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# BST 210 Lab: Week 12-13

## Survival Analysis

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### Introduction to Survival

In some studies, the outcome of interest is the time to event (survival time to failure). We can perform survival analysis to estimate the distribution of survival times, test equality of two or more survival distributions, or estimate the relationships between predictors and the survival time while control for other covariates.

Let  $T_i$  be the time to event for patient  $i$ . There exist four interrelated functions which categorize the distribution of  $T_i$ :

1. **Survival function:**

$$S(t) = P(T_i > t) = 1 - P(T_i \leq t) = 1 - F(t)$$

This is the probability of not having event until time  $t$ . Note  $S(0) = 1$ , and  $S(t) \in [0, 1], t \in [0, \infty)$ .

2. **Cumulative distribution function:**  $F(t) = P(T_i \leq t) = 1 - S(t)$ .

3. **Probability density function:**  $f(t) = F'(t) = -S'(t)$ .

4. **Hazard function:**

$$h(t) = \lim_{\Delta t \rightarrow 0} \frac{P(t \leq T_i \leq t + \Delta t | T_i \geq t)}{\Delta t} = \frac{f(t)}{S(t)}$$

This instantaneous rate of failure at time  $t$ , given that a subject has survived to time  $t$ . Note that  $h(t) \geq 0$  but does not have an upper bound because it is not a probability.<sup>1</sup>

### Censoring

Theoretically, there is no reason why we cannot use analyses/tests/regression methods we learned previously in this class for time to event outcomes, since it is a continuous outcome and  $t$ . The unique feature of survival analysis (and why there is a field dedicated to this type of data) is how it handles censorship.

A patient is **censored** if we do not observe the time of their outcome. This could be due to many different factors but is essentially a missing data problem (we cannot observe  $T$ ). Unlike most missing data problems, however, survival data is unique in that the analyses we have developed can incorporate information from censored patients. Let's denote the censoring time as  $C$ .

However, survival analysis is dependent on the assumption of **noninformative censoring**, which says that the event of censoring must be independent of the time of event.

**Question:** What situations would the noninformative censoring assumption be invalid?

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<sup>1</sup>Hernán, Miguel A. "The hazards of hazard ratios." *Epidemiology* (Cambridge, Mass.) 21.1 (2010): 13.

Censoring comes in three flavors:

1. **Left censoring:** A patient is not observed before a certain time point, and may have experienced the event already, (i.e.  $T < C$ ). This time point is usually their recruitment into the study.
2. **Interval censoring:** A patient experiences an event within an unobserved interval, but the exact time within the interval is unknown. Most survival data suffers from interval censoring, simply because it is difficult to collect information on the exact time an event occurs. The severity of interval censoring depends on the frequency of follow-up and effort made to find supplemental information (death certificates etc.) Commonly, statisticians can assume events all occur right after the last follow-up, right before the current follow-up or in the middle. This does not change the outcome of the analysis with regards to hypothesis testing (will change the outcome with regards to estimated survival time, especially if the intervals are large), as long as the assumptions are consistent.
3. **Right censoring:** A patient is not observed experiencing the event, but may have the event after their observation period is over (i.e.  $T > C$ ). Right censoring is the most common "problem" in survival analysis because it cannot be designed or assumed away and must be dealt with through analysis. Common causes of right censoring include dropout, study conclusion and death.

## Nonparametric survival analysis: Kaplan-Meier estimator

The **Kaplan-Meier** estimator (aka **product-limit** estimator) is a **non-parametric** estimator for survival curves. Using the K-M estimator, we do not have to make any distributional assumptions about survival times. This method is intuitive, but has limitations in that we cannot account for covariate data.

Assume that in your data, there were  $m$  distinct times when events occurred, which we call  $t_1 < t_2 < \dots < t_m$ . Then we can go back and look at each time  $t_j$  to see the number of events that occurred  $d_j$ , as well as the total number of people present in the data at that time  $n_j$ . We say these people are in the 'risk set' at time  $t_j$ . Don't forget, the people who experienced events are still included in  $n_j$ , because they were at risk when they experienced the event!

