CURVE-FITTING METHODS AND THE MESSAGES THEY SEND LINEAR QUADRATIC LOGARITHMIC . "HEY, I DID A "I WANTED A CURVED "LOOK, IT'S REGRESSION." LINE, SO I MADE ONE TAPERING OFF!" **WITH MATH.**" EXPONENTIAL LOESS LINEAR, NO SLOPE "I'M SOPHISTICATED, NOT "LOOK, IT'S GROWING "I'M MAKING A UNCONTROLLABLY!" LIKE THOSE BUMBLING SCATTER PLOT BUT POLYNOMIAL PEOPLE." I DON'T WANT TO." CONFIDENCE LOGISTIC PIECEWISE INTERVAL "I NEED TO CONNECT THESE "LISTEN, SCIENCE IS HARD. "I HAVE A THEORY, TWO LINES, BUT MY FIRST IDEA BUT I'M A SERIOUS AND THIS IS THE ONLY DIDN'T HAVE ENOUGH MATH." PERSON DOING MY BEST." DATA I COULD FIND." CONNECTING AD-HOC HOUSE OF LINES FILTER CARD5 "I HAD AN IDEA FOR HOW "I CLICKED 'SMOOTH "AS YOU CAN SEE, THIS LINES' IN EXCEL." TO CLEAN UP THE DATA. MODEL SMOOTHLY FITS WHAT DO YOU THINK?" THE- WAIT NO NO DON'T EXTEND IT AAAAAA!!"

Censoring, Survival, & Hazards

Introduction

What is Survival Analysis?

In survival analysis, we are interested in the **time until an event occurs**, or **failure time**.

Event is a qualitative change that can be tied to a specific point in time.

Originally designed to study the occurrence of death in medical studies – hence *survival analysis*.

Other names:

- Time-to-event analysis
- Duration analysis
- Failure time analysis
- Reliability analysis

"Time-to-Event" Data?

In survival analysis, "time" generally refers to **tenure** rather than actual calendar time.

The "event" is some specific outcome of interest:

- Customer cancel service
- Customer make another purchase
- Patient develops disease

Logistic regression: "Did it happen?"

Survival analysis: "How long did it take to happen?"

Numeric Target – Linear Regression?

Biggest problem with using OLS for time-to-event data is **censoring** – for some observations, the event may never occur (or hasn't occurred yet).

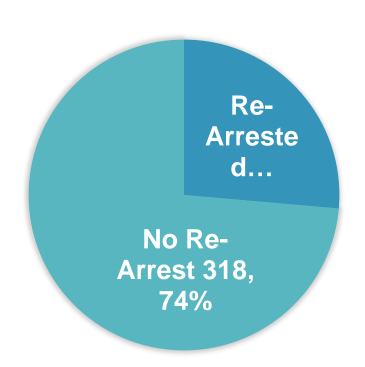
Other problems with OLS:

- Tenure is always positive problem satisfying normality assumption
- Risk of failure may change over time

Maryland Recidivism Data Set

Study from 1970's following men for one year after being released from Maryland state prisons

Of the 432 men, 114 were re-arrested within one year



Data Structure

In survival analysis, the target variables is actually two pieces – one continuous and one categorical:

- 1. Time: the tenure for an observation (continuous)
- 2. Event/status: At the end of that time, what happened? (categorical)

Observation	Time (Week)	Status (Re-Arrest?)
1	20	1
2	17	1
3	25	1
4	52	0
5	52	0
6	52	0
7	23	1

Data Structure

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5	52	0
6	52	0
7	23	1

Maryland Recidivism Data Set

Model the association between various factors and length of time before re-arrest.

Target:

- week: week of arrest week = 52 if censored (not arrested)
- arrest: indicator for arrest (1 = yes, 0 = no)

Maryland Recidivism Data Set

Model the association between various factors and length of time before re-arrest.

Predictors:

- fin: received financial aid upon release (1 = yes, 0 = no)
- age: age at time of release (years)
- race: indicator for *Black* (1 = yes, 0 = no)
- wexp: indicator of prior work full-time work experience prior to incarceration (1 = yes, 0 = no)
- mar: married at time of release (1 = yes, 0 = no)
- paro: released on parole (1 = yes, 0 = no)
- prio: number of prior convictions

Time & censoring

The Meaning of Time

Survival analysis has a few things that set it apart from any other statistical modeling you've seen in this program so far:

- Time to an event
- Censoring

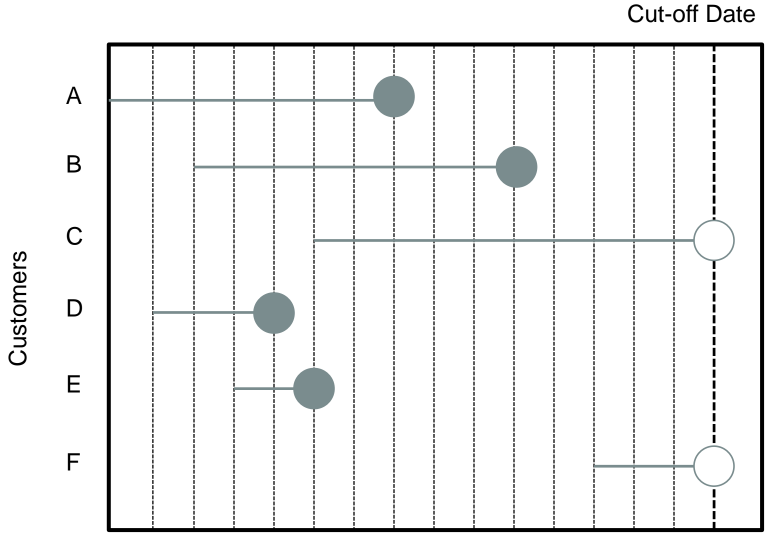
Just like most models, survival analysis depends on certain assumptions.

