## Part III: Survival Models

## **Load Packages**

Again, we must load the packages that will be used in the first part of this workshop.

```
library(pastecs, quietly = TRUE)
library(lm.beta, quietly = TRUE)
library(lmtest, quietly = TRUE)
library(foreign, quietly = TRUE)
library(lattice, quietly = TRUE)
library(lme4, quietly = TRUE)
library(nlme, quietly = TRUE)
library(survival, quietly = TRUE)
library(dplyr, quietly = TRUE)
library(ggfortify, quietly = TRUE)
library(surviminer, quietly = TRUE)
library(rms, quietly = TRUE)
library(rms, quietly = TRUE)
```

## Introduction

Survival models concerns the analysis of the time for an event to occur. The response variable is the time for the event to occur. The event is generally called "death."

#### **Definitions**

Survival models involve two functions:

- S(t): the survival function. S(t) is probability that death has not occurred until after time, t.
- $\lambda(t)$ : the hazard function.

$$\lambda(t) = \frac{\text{probability of dying in at time, } t}{\text{probability of survival until time, } t} \approx \frac{\text{number of people who died at time, } t}{\text{number of people who lived until time, } t}$$

 $\lambda(t)$  measures the likelihood of death in a very small time interval, t and t + dt. It is a measure of risk.

•  $\Lambda(t)$ : the cumulative hazard. It is total hazard from 0 to time, t.

#### Note that these two functions are related:

• 
$$\lambda(t) \leftrightarrow S(t)$$
 
$$\lambda(t) = -\frac{d}{dt} \log S(t)$$

• 
$$\lambda(t) \leftrightarrow \Lambda(t)$$
 
$$\Lambda(t) = \int_0^t \lambda(t) dt$$

•  $S(t) \leftrightarrow \Lambda(t)$  and  $S(t) \leftrightarrow \lambda(t)$ 

$$S(t) = \exp\left(-\Lambda(t)\right) = \exp\left(-\int_0^t \lambda(t) \, dt\right).$$

#### Censoring

Like most models, survival models succeptible to imperfect data. Let's say a subject is recorded for a study up until a time,  $t^*$ . After time  $t^*$ , the subject may decide not to continue with study or it is not possible to locate the subject. Many things could have caused a lack of follow up. This subject is called *censored*. While it maybe reasonable to discard this data point, the censored data actually contains information that we know the event has not occurred prior to  $t^*$ . This gives more information to our model about time prior to  $t^*$  than if we were to discard the censored data.

## Data

#### Description

We will be working the **colon** data set. This data comes from one of the first successful trials of a drug for colon cancer. The recurrence and death times are recorded for all patients in the study.

The colon dataset has the following columns:

- id: id
- study: 1 for all patients
- rx: Treatment Obs(ervation), Lev(amisole), Lev(amisole)+5-FU. Levamisole is a low-toxicity compound previously used to treat worm infestations in animals; 5-FU is a moderately toxic (as these things go) chemotherapy agent.
- sex: 0 = female, 1 = male
- age: age of the patient
- obstruct: 0 = if tumour did not obstructed colon, 1 = if tumour obstructed colon
- perfor: perforation of colon
- adhere: adherence to nearby organs
- nodes: number of lymph nodes with detectable cancer
- time: days until event or censoring
- status: censoring status
- differ: differentiation of tumour (1=well, 2=moderate, 3=poor)
- extent: Extent of local spread (1=submucosa, 2=muscle, 3=serosa, 4=contiguous structures)
- surg: time from surgery to registration (0=short, 1=long)
- node4: more than 4 positive lymph nodes
- etype: event type: 1=recurrence,2=death

# attach(colon) head(colon)

```
##
                      rx sex age obstruct perfor adhere nodes status differ
## 1
      1
              1 Lev+5FU
                               43
                                           0
                                                   0
                                                           0
                                                                  5
                                                                          1
                                                                                  2
##
      1
              1 Lev+5FU
                               43
                                           0
                                                   0
                                                           0
                                                                  5
                                                                          1
                                                                                  2
                           1
##
   3
      2
                               63
                                           0
                                                   0
                                                           0
                                                                  1
                                                                          0
                                                                                  2
              1 Lev+5FU
                           1
## 4
      2
              1 Lev+5FU
                           1
                               63
                                           0
                                                   0
                                                           0
                                                                  1
                                                                          0
                                                                                  2
## 5
                                           0
                                                                                  2
      3
              1
                    Obs
                               71
                                                   0
                                                           1
                                                                  7
                            0
                                                                          1
   6
              1
                    Obs
                                                                                  2
##
                            0
##
     extent surg node4 time etype
## 1
           3
                 0
                        1 1521
                                     2
                 0
                           968
## 2
           3
                                     1
## 3
           3
                 0
                        0 3087
                                     2
           3
                 0
                        0 3087
                                     1
## 4
                                     2
## 5
           2
                 0
                        1
                           963
           2
                 0
## 6
                        1
                           542
                                     1
```

### Subsetting data and converting data

We will be studying the recurrence event of colon cancer.

```
colon_subset_recurrence = colon[colon$etype==1,]
```

Some survival models can only handle variables encoded in 0 and 1. We need to convert continuous variables, such as age and nodes, to 0 and 1.

If the binary variables are stored as numeric variables, the survival models will treat the explanatory variables as continuous variables rather than as discrete variables.

```
sapply(colon,class)
```

```
##
          id
                                                  age obstruct
                 study
                               rx
                                        sex
## "numeric" "numeric"
                        "factor" "numeric" "numeric" "numeric" "numeric"
##
                 nodes
                           status
                                     differ
                                               extent
                                                                     node4
      adhere
                                                            surg
## "numeric" "numeric" "numeric" "numeric" "numeric" "numeric" "numeric"
##
        time
## "numeric" "numeric"
```

Many discrete variables are stored as numeric variables. We have to convert these columns to factor.