Then, given that someone is in the risk set at time  $t_j$ , their probability of experiencing the event at that time is simply  $\hat{p}_j = \frac{d_j}{n_j}$ , which in turn means that given a person remained in the risk set until time  $t_j$ , their probability of survival beyond time  $t_j$  is  $\hat{q}_j = 1 - \frac{d_j}{n_j} = \frac{n_j - d_j}{n_j}$ .

The Kaplan-Meier estimator simply multiplies these running conditional survival probabilities together up through each event time. The empirical probabilities at each time  $t$  follow the formula

$$\hat{S}(t) = \prod_{j=1}^k \hat{q}_j = \prod_{j=1}^k \frac{n_j - d_j}{n_j}, \quad t_k \leq t < t_{k+1}$$

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

Let's do an example: let the survival times of 20 patients be: 1, 1+, 2, 3, 4, 4, 4, 5, 5+, 5+, 6, 6, 6+, 7, 7, 7, 8, 8+, 8+, 9. Where '+' indicates censoring (if an observation is designated '1+' it means that they had their event after time period 1). Fill out the following table and graph this curve by hand.

<b>k</b>	<b>Risk Set <math>n_k</math></b>	<b>Events <math>d_k</math></b>	<b>Conditional Survival <math>\hat{q}_k = 1 - \frac{d_k}{n_k}</math></b>	<b>Unconditional Survival <math>\hat{S}(t_k) = \prod_{j=1}^k \hat{q}_j</math></b>
<b>1</b>	20	1	19/20	(19/20) = 0.95
<b>2</b>	18	1	17/18	(19/20)(17/18) = 0.897
<b>3</b>	17	1	16/17	(19/20)(17/18)(16/17) = 0.844
<b>4</b>	16	3	13/16	(19/20)...(13/16)=0.686
<b>5</b>	13	1	12/13	(19/20)...(12/13)=0.633
<b>6</b>	10	2	8/10	(19/20)...(8/10)=0.507
<b>7</b>	7	3	4/7	(19/20)...(4/7)=0.290
<b>8</b>	4	1	3/4	(19/20)...(3/4)=0.217
<b>9</b>	1	1	0	0

Draw a survival curve based on this table:

## Survival Analysis in Stata

The Sorbinil Retinopathy Trial (SRT) was a randomized, double-blind, placebo controlled trial testing whether the aldose reductase inhibitor, sorbinil, would reduce rates of progression in complications of diabetes, with a primary focus on diabetic retinopathy (Arch Ophthalmol 1990; 108:1234-1244). Between August 1983 and October 1986, the SRT randomized 497 type I diabetic patients at 11 clinical centers. The primary outcome of the trial was worsening of a patient's retinopathy by two or more steps, based on grading fundus photographs. The file `srt.dat` contains data from 478 randomized participants with some follow-up and complete information on important baseline covariates.

Variables on the file `srt.dat` are: `id` (a subject id), `sorb` (randomized treatment assignment, 1=sorbinil, 0=placebo), `tgh` (total glycosylated hemoglobin in percent), `dur` (duration of diabetes in years since diagnosis), `sex` (2=female, 1=male), `fup` (duration of follow-up in years until progression of diabetic retinopathy or end of follow-up), and `status` (1=diabetic retinopathy progressed, 0=no progression).

Read the data into Stata, prepare them for survival analysis, and compare the distributions of patient characteristics by treatment group. Note that women were less likely to be included because sorbinil was teratogenic.

```
import delimited srt.csv

*Summarize by treatment group
sort sorb
by sorb: su
-----
-> sorb = 0
```

Variable	Obs	Mean	Std. Dev.	Min	Max
id	241	3.46e+08	9.58e+07	1.80e+08	6.13e+08
sorb	241	0	0	0	0
tgh	241	11.81743	1.920966	8.35	18.1
dur	241	6.875259	3.676774	1	16
sex	241	1.261411	.4403177	1	2
fup	241	2.974056	.8576834	.9747	4.4983
status	241	.2738589	.4468655	0	1

```
-----
-> sorb = 1
```

