

# STAD—Survival analysis

Mathieu Ribatet—Full Professor of Statistics



## References

---

- [1] D. Collett. *Modelling Survival Data in Medical Research*. Chapman and Hall/CRC, 3rd edition, 2014.
- [2] J. P. Klein and M. L. Moeschberger. *Survival analysis*. Springer–Verlag, 2nd edition, 2003.
- [3] D. G. Kleinbaum and M. Klein. *Survival analysis: A self learning text*. Springer–Verlag, New–York, 3rd edition, 2012.

# What is survival analysis?

---

- Survival analysis is the analysis of times from a time origin until the occurrence of some event or end-point.
- Typical examples are:
  - recruitemet of an individuals (time origin) for an experimental study and first occurence of some symptoms
  - begining of a stress testing (time origin) up to default of an electrical component
- Essentialy you can use survival analysis as long as the data you analyzed are durations.
- It is widely used in medical research but (unfortunately) less used in quality control.

# What is survival analysis?

---

- Survival analysis is the analysis of times from a time origin until the occurrence of some event or end-point.
- Typical examples are:
  - recruitemet of an individuals (time origin) for an experimental study and first occurence of some symptoms
  - begining of a stress testing (time origin) up to default of an electrical component
- Essentialy you can use survival analysis as long as the data you analyzed are durations.
- It is widely used in medical research but (unfortunately) less used in quality control.

☞ Hence do not panick if I use medical datasets and I mainly use “medical phrasing” ;-)

# Special features of survival data

---

- We **cannot** use standard statistical techniques for survival data due to
  - **asymmetry**, e.g., positively skewed, since durations are obviously positive<sup>1</sup>
  - **censoring**, i.e., some observations are not “real” observations (I will be more specific later)
- We thus need a new framework for these data!

---

<sup>1</sup>one could take the log of the data and analyze the transformed data but it is more sensible to work on the original scale

## Examples

---

genfan Failure times of diesel engine fans

imotor Failure times of motor insulation

ovarian Survival times on two different treatments for ovarian cancer

Rossi Time to no recidivism when there is financial support or no financial support on releas

## Examples

---

genfan Failure times of diesel engine fans

imotor Failure times of motor insulation

ovarian Survival times on two different treatments for ovarian cancer

Rossi Time to no recidivism when there is financial support or no financial support on releas

ours [we will create our own data](#)

## Examples

---

genfan Failure times of diesel engine fans

imotor Failure times of motor insulation

ovarian Survival times on two different treatments for ovarian cancer

Rossi Time to no recidivism when there is financial support or no financial support on releas

ours we will create our own data in a few moment!

# The genfan dataset

---

Data set ‘genfan’: Time to failure of 70 diesel engine fans.

- ‘hours’: hours of service
- ‘status’: 1=failure, 0=censored

```
> head(genfan)
  hours status
1    450     1
2    460     0
3   1150     1
4   1150     1
5   1560     0
6   1600     1
```

# The imotor dataset

---

Data set ‘imotor’: Breakdown of motor insulation as a function of temperature.

- temp: temperature of the test
- time: time to failure or censoring
- status: 0=censored, 1=failed

```
> head(imotor)
  temp time status
1 150 8064     0
2 150 8064     0
3 150 8064     0
4 150 8064     0
5 150 8064     0
6 150 8064     0
```

# The ovarian dataset

---

Ovarian Cancer Survival Data

Description:

Survival in a randomised trial comparing two treatments for ovarian cancer

Format:

futime: survival or censoring time  
fustat: censoring status  
age: in years  
resid.ds: residual disease present (1=no,2=yes)  
rx: treatment group  
ecog.ps: ECOG performance status (1 is better, see reference)

head(ovarian)

	futime	fustat	age	resid.ds	rx	ecog.ps
1	59	1	72.3315	2	1	1
2	115	1	74.4932	2	1	1
3	156	1	66.4658	2	1	2
4	421	0	53.3644	2	2	1
5	431	1	50.3397	2	1	1
6	448	0	56.4301	1	1	2

# The Rossi dataset

---

Description:

Time to recidivism on 432 convicts who were released from Maryland in the 70s (half of them get financial support)

Format:

week: time to recidivism or censoring time  
arrest: arrtested or not, i.e., censoring status  
fin: is there a financial support  
age: age (years) at release  
race: black or other  
wexp: has a working experience after release  
mar: is married on release  
paro: was released on parole  
prio: number of prior conviction  
educ: a factor encoding the level of education

> head(data)

	week	arrest	fin	age	race	wexp	mar	paro	prio	educ
1	20	1	no	27	black	no	not married	yes	3	3
2	17	1	no	18	black	no	not married	yes	8	4
3	25	1	no	19	other	yes	not married	yes	13	3
4	52	0	yes	23	black	yes	married	yes	1	5

# Let's introduce ourselves

---

- Who am I?
- Who are you?
- What do you know about statistics and probability?

## The Kaplan–Meier theatre (from Thomas A. Gerds)

---

- We are all on the Titanic, and the Titanic is going down. Once under water, you would have to hold your breath. But how long can you do this?
- Pick up a sheet of paper and write:
  - an unique identifier
  - your sex, smoking status, un/like sport
- Pick up your smartphone and open a timer
- The Titanic is about to sink and as I say your personal id you should start holding your breath
- If you stop holding your breath at some time, write it down on your sheet of paper
- If you hold your breath until I said stop, just write the current time on your timer adding a plus after it, e.g., 52.3+.

# Let's talk about our data

---

▷ 1. Preliminaries

2. Non parametric  
estimation

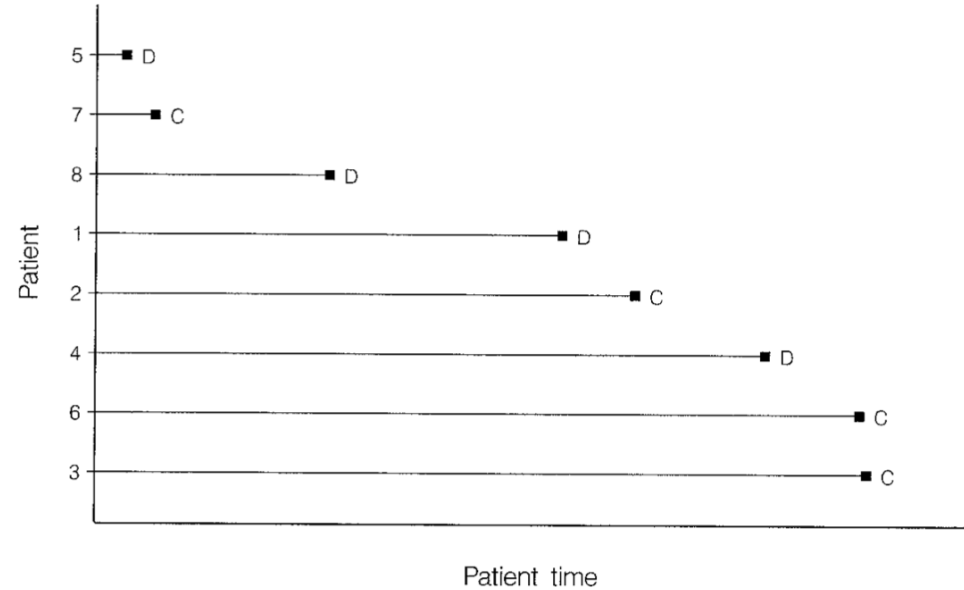
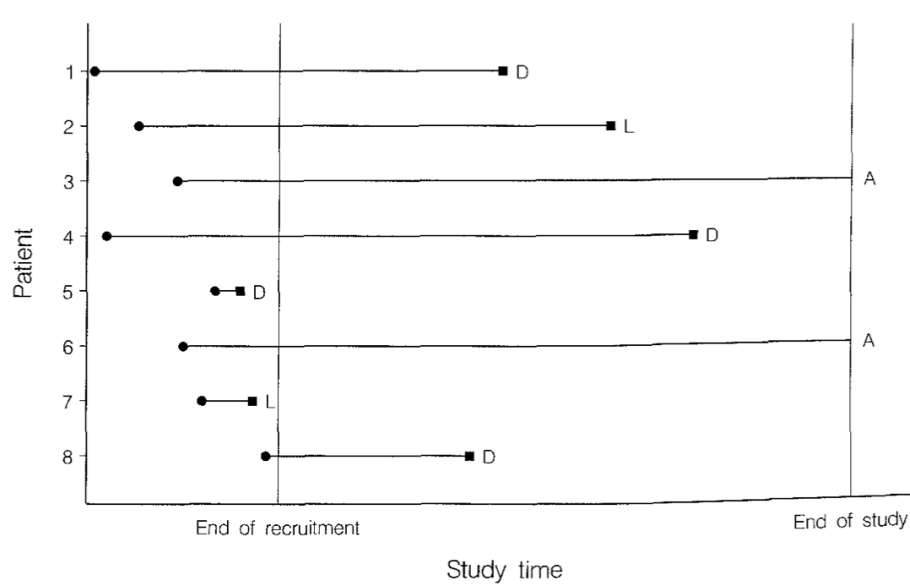
3. Coping with  
covariates

4. Time to recidivism

Conclusion

# 1. Preliminaries

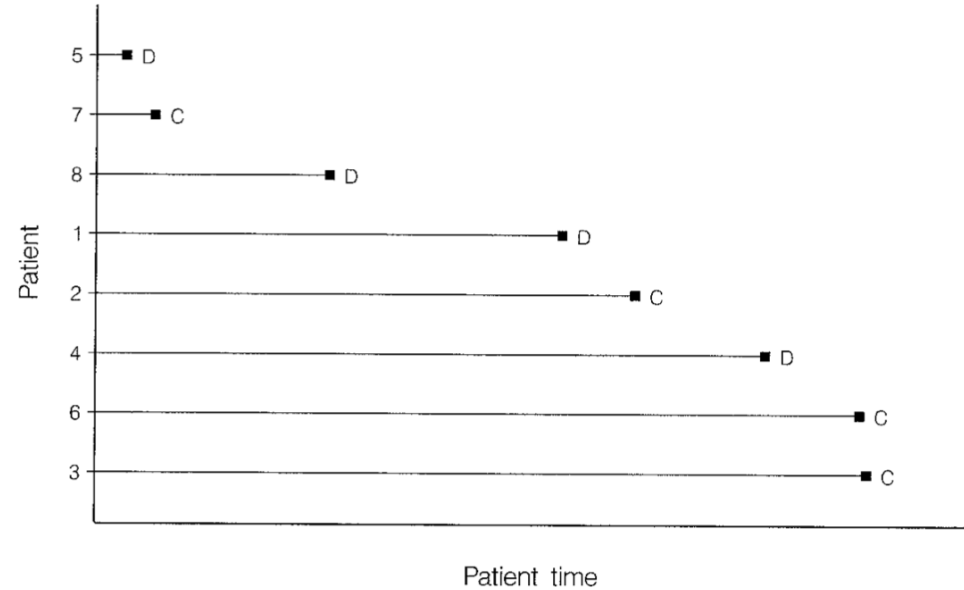
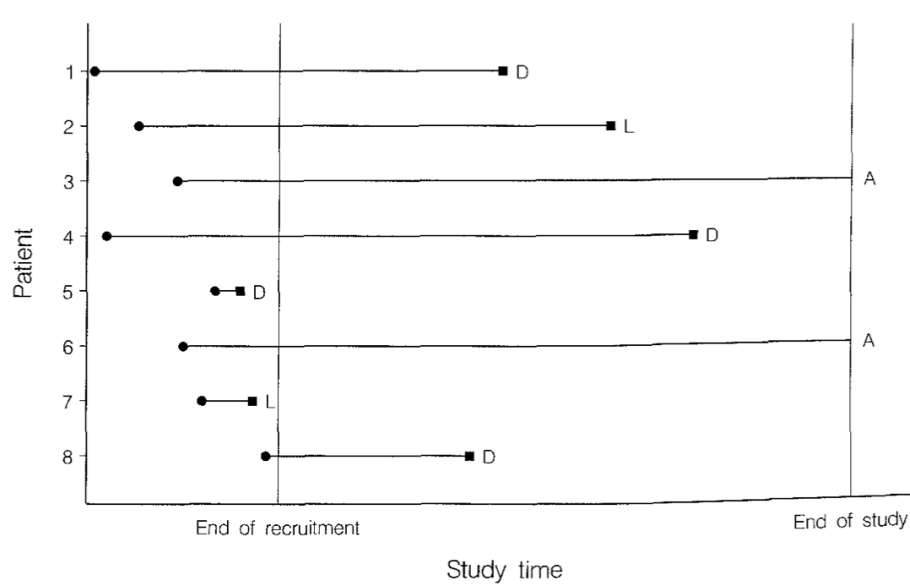
# Patient time and study time



**Figure 1:** Illustration of patient and study times. Left: Study time. (D: death, L: lost to follow-up, A: alive). Right: Patient time (C: right censored).

- Often patients are not all recruited at the same time
- We have to pay attention to the difference between **study** and **patient** time

# Patient time and study time



**Figure 1:** Illustration of patient and study times. Left: Study time. (D: death, L: lost to follow-up, A: alive). Right: Patient time (C: right censored).

- Often patients are not all recruited at the same time
- We have to pay attention to the difference between **study** and **patient** time
- Statistical modelling is often made on **patient time**

## p.d.f., c.d.f. and co

---

- Let  $T$  be a continuous **positive** random variable having a **probability density function**  $f$ , i.e.,

$$f(t) \geq 0, \quad \int_0^\infty f(t)dt = 1.$$

- The **cumulative distribution function** of  $T$  is

$$F(t) = \Pr(T \leq t) = \Pr(T < t) = \int_0^t f(u)du,$$

and gives the **probability that the survival time is less than  $t$ .**

- The **survival function** of  $T$  is

$$S(t) = \Pr(T > t) = \int_t^\infty f(u)du = 1 - F(t),$$

and gives the **probability that the survival time is greater than  $t$ .**

## Hazard rate and cumulative hazard

---

- In survival analysis we often talk about the hazard rate

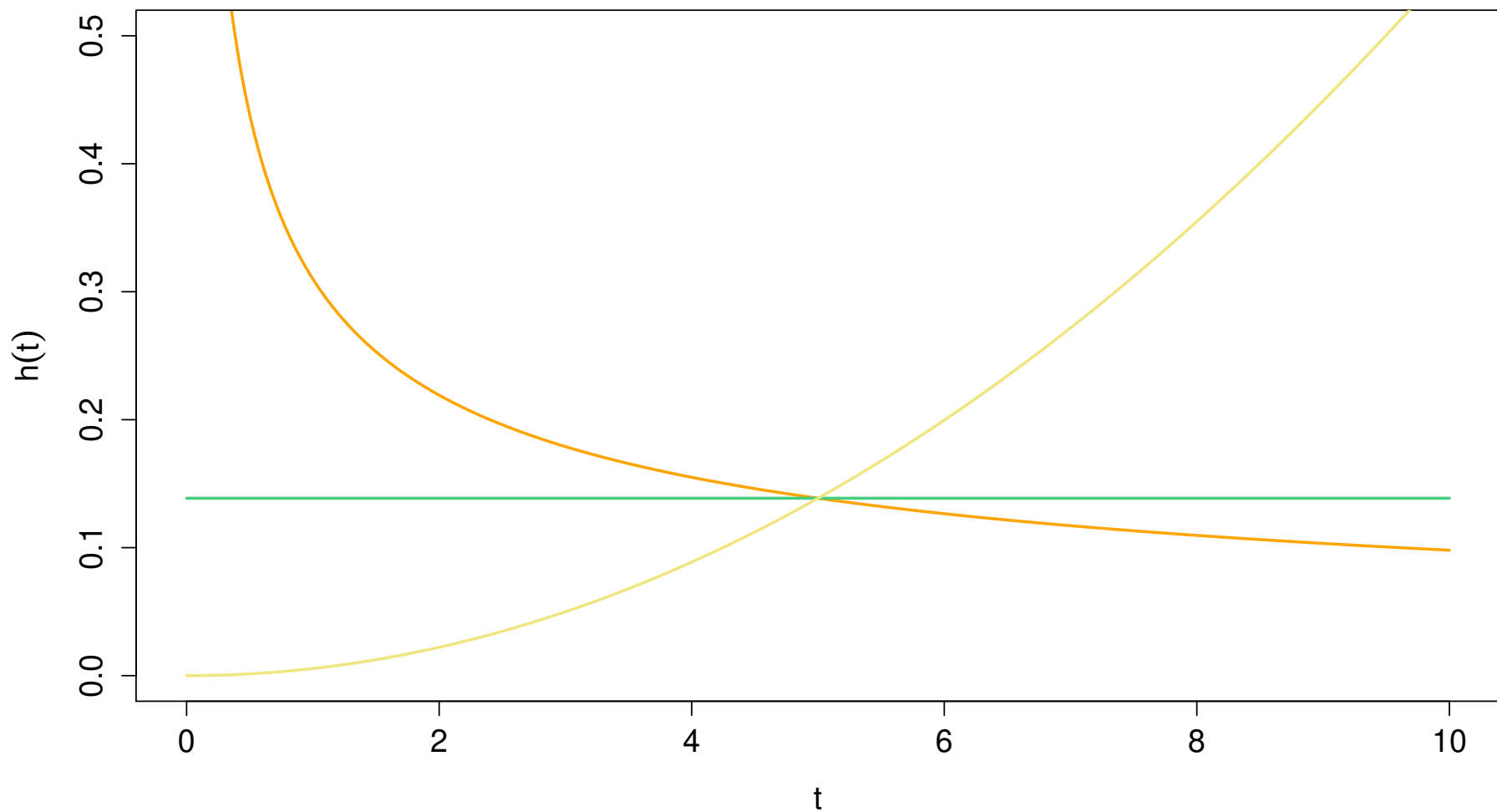
$$h(t) = \lim_{\Delta t \downarrow 0} \frac{\Pr(t \leq T < t + \Delta t \mid T \geq t)}{\Delta t},$$

and the cumulative hazard

$$H(t) = \int_0^t h(u) du.$$

**Exercise 1.** If  $T$  has density  $f$ , show that we have for all  $t > 0$

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



**Figure 2:** Plot of three different hazard rate (obtained from a Weibull distribution).

# Censoring

---

**Definition 1.** An observation is said to be **censored** when the **end-point** has not been observed.

**Example 1.** lost to follow-up, child knows how to read before school, component still OK as the stress testing phase ends, end-point is not related to our analysis...

