Applied Survival Analysis Using R Chapter 3: Nonparametric Survival Curve Estimation

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Kaplan-Meier estimator

 In this chapter we will discuss non-parametric estimators of the survival function

Kaplan-Meier estimator

KM estimator is the product over the failure times of the conditional probabilities of surviving to the next failure time.

Formally, it given by:

$$\hat{S}(t) = \prod_{t_i \le t} (1 - \hat{q}_i) = \prod_{t_i \le t} (1 - \frac{d_i}{n_i})$$
 (1)

• where n_i is the number of subjects at risk at time t_i , and d_i is the number of individuals who fail at that time, so $q_i = \frac{d_i}{n_i}$ is failure probability

Example

• The example data in Table 1.1 may be used to illustrate the construction of the *KM estimate*, as shown in Table 3.1.

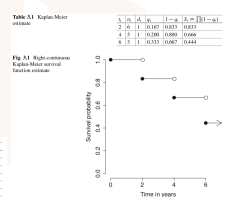


Table 1.1 Survival data

Patient	Survtime	Status
l .	7	0
2	6	1
3	6	0
1	5	0
5	2	1
	4	1

Get Variance By Delta Method

• To obtain confidence limits for the product-limit estimator, we first use what is known as the "delta method" to obtain the variance of $log(\hat{S}(t))$

$$\operatorname{Var}(\log(\hat{S}(t_k))) = \sum_{t_i \le t} \operatorname{Var}(\log(1 - \hat{q}_i)) \approx \sum_{t_i \le t} \frac{d_i}{n_i(n_i - d_i)}$$
(2)

• To get the variance of $\hat{S}(t)$ itself, we use the delta method again to obtain:

$$\operatorname{Var}(\hat{S}(t_k)) \approx [\hat{S}(t)]^2 \sum_{t_i < t} \frac{d_i}{n_i(n_i - d_i)}$$
(3)



log-log transformation

• A more satisfying approach is to find confidence intervals for the complementary log-log transformation of $\hat{S}(t)$ as follows:

$$\operatorname{Var}(\log([-\log \hat{S}(t_k)]) \approx \frac{1}{[\log \hat{S}(t)]^2} \sum_{t_i \leq t} \frac{d_i}{n_i(n_i - d_i)}$$
(4)



• To obtain estimates of the *Kaplan-Meier estimator* in R for the data in Table 1.1, we first load the "survival" library, and then enter the data

```
1 > library(survival)

2 > tt <- c(7,6,6,5,2,4)

3 > cens <- c(0,1,0,0,1,1)

4 > Surv(tt, cens)

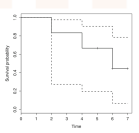
5 [1] 7+ 6 6+ 5+ 2 4
```

For the estimation itself we use the "survfit" function

R

• To see the full *Kaplan-Meier estimate*, and plot it, we use the "summary" and "plot" functions:

```
summary (result.km)
time
       n.risk
                 n.event.
                          survival
                                      std.err
                                                 lower95%CT
                                                               upper95%CI
                         0.833
                                      0.152
                                                   0.2731
                                                                 0.975
                         0.667
                                      0.192
                                                   0.1946
                                                                 0.904
                         0.444
                                      0.222
                                                   0.0662
                                                                 0.785
  plot(result.km)
```



Nelson-Aalen Estimator

- Based on the relationship of S(t) and h(t).
- An estimate of the cumulative hazard function is the sum of the estimated hazards up to a time t_i:

$$H(t) = \sum_{t_i < t} \frac{d_i}{n_i} \tag{5}$$

and the survival function estimate is simply

$$S(t) = e^{-H(t)} \tag{6}$$

• The Nelson-Aalen estimate may be obtained using the "survfit" function with the option type = "fh"

```
result.fh <- survfit(Surv(tt, cens) ~ 1, conf.type="log-log",
type="fh")
> summary(result.fh)
[1]
time
      n.risk
                n.event
                         survival
                                    std.err
                                              lower95%CT
                                                            upper95%CI
          6
                         0.846
                                    0.155
                                                  0.2401
                                                               0.981
          5
                         0.693
                                     0.200
                                                 0.1799
                                                               0.925
                         0.497
                                     0.248
                                                  0.0585
                                                               0.841
```

Median and CI for Median

- Formally, the median survival time may be defined as: $\hat{t}_{med} = \inf\{t : \hat{S}(t) \le 0.5\}$
- To find a 1α confidence interval for the median:

$$-z_{\alpha/2} \le \frac{g\{\hat{S}(t)\} - g(0.5)}{\sqrt{\text{Var}[g\{\hat{S}(t)\}]}} \le z_{\alpha/2}$$
 (7)

• where $g(\mu) = \log[-\log(\mu)]$ and $\operatorname{Var}[g\{\hat{S}(t)\}]$ is given by equation (4)

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Left Truncation

Left Truncation

Instead of examining the time from entry into the clinical trial until censoring or death, let us use as the time origin the time of diagnosis.