Variable	Obs	Mean	Std. Dev.	Min	Max
id	237	3.51e+08	1.00e+08	1.80e+08	6.13e+08
sorb	237	1	0	1	1
tgh	237	11.93734	2.063994	8.2	23.05
dur	237	6.797732	3.346496	1.375	15.9375
sex	237	1.240506	.4282955	1	2
fup	237	2.932503	.9030455	.9692	4.5366
status	237	.2700422	.444921	0	1

Note that we need to create a new variable to denote which observations are censored:

```
*Set up survival analysis in stata
stset fup, id(id) failure(status)
```

We can get compute the  $\hat{S}(t)$ , its standard error (Greenwood's method) and its confidence interval (complementary log-log transformation). If you use Greenwood's method to compute the confidence interval, make sure you truncate the results so that the range is within  $[0, 1]$ .

Let's first get the K-M estimates for the males in Stata:

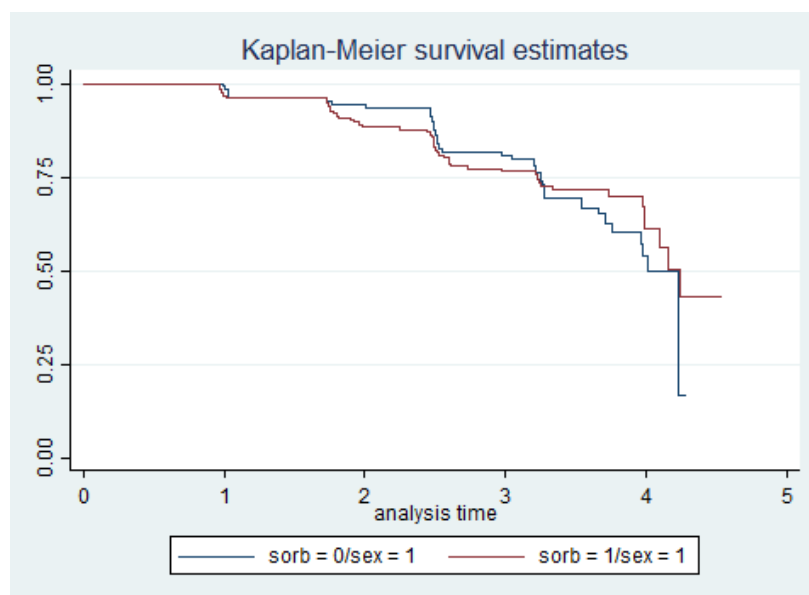
```
. sts list if sex==1, by(sorb sex)

      failure _d:  status
analysis time _t:  fup
              id:  id
```

Time	Beg. Total	Fail	Net Lost	Survivor Function	Std. Error	[95% Conf. Int.]	
-----							
sorb=0 sex=1							
.9747	178	0	1	1.0000	.	.	.
.9938	177	1	0	0.9944	0.0056	0.9606	0.9992
. . .							
4.23	2	1	0	0.1659	0.1375	0.0118	0.4833
4.29	1	0	1	0.1659	0.1375	0.0118	0.4833
sorb=1 sex=1							
.9692	180	1	0	0.9944	0.0055	0.9612	0.9992
.9719	179	1	0	0.9889	0.0078	0.9563	0.9972
. . .							
4.501	2	0	1	0.4321	0.0985	0.2395	0.6110
4.537	1	0	1	0.4321	0.0985	0.2395	0.6110
-----							

And we can plot the K-M curve for sorbinil and placebo groups separately (only for males):

```
*Plot Kaplan-Meier estimate
sts graph if sex==1, by(sorb sex)
```



**Question:** What conclusion can you draw from these two survival curves?

## Testing differences: log-rank test

The **log-rank test** is a non-parametric test of the difference in survival curves:

$$H_0 : S_1(t) = S_2(t) \text{ for all } t$$

$$H_1 : S_1(t) \neq S_2(t) \text{ for at least one value of } t$$

It is a generalization of the Mantel-Haenszel test and requires the following steps:

1. Create a 2x2 table for each observed failure time  $t_k, k = 1, 2, \dots, m$ , where  $m$  is the number of distinct failure times.
2. At each time point  $t_k$ , note the number of events ( $d_{jk}$ ) and patients in the risk set ( $n_{jk}$ ) for each group  $j = 1, 2$ . Let  $d_k$  be the total number of events at  $t_k$  and  $n_k$  be the total number of patients in the risk set.
3. Calculate  $O = \sum_{k=1}^m d_{1k}$ ,  $E = \sum_{k=1}^m \frac{n_{1k}d_k}{n_k}$  and  $V = \sum_{k=1}^m \frac{n_{1k}n_{2k}d_k(n_k - d_k)}{n_k^2}$ .
4. Compare  $\frac{(O-E)^2}{V}$  to a chi-square distribution with 1 degree of freedom ( $\chi_1^2$ ).

The restrictions of this test are that  $V$  need to be larger than 5. Also, the test is the most powerful when the proportional hazards assumption holds (i.e. the survival/hazard curves of the two groups do not cross, with  $h_2(t)/h_1(t) = c$ ,  $c$  is some constant). If this assumption is violated, the test is still valid but became underpowered.

We will perform an log-rank test on our Stata dataset as an example, but keep in mind that the survival curves we are comparing do cross.

```
. sts test sorb if sex==1
```

```
      failure _d:  status
analysis time _t:  fup
              id:  id
```

Log-rank test for equality of survivor functions

sorb		Events observed	Events expected
0		50	47.49
1		49	51.51
Total		99	99.00

```
chi2(1) =      0.26
Pr>chi2 =      0.6108
```

Also, you can perform stratified log-rank tests to adjust for sex:

```
. sts test sorb, strata(sex)
```

```
      failure _d:  status
analysis time _t:  fup
              id:  id
```

Stratified log-rank test for equality of survivor functions

sorb		Events observed	Events expected(*)
0		66	64.32
1		64	65.68
Total		130	130.00

(\*) sum over calculations within sex

```
chi2(1) =      0.09
Pr>chi2 =      0.7672
```

## Parametric survival analysis

Kaplan-Meier survival curves and Log-Rank test compares the survival data in two groups using completely non-parametric methods. But what if we wanted to measure the change in survival across a continuous covariate? Or across multiple covariates at once?

**Exponential survival regression model**

Let  $X_{i1}, \dots, X_{ip}$  denote  $p$  covariates for patient  $i$ . If we assume that  $T_i|X_{i1}, \dots, X_{ip}$  follows a exponential distribution with parameter  $\lambda$ :

$$f(t) = \lambda \exp(-\lambda t)$$

The corresponding hazard function is:

$$h(t) = \lambda$$

which is independent of  $t$ . Then we can build a regression model:

$$h(t|X_1, \dots, X_p) = \exp\{\beta_0 + \beta_1 X_1 + \dots + \beta_p X_p\}$$

We define the baseline hazard function as hazard function when all covariates are 0. In exponential survival regression,  $h_0 = h_0(t) = h(t|X_1 = 0, \dots, X_p = 0) = \exp(\beta_0)$ , then the model can be rewritten as:

$$h(t|X_1, \dots, X_p) = h_0 \exp\{\beta_1 X_1 + \dots + \beta_p X_p\}$$

or equivalently,

$$\log \left\{ \frac{h(t|X_1, \dots, X_p)}{h_0} \right\} = \beta_1 X_1 + \dots + \beta_p X_p$$

**Question:** Can you find any connection between this model and a Poisson regression model?



Let's fit an exponential survival regression model in Stata:

```
. streg i.sorb i.sex tgh dur, distribution(exponential)
```

failure \_d: status  
 analysis time \_t: fup  
 id: id

Iteration 0: log likelihood = -326.78183  
 Iteration 1: log likelihood = -312.36888  
 Iteration 2: log likelihood = -309.78515  
 Iteration 3: log likelihood = -309.77591  
 Iteration 4: log likelihood = -309.77591

