Fundamentals of Statistical Modeling (VT20)

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Lab 4

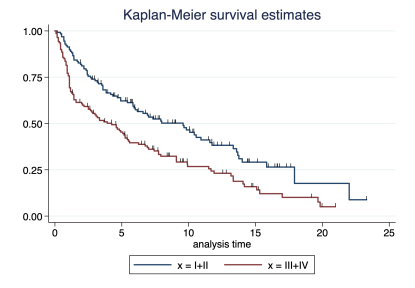
Load the dataset and the mlci command

```
. version 14
. use https://raw.githubusercontent.com/anddis/fsm/master/data/lab4_1.dta, clear
. run https://raw.githubusercontent.com/anddis/fsm/master/do/mlci.do
```

Exercise 1

We use data from 267 patients diagnosed with oral cancer. We measured time to death (y) in subjects with low-grade cancer (x = 0) and high-grade cancer (x = 1). Some survival times are censored (d = 0). First, we plot Kaplan-Meier estimates of the survival functions.

```
. stset y, fail(d)
    failure event: d != 0 & d < .
obs. time interval: (0, y]
exit on or before: failure
       267 total observations
         0 exclusions
       267 observations remaining, representing
       184 failures in single-record/single-failure data
 1,620.864 total analysis time at risk and under observation
                                               at risk from t =
                                                                        0
                                    earliest observed entry t =
                                                                        0
                                                                   23.258
                                         last observed exit t =
. sts graph, by(x) cens(single) name(km, replace)
        failure _d: d
  analysis time _t: y
. graph export km.png, replace
(file km.png written in PNG format)
```



We consider a log-logistic model (AFT) for f(y|x) (see slides 112). Estimate the model's parameters. Constrain the parameter λ to be positive. Take into account right censoring (see slide 76).

The PDF of a (standard) logistic distribution is:

$$f(y) = \frac{\exp(-y)}{(1 + \exp(-y))^2}$$

while the Survival function is:

$$S(y) = 1 - \frac{1}{1 + \exp(-y)}$$

```
. local lambda = "exp({theta})"
. local G = "(log(y)-({beta0}+{beta1}*x))/`lambda`"
. local g = "1/(`lambda'*y)"
. local f = \exp(-(G'))/((1+\exp(-(G')))^2)*g''
. local S = "1-1/(1+exp(-(`G')))"
. mlexp ((d==1)*ln(`f') + (d==0)*ln(`S'))
initial:
               log likelihood = -696.29352
               log likelihood = -641.95479
alternative:
rescale:
               log\ likelihood = -641.95479
rescale eq:
               log likelihood = -573.91032
               log\ likelihood = -573.91032
Iteration 0:
Iteration 1:
               log likelihood =
               log likelihood = -572.24452
Iteration 2:
Iteration 3:
               log\ likelihood = -572.24452
Maximum likelihood estimation
Log likelihood = -572.24452
                                                 Number of obs
```

	Coef.	Std. Err.	z	P> z	[95% Conf.	Interval]
/beta0	2.051811	.1455026	14.10	0.000	1.766631	2.336991
/beta1	8090195	.1993996	-4.06	0.000	-1.199836	4182034
/theta	1109087	.060722	-1.83	0.068	2299216	.0081042

```
. mlci exp /beta1
```

.4452944 95% CI: .3012437, .6582283

. mlci exp /theta

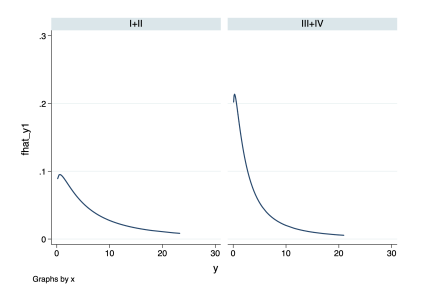
.8950204 95% CI: .7945959, 1.008137

Plot the estimated densities $\hat{f}(y|x)$.

