Fundamentals of Statistical Modeling (VT20)

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Lab 2

Load the dataset and the mlci command

- . version 14
- . use https://raw.githubusercontent.com/anddis/fsm/master/data/lab2.dta, clear
- . run https://raw.githubusercontent.com/anddis/fsm/master/do/mlci.do

Install the rcsgen command (you need to be connected to the Internet)

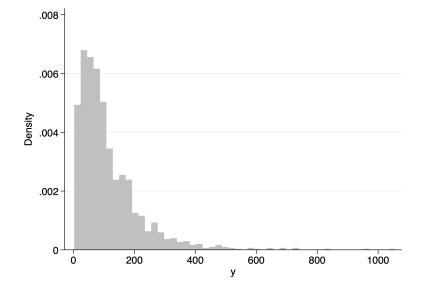
. cap net install http://fmwww.bc.edu/RePEc/bocode/r/rcsgen.pkg

Exercise 1

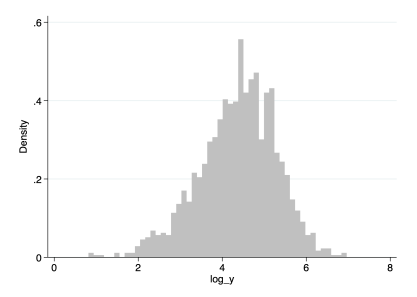
This dataset contains information on the blood concentration of a biomarker (y) in a random sample of 1432 subjects. Take a look at the histogram. What can we say about the distribution of this biomarker?

Plot also the histogram of log(y). How does the distribution of the biomarker after logarithmic transform look like?

```
. hist y, bin(50) name(p1, replace)
(bin=50, start=2.2592716, width=21.079531)
. graph export p1.png, replace
(file p1.png written in PNG format)
```



```
. gen log_y = log(y)
. hist log_y, bin(50) name(p2, replace)
(bin=50, start=.8150425, width=.12294848)
. graph export p2.png, replace
(file p2.png written in PNG format)
```



We assume that f(y) is gamma (see Lab 1). Estimate the parameters α and β using the gammaden() function. Fix the location parameter g (the third argument of the gammaden() function) to be equal to 0. Constrain α and β to be positive.

Note: the parameters α and β are not interpretable. We can reparametrise the gamma distribution so that one parameter is equal to its mean. This is described in the Extra material for Lab 2.

1,432

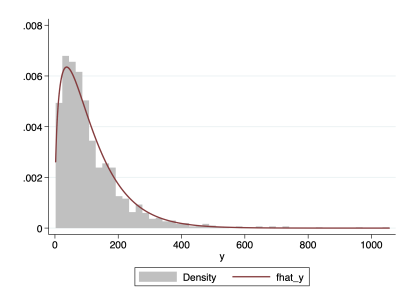
```
. local f = "gammaden(exp({theta1}), exp({theta2}), 0, y)"
. mlexp (log(`f'))
initial:
               log likelihood =
                                    -<inf>
                                             (could not be evaluated)
feasible:
               log likelihood = -96414.257
               log\ likelihood = -13891.173
rescale:
rescale eq:
               log likelihood = -13891.173
               log likelihood = -13891.173
Iteration 0:
Iteration 1:
               log\ likelihood = -8165.6417
               log likelihood = -8160.8897
Iteration 2:
Iteration 3:
               log likelihood = -8160.8781
Iteration 4:
               log likelihood = -8160.8781
Maximum likelihood estimation
Log likelihood = -8160.8781
                                                 Number of obs
```

	Coef.	Std. Err.	z	P> z	[95% Conf.	Interval]
/theta1 /theta2	.3906123 4.349872	.033984	11.49 107.83	0.000	.3240049 4.270804	.4572196 4.42894

```
. mlci exp /theta1
1.477885 95% CI: 1.382654, 1.579676
. mlci exp /theta2
77.46854 95% CI: 71.57918, 83.84246
```

Plot the estimated density $\hat{f}(y)$ over the sample histogram

```
. gen fhat_y = gammaden(exp(_b[/theta1]), exp(_b[/theta2]), 0, y)
. tw (hist y, bin(50)) (line fhat_y y, sort), name(p3, replace) legend(rows(1))
. graph export p3.png, replace
(file p3.png written in PNG format)
```