Exponential regression -- log relative-hazard form

No. of subjects =	478	Number of obs =	478
No. of failures =	130		
Time at risk =	1411.750697		
		LR chi2(4) =	34.01
Log likelihood =	-309.77591	Prob > chi2 =	0.0000

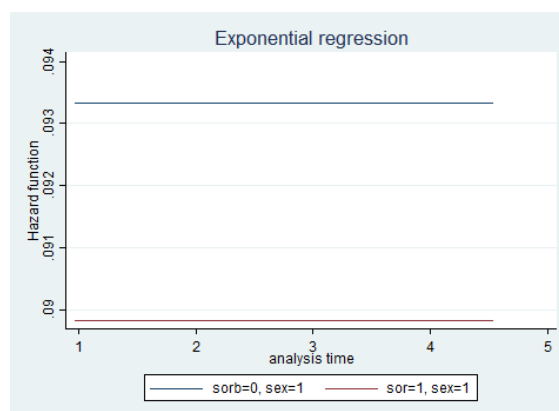
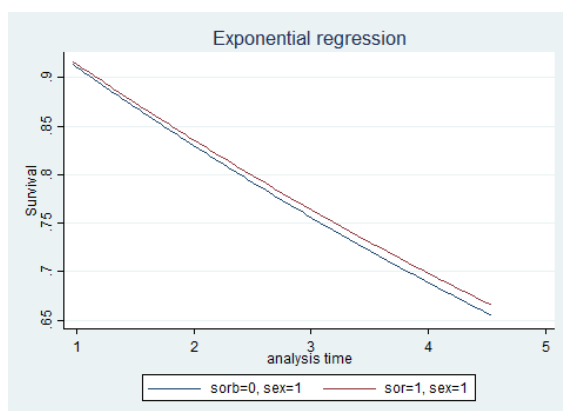
  

	_t	Haz. Ratio	Std. Err.	z	P> z	[95% Conf. Interval]
1.sorb		.9624412	.1693349	-0.22	0.828	.681728 1.358743
2.sex		.7256609	.1520984	-1.53	0.126	.481196 1.094323
tgh		1.237219	.0457712	5.75	0.000	1.150685 1.330262
dur		1.083351	.0275497	3.15	0.002	1.030678 1.138715
_cons		.0043087	.0023062	-10.18	0.000	.0015092 .012301

**Question:** Interpret the intercept and the coefficient for sorb.

Plot hazard function:

```
stcurve, survival at1(sorb=0, sex=1) at2(sor=1, sex=1)
stcurve, hazard at1(sorb=0, sex=1) at2(sor=1, sex=1)
```



**Question:** If you adjust for multiple covariates in your regression, you need to specify the values for those covariates when you plot the survival/hazard/cumulative probability functions. What values would you use?

## Weibull survival regression model

In exponential survival regression, we assumed the hazard is constant over time, which is hardly true in real life. Weibull survival regression model is one of the more flexible model where we assume  $T$  follows a Weibull distribution and the hazard function varies over time. The Weibull model can be written as:

$$h(t|X_1, \dots, X_p) = h_0(t) \exp(\beta_1 X_1 + \dots + \beta_p X_p)$$

where

$$h_0(t) = \exp(\beta_0) \gamma t^{\gamma-1} = \lambda \gamma t^{\gamma-1}$$

$\lambda$  is called the scale parameter, and  $\gamma$  is called the shape parameter. Specifically, the baseline hazard increases over time when  $\gamma > 1$ , decreases when  $0 < \gamma < 1$ , and remain constant when  $\gamma = 1$  (the model reduces to a exponential survival regression model).

```
. streg i.sorb i.sex tgh dur, distribution(weibull)

      failure _d:  status
analysis time _t:  fup
             id:  id

Fitting constant-only model:

Iteration 0:  log likelihood = -326.78183
Iteration 1:  log likelihood = -275.67984
Iteration 2:  log likelihood = -267.73983
Iteration 3:  log likelihood = -267.61483
```

```
Iteration 4:  log likelihood = -267.61481
Iteration 5:  log likelihood = -267.61481
```