The factor takes as arguments:

- the dicrete data in the first argument
- level is current coding the discrete data. This is an optional argument.
- label is the encoding that you would like to change to discrete data. This is an optional argument. Use this argument if you would to change the labeling of the discrete data.

```
colon_subset_recurrence$age.ds <- factor(colon_subset_recurrence$age.ds,</pre>
                                           levels= c("0","1"),
                                           labels=c("<60",">60"))
colon_subset_recurrence$node4 <- factor(colon_subset_recurrence$node4,</pre>
                                             levels= c("0","1"),
                                              labels=c("<4",">4"))
colon_subset_recurrence$sex <- factor(colon_subset_recurrence$sex,</pre>
                                        levels= c("0","1"), labels=c("F","M"))
colon_subset_recurrence$obstruct <- factor(colon_subset_recurrence$obstruct,</pre>
                                             levels= c("0","1"),
                                             labels=c("no obstruct","obstruct"))
colon_subset_recurrence$adhere <- factor(colon_subset_recurrence$adhere,</pre>
                                           levels= c("0","1"),
                                           labels=c("no adhere","adhere"))
colon_subset_recurrence$perfor <- factor(colon_subset_recurrence$perfor,</pre>
                                           levels= c("0","1"),
                                           labels=c("no perfor", "perfor"))
colon_subset_recurrence$differ <- factor(colon_subset_recurrence$differ,</pre>
                                           levels= c("1","2","3"),
                                           labels=c("well","mod","poor"))
colon_subset_recurrence$extent <- factor(colon_subset_recurrence$extent,</pre>
                                           levels= c("1","2","3","4"),
```

Now, let's take a look at the data.

```
head(colon_subset_recurrence)
```

```
##
      id study
                                  obstruct
                                              perfor
                                                        adhere nodes status
                   rx sex age
## 2
             1 Lev+5FU
                        M 43 no obstruct no perfor no adhere
                                                                          1
## 4
      2
                        M 63 no obstruct no perfor no adhere
                                                                          0
             1 Lev+5FU
## 6
      3
            1
                   0bs
                        F
                           71 no obstruct no perfor
                                                        adhere
                                                                   7
                                                                          1
## 8
      4
             1 Lev+5FU
                        F 66
                                  obstruct no perfor no adhere
                                                                   6
                                                                          1
## 10 5
             1
                   Obs
                        M 69 no obstruct no perfor no adhere
                                                                  22
                                                                          1
## 12 6
             1 Lev+5FU
                        F 57 no obstruct no perfor no adhere
                                                                   9
                                                                          1
##
      differ extent surg node4 time etype age.ds
                                              <60
## 2
        mod serosa short
                             >4
                                968
        mod serosa short
                             <4 3087
                                         1
                                              >60
                                              >60
## 6
        mod muscle short
                             >4 542
                                         1
## 8
        mod serosa long
                             >4
                                 245
                                         1
                                              >60
## 10
        mod serosa long
                             >4 523
                                              >60
## 12
        mod serosa short
                                 904
                                              <60
                             >4
                                         1
```

#### Surv Object

The Surv function takes as input the time and censoring status (0 or 1) of a data point. It returns a object that packages together time and censoring status.

```
surv <-with(colon_subset_recurrence, Surv(time,status))
head(surv)</pre>
```

```
## [1] 968 3087+ 542 245 523 904
```

The + at the end of the time indicates that the data point was censored.

## Kalpan-Meier Estimator

First, let  $t_i$  be the *i*th recorded time in the data. That is,  $t_1$  is the 1st recorded time,  $t_2$  is the 2nd recorded time, ...,  $t_{20}$  is the 20th recorded, etc.

Kalpan-Meier assumes that the survival function can be estimated as

$$\hat{S}(t) = \prod_{\text{for } i: t_i < t} \left( 1 - \frac{d_i}{n_i} \right)$$

where  $d_i$  is the number of persons that "died" after time  $t_i$  and  $n_i$  is the number of uncensored persons that have lived up to  $t_i$ .

#### Kalpan-Meier Estimator for the entire data

To fit  $\hat{S}(t) = \prod_{\text{for } i: t_i \le t} 1 - \frac{d_i}{n_i}$  to the entire data, we use the command below.

```
km_fit <- survfit(surv~1, data=colon_subset_recurrence)</pre>
```

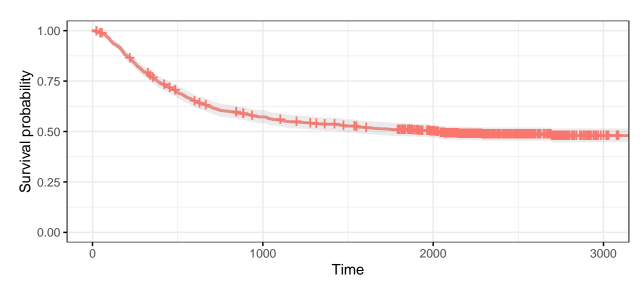
We can return a summary of the  $\hat{S}(t)$  at certain time points. summary(km\_fit) will return a summary km\_fit for all time points in the data. Since the data set is large, I do not run the command, summary(km\_fit). However, you are free to do this on your own.

```
summary(km_fit,times=c(1,10,20,30,40,50))
```

```
## Call: survfit(formula = surv ~ 1, data = colon subset recurrence)
##
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
             929
                       0
                             1.000 0.00000
                                                    1.000
                                                                  1.000
       1
             927
                       2
                             0.998 0.00152
                                                    0.995
##
      10
                                                                  1.000
##
      20
             926
                             0.996 0.00215
                                                    0.991
                                                                  1.000
##
      30
             922
                       1
                             0.995 0.00240
                                                    0.990
                                                                  0.999
             919
##
      40
                       4
                             0.990 0.00322
                                                    0.984
                                                                  0.997
##
      50
             914
                       3
                             0.987 0.00371
                                                    0.980
                                                                  0.994
```

There is a convience function ggsurvplot that generates a plot for a survfit object. conf.int = TRUE shows the confidence interval around the estimate. risk.table = TRUE shows a tabulation of risk below  $\hat{S}(t)$ .