- There are 3 different type of censoring:
  - right censoring where we observed  $T = \min(C, T_*)$ ;
  - left censoring where we observed  $T = \max(C, T_*)$ ;
  - and interval censoring where we just know  $T_* \in [A, B]$ .

# Censoring

---

**Definition 1.** An observation is said to be **censored** when the **end-point** has not been observed.

**Example 1.** lost to follow-up, child knows how to read before school, component still OK as the stress testing phase ends, end-point is not related to our analysis...

- There are 3 different type of censoring:
  - right censoring where we observed  $T = \min(C, T_*)$ ;
  - left censoring where we observed  $T = \max(C, T_*)$ ;
  - and interval censoring where we just know  $T_* \in [A, B]$ .

☞ The most common censoring is by far **right censoring**.

---

1. Preliminaries

2. Non parametric  
▷ estimation

---

3. Coping with  
covariates

---

4. Time to recidivism

---

Conclusion

---

## 2. Non parametric estimation

# Non parametric statistics

---

- Most often stochastic modelling assumes that the random variable  $T$  under study has a **pre-specified distribution**, e.g.,  $T \sim \text{Log-Normal}$ .
- Non parametric statistics make no **distributional assumptions** on  $T$
- The price to pay is most often the speed of convergence, statistical power, e.g., larger variance of the estimator...
- However **non parametric procedures** are appealing as they can be used later for **model checking**.

## Naive estimate

---

- Remember the empirical distribution function

$$\hat{F}(t) = \frac{1}{n} \sum_{i=1}^n 1_{\{T_i \leq t\}}, \quad t > 0,$$

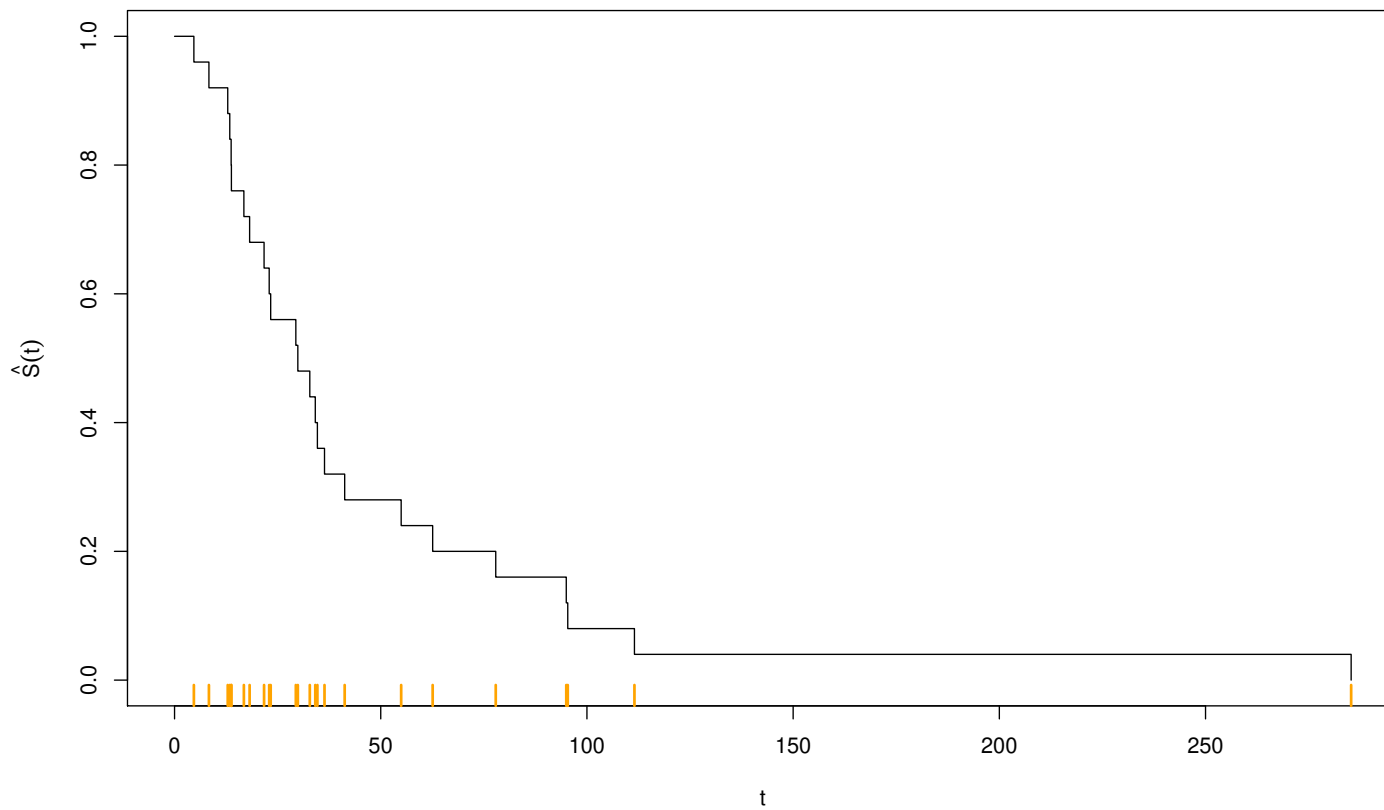
where  $T_i \stackrel{\text{iid}}{\sim} F$ .

- The law of large number implies, provided  $\mathbb{E}(T) < \infty$ ,

$$\hat{F}(t) \longrightarrow \mathbb{E}[1_{\{T \leq t\}}] = F(t), \quad n \rightarrow \infty, \quad t > 0.$$

- Equivalently we can estimate the survivor function from

$$\hat{S}(t) = 1 - \hat{F}(t) = \frac{1}{n} \sum_{i=1}^n 1_{\{T_i > t\}}.$$



**Figure 3:** An example of a naive estimate of the survival function.

- It is a **step function** (actually cadlag).
- We always have  $\hat{S}(0) = 1$  and  $\hat{S}(\max T_i) = 0$ .

# There's a trap!

---

- The above estimator is usually **consistent** and **unbiased**.
- But it is not the case anymore when there are **censored observations**!
- Why? Essentially because some  $T_i \not\sim F\dots$
- We need to use different estimators for  $S(t)$ :
  - Kaplan–Meier estimator
  - Nelson–Aalen estimator
  - Life–table estimator
- We will cover these estimators in turn.

## Towards the Kaplan–Meier estimator

---

- Consider the **order statistics** (without ties) of  $n$  durations (censored or not)  
 $t_1, \dots, t_n$ , i.e.,  $0 = t_{(0)} < t_{(1)} < \dots < t_{(n)}$ .
- We thus have for any  $j \in \{1, \dots, n\}$

$$\begin{aligned}\Pr(T > t_{(j)}) &= \Pr(T > t_{(j)}, T > t_{(j-1)}) \\ &= \Pr(T > t_{(j)} \mid T > t_{(j-1)}) \Pr(T > t_{(j-1)}) \\ &= \dots \\ &= \prod_{\ell=1}^j \Pr(T > t_{(\ell)} \mid T > t_{(\ell-1)}).\end{aligned}$$

- The Kaplan–Meier estimator uses this decomposition and plugs in empirical estimator for these conditional probabilities

## Kaplan–Meier estimator

---

- Each conditional probability  $p_\ell = \Pr(T > t_{(\ell)} \mid T > t_{(\ell-1)})$  is easily estimated from its **empirical counterpart**

$$\hat{p}_\ell = 1 - \frac{d_\ell}{n_\ell}, \quad d_\ell = \# \text{ deaths at } t_{(\ell)}, \quad n_\ell = \# \text{ at risk at } t_{(\ell)}^-.$$

- This gives the **Kaplan–Meier estimator**

$$\hat{S}_{KM}(t) = \prod_{i: T_i \leq t} \left(1 - \frac{d_i}{n_i}\right), \quad t > 0.$$

## Application: failure times (hours) of diesel engine fans

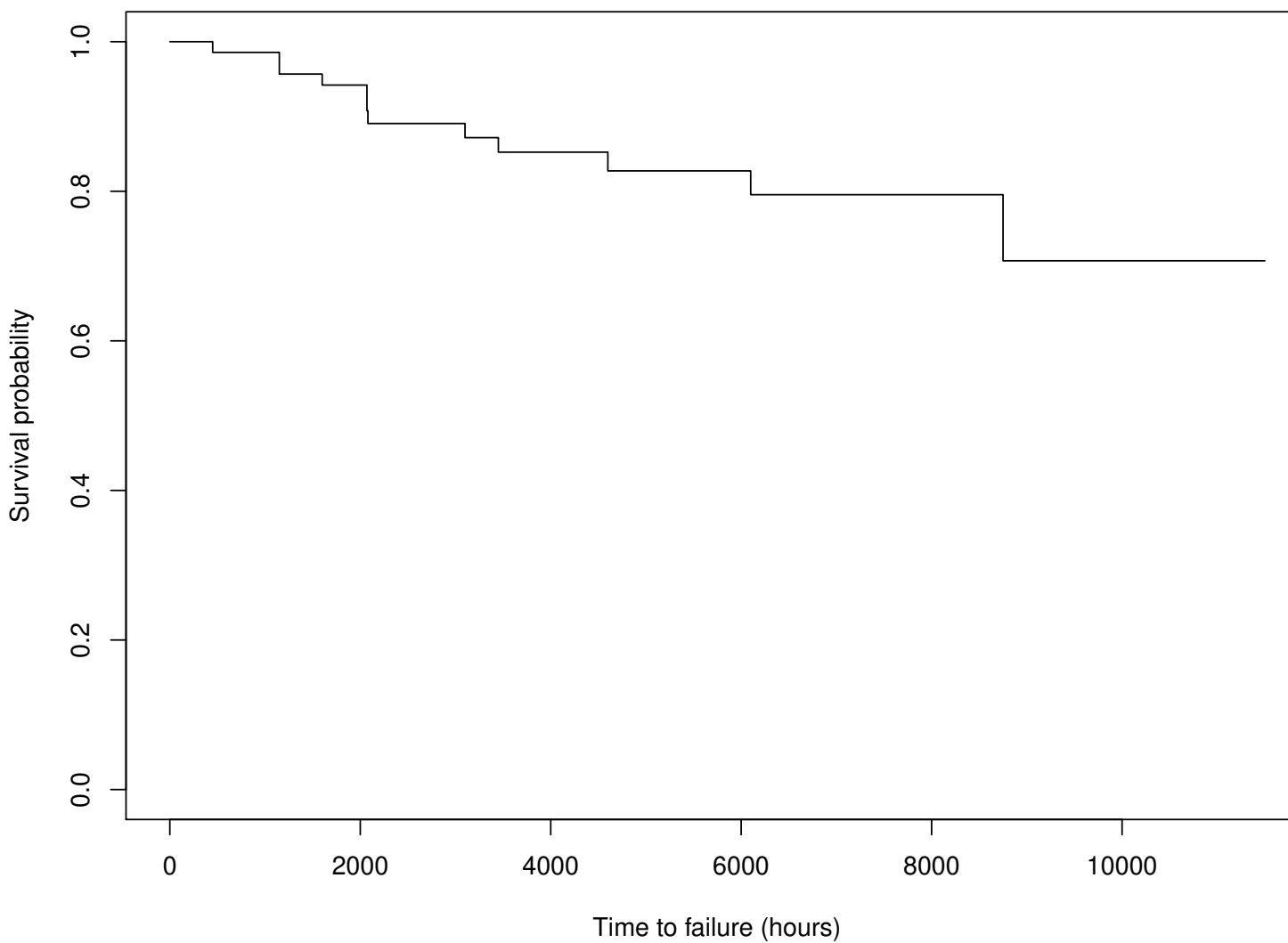
450	1850+	2030+	3000+	4150+	4300+	5000+	6450+	8100+	8750+
460+	1850+	2070	3000+	4150+	4600	5000+	6450+	8200+	9400+
1150	1850+	2070	3100	4150+	4850+	6100+	6700+	8500+	9900+
1150	1850+	2080	3200+	4150+	4850+	6100	7450+	8500+	10100+
1560+	1850+	2200+	3450	4300+	4850+	6100+	7800+	8500+	10100+
1600	2030+	3000+	3750+	4300+	4850+	6100+	7800+	8750+	10100+
1660+	2030+	3000+	3750+	4300+	5000+	6300+	8100+	8750	11500+

## Application: failure times (hours) of diesel engine fans

450	1850+	2030+	3000+	4150+	4300+	5000+	6450+	8100+	8750+
460+	1850+	2070	3000+	4150+	4600	5000+	6450+	8200+	9400+
1150	1850+	2070	3100	4150+	4850+	6100+	6700+	8500+	9900+
1150	1850+	2080	3200+	4150+	4850+	6100	7450+	8500+	10100+
1560+	1850+	2200+	3450	4300+	4850+	6100+	7800+	8500+	10100+
1600	2030+	3000+	3750+	4300+	4850+	6100+	7800+	8750+	10100+
1660+	2030+	3000+	3750+	4300+	5000+	6300+	8100+	8750	11500+

**Table 1:** Kaplan–Meier estimate of the survival function on the genfan dataset.

Time	$n_{\text{risk}}$	$n_{\text{event}}$	$\hat{S}_{KM}$
0	70	0	1.00
450	70	1	0.99
1150	68	2	0.96
1600	65	1	0.94
2070	55	2	0.91
2080	53	1	0.89
3100	47	1	0.87
3450	45	1	0.85
4600	34	1	0.83
6100	26	1	0.80
8750	9	1	0.71



**Figure 4:** Kaplan–Meier estimate for the genfan dataset.

## Ties in Kaplan–Meier

- We may have ties, e.g.,  $t_1 = t_2$ , and in such situations we have
  - $d_1 = 2$  if obs. 1 and 2 are “death”
  - if obs. 1 is censored and 2 isn’t, we assume that “death” happens before

**Table 2:** Kaplan–Meier estimate of the survival function on the genfan dataset.

Time	$n_{\text{risk}}$	$n_{\text{event}}$	$\hat{S}_{KM}$
0	70	0	1.00
450	70	1	0.99
1150	68	2	0.96
1600	65	1	0.94
2070	55	2	0.91
2080	53	1	0.89
3100	47	1	0.87
3450	45	1	0.85
4600	34	1	0.83
6100	26	1	0.80
8750	9	1	0.71

## Delta–method

---

**Theorem 1.** Let  $X_1, X_2, \dots$  a sequence of random variables such that

$$\sqrt{n}(X_n - \theta) \xrightarrow{d.} N(0, \sigma^2), \quad n \rightarrow \infty,$$

for some fixed values  $\theta \in \mathbb{R}$ . Let  $g$  be a function such that  $g'(\theta)$  exists and is non null. Then

$$\sqrt{n}(g(X_n) - g(\theta)) \xrightarrow{d.} N(0, g'(\theta)^2 \sigma^2), \quad n \rightarrow \infty.$$

*Proof.* Taylor expansion + Continuous mapping theorem + Slutsky's lemma □

## Delta–method

---

**Theorem 1.** Let  $X_1, X_2, \dots$  a sequence of random variables such that

$$\sqrt{n}(X_n - \theta) \xrightarrow{d.} N(0, \sigma^2), \quad n \rightarrow \infty,$$

for some fixed values  $\theta \in \mathbb{R}$ . Let  $g$  be a function such that  $g'(\theta)$  exists and is non null. Then

$$\sqrt{n}(g(X_n) - g(\theta)) \xrightarrow{d.} N(0, g'(\theta)^2 \sigma^2), \quad n \rightarrow \infty.$$

*Proof.* Taylor expansion + Continuous mapping theorem + Slutsky's lemma

□

☞ The delta–method is very useful when one have to compute (asymptotic) variance of a transformation of an estimator

## Simple application

---

**Exercise 2.** Let  $X_1, X_2, \dots$  be independent copies of a random variable  $X$  such that  $\mu = \mathbb{E}(\log X)$  and  $\sigma^2 = \text{Var}(\log X) < \infty$ . How would you get an approximation, i.e., as long as  $n$  is large enough, for the variance of the geometric mean of those  $X_i$ 's?

## Variance of Kaplan–Meier

---

$$\hat{S}_{KM}(t) = \prod_{i: T_i \leq t} \left(1 - \frac{d_i}{n_i}\right), \quad t > 0.$$

- Computing the variance of a product is a mess but that of a sum of i.i.d. random variable is easy.

