

STAT347: Generalized Linear Models

Lecture 14

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Today's topics:

- Survival analysis
 - Examples of survival analysis datasets
 - Basic concepts in survival analysis: survival function, hazard rate, censoring
 - Kaplan-Meier estimator of the survival function

Example 1:
Northern California Oncology Group (NCOG) study

- Two treatments for head and neck cancer:
Arm A: Chemotherapy; Arm B: Chemotherapy + Radiation
- Data: censored survival time in days
'+' indicate patients still alive on their final day of observation

Arm A: Chemotherapy										
7 129 160 279+ 1101	34 133 165 297 1116+	42 133 173 319+ 1146	63 139 176 405 1226+	64 140 185+ 417 1349+	74+ 140 218 420 1412+	83 146 225 440 1417	84 149 241 523	91 154 248 523+	108 157 273 583	112 160 277 594
Arm B: Chemotherapy+Radiation										
37 146 319 817 2297+	84 155 339 1092+	92 159 432 1245+	94 169+ 469 1331+	110 173 519 1557	112 179 528+ 1642+	119 194 547+ 1771+	127 195 613+ 1776	130 209 633 1897+	133 249 725 2023+	140 281 759+ 2146+

Example 1:

Northern California Oncology Group (NCOG) study

- Two treatments for head and neck cancer:
Arm A: Chemotherapy; Arm B: Chemotherapy + Radiation
- Data: censored survival time in days
+ indicate patients still alive on their final day of observation

Main questions:

- Is the Arm B more effective treatment than Arm A?
- Instead of just compare the mean survival time, we would like to know more information about the survival time distribution (the survival curve)
- How to deal with “lost to follow-up” (censoring)?

Example 2: duration of nursing home stay

- Goal: assess the effects of different financial incentives on length of stay.
- Treated nursing homes received higher per diems for Medicaid patients, and bonuses for improving a patient's health and sending them home.
- Study included 1601 patients admitted between May 1, 1981 and April 30, 1982.

Measured variables:

- LOS - Length of stay of a resident (in days)
- AGE - Age of a resident
- RX - Nursing home assignment (1:bonuses, 0:no bonuses)
- gender, age, married or not, health status
- CENSOR - Censoring indicator (1:censored, 0:discharged)

Goal: treatment effect on stay length after adjusting for other covariates and censoring?

Basic concepts

- Survival time: T is a random non-negative variable, the duration from the start of treatment to death.
 - Continuous: T has a density function $f(t)$
 - Discrete: $T \in \{0, 1, 2, 3, \dots\}$, $f_i = P(T = i)$
- Survival function/curve: $S(t) = P(T > t)$
 - Continuous: $S(t) = \int_t^\infty f(t') dt'$
 - Discrete: $S_i = \sum_{j>i} f_j$
- Hazard rate/function: $h(t) = f(t)/S(t)$ (or $h_i = f_i/s_{i-1}$ for discrete T)
- Accumulative hazard function: $H(t) = \int_0^t h(t)$ (or $H_i = \sum_{j \leq i} h_j$ for discrete T)

Basic concepts

- The survive function and hazard rate provide more information than $E(T)$.
- An important fact is that knowing one of the three functions of $H(t)$, $h(t)$ and $S(t)$ will enable inferring the other two functions.
- For discrete T :

$$S_i = \prod_{j=0}^i P[T \geq j + 1 \mid T \geq j] = \prod_{j=0}^i (1 - h_j)$$

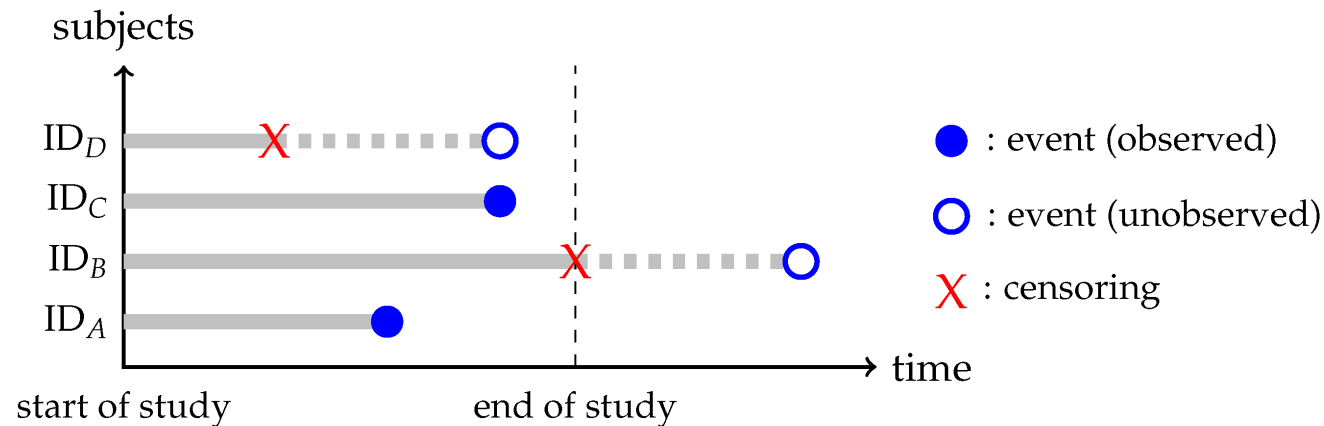
- For continuous T :