Table 3.3 Data from Table 1.1, with the addition of the time of diagnosis

Patient	Diagnosis	Survtime	Censor	SurvtimeDiag
1	-2	7	0	9
2	-5	6	1	11
3	-3	6	0	9
4	-3	5	0	8
5	-2	2	1	4
6	-5	4	1	9
X	-4	-2	1	

The time units are still the same, with time 0 indicating the time of entry into the trial and the time "Diagnosis" indicating the prior time of diagnosis. The new variable "SurvtimeDiag" denotes the time from diagnosis until censoring or death. The variables "Survtime" and "Censor" are as they were in Table 1.1. The new "Patient X" is a hypothetical patient with a short time from diagnosis until death. Practically speaking, such a patient is never observed; even if we somehow had a record of his diagnosis and early death, we could not possibly know for certain if that person would have entered the trial had he lived long enough. Such patients with short survival times are less likely to be enrolled in the trial than other patients, resulting in length-biased sampling

Figure

Fig. 3.8 Data from Table 1.1, now with diagnosis times

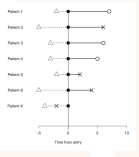
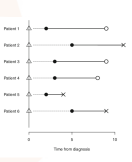


Fig. 3.9 Time from diagnosis to death. Entry into the clinical trial is denoted by solid circles. The dashed lines are "left truncation" times. Had the event occurred during these intervals, the patient would not have been

observed



 We have used the terms "tm.enter" and "tm.exit" for the left truncation and survival times, respectively.

R

```
> tt <- c(7, 6, 6, 5, 2, 4)
 > status <- c(0, 1, 0, 0, 1, 1)
  > backTime <- c(-2, -5, -3, -3, -2, -5)
4 > tm.enter <- -backTime
  > tm.exit <- tt - backTime
  > result.left.trunc.km <- survfit(Surv(tm.enter, tm.exit, status,</pre>
7 type="counting") ~ 1, conf.type = "none")
  > summary(result.left.trunc.km)
  [1]
10 time n.risk n.event entered censored survival std.err
                                               0.750
                                                         0.217
                                                0.562
                                                          0.230
                                                0.000
                                                         NAN
13
  > result.left.trunc.naa <- survfit(Surv(tm.enter, tm.exit, status,
  type="counting") ~ 1, type="fleming-harrington", conf.
15
16 type="none")
  > summary(result.left.trunc.naa)
  [1]
18
  time n.risk n.event entered censored
                                              survival std.err
19
                                               0.779
                                                          0.225
20
                                               0.607
                                                           0.248
21
22 | 11
                                               0.223
                                                            Inf
```

Data

- A serious problem arises with left-truncated data if the risk set becomes empty at an early survival time.
- Consider for example the *Channing House data*, "ChanningHouse".
- This data is subject to *left truncation* because subjects who die at older ages are more likely to have enrolled in the center than patients who died at younger ages.

1	>	head (Chanr	ningHou	ıse)			
2	[1	l] sex	entry	exit	time	cens	
3	1	Male	782	909	127	1	
4	2	Male	1020	1128	108	1	
5	3	Male	856	969	113	1	
6	4	Male	915	957	42	1	
7	5	Male	863	983	120	1	
8	6	Male	906	1012	106	1	



Transform data and using KM and NAA estimator

Transform the data

```
>ChanningHouse <- within(ChanningHouse,
entryYears <- entry/12exitYears <- exit/12)
>ChanningMales <- ChanningHouse[ChanningHouse$sex == "Male",]
```

KM estimator

```
1 >result.km <- survfit(Surv(entryYears, exitYears, cens,
2 type="counting") ~ 1, data=ChanningMales)
3 >plot(result.km, xlim=c(64, 101), xlab="Age",
4 ylab="Survival probability", conf.int=F)
```

NAA estimator

```
1 >result.naa <- survfit(Surv(entryYears, exitYears, cens,
2 type="counting") ~ 1, type="fleming-harrington",
3 data=ChanningMales)
4 >lines(result.naa, col="blue", conf.int=F)
```

KM.68 and Plot

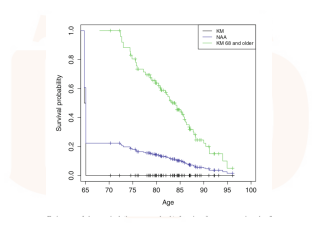
Men reach the age of 68, using the "start.time" option:

```
>result.km.68 <- survfit(Surv(entryYears, exitYears, cens, type="counting") ~ 1, start.time=68, data=ChanningMales)
```

Plot

```
1 > lines(result.km.68, col="green", conf.int=F)
2 > legend("topright", legend=c("KM", "NAA", "KM 68 and older"),
3 lty=1, col=c("black", "blue", "green"))
```

Plot



Apparently, the KM 68 and older is much better behaved.