When Does Time Start?

Create an artificial world in which everyone "starts" at the same time.

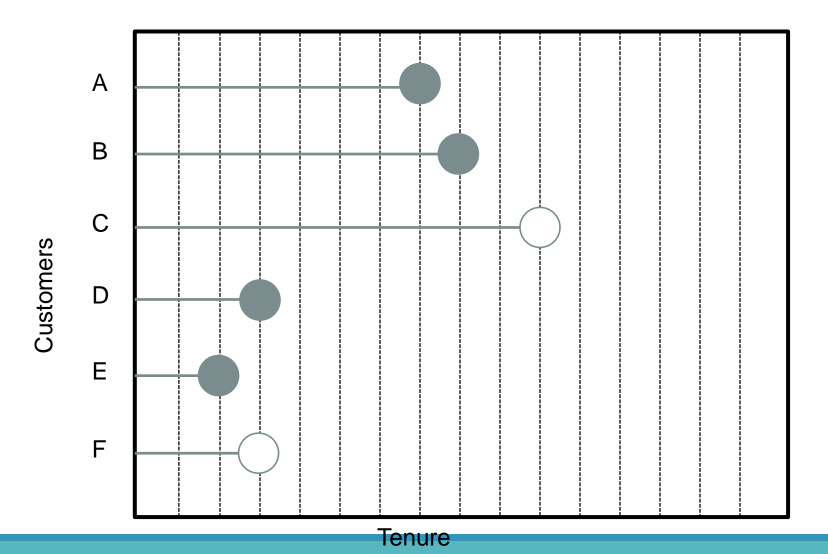
Not actually interested in time, but tenure.

Time vs. Tenure



Calendar Time

Time vs. Tenure



When Does Time Start?

Create an artificial world in which everyone "starts" at the same time.

Not actually interested in time, but tenure.

Choice of starting point isn't always obvious:

- Time since exposure to disease vs. developing disease
- Time since diagnosis vs. surgery vs. treatment
- Time since another event
- Time until car dies from production vs. purchase vs. last repair

Observed Time & Status

Interested in time to event *T*, but we can not observe this for all observations.

These observations are **censored**.

The "time" we actually observe for each observation i is the minimum between T_i and C_i :

- \circ T_i is the time until the event
- C_i is the censoring time

Need another "status" variable to tell us which one we observe for each observation.

Data Structure

In survival analysis, the target variables is actually two pieces – one continuous and one categorical:

- 1. Time: the tenure for an observation (continuous)
- 2. Event/status: At the end of that time, what happened? (categorical)

Observation	Time (Week)	Status (Re-Arrest?)
1	20	1
2	17	1
3	25	1
4	52	0
5	52	0
6	52	0
7	23	1

Censored IS NOT Missing

Do not know the actual time to event T_i for censored observations.

Do know that for some amount of time – namely, C_i – the event has not occurred.

• Provides **some** but not all information about T_i .

Censored data is **incomplete**, but not missing.

Ignoring censored observations would be falsely acting as if we know nothing about T_i .

Type I vs. Type II Censoring

Type I – there is a specific end time *c*, and any subject that hasn't had the event by time *c* is censored (most common).

Type II – time goes until a certain (pre-specified) number of events have occurred, and any subjects who haven't had the event by that time are censored.

Right, Left and Interval Censoring

If an observation is **right censored**, then T > c. This is what normally happens.

Example:

Clinical trial ends, and patient is still alive.

If an observation is **left censored**, then all we know is that T < c.

Example:

 Became a customer more than 3 years ago. Implemented new customer tracking system, but current customers were around before.

Right, Left and Interval Censoring

Interval censoring combines both right and left censoring where a < T < b.

Example:

 Person tests negative during appointment at a, but positive during appointment at b. So time developing disease is between a and b.

Assumption for Right Censoring

Censoring is NONINFORMATIVE

 This means that patients who are censored should have the same future risk for the occurrence of the event of interest, conditional on exposure, as those who continue to be followed ("assumption of independent censoring")

Survival Function

Summarizing Survival Data

Interested in the event time T.

Unique challenges to summarizing information about *T*:

- Are means/variances useful for skewed distributions such as time?
- In the presence of censoring, can we even estimate means and variances without actually knowing all the true values of T?

Survival analysis described in **two** major quantities:

- Survival Function
- Hazard Function

Survival Function

Survival function: probability of surviving **beyond** time *t*.

$$S(t) = P(T > t)$$

Properties:

- Always starts at 1 (or 100%).
- Never increases.
- Bounded below by 0 (or 0%).

Survival curves used to be the only method in survival analysis (1958: Edward L. Kaplan and Paul Meier).

Kaplan-Meier Method

Estimating the survival function:

 Want to estimate the proportion of individuals "still alive" at any given time t.

$$\hat{S}(t) = \prod_{k \le t} \left(1 - \frac{d_k}{r_k} \right) \qquad \text{# events occurring at time } t$$

Kaplan-Meier Method

Estimating the survival function:

 Want to estimate the proportion of individuals "still alive" at any given time t.

$$\hat{S}(t) = \prod_{k \le t} \left(1 - \frac{d_k}{r_k} \right)$$
events occurring at time t
before time t (**risk set**)

Kaplan-Meier Method

Estimating the survival function:

 Want to estimate the proportion of individuals "still alive" at any given time t.

$$\hat{S}(t) = \prod_{k \le t} \left(1 - \frac{d_k}{r_k} \right)$$
events occurring at time t
observations available right before time t (**risk set**)

Kaplan and Meier showed it was the maximum likelihood estimate for the nonparametric estimation of the survival curve.