We assume that f(y) is log-normal distributed. That is, we assume that the biomarker is standard normal distributed after we apply the transform

$$G(y) = (\log(y) - \mu)/\sigma$$

The derivative of G(y) with respect to y is

$$G'(y) = g(y) = 1/(y\sigma).$$

Estimate the parameters μ and σ . Constrain σ to be positive.

```
. local sigma = "exp({theta})"
. local G = "(log(y) - {mu}) / `sigma'"
. local g = "(1 / y / `sigma´)"
. local f = "normalden(`G´)*`g´"
. mlexp (log(`f´))
               log likelihood = -21814.225
initial:
               log likelihood = -12440.421
alternative:
               log likelihood = -10178.274
rescale:
               \log likelihood = -8264.4966
rescale eq:
               \log likelihood = -8264.4966
Iteration 0:
               log likelihood = -8198.7786
Iteration 1:
               \log likelihood = -8159.1222
Iteration 2:
               log likelihood = -8158.8813
Iteration 3:
Iteration 4:
               log likelihood = -8158.8812
Maximum likelihood estimation
Log likelihood = -8158.8812
```

	Coef.	Std. Err.	z	P> z	[95% Conf.	Interval]
/mu /theta	4.365484 0868798		180.19 -4.65		4.318 1235034	4.412968 0502561

. mlci exp /theta .9167873 95% CI: .8838186, .9509858

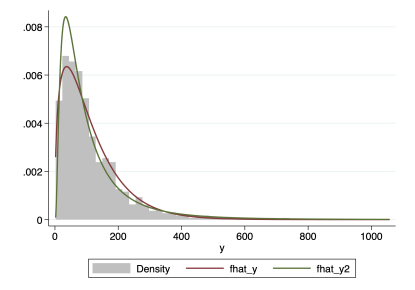
Compare the likelihood with that from the gamma model

Plot the estimated density $\hat{f}(y)$ over the sample histogram

Number of obs

1,432

```
. gen fhat_y2 = normalden((log(y) - _b[/mu]) / exp(_b[/theta]))*(1 / y / exp(_b[/theta]))
. tw (hist y, bin(50)) (line fhat_y fhat_y2 y, sort), name(p4, replace) legend(rows(1))
. graph export p4.png, replace
(file p4.png written in PNG format)
```



We make the transform G(y) more flexible using polynomials. Consider the transform

$$G(y) = (\log(y) + \eta \log(y)^2 - \mu)/\sigma$$

The derivative of G(y) with respect to y is

$$G'(y) = g(y) = (1 + 2\eta \log(y)) / (\sigma y)$$

Estimate the parameters μ, σ, η . Constrain σ to be positive.

```
. local sigma = "exp({theta})"
. local G = "(\log(y)+\{eta\}*\log(y)^2 - \{mu\}) / \text{`sigma'}"
. local g = "(1 + \{eta\}*2*log(y)) / (`sigma'*y)"
. local f = "normalden(`G´)*`g´"
. mlexp (log(`f'))
               log likelihood = -21814.225
initial:
               \log likelihood = -62186.361
alternative:
               log likelihood = -21814.225
rescale:
               log likelihood = -21814.225
rescale eq:
Iteration 0:
               \log likelihood = -21814.225
                                             (not concave)
               \log likelihood = -10891.752
Iteration 1:
                                             (not concave)
Iteration 2:
               \log likelihood = -8663.3365
               log likelihood = -8355.3282
Iteration 3:
               log likelihood = -8210.2164
Iteration 4:
               \log = -8167.6715
Iteration 5:
               log\ likelihood = -8151.9226
Iteration 6:
Iteration 7:
               log likelihood = -8142.8803
               log likelihood = -8139.5952
Iteration 8:
Iteration 9:
               log\ likelihood = -8138.8787
               log likelihood = -8138.5163
Iteration 10:
               log likelihood = -8138.5033
Iteration 11:
Iteration 12:
               log likelihood = -8138.5022
Iteration 13: log likelihood = -8138.5022
Maximum likelihood estimation
Log likelihood = -8138.5022
                                                 Number of obs
```

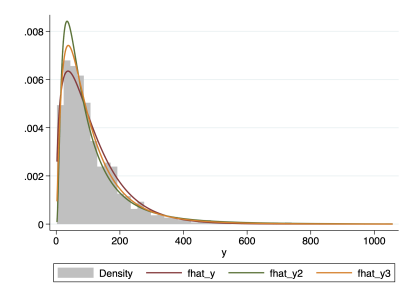
1,432

	Coef.	Std. Err.	z	P> z	[95% Conf.	Interval]
/eta	. 1804595	.0689773	2.62	0.009	.0452665	.3156524
/mu	7.956257	1.37386	5.79	0.000	5.263542	10.64897
/theta	.8361374	.2318207	3.61	0.000	.3817772	1.290498

