

ReportRevisited

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Statistical Analysis of Covariates

1. Age

The age of the patient in days.

1.1 Model selection

The data seems to follow a poisson distribution $Poi(\lambda)$. Using the non informative Jeffreys prior, we can derive that the posterior for the parameter λ is Gamma distributed.

1.2 Results

The posterior distribution is:

```
## Gamma( 5700066 , 312 )
```

With mean and variance:

```
## mean: 18269.44 , variance: 58.55591
```

The 95% HDI interval calculates to:

```
##      lower      upper
## 18254.57 18284.63
## attr(,"credMass")
## [1] 0.95
```

2. Sex

The sex of the patients is encoded in a binary variable, where 0 means *male* and 1 means *female*.

2.1 Model selection

We assume a Bernoulli model $Ber(\theta)$ for the sex of the patient conditional on one parameter θ , the probability of the patient to be female. The density function is given by

$$f(x|\theta) = \theta^x(1 - \theta)^{1-x}, \quad (1)$$

where $x \in \{0, 1\}$. As a prior distribution for θ we use the natural conjugate family of the Bernoulli distribution, namely the Beta distribution, $Beta(a, b)$, with two shape parameters $a = b = 2$ to give more weight to the middle of the interval $[0, 1]$, knowing how females and males are represented in the general population. The density is given by

$$h(\theta) = \frac{\Gamma(a+b)}{\Gamma(a)\Gamma(b)} \theta^{a-1} (1-\theta)^{b-1}, \quad (2)$$

for $\theta \in [0, 1]$.

2.2 Results

From the given dataset we get the sample size n and the sum of the observations s :

```
n<-length(mydata$V6[!is.na(mydata$V6)])
s<-sum(mydata$V6)
n
```

```
## [1] 312
```

```
s
```

```
## [1] 276
```

Therefore the posterior distribution is $Beta(2 + s, 2 + n - s)$, which turns out to be $Beta(278, 38)$. From that we get

```
## Posterior mean: 0.8797468
```

```
## Posterior mode: 0.8821656
```

```
## Centered 95% Confidence Interval: [ 0.8417454 , 0.9132003 ]
```

And the HPD confidence Interval calculates to:

```
tst<-rbeta(1e5,278,38)
hdi(tst)
```

```
##      lower      upper
## 0.8440046 0.9149565
## attr(,"credMass")
## [1] 0.95
```

3 Ascictes

3.1 Model selection

Since the Ascictes - covariate has a 0-1 outcome we can assume that it is Bernoulli distributed with parameter θ . A natural conjugate prior for the Bernoulli distribution is the Beta distribution. The posterior beta distribution for the parameter is given by

$$Beta(\theta|a + \sum_{i=1}^n x_i, b + n - \sum_{i=1}^n x_i)$$

3.2 Results

The Beta distribution reduces to a simple uniform distribution. Since we have minimal prior knowledge about the parameters we will continue with a prior distribution that is uniformly distributed.

The posterior distribution has the following calculated information:

```
## Posterior mean: 0.08227848  
  
## Posterior mode: 0.07961783  
  
## Centered 95% Confidence Interval: [ 0.05235453 , 0.1119428 ]
```

With the HPD interval:

```
##      lower      upper  
## 0.05079471 0.11003097  
## attr(,"credMass")  
## [1] 0.95
```

4. Hepatomegaly

The presence of hepatomegaly is encoded in a Binary variable, where 1 means hepatomegaly is present.

4.1 Model selection

We assume a Bernoulli model $Ber(\theta)$ for the presence of hepatomegaly in the patient, conditional on one parameter θ , the probability of the presence of hepatomegaly in the patient. The density function is given as stated earlier. As a prior distribution for θ we use the natural conjugate family of the Bernoulli distribution, namely the Beta distribution, $Beta(a, b)$, with two shape parameters $a = b = 1$, because we have no prior information. The density is given as above.

4.2 Results

From the given dataset we get the sample size n and the sum of the observations s :

```
n<-length(mydata$V8[!is.na(mydata$V8)])  
s<-sum(mydata$V8)  
n
```

```
## [1] 312
```

```
s
```

```
## [1] 160
```

Therefore the posterior distribution is $Beta(1 + s, 1 + n - s)$, which turns out to be $Beta(161, 153)$. From that we get

```
## Posterior mean: 0.5126582
```

```
## Posterior mode: 0.5127389
```

```
## Centered 95% Confidence Interval: [ 0.4575015 , 0.5678225 ]
```

And the HPD confidence interval calculates to:

```
tst<-rbeta(1e5,161,153)
hdi(tst)
```

```
##      lower      upper
## 0.4574971 0.5675879
## attr(,"credMass")
## [1] 0.95
```

5. Spiders

The presence of spiders is encoded in a Binary variable, where 1 means spiders are present.