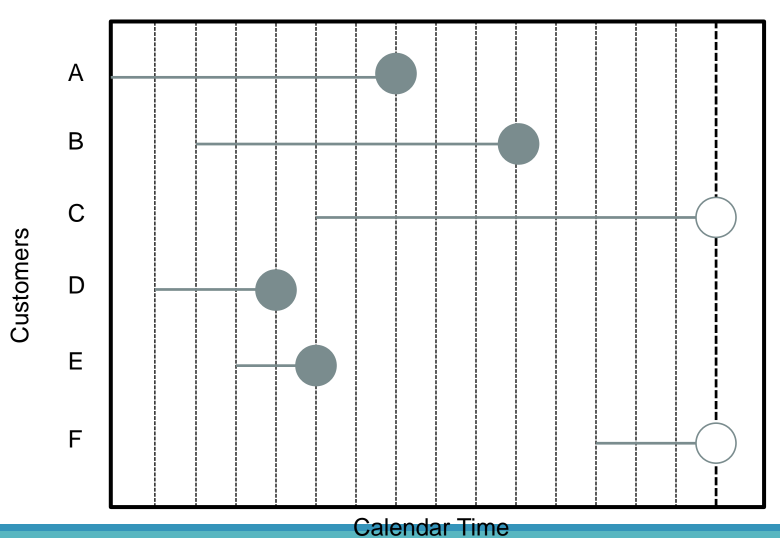
At the beginning (t = 0), all observations are at risk $(r_0 = n)$ and no events have occurred $(d_0 = 0)$:

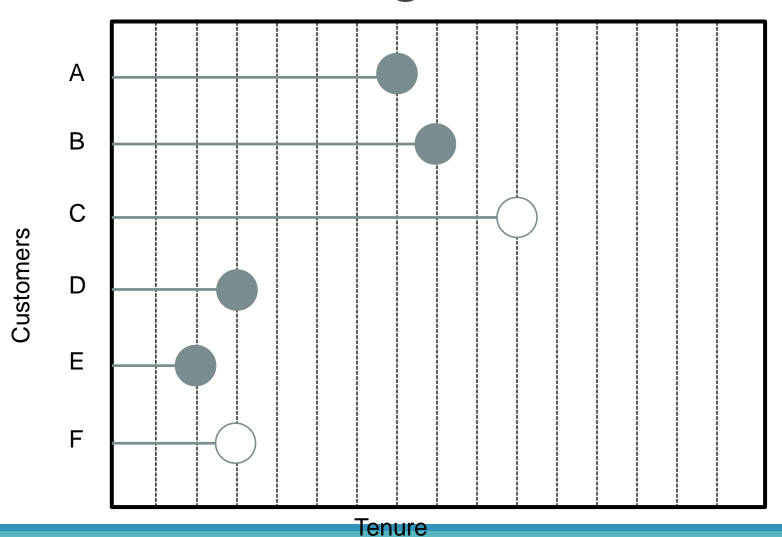
$$\hat{S}(t) = \prod_{k \le t} \left(1 - \frac{d_k}{r_k} \right) = \left(1 - \frac{0}{n} \right) = 1$$

Start with S(0) = 1 and step forward in time, reducing $\hat{S}(t)$ by a factor of $\left(1 - \frac{d_t}{r_t}\right)$ at each time period:

$$\hat{S}(1) = S(0) \times \left(1 - \frac{d_1}{r_1}\right)$$

$$\hat{S}(2) = \hat{S}(1) \times \left(1 - \frac{d_2}{r_2}\right)$$





Time = 0:

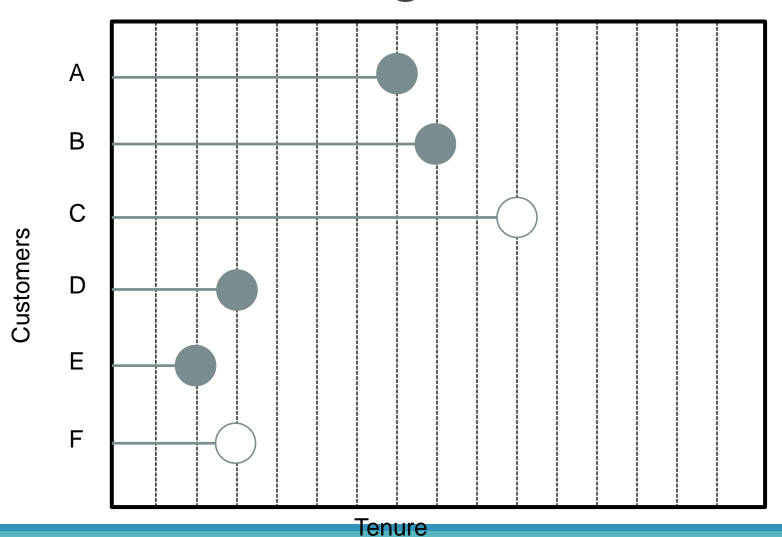
$$\hat{S}(0) = 1$$

Time = 1:

$$\hat{S}(1) = S(0) \times \left(1 - \frac{0}{6}\right) = 1$$

Time = 2:

$$\hat{S}(2) = \hat{S}(1) \times \left(1 - \frac{1}{6}\right) = 0.8333$$



Time = 3:

$$\hat{S}(3) = \hat{S}(2) \times \left(1 - \frac{1}{5}\right) = 0.833 \times 0.80 = 0.667$$