# Variance of Kaplan–Meier

$$\hat{S}_{KM}(t) = \prod_{i: T_i \leq t} \left(1 - \frac{d_i}{n_i}\right), \quad t > 0.$$

- Computing the variance of a **product** is a mess but that of a **sum** of i.i.d. random variable is easy. Idea: take the log and use Delta–method!
- Doing so we get the **Greenwood formula**

$$\widehat{\text{Var}}(\hat{S}_{KM}(t)) = \hat{S}_{KM}(t)^2 \sum_{i: T_i \leq t} \frac{d_i}{n_i(n_i - d_i)}.$$

- Since Kaplan–Meier is asymptotically normal, a **pointwise confidence interval** for  $S(t)$  of level  $\alpha$  is

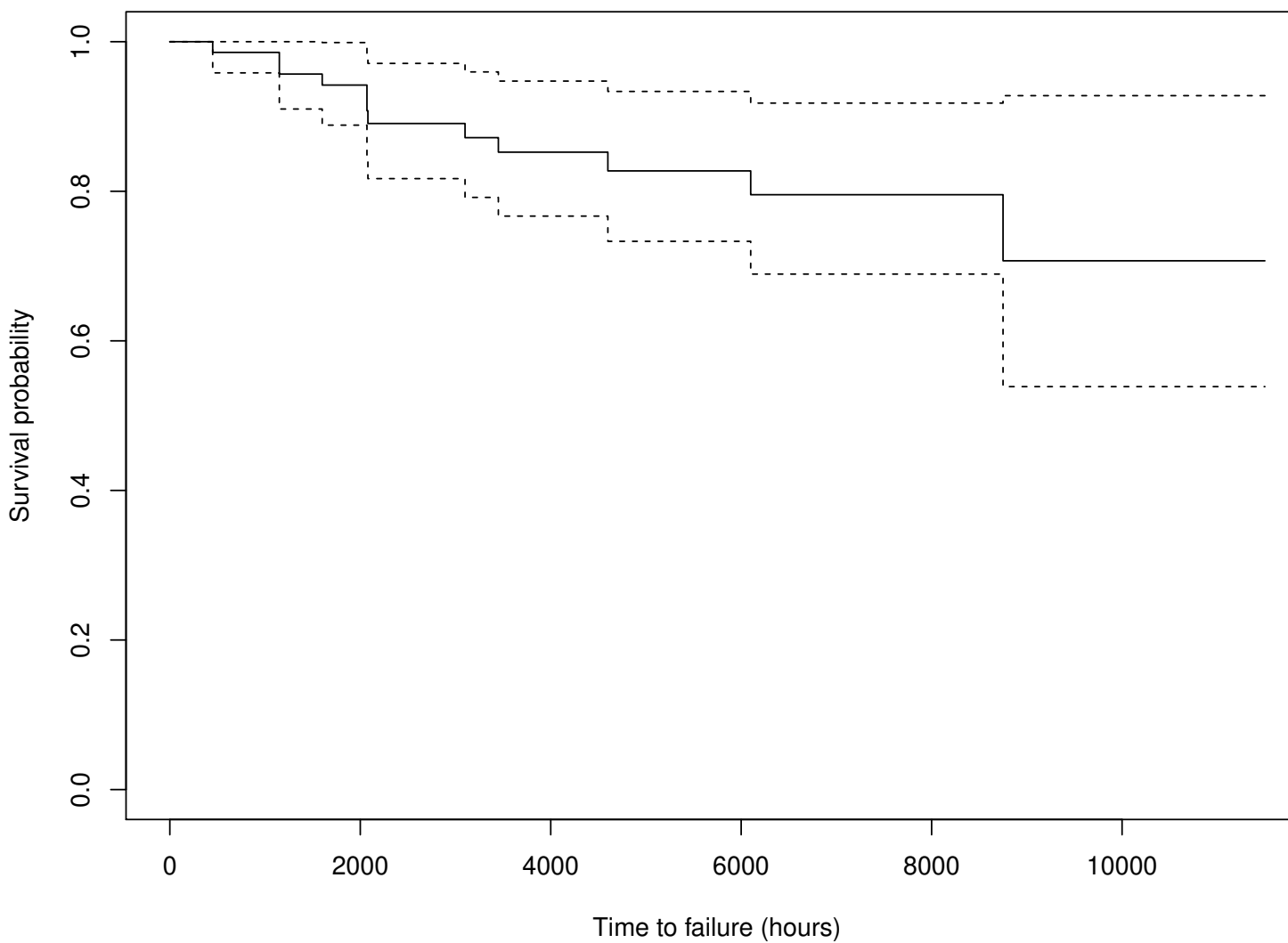
$$\left[ \hat{S}_{KM}(t) - z_{1-(1-\alpha)/2} \sqrt{\widehat{\text{Var}}(\hat{S}_{KM}(t))}, \hat{S}_{KM}(t) + z_{1-(1-\alpha)/2} \sqrt{\widehat{\text{Var}}(\hat{S}_{KM}(t))} \right],$$

e.g.,  $z_{1-(1-\alpha)/2} = 1.96$  when  $\alpha = 95\%$ .

# Application: failure times (hours) of diesel engine fans

**Table 3:** Kaplan–Meier estimate of the survival function on the genfan dataset [with confidence intervals](#).

Time	$n_{\text{risk}}$	$n_{\text{event}}$	$\hat{S}_{KM}$	Std. err.	lower 95% CI	upper 95% CI
450	70	1	0.99	0.01	0.96	1.00
1150	68	2	0.96	0.02	0.91	1.00
1600	65	1	0.94	0.03	0.89	1.00
2070	55	2	0.91	0.04	0.84	0.98
2080	53	1	0.89	0.04	0.82	0.97
3100	47	1	0.87	0.04	0.79	0.96
3450	45	1	0.85	0.05	0.77	0.95
4600	34	1	0.83	0.05	0.73	0.93
6100	26	1	0.80	0.06	0.69	0.92
8750	9	1	0.71	0.10	0.54	0.93



**Figure 5:** Kaplan–Meier estimate for the genfan dataset.

## Derivation (quite) of the Greenwood's formula

---

$$\hat{S}_{KM}(t) = \prod_{i: T_i \leq t} \left(1 - \frac{d_i}{n_i}\right) = \prod_{i: T_i \leq t} \hat{p}_i \iff \log \hat{S}_{KM}(t) = \sum_{i: T_i \leq t} \log \hat{p}_i,$$

where  $\hat{p}_i = (n_i - d_i)/n_i$  is an estimator of  $p_i$  the “true” probability to survive in the  $i$ -th interval. So...

## Wait a minute...

$$[\hat{S}_{KM}(t) - \Delta_\alpha, \hat{S}_{KM}(t) + \Delta_\alpha]$$



- One way to fix this is to set the lower bound to  $\max\{0, \hat{S}_{KM}(t) - \Delta_\alpha\}$  and the upper bound to  $\min\{1, \hat{S}_{KM}(t) + \Delta_\alpha\}$ .
- Another strategy is to use the **log-log survivorship function**, i.e.,

$$\hat{S}_{KM}(t) \longmapsto \log\{-\log \hat{S}_{KM}(t)\},$$

which is a **one-one mapping** that maps  $[0, 1]$  to  $\mathbb{R}$ .

- Again using our best friend Delta-method, we will have a confidence interval that always belongs to  $[0, 1]$ .

- 
- The Greenwood formula uses the following estimator

$$\widehat{\text{Var}}(\log \hat{S}_{KM}(t)) = \sum_{i: T_i \leq t} \frac{d_i}{n_i(n_i - d_i)}$$

- This suggests following estimator for  $\log\{-\log S(t)\}$

$$\Delta^2 := \widehat{\text{Var}}(\log\{-\log \hat{S}_{KM}(t)\}) = \left( -\frac{1}{\log \hat{S}_{KM}(t)} \right)^2 \widehat{\text{Var}}(\log \hat{S}_{KM}(t)).$$

- A symmetric confidence interval for  $\log\{-\log S(t)\}$  is thus

$$\left[ \log\{-\log \hat{S}_{KM}(t)\} - z_{1-(1-\alpha)/2} \Delta, \log\{-\log \hat{S}_{KM}(t)\} + z_{1-(1-\alpha)/2} \Delta \right],$$

- 
- Recall that

$$IC = \left[ \log\{-\log \hat{S}_{KM}(t)\} - z_{1-(1-\alpha)/2}\Delta, \log\{-\log \hat{S}_{KM}(t)\} + z_{1-(1-\alpha)/2}\Delta \right],$$

which by construction satisfies

$$\Pr(\log\{-\log S(t)\} \in IC) \rightarrow \alpha, \quad n \rightarrow \infty.$$

- Now switching back to the original scale we get, i.e., applying pointwise  $f : x \mapsto \exp\{-\exp(x)\}$ , we get

$$\Pr\{S(t) \in f(IC)\} \rightarrow \alpha, \quad n \rightarrow \infty,$$

where

$$f(IC) = \left[ \hat{S}_{KM}(t)^{\exp(z_{1-(1-\alpha)/2}\Delta)}, \hat{S}_{KM}(t)^{\exp(-z_{1-(1-\alpha)/2}\Delta)} \right]$$

## Some key features of Kaplan–Meier

---

- If we have no censoring, Kaplan–Meier corresponds to the usual empirical survivor function
- Let  $T_{(n)}$  be the largest observation. There are two cases:
  - it is an event, then the last term is obviously

$$1 - \frac{1}{1} = 0,$$

and  $\hat{S}_{KM}(t) = 0$  for all  $t \geq T_{(n)}$

- it is a censored observation, then the last term will be strictly positive and  $\hat{S}_{KM}(t) > 0$  for all  $t \geq T_{(n-1)}^2$

---

<sup>2</sup>where we suppose that  $T_{(n-1)}$  is an event otherwise move to the last event.

## Life-table estimator

- The life-table estimator, a.k.a. the actuarial estimator, consists in discretizing the time domain  $[0, D]$  rather than using the observed time events.
- Let  $D = \cup_{j=1}^m I_j$ ,  $I_j = [t'_{j-1}, t'_j)$ , where  $t_0 = 0 < t_1 < \dots < t_{m-1} < t_m = \infty$ , and denote

$$n_j = \# \text{ alive at } t'_j, \quad d_j = \# \text{ deaths in } I_j, \quad c_j = \# \text{ censoring in } I_j.$$

- The life-table estimator assumes that within  $I_j$  the censoring process is uniform over  $I_j$  so that the expected number of observation at risk within  $I_j$  is  $\tilde{n}_j = n_j - c_j/2$ .
- This leads to the life-table estimator

$$\hat{S}_{LF}(t) = \prod_{i: T_i \leq t} \left(1 - \frac{d_i}{\tilde{n}_i}\right), \quad \widehat{\text{Var}}\{\hat{S}_{LF}(t)\} = \hat{S}_{LF}(t) \sum_{i: T_i \leq t} \frac{d_i}{\tilde{n}_i(\tilde{n}_i - d_i)}.$$

## Nelson–Aalen estimator

---

- The Nelson–Aalen estimator is an estimator of the cumulative hazard, i.e.,

$$H(t) = \int_0^t \lambda(u)du, \quad t \geq 0.$$

- The mathematical foundations for this estimator is quite technical (based on point processes and martingales) and beyond the scope of this lecture.
- The Nelson–Aalen estimator is given by

$$\hat{H}_{NA}(t) = \sum_{i: T_i \leq t} \frac{d_i}{n_i}, \quad t \geq 0,$$

and an estimator for its variance is

$$\widehat{\text{Var}}\{\hat{H}_{NA}(t)\} = \sum_{i: T_i \leq t} \frac{d_i}{n_i^2}.$$

## New from old

---

- The Nelson–Aalen estimator can be used to estimate the **survivor function** since  $S(t) = \exp\{-H(t)\}$ ,  $t \geq 0$ .
- This leads to the **Flemming–Harington estimator**

$$\hat{S}_{FH}(t) = \exp\{-\hat{H}_{NA}(t)\}, \quad t \geq 0.$$

- Conversely one can get an estimator of the cumulative hazard using the Kaplan–Meier and life-table estimators

$$\hat{H}_{KM}(t) = -\log \hat{S}_{KM}(t), \quad \hat{H}(t) = -\log \hat{S}_{LT}(t), \quad t \geq 0.$$

**Exercise 3.** Try to give an estimate of the variance of the Flemming–Harington estimator.

## As an aside

---

- First note that

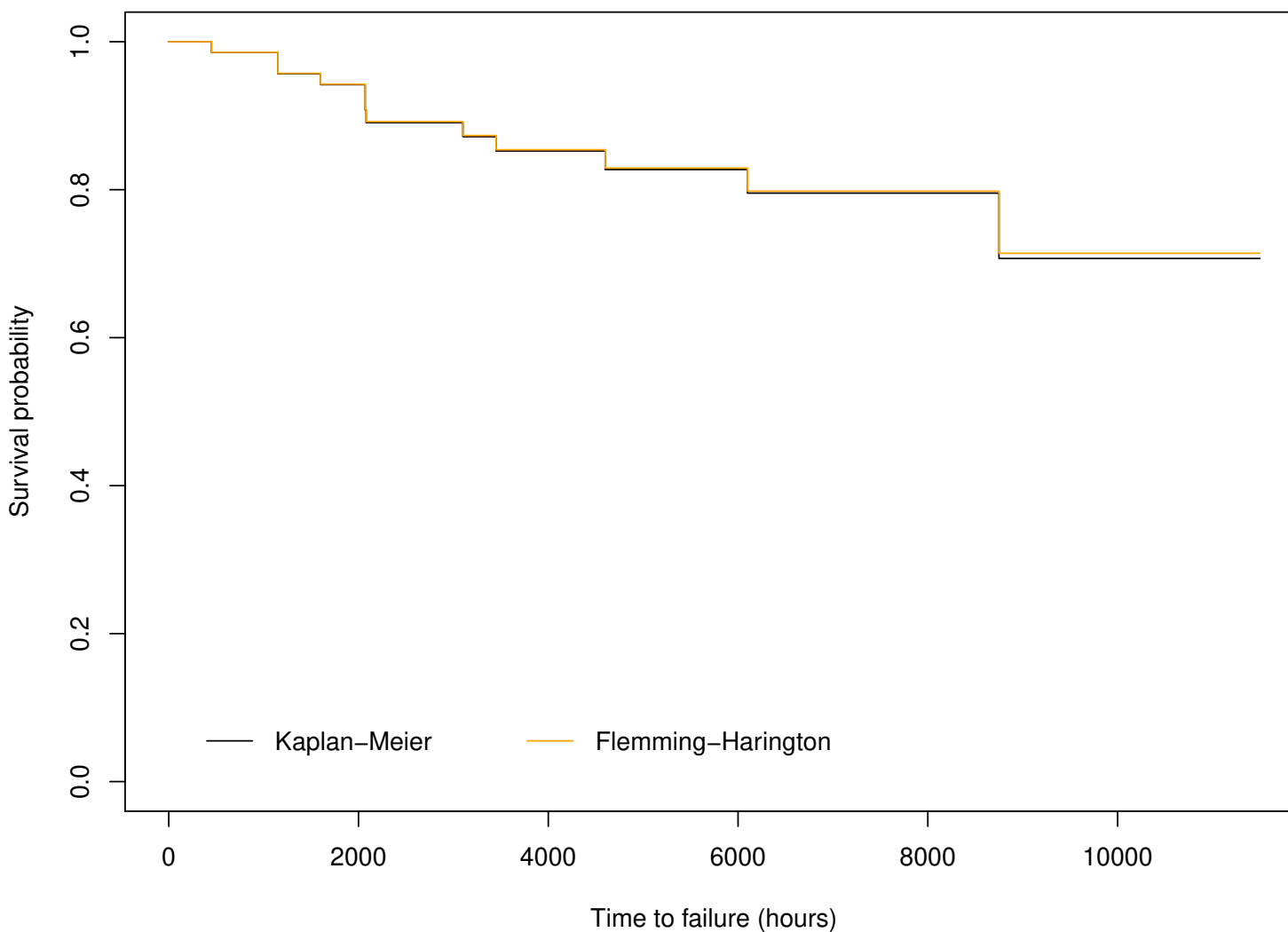
$$\hat{S}_{FH}(t) = \exp\{-\hat{H}_{NA}(t)\} = \exp\left(-\sum_{i: T_i \leq t} \frac{d_i}{n_i}\right) = \prod_{i: T_i \leq t} \exp\left(-\frac{d_i}{n_i}\right).$$

- Since  $\exp(-x) = 1 - x + o(x)$ , we thus have

$$\hat{S}_{FH}(t) \approx \prod_{i: T_i \leq t} \left(1 - \frac{d_i}{n_i}\right) := \hat{S}_{KM}(t),$$

as long as  $d_i \ll n_i$  (which will be true except for the largest survival times).

- We can go even further by recalling that  $\exp(x) \geq 1 - x$  for all  $x \in \mathbb{R}$ . Hence we **always** have  $\hat{S}_{FH}(t) \geq \hat{S}_{KM}(t)$  for all  $t \geq 0$ .



**Figure 6:** Comparison of the Kaplan–Meier and Flemming–Harington estimates for the genfan dataset. As expected  $\hat{S}_{FH}(t) \geq \hat{S}_{KM}(t)$  and both estimates are close as  $d_i \ll n_i$ .

## Estimating the median of survival times

---

- Since the distribution of survival times is often positively skewed, it is preferable to use the median rather than the expectation as a measure of location.
- The median is easily estimating from  $\hat{S}(t)$  (whatever estimator you chose).
- By definition, the median, denoted by  $t_{0.5}$ , satisfies  $S(t_{0.5}) = 0.5$  and we just substitute  $S$  for  $\hat{S}$ .
- Some care is needed though, since  $\hat{S}$  is a step function, and we use

$$\hat{t}_{0.50} = \min\{t_i : \hat{S}(t_i) < 0.5\}.$$

- In the very unlucky situation where  $\hat{S}(t) = 0.5$  for  $t \in [t_{(i)}, t_{(i+1)}]$ , the convention is to set

$$\hat{t}_{0.50} = \frac{t_{(i)} + t_{(i+1)}}{2}.$$

## Estimating quantiles of survival times

---

- The above methodology easily extends to quantiles of arbitrary order  $p$ ,  $0 < p < 1$ .
- By definition, the quantile of order  $p$ , denoted by  $t_p$ , satisfies  $S(t_p) = 1 - p$ .
- Hence an estimator for  $t_p$  is

$$\hat{t}_p = \min\{t_i : \hat{S}(t_i) < 1 - p\}.$$

- It may happen that  $\hat{S}(t) > 1 - p$  for all  $t \geq 0$ . In such situation,  $\hat{t}_p$  is left undefined.