5.1 Model selection

We assume a Bernoulli model $Ber(\theta)$ for the presence of spiders in patients conditional on one parameter θ , the probability of the presence of spiders in the patient. The density function is given as stated earlier. As a prior distribution for θ we use the natural conjugate family of the Bernoulli distribution, namely the Beta distribution, $Beta(a, b)$, with two shape parameters $a = b = 1$, because we have no prior information. The density is given as above.

5.2 Results

From the given dataset we get the sample size n and the sum of the observations s :

```
n<-length(mydata$V9[!is.na(mydata$V9)])
s<-sum(mydata$V9)
n
```

```
## [1] 312
```

```
s
```

```
## [1] 90
```

Therefore the posterior distribution is $Beta(1 + s, 1 + n - s)$, which turns out to be $Beta(91, 223)$. From that we get

```
## Posterior mean: 0.2911392
```

```
## Posterior mode: 0.2898089
```

```
## Centered 95% Confidence Interval: [ 0.2410228 , 0.341131 ]
```

And the HPD confidence interval calculates to:

```
tst<-rbeta(1e5,91,223)
hdi(tst)
```

```
##      lower      upper
## 0.2411227 0.3411279
## attr("credMass")
## [1] 0.95
```

6. Edema

The presence of edema is coded into (0, 0.5, 1). Where 0 is the class corresponding to no edema and no diuretic therapy for edema. The class 0.5 corresponds to edema present without diuretics, or edema resolved by diuretics. Lastly the class 1 corresponds to edema despite diuretic therapy.

```
##  0 0.5  1
## 263  29  20
```

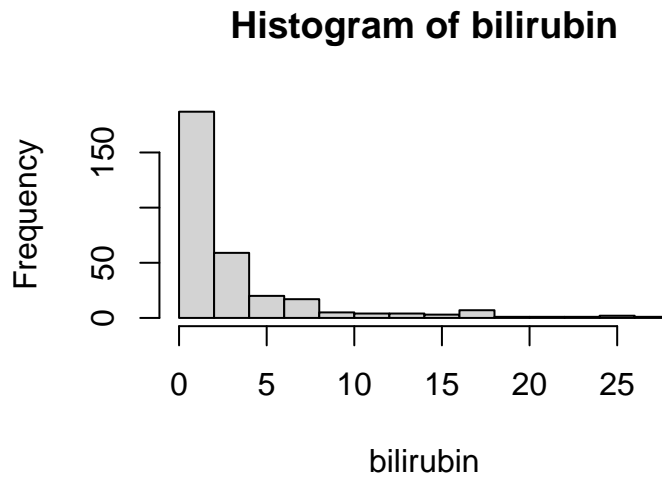
We observe from the frequencies that most of the patients don't have edema and do not go to diuretic therapy for edema.

7. Bilirubin

The serum bilirubin of the patients is given in mg/dl.

7.1 Model Selection

We assume by inspecting the histogram plot,



that the data follows a exponential distribution with parameter λ . Density is given by

$$f(x|\lambda) = \lambda e^{-\lambda x} \quad (3)$$

As a prior for λ we use, the jeffreys non-informative prior, namely: $h(\lambda) \propto \frac{1}{\lambda}$.

7.2 Results

From the data we get the number of samples n and the sum of the samples s as

```
## [1] 312
```

```
## [1] 1015.9
```

That means the posterior distribution for λ is $Gamma(n, s)$. Which turns out to be $Gamma(312, 1015.9)$. From that we get

```
## Posterior mean: 0.3071168
```

```
## Posterior mode: 0.3061325
```

```
## Centered 95% Confidence Interval: [ 0.2739805 , 0.3421174 ]
```

And the HPD confidence interval calculates to:

```
tst<-rgamma(1e5,312,1015.9)
hdi(tst)
```

```
##      lower      upper
## 0.2742943 0.3421982
## attr(,"credMass")
## [1] 0.95
```

8. Cholesterol

8.1 Model selection

We assume that the data is sampled from a poisson, $Poi(\lambda)$, distribution, and we use the non informative Jeffreys prior for the rate parameter.