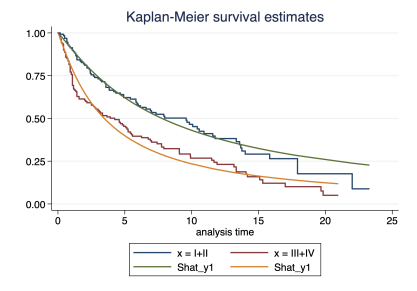
. gen fhat_y1 = $\exp(-((\ln(y)-(_b[/beta0]+_b[/beta1]*x))/\exp(_b[/theta])))/((1+exp(-(\ln(y)-(_b[/beta0]+_b[/beta1]*x))/exp(_b[/theta])))^2)*1/(exp(_b[/theta])*y)$

267

```
. tw (line fhat_y1 y, sort), name(p1, replace) by(x) ylabel(0(0.1)0.3) . graph export p1.png, replace (file p1.png written in PNG format)
```



Plot the estimated survival functions $\hat{S}(y|x)$ together with the Kaplan-Meier estimates.



Exercise 2

Use a RCS transformation of time to death to make the log-logistic model more flexible (see slide 118).

Generate a RCS transform of y (V2) and its derivative (v2) using rcsgen. Estimate the model's parameters. Constrain the parameter λ to be positive

```
. rcsgen y, gen(V) dgen(v) df(2)
```

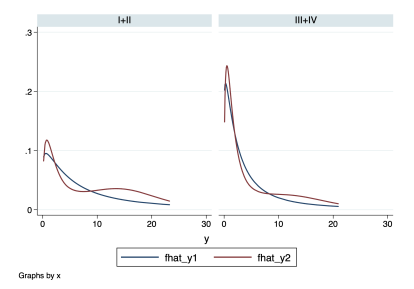
```
. local f = \exp(-(G'))/((1+\exp(-(G')))^2)*g''
     . local S = "1-1/(1+exp(-(`G')))"
     . mlexp ((d==1)*ln(`f') + (d==0)*ln(`S'))
     initial:
                   log likelihood = -696.29352
                   log likelihood = -696.29352
     final:
     rescale:
                   log likelihood = -696.29352
                   log likelihood = -696.29352
     Iteration 0:
                                              (not concave)
                   log likelihood = -596.84712
     Iteration 1:
                                              (not concave)
     Iteration 2:
                   log\ likelihood = -573.66527
                   log likelihood = -566.46684
     Iteration 3:
                   log likelihood = -562.42408
     Iteration 4:
                   \log = -562.22973
     Iteration 5:
                   \log likelihood = -562.22929
     Iteration 6:
     Iteration 7:
                   \log likelihood = -562.22929
     Maximum likelihood estimation
     Log likelihood = -562.22929
                                                  Number of obs
                                                                            267
                        Coef.
                               Std. Err.
                                                  P>|z|
                                                            [95% Conf. Interval]
            /eta1
                    -.1457699
                                .0290741
                                           -5.01
                                                  0.000
                                                           -.2027541
                                                                      -.0887858
                    -.0008936
            /eta2
                               .0001706
                                           -5.24
                                                  0.000
                                                           -.0012279
                                                                      -.0005592
           /beta0
                     1.334615
                                .2061882
                                           6.47
                                                  0.000
                                                            .9304931
                                                                       1.738736
           /beta1
                    -.5995634
                                .1623232
                                           -3.69
                                                  0.000
                                                            -.917711
                                                                      -.2814157
           /theta
                    -.3866109
                               .1145084
                                                  0.001
                                                                      -.1621786
                                           -3.38
                                                           -.6110432
     . mlci exp /beta1
      .5490513
               95% CI: .3994323, .7547145
     . mlci exp /theta
      .6793554
                95% CI: .5427843, .8502893
     . test [eta1]_cons [eta2]_cons
      (1) [eta1]_{cons} = 0
      (2) [eta2]_{cons} = 0
                chi2( 2) =
                            28.76
             Prob > chi2 =
                             0.0000
Plot the estimated densities \hat{f}(y|x)
     > ta1]*y+_b[/eta2]*V2-(_b[/beta0]+_b[/beta1]*x))/exp(_b[/theta])))^2)*(1/y+_b[/eta1]+_b[/eta2]*v2)/(exp(_b[/theta]))
     . tw (line fhat_y1 fhat_y2 y, sort), name(p3, replace) by(x) ylabel(0(0.1)0.3) legend(rows(1))
      . graph export p2.png, replace
     (file p2.png written in PNG format)
```