[.] mlci exp /theta 2.307437 95% CI: 1.464886, 3.634595

Plot the estimated density $\hat{f}(y)$ over the sample histogram

```
. gen fhat_y3 = normalden((log(y)+_b[/eta]*log(y)^2 - _b[/mu])/exp(_b[/theta])) * ///
> (1+_b[/eta]*2*log(y)) / (exp(_b[/theta]) * y)
. tw (hist y, bin(50)) (line fhat_y fhat_y2 fhat_y3 y, sort), name(p5, replace) legend(rows(1))
. graph export p5.png, replace
(file p5.png written in PNG format)
```



Exercise 5

Instead of a quadratic term, we add two restricted cubic splines transforms of $\log(y)$: $V_2(\log(y))$ and $V_3(\log(y))$. We consider the transform

$$G(y) = (\log(y) + \eta_1 V_2(\log(y)) + \eta_2 V_3(\log(y)) - \mu)/\sigma$$

The derivative of G(y) with respect to y is

$$G'(y) = g(y) = (1 + \eta_1 v_2(\log(y)) + \eta_2 v_3(\log(y))) / (\sigma y)$$

Estimate the parameters μ , σ , η_1 , η_2 . Constrain σ to be positive. Jointly test the 2 parameters η_1 , η_2 to assess whether adding the 2 RCS transforms improves the fit of this model with respect to the "basic" log-normal model (see Exercise 3).

```
. rcsgen log_y, gen(V) dgen(v) df(3)
Variables V1 to V3 and v1 to v3 were created
. local sigma = "exp({theta})"
. local G = "(log(y)+{eta1}*V2+{eta2}*V3-{mu})/`sigma´"
. local g = "(1+{eta1}*v2+{eta2}*v3)/(`sigma´*y)"
. local f = "normalden(`G´)*`g´"
. mlexp (log(`f´))
```

```
log likelihood = -21814.225
initial:
               \log likelihood = -21814.225
final:
rescale:
               \log likelihood = -21814.225
               log\ likelihood = -21814.225
                                             (not concave)
Iteration 0:
Iteration 1:
               log likelihood = -15749.959
                                             (not concave)
Iteration 2:
               log likelihood = -11575.393
                                             (not concave)
               log likelihood = -8930.8791
Iteration 3:
                                             (not concave)
Iteration 4:
               log likelihood = -8353.802
               log \ likelihood = -8226.7656
Iteration 5:
Iteration 6:
               log\ likelihood = -8161.3477
               log likelihood = -8141.3375
Iteration 7:
               log likelihood = -8137.886
Iteration 8:
Iteration 9:
               log likelihood = -8137.333
               log likelihood = -8137.3305
Iteration 10:
Iteration 11:
               log likelihood = -8137.3305
```

Maximum likelihood estimation

Log likelihood = -8137.3305

/eta1 /eta2

/mu /theta

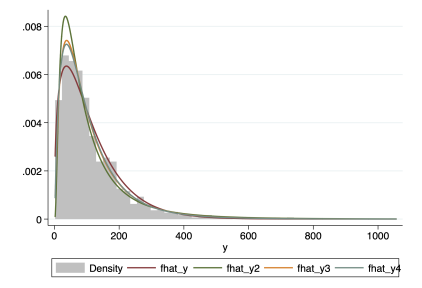
Coef.	Std. Err.	z	P> z	[95% Conf.	Interval]
0080713	.0421601	-0.19	0.848	0907036	.074561
0259272	.0451895	-0.57	0.566	114497	.0626426
5.045942	.2441727	20.67	0.000	4.567372	5.524512
.2911407	.1009571	2.88	0.004	.0932684	.489013

Number of obs

1,432

Plot the estimated density $\hat{f}(y)$ over the sample histogram

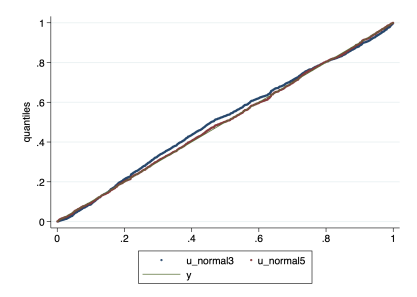
```
. gen fhat_y4 = normalden((log(y)+_b[/eta1]*V2+_b[/eta2]*V3 - _b[/mu])/exp(_b[/theta])) * ///
> (1+_b[/eta1]*v2+_b[/eta2]*v3) / (exp(_b[/theta]) * y)
. tw (hist y, bin(50)) (line fhat_y fhat_y2 fhat_y3 fhat_y4 y, sort), name(p6, replace) legend(rows(1))
. graph export p6.png, replace
(file p6.png written in PNG format)
```



