

Tips and Tricks for Communicating Survival Analysis Results to Clinical Collaborators

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R/Medicine Conference

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Memorial Sloan Kettering
Cancer Center

 COLUMBIA UNIVERSITY | MAILMAN SCHOOL
of PUBLIC HEALTH
BIostatISTICS

The most common questions in cancer research relate to disease survival

RESEARCH ARTICLE

Ten-year experience with ophthalmic artery chemosurgery: Ocular and recurrence-free survival

Jasmine H. Francis^{1,2†}, Ariana M. Levin^{2†}, Emily C. Zabor¹, Y. Pierre Gobin², David H. Abramson^{1,2}

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JOURNAL OF CLINICAL ONCOLOGY

ORIGINAL REPORT

Preexisting Cardiovascular Risk and Subsequent Heart Failure Among Non-Hodgkin Lymphoma Survivors

Talya Salis, Emily C. Zabor, Peter de Nully Brown, Susanne Okoberg Dalton, Nirupa J. Raghunathan, Matthew J. Maitland, Richard Steingart, Andrew J. Vickers, Peter Sørensen Munksgaard, Kevin C. Oeffinger, and Christoffer Johansen

Ann Surg Oncol (2017) 24:3141–3147
DOI: 10.1245/s10434-017-5965-5

Annals of
SURGICAL ONCOLOGY
OFFICIAL JOURNAL OF THE SOCIETY OF SURGICAL ONCOLOGY

ORIGINAL ARTICLE – BREAST ONCOLOGY

Oncologic Outcomes After Treatment for MRI Occult Breast Cancer (pT0N+)

Damian P. McCartan, MD¹, Emily C. Zabor, MS², Monica Morrow, MD¹, Kimberly J. Van Zee, MS¹, Mahmoud B. El-Tamer, MD¹

Personalized Medicine and Imaging

Clinical
Cancer
Research

DNA Damage Response and Repair Gene Alterations Are Associated with Improved Survival in Patients with Platinum-Treated Advanced Urothelial Carcinoma

Min Yuen Teo¹, Richard M. Bambury², Emily C. Zabor³, Emmet Jordan¹, Hikmat Al-Ahmadie⁴, Mariel E. Boyd⁵, Nancy Bouvier⁶, Stephanie A. Mullane⁶, Eugene K. Cha⁷, Nitin Roper⁸, Irina Ostrovnaya¹, David M. Hyman⁹, Bernard H. Bochner⁷, Maria E. Arcila⁴, David B. Solit¹, Michael F. Berger⁵, Dean F. Bajorin¹, Joaquin Bellmunt⁴, Gopakumar Iyer¹, and Jonathan E. Rosenberg¹

Survival analysis is a complex statistical procedure, so communication with collaborators is key

Strategies for conveying important information about survival analysis:

- ☐ Be ready with **examples** to explain complex ideas
- ☐ Use detailed **graphics** to convey information
- ☐ Accompany numbers such as p-values and hazard ratios with detailed **interpretations**

Example of a dataset with censored data for a clinical application

The `lung` dataset is available from the `survival` package in R. The data contain subjects with advanced lung cancer from the North Central Cancer Treatment Group.

Variable descriptions, from the documentation:

- `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 (0=good 5=dead)
- `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

Example of a dataset with censored data for a clinical application

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Example of a dataset with censored data for a clinical application

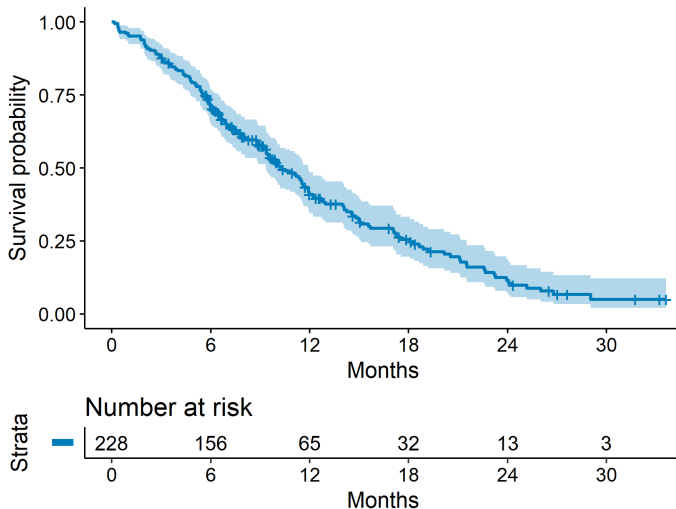
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Variable descriptions, from the documentation:

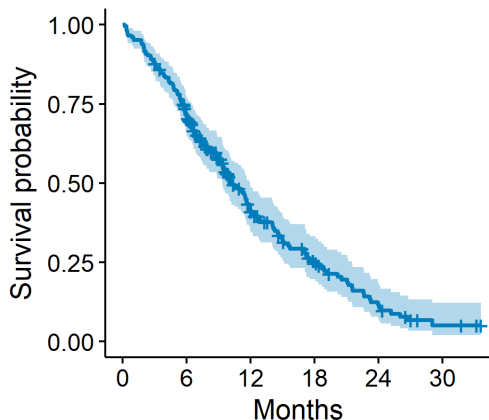
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The Kaplan-Meier survival function forms the basis of most survival analyses

Using the `ggsurvplot` function from the `survminer` package:

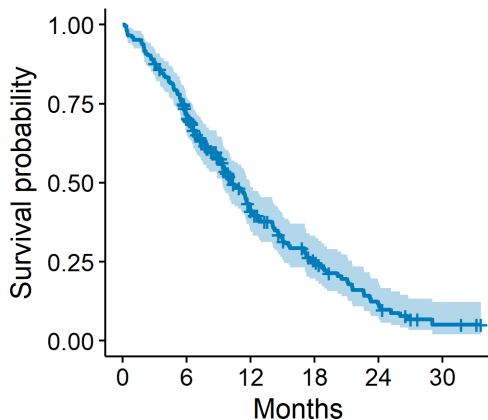


First just describe what is on the plot



- The x-axis is time and the y-axis is the survival function
- Step function where each step down represents a time at which one or more events occurred
- Censored subjects are usually denoted by tick marks

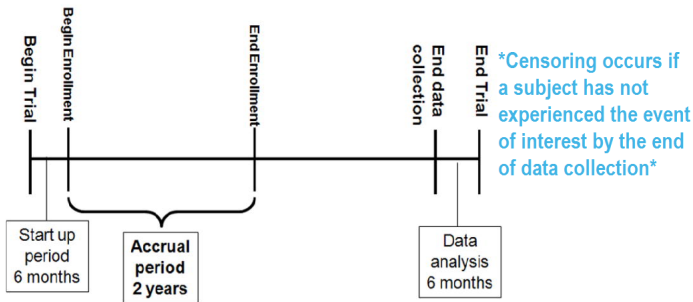
If appropriate give a basic description of the underlying calculations



The survival function is calculated at each time as the ratio of subjects who did not experience the event by that time to the total number of subjects still at risk at that time

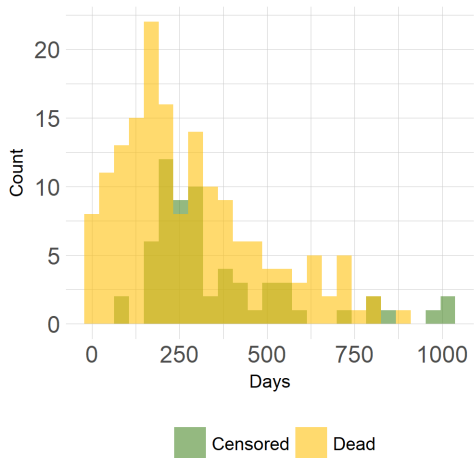
Question from collaborators: What is censoring?

In the context of a clinical trial:



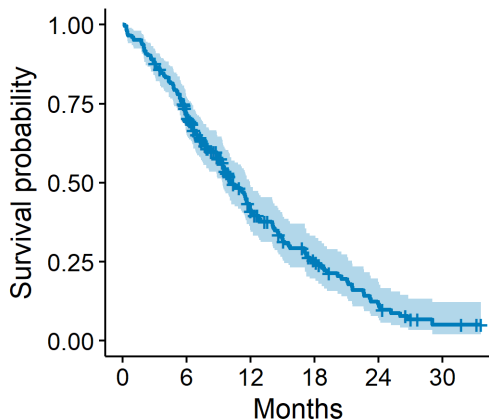
RICH JT, NEELY JG, PANIELLO RC, VOELKER CCJ, NUSSENBAUM B, WANG EW. A PRACTICAL GUIDE TO UNDERSTANDING KAPLAN-MEIER CURVES. Otolaryngology head and neck surgery: official journal of American Academy of Otolaryngology Head and Neck Surgery. 2010;143(3):331-336. doi:10.1016/j.otohns.2010.05.007.