Fitting full model:

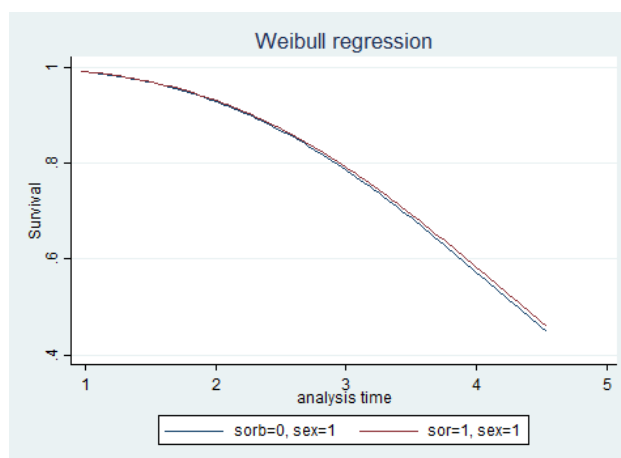
```
Iteration 0:  log likelihood = -267.61481
Iteration 1:  log likelihood = -248.51025
Iteration 2:  log likelihood = -243.26574
Iteration 3:  log likelihood = -243.21947
Iteration 4:  log likelihood = -243.21945
Iteration 5:  log likelihood = -243.21945
```

Weibull regression -- log relative-hazard form

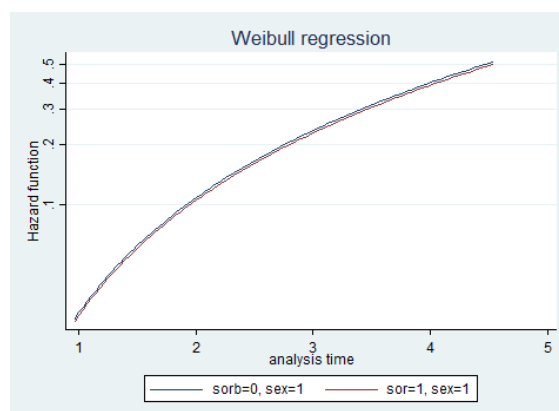
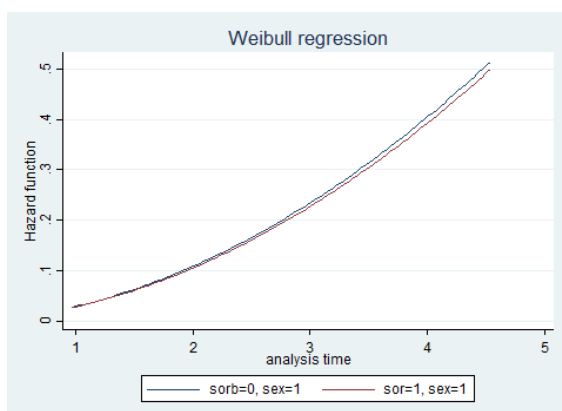
```
No. of subjects =          478          Number of obs   =          478
No. of failures =          130
Time at risk    = 1411.750697
Log likelihood   = -243.21945          LR chi2(4)        =          48.79
                                      Prob > chi2         =          0.0000
```

	_t	Haz. Ratio	Std. Err.	z	P> z	[95% Conf. Interval]	
1.sorb		.970657	.1705529	-0.17	0.865	.6878634	1.369712
2.sex		.6384986	.134474	-2.13	0.033	.4225603	.9647865
tgh		1.31143	.0506003	7.03	0.000	1.215913	1.414451
dur		1.096829	.0295709	3.43	0.001	1.040376	1.156346
_cons		.0002114	.0001371	-13.05	0.000	.0000593	.0007535
/ln_p		1.066101	.074717	14.27	0.000	.9196587	1.212544
p		2.904036	.2169808			2.508434	3.362026
1/p		.3443484	.0257287			.2974397	.3986551

```
stcurve, survival at1(sorb=0, sex=1) at2(sor=1, sex=1)
```



```
stcurve, hazard at1(sorb=0, sex=1) at2(sor=1, sex=1)
stcurve, hazard at1(sorb=0, sex=1) at2(sor=1, sex=1) yscale(log)
```



Both exponential and Weibull models are parametric because  $h_0(t)$  has a specified function. Also, you can choose other distributions to build regression models (e.g. gamma distribution).