8.2 Results

The posterior distribution is:

```
## Gamma( 104941.5 , 312 )
```

With mean and variance:

```
## mean: 336.351 , variance: 1.078048
```

The 95% HDI interval calculates to:

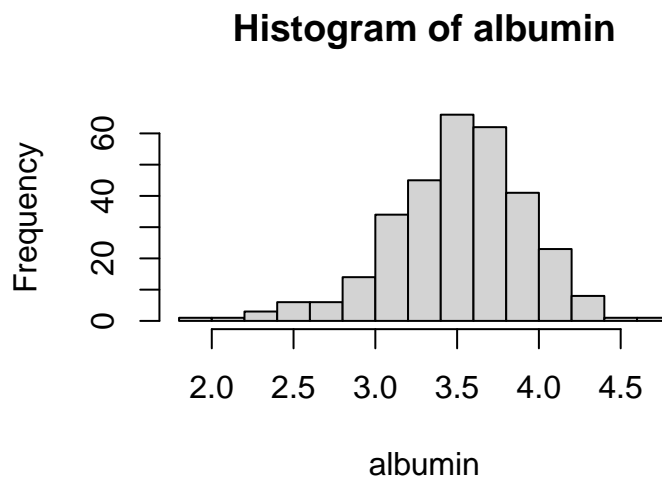
```
##      lower      upper  
## 334.2887 338.3643  
## attr(,"credMass")  
## [1] 0.95
```

9. Albumin

The Albumin is given in mg/dl.

9.1 Model selection

By the histogram plot of the data we see,



that albumin could be gamma distributed with shape and rate parameters a and b . We assume prior independence between a and b and use the marginal prior distributions $Gamma(0.001, 0.001)$ for both of them.

9.2 Results

Using OpenBUGS and MCMC methods, we get posterior information about the parameters a and b :

```
n<-length(albumin[!is.na(albumin)])
X<-albumin
data1<-list("X","n")
params<-c("a" , "b")
inits<-list(a=1,b=1)
fit1<-bugs(data=data1,inits=list(inits),parameters.to.save=params,"model_albu.txt",n.chains=1, n.iter=20000)
fit1$summary
```

```
##              mean          sd   2.5%   25%   50%   75%   97.5%
## a           66.32709 4.995273  57.66  62.76  66.30  69.38  77.1405
## b           18.84450 1.424988  16.37  17.83  18.84  19.71  21.9400
## deviance    365.10028 1.880193 363.30 363.70 364.50 365.80 370.1000
```

And the HPD confidence interval for a calculates to:

```
## lower upper
## 56.88 76.12
## attr("credMass")
## [1] 0.95
```

whereas the HPD confidence interval for b is

```
## lower upper
## 16.19 21.69
## attr("credMass")
## [1] 0.95
```

10. Urine

Urine copper in ug/day

10.1 Model selection

By observing the data we see that it seems to follow a poisson distribution.

10.2 Results

Using the non informative jeffreys prior we get the following posterior results:

The posterior distribution is:

```
## Gamma( 30271.5 , 312 )
```


With mean and variance:

```
## mean: 97.02404 , variance: 0.3109745
```

The 95% HDI interval calculates to:

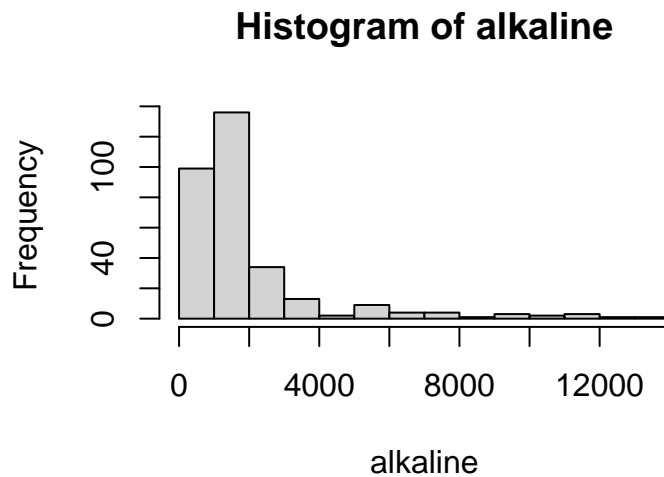
```
##      lower      upper
## 95.93014 98.10755
## attr(,"credMass")
## [1] 0.95
```

11. Alkaline

The data contains the units of alkaline phosphatase per liter of the patients.

11.1 Model selection

Since the units of alkaline per liter are integers, we assume that it is a counting process. Therefore we want to assume, that the data is poisson distributed conditional on one parameter λ . The histogram plot justifies our assumptions:



The density function of a single observation is given as

$$f(x|\lambda) = e^{-\lambda} \frac{\lambda^x}{x!} \quad (4)$$

As a prior for λ we use the natural conjugate prior of the poisson distribution which is the gamma distribution. To not give a lot of prior information, we will use $\text{Gamma}(0.001, 0.001)$.