Question from collaborators: Why do I need specialized methods to analyze time-to-event data?



1. Censored subjects still provide information so must be appropriately included in the analysis
2. The distribution of follow-up times is skewed

Question from collaborator: Can I report the percentage of events out of the total study population?

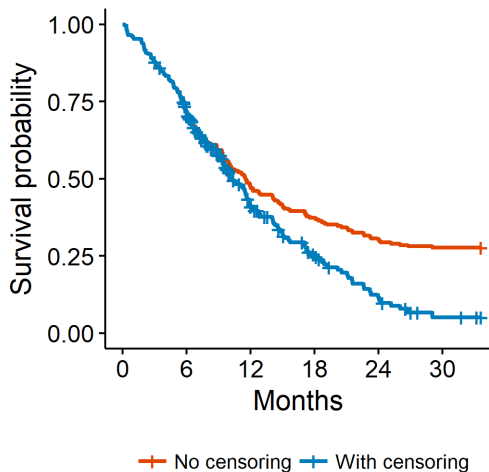


In the lung data this would lead to an estimate of survival probability at the end of the study of $1 - \frac{165}{228} = 0.28$.

But this is **incorrect**.

Why?

It is clear that the survival curve is lower when there is censoring during follow-up

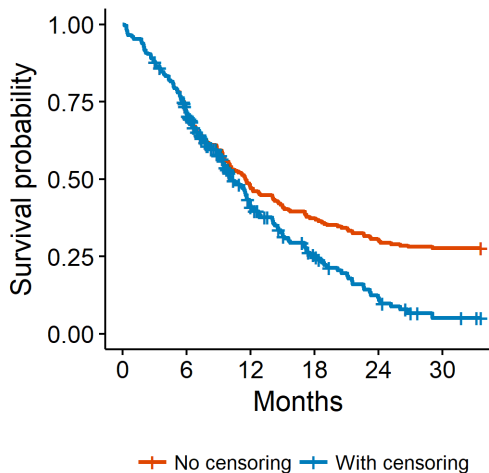


Imagine two studies, each with 228 subjects.

There are 165 deaths in each study.

The orange study has no censoring during follow-up, the blue study has subjects censored throughout (the true lung data).

When there is not censoring during follow-up, the survival estimate is straightforward



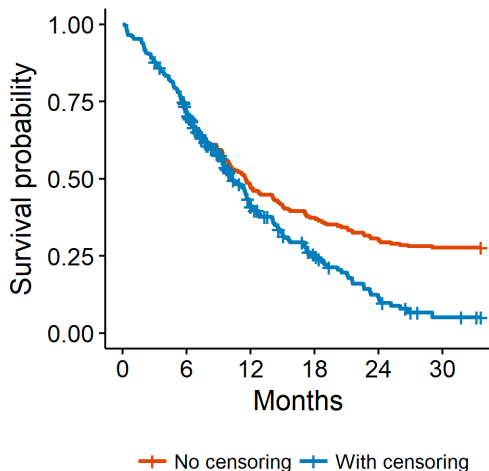
In the **orange** study everyone is followed until the last follow-up time (i.e. no censoring during follow-up).

The survival probability at last follow-up is 0.28

This is equivalent to an estimating the raw percent of patients still alive:

$$1 - \frac{165}{228} = 0.28$$

When there is censoring during follow-up, a naive estimate will be incorrect

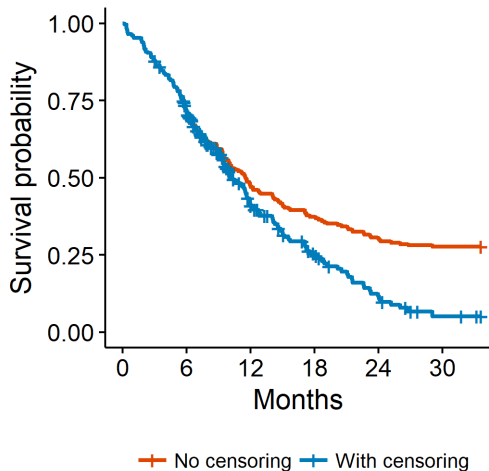


The blue study has 63 subjects censored during follow-up. Still 165 subjects died.