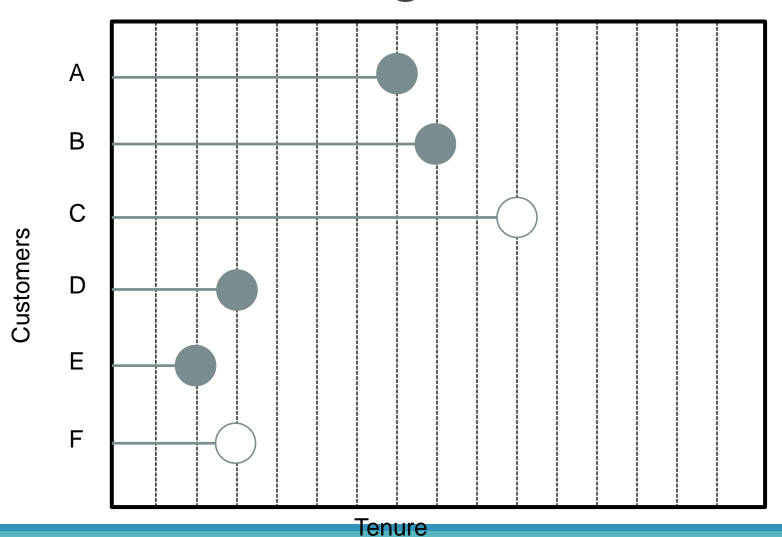
Time = 4:

$$\hat{S}(4) = \hat{S}(3) \times \left(1 - \frac{0}{3}\right) = 0.667$$

Time = 5:

$$\hat{S}(5) = \hat{S}(4) \times \left(1 - \frac{0}{3}\right) = 0.667$$

Calculating K-M Estimate



Calculating K-M Estimate

Time = 6:

$$\hat{S}(6) = \hat{S}(5) \times \left(1 - \frac{0}{3}\right) = 0.667$$

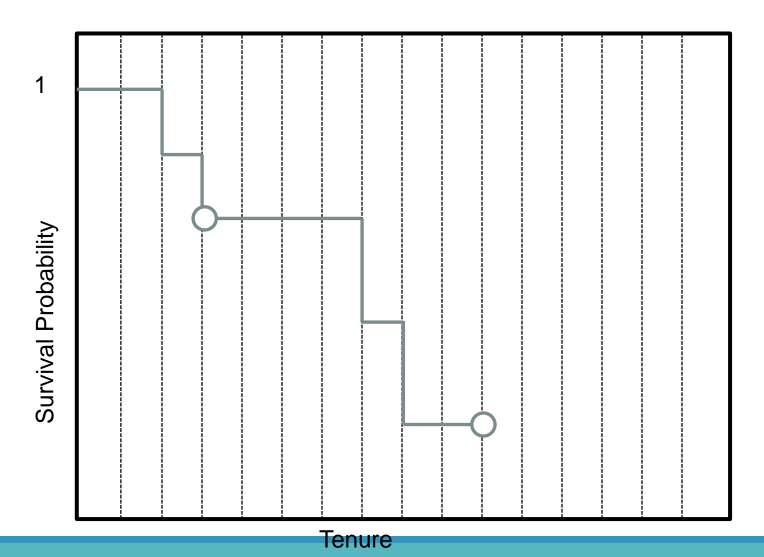
Time = 7:

$$\hat{S}(7) = \hat{S}(6) \times \left(1 - \frac{1}{3}\right) = 0.667 \times 0.667 = 0.444$$

Time = 8:

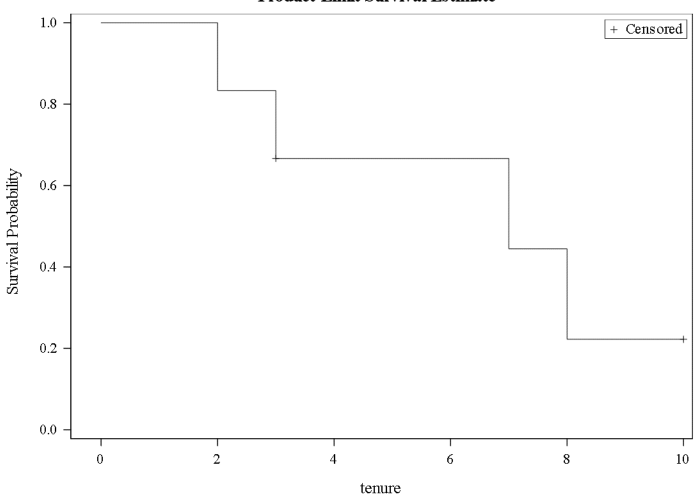
$$\hat{S}(8) = \hat{S}(7) \times \left(1 - \frac{1}{2}\right) = 0.444 \times 0.5 = 0.222$$

Visualizing K-M Estimate



Visualizing K-M Estimate

Product-Limit Survival Estimate



Summary Statistics

Due to censoring, the mean is difficult to estimate, but the median is still valid as long as the event occurs for at least half of the sample.

The median (also called **half-life**) is the time t that $\hat{S}(t)$ drops below 0.5 (or 50%).

Half-life interpretation: 50% of observations survive beyond time *t*.