## Semi-parametric survival analysis: Cox regression model

The **Cox regression model** (aka Cox proportional hazard model) is a semi-parametric regression model for survival data. It is semi-parametric in that it makes parametric assumptions about a linear combination of covariates, but the baseline hazard is not required to be a known distribution:

$$h(t|X_1, \dots, X_p) = h_0(t) \exp(\beta_1 X_1 + \dots \beta_p X_p)$$

To avoid estimating  $h_0(t)$ , the  $\beta$ s are computed by maximizing the partial likelihood. When there are tied events (more than 1 events happen at an observed failure time  $t_k$ ), the computation of partial likelihood becomes tricky. There are a few different ways to handle ties, such as exact partial likelihood, Efron method and Breslow method. Among these three, exact partial likelihood is the most accurate but computationally expensive, while Breslow's method tend to be the least accurate but computationally efficient.

Let's fit a full Cox regression with our variables of interest in Stata.

```

*Full Cox Model
stcox i.sorb i.sex tgh dur
estimates store fullmodel

failure _d: status
analysis time _t: fup
id: id

Iteration 0:  log likelihood = -710.06399
Iteration 1:  log likelihood = -687.1304
Iteration 2:  log likelihood = -685.80968
Iteration 3:  log likelihood = -685.803
Iteration 4:  log likelihood = -685.803
Refining estimates:
Iteration 0:  log likelihood = -685.803

Cox regression -- Breslow method for ties

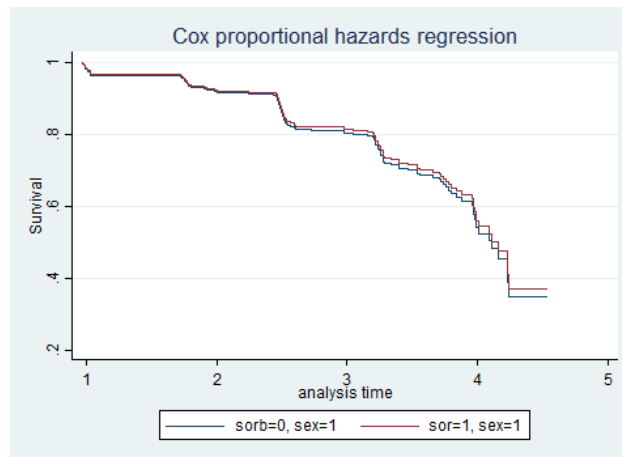
No. of subjects =          478                Number of obs   =          478
No. of failures =          130
Time at risk   =  1411.750697
Log likelihood  =    -685.803                LR chi2(4)        =          48.52
                                                Prob > chi2        =          0.0000

-----+-----
      _t | Haz. Ratio   Std. Err.      z    P>|z|    [95% Conf. Interval]
-----+-----
      1.sorb |   .9436001   .1662714    -0.33   0.742    .6680332    1.33284
      2.sex  |   .6489158   .1361722    -2.06   0.039    .4300979    .97906
      tgh    |    1.3102    .0504697     7.01   0.000    1.214923    1.412949
      dur    |    1.097906   .0300962     3.41   0.001    1.040476    1.158507
-----+-----

```

**Question:** Interpret the coefficient for sex.

```
stcurve, survival at1(sorb=0, sex=1) at2(sor=1, sex=1)
```



**Question:** Can you graph the hazard functions for males in each of two treatment groups solely based on the Cox regression outputs shown above? Why? Can you do that for exponential or Weibull survival regression?

```
stcurve, hazard at1(sorb=0, sex=1) at2(sor=1, sex=1)
stcurve, hazard at1(sorb=0, sex=1) at2(sor=1, sex=1) yscale(log)
```