Let's assess the goodness-of-fit of the log-normal model with RCS transforms (see Exercise 5) and of the log-normal model (see Exercise 3) using a quantile plot.

```
. gen u_normal5 = normal((log(y)+_b[/eta1]*V2+_b[/eta2]*V3 - _b[/mu])/exp(_b[/theta]))
. // Re-fit log-normal model (Exercise 3)
. local sigma = "exp({theta})"
. local G = "(log(y) - {mu}) / `sigma`"
. local g = "(1 / y / `sigma´)"
. local f = "normalden(`G´)*`g´"
. mlexp (log(`f´))
initial:
               log likelihood = -21814.225
               \log = -12440.421
alternative:
               log likelihood = -10178.274
rescale:
rescale eq:
               log likelihood = -8264.4966
               log likelihood = -8264.4966
Iteration 0:
               \log likelihood = -8198.7786
Iteration 1:
               log likelihood = -8159.1222
Iteration 2:
               log likelihood = -8158.8813
Iteration 3:
Iteration 4:
               log\ likelihood = -8158.8812
Maximum likelihood estimation
Log likelihood = -8158.8812
                                                 Number of obs
                                                                           1,432
                    Coef.
                            Std. Err.
                                            z
                                                 P>|z|
                                                            [95% Conf. Interval]
                                                                       4.412968
                 4.365484
                             .0242269
                                        180.19
                                                 0.000
                                                              4.318
         /mu
      /theta
                -.0868798
                            .0186859
                                         -4.65
                                                 0.000
                                                          -.1235034
                                                                       -.0502561
```

```
. gen u_normal3 = normal((log(y) - _b[/mu])/exp(_b[/theta]))
.
. qplot u_normal3 u_normal5, addplot(function y = x, lw(medthin)) name(p7, replace) ///
> msym(0h 0h) msize(tiny tiny)
. graph export p7.png, replace
(file p7.png written in PNG format)
```



Extra: Exercise 7

We now assume that f(y) is gamma-distributed after square root transform.

$$G(y) = \sqrt{y}$$

The derivative is

$$G'(y) = g(y) = 0.5/\sqrt{y}$$

Estimate the parameters α and β using the gammaden() function. Fix the location parameter g to be equal to 0. Constrain α and β to be positive. Compare the likelihood with that form the log-normal and gamma models

```
. local G = "sqrt(y)"
. local g = "(0.5 / sqrt(y))"
. local f = \text{"gammaden}(\exp(\{\text{theta1}\}), \exp(\{\text{theta2}\}), 0, G') * G') * G')
. mlexp (log(`f'))
initial:
                log likelihood = -18140.526
                log likelihood = -11624.987
alternative:
rescale:
                \log likelihood = -8442.1703
                log likelihood = -8442.1703
rescale eq:
Iteration 0:
                log likelihood = -8442.1703
                \log = -8185.3624
Iteration 1:
Iteration 2:
                log\ likelihood = -8138.5993
Iteration 3:
                \log likelihood = -8138.2943
                log likelihood = -8138.2942
Iteration 4:
Maximum likelihood estimation
Log likelihood = -8138.2942
                                                   Number of obs
                                                                               1,432
                              Std. Err.
                                                               [95% Conf. Interval]
                                                   P>|z|
                     Coef.
                                              z
     /theta1
                  1.652508
                              .0362406
                                           45.60
                                                   0.000
                                                              1.581478
                                                                           1.723538
                                                                            .7036256
     /theta2
                   .6290655
                              .0380415
                                           16.54
                                                   0.000
                                                               .5545055
```

```
. mlci exp /theta1
5.220056 95% CI: 4.862136, 5.604323
. mlci exp /theta2
1.875857 95% CI: 1.74108, 2.021067
```

Plot the estimated density $\hat{f}(y)$ over the sample histogram. Visually compare the estimated density from the lognormal + splines model with the density from the gamma model after square root transform. What do you conclude?

```
. gen fhat_y5 = gammaden(exp(_b[/theta1]), exp(_b[/theta2]), 0, sqrt(y))*(.5 / sqrt(y))
. tw (hist y, bin(50)) (line fhat_y fhat_y2 fhat_y3 fhat_y4 fhat_y5 y, sort), name(p8, replace) legend(rows(1))
. graph export p8.png, replace
(file p8.png written in PNG format)
```