## Variance of $\hat{t}_p$

---

- To get an estimator for  $\text{Var}(\hat{t}_p)$ , we start by noticing that

$$\text{Var}\{\hat{S}(\hat{t}_p)\} \approx (-f(t_p))^2 \text{Var}(\hat{t}_p).$$

- Hence we can derive an estimator for  $\text{Var}(\hat{t}_p)$

$$\widehat{\text{Var}}(\hat{t}_p) = \frac{1}{\hat{f}(\hat{t}_p)^2} \widehat{\text{Var}}\{\hat{S}(\hat{t}_p)\}.$$

- In the above estimator,  $\widehat{\text{Var}}\{\hat{S}(\hat{t}_p)\}$  is obtained from the Kaplan–Meier or Fleming–Harrington variance, while  $\hat{f}(\hat{t}_p)$  is typically estimated using **finite differences** on  $\hat{S}$ .
- Confidence intervals are done using the **usual way**, i.e.,

$$\left[ \hat{t}_p - z_{1-(1-\alpha)/2} \sqrt{\widehat{\text{Var}}(\hat{t}_p)}, \hat{t}_p + z_{1-(1-\alpha)/2} \sqrt{\widehat{\text{Var}}(\hat{t}_p)} \right]$$

## Comparison of two groups of survival data

---

- Often one may wonder if condition A yields larger survival times than condition B.
- One (non rigorous) way is to compare the two estimated survival function  $\hat{S}_A$  and  $\hat{S}_B$ .
- The formal way to do it is within the (statistical) hypothesis testing framework

## Hypothesis testing (reminder or not)

---

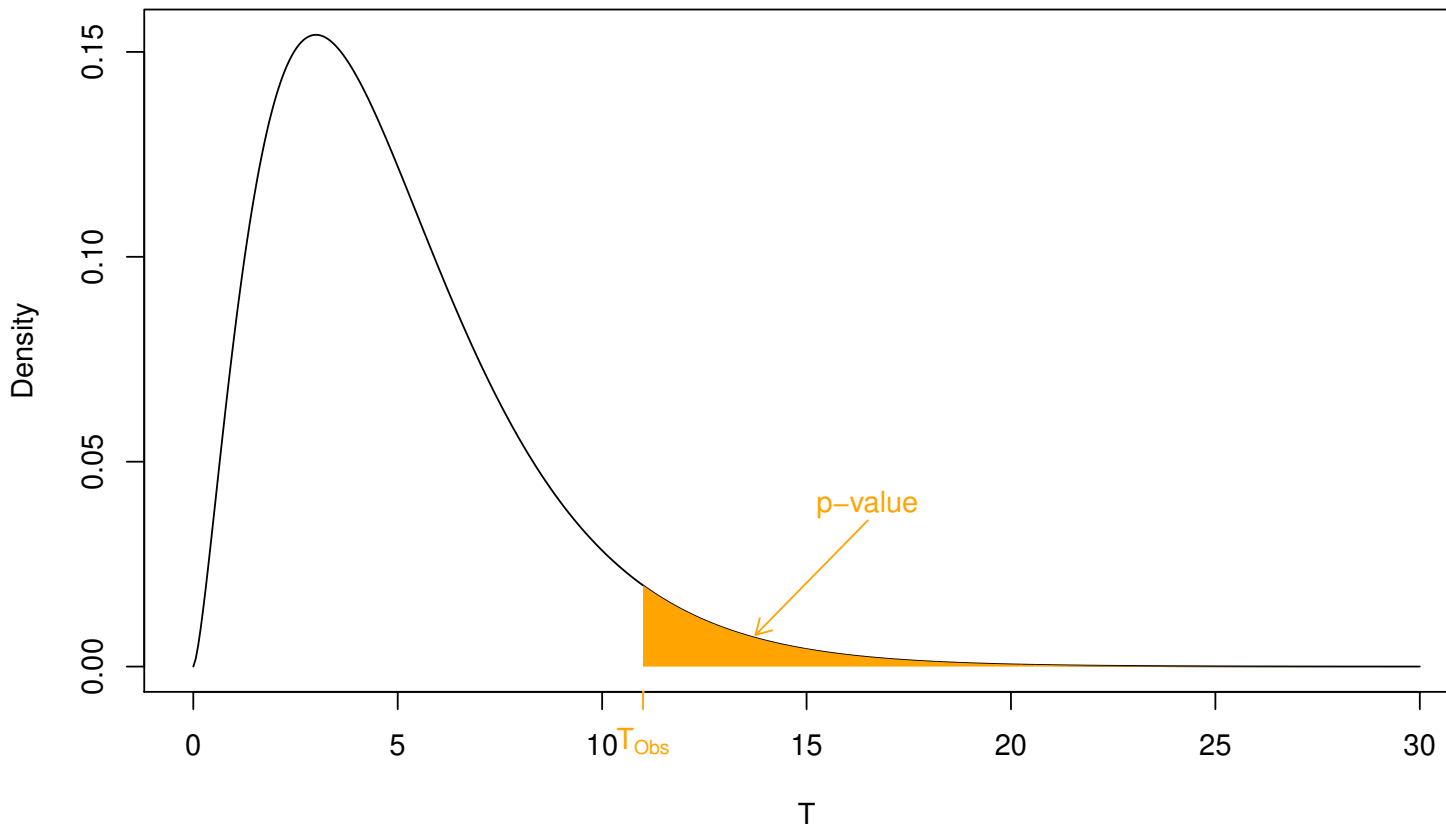
- A statistical hypothesis testing consists of 3 ingredients:
  - Hypothesis: the null hypothesis  $H_0$  and its alternative  $H_1$  (usually the complement of  $H_0$ ).
  - A test statistics  $T$  whose distribution is known under the null  $H_0$
  - A binary decision rule indicating if we
    - ▷ are not able to reject  $H_0$  in favor of  $H_1$
    - ▷ reject  $H_0$  in favor of  $H_1$ .
- The decision rule is always defined from a Type I error  $\alpha \in (0, 1)$ , e.g.,  $\alpha = 5\%, 10\%$  which corresponds to

$$\alpha = \Pr(\text{decide in favor of } H_1 \mid H_0 \text{ is true})$$

- The type I error  $\alpha$  is set by the user while the Type II error  $\beta$  is unknown

$$\beta = \Pr(\text{not able to reject } H_0 \mid H_1 \text{ is true})$$

# Plot of the decision rule



**Figure 7:** Illustration of the decision rule for a hypothesis test.

## Towards the log–rank test

- Suppose we have two conditions  $A$  and  $B$  and that we focus on a specific time event  $t_{(i)}$ . We summarize the data as follow

Group	Number of death at $t_{(i)}$	Number of surviving beyond $t_{(i)}$	Number at risk just before $t_{(i)}$
$A$	$d_{Ai}$	$n_{Ai} - d_{Ai}$	$n_{Ai}$
$B$	$d_{Bi}$	$n_{Bi} - d_{Bi}$	$n_{Bi}$
Total	$d_i$	$n_i - d_i$	$n_i$

- The hypothesis are  $H_0: S_A = S_B$  versus  $H_1: S_A \neq S_B$ .
- Now under  $H_0$ , i.e., independence on the conditions  $A$  or  $B$ ,<sup>3</sup> we have

$$d_{Ai} \sim \text{HyperGeometric}(n_i, d_i, n_{Ai}),$$

where here  $n_i$  is the population size,  $d_i$  is the number of successes within the population and  $n_{Ai}$  the number of trials (without replacement).

---

<sup>3</sup>treating the margins of the table as fixed

# The hypergeometric distribution

---

- The hypergeometric distribution is a discrete distribution that models the number of successes in successive trials without replacement.
- To be more specific, the setting is the following:
  - A finite population of size  $N$
  - Among those  $N$  units,  $K$  have a desired feature (success if drawn)
  - The number of trials  $n$ .
- We thus write  $X \sim \text{HyperGeom}(N, K, n)$  and we have

$$\Pr(X = k) = \frac{\binom{K}{k} \binom{N-K}{n-k}}{\binom{N}{n}}, \quad k \in \{0, \dots, K\},$$

which makes sense since

$$\binom{N}{n} = \binom{K}{k} = \binom{N-K}{n-k} =$$

## Towards the log–rank test (2)

---

- If  $d_{Ai} \sim \text{HyperGeometric}(n_i, d_i, n_{Ai})$ , we have

$$\mu_{Ai} := \mathbb{E}(d_{Ai}) = \frac{n_{Ai}d_i}{n_i}, \quad \sigma_i^2 := \text{Var}(d_{Ai}) = \frac{n_{Ai}n_{Bi}d_i(n_i - d_i)}{n_i^2(n_i - 1)}.$$

- Now it makes sense to sum across all time events, leading to

$$U_L = \sum_{i=1}^r (d_{Ai} - \mu_{Ai})$$

## Towards the log–rank test (2)

---

- If  $d_{Ai} \sim \text{HyperGeometric}(n_i, d_i, n_{Ai})$ , we have

$$\mu_{Ai} := \mathbb{E}(d_{Ai}) = \frac{n_{Ai}d_i}{n_i}, \quad \sigma_i^2 := \text{Var}(d_{Ai}) = \frac{n_{Ai}n_{Bi}d_i(n_i - d_i)}{n_i^2(n_i - 1)}.$$

- Now it makes sense to sum across all time events, leading to

$$U_L = \sum_{i=1}^r (d_{Ai} - \mu_{Ai}) = \sum_i (\text{Observed} - \text{Expected}),$$

where  $r$  is the number of event.

## Towards the log–rank test (2)

---

- If  $d_{Ai} \sim \text{HyperGeometric}(n_i, d_i, n_{Ai})$ , we have

$$\mu_{Ai} := \mathbb{E}(d_{Ai}) = \frac{n_{Ai}d_i}{n_i}, \quad \sigma_i^2 := \text{Var}(d_{Ai}) = \frac{n_{Ai}n_{Bi}d_i(n_i - d_i)}{n_i^2(n_i - 1)}.$$

- Now it makes sense to sum across all time events, leading to

$$U_L = \sum_{i=1}^r (d_{Ai} - \mu_{Ai}) = \sum_i (\text{Observed} - \text{Expected}),$$

where  $r$  is the number of event.

- It can be shown that  $U_L$  is approximately normal with mean 0, hence

$$T = \frac{U_L}{\sqrt{\sum_{i=1}^r \sigma_i^2}} \stackrel{\cdot}{\sim} N(0, 1) \quad \text{or equivalently} \quad T^2 = \frac{U_L^2}{\sum_{i=1}^r \sigma_i^2} \stackrel{\cdot}{\sim} \chi_1^2.$$

## The log–rank test

---

- The log–rank test set the hypothesis to

$$H_0: t \mapsto S_A(t) - S_B(t) \equiv 0 \quad \text{vs.} \quad H_1: t \mapsto S_A(t) - S_B(t) \not\equiv 0$$

- It consists in the following steps:
  - Fix the Type I error to some level  $\alpha$ ;
  - Compute the **observed** test statistics  $T_{Obs}^2 = U_L^2 / \sum_{i=1}^r \sigma_i^2$  using the data;
  - Compute the associated **p-value** =  $\Pr(\chi_1^2 > T_{Obs}^2)$ ;
  - Apply the decision rule, i.e., reject  $H_0$  in favor of  $H_1$  if  $p\text{-value} < \alpha$ .

## The log–rank test

---

- The log–rank test set the hypothesis to

$$H_0: t \mapsto S_A(t) - S_B(t) \equiv 0 \quad \text{vs.} \quad H_1: t \mapsto S_A(t) - S_B(t) \not\equiv 0$$

- It consists in the following steps:
  - Fix the Type I error to some level  $\alpha$ ;
  - Compute the **observed** test statistics  $T_{Obs}^2 = U_L^2 / \sum_{i=1}^r \sigma_i^2$  using the data;
  - Compute the associated **p-value** =  $\Pr(\chi_1^2 > T_{Obs}^2)$ ;
  - Apply the decision rule, i.e., reject  $H_0$  in favor of  $H_1$  if  $p\text{-value} < \alpha$ .

*Remark.* This test is known as the log–rank test but has several names including Mantel–Haenszel, Mantel–Cox, Peto–Mantel–Haenszel.

## The Wilcoxon test

---

- The Wilcoxon test, sometimes called Breslow test, is similar to the logrank test, i.e.,

$$H_0: t \mapsto S_A(t) - S_B(t) \equiv 0 \quad \text{vs.} \quad H_1: t \mapsto S_A(t) - S_B(t) \not\equiv 0,$$

but uses a different test statistic.

- More precisely the test statistic is now  $T = U_W^2/V_W$  where

$$U_W = \sum_{i=1}^r n_i(d_{Ai} - \mu_{Ai}), \quad V_W = \sum_{i=1}^r n_i^2 \sigma_i^2,$$

and under the null  $T \sim \chi_1^2$ .

- We keep on doing the test as for that of the log–rank.

## Log–rank vs. Wilcoxon

---

- There is a subtle difference in the test statistics of these two tests.
- To be more precise we have

$$U_L = \sum_{i=1}^r (d_{Ai} - \mu_{Ai}) \quad \text{and} \quad U_W = \sum_{i=1}^r n_i (d_{Ai} - \mu_{Ai})$$

- Hence the Wilcoxon approach puts less emphasis on the largest survival times (since in that case  $n_i$  are smaller)
- In practice we often use the **log–rank test** unless the assumption of **proportional hazards** (to be defined later on) is completely flawed.

## Generalization of Wilcoxon

---

- We can embed both log–rank and Wilcoxon test statistics using weights, i.e.,

$$U_G = \sum_{i=1}^r w(t_{(i)})(d_{Ai} - \mu_{Ai}),$$

where  $w(t_{(i)})$  are positive weights.

- Common choices for  $w(t_{(i)})$  are:
  - $w(t_{(i)}) = 1$  leading to the log–rank test
  - $w(t_{(i)}) = n_i$  leading to the Wilcoxon test
  - $w(t_{(i)}) = \hat{S}(t_{(i)})$  leading to the Peto & Peto test
  - $w(t_{(i)}) = \hat{S}(t_{(i)})^\rho$ ,  $0 \leq \rho \leq 1$ , leading to the Flemington–Harrington test

## Extension to more than 2 groups

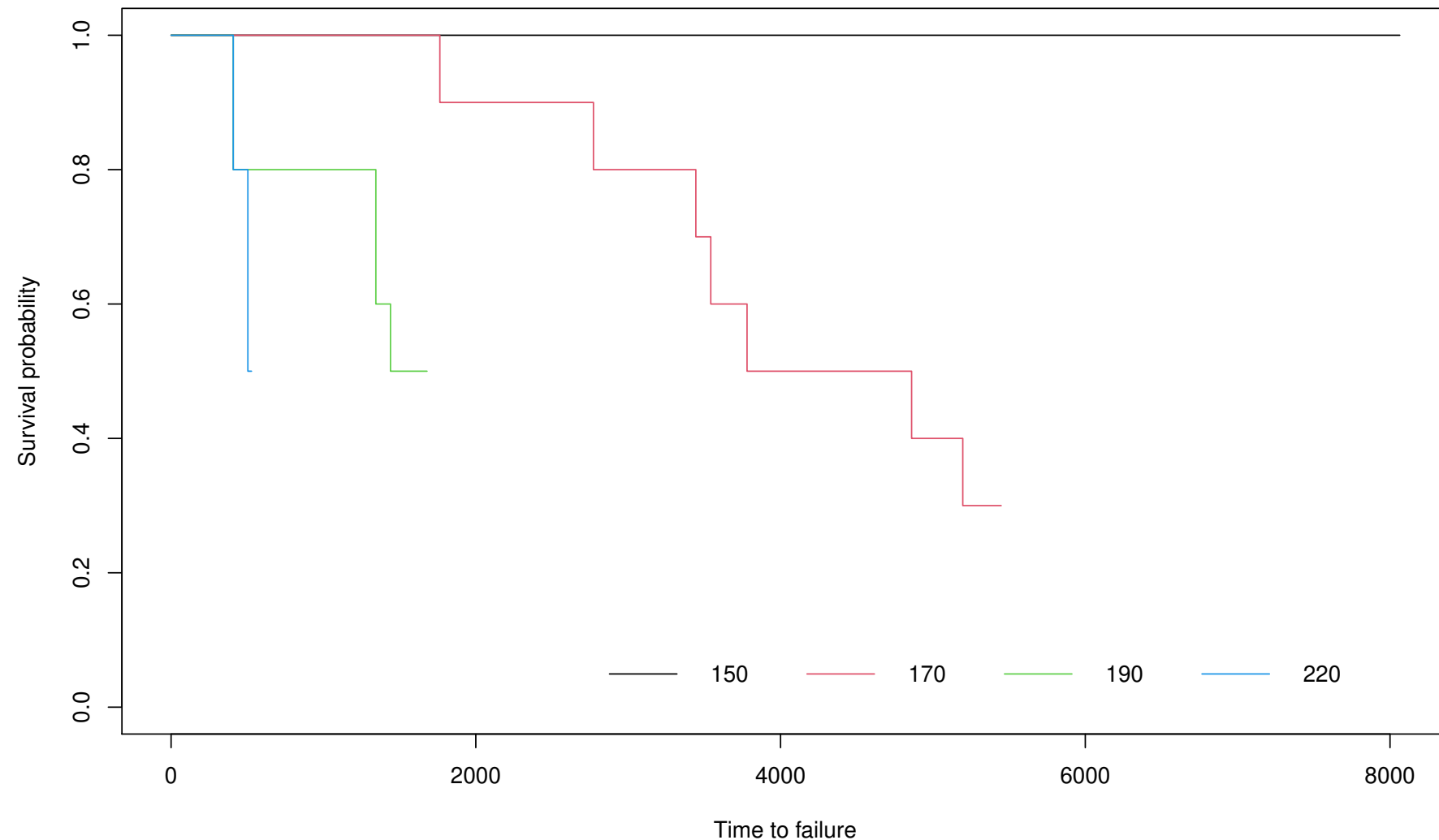
---

- In many application we may have more than 2 conditions  $A$  and  $B$
- Fortunately both the log–rank and the Wilcoxon tests extend easily to such situations.
- Suppose that we now have  $G > 2$  conditions. Similarly to what we just did, we define

$$U_{L,g} = \sum_{i=1}^r (d_{gi} - \mu_{gi}), \quad U_{W,g} = \sum_{i=1}^r n_i (d_{gi} - \mu_{gi}), \quad g = 1, \dots, G.$$

- The only difficulty is that we now have to compute covariances between  $U_{L,g}$  and  $U_{L,g'}$ . As the formula are a bit tedious, I skip them.
- All you have to know is that the test statistic satisfies under the null  $T \sim \chi_{G-1}^2$ .

# The imotor dataset



**Figure 8:** Kaplan–Meier estimate of the time to failure of motor insulation at different temperatures (*imotor* dataset).