Survival Function – R

Data set: tenure censored

simple.s=Surv(time=simple\$tenure,event=simple\$censored)

simple.s [1] 7 8 10+ 3 2 3+

Kaplan-Meier

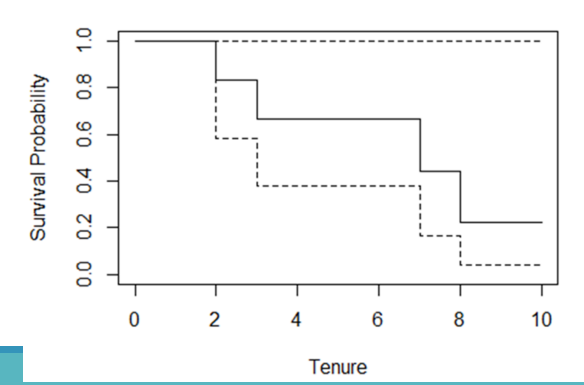
```
simple_km=survfit(Surv(time = tenure, event = censored)~1, data = simple) summary(simple_km)
```

time	n.risl	k n.eve	ent survival	std.err	lower 95% CI
2	6	1	0.833	0.152	0.5827
3	5	1	0.667	0.192	0.3786
7	3	1	0.444	0.222	0.1668
8	2	1	0.222	0.192	0.0407

Survival Curve

```
plot(simple_km, main = "Survival Function", xlab = "Tenure",
    ylab = "Survival Probability")
```

Survival Function

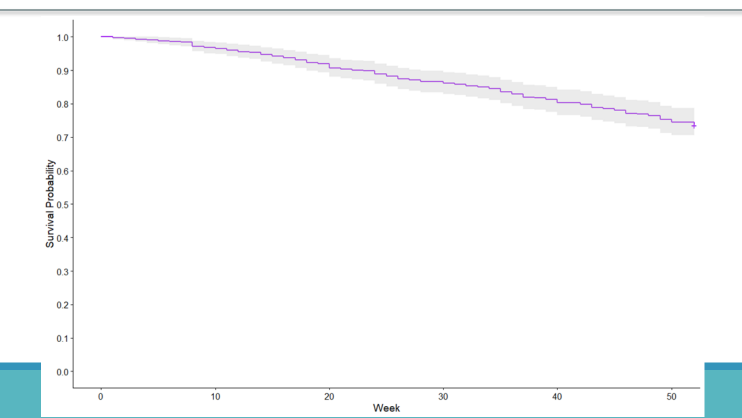


K-M for Recid

```
Recid.fit = survfit(Surv(time = week, event = arrest ~ 1,
data = recid)
summary(recid.fit)
```

```
time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
           432
##
       1
                          0.998 0.00231
                                               0.993
                                                            1.000
                          0.995 0.00327
           431
                                                            1,000
##
                                               0.989
##
           430
                          0.993 0.00400
                                               0.985
                                                            1.000
                     1 0.991 0.00461
##
           429
                                               0.982
                                                            1,000
##
         428
                     1 0.988 0.00515
                                               0.978
                                                           0.999
##
         427
                     1 0.986 0.00563
                                               0.975
                                                           0.997
                     1 0.984 0.00607
##
          426
                                               0.972
                                                           0.996
##
           425
                          0.972 0.00791
                                               0.957
                                                            0.988
                                                            0.779
     52
           322
                          0.736 0.02121
                                               0.696
##
```

Survival Function – R

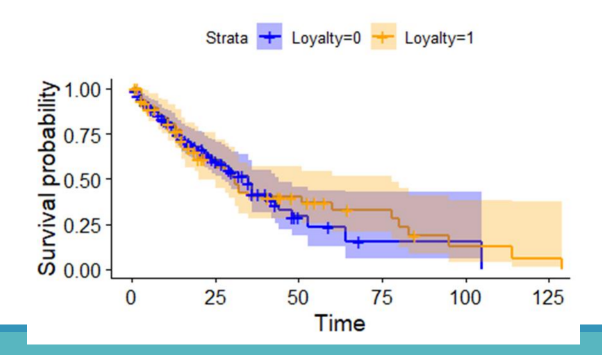


Stratified Analysis

Stratified Analysis

Can also create separate/stratified curves by group.

Different curves result in different estimates for each group.



Stratified Analysis

R provides 2 tests that each have the same null hypothesis – all survival curves are **equal** (alternative is that at least one curve is different).

- 1. Log-rank test (developed by Mantel-Haenszel)
- 2. Wilcoxon test

Log-rank Tests

The **log-rank test** combines all the information from the K-M estimate at times where events occur.

Similar to the Mantel-Haenszel tests for association from categorical data.

At time t	# Events	# Non-events	Total
Group 1	$d_{1,t}$	$r_{1,t}-d_{1,t}$	$r_{1,t}$
Group 2	$d_{2,t}$	$r_{2,t}-d_{2,t}$	$r_{2,t}$
Total	d_t	$r_t - d_t$	r_t

Comparing Survival Function

Log-Rank test:

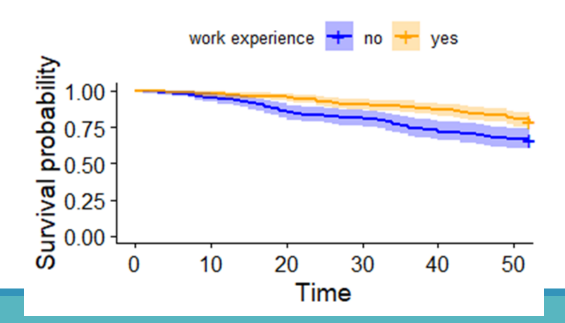
For each group, calculate expected events and compare to observe events ($(O-E)^2/E$. This is a χ^2 statistic with k-1 df (k is the number of groups being compared!). This is the statistic when we set "Rho = 0")

Wilcoxon test (places larger emphasis on earlier event times):

Similar to Log-Rank test except that we now use weights. This is what happens when "Rho = 1").