#### Number at risk



## Kalpan-Meier Estimator for the data divided into obstruct and no obstruct

colon\_subset\_recurrence can be divided two data sets by the obstruct column. Those patients whose colons are obstructed by the tumour and those whose colons aren't. We can fit to each data partition to a Kalpan-Meier Estimator

$$\hat{S}_{\text{obstruct}}(t) = \prod_{\substack{\text{for } i: t_i \leq t \\ \text{obstruct}_i = \text{obstruct}}} \left(1 - \frac{d_i}{n_i}\right)$$

$$\hat{S}_{\text{no obstruct}}(t) = \prod_{\substack{\text{for } i: t_i \leq t \\ \text{obstruct} = \text{no obstruct}}} \left(1 - \frac{d_i}{n_i}\right)$$

to the entire data. To do this, we use the command below.

km\_fit <- survfit(surv~obstruct, data=colon\_subset\_recurrence)</pre>

summary(km\_fit,times=c(1,10,20,30,40,50))

## Call: survfit(formula = surv ~ obstruct, data = colon\_subset\_recurrence) ## ## obstruct=no obstruct time n.risk n.event survival std.err lower 95% CI upper 95% CI ## ## 1 749 0 1.000 0.00000 1.000 1.000 1 ## 10 748 0.999 0.00133 0.996 1.000 0 0.999 0.00133 ## 20 748 0.996 1.000 ## 30 746 1 0.997 0.00189 0.994 1.000 40 ## 745 1 0.996 0.00231 0.991 1.000 ## 50 742 0.992 0.00326 0.986 0.998

```
##
##
                     obstruct=obstruct
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
       1
             180
                        0
                              1.000 0.00000
                                                      1.000
                                                                     1.000
##
      10
             179
                        1
                              0.994 0.00554
                                                      0.984
                                                                     1.000
      20
                        2
                              0.983 0.00954
                                                      0.965
                                                                     1.000
##
             178
      30
                        0
                              0.983 0.00954
                                                      0.965
##
             176
                                                                     1.000
             174
                        3
##
      40
                              0.967 0.01342
                                                      0.941
                                                                    0.993
##
      50
             172
                        0
                              0.967 0.01342
                                                      0.941
                                                                     0.993
ggsurvplot(km_fit, data = colon_subset_recurrence,
            pval = TRUE, conf.int = TRUE,
            risk.table = TRUE, ggtheme = theme_bw(),
            risk.table.col = "strata")
                                               obstruct=no obstruct
                                                                       obstruct=obstruct
                 1.00
Survival probability
                 0.75
                 0.50
                 0.25
                           p = 0.036
                 0.00
                                               1000
                                                                      2000
                                                                                             3000
                         0
                                                          Time
                      Number at risk
                                               428
   obstruct=no obstruct
                                                                      323
                                               88
                                                                       71
                         ò
                                               1000
                                                                      2000
                                                                                             3000
```

The p-value in the plot comes the log-rank hypothesis test which allows us to compare a set of Kaplan-Meier estimators. The null hypothesis is that there is no significant different between the Kaplan-Meier estimators. Since p < 0.05, we reject the null hypothesis.

Time

We also do the log-rank hypothesis test using the survdiff function.

```
p_value <- survdiff(surv~obstruct, data=colon_subset_recurrence)</pre>
print(p_value)
## survdiff(formula = surv ~ obstruct, data = colon_subset_recurrence)
##
                           N Observed Expected (O-E)^2/E (O-E)^2/V
                                          386.2
                                                                  4.4
## obstruct=no obstruct 749
                                   369
                                                     0.768
## obstruct=obstruct
                         180
                                    99
                                           81.8
                                                     3.628
                                                                  4.4
```

```
##
   Chisq= 4.4 on 1 degrees of freedom, p= 0.04
survdiff(surv~adhere + obstruct,data=colon_subset_recurrence)
## survdiff(formula = surv ~ adhere + obstruct, data = colon_subset_recurrence)
##
##
                                             N Observed Expected (0-E)^2/E
## adhere=no adhere, obstruct=no obstruct 642
                                                    306
                                                           335.6
                                                                       2.61
## adhere=no adhere, obstruct=obstruct
                                                     80
                                                            69.5
                                                                       1.59
                                           152
## adhere=adhere, obstruct=no obstruct
                                           107
                                                     63
                                                            50.6
                                                                       3.02
## adhere=adhere, obstruct=obstruct
                                            28
                                                     19
                                                            12.3
                                                                       3.68
##
                                           (0-E)^2/V
## adhere=no adhere, obstruct=no obstruct
                                                9.24
## adhere=no adhere, obstruct=obstruct
                                                1.86
## adhere=adhere, obstruct=no obstruct
                                                3.39
  adhere=adhere, obstruct=obstruct
                                                3.79
##
   Chisq= 10.9 on 3 degrees of freedom, p= 0.01
```

#### Kalpan-Meier Estimator for the data divided into adhere and no adhere

colon\_subset\_recurrence can be divided two data sets by the adhere column. Those patients whose colons are obstructed by the tumour and those whose colons aren't. We can fit to each data partition to a Kalpan-Meier Estimator

$$\hat{S}_{\text{adhere}}(t) = \prod_{\substack{\text{for } i: t_i \leq t\\ \text{adher}_i = \text{adhere}}} \left(1 - \frac{d_i}{n_i}\right)$$

$$\hat{S}_{\text{no adhere}}(t) = \prod_{\substack{\text{for } i: t_i \leq t \\ \text{adhered adhere}}} \left(1 - \frac{d_i}{n_i}\right)$$

to the entire data. To do this, we use the command below.

3

##

##

50

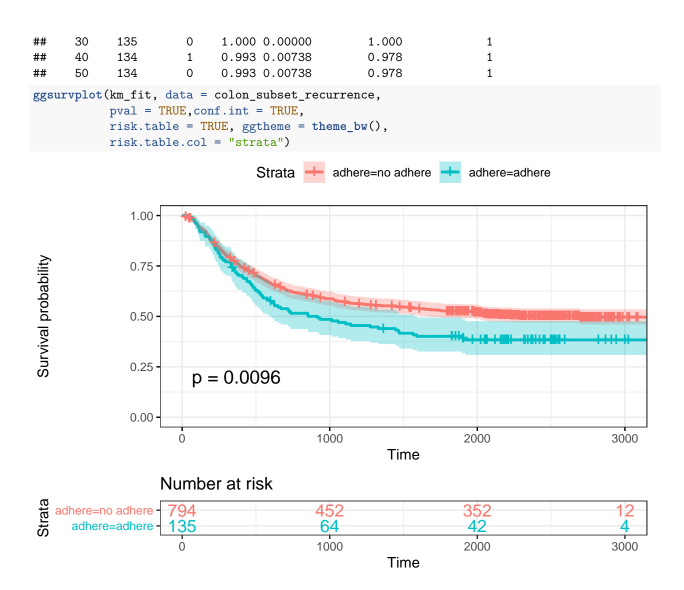
780

```
km_fit <- survfit(surv~adhere, data=colon_subset_recurrence)</pre>
summary(km_fit,times=c(1,10,20,30,40,50))
## Call: survfit(formula = surv ~ adhere, data = colon_subset_recurrence)
##
                    adhere=no adhere
##
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
       1
            794
                       0
                            1.000 0.00000
                                                   1.000
                                                                 1.000
            792
                       2
                                                   0.994
##
      10
                            0.997 0.00178
                                                                 1.000
##
      20
            791
                       2
                            0.995 0.00251
                                                   0.990
                                                                 1.000
##
      30
            787
                       1
                            0.994 0.00281
                                                   0.988
                                                                 0.999
##
      40
            785
                       3
                            0.990 0.00355
                                                   0.983
                                                                 0.997
```