Variables V1 to V2 and v1 to v2 were created

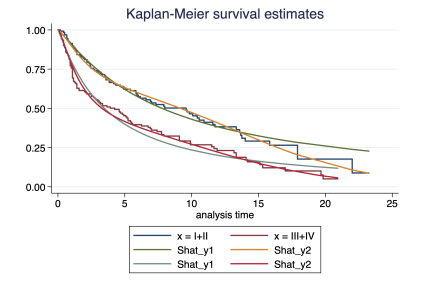
. local g = $(1/y+\{eta1\}+\{eta2\}*v2)/`lambda´"$

. local G = "(log(y)+{eta1}*y+{eta2}*V2-({beta0}+{beta1}*x))/`lambda´"

. local lambda = "exp({theta})"



Plot the estimated survival functions $\hat{S}(y|x)$ over the Kaplan-Meier estimates.



Exercise 3

We consider a Weibull model (PH) (see slide 124). Estimate the model's parameters. Constrain the parameter k to be positive.

But first: we need to extend the likelihood on slide 123 in order to accommodate the censored observations.

The log-likelihood, in the presence of right censoring, is (see slides 76):

$$\log[L(\theta)] = \sum_{i=1}^{n} I(d_i = 1) \log[f(z_i)] + I(d_i = 0) \log[Sz_i].$$

Knowing that f(y) = S(y)h(y) and $S(y) = \exp(-H(y))$ (see slide 121), we can rewrite it as

$$\log[L(\theta)] \doteq \sum_{i=1}^{n} I(d_i = 1)(\log[\exp(-H(z_i))] + \log[h(z_i)]) + I(d_i = 0)\log[\exp(-H(z_i))] = \sum_{i=1}^{n} I(d_i = 1)\log[h(z_i)] - H(z_i).$$

This equation justifies the form of the log-likelihood passed to mlexp in Exercise 4 and 5.

What's the interpretation of β_1 ?

```
. local k = "exp({theta})"
. local G = \text{``k'*log(y)+\{beta0\}+\{beta1\}*x''}
. local g = "(k'/y)"
. local H = "exp(`G')"
. local h = "'H'*g'"
. mlexp ((d==1)*(ln(`h')-`H') + (d==0)*(-`H'))
initial:
               log likelihood = -1620.864
               log likelihood = -659.98633
alternative:
               \log = -659.98633
rescale:
               log \ likelihood = -605.31316
rescale eq:
Iteration 0:
              \log likelihood = -605.31316
               log likelihood = -575.24733
Iteration 1:
Iteration 2:
               log\ likelihood = -573.66495
Iteration 3:
               \log \frac{1}{100} likelihood = -573.66061
Iteration 4:
               log likelihood = -573.66061
Maximum likelihood estimation
Log likelihood = -573.66061
                                                  Number of obs
                                                                               267
                     Coef.
                             Std. Err.
                                             z
                                                  P>|z|
                                                             [95% Conf. Interval]
      /theta
                 -.1586235
                              .0610589
                                          -2.60
                                                  0.009
                                                            -.2782968
                                                                         -.0389502
```

-13.29

3.58

0.000

0.000

.1611149

.1499405

Plot the estimated densities $\hat{f}(y|x)$

-2.141021

.5370332

/beta0

/beta1

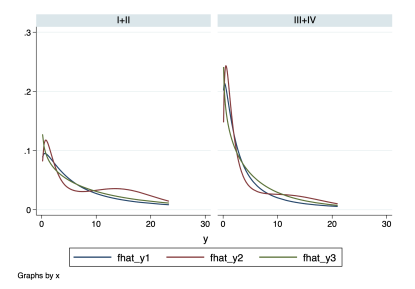
```
. gen fhat_y3 = exp(ln(exp(exp(_b[/theta])*log(y)+_b[/beta0]+_b[/beta1]*x)*(exp(_b[/theta])/y))-exp(exp(_b[/theta])*log(
> y)+_b[/beta0]+_b[/beta1]*x))
. tw (line fhat_y1 fhat_y2 fhat_y3 y, sort), name(p3, replace) by(x) ylabel(0(0.1)0.3) legend(rows(1))
. graph export p3.png, replace
(file p3.png written in PNG format)
```