## The imotor dataset (2)

---

```
> survdiff(Surv(time, status) ~ temp, imotor)## <-- log--rank
```

	N	Observed	Expected	(O-E)^2/E	(O-E)^2/V
temp=150	10	0	7.11	7.108	14.083
temp=170	10	7	5.76	0.269	0.431
temp=190	10	5	2.47	2.595	3.657
temp=220	10	5	1.67	6.667	9.407

Chisq= 23 on 3 degrees of freedom, p= 4e-05

```
> survdiff(Surv(time, status) ~ temp, imotor, rho = 1)## Wilcoxon
```

	N	Observed	Expected	(O-E)^2/E	(O-E)^2/V
temp=150	10	0.00	5.21	5.212054	12.03887
temp=170	10	4.38	4.44	0.000702	0.00138
temp=190	10	4.42	2.25	2.090372	3.21890
temp=220	10	4.70	1.60	6.006250	8.83022

Chisq= 19.9 on 3 degrees of freedom, p= 2e-04

## Stratified test

---

- In some cases you may have survival times depending on some features that you assume **irrelevant to the purpose of your study** but may have an **undesirable side effects on survival times**
- To annihilate possible side effects one may use **stratified tests**.
- An example of stratification is for instance if in our `imotor` dataset we have a variable indicating the location of the technical centre where was conducted the experiment. Each center is thus a **stratum**. We may not be interested in analyzing center's behavior but rather the overall failure of motor insulation.
- Shortly stratified test just compute the same statistics (log-rank or Wilcoxon) on each stratum and add them together leading (assuming independence) to get

$$T = \frac{\sum_{s \in \text{Strata}} U_{L/W,s}^2}{\sum_{s \in \text{Strata}} V_{L/W,s}} \sim \chi_1^2$$

# Lung cancer

```
> head(lung)
```

	inst	time	status	age	sex	ph.ecog	ph.karno	pat.karno	meal.cal	wt.loss
1	3	306	2	74	1	1	90	100	1175	NA
2	3	455	2	68	1	0	90	90	1225	15
3	3	1010	1	56	1	0	90	90	NA	15
4	5	210	2	57	1	1	90	60	1150	11
5	1	883	2	60	1	0	100	90	NA	0
6	12	1022	1	74	1	1	50	80	513	0

- inst: Institution code  
time: Survival time in days  
status: censoring status 1=censored, 2=dead  
age: Age in years  
sex: Male=1 Female=2  
ph.ecog: ECOG performance score as rated by the physician.  
0=asymptomatic, 1= symptomatic but completely ambulatory, 2= in bed <50% of the day, 3= in bed > 50% of the day but not bedbound, 4 = bedbound  
ph.karno: Karnofsky performance score (bad=0-good=100) rated by physician  
pat.karno: Karnofsky performance score as rated by patient  
meal.cal: Calories consumed at meals  
wt.loss: Weight loss in last six months

## Lung cancer (2)

---

```
> survdiff(Surv(time, status)~sex + strata(inst), lung)
Call:
survdiff(formula = Surv(time, status) ~ sex + strata(inst), data = lung)
```

n=227, 1 observation deleted due to missingness.

	N	Observed	Expected	$(O-E)^2/E$	$(O-E)^2/V$
sex=1	137	111	93.9	3.10	8.52
sex=2	90	53	70.1	4.16	8.52

Chisq= 8.5 on 1 degrees of freedom, p= 0.004

- What are we testing here?
- What can you conclude?

1. Preliminaries

2. Non parametric  
estimation

3. Coping with  
▷ covariates

Illustration on our  
data set

4. Time to recidivism

Conclusion

### **3. Coping with covariates**

- In many situations, you have access to additional information.
- Such “extra” variables are called covariates or features.<sup>4</sup>
- We saw some techniques on how to use such covariates, e.g., log–rank, stratification, . . .
- It was a bit limited. Let’s try to do better, e.g., which group of covariates impact most the survival?
- A widely used framework for this is known as the Cox proportional hazards model

---

<sup>4</sup>You can easily discriminate statisticians and non-statistician from these two words ;-)

## Towards Cox proportional risk model

---

- Suppose we have a standard and new process for engineering something.
- Let  $h_s(t)$  and  $h_n(t)$  be their corresponding hazard rates.
- We assume that these hazard rates are proportional, i.e., there exists a positive constant  $\psi$  such that

$$h_n(t) = \psi h_s(t), \quad t > 0.$$

- Two situations may arise:
  - $\psi < 1$  and the hazard rate for the new process is smaller than the standard one. improvement.
  - $\psi > 1$  and the hazard rate for the new process is larger than the standard one. standard process superior.

## Cox proportional risk model

---

- Suppose we have  $n$  observations  $T_1, \dots, T_n$  “tied” with some additional features  $\mathbf{x}_i = (x_{1,i}, \dots, x_{p,i})^\top$ ,  $i = 1, \dots, n$ .
- The Cox proportional risk model is defined by

$$h_i(t) = \exp(\mathbf{x}_i^\top \boldsymbol{\beta}) h_0(t),$$

where  $\boldsymbol{\beta}$  is a parameter to be estimated,  $h_i$  is the hazard rate for the  $i$ -th individuals and  $h_0$  is a **baseline hazard rate**.

- Note that there is **no intercept** since

$$\exp(\beta_0 + \mathbf{x}_i^\top \boldsymbol{\beta}) h_0(t) = \exp(\mathbf{x}_i^\top \boldsymbol{\beta}) \exp(\beta_0) h_0(t) = \exp(\mathbf{x}_i^\top \boldsymbol{\beta}) \tilde{h}_0(t),$$

i.e., intercept is included within the baseline hazard.

## Different type of features

---

- There are two main types for the features  $x_j$ ,  $j = 1, \dots, p$ :
  - variates correspond to variables which take numerical values, e.g., age, number of components.
  - factors correspond to variables which have a finite number of possible outcome called levels, e.g., sex, color.

## Features that are variates

---

- If all features are variates, then the baseline hazard  $h_0$  corresponds to the case where the variates are all 0.
- Further, since

$$\exp(x_1\beta_1 + \cdots + (x_j + 1)\beta_j + \cdots + x_p\beta_p)h_0(t) = \exp(\beta_j)h_i(t),$$

the  $\exp(\beta_j)$  quantify the impact of one unit of increase of the  $j$ -th feature in the hazard rate when all other features being fixed to the same values.

- Consequently, three cases may arise:
  - $\exp(\beta_j) = 1$  (or  $\beta_j = 0$ ) no effect
  - $\exp(\beta_j) > 1$  (or  $\beta_j > 0$ ) increase in hazard
  - $\exp(\beta_j) < 1$  (or  $\beta_j < 0$ ) decrease in hazard

## Features that are factors

---

- Consider the case of a **single feature  $x$**  that is a **factor** with  $\ell + 1$  levels  $\alpha_0, \dots, \alpha_\ell$ , e.g., brown, blue, green for eyes.
- It is often most convenient to use the **one hot encoding**, i.e., work with the new feature

$$\tilde{x} = (1_{\{x=\alpha_0\}}, \dots, 1_{\{x=\alpha_\ell\}}).$$

- To define appropriately the Cox model, one level has to be set as the **reference level**. Typically the first level is used as reference.
- Hence in this case, Cox model now writes

$$h_i(t) = \exp(\beta_1 \tilde{x}_1 + \dots + \beta_\ell \tilde{x}_\ell) h_0(t),$$

and  $\exp(\beta_j)$  quantifies how changes the hazard rate as we move from the reference level  $\alpha_0$  to level  $\alpha_j$ .

# Illustration on our dataset

---

# Interaction

---

- Suppose now that we have **two** factors  $x_1$  and  $x_2$  (with respective levels  $\alpha_j^{(1)}$  and  $\alpha_k^{(2)}$ )
- We may be tempted to use an **interaction** between these two factors, i.e., consider the impact a factor with levels  $(\alpha_i^{(1)}, \alpha_j^{(2)})_{j,k}$ .
- Some care is needed though, it is really an **interaction effect** if both factors  $x_1$  and  $x_2$  are used in the formulation of  $h_i(t)$ . We call it the **main effects** of  $x_1$  and  $x_2$ .
- If only  $x_1$  is included in the model, but not  $x_2$ , then the term  $(\alpha_j^{(1)}, \alpha_k^{(2)})$  is the **effect of  $x_2$  nested within  $x_1$**
- If both  $x_1$  and  $x_2$  are excluded from the model, then the term  $(\alpha_j^{(1)}, \alpha_k^{(2)})$  can be thought as a new factor.

# Illustration on our dataset

---

## Mixed term

---

- We may also be tempted to check the effect of the combination of a factor  $x_f$  and a variate  $x_v$ . Such a term is called a **mixed term**.
- Some care is needed in defining such mixed term, i.e., identifiability problems, but decent statistical software should manage it appropriately.

# Fitting the proportional hazards model

- The model is fit by maximizing the likelihood and Sir D. Cox showed an important result, the terms  $\exp(\beta^\top \mathbf{x})$  and  $h_0(t)$  can be fit separately.
- Hence if interest is only on the effect of features on the hazard rate, we need to maximize Cox's partial likelihood

$$L(\beta) = \prod_{\substack{i: t_{(i)} \text{ is} \\ \text{an event}}} \frac{\exp(\beta^\top x_i)}{\sum_{j \in R(t_{(i)})} \exp(\beta^\top x_j)},$$

where  $x_i$  is the vector of features for the individuals who dies at time  $t_{(i)}$  and  $R(t_{(i)})$  is the set of indices of individuals still at risk at time  $t_{(i)}$ .

- There are two interesting points in this partial likelihood:
  - First individuals that are censored do not contribute to the numerator but only to the denominator;
  - Second survival times  $t_{(i)}$  do not explicitly appear in  $L(\beta)$  and only the ranking is needed (to compute the set  $R(t_{(i)})$ ).

## Maximization of the partial likelihood

---

- As usual, we actually do not maximize the partial likelihood  $L(\beta)$  but minimize the **negative (partial) log-likelihood**

$$\hat{\beta} = \arg \min_{\beta \in \mathbb{R}^p} -\log L(\beta)$$

- There is no **closed form solution** to this problem and numerical optimization techniques are needed.
- Typically the objective function is minimized using a **Newton–Raphson algorithm** which is known to converge in few iterations (for this problem).

## Ties in the partial likelihood

---

- Cox proportional hazards model assumes that the hazard rate is **continuous** and in theory we cannot have **ties**. In practice, though, ties may occur due to rounding.
- Handling ties appropriately results in a too CPU demanding objective function, so different approximations have been suggested such as:

### Breslow

$$\tilde{L}(\boldsymbol{\beta}) = \prod_{\substack{i: t_{(i)} \text{ is} \\ \text{an event}}} \frac{\exp(\boldsymbol{\beta}^\top s_i)}{\left\{ \sum_{j \in R(t_{(i)})} \exp(\boldsymbol{\beta}^\top x_j) \right\}^{d_i}},$$

where  $s_i = \sum_{j: t_{(j)}=t_{(i)}} x_j$  and  $d_i$  is the number of death at time  $t_{(i)}$ .

## Ties in the partial likelihood

---

- Cox proportional hazards model assumes that the hazard rate is **continuous** and in theory we cannot have **ties**. In practice, though, ties may occur due to rounding.
- Handling ties appropriately results in a too CPU demanding objective function, so different approximations have been suggested such as:

### Efron

$$\tilde{L}(\boldsymbol{\beta}) = \prod_{\substack{i: t_{(i)} \text{ is} \\ \text{an event}}} \frac{\exp(\boldsymbol{\beta}^\top s_i)}{\prod_{k=1}^{d_i} \left\{ \sum_{j \in R(t_{(i)})} \exp(\boldsymbol{\beta}^\top x_j) - (k-1)d_i^{-1} \sum_{\ell \in D(t_{(i)})} \exp(\boldsymbol{\beta}^\top x_\ell) \right\}},$$

where  $D(t_{(i)})$  is the set of indices of individuals who died at time  $t_{(i)}$ .

## Ties in the partial likelihood

---

- Cox proportional hazards model assumes that the hazard rate is continuous and in theory we cannot have ties. In practice, though, ties may occur due to rounding.
- Handling ties appropriately results in a too CPU demanding objective function, so different approximations have been suggested such as:

**Cox**

$$\tilde{L}(\beta) = \prod_{i: t_{(i)} \text{ is an event}} \frac{\exp(\beta^\top s_i)}{\sum_{j \in R(t_{(i)}; d_i)} \exp(\beta^\top x_j)},$$

where  $R(t_{(i)}; d_i)$  is the set of all possible draws of  $d_i$  individuals without replacement among  $R(t_{(i)})$ .

# Model selection

---

- Before starting what we are about to learn is **not** specific to Cox's model but **can be applied as long as you work with the maximum likelihood estimator.**
- **Model selection** consists in selecting the best model among  $N$  competitive fitted models  $M_1, \dots, M_N$ .
- To decide which model performs bests, we need a **metric** to be able to rank each of them.
- There are three main roads for this:
  - **validation methods**, e.g., cross-validation, data set splitting (train / validation / test)
  - **Hypothesis testings**, e.g., Wald test, likelihood ratio
  - **Information criteria**, e.g., Akaike / Bayesian Information Criterion
- Typically we won't have huge dataset so I will focus on the last two only<sup>5</sup>

---

<sup>5</sup>You can find details on validation methods in other lecture I give ;-)

## Asymptotic of the maximum likelihood estimator

- We start by recalling the very desirable asymptotic property<sup>6</sup> of the maximum likelihood estimator  $\hat{\theta}$

$$\sqrt{n}(\hat{\theta} - \theta_0) \xrightarrow{d.} N\left\{0, -H(\theta_0)^{-1}\right\}, \quad n \rightarrow \infty,$$

where  $H(\theta_0) = \mathbb{E}\{\nabla^2 \log f(X; \theta_0)\}$  where  $X$  is a single random variable from the true model  $f(\cdot; \theta_0)$ .

- Stating it less formally, we can argue that for  $n$  large enough

$$\hat{\theta} \stackrel{\text{d.}}{\sim} N(\theta_0, \Sigma),$$

with  $\Sigma = \left(-\nabla^2 \ell(\hat{\theta})\right)^{-1}$ , i.e., the inverse of the Hessian matrix of the negative log-likelihood at its minimum.

---

<sup>6</sup>If regularity conditions hold...

## Wald test

---

- The Wald test tests if a given component of the true parameter  $\theta$ , say the  $j$ -th, is 0 or not, i.e.,

$$H_0: \theta_j = 0 \quad \text{vs.} \quad H_1: \theta_j \neq 0$$

- The test statistic satisfies under the null  $H_0$

$$T_W = \frac{\hat{\theta}_j}{\text{Standard error}(\hat{\theta}_j)} \xrightarrow{\text{d.}} N(0, 1), \quad n \rightarrow \infty.$$

- We then take the decision rule as usual.

# Illustration on our data set

---

## Likelihood ratio test

---

- The likelihood ratio test check whether it is preferable to opt for the nested model or the more general model.
- Recall that a model  $M_A$  is said to be nested within model  $M_B$  if  $M_A$  is a special case of  $M_B$ , e.g., the exponential distribution is nested within the Gamma family.
- The test is formalized as follows

$$H_0 : \text{nested model is correct} \quad \text{vs.} \quad H_1 : \text{general model is correct}$$

- The test statistic satisfies under the null  $H_0$

$$T_{LR} = -2 \left\{ \ell(\hat{\theta}_{\text{gen}}) - \ell(\hat{\theta}_{\text{nest}}) \right\} \xrightarrow{\text{d.}} \chi_p^2, \quad n \rightarrow \infty,$$

where  $\hat{\theta}_{\text{gen}}$  and  $\hat{\theta}_{\text{nest}}$  are the maximum likelihood estimators for the general and nested models respectively and  $p = \dim(\theta_{\text{gen}}) - \dim(\theta_{\text{nest}})$ .

# Illustration on our data set

---

## Wald or likelihood ratio?

---

- The Wald and likelihood ratio tests share the same goal.
- They can be used for model selection or to test whether a feature is statistically relevant.
- They usually point to the same answer
- However for small sample size, e.g.,  $n \leq 30$ , Wald is known to perform poorly.
- It is thus often recommended to use the likelihood ratio test.

# Information criterion

---

- Likelihood ratio / Wald tests are useful for **nested models**
- If we have **non nested models** we can use the following information criterion:

**Akaïke**  $AIC(M) = -2\ell(\hat{\theta}_M) + 2 \dim(\hat{\theta}_M)$

**Schwarz**  $BIC(M) = -2\ell(\hat{\theta}_M) + \dim(\hat{\theta}_M) \log n,$

where  $\hat{\theta}_M$  is the maximum likelihood estimator for model  $M$ .

- Both information criterion is a tradeoff between **model's goodness of fit** and **model's complexity**
- Since  $\log n > 2$  when  $n \geq 8$ ,  $BIC$  selects simpler model than  $AIC$ . Further, model selection using  $AIC$  is **inconsistent**.
- It is often sensible to rely on  $BIC$

# Information criterion

---

- Likelihood ratio / Wald tests are useful for **nested models**
- If we have **non nested models** we can use the following information criterion:

**Akaïke**  $AIC(M) = -2\ell(\hat{\theta}_M) + 2 \dim(\hat{\theta}_M)$

**Schwarz**  $BIC(M) = -2\ell(\hat{\theta}_M) + \dim(\hat{\theta}_M) \log n,$

where  $\hat{\theta}_M$  is the maximum likelihood estimator for model  $M$ .

- Both information criterion is a tradeoff between **model's goodness of fit** and **model's complexity**
  - Since  $\log n > 2$  when  $n \geq 8$ ,  $BIC$  selects simpler model than  $AIC$ . Further, model selection using  $AIC$  is **inconsistent**.
  - It is often sensible to rely on  $BIC$
- ☞ In practice we compute  $AIC$  and  $BIC$  for various models and select the one with the smallest information criterion value.

# Model checking

---

- So far we learn the basic steps for the statistical modelling of survival data, i.e.,
  - a general framework, i.e., Cox's proportional hazards model
  - an estimator, i.e., the maximum (partial) likelihood estimator
  - how to get the best model from several competitive models, i.e., likelihood ratio, BIC
- However...

# Model checking

- So far we learn the basic steps for the statistical modelling of survival data, i.e.,
  - a general framework, i.e., Cox's proportional hazards model
  - an estimator, i.e., the maximum (partial) likelihood estimator
  - how to get the best model from several competitive models, i.e., likelihood ratio, BIC
- However...



# Model checking

- So far we learn the basic steps for the statistical modelling of survival data, i.e.,
  - a general framework, i.e., Cox's proportional hazards model
  - an estimator, i.e., the maximum (partial) likelihood estimator
  - how to get the best model from several competitive models, i.e., likelihood ratio, BIC
- However... you still have a poor model



# Model checking

- So far we learn the basic steps for the statistical modelling of survival data, i.e.,
  - a general framework, i.e., Cox's proportional hazards model
  - an estimator, i.e., the maximum (partial) likelihood estimator
  - how to get the best model from several competitive models, i.e., likelihood ratio, BIC
- However... you still have a poor model



👉 We need to check if the best model is actually good!

## Evaluating model adequacy

---

- Model adequacy is most often based on graphical investigation where we compare **what we observed** to **what is expected** from the fitted model: a good match indicates that the model is sensible.
- One way to do it is by analyzing **residuals**. Roughly speaking, a generic definition is

$$\text{residuals}_i = \text{Observation}_i - \text{Prediction}_i$$

- For instance, in the linear model  $Y = X^\top \beta + \varepsilon$ , we have

$$\text{residuals}_i = Y_i - X_i^\top \hat{\beta},$$

- and we usually compare those residuals to a centered Gaussian distribution.
- For the Cox's proportional model, it is a bit more tricky (in part due to censoring) and a first attempt was to analyze **Cox–Snell residuals**.