11.2 Results

From our data we get the sample size n and the sum s over the sample:

```
## [1] 312
```

```
## [1] 618588.6
```

The posterior distribution for λ is given by $\text{Gamma}(s + 0.001, n + 0.001)$ which in our case results to $\text{Gamma}(618588.601, 312.001)$. This yields to:

```
## Posterior mean: 1982.649
```

```
## Posterior mode: 1982.646
```

```
## Centered 95% Confidence Interval: [ 1977.712 , 1987.593 ]
```

And the HPD confidence interval calculates to:

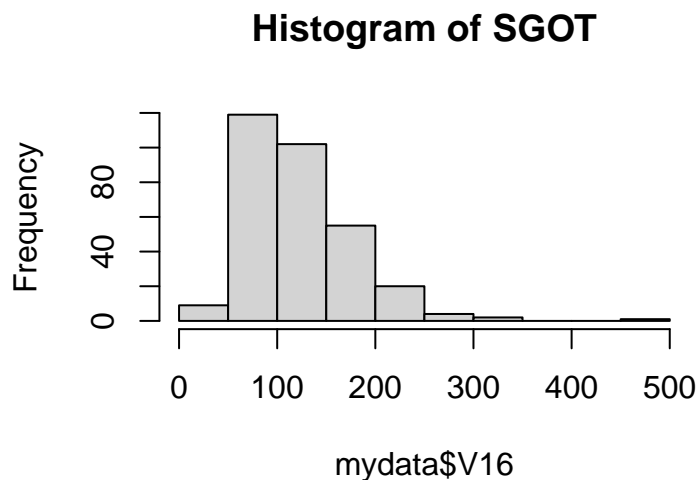
```
tst<-rgamma(1e5,618588.601,312.001)
hdi(tst)
```

```
##      lower      upper
## 1977.692 1987.606
## attr(,"credMass")
## [1] 0.95
```

12 SGOT

Data provided in U/ml

12.1 Model selection



By inspecting the data and the histogram plot we can make the assumption that the data is sampled from a Gamma distribution with parameters a and b . Using MCMC methods included in the OpenBUGS library we can estimate the posterior parameters. We assume prior parameters $a = b = 0.001$

12.2 Results

```
##              mean          sd        2.5%        25%        50%        75%
## a          5.2262694 0.396605907    4.49295    4.94775 5.207e+00    5.4810
## b          0.0426461 0.003383706    0.03634    0.04029 4.252e-02    0.0448
## deviance 3330.5929000 1.910794260 3329.00000 3329.00000 3.330e+03 3331.0000
##              97.5%
## a          6.042025
## b          0.049610
## deviance 3336.0000000
```

with the following HDI for the parameters a and b respectively:

```
hdi(fit1$sims.list$a)
```

```
## lower upper
## 4.467 6.012
## attr(,"credMass")
## [1] 0.95
```

```
hdi(fit1$sims.list$b)
```

```
## lower upper
## 0.03603 0.04927
## attr(,"credMass")
## [1] 0.95
```

13. Triglicerides

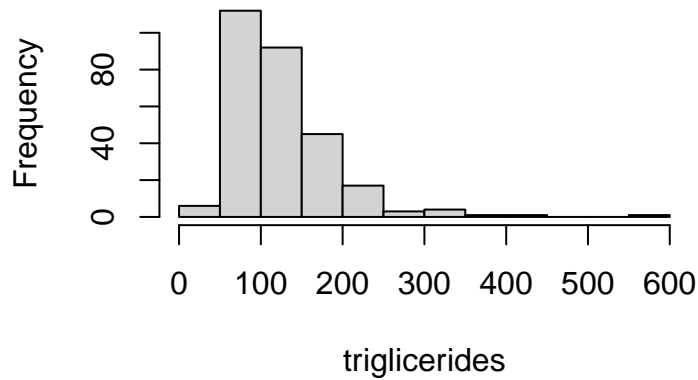
Triglicerides of the patients in mg/dl.

13.1 Model selection

By the histogram plot of the data we see,

```
## Warning: NAs introduced by coercion
```

Histogram of triglicerides



that triglicerides could be gamma distributed with shape and rate parameters a and b . We assume prior independence between a and b and use the marginal prior distributions $Gamma(0.001, 0.001)$ for both of them.

13.2 Results

By OpenBUGS and MCMC methods we get posterior information about the parameters a and b :

```
##           mean          sd        2.5%        25%        50%        75%
## a      4.742428e+00  0.410163943  3.988975e+00  4.457e+00  4.7310e+00  5.010e+00
## b      3.775649e-02  0.003443284  3.136975e-02  3.535e-02  3.7645e-02  4.001e-02
## deviance 2.732715e+03  1.974568784  2.731000e+03  2.731e+03  2.7320e+03  2.734e+03
##           97.5%
## a           5.58705
## b           0.04484
## deviance 2738.00000
```

And the HPD confidence interval for a calculates to:

```
## lower upper
## 3.939 5.522
## attr(,"credMass")
## [1] 0.95
```

whereas the HPD confidence interval for b is

```
## lower upper
## 0.03086 0.04419
## attr(,"credMass")
## [1] 0.95
```

14. Platelets

Data given in per cubic ml / 1000.

14.1 Model selection

The data resembles a poisson distribution as it is measured in integers. Using the non informative prior we can calculate the posterior gamma distribution for the rate parameter.