The blue curve is clearly lower than the orange curve, especially at the end.

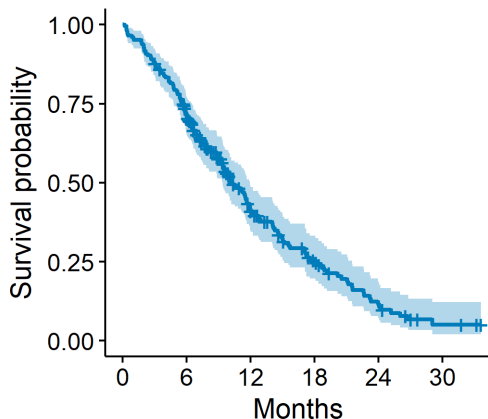
The survival probability at end of study = something < 0.28 (actually 0.005)

Ignoring censoring leads to an overestimate of the overall survival probability



This occurs because the censored subjects only contribute information for **part** of the follow-up time, and then they fall out of the denominator, thus pulling down the cumulative probability of survival.

Question from collaborator: What is the 1-year survival probability?

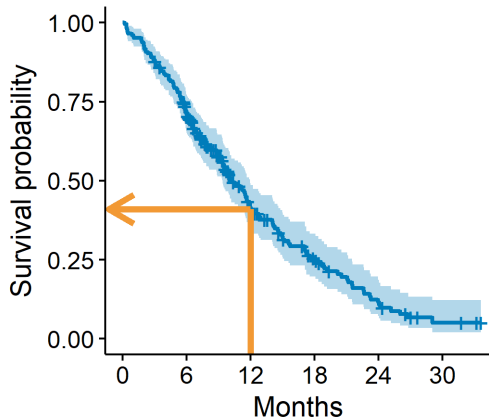


In the lung data the 1-year survival probability is 0.41

But where does this number come from?

And what does it mean?

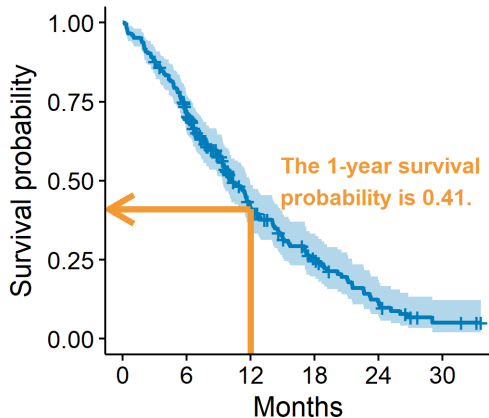
We start by showing what the 1-year survival probability is on the survival curve



The 1-year survival probability is the **probability on the y-axis** corresponding to **1-year on the x-axis**

The `geom_segment` function from `ggplot2` can help with this

Next we annotate the plot with text to state the probability alongside the curve



The glue function from the glue package provides an easy way to reproducibly annotate plots

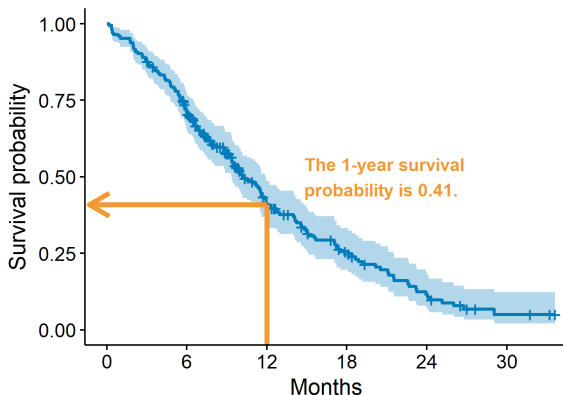
Additionally create reproducible text to make sure numbers are being correctly interpreted