## Cox–Snell residuals

---

- The motivation on the use of Cox–Snell residuals is based on the fact that

$$T_* \text{ has cumulative hazard } H \implies H(T_*) \sim \text{Exp}(1).$$

- However since some observation may be right censored we also use the fact that

$$\mathbb{E}(H(T_*) \mid H(T_*) \geq H(C)) = H(C) + 1.$$

- Having fitted a Cox's model, it is thus sensible to define the Cox–Snell residuals as follows

$$r_{CS,i} = \begin{cases} \exp(\hat{\beta}x_i)\hat{H}_O(T_i), & \text{if event} \\ \exp(\hat{\beta}x_i)\hat{H}_O(T_i) + 1, & \text{if right censored,} \end{cases}$$

and to compare it to an  $\text{Exp}(1)$  using a QQ–plot.

- Unfortunately, such a procedure is not very efficient in detecting departure from the fitted model.
- This is one reason they are not even implemented within the `survival` package
- To bypass this hurdle, different type of residuals have been suggested:
  - the **martingale** residuals
  - the **deviance** residuals
  - the **Schoenfeld** residuals
- We will cover these different types in turn.

## The martingale residuals

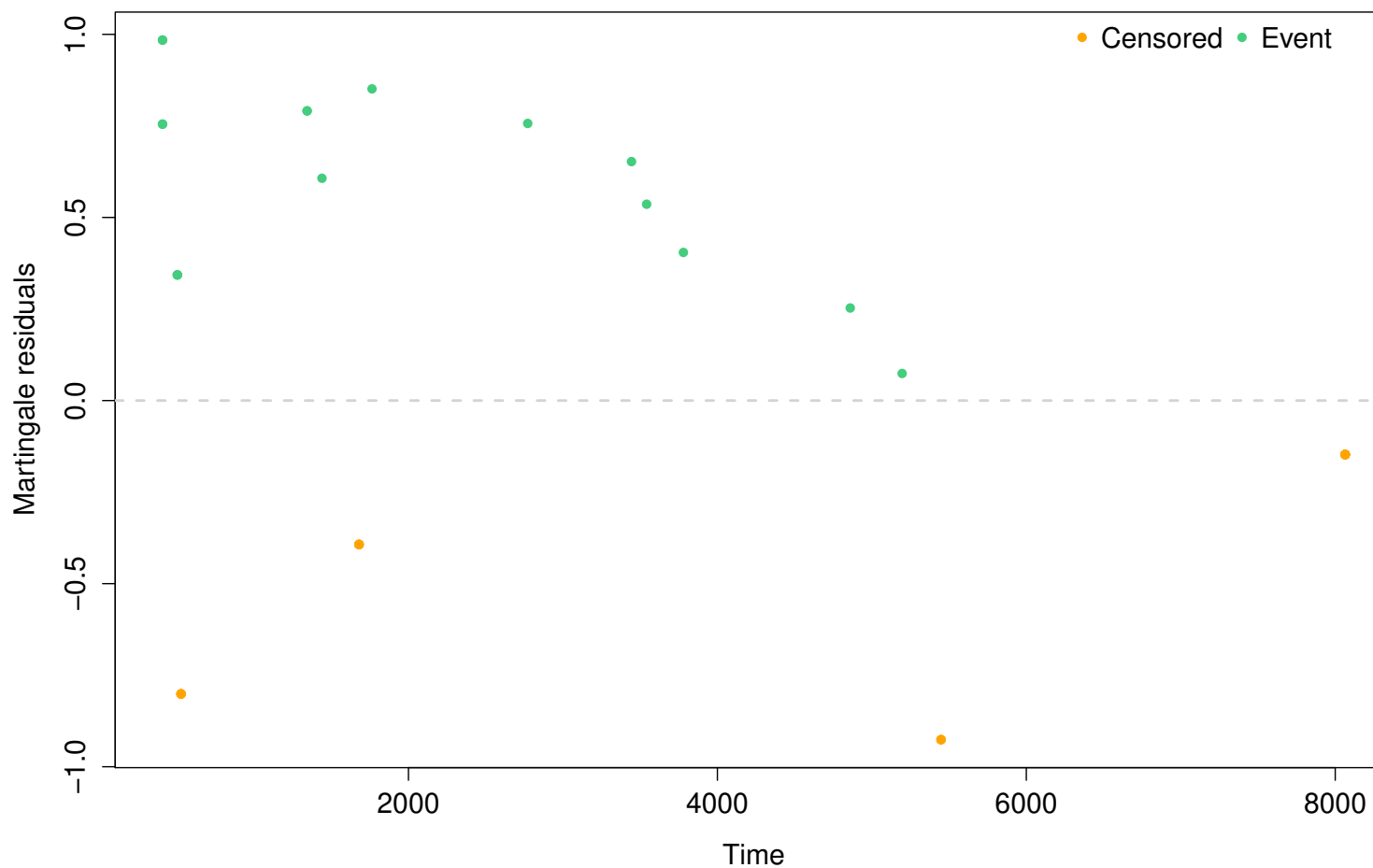
---

- The **martingale residuals** are defined by

$$\begin{aligned} r_{M,i} &= \delta_i - r_{CS,i} \\ &= \text{observed} - \text{model prediction}, \end{aligned}$$

where  $\delta_i = 1$  if  $T_i$  is an event and 0 otherwise.

- As a consequence,
  - positive values indicate that the event actually occur sooner than what the model says
  - negative values indicate that the event actually occur later than what the model says (or censored)
- In particular one can identify **which individuals** is poorly fit



**Figure 9:** Martingale residuals for the *imotor* data set.

## Properties of martingale residuals

---

- The martingale residuals have nice properties:
  - $\mathbb{E}(R_{M,i}) = 0$
  - $\text{Cov}(R_{M,i}, R_{M,j}) = 0$ , for  $i \neq j$
  - $\sum_{i=0}^n r_{M,i} = 0$
- but also undesirable ones:
  - skewed since  $R_{M,i} \in (-\infty, 1]$
  - always negative for censored observation, i.e., when  $\delta_i = 0$ .

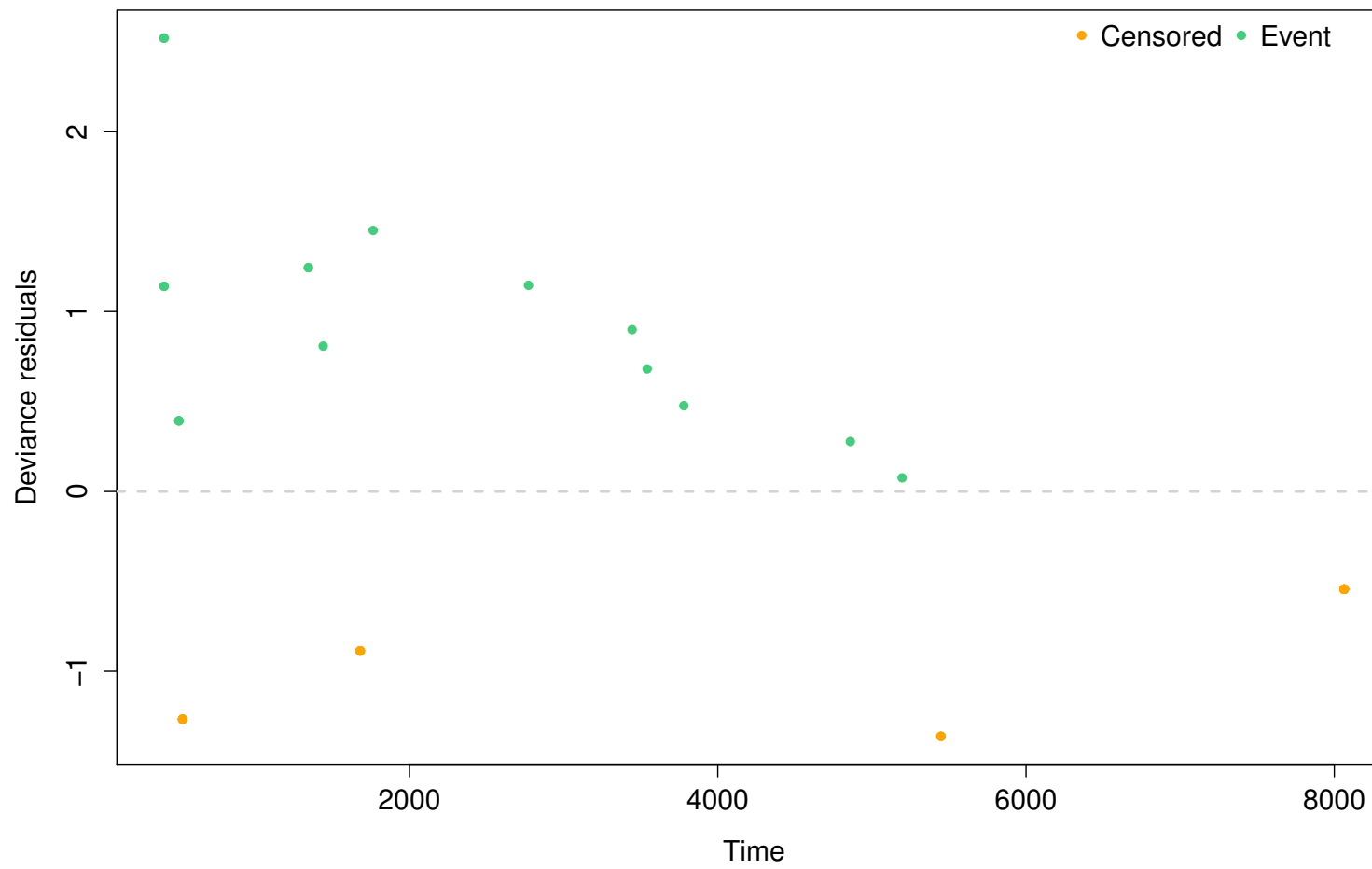
## The deviance residuals

---

- The deviance residuals can be thought as a **symmetrization** of the martingale residuals.
- They are defined by

$$r_{D,i} = \text{sign}(r_{M,i}) \sqrt{-2 \{ r_{M,i} + \delta_i \log(\delta_i - r_{M,i}) \}},$$

and, by construction, should be approximately symmetric around 0.



**Figure 10:** Deviance residuals for the *imotor* data set.

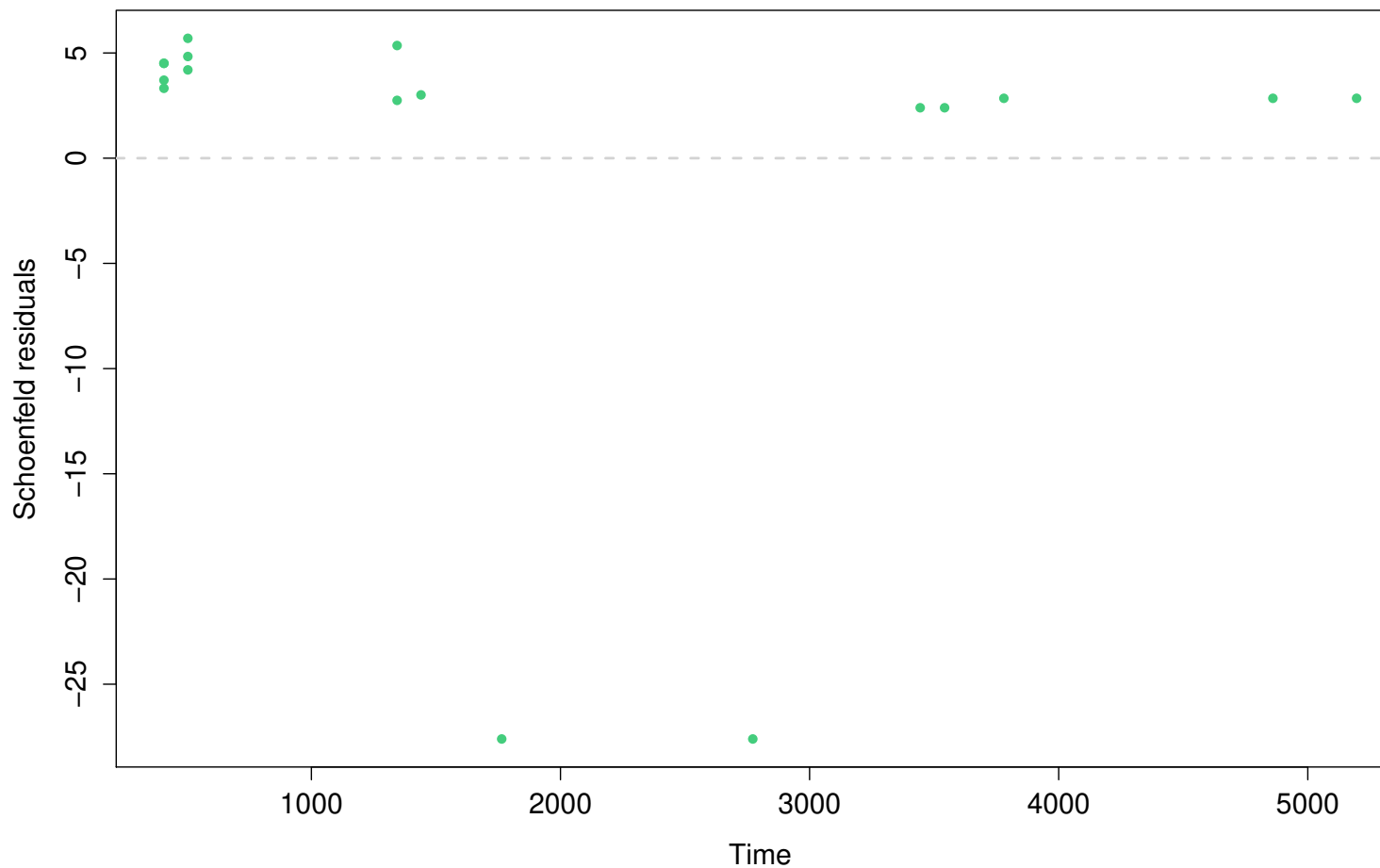
## Schoenfeld residuals

---

- The Schoenfeld residuals differ significantly from the above residuals as, for a fixed individual, they lead to one value for each feature  $x_1, \dots, x_p$ .
- They are given by

$$r_{S,ij} = \delta_i(x_{ij} - \hat{a}_{i,j}), \quad \hat{a}_{i,j} = \frac{\sum_{\ell: T_\ell \geq T_i} x_{\ell j} \exp(\hat{\beta}^\top \mathbf{x}_\ell)}{\sum_{\ell: T_\ell \geq T_i} \exp(\hat{\beta}^\top \mathbf{x}_\ell)}$$

- Clearly  $r_{S,ij} = 0$  for censored observations (since  $\delta_i = 0$ ) and it is common to report values only for events.
- Now if the last observation is an event then  $\hat{a}_{i,j} = x_{ij} \Rightarrow r_{S,ij} = 0$ .
- We also have the desirable properties:
  - $\mathbb{E}(R_{S,ij}) = 0$
  - $\sum_{i=0}^n r_{S,ij} = 0$



**Figure 11:** Schoenfeld residuals for the *imotor* data set (for our single feature *temp*).

## Take home message about residuals

---

- Graphical inspection of residuals is an important stage to identify
  1. possible **outliers**, i.e., largest residuals in absolute values, and to understand why such a departure.
  2. possible useful **transformation of the features**, i.e., taking  $\log temp$  rather than  $temp$ .
- **Objective 1** is usually done by plotting residuals w.r.t. individual's index or time
- **Objective 2** is usually done by plotting residuals w.r.t. to a, possibly new, covariate.

## Testing proportional hazards

---

- In the last slides we talk about the Cox's proportional hazards model which, as its name suggests, assumes that hazards are proportional, i.e.,

$$\frac{h_i(t)}{h_j(t)} = \frac{\exp(x_i^\top \beta)}{\exp(x_j^\top \beta)}, \quad i \neq j, \quad t > 0.$$

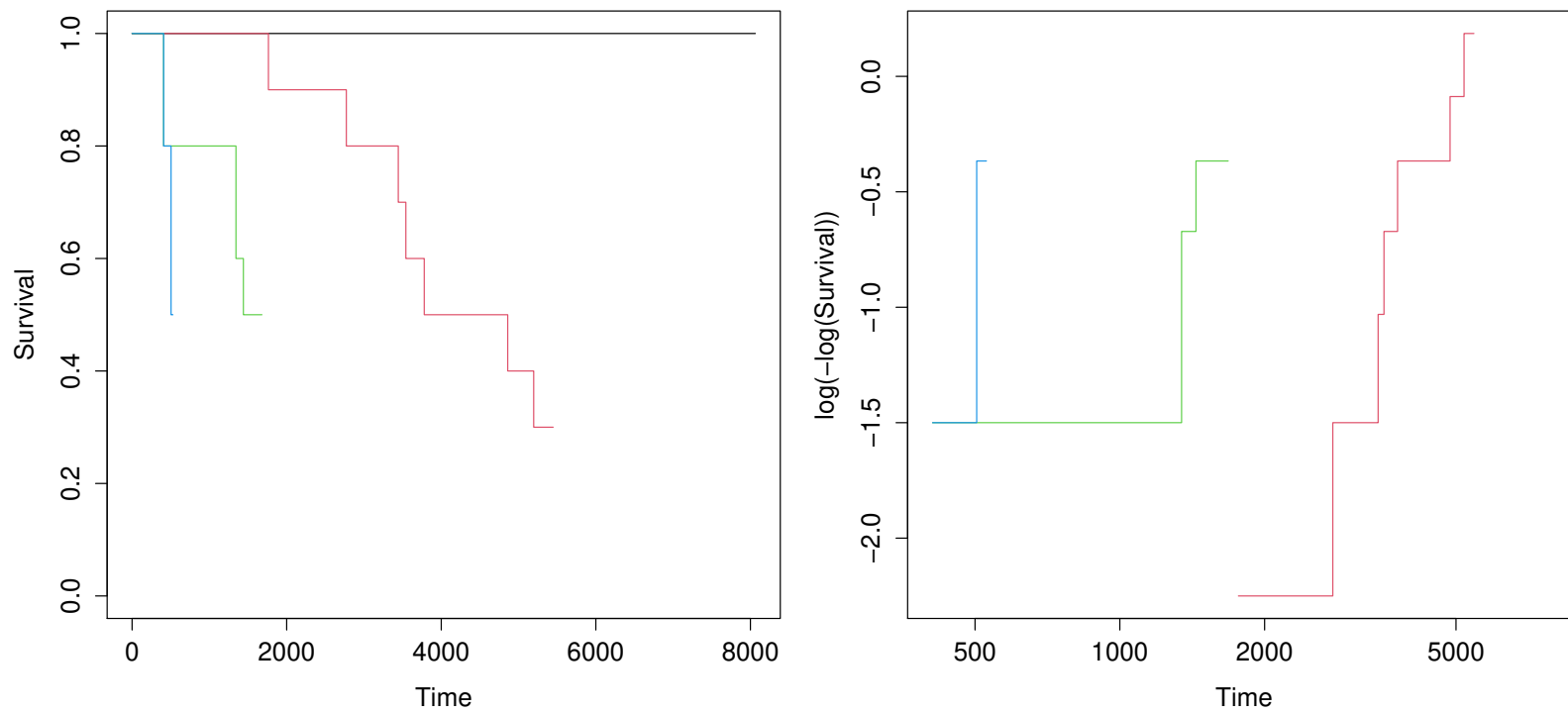
- What if this assumption is not supported by the data? Check!
- Since  $S(t) = \exp\{-\int_0^t h(u)du\}$ , we thus have

$$S_i(t) = \exp\{-\exp(x_i^\top \beta)H_0(t)\} = S_0(t)^{\exp(x_i^\top \beta)},$$

i.e., survival curves should not cross or, equivalently,

$$\log\{-\log S_i(t)\} = x_i^\top \beta + \log H_0(t),$$

i.e., the log-log survivorships should be parallel.