Stratified Analysis – R

```
Recid.KP = survfit(Surv(week, arrest) ~ wexp,data=recid)
ggsurvplot(Recid.KP,data=recid,palette = c("blue","orange"),conf.int = T,
legend.title = "work experience", legend.labs = c("no", "yes"))
```

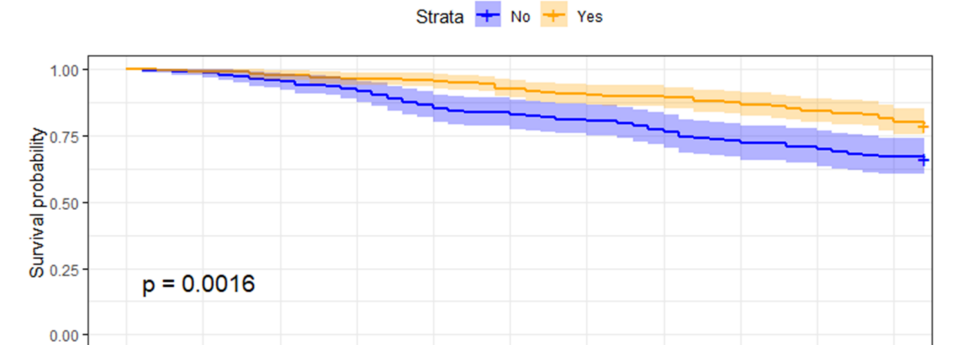


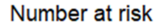
Stratified Analysis – R

```
survdiff(Surv(week, arrest) ~ wexp, data=recid, rho=0)
```

```
## N Observed Expected (0-E)^2/E (0-E)^2/V ## wexp=0 185 62 45.6 5.91 9.91 ## wexp=1 247 52 68.4 3.94 9.91 ## Chisq= 9.9 on 1 degrees of freedom, p= 0.002
```

```
ggsurvplot(Recid.KP, data = recid, size = 1, palette =c("blue","orange"),
  conf.int = TRUE, pval = TRUE, risk.table = TRUE, risk.table.col =
"wexp", legend.labs =c("No", "Yes"), risk.table.height = 0.25,
  ggtheme = theme_bw() )
```







Time

Hazard function

Hazard Function

In survival analysis we also use the **hazard function** to summarize the data.

There are two common types of hazard functions:

1. Hazard Probabilities:

$$h(t) = P(t < T < t + 1 \mid T > t)$$

2. Hazard Rates:

$$h(t) = \lim_{\Delta t \to 0} \frac{P(t < T < t + \Delta t \mid T > t)}{\Delta t}$$

Hazard Function

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There are two common types of hazard functions:

1. Hazard Probabilities:

$$(h(t)) = P(t < T < t + 1 | T > t)$$

Hazard Rates:

$$h(t) = \lim_{\Delta t \to 0} \frac{P(t < T < t + \Delta t \mid T > t)}{\Delta t}$$

Both are denoted the same way in different texts!

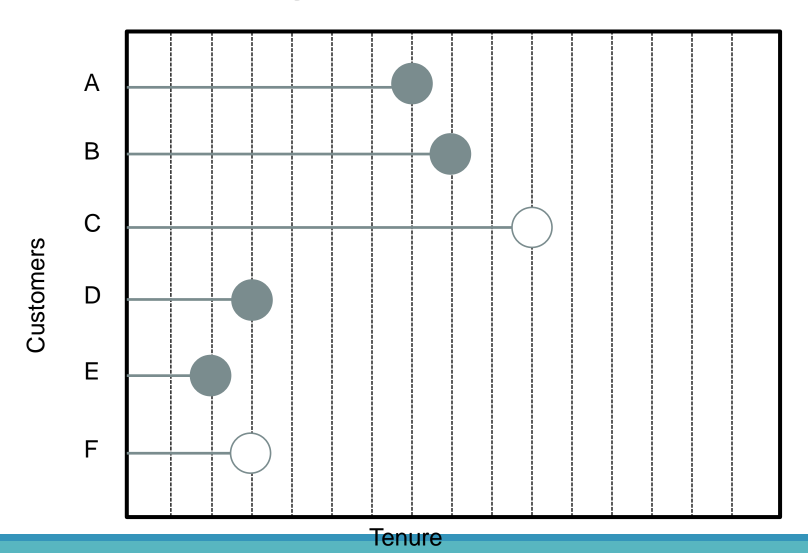
Hazard Probabilities

Hazard probabilities are very useful and common in business settings.

Example:

- A customer has survived for a certain length of time,
 so the customer's tenure is t.
- What is the probability that the customer leaves before t+1?

$$h(t) = P(t < T < t + 1 \mid T > t) = \frac{a_t}{r_t}$$



Time = 0:

$$h(0) = 0$$

Time
$$= 1$$
:

$$h(1) = \frac{0}{6} = 0$$

Time
$$= 2$$
:

$$h(2) = \frac{1}{6} = 0.1667$$

Time
$$= 3$$
:

$$h(3) = \frac{1}{5} = 0.2$$

Time
$$= 4$$
:

$$h(4) = \frac{0}{3} = 0$$

Time
$$= 5$$
:

$$h(5) = \frac{0}{3} = 0$$

Time = 3:

$$h(3) = \frac{1}{5} = 0.2$$

$$h(3) = \frac{1}{5} = 0.2$$
 OR $h(3) = \frac{1}{4.5} = 0.222$

Time = 4:

$$h(4) = \frac{0}{3} = 0$$

Time = 5:

$$h(5) = \frac{0}{3} = 0$$

Time = 6:

$$h(6) = \frac{0}{3} = 0$$

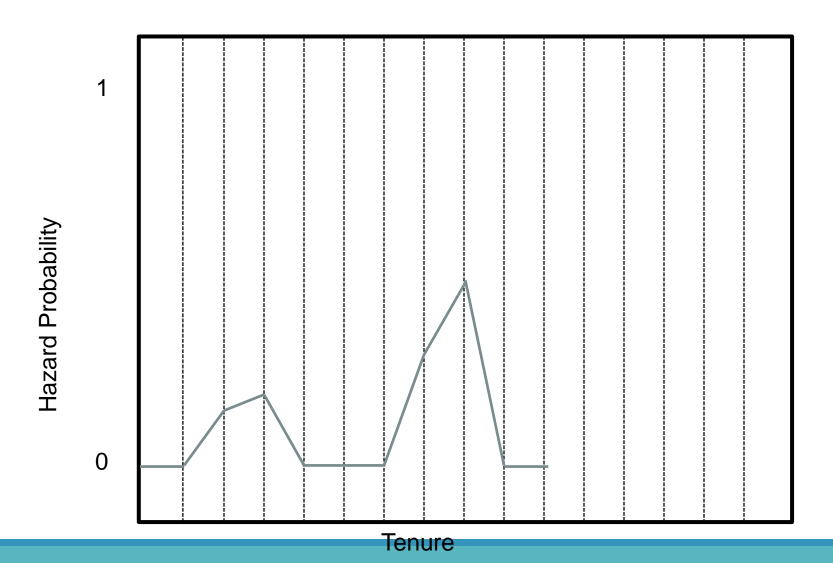
Time = 7:

$$h(7) = \frac{1}{3} = 0.333$$

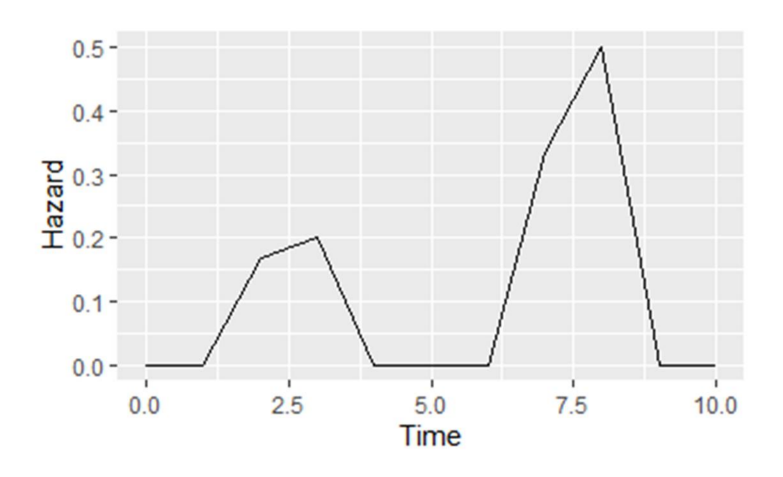
Time = 8:

$$h(8) = \frac{1}{2} = 0.5$$

Visualizing Hazard Probabilities



Visualizing Hazard Probabilities



Hazard Rates

Hazard rates have a slightly different interpretation than the hazard probabilities because they are limits of conditional probabilities.

$$h(t) = \lim_{\Delta t \to 0} \frac{P(t < T < t + \Delta t \mid T > t)}{\Delta t}$$

The hazard rate is the **instantaneous event rate** for the risk set at time *t*.

• Given survival up until time t, it is the rate of events in the interval $[t, t + \Delta t]$.

Hazard Rates

Hazard rates have a slightly different interpretation than the hazard probabilities because they are limits of conditional probabilities.

$$h(t) = \lim_{\Delta t \to 0} \frac{P(t < T < t + \Delta t \mid T > t)}{\Delta t}$$

The hazard rate is the **instantaneous event rate** for the risk set at time *t*.

Bounded below by 0, but are NOT bounded above by 1!

Hazard Probability

Hazard probability can be interpreted as "Assuming the event has not occurred yet, this is the probability the event occurs by the next time point".

Examples:

- Say the hazard probability for some point in time for contracting a sinus infection is 0.2 with a time measured in months.
- "Assuming the individual has not contracted a sinus infection, the probability s/he contracts a sinus infection by next month is 0.2."

Hazard Rates – Inverse

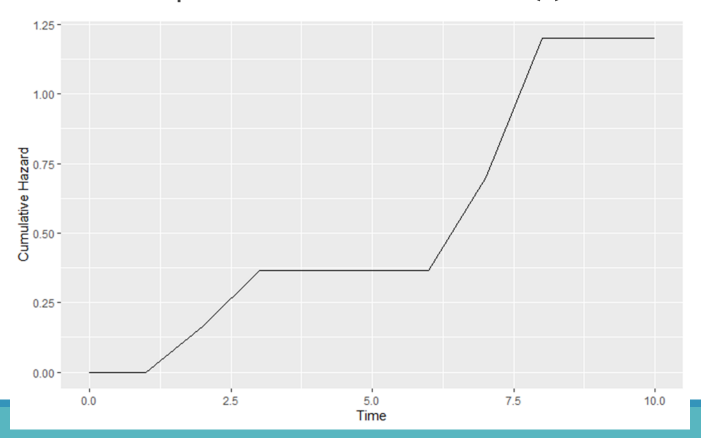
The interpretation of the inverse of the hazard function is the length of time before the next occurrence.