14.2 Results

The posterior distribution is:

```
## Gamma( 80676.5 , 312 )
```

With mean and variance:

```
## mean: 258.5785 , variance: 0.8287773
```

The 95% HDI interval calculates to:

```
##      lower      upper
## 256.8118 260.3795
## attr(,"credMass")
## [1] 0.95
```

15. Prothrombin

Prothrombin data given in seconds.

15.1 Model selection

By inspecting the data and histogram we can assume that it follows a Gamma distribution. Again using MCMC methods with the OpenBUGS library we can estimate the posterior information on the parameters.

15.2 Results

```
##           mean      sd      2.5%    25%    50%    75%    97.5%
## a      125.61267 9.1158275 107.90000 119.40 125.30 131.60 143.6025
## b       11.71168 0.8513285  10.04975  11.13  11.68  12.27  13.4000
## deviance 861.41920 1.8145938 859.60000 860.10 860.90 862.20 866.1000
```

The 95% HDI for the a - parameter:

```
## lower upper
## 109.6 144.7
## attr(,"credMass")
## [1] 0.95
```

The 95% HDI for the b-parameter:

```
## lower upper
## 10.21 13.49
## attr(,"credMass")
## [1] 0.95
```

16. Histologic stage

The Histologic stage of the disease is a number in $\{1, 2, 3, 4\}$, where the stage increases with severeness. We will give here the frequencies of the stages in the dataset.

```
##      1      2      3      4
##    16    67   120   109
```

We see that, most patients have been diagnosed in the last to stages of the disease.

Weibull Survival Analysis

We will use a survival model, based on a hazard function, conditional on regression parameters and dependent on (now assumed) deterministic covariates. The hazard function is given by

$$\lambda(t|\alpha, \beta, \delta) = \delta \alpha t^{\alpha-1} e^{\beta^T z} \quad (5)$$

where z is the covariate vector. As prior distribution for the regression parameters β we will use normal distributions with 0 mean and $\sigma^2 = 1000$. For the parameters α and δ we use *Gamma*(0.001, 0.001) prior distribution. Prior independence is assumed. MCMC methods and OpenBUGS help us to get inference about our parameters:

```
##              mean          sd          2.5%          25%          50%
## alpha      2.174216e+01 3.279157e-01 21.230000 2.154e+01 21.660000
## beta[1]    7.799742e+00 1.180215e+00  6.199000 6.517e+00  7.912000
## beta[2]   -2.259021e-02 2.066046e-04 -0.023050 -2.269e-02 -0.022580
## beta[3]   -2.748310e+01 2.422248e+00 -30.920000 -2.956e+01 -28.270000
## beta[4]    4.224655e+01 5.318714e+00 36.200000 3.697e+01 40.950000
## beta[5]   -9.287410e+00 1.300432e+00 -11.160000 -1.046e+01 -9.569000
## beta[6]    1.861271e+00 1.731042e+00 -1.003000 5.056e-01  1.479000
## beta[7]    3.613975e+01 4.380977e+00 28.350000 3.308e+01 37.680000
## beta[8]    4.166480e+00 1.274159e-01  3.901000 4.090e+00  4.230000
## beta[9]   -2.383417e-02 3.570780e-03 -0.028280 -2.757e-02 -0.024730
## beta[10]   1.305721e+00 2.105063e-01  0.921900 1.137e+00  1.284488
## beta[11]   3.482789e-02 5.964091e-03  0.022470 2.999e-02  0.035490
## beta[12]   3.496828e-03 9.524787e-05  0.003286 3.463e-03  0.003507
## beta[13]   3.966723e-02 1.293091e-02  0.018890 2.672e-02  0.038270
## beta[14]   1.694033e-02 4.875011e-03  0.009749 1.232e-02  0.016340
## beta[15]   6.666559e-02 3.990902e-03  0.058490 6.381e-02  0.066980
## beta[16]   2.283336e+00 1.894305e-01  1.947000 2.145e+00  2.351000
## beta[17]   7.395990e+00 4.670291e-01  6.368000 7.128e+00  7.461000
## delta      8.400426e+03 2.982289e+03 3532.899945 6.224e+03 8142.499985
##              75%          97.5%
## alpha      2.195e+01    22.440000
## beta[1]    8.706e+00     9.579000
## beta[2]   -2.244e-02    -0.022370
## beta[3]   -2.598e+01   -22.090000
## beta[4]    4.619e+01    51.510000
## beta[5]   -7.931e+00    -7.351000
## beta[6]    3.164e+00     5.114000
## beta[7]    4.019e+01    41.210000
## beta[8]    4.264e+00     4.296000
```

```
## beta[9] -2.046e-02 -0.017640
## beta[10] 1.448e+00 1.806324
## beta[11] 4.028e-02 0.043520
## beta[12] 3.584e-03 0.003609
## beta[13] 5.297e-02 0.056200
## beta[14] 2.210e-02 0.024500
## beta[15] 6.847e-02 0.074350
## beta[16] 2.432e+00 2.517000
## beta[17] 7.557e+00 8.373000
## delta 1.021e+04 15100.499677
```