**Figure 12:** Graphical assessment of the validity of the proportional hazards assumption on the *imotor* dataset.

## Schoenfeld residuals are back

---

- Consider an extended Cox's model with time-dependent coefficients, i.e.,

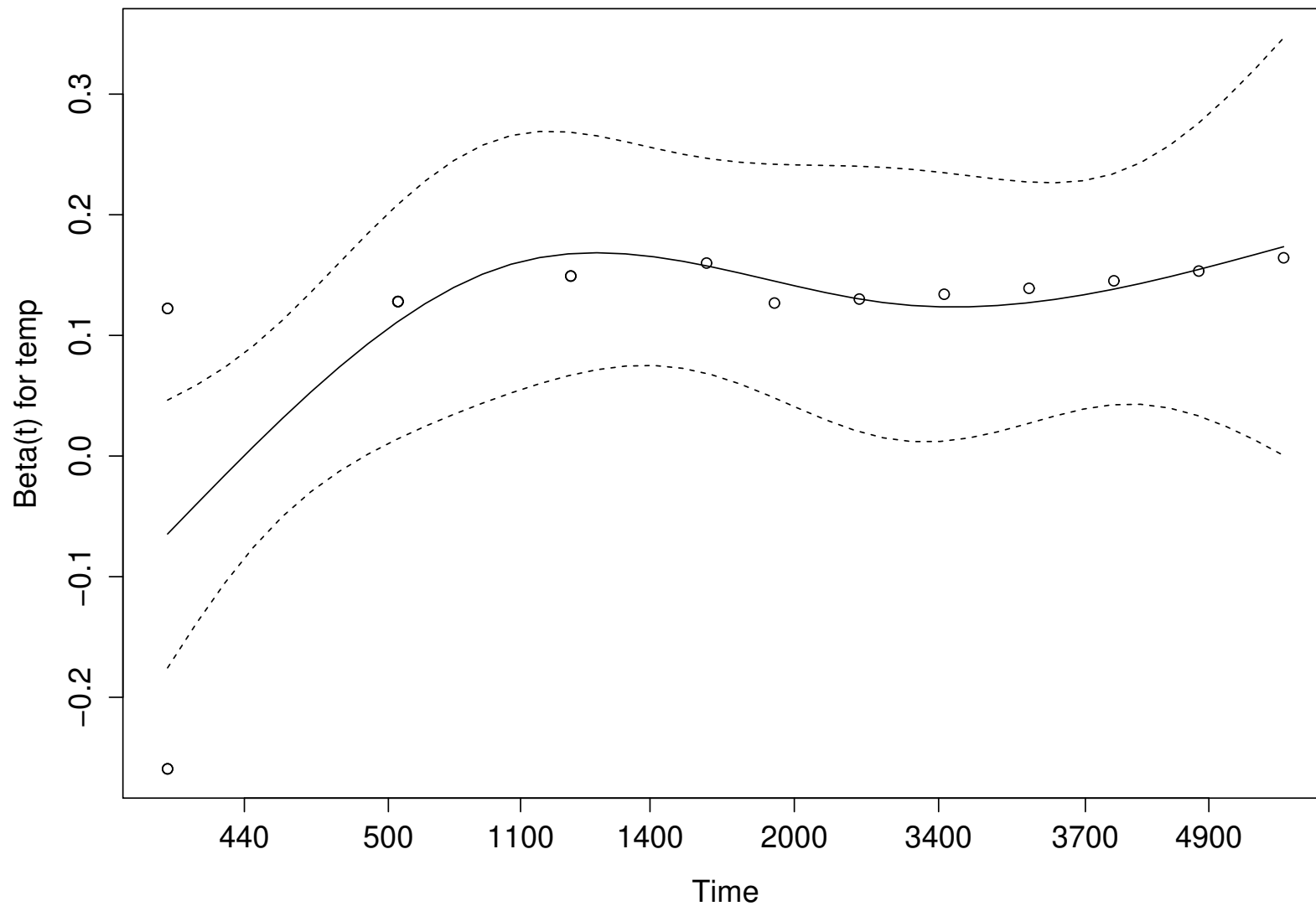
$$h_i(t) = h_0(t) \exp \left\{ x_i^\top \beta(t) \right\}.$$

- With such a model, the influence of a given feature, say  $x_j > 0$ , may vary with time: if  $\beta_j$  decreases with  $t$ , then  $x_j$  has less and less influence.
- Proportional hazards assumption corresponds to the case where  $\beta(t) \equiv \beta$ .
- It can be shown that

$$\beta_j(T_i) \approx \hat{\beta}_j + r_{S,ij},$$

where  $\hat{\beta}$  is the usual estimator of the Cox's model and  $r_{S,ij}$  are the (scaled) Schoenfeld residuals.

- The above equation suggests to plot  $\hat{\beta}_j + r_{S,ij}$  versus time (or a some function of time  $g(t)$ ) and see whether it is constant or not



**Figure 13:** Graphical assessment of the validity of the proportional hazards assumption on the imotor dataset.

1. Preliminaries

2. Non parametric  
estimation

3. Coping with  
covariates

4. Time to  
recidivism

Conclusion

## **4. Time to recidivism**

## Time to recidivism (Rossi data set)

---

- Recidivism is the event re-incarceration after release from prison
- A randomized study with 52 weeks of follow-up collected information of the following variables:

`fin` financial support vs. no financial support after release

`week` Time in week to either re-arrest or censoring

`arrest` 1 = arrest during the follow-up, 0 = no arrest

`age` Age (years) at the time of release

`race` A factor with levels black and other

`wexp` A factor with levels yes/no if work experience prior to incarceration

`mar` A factor with levels (not) married if (not) married at the time of release

`paro` A factor with levels yes/no if released on parole

`prio` Number of prior conviction

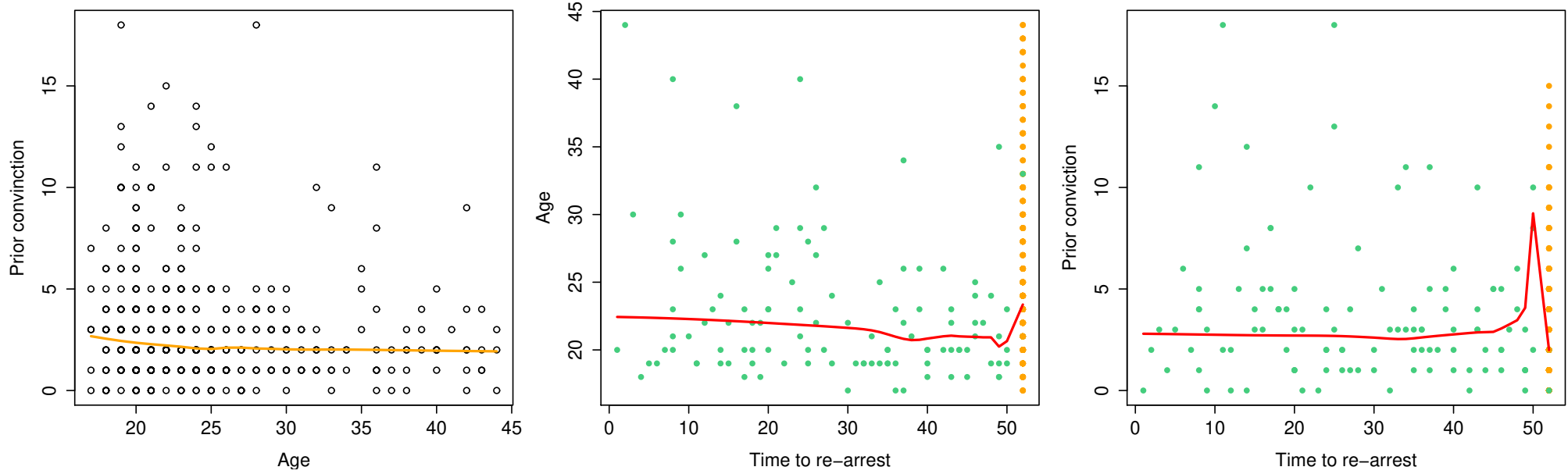
`educ` Categorical variable coded numerically with codes 2 (grade  $\leq 6$ ), 3 (grades 6–9), 4 (grades 10–11), 5 (grade 12) or 6 (post-secondary)

```

> head(data)
  week arrest fin age race wexp      mar paro prio educ
1 20     1 no 27 black no not married yes   3   3
2 17     1 no 18 black no not married yes   8   4
3 25     1 no 19 other yes not married yes 13   3
4 52     0 yes 23 black yes married yes   1   5
5 52     0 no 19 other yes not married yes   3   3
6 52     0 no 24 black yes not married no    2   4

> attach(data)
> table(fin)                                > table(paro)
no yes
216 216
> table(race)                               no yes
black other
379 53
> table(wexp)                                165 267
no yes
185 247
> table(mar)                                 > table(educ)
married not married
53     379
                                         2   3   4   5   6
                                         24 239 119 39 11
                                         > summary(age)
                                         Min. 1st Qu. Median Mean 3rd Qu. Max.
                                         17.0 20.0 23.0 24.6 27.0 44.0
                                         > summary(prio)
                                         Min. 1st Qu. Median Mean 3rd Qu. Max.
                                         0.000 1.000 2.000 2.984 4.000 18.000

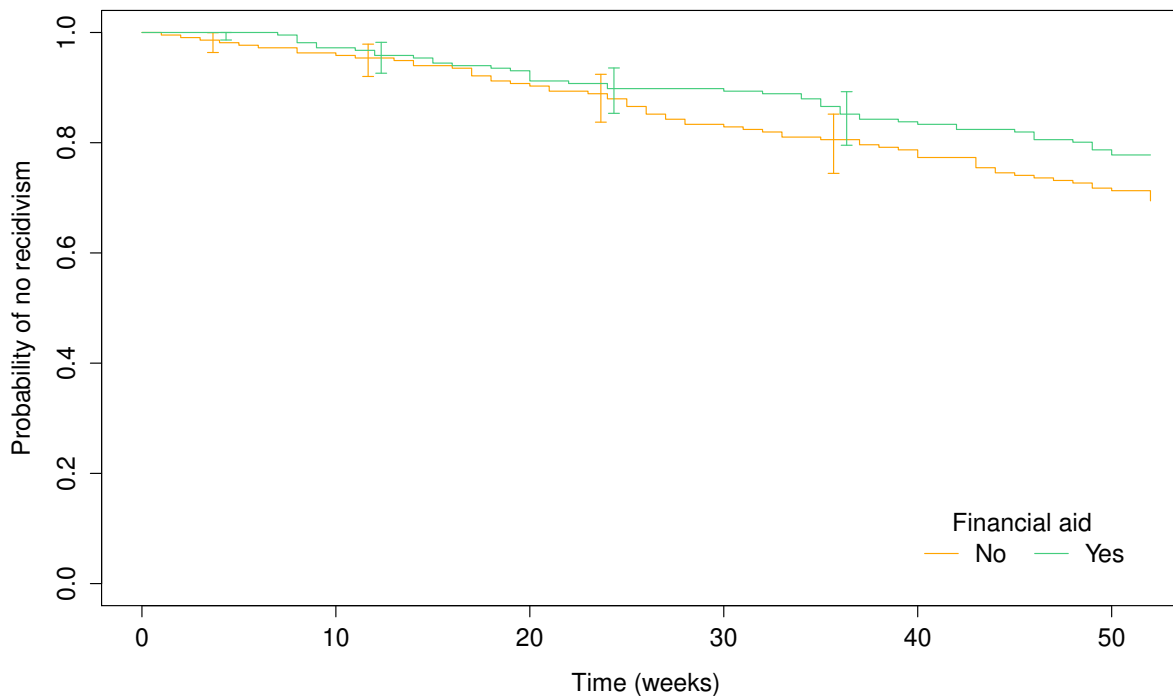
```



**Figure 14:** Some exploratory plots for the Rossi dataset.

- No clear trend of features w.r.t. to time to recidivism.

# Probability to no recidivism



**Figure 15:** Kaplan–Meier estimates according to the financial support status.

- The curve corresponding to no financial support is always below, i.e., **recidivism appears to occur sooner w/o financial aid**
- This point has to be **mitigated** as confidence intervals overlap.
- Further other features may be related to this behavior

# Is there an effect of financial support?

---

```
> survdiff(Surv(week,arrest)~fin, data = data)
```

	N	Observed	Expected	$(O-E)^2/E$	$(O-E)^2/V$
fin=no	216	66	55.6	1.96	3.84
fin=yes	216	48	58.4	1.86	3.84

Chisq= 3.8 on 1 degrees of freedom, p= 0.05

```
> survdiff(Surv(week,arrest)~fin, data = data, rho = 1)
```

	N	Observed	Expected	$(O-E)^2/E$	$(O-E)^2/V$
fin=no	216	57.5	48.5	1.68	3.75
fin=yes	216	41.8	50.9	1.60	3.75

Chisq= 3.7 on 1 degrees of freedom, p= 0.05

- The  $p$ -value is unfortunately on the boundary of the decision rule for a Type I error of  $\alpha = 5\%$ , i.e., no clear cut decision if an effect.
- stratified versions of the test** (not shown) gives the same results ( $p$ -values ranging from 0.04 to 0.05).

## Cox's regression

```
> (fit <- coxph(Surv(week,arrest) ~ fin + age + race + wexp + mar +  
paro + prio + factor(educ), data = data))
```

	coef	exp(coef)	se(coef)	z	p
finyes	-0.40265	0.66855	0.19295	-2.087	0.03691
age	-0.05141	0.94989	0.02220	-2.316	0.02055
raceother	-0.36151	0.69663	0.31216	-1.158	0.24683
wexpyes	-0.12002	0.88690	0.21346	-0.562	0.57393
marnot married	0.42362	1.52749	0.38216	1.108	0.26765
paroyes	-0.09822	0.90645	0.19587	-0.501	0.61604
prio	0.07944	1.08268	0.02935	2.707	0.00679
factor(educ)3	0.59335	1.81004	0.51961	1.142	0.25349
factor(educ)4	0.32844	1.38880	0.54368	0.604	0.54577
factor(educ)5	-0.12098	0.88605	0.67519	-0.179	0.85780
factor(educ)6	-0.40696	0.66567	1.12333	-0.362	0.71714

Likelihood ratio test=38.68 on 11 df, p=6.013e-05

n= 432, number of events= 114

- Note the use of `factor` for feature `educ`!
- According to Wald's test, some features may be dropped
- We need to perform model selection

## One example of likelihood ratio test

---

- According to the previous output, Wald's test suggest to drop feature paro
- Is it consistent with likelihood ratio test?

```
> fit2 <- update(fit, . ~ . - paro)
> anova(fit, fit2)
Analysis of Deviance Table
Cox model: response is Surv(week, arrest)
Model 1: ~ fin + age + race + wexp + mar + paro + prio + factor(educ)
Model 2: ~ fin + age + race + wexp + mar + prio + factor(educ)
loglik  Chisq Df P(>|Chi|)
1 -656.04
2 -656.17 0.2499  1      0.6171
```

- We have a  $p$ -value very close to that of the Wald's test (as often)
- Conclusion is we can drop feature paro

## The case of factor with more than 2 levels

---

- Some care is needed when deciding if one should drop educ or not
- Basically there are two main paths:
  - either we completely drop the feature using likelihood ratio test
  - or we try to merge some levels as shown below

```
> educ2 <- factor(educ)
> levels(educ2) <- list('2-3' = 2:3, '4-6' = 4:6)
> table(educ2)
educ2
2-3 4-6
263 169
> fit3 <- update(fit2, . ~ . - factor(educ) + factor(educ2))
> AIC(fit2, fit3)##these are *not* nested so use AIC not LRT
      df      AIC
fit2 10 1332.335
fit3  7 1329.282
```

- Here the drop in AIC is about 3 which (very!) slightly better
- Anyway, the feature educ is not significant (not shown)