0.978

0.994

0.986 0.00416



# Kalpan-Meier Estimator for the data divided into (adhere, obstruct), (adhere, no obstruct), (no adhere, obstruct) and (no adhere, no obstruct)

colon\_subset\_recurrence can be divided in any amount by the explanatory variables Let's consider breaking up the data based on a patient's obstruction and adherence status. We can fit to each data partition to a Kalpan-Meier Estimator

$$\hat{S}_{\text{adhere,obstruct}}(t) = \prod_{\substack{\text{for } i: t_i \leq t \\ \text{adher}_i = \text{adhere} \\ \text{obstruct}}} \left(1 - \frac{d_i}{n_i}\right)$$

$$\hat{S}_{\text{no adhere,obstruct}}(t) = \prod_{\substack{\text{for } i: t_i \leq t \\ \text{adher}_i = \text{no adhere} \\ \text{obstruct}}} \left(1 - \frac{d_i}{n_i}\right)$$

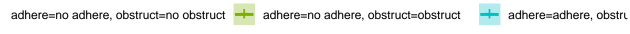
$$\hat{S}_{\text{adhere,no obstruct}}(t) = \prod_{\substack{\text{for } i: t_i \leq t \\ \text{adher}_i = \text{adhere}} \\ \text{obstruct}}} \left(1 - \frac{d_i}{n_i}\right)$$

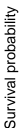
obstruct<sub>i</sub>=no obstruct

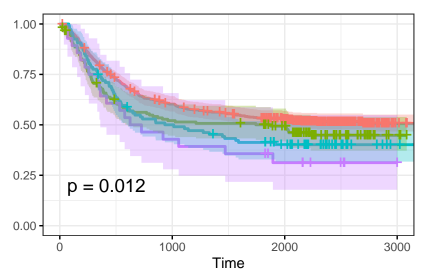
$$\hat{S}_{\text{no adhere,no obstruct}}(t) = \prod_{\substack{\text{for } i: t_i \leq t\\ \text{adher}_i = \text{no adhere}\\ \text{obstruct}_i = \text{no obstruct}}} \left(1 - \frac{d_i}{n_i}\right)$$

to the entire data. To do this, we use the command below.

```
km_fit <- survfit(surv~adhere + obstruct, data=colon_subset_recurrence)</pre>
summary(km_fit,times=c(1,10,20,30,40,50))
## Call: survfit(formula = surv ~ adhere + obstruct, data = colon_subset_recurrence)
##
##
                    adhere=no adhere, obstruct=no obstruct
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
                             1.000 0.00000
                                                    1.000
##
       1
             642
                        0
                                                                  1.000
##
      10
             641
                             0.998 0.00156
                                                    0.995
                                                                  1.000
                        1
##
      20
             641
                        0
                             0.998 0.00156
                                                    0.995
                                                                  1.000
##
      30
             639
                        1
                             0.997 0.00220
                                                    0.993
                                                                  1.000
##
      40
             638
                             0.995 0.00269
                                                    0.990
                                                                  1.000
                        1
##
      50
             635
                        3
                             0.991 0.00380
                                                    0.983
                                                                  0.998
##
                    adhere=no adhere, obstruct=obstruct
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
##
       1
             152
                       0
                             1.000 0.00000
                                                    1.000
                                                                  1.000
             151
                             0.993 0.00656
                                                    0.981
                                                                  1.000
##
      10
                        1
##
      20
             150
                       2
                             0.980 0.01128
                                                    0.958
                                                                  1.000
      30
                             0.980 0.01128
##
             148
                        0
                                                    0.958
                                                                  1.000
##
      40
             147
                        2
                             0.967 0.01451
                                                    0.939
                                                                  0.996
      50
                        0
                             0.967 0.01451
##
             145
                                                    0.939
                                                                  0.996
##
##
                    adhere=adhere, obstruct=no obstruct
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
       1
             107
                        0
                                 1
                                          0
                                                        1
                                                                       1
                        0
##
      10
             107
                                 1
                                          0
                                                        1
                                                                       1
##
      20
             107
                        0
                                          0
                                                                       1
                                                        1
##
      30
             107
                        0
                                          0
                                 1
                                                        1
                                                                       1
##
      40
             107
                        0
                                 1
                                          0
                                                        1
                                                                       1
      50
             107
                        0
                                 1
                                          0
##
                                                        1
                                                                       1
##
##
                    adhere=adhere, obstruct=obstruct
##
    time n.risk n.event survival std.err lower 95% CI upper 95% CI
##
       1
              28
                       0
                             1.000 0.0000
                                                    1.000
                                                                       1
##
      10
              28
                        0
                             1.000 0.0000
                                                    1.000
                                                                       1
      20
              28
                        0
                             1.000
                                    0.0000
##
                                                    1.000
                                                                       1
##
      30
              28
                        0
                             1.000
                                    0.0000
                                                    1.000
                                                                       1
##
      40
              27
                        1
                             0.964
                                    0.0351
                                                    0.898
                                                                       1
##
              27
                       0
                             0.964
                                    0.0351
                                                    0.898
      50
                                                                       1
ggsurvplot(km_fit, data = colon_subset_recurrence,
           pval = TRUE, conf.int = TRUE,
           risk.table = TRUE, ggtheme = theme_bw(),
            risk.table.col = "strata")
```