As part of a reproducible workflow that includes writing reports in R Markdown, the `glue_data` function from the `glue` package can easily print reproducible text as a corresponding description:

```
fit1 <- survival::survfit(  
  survival::Surv(time, status) ~ 1, data = lung)  
glue::glue_data(summary(fit1, times = 365.25),  
  "The survival probability at 1-year is ",  
  "{round(surv, 2)} ",  
  "(95% CI: ", "{round(lower, 2)} - ", "{round(upper, 2)}",  
  ")\n which represents the estimated proportion of ",  
  "patients who\n survived beyond 1 year.")
```

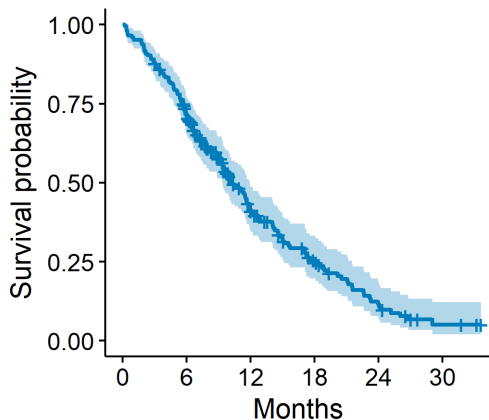
```
## The survival probability at 1-year is 0.41 (95% CI: 0.34 - 0.49),  
## which represents the estimated proportion of patients who  
## survived beyond 1 year.
```

Finally put everything together into a report to our collaborator about the 1-year survival probability



```
## The survival probability at 1-year is 0.41 (95% CI: 0.34 - 0.49),  
## which represents the estimated proportion of patients who  
## survived beyond 1 year.
```

Question from collaborator: What is the median survival time?

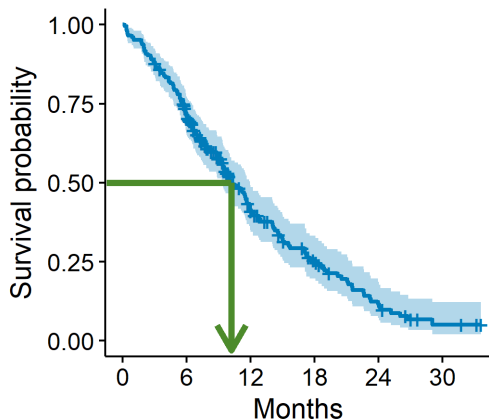


In the lung data the median survival time is 310 days

But where does this number come from?

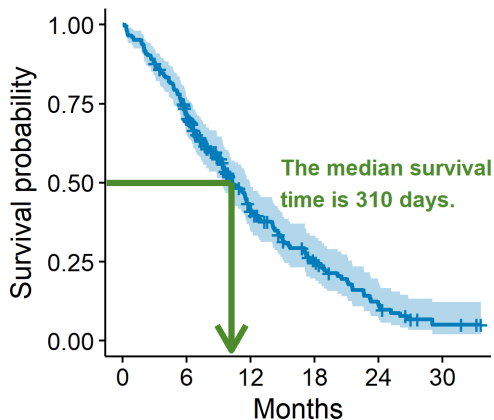
And what does it mean?

We start by showing what the median survival time is on the survival curve



The median survival time is the **time on the x-axis** corresponding to a **survival probability of 0.5** on the y-axis.

Next we annotate the plot with text to state the time estimate alongside the curve



We again use the `glue` function from the `glue` package to annotate our plot with reproducible text

Inline R code can also be used to incorporate reproducible text into reports alongside graphical examples

First define the `survfit` summary object, using the `survival` package:

```
fit1 <- survival::survfit(survival::Surv(time, status) ~ 1,  
                          data = lung)  
res <- summary(fit1)$table
```

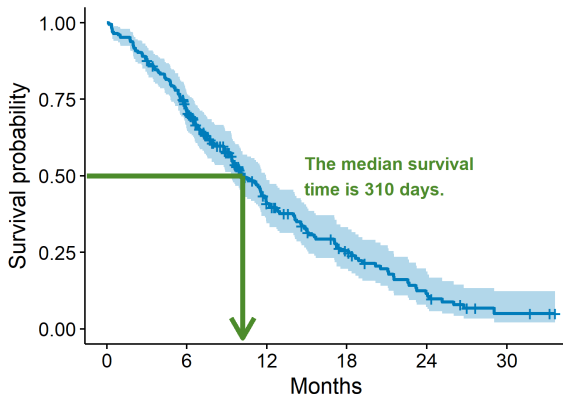
Then:

The median survival time is ``r round(res["median"])` days (95% CI: ``r round(res["0.95LCL"])` - ``r round(res["0.95UCL"])`), which represents the estimated point in time that half of subjects will live beyond.