By applying Heidelberg and Welchs method to decide whether the simulated values from the markov chain come from its stationary distribution we get

```
##
## Stationarity start p-value
## test iteration
## alpha passed 1 0.1035
## beta[1] passed 1 0.1408
## beta[2] passed 1 0.0608
## beta[3] passed 1 0.1588
## beta[4] passed 1 0.0855
## beta[5] passed 1 0.3365
## beta[6] passed 1 0.1565
## beta[7] passed 1 0.0845
## beta[8] passed 1 0.0843
## beta[9] passed 1 0.1377
## beta[10] passed 1 0.0556
## beta[11] passed 1 0.1836
## beta[12] passed 1 0.0764
## beta[13] passed 1 0.1897
## beta[14] passed 1 0.1843
## beta[15] passed 1 0.2674
## beta[16] passed 1 0.1231
## beta[17] passed 1 0.1515
## delta passed 1 0.9920
##
## Halfwidth Mean Halfwidth
## test
## alpha passed 21.7422 8.90e-03
## beta[1] passed 7.7997 3.18e-02
## beta[2] passed -0.0226 5.73e-06
## beta[3] passed -27.4831 6.71e-02
## beta[4] passed 42.2465 1.47e-01
## beta[5] passed -9.2874 3.52e-02
## beta[6] passed 1.8613 4.80e-02
## beta[7] passed 36.1397 1.21e-01
## beta[8] passed 4.1665 3.53e-03
## beta[9] passed -0.0238 9.90e-05
## beta[10] passed 1.3057 5.83e-03
## beta[11] passed 0.0348 1.61e-04
## beta[12] passed 0.0035 2.64e-06
## beta[13] passed 0.0397 3.48e-04
## beta[14] passed 0.0169 1.31e-04
```

```
## beta[15] passed      0.0667 1.11e-04
## beta[16] passed      2.2833 5.25e-03
## beta[17] passed      7.3960 1.29e-02
## delta   passed      8400.4258 8.27e+01
```

To assess the quality of our model, we compute standardized predictive residuals for uncensored data. For a uncensored survival time $y(z)$, dependent of a covariate vector z , and the posterior predictive $Y(z)|x$, where x is the data used in the bayesian analysis, we define the standardized predictive residual as

$$d = \frac{y(z) - E(Y(z)|x)}{\sqrt{Var(Y(z)|x)}} \quad (6)$$

```
d<-abs((t[cens==1]-fit$mean$Y[cens==1])/fit$sd$Y[cens==1])
sum(d)
```

```
## [1] 924.7143
```

The sum of the absolute values of the predictive residuals indicates the quality of the model. Now we will remove covariates with small regression coefficient means, trying to find better model fits. For the second model we kick out covariates for Age, Spiders and Cholesterol:

##	mean	sd	2.5%	25%	50%
## alpha	1.649426e+00	1.052831e-01	1.461975e+00	1.57700e+00	1.6460e+00
## beta[1]	-2.252330e-01	1.917173e-01	-6.177000e-01	-3.44100e-01	-2.1810e-01
## beta[2]	-6.350418e-01	2.853519e-01	-1.158000e+00	-8.25200e-01	-6.1830e-01
## beta[3]	1.363722e-01	3.443228e-01	-5.235325e-01	-1.03800e-01	1.4230e-01
## beta[4]	4.627324e-02	2.333311e-01	-3.707000e-01	-1.22600e-01	3.5020e-02
## beta[5]	8.125997e-01	3.626407e-01	6.101000e-02	5.78000e-01	8.1200e-01
## beta[6]	8.961414e-02	2.218430e-02	4.581649e-02	7.57300e-02	9.0530e-02
## beta[7]	-1.112314e+00	1.375734e-01	-1.375000e+00	-1.22425e+00	-1.1020e+00
## beta[8]	2.601600e-03	1.287428e-03	2.668000e-04	1.65200e-03	2.6230e-03
## beta[9]	-8.693655e-06	2.324338e-05	-5.488000e-05	-3.48800e-05	-7.2280e-06
## beta[10]	4.443800e-03	1.676903e-03	1.236000e-03	3.23900e-03	4.5100e-03
## beta[11]	-1.503449e-03	1.128316e-03	-3.870000e-03	-2.22500e-03	-1.4310e-03
## beta[12]	7.945353e-04	8.782255e-04	-1.221000e-03	2.35600e-04	9.0440e-04
## beta[13]	3.552868e-01	8.420755e-02	2.295000e-01	2.87900e-01	3.4320e-01
## beta[14]	3.650015e-01	1.386479e-01	9.645000e-02	2.69800e-01	3.6380e-01
## delta	4.836558e-07	5.736641e-07	3.936725e-08	1.19850e-07	2.6985e-07
##	75%	97.5%			
## alpha	1.717000e+00	1.88100e+00			
## beta[1]	-1.080000e-01	2.14200e-01			
## beta[2]	-4.467000e-01	-9.81300e-02			
## beta[3]	3.883000e-01	7.80700e-01			
## beta[4]	1.976000e-01	5.17400e-01			
## beta[5]	1.069000e+00	1.51510e+00			
## beta[6]	1.034000e-01	1.32000e-01			
## beta[7]	-9.895000e-01	-8.83800e-01			
## beta[8]	3.518000e-03	4.96900e-03			
## beta[9]	5.955000e-06	3.55400e-05			
## beta[10]	5.478000e-03	7.92200e-03			
## beta[11]	-6.948000e-04	3.05300e-04			
## beta[12]	1.408000e-03	2.33400e-03			
## beta[13]	4.490000e-01	4.95600e-01			
## beta[14]	4.535748e-01	6.45800e-01			
## delta	6.129748e-07	2.08115e-06			


```
d<-abs((t[cens==1]-fit2$mean$Y[cens==1])/fit2$sd$Y[cens==1])
sum(d)
```

```
## [1] 86.33356
```