-2.4568

.2431551

-1.825241

.8309113



Exercise 4

We now use a RCS transformation of time to death to make the Weibull model more flexible (see slide 128).

```
. local k = "exp({theta})"
. local G = \text{``k'*log(y)+\{eta1\}*y+\{eta2\}*V2+\{beta0\}+\{beta1\}*x''}
. local g = "(k'/y+\{eta1\}+\{eta2\}*v2)"
. local H = "exp(`G')"
. local h = "'H'*g'"
. mlexp ((d==1)*(ln(`h')-`H') + (d==0)*(-`H'))
               log likelihood = -1620.864
initial:
               log likelihood =
final:
                                 -1620.864
               log likelihood = -1620.864
rescale:
Iteration 0:
               log likelihood = -1620.864
               log likelihood = -582.7029
Iteration 1:
                                             (not concave)
Iteration 2:
               log likelihood =
                                  -568.651
               log likelihood = -567.19767
Iteration 3:
Iteration 4:
               log likelihood = -565.12405
Iteration 5:
               log\ likelihood = -565.06182
               log likelihood = -565.06137
Iteration 6:
Iteration 7:
               log\ likelihood = -565.06137
Maximum likelihood estimation
Log likelihood = -565.06137
                                                  Number of obs
```

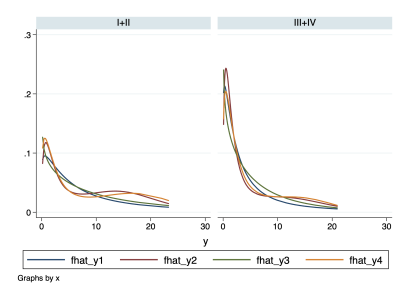
267 Coef. Std. Err. z P>|z| [95% Conf. Interval] .0193952 . 2395006 .1123007 2.13 0.033 .4596059 /theta /eta1 -.1837425 .0466373 -3.940.000 -.2751499 -.092335 -.0008603 .000215 -4.00 0.000 -.0012817 -.000439 /eta2 /beta0 -1.909837 .1633405 -11.69 0.000 -2.229978 -1.589695 .5297763 .1499195 3.53 0.000 .2359395 .823613 /beta1

Plot the estimated densities $\hat{f}(y|x)$

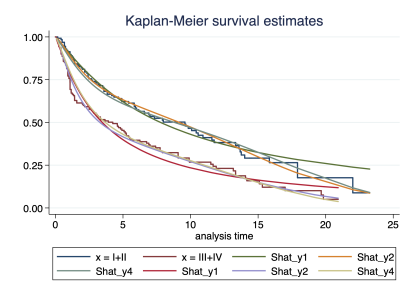
```
. gen fhat_y4 = \exp(\ln(\exp(\exp(_b[/theta])*\log(y)+_b[/eta1]*y+_b[/eta2]*V2+_b[/beta0]+_b[/beta1]*x)*(\exp(_b[/theta])/y+_b[/eta1]+_b[/eta2]*v2))-\exp(\exp(_b[/theta])*\log(y)+_b[/eta1]*y+_b[/eta2]*V2+_b[/beta0]+_b[/beta1]*x))
```

[.] tw (line fhat_y1 fhat_y2 fhat_y3 fhat_y4 y, sort), name(p4, replace) by(x) ylabel(0(0.1)0.3) legend(rows(1))

. graph export p4.png, replace (file p4.png written in PNG format)



Plot the estimated survival functions $\hat{S}(y|x) = \exp(-H(y|x))$ over the Kaplan-Meier estimates.