On knitting this will print our desired description:

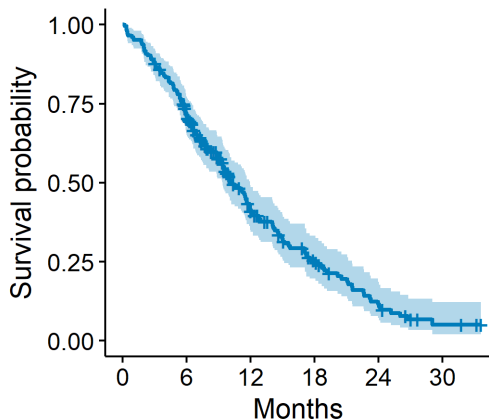
The median survival time is 310 days (95% CI: 285 - 363), which represents the estimated point in time that half of subjects will live beyond.

Finally put everything together into a report to our collaborator about the median survival time



The median survival time is 310 days (95% CI: 285 - 363), which represents the estimated point in time that half of subjects will live beyond.

Question from collaborator: Why can't I just estimate the median among those who had the event?

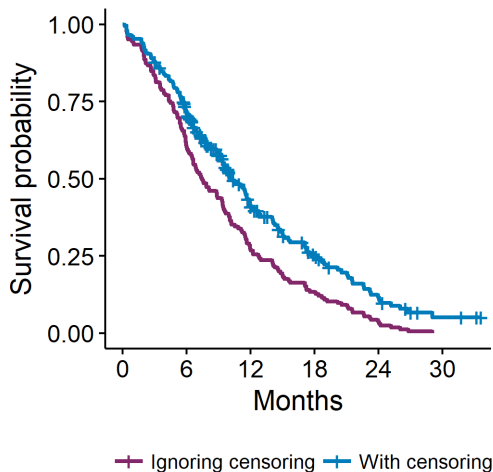


In the lung data this would lead to an estimate of median survival time of 226 days.

But this is **incorrect**.

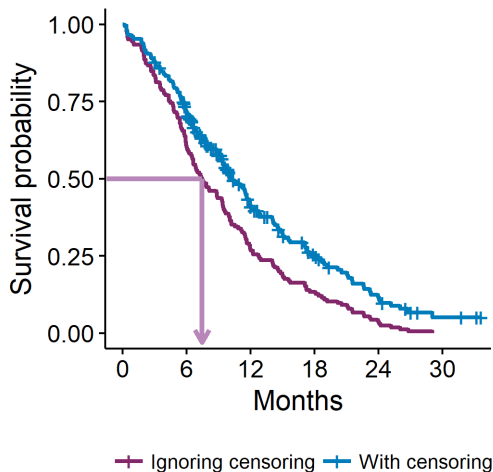
Why?

The true survival curve falls above a curve excluding the censored subjects



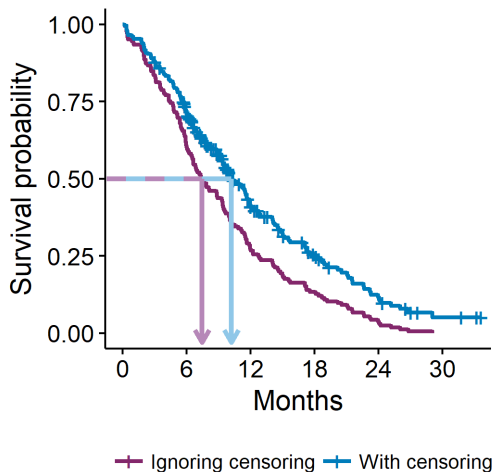
This graphical example compares the true survival curve in blue, and the curve excluding censored subjects in purple.