We see that our model improved a lot! We continue by neglecting, the used drug and the presence of asictes to get another model:

```
##           mean          sd          2.5%          25%          50%
## alpha      1.581573e+00 1.089669e-01 1.374000e+00 1.502e+00 1.5840e+00
## beta[1]    -5.828211e-01 2.793994e-01 -1.121000e+00 -7.695e-01 -6.0310e-01
## beta[2]    -1.293594e-02 2.500077e-01 -4.964000e-01 -1.837e-01 -1.5580e-02
## beta[3]     9.077950e-01 3.483444e-01 2.488875e-01 6.857e-01 8.9550e-01
## beta[4]     9.003871e-02 2.174679e-02 4.620000e-02 7.606e-02 9.0430e-02
## beta[5]    -6.600518e-01 2.015973e-01 -1.041000e+00 -7.971e-01 -6.0830e-01
## beta[6]     2.522762e-03 1.084539e-03 6.682000e-04 1.666e-03 2.5380e-03
## beta[7]     1.181760e-05 4.271823e-05 -8.048000e-05 -2.116e-05 3.1050e-05
## beta[8]     3.948636e-03 1.765410e-03 5.150000e-04 2.609e-03 4.0220e-03
## beta[9]    -1.073597e-03 1.220436e-03 -4.014000e-03 -1.969e-03 -1.1710e-03
## beta[10]    8.993215e-04 1.126889e-03 -1.229000e-03 1.418e-04 8.4730e-04
## beta[11]    3.469549e-01 7.395093e-02 1.973950e-01 3.058e-01 3.4630e-01
## beta[12]    5.915423e-01 2.030411e-01 2.857000e-01 4.444e-01 5.4790e-01
## delta      7.142071e-08 8.580466e-08 2.585975e-09 1.147e-08 4.1405e-08
##           75%          97.5%
## alpha      1.65425e+00 1.805000e+00
## beta[1]    -4.04600e-01 3.287000e-02
## beta[2]     1.50200e-01 4.822000e-01
## beta[3]     1.13500e+00 1.608000e+00
## beta[4]     1.04900e-01 1.324000e-01
## beta[5]    -5.57700e-01 -1.980000e-01
## beta[6]     3.41300e-03 4.503025e-03
## beta[7]     4.07300e-05 6.642000e-05
## beta[8]     5.01700e-03 7.350000e-03
## beta[9]    -1.97300e-04 1.089000e-03
## beta[10]    1.63700e-03 3.028000e-03
## beta[11]    4.15500e-01 4.585000e-01
## beta[12]    6.91100e-01 1.077000e+00
## delta      9.94150e-08 3.298324e-07
```

```
d<-abs((t[cens==1]-fit3$mean$Y[cens==1])/fit3$sd$Y[cens==1])
sum(d)
```

```
## [1] 84.9674
```