## Number at risk



survdiff(surv~adhere + obstruct,data=colon\_subset\_recurrence)

```
## Call:
## survdiff(formula = surv ~ adhere + obstruct, data = colon_subset_recurrence)
##
##
                                             N Observed Expected (0-E)^2/E
## adhere=no adhere, obstruct=no obstruct 642
                                                    306
                                                            335.6
                                                                       2.61
## adhere=no adhere, obstruct=obstruct
                                                     80
                                                            69.5
                                                                       1.59
                                           152
                                                                       3.02
## adhere=adhere, obstruct=no obstruct
                                           107
                                                     63
                                                            50.6
## adhere=adhere, obstruct=obstruct
                                            28
                                                     19
                                                            12.3
                                                                       3.68
##
                                           (0-E)^2/V
## adhere=no adhere, obstruct=no obstruct
                                                9.24
## adhere=no adhere, obstruct=obstruct
                                                1.86
## adhere=adhere, obstruct=no obstruct
                                                3.39
## adhere=adhere, obstruct=obstruct
                                                3.79
##
   Chisq= 10.9 on 3 degrees of freedom, p= 0.01
```

## Cox Proportional Hazard

In the limit of large data, the Kaplan-Meier estimator converges to true survival function. However, the Kaplan-Meier has two disadvantages:

- it cannot effectively accommodate continuous data
- it is non-parameteric this means that given a data point, we cannot predict their life trajectory from data. This will be seen more clearly later in this section.

Rather than estimating survival function at each time interval, the Cox Proportional Hazard assumes that hazard function is an exponentiated linear function of explanatory variables. That is,

$$\lambda_i(t) = \lambda_0(t) \exp \left(\beta_1 X_{1i} + \dots + \beta_n X_{ni}\right).$$

 $\lambda_0(t)$  is called the baseline function.  $\lambda(t) = \lambda_0(t)$  when  $X_{1i} = X_{2i} = \cdots = X_{ni} = 0$ .

The Cox Proportional Hazard models the effects of the covariates on the baseline function. It assumes that the ratio of hazards are independent of time. The baseline function is generally unknown. However, the effects of the covariates can still be determined regardless of the baseline function. The  $\beta_i$ 's is calculated using partial maximum likelihood. Avoiding the estimation of  $\lambda_0(t)$  prevents accumulation of errors in a unknown function.

Note that the Cox Proportional Hazard does not solve all the problems of the Kaplan-Meier estimator. Cox Proportional Hazard has one (or 1/2) disadvantage:

• it is semi-parametric. Given a data point, we can estimate the effect of a covariate on the baseline function. However, we cannot predict the life trajectory of data point unless we know  $\lambda_0(t)$ .

## Cox Proportional Hazard for $X_1 = surg$

Given only one covariate, our Cox Proportional Hazard function takes the form

$$\lambda_i(t) = \lambda_0(t) \exp(\beta_1 X_{1i}).$$

where

$$X_{1i} = \begin{cases} 1 & \text{if surgery time of ith data point is long} \\ 0 & \text{otherwise} \end{cases}.$$

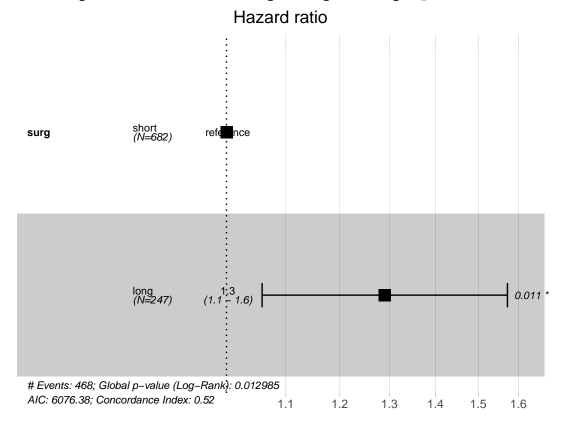
#### Learning Cox Proportional Hazard model

We fit the Cox Proportional Hazard model accordingly.

```
## Call:
## coxph(formula = surv ~ surg, data = colon_subset_recurrence)
##
## n= 929, number of events= 468
##
## coef exp(coef) se(coef) z Pr(>|z|)
## surglong 0.2549 1.2903 0.1008 2.529 0.0114 *
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
            exp(coef) exp(-coef) lower .95 upper .95
                                     1.059
## surglong
                 1.29
                           0.775
                                                1.572
##
## Concordance= 0.523 (se = 0.01)
## Likelihood ratio test= 6.17 on 1 df,
                                           p=0.01
## Wald test
                        = 6.39
                                on 1 df,
                                           p=0.01
## Score (logrank) test = 6.43
                                on 1 df,
                                           p=0.01
coef(cox)
##
   surglong
## 0.2548703
ggforest(cox, data = colon_subset_recurrence)
```

## Warning: Removed 1 rows containing missing values (geom\_errorbar).



#### **Testing Proportionality Assumption**

The Cox proportionality hazard model assumes that ratio of the hazards are constant over time. If ratio of the hazards are constant over time, then covariates and their effects must also be constant over time. If this assumption is violated, then one might get strange results (such as the crossing of Kaplan-Meier curves).

To test for proportionality hazard assumption, we use the cox.zph function. cox.zph takes a coxph model as input and returns a p-value to determine whether the proportionality hazard assumption was voilated for each covariate. cox.zph tests the null hypothesis that there are no time dependent relationships in the covariates and their effects.