Examples:

- Hazard for some point in time for contracting a sinus infection is 0.2 with a time measured in months.
- "Assuming an individual has not yet contracted a sinus infection, we expect this individual to go 5 (= 1/0.2) months before contracting a sinus infection (assuming the hazard stays constant)."

Cumulative Hazard Probability

The cumulative hazard probability is just the total hazard rate up until time t – denoted $\Lambda(t)$.



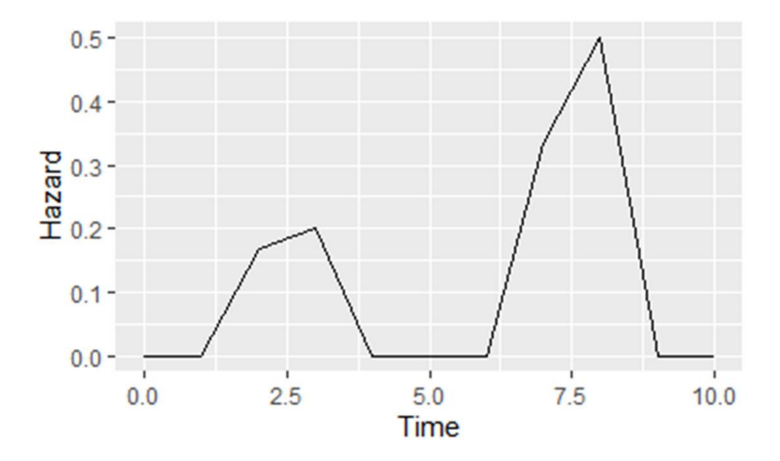
summary(simple_km)

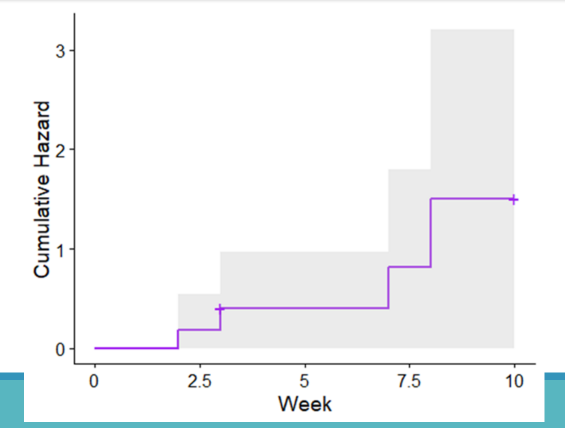
```
## Call: survfit(formula = Surv(time = tenure, event = (censored == 0))
     1, data = simple)
##
##
   time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
                    0.833 0.152
                                     0.5827
##
## 3 5
                 1 0.667 0.192 0.3786
                                                   1
## 7 3 1 0.444 0.222 0.1668
                 1 0.222 0.192
                                    0.0407
##
```

```
simple_km$hp = simple_km$n.event/simple_km$n.risk
print(simple_km$hp)
```

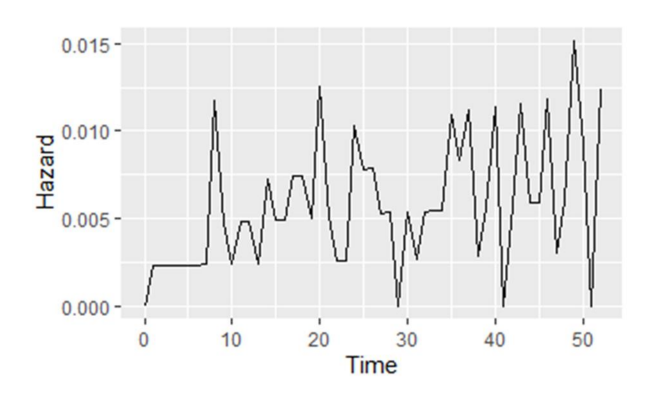
```
## [1] 0.1666667 0.2000000 0.3333333 0.5000000 0.0000000
```

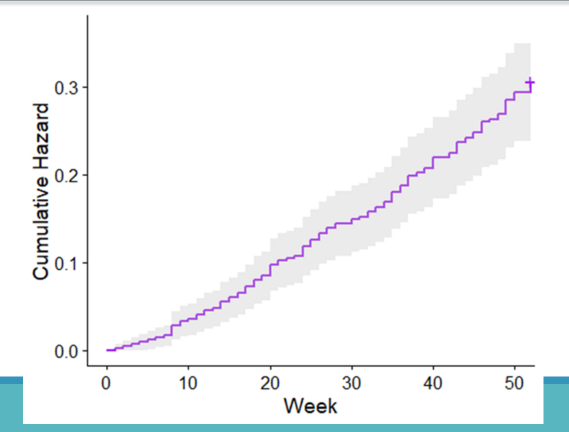
```
h= simple_km$n.event/simple_km$n.risk
index.h=rep(0,length=(max(simple$tenure)+1)) #Need to add 0
index.h[(simple_km$time)+1]=h #Because of 0
haz.plot=data.frame(cbind(seq(0,max(simple$tenure)), index.h))
colnames(haz.plot)=c("Time","Hazard")
ggplot(haz.plot,aes(x=Time,y=Hazard))+geom_line()
```





Hazard Functions – Recid data





Survival and Hazard Relationship

The survival, hazard, and cumulative hazard functions are all directly related:

$$\circ S(t) = e^{-\Lambda(t)}$$

$$b(t) = -\frac{d}{dt}\log S(t) = \frac{f(t)}{S(t)}$$

These three quantities are all different ways of describing the same distribution; if you know one of them, you can compute the others.