We see that our model improved a little! We continue by neglecting the sex, presence of hepatomegaly, alkaline, triglicerides and platelets:

```
##           mean          sd          2.5%          25%          50%          75%
## alpha      1.205862e+00 0.108954588 1.048000 1.092249 1.1860000 1.321000
## beta[1]    2.282041e-01 0.042702805 0.173300 0.194000 0.2177000 0.272000
## beta[2]    1.280806e-01 0.006933680 0.114300 0.121200 0.1301000 0.134100
```

```
## beta[3] -6.530799e-01 0.062296408 -0.734200 -0.717500 -0.6456000 -0.584300
## beta[4] 3.699941e-03 0.001037102 0.001572 0.003102 0.0036120 0.004464
## beta[5] 2.176426e-05 0.001763321 -0.003461 -0.001087 0.0003246 0.001032
## beta[6] -4.412231e-01 0.044021390 -0.503700 -0.479000 -0.4541000 -0.399100
## beta[7] 3.896612e-01 0.039671153 0.332200 0.348300 0.3845000 0.431800
## delta 6.029195e-03 0.002624320 0.002445 0.004107 0.0055660 0.007963
## 97.5%
## alpha 1.348000
## beta[1] 0.290800
## beta[2] 0.137500
## beta[3] -0.563400
## beta[4] 0.005717
## beta[5] 0.004004
## beta[6] -0.369300
## beta[7] 0.439000
## delta 0.011760
```

```
d<-abs((t[cens==1]-fit4$mean$Y[cens==1])/fit4$sd$Y[cens==1])
sum(d)
```

```
## [1] 81.96315
```

We see that our model improved a little! For our last model we neglect, SGOT, and we get:

```
##          mean          sd      2.5%      25%      50%      75%
## alpha  1.165389e+00 1.075848e-02 1.146e+00 1.158e+00 1.168e+00 1.175e+00
## beta[1] -1.199254e-01 1.905851e-03 -1.230e-01 -1.224e-01 -1.192e-01 -1.190e-01
## beta[2] 1.576088e-01 1.082697e-03 1.555e-01 1.573e-01 1.575e-01 1.582e-01
## beta[3] -1.389908e-01 7.011369e-04 -1.401e-01 -1.397e-01 -1.391e-01 -1.384e-01
## beta[4] 3.828370e-03 5.075833e-04 3.120e-03 3.353e-03 3.589e-03 4.216e-03
## beta[5] -6.459523e-02 1.587148e-03 -6.605e-02 -6.590e-02 -6.589e-02 -6.254e-02
## beta[6] -1.016850e-01 2.207998e-03 -1.038e-01 -1.037e-01 -1.025e-01 -1.002e-01
## delta 8.596105e-05 7.375110e-06 7.845e-05 8.265e-05 8.279e-05 8.602e-05
## 97.5%
## alpha 1.1760000
## beta[1] -0.1178000
## beta[2] 0.1592000
## beta[3] -0.1380000
## beta[4] 0.0049650
## beta[5] -0.0623400
## beta[6] -0.0981600
## delta 0.0001019
```

```
d<-abs((t[cens==1]-fit5$mean$Y[cens==1])/fit5$sd$Y[cens==1])
sum(d)
```

```
## [1] 95.40789
```

Our model got worse! So we stop at the previous model and keep that as our survival model. Our remaining covariates are Edema, Bilirubin, Albumin, Urin copper, SGOT, prothrombin time and histologic stage. In the after-transplantation lifetime study, those covariates also play the most important role (except SGOT and urine copper)

Appendix

Bernoulli/Beta

A natural conjugate prior for the Bernoulli distribution is the Beta distribution.

$$\begin{aligned}f(x_i|\theta) &= \theta^{x_i}(1-\theta)^{1-x_i} \\L(\mathbf{x}|\theta) &= \theta^{\sum_{i=1}^n x_i}(1-\theta)^{n-\sum_{i=1}^n x_i} \\h(\theta) &= \text{Beta}(a, b)\end{aligned}$$

We proceed by calculating the posterior distribution for θ

$$\begin{aligned}h(\theta|\mathbf{x}) &\propto L(\mathbf{x}|\theta)h(\theta) = \theta^{\sum_{i=1}^n x_i}(1-\theta)^{n-\sum_{i=1}^n x_i} \frac{1}{B(a, b)} \theta^{a-1}(1-\theta)^{b-1} I(0 < \theta < 1) \\&\propto \text{Beta}(\theta|a + \sum_{i=1}^n x_i, b + n - \sum_{i=1}^n x_i)\end{aligned}$$

Poisson/Gamma

If our data X_1, \dots, X_n are iid $\text{Poisson}(\lambda)$ distributed then a $\text{gamma}(\alpha, \beta)$ prior on λ is a conjugate prior. The Likelihood function is:

$$L(\lambda|\mathbf{x}) = \prod_{i=1}^n \frac{e^{-\lambda} \lambda^{\sum_{i=1}^n x_i}}{x_i!} = \frac{e^{-\lambda} \lambda^{\sum_{i=1}^n x_i}}{\prod_{i=1}^n x_i!}$$

Our gamma prior has the expression:

$$h(\lambda) = \frac{\beta^\alpha}{\Gamma(\alpha)} \lambda^{\alpha-1} e^{-\beta\lambda