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April 11, 2022





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- 2 Survival function
- 3 Kaplan-Meier estimator
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- 5 Cox's proportional hazards model







We can measure **time** in:

years

Time-to-event data

- months
- seconds

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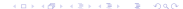
years

Time-to-event data •0000

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The **event** could be:

- death from disease
- product failure
- losing a customer







What is time-to-event (TTE) data?

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must be a binary variable







What is time-to-event (TTE) data?

We can measure time in:

- years
- months
- seconds

The **event** could be:

- death from disease
 product failure
 must be a binary variable
- losing a customer

yes/no

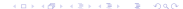
TTE data consists of (time, event) tuples.



Time-to-event (TTE) data

TTE analysis is also known as:

- survival analysis
- failure time analysis
- reliability theory (engineering)
- duration modelling (economics)
- event history analysis (sociology)







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Use cases for TTE analysis:

- clinical research
- customer analytics (churn)
- hardware (equipment failure)

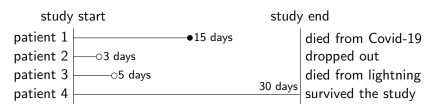


A randomised controlled trial (n = 4) was conducted to assess the efficacy of drug ABC in treating Covid-19. This is what happened to the patients:

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| patient | received ABC? | outcome |
|---------|---------------|--------------------------------------|
| 1 | yes | died from Covid-19 on day 15 |
| 2 | no | dropped out of the study after day 3 |
| 3 | yes | died by a lightning stroke on day 5 |
| 4 | no | survived the study (30 days) |

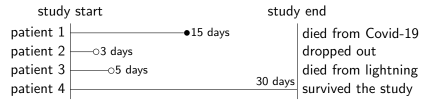
Example: Covid-19 treatment trial







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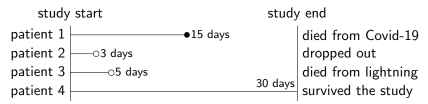


The **time** is the number of days since testing positive for Covid-19. The **event** is whether the patient died due to Covid-19.





Example: Covid-19 treatment trial

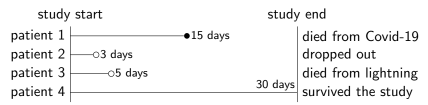


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| | • | | |
|--------------------|---------|------|-------|
| Time-to-event data | | | |
| | patient | time | event |
| | 1 | 15 | yes |
| | 2 | ? | ? |
| | 3 | ? | ? |
| | 4 | ? | no |



Example: Covid-19 treatment trial



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| Time-to-event data | | | | |
|--------------------|---------|---------|-------|--|
| | patient | time | event | |
| | 1 | 15 | yes | |
| | 2 | [0, 3] | no | |
| | 3 | [0, 5) | no | |
| | 4 | [0, 30] | no | |



> **Censoring** occurs when we have some information about an individual's survival time, but don't know the exact time. Possible reasons include

- not experiencing the event before the study concludes;
- getting lost to follow-up during the study period;
- withdrawing from the study.







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We just saw examples of right-censored data.







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Left censoring happens if the individual observed the event before the start of the study. This is often very hard to deal with and therefore not included in the study.

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Survival function

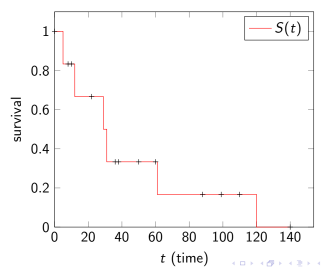
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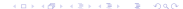
$$S(t) = \Pr(T > t)$$



Survival curve



The **Kaplan-Meier estimator** provides a non-parametric estimate of the survival function S(t) using the survival curve.





Modelling the survival function

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Kaplan-Meier estimator

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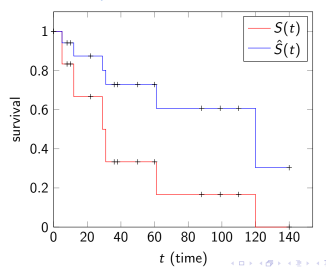
$$\hat{S}(t) = \prod_{i:t_i \le t} \left(1 - \frac{d_i}{n_i}\right)$$

where

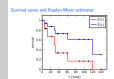
- t_i is an event time
- di is the number of deaths at time ti
- n_i is the number of individuals known to have survived until t_i



Survival curve and Kaplan-Meier estimator







Survival curve and Kaplan-Meier estimator

- When there is no censoring, $S(t) = \hat{S}(t)$.
- Commonly used to compare two study populations.
- Does not control for covariates.

