
- $\bullet$  Therefore, in estimating TP(c,t) and FP(c,t) we need to account for censoring



 $\bullet$  To estimate both TP(c,t) and FP(c,t) we only require an estimate of the bivariate survival function

$$S(c,t) = \Pr(M > c, T^* > t) = \int_0^\infty S(t \mid M = m) \, dF_M(m)$$

where  $F_M(m)$  is the cdf for the marker M

• We will use the Nearest Neighbor Estimation method

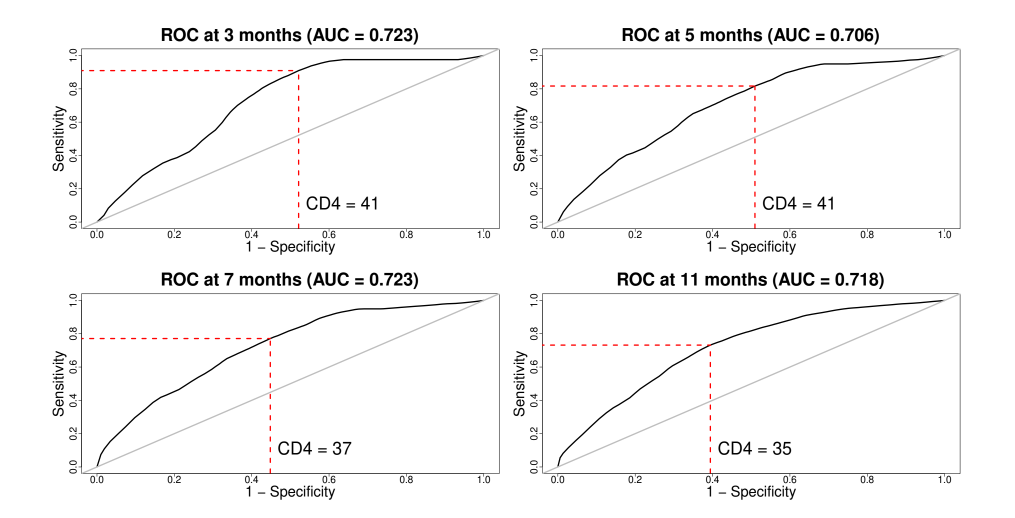
$$\hat{S}_{\lambda}(c,t) = \frac{1}{n} \sum_{i} \hat{S}_{\lambda}(t \mid M = m_i) I(m_i > c)$$

where  $\hat{S}_{\lambda}(t \mid M = m_i)$  is a smooth estimator of the conditional survival function  $\lambda$  controls the degree of smoothness



- Example: in the AIDS data set we have seen that the baseline CD4 cell count is highly associated with the risk of the death
  - ▷ but how good is CD4 cell count in discriminating between patients?
- Which follow-up time is most important?
  - ▷ medical importance
  - ▷ we illustrate time-dependent ROCs and AUCs at 3, 5, 7 and 11 months of follow-up
  - $\triangleright$  we also include an estimate of the 'optimal' cutoff point using the Youden index (Sens + Spec 1)







R> To calculate time-dependent ROC curves in we need to make use of function survivalROC() from package survivalROC

```
# ROC curve at 11 months
sroc <- with(aids.id,
    survivalROC(Time, death, -CD4, predict.time = 11,
    span = 0.55*length(Time)^(-0.20))
)</pre>
```

```
sroc$AUC # area under ROC curve
```



- Obtaining survival probabilities from a Cox model
  - $\triangleright$  the Cox model makes no assumption for the baseline survival function
  - b in order to estimate survival probabilities this baseline survival function needs to be estimated
  - ▷ Breslow estimator (extension from the univariate case)
- Stratified Cox models
  - ▷ the baseline hazard of an event could be different between strata (e.g., hospitals)
  - > categorical covariate do not satisfy the PH assumption
  - > a simple extension of the Cox model is to consider a different baseline hazard per stratum
  - $\triangleright$  disadvantage: we obtain no p-value for the stratification factor