Since the p value is greater than 0.05, we fail to reject the null hypothesis

#### **Model Selection**

where

```
anova(cox)

## Analysis of Deviance Table

## Cox model: response is surv

## Terms added sequentially (first to last)

##

## loglik Chisq Df Pr(>|Chi|)

## NULL -3040.3

## surg -3037.2 6.1712 1 0.01299 *

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

## Cox Proportional Hazard for $X_1 = \text{surg}$ , $X_2 = \text{adher}$

Given only two covariate, our Cox Proportional Hazard function takes the form

$$\lambda_i(t) = \lambda_0(t) \exp(\beta_1 X_{1i} + \beta_2 X_{2i}).$$
1 if surgery time of i th data point is less than 1 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 2 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point is less than 3 in the surgery time of its data point i

 $X_{1i} = \begin{cases} 1 & \text{if surgery time of i th data point is long} \\ 0 & \text{otherwise} \end{cases}$ 

 $X_{2i} = \begin{cases} 1 & \text{if the i th data point has adherence to other organs} \\ 0 & \text{otherwise} \end{cases}$ 

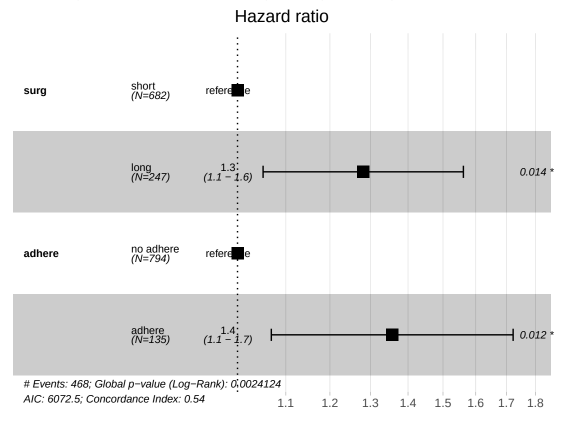
#### Learning Cox Proportional Hazard model

We fit the Cox Proportional Hazard model accordingly.

```
## Call:
## coxph(formula = surv ~ surg + adhere, data = colon_subset_recurrence)
##
##
     n= 929, number of events= 468
##
##
                  coef exp(coef) se(coef)
                                               z Pr(>|z|)
## surglong
                0.2481
                          1.2816
                                   0.1008 2.460
                                                   0.0139 *
## adhereadhere 0.3053
                          1.3570
                                   0.1217 2.508
                                                   0.0121 *
## ---
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
                exp(coef) exp(-coef) lower .95 upper .95
                    1.282
                              0.7803
                                         1.052
                                                   1.562
## surglong
                    1.357
                              0.7369
## adhereadhere
                                         1.069
                                                   1.723
##
## Concordance= 0.538 (se = 0.012)
## Likelihood ratio test= 12.05 on 2 df,
                                            p=0.002
## Wald test
                        = 12.72 on 2 df,
                                            p=0.002
## Score (logrank) test = 12.81 on 2 df,
                                            p=0.002
coef(cox)
##
       surglong adhereadhere
##
      0.2480832
                   0.3052677
ggforest(cox, data = colon_subset_recurrence)
```

## Warning: Removed 2 rows containing missing values (geom\_errorbar).



#### **Testing Proportionality Assumption**

```
test.ph <- cox.zph(cox)
test.ph

## rho chisq p
## surglong 0.0486 1.110 0.292
## adhereadhere 0.0403 0.764 0.382
## GLOBAL NA 1.931 0.381
```

Since the p value is greater than 0.05, we fail to reject the null hypothesis

#### **Model Selection**

where

## ---

```
anova(cox)

## Analysis of Deviance Table

## Cox model: response is surv

## Terms added sequentially (first to last)

##

## loglik Chisq Df Pr(>|Chi|)

## NULL -3040.3

## surg -3037.2 6.1712 1 0.01299 *

## adhere -3034.2 5.8831 1 0.01529 *

## ---

## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

## Cox Proportional Hazard for $X_1 = \text{surg}$ , $X_2 = \text{adher}$ , $X_3 = \text{nodes}$

Given only three covariate, our Cox Proportional Hazard function takes the form

$$\lambda_i(t) = \lambda_0(t) \exp\left(\beta_1 X_{1i} + \beta_2 X_{2i} + \beta_3 X_{3i}\right).$$
 
$$X_{1i} = \begin{cases} 1 & \text{if surgery time of i th data point is long} \\ 0 & \text{otherwise} \end{cases},$$
 
$$X_{2i} = \begin{cases} 1 & \text{if the i th data point has adherence to other organs} \\ 0 & \text{otherwise} \end{cases}$$

and  $X_{3i}$  is number of nodes of the i th data point.

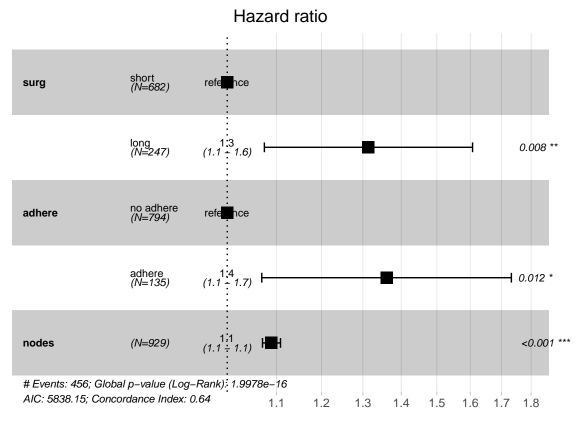
#### Learning Cox Proportional Hazard model

We fit the Cox Proportional Hazard model accordingly.

```
cox <- coxph(surv ~ surg + adhere + nodes,</pre>
             data=colon subset recurrence)
summary(cox)
## Call:
## coxph(formula = surv ~ surg + adhere + nodes, data = colon_subset_recurrence)
##
##
     n= 911, number of events= 456
      (18 observations deleted due to missingness)
##
##
                   coef exp(coef) se(coef)
##
                                                z Pr(>|z|)
                0.27318
                          1.31414 0.10273 2.659 0.00783 **
## surglong
## adhereadhere 0.30821
                          1.36099
                                   0.12317 2.502 0.01234 *
## nodes
                0.08562
                         1.08939 0.00888 9.642 < 2e-16 ***
```

```
## Signif. codes: 0 '***' 0.001 '**' 0.01 '*' 0.05 '.' 0.1 ' ' 1
##
##
               exp(coef) exp(-coef) lower .95 upper .95
                             0.7610
                                        1.074
## surglong
                   1.314
                                                  1.607
                             0.7348
## adhereadhere
                   1.361
                                        1.069
                                                  1.733
## nodes
                   1.089
                             0.9179
                                        1.071
                                                  1.109
## Concordance= 0.636 (se = 0.013)
                                          p=<2e-16
## Likelihood ratio test= 76.21 on 3 df,
## Wald test
                       = 102.7 on 3 df, p=<2e-16
## Score (logrank) test = 105.5 on 3 df, p=<2e-16
coef(cox)
##
      surglong adhereadhere
                                   nodes
##
    0.27318255
                 0.30821401
                              0.08561674
ggforest(cox, data = colon_subset_recurrence)
```