Ignoring censoring leads to an underestimate of the median survival time



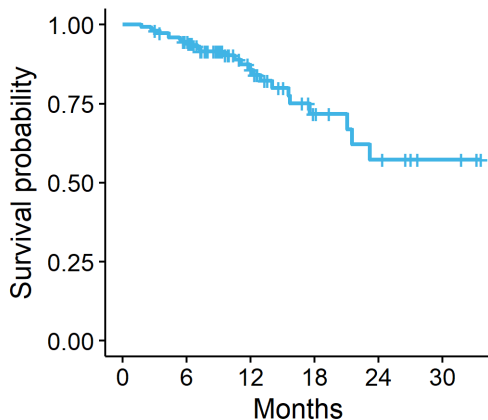
- The **purple** group ignores censored subjects.
- This is equivalent to the naive estimate of median time to death **among subjects who died** of 226 days

Censored subjects contribute information because we know their event occurred after the censoring time



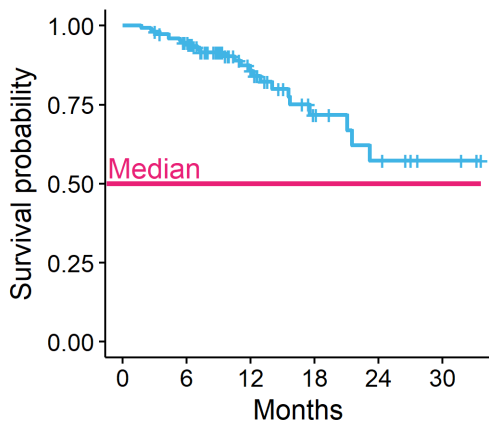
- The blue group includes the censored subjects.
- The median survival time is clearly longer, at 310 days.

Question from collaborator: What does it mean when you say median survival is not reached?



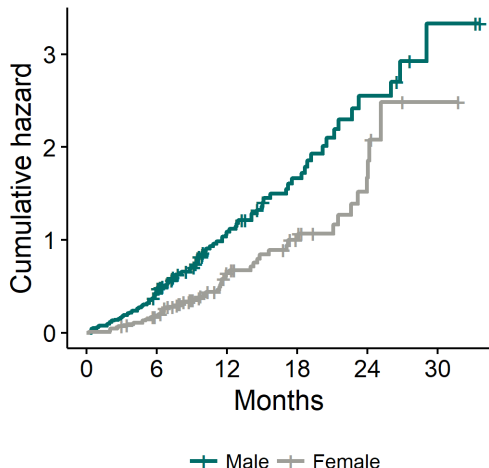
- Survival curve based on a sample of lung data subjects
- In these data, we would report a median survival time of NA

The survival curve must cross the survival probability of 0.5 before median survival time has been reached



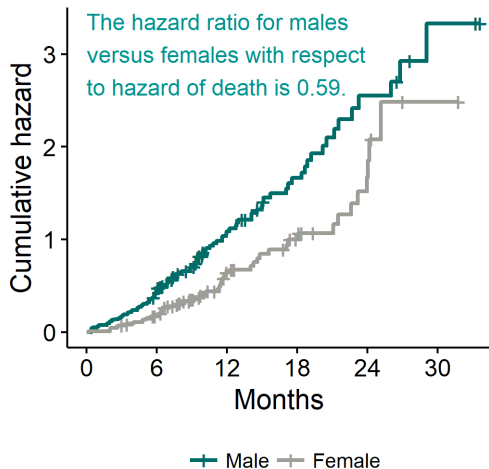
- By the end of the study, we have **not observed** the point in time that half the subjects will survive beyond
- Subjects would need to be followed longer, so that more deaths could occur, before we could observe median survival time

Question from collaborator: What is a hazard ratio?



- The hazard ratio (HR) represents the difference in hazards between two groups at any particular point in time
- $HR = 0.59$ comparing male to female with respect to hazard of death in the lung data

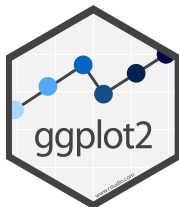
Hazard ratios apply at any given point in time, which is what differentiates them from relative risks



- $HR = 0.5$ would mean that at any particular time, half as many females as males are dying
- So our $HR = 0.59$ implies that around 0.4 times as many females are dying as males, at any given time

Essential R packages


- survival
- survminer
- ggplot2
- glue
- rmarkdown
- knitr





Thank you

Slides available at: <https://github.com/zabore/r-medicine>

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