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

Concept of censoring

Censoring

- We may not be able to observe every T_i where i is an individual.
- Censoring can occur when
 - the study ends, some individual have not had the event yet (still alive)
 - Some individuals dropout or get lost in the middle of the study.



- Typically, individuals do not enter the study at the same time
 - Not a concern as T_i is the length of duration
 - can adjust for starting time by add it as a covariate

Concept of censoring

Denote each sample's censoring time as C_1, C_2, \dots, C_n . Then what we can actually observe for each sample are $Y_i = \min(T_i, C_i)$ and an indicator of whether censoring occurs:

$$\delta_i = \begin{cases} 0 & \text{if } T_i \leq C_i \text{ (observed death)} \\ 1 & \text{Otherwise} \end{cases}$$

When each sample also has its covariate, what we observe can be denoted as (Y_i, X_i, δ_i) for $i = 1, 2, \dots, n$.

Throughout the class, we only consider **non-informative censoring**, which is basically requiring

$$T_i \perp C_i \mid X_i$$

which means that the censoring time is not associated with the survival time, at least conditioning on other known covariates X_i .

Estimating the survival function

- We consider the scenario with no observed covariates X_i and the survival time T_i are i.i.d.
- A non-parametric way with no censoring

$$\hat{S}_n(t) = \frac{1}{n} \sum_i 1_{T_i > t}$$

- This does not work if there are censored data
- Example:

survival times: 1, 1, 2, 2+, 3+, 4, 4, 5, 5, 8, 8, 8, 8, 11, 11, 12, 12, 15, 17, 22, 23

We don't know how to estimate $S(3)$ from the empirical cdf approach

Kaplan-Meier estimator

- Assume we have discrete time points
- Make use of the equation:

$$S_i = \prod_{j=0}^i P[T \geq j+1 \mid T \geq j] = \prod_{j=0}^i (1 - h_j)$$

- How to estimate a hazard rate h_i ? For time bin i , assume
 - r_i samples that are still alive at the beginning of this time bin
 - d_i death during this time bin
 - c_i drop-outs at the end of this time bin
 - No drop-outs during this time bin

$$d_i \sim \text{Bernoulli}(r_i, h_i) \qquad \hat{h}_i = \frac{d_i}{r_i}$$

- Kaplan-Meier estimator

$$\hat{S}_i = \prod_{j \leq i} (1 - \hat{h}_j)$$

Kaplan-Meier estimator

- For continuous T , we can discretize time into bins and make the bin size smaller and smaller
- The Kaplan-Meier estimator in the limiting case becomes

$$\hat{S}(t) = \prod_{j: \tau_j \leq t} \frac{r_j - d_j}{r_j}$$

where $\{\tau_1, \tau_2, \dots, \tau_K\}$ is the set of K distinct uncensored failure times observed in the sample, d_j is the number of death at τ_j and r_j is the total number of people who are at risk right before τ_j .

- The above formula also works for discrete time points

Variance of $\hat{S}(t)$

- The estimates $\hat{h}_1, \dots, \hat{h}_K$ are not independent: $r_{j+1} = r_j - d_j - c_j$, $\hat{h}_i = \frac{d_i}{r_i}$

The Greenwood formula for estimating the uncertainty in $\hat{S}(t)$:

$$\log \hat{S}(t) = \sum_{j: \tau_j \leq t} \log(1 - \hat{h}_j)$$

Using the Delta method

$$\begin{aligned} \log \hat{S}(t) &\approx \sum_{j: \tau_j \leq t} \left[\log(1 - h_j) - \frac{1}{1 - h_j} (\hat{h}_j - h_j) \right] \\ &= \text{Const} - \sum_{j: \tau_j \leq t} \frac{1}{1 - h_j} (\hat{h}_j - h_j) \end{aligned}$$

- Though the estimates $\hat{h}_1, \dots, \hat{h}_K$ are not independent, we always have $E[\hat{h}_i - h_i | \hat{h}_1, \dots, \hat{h}_{i-1}] = 0$