## Warning: Removed 2 rows containing missing values (geom\_errorbar).



#### Testing Proportionality Assumption

```
test.ph <- cox.zph(cox)
test.ph

## rho chisq p
## surglong 0.0590 1.593 0.207
## adhereadhere 0.0307 0.430 0.512
```

```
## nodes -0.0460 0.601 0.438
## GLOBAL NA 2.815 0.421
```

Since the p value is greater than 0.05, we fail to reject the null hypothesis

#### **Model Selection**

```
anova(cox)
## Analysis of Deviance Table
## Cox model: response is surv
## Terms added sequentially (first to last)
##
##
                   Chisq Df Pr(>|Chi|)
          loglik
## NULL
         -2954.2
         -2951.6 5.1725 1
                               0.02295 *
## surg
## adhere -2948.6 6.0414 1
                               0.01397 *
## nodes -2916.1 64.9922 1 7.519e-16 ***
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
```

#### **Estimating Survival Curve**

It is possible to estimate the survival curve for the Cox Proportional Model as long as we have some estimate for  $\lambda_0(t)$ . One way to estimate  $\lambda_0(t)$  from data is to use formula:

$$\lambda_0(t_i) \approx \frac{d_i}{\sum_{s \in R_i} \exp(\beta_1 X_{1s} + \dots + \beta_n X_{ns})}$$

where  $d_i$  is the number of deaths in at time  $t_i$ ,  $R_i$  is set of persons alive after  $t_i$  and  $X_{ij}$  is the *i*th explanatory variable of the *j*th person.

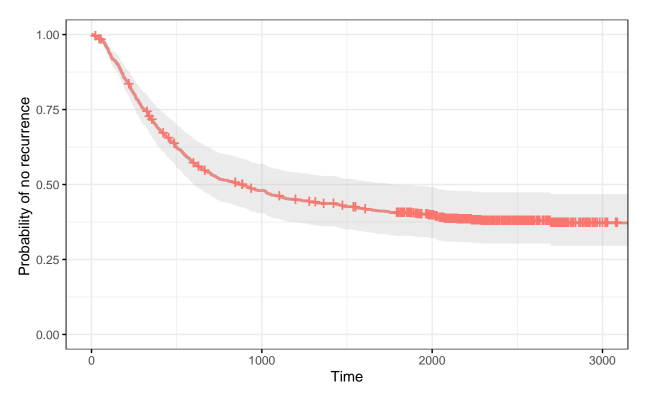
Now let's create some data point. This data point will have the surg set to short, adhere set to no adhere, nodes set to 5 and extent set to serosa.

Using the survfit function, we can generate an object which will be used for plotting. survfit takes as argument:

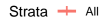
- first argment: cox proportional hazard model fit with coxph
- second argment: the data point in question. It must have the same explanatory variables as the model in the first argument
- data: the data set used to fit the coxph object.

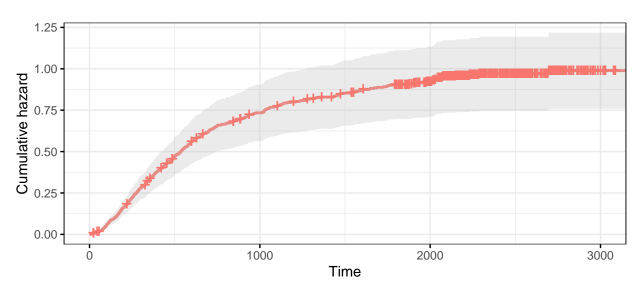
We then use the ggsurvplot function to plot the estimate of the survival curve from survfit fit object.

## Strata + All



We can also use the <code>ggsurvplot</code> function to plot the estimate of the cumulative hazard curve from <code>survfit</code> fit object







#### Accelerated failure time models

Accelerated failure time model assume that the log time for an event to occur is a function of the covariates of the data. That is,

$$\log T_i = \beta_1 X_{1i} + \dots + \beta_n X_{ni} + \varepsilon_i$$

where  $\varepsilon$  is a random error term that follows a distribution.

This is called an accelerated failure model since covariates can scale the base time distribution,  $T_0$ , by their effects.

$$T_i = T_0 \exp(\beta_1 X_{1i} + \dots + \beta_n X_{ni})$$

where  $T_0 = \exp(\varepsilon_i)$ .

There is difference between proportional hazard models (PH) and accelerated failure time models (AFT). The effect of the covariates in PH models act multiplicately on the base hazard. However, in AFT models, these effects act multiplicately on the base time. Despite this difference, it is possible that AFT models are also PH models.

We use the function, survreg, to fit accelerated failure time models. The argument, dist, specifies the distribution which implies the form of  $\lambda_0(t)$ . We will be considering:

- exponential models, dist="exponential"
- weibull models, dist="weibull"
- lognormal models, dist="lognormal"

These are fully parameteric model and are thus a suitable alterative Kaplan-Meier estimators and Cox Proportional Hazard models. However, AFT assume the function form  $T_0$  and, thus, the baseline hazard and the baseline survival functions. Incorrect assumptions introduce errors in our modeling.

#### Exponential models

Exponential accelerated failure time models are also proportional hazard models. Exponential accelerated failure time models assume that  $T_0$  follows a exponential distribution with parameter  $\lambda$ .

From our definitions of terms and with some probability theory (not covered), the hazard and survival function of an exponential AFT models are

$$\lambda_i(t) = \lambda \exp\left(\beta_1 X_{1i} + \dots + \beta_n X_{ni}\right)$$

and 
$$S_i(t) = \exp(\beta_1 X_{1i} + \dots + \beta_n X_{ni}) S_0(t), S_0(t) = \exp(-\lambda t).$$

As proportional hazard model, exponential accelerated failure time models assumes that the baseline hazard is constant,  $\lambda_0(t) = \lambda$ .