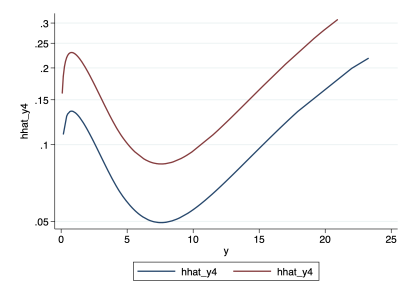


Plot the model-based estimated hazards functions $\hat{h}(y|x)$. Use a log scale for the vertical axis to visually check that the model-based hazard functions are actually proportional. **Important**: the hazard functions are proportional because we forced them to be so (using a Weibull PH model)!

Now interpret them, if you can :-)

```
. gen hhat_y4 = \exp(\exp(_b[/theta])*\log(y)+_b[/eta1]*y+_b[/eta2]*V2+_b[/beta0]+_b[/beta1]*x) * /// > (\exp(_b[/theta])/y+_b[/eta1]+_b[/eta2]*v2)
. tw (line hhat_y4 y if x == 0, sort) (line hhat_y4 y if x == 1, sort), ///
```

```
> yscale(log) name(p5, replace)
. graph export p5.png, replace
(file p5.png written in PNG format)
```



Extra

Can we fit a so-called "flexible parametric survival model" using the tools we've learned so far? Of course. To us, it's just one possible way of modeling y.

Note that here we apply RCS transforms to $z = \log(y)$ instead of y.

(You'll need to install the command stpm2, first)

- . cap net install stpm2, from(http://fmwww.bc.edu/RePEc/bocode/s)
- . stpm2 x, df(3) scale(h) noorthog nolog

Log likelihood = -395.20707Number of obs 267 Coef. Std. Err. z P>|z| [95% Conf. Interval] xb .5302016 .1499414 3.54 0.000 .2363219 .8240813 0.000 2.73159 2.104715 .3198397 6.58 1,477841 _rcs1 _rcs2 .2025602 .0496948 4.08 0.000 .1051602 . 2999602 -.2002837 .0547224 -3.66 0.000 -.3075376 -.0930298 _rcs3 -1.165862 .2565388 -1.668668 -.6630549 _cons -4.540.000

```
di e(ln_bhknots)
```

-2.465104022491821 .223943231484774 1.641711472984396 3.091360584567398

```
. gen double z = log(y)
```

. rcsgen z, gen(z_rcs) dgen(z_d_rcs) knots(-2.465104022491821 .223943231484774 1.641711472984396 3.091360584567398) Variables z_rcs1 to z_rcs3 and z_d_rcs1 to z_d_rcs3 were created

- . local G = " $\{eta0\}*z+\{eta1\}*z_rcs2+\{eta2\}*z_rcs3+\{beta0\}+\{beta1\}*x$ "
- . local g = "($\{eta0\}+\{eta1\}*z_d_rcs2+\{eta2\}*z_d_rcs3$)"
- . local H = "exp(`G')"
- . local h = "'H'*g'"
- . mlexp ((d==1)*ln(`h')-`H'), from(eta0=1 eta1=0 eta2=0 beta0=0 beta1=0)

Iteration 0: log likelihood = -1453.6616 (not concave)

Iteration 1: log likelihood = -513.69571
Iteration 2: log likelihood = -440.91209

Iteration 3: log likelihood = -402.04626

¹Royston, P., & Parmar, M. K. (2002). Flexible parametric proportional-hazards and proportional-odds models for censored survival data, with application to prognostic modelling and estimation of treatment effects. Statistics in medicine, 21(15), 2175-2197.

Iteration 4: log likelihood = -395.46006 Iteration 5: log likelihood = -395.20744 Iteration 6: log likelihood = -395.20707 Iteration 7: log likelihood = -395.20707

Maximum likelihood estimation

Log likelihood = -395.20707 Number of obs = 267

	Coef.	Std. Err.		P> z	[95% Conf.	Tn+on11
	Coei.	Std. Err.	Z	P> Z	[95% CONI.	Interval
/eta0	2.104716	.3198539	6.58	0.000	1.477814	2.731618
/eta1	.2025603	.0496982	4.08	0.000	.1051535	.2999671
/eta2	2002838	.0547262	-3.66	0.000	3075452	0930224
/beta0	-1.165861	.2565491	-4.54	0.000	-1.668688	6630343
/beta1	.5302016	.1499414	3.54	0.000	.2363219	.8240813