- The partial sums form a martingale
- $\hat{h}_1, \dots, \hat{h}_K$ are pairwise uncorrelated

Variance of $\hat{S}(t)$

- When calculating the variance, we can treat $\hat{h}_1, \dots, \hat{h}_K$ as “independent” and K as fixed.

$$\begin{aligned}\widehat{\text{Var}}\left(\log \hat{S}(t)\right) &\approx \sum_{j:\tau_j \leq t} \left(\frac{1}{1 - \hat{h}_j}\right)^2 \widehat{\text{Var}}(\hat{h}_j) \\ &= \sum_{j:\tau_j \leq t} \frac{\hat{h}_j}{(1 - \hat{h}_j)r_j} = \sum_{j:\tau_j \leq t} \frac{d_j}{(r_j - d_j)r_j}\end{aligned}$$

Using Delta method on $\hat{S}(t) = e^{\log \hat{S}(t)}$, we get

$$\begin{aligned}\widehat{\text{Var}}\left(\hat{S}(t)\right) &= [\hat{S}(t)]^2 \widehat{\text{Var}}\left(\log(\hat{S}(t))\right) \\ &= [\hat{S}(t)]^2 \sum_{j:\tau_j \leq t} \frac{d_j}{(r_j - d_j)r_j}\end{aligned}$$

- Under some conditions, we can also have CLT of $\log \hat{S}(t)$

Comparison between two survival survival curves

- In the NCOG data, we may want to know if the whole survival curve of Arm B is significantly larger than the whole curve of Arm A.
- Here, we consider testing for the simple null hypothesis

$$H_0 : S_1(t) \equiv S_2(t)$$

This tests if the two curves are exactly the same

The Cochran-Mantel-Haenszel log-rank test

- Assume we have discrete time points
- For each discrete survival time i ,
 - We observe r_{i1} and r_{i2} samples that are still alive at the beginning of this time bin for each group respectively
 - Observe d_{i1} and d_{i2} death during this time bin for two groups respectively.
 - Assume that drop-outs happen at the end of each time bin. (so we don't need to consider it)

	death	alive	total at risk
Group 1	d_{i1}	$r_{i1} - d_{i1}$	r_{i1}
Group 2	d_{i2}	$r_{i2} - d_{i2}$	r_{i2}
Total	d_i	$r_i - d_i$	r_i

The Cochran-Mantel-Haenszel log-rank test

The Cochran-Mantel-Haenszel log-rank test is to test whether the group has no effect on death rate in each table. If the margins of this table are considered fixed, then under H_0 , d_{i1} follows a Hypergeometric distribution, with (check the Wikipedia page)

$$E(d_{i1}) = \frac{d_i}{r_i} r_{i1}, \quad \text{Var}(d_{i1}) = \frac{r_{i1} r_{i2} d_i (r_i - d_i)}{r_i^2 (r_i - 1)}$$

The log-rank test statistics is

$$X_{CMH}^2 = \frac{\{\sum_i (d_{i1} - r_{i1} d_i / r_i)\}^2}{\sum_i r_{i1} r_{i2} d_i (r_i - d_i) / [r_i^2 (r_i - 1)]}$$

- Compare X_{CMH}^2 with a χ_1^2 distribution to get p-value

The Cochran-Mantel-Haenszel log-rank test

For continuous survival time, we can make the bin finer and finer, and in the limit, the Cochran-Mantel-Haenszel log-rank test statistics is

$$X_{CMH}^2 = \frac{\left\{ \sum_{j=1}^K (d_{j1} - r_{j1}d_j/r_j) \right\}^2}{\sum_{j=1}^K r_{j1}r_{j2}d_j(r_j - d_j)/[r_j^2(r_j - 1)]}$$

where $\{\tau_1, \tau_2, \dots, \tau_K\}$ is the set of K distinct uncensored failure times observed in the sample including both two groups, d_{j1} and d_{j2} are the number of death at τ_j for each group respectively, and r_{j1} and r_{j2} are the total number of people who are at risk right before τ_j for each group respectively. $r_j = r_{j1} + r_{j2}$ and $d_j = d_{j1} + d_{j2}$.

Some remarks

- The asymptotics work when the total number of samples n goes to ∞ , so we can have either a fixed K or a growing number of K
- For each 2×2 table, there can be many different tests for the group effect or death, for example testing for the odds ratio being 1 with a logistic regression, the challenge is to combine K different tables and have valid inference when each y_j is very small (exactly 1 when there is no tie).
- The CMH log-rank test is powerful when the survive curves does not across each other. It is most powerful when $h_2(t) = \alpha h_1(t)$
- the Log-rank test is non-parametric, and only depends on the ranks

Data example

- Check Example10 R notebook