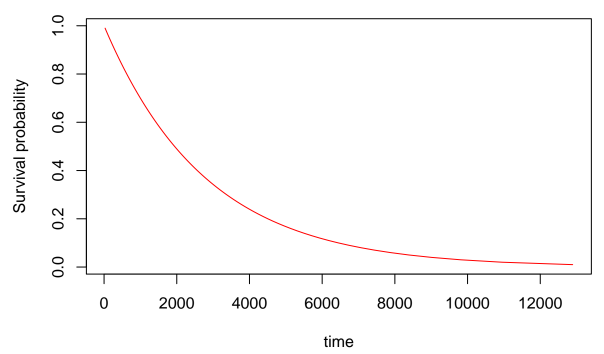
#### Learning Exponential models

survreg learns the parameter value,  $\lambda$ , and the regression coefficients. As an example, we will be consider the model: surv ~ 1 + surg + adhere + nodes.

```
survregExp <- survreg(surv ~ 1 + surg + adhere + nodes,</pre>
                          dist="exponential",data=colon_subset_recurrence)
summary(survregExp)
##
## Call:
## survreg(formula = surv ~ 1 + surg + adhere + nodes, data = colon_subset_recurrence,
      dist = "exponential")
##
                 Value Std. Error
## (Intercept) 8.45944
                         0.07338 115.29 <2e-16
## surglong
             -0.32521 0.10289 -3.16 0.0016
## nodes
              -0.10446 0.00878 -11.90 <2e-16
##
## Scale fixed at 1
##
## Exponential distribution
## Loglik(model) = -4024.3
                         Loglik(intercept only) = -4078.5
## Chisq= 108.33 on 3 degrees of freedom, p= 2.5e-23
## Number of Newton-Raphson Iterations: 5
## n=911 (18 observations deleted due to missingness)
Therefore, \lambda = \exp(8.45944).
```

#### **Estimating Survival Curve**

## **Exponential AFT Model**



Weibull models

Weibull accelerated failure time models are also proportional hazard models. Weibull accelerated failure time models assume that  $T_0$  follows a Weibull distribution with parameters,  $\lambda$  and  $\gamma$ .

##

From our definitions of terms and with some probability theory (not covered), the hazard and survival function of a Weibull AFT models are

$$\lambda_i(t) = \lambda \gamma t^{\gamma - 1} \exp(\beta_1 X_{1i} + \dots + \beta_n X_{ni}).$$

and 
$$S_i(t) = \exp(\beta_1 X_{1i} + \dots + \beta_n X_{ni}) S_0(t), S_0(t) = \exp(-(\lambda t)^{\gamma}).$$

As proportional hazard model, Weibull accelerated failure time models assumes that the baseline hazard is constant,  $\lambda \gamma t^{\gamma-1}$ . One can see that exponential accelerated failure time models are a special case of Weibull accelerated failure time models with  $\gamma = 1$ .

#### Learning Weibull models

survreg learns the parameter value,  $\lambda$  and  $\gamma$ , and the regression coefficients.

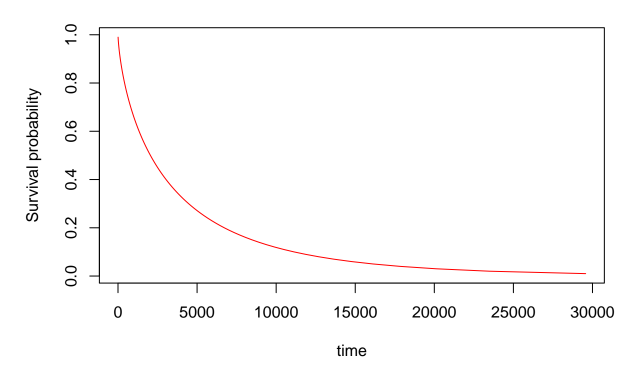
As an example, we will be consider the model: surv ~ 1 + surg + adhere + nodes for all the accelerated time models.

```
##
## Call:
## survreg(formula = surv ~ 1 + surg + adhere + nodes, data = colon_subset_recurrence,
```

```
dist = "weibull")
##
##
                  Value Std. Error
                                        z
                 8.7993 0.1155 76.21 <2e-16
## (Intercept)
## surglong
               -0.4156
                            0.1454 -2.86 0.0042
## adhereadhere -0.4629
                            0.1743 -2.66 0.0079
## nodes -0.1315
                            0.0131 -10.04 <2e-16
## Log(scale)
                0.3432
                            0.0414 8.30 <2e-16
##
## Scale= 1.41
##
## Weibull distribution
## Loglik(model) = -3984.6 Loglik(intercept only) = -4028.2
## Chisq= 87.17 on 3 degrees of freedom, p= 8.9e-19
## Number of Newton-Raphson Iterations: 5
## n=911 (18 observations deleted due to missingness)
Therefore,
                                      \gamma = \exp(0.3432)
                                     \lambda = \exp(8.7993 \times \gamma)
```

#### **Estimating Survival Curve**

## Weibull AFT Model



#### Log-normal models

Log-normal accelerated failure time models assume that  $T_0$  follows a log normal distribution with a scale parameter. Log-normal accelerated failure time models are not proportional hazard models.

The hazard and survival function of a log-normal AFT models are a bit complicated so they will not be shown here.

#### Learning Log-normal models

survreg learns the parameter value,  $\lambda$  and  $\gamma$ , and the regression coefficients.

As an example, we will be consider the model: surv ~ 1 + surg + adhere + nodes for all the accelerated time models.

```
##
## Call:
  survreg(formula = surv ~ 1 + surg + adhere + nodes, data = colon_subset_recurrence,
##
       dist = "lognormal")
##
                  Value Std. Error
                                        z
## (Intercept)
                 8.3066
                             0.1223 67.93 <2e-16
## surglong
                -0.3521
                             0.1525 - 2.31
                                           0.021
## adhereadhere -0.4685
                             0.1876 - 2.50
                                          0.013
                -0.1613
## nodes
                             0.0182 -8.86 <2e-16
## Log(scale)
                 0.6127
                             0.0371 16.49 <2e-16
##
```

```
## Scale= 1.85
##
## Log Normal distribution
## Loglik(model)= -3940.5 Loglik(intercept only)= -3983.6
## Chisq= 86.09 on 3 degrees of freedom, p= 1.5e-18
## Number of Newton-Raphson Iterations: 3
## n=911 (18 observations deleted due to missingness)
```

#### **Estimating Survival Curve**

# **Log Normal AFT Model**