- After a stage of (careful) model selection, we finally end up with

```
> summary(bestFit)
Call:
coxph(formula = Surv(week, arrest) ~ fin + age + prio, data = data)

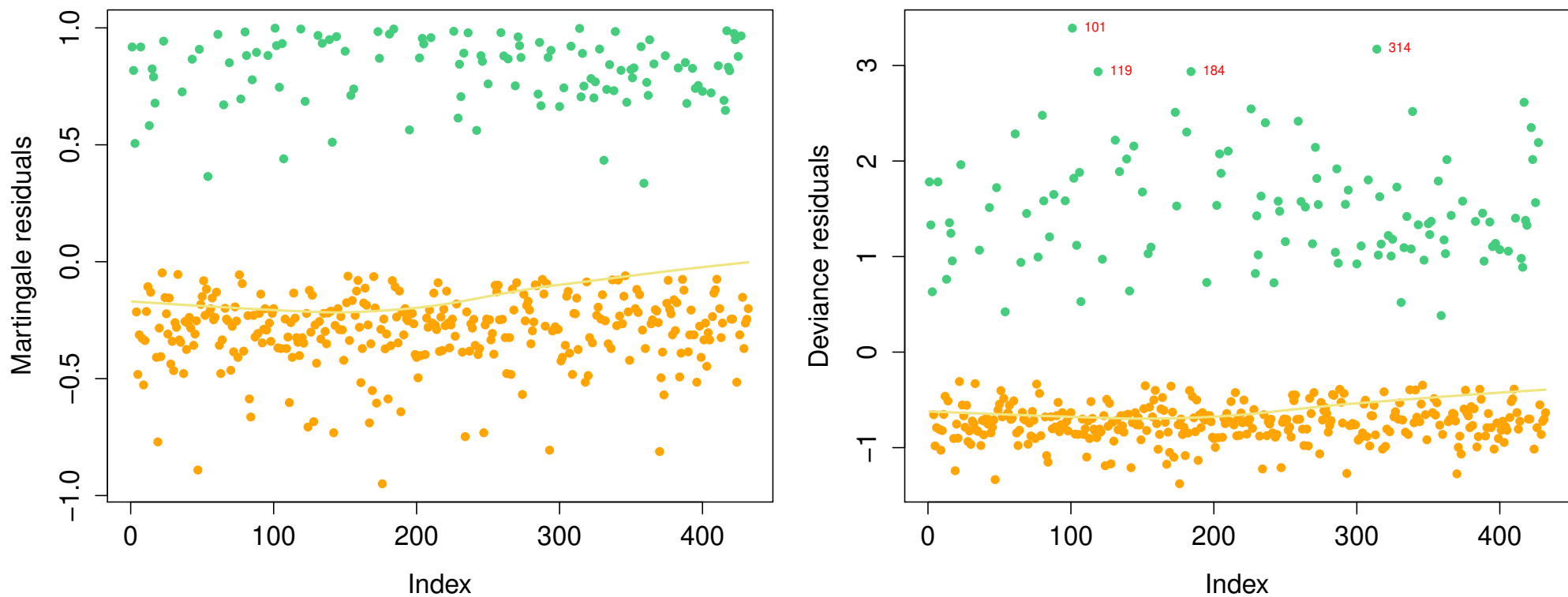
n= 432, number of events= 114

            coef exp(coef) se(coef)      z Pr(>|z|)
finyes -0.34695   0.70684  0.19025 -1.824 0.068197 .
age     -0.06711   0.93510  0.02085 -3.218 0.001289 **
prio    0.09689   1.10174  0.02725  3.555 0.000378 ***
---
Signif. codes:  0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1

            exp(coef) exp(-coef) lower .95 upper .95
finyes     0.7068      1.4148    0.4868    1.0263
age        0.9351      1.0694    0.8977    0.9741
prio       1.1017      0.9077    1.0444    1.1622

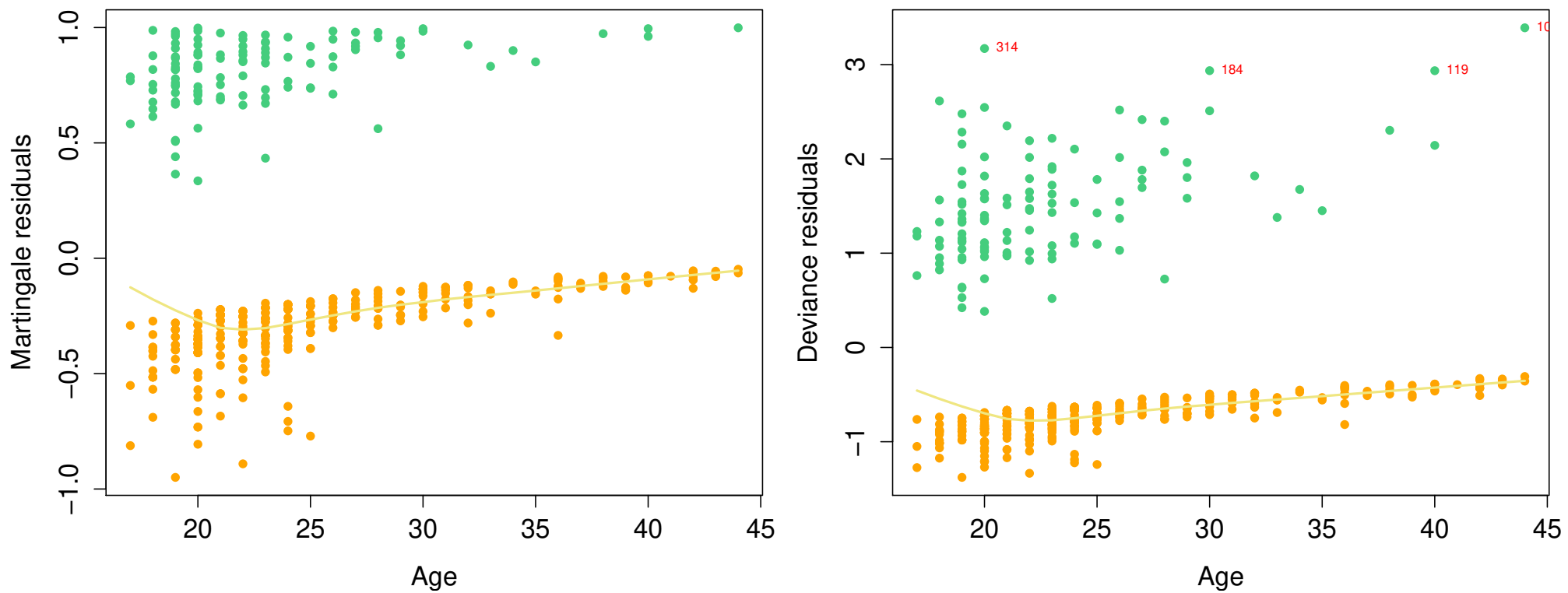
Concordance= 0.63  (se = 0.027 )
Likelihood ratio test= 29.05  on 3 df,  p=2e-06
Wald test             = 27.94  on 3 df,  p=4e-06
Score (logrank) test = 29.03  on 3 df,  p=2e-06
```

# Residuals analysis, i.e.,



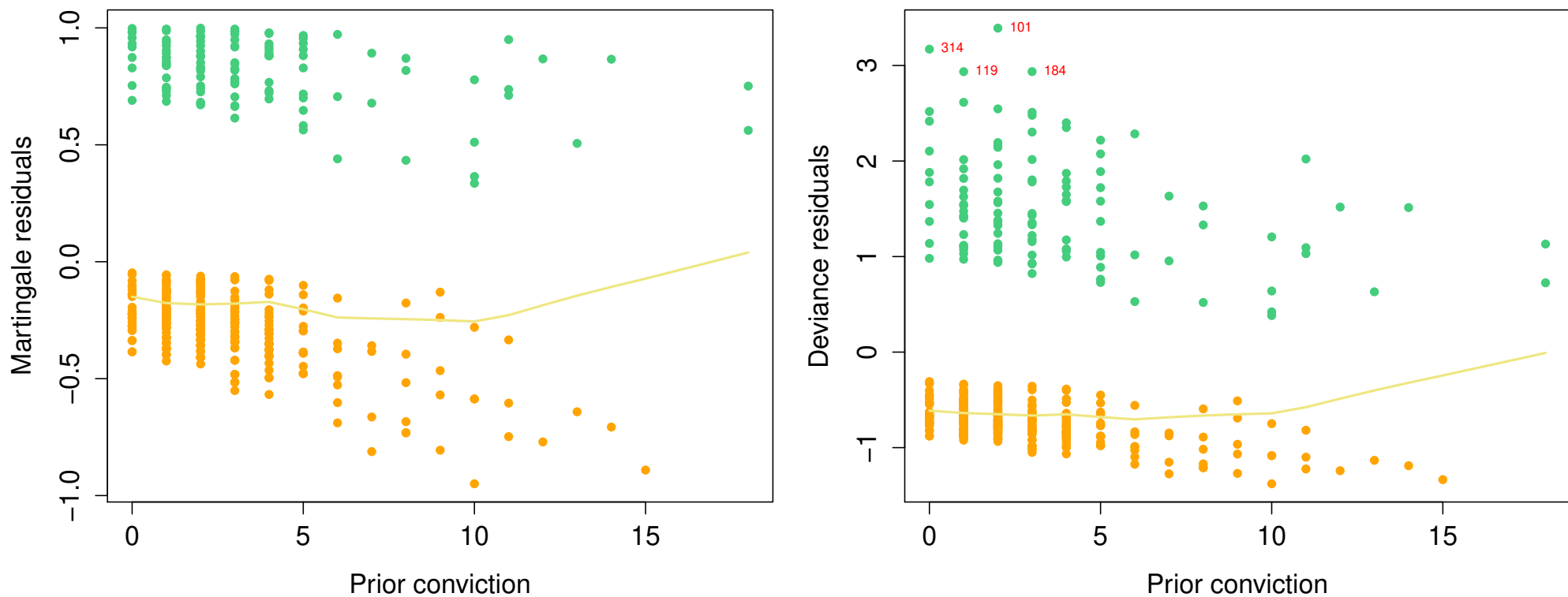
**Figure 16:** Martingale and deviance residuals for our best fitted model.

# Residuals analysis, i.e.,



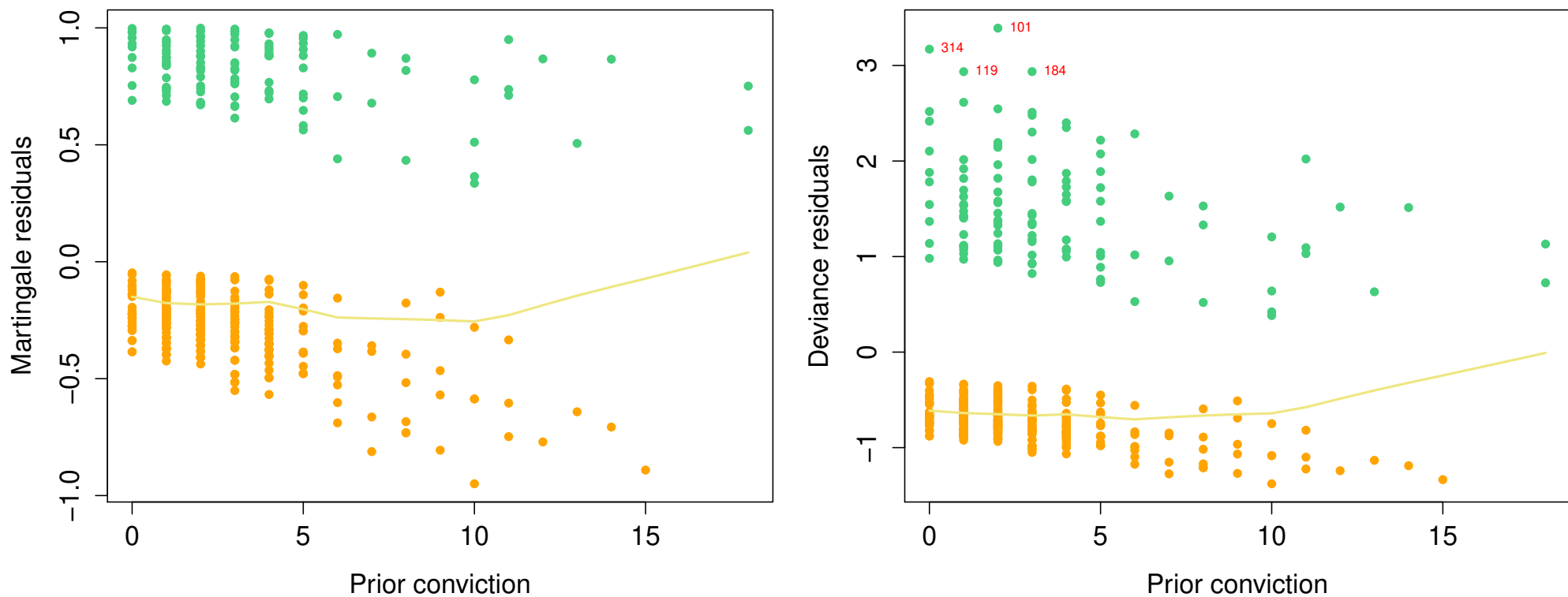
**Figure 16:** Martingale and deviance residuals for our best fitted model.

# Residuals analysis, i.e.,



**Figure 16:** Martingale and deviance residuals for our best fitted model.

# Residuals analysis, i.e.,



**Figure 16:** Martingale and deviance residuals for our best fitted model.

- No clear trend in residuals except for age where there is slight upward trend
- We may investigate the 4 outliers and check if **influentials** (not done but OK)

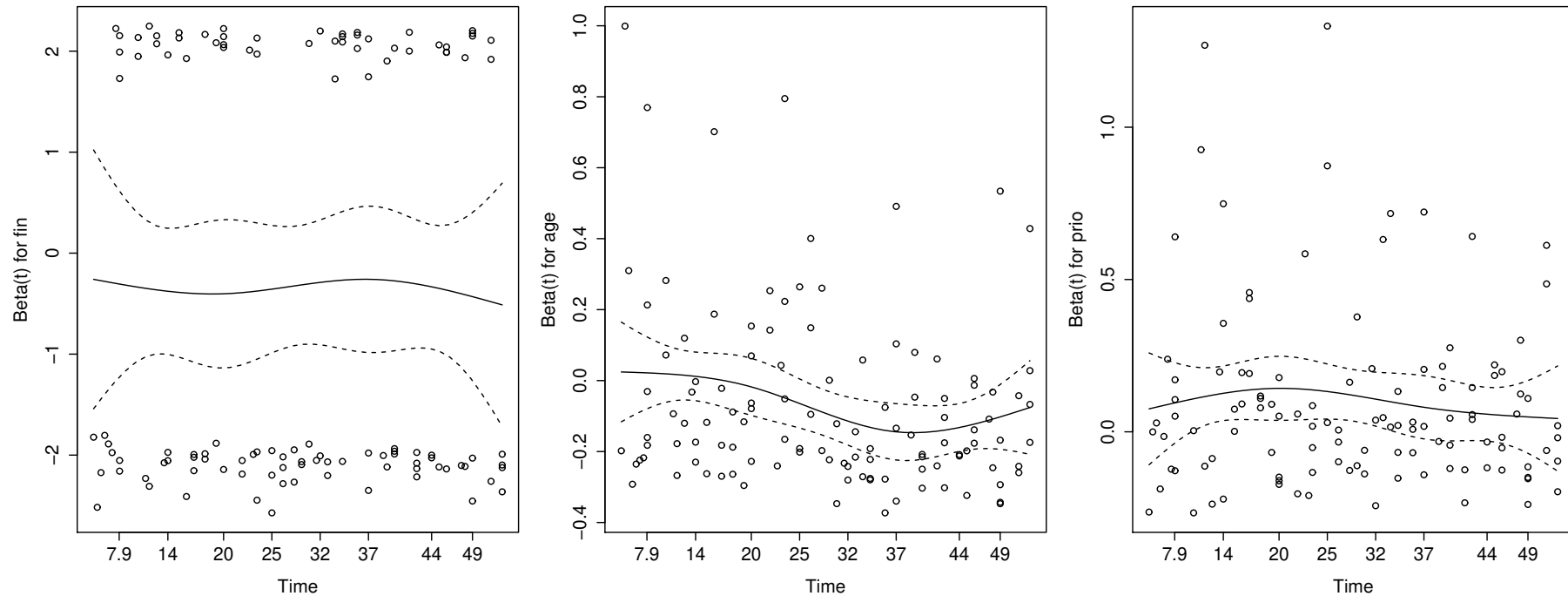
# Outliers

---

```
> idx.outliers <- which(residuals(fit, "deviance") > 2.7)
> data[idx.outliers,]
   week arrest fin age race wexp          mar paro prio educ
101    2      1  no  44 black  yes not married  yes     2     5
119    8      1 yes  40 black  yes not married  yes     1     5
184    3      1  no  30 black  no not married  yes     3     3
314    1      1  no  20 black  no not married   no     0     3
```

- They correponds to (very) early recidivism

# Proportional hazards assumption checking



**Figure 17:** Graphical assessment of the proportional hazards assumption.

- The assumption seems OK except maybe for feature age but that's probably related to our outliers
- So we can safely state that our model is rather decent!
- We are (at last!) able to interpret our results

# Interpretation (all other things being equal!)

```
> summary(bestFit)
Call:
coxph(formula = Surv(week, arrest) ~ fin + age + prio, data = data)

            coef exp(coef) se(coef)      z Pr(>|z|)
finyes -0.34695   0.70684  0.19025 -1.824 0.068197 .
age     -0.06711   0.93510  0.02085 -3.218 0.001289 **
prio    0.09689   1.10174  0.02725  3.555 0.000378 ***
---
Signif. codes:  0 ‘***’ 0.001 ‘**’ 0.01 ‘*’ 0.05 ‘.’ 0.1 ‘ ’ 1

            exp(coef) exp(-coef) lower .95 upper .95
finyes     0.7068      1.4148    0.4868    1.0263
age        0.9351      1.0694    0.8977    0.9741
prio       1.1017      0.9077    1.0444    1.1622
```

- Financial support **decreases risk by 30%** (the 95% CI upper bound is slightly over 1 though)
- Being release one year older **decrease the risk by 6.5%** (I'm getting too old for this...)
- Having one more prior conviction **increases the risk by 10%**

# Probability of no recidivism prediction

---



**Figure 18:** Click on me to see the evolution of the probability of no recidivism as the number of prior conviction increases.

---

1. Preliminaries

2. Non parametric  
estimation

3. Coping with  
covariates

4. Time to recidivism

▷ Conclusion

---

# Conclusion

- 
- This lecture was pretty dense (w.r.t. the time slots allowed) but many interesting stuffs were not discussed at all.
  - Of particular interest are the following:
    - time dependent features (extended Cox's model)
    - parametric survival models, e.g., Weibull distribution.
    - multiple state models, e.g., alive and tumour-free, alive and tumour present, dead.
    - frailty models, e.g., including random effects in Cox's model.

- 
- This lecture was pretty dense (w.r.t. the time slots allowed) but many interesting stuffs were not discussed at all.
  - Of particular interest are the following:
    - time dependent features (extended Cox's model)
    - parametric survival models, e.g., Weibull distribution.
    - multiple state models, e.g., alive and tumour-free, alive and tumour present, dead.
    - frailty models, e.g., including random effects in Cox's model.

THAT'S IT! I HOPE YOU ENJOYED THIS  
LECTURE!