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What does survival depend on?

Recall the survival function S(t) = Pr(T > t) as the probability that an individual will survive past time t.



Recall the survival function $S(t) = \Pr(T > t)$ as the probability that an individual will survive past time t. Let's assume that S(t)depends on

- 1 the **baseline hazard function** (how risk of event occurence changes over time at baseline covariates); and
- 2 the effect parameters (how hazard varies due to the covariates), also known as the partial hazard.



Cox's proportional hazards model uses both factors to provide a semi-parametric estimate of the hazard function $\lambda(t)$ conditioned on the covariates \mathbf{x} .

Cox's proportional hazards model

$$\lambda(t|\mathbf{x}) = \overbrace{\lambda_0(t)}^{ ext{baseline}} \underbrace{\exp\left(\sum_{i=1}^n \beta_i \mathbf{x}_i\right)}^{ ext{partial hazard}}$$

—Cox's proportional hazards model



- $\lambda_0(t)$ is a population-level baseline hazard that changes over time (for a reference individual with zeroed covariates).
- The partial hazard is a linear function of the covariates that is exponentiated. Each coefficient β_i is the relative risk associated with covariate \mathbf{x}_i .

Proportional hazards assumption

The model assumes fixed **proportional hazards**, i.e. the hazard for an individual i in proportion to the hazard of any other individual i is fixed over time.





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Therefore.

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the baseline hazard λ₀(t) is independent of the covariates, and
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The so-called **extended Cox model** allows the partial hazard to vary with time, and therefore no longer satisfies the proportional hazards assumption.

Partial likelihood

For each individual i. let

- T_i be a possibly censored survival time random variable, and
- X; denote the covariates.

Further, let the **risk set** $\mathcal{R}(t) = \{i : T_i \geq t\}$ be the set of individuals that are "at risk" at time t.

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Maximising this function allows us to estimate the parameters β .

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$$L(\beta) = \prod_{j=1}^{N} \Pr(\text{individual } j \text{ dies } | \text{ one death from } \mathcal{R}(T_j))$$



 $L(\beta) = \prod Pr (individual i dies | one death from <math>R(T_i)$)

• $L_j(\beta)$ is a *partial* likelihood because it considers only patients who died, not those that are censored.

Partial likelihood formula

$$L(\beta) = \prod_{j=1}^{N} \Pr\left(\text{individual } j \text{ dies } | \text{ one death from } \mathcal{R}(T_j)\right)$$

$$= \dots$$

$$= \prod_{j=1}^{N} \frac{\lambda(T_j | \mathbf{X}_j)}{\sum_{k \in \mathcal{R}(T_j)} \lambda(T_j | \mathbf{X}_k)}$$

$$= \prod_{j=1}^{N} \frac{\lambda_0(T_j) \exp\left(\beta \mathbf{X}_j\right)}{\sum_{k \in \mathcal{R}(T_j)} \lambda_0(T_j) \exp\left(\beta \mathbf{X}_k\right)}$$

$$= \prod_{j=1}^{N} \frac{\exp\left(\beta \mathbf{X}_j\right)}{\sum_{k \in \mathcal{R}(T_j)} \exp\left(\beta \mathbf{X}_k\right)}$$



Parameter estimation

We can estimate the parameters β by minimizing the negative partial log-likelihood, i.e. $-\log L(\beta)$, by taking the partial derivatives with respect to the parameters $oldsymbol{eta}$ and solving for the minimum using e.g. the Newton-Raphson algorithm.

Hazard ratios

The fraction used to express the proportional hazards assumption is actually the **hazard ratio**, measuring the risk of individual i relative to individual *j*:

$$HR = rac{\lambda(t|\mathbf{X}_i)}{\lambda(t|\mathbf{X}_i)} = \exp\left(eta(\mathbf{X}_i - \mathbf{X}_j)
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We may be interested in the relative risk associated with a particular covariate c, specifically the risk of said covariate having value c_i compared to c_i . Consider two dummy individuals i and jdiffering only in the c^{th} covariate, i.e. $\mathbf{X}_{i,k} = \mathbf{X}_{i,k}$ for $k \neq c$. Then the relative risk associated with c_i compared to c_i is

$$HR = \exp(\beta_c(c_i - c_i))$$
.



- HR = 1: no effect.
- HR > 1: increase in hazard
- HR < 1: reduction in hazard