- Time-dependent covariates
  - in many cases we are interested in the effect of covariates whose values changes with time (e.g., time-dependent treatment dose, blood values, etc.)
  - > important distinction: external vs internal time-dependent covariates
  - > external time-dependent covariates can be easily handled within the framework of the extended Cox model
  - $\triangleright$  internal covariates are more difficult and require specialized statistical models  $\rightarrow$  joint modelling of longitudinal and time-to-event data
  - b if internal covariates are treated as external (using the extended Cox model) we may encounter spurious results



- Clustered event data
  - ▷ clustered event times occur frequently (e.g., patients within hospitals)
  - > subjects in the same cluster are expected to be correlated
  - $\triangleright$  these correlations must be taken into account in the analysis
    - \* marginal models (adjusted variance using jackknife)
    - \* frailty models (latent variables)



- Competing risks
  - > often in clinical studies we are interested in multiple endpoints
  - ▷ if the distributions of failure times from different causes are independent, then proceed as usual
  - $\triangleright$  if not, then more care is required: work with
    - \* cause-specific hazards
    - \* cumulative incidence function
    - \* Cox model on cause-specific hazards (be careful of the derived interpretations)



- Discrimination
  - ▷ we aim at discriminating between patients of high and low risk of having the event
  - ROC methodology, estimating the True Positive and False Positive rates
  - b we require special definitions in the survival setting due to censoring and time dimension
  - $\triangleright$  in the estimation of the accuracy measures we need to account for censoring

 $\mathbf{Part}\ \mathbf{VI}$ 

**Closing: Review of Key Points in Survival Analysis** 



- We will learn which are the special characteristics of event time data and why they require special treatment (from a statistical point of view)
- From the course it will become clear
  - > which statistical tools are applicable for this kind of data
  - ▷ which are their advantages and disadvantages
  - ▷ which are the optimal inferential strategies
- What is there further in survival analysis than what we will cover in this course



- Time-to-event data exhibit
  - $\triangleright$  skewed distributions
  - ▷ censoring and/or truncation
- Statistical tools applicable to survival data
  - ▷ Kaplan-Meier estimate of the survival function
  - Log-rank and Peto & Peto modified Gehan-Wilcoxon tests can be used to test whether the survival functions of 2 groups differ statistically significantly
  - > AFT and Cox models can be used to account for effect of more than one explanatory variables in the time-to-event



- Modelling strategies
  - b think carefully about the purpose of modelling (i.e., prediction, effect estimation or hypothesis testing)
  - $\triangleright$  consider which explanatory variables you want to include in the model
  - > relax the linearity assumption of quantitative predictors (splines)
  - > include meaningful interaction terms
  - $\triangleright$  use residuals to check model assumptions
  - ▷ use likelihood ratio tests for hypothesis testing
  - $\triangleright$  use effect plots to communicate the results of the model



- Extending the Cox model
  - ▷ expected survival
  - $\triangleright$  stratification
  - b time-dependent covariates (external & internal)
  - > clustered event times (marginal & frailty models)
  - > competing risks (dependent & independent)



- The last three i.e.,
  - ▷ time-dependent covariates,
  - $\triangleright$  clustered event times, and
  - ▷ competing risks
  - have been briefly covered
- Their full treatment requires more advanced of theoretical statistics and therefore it falls outside the scope of this course



- By now you should have a clear view on the different survival analysis approaches, and how they should be used in practice.
- However, as we have seen throughout this course, statistical analysis is based on assumptions – if these assumptions are seriously violated, we may obtain spurious results
- This is especially the case when we deal with more complex models such as the one required for time-dependent covariates, clustered event times and competing risks



• Therefore, whenever you do not feel sure about the correct type of analysis for a specific research question at hand, consult a local statistician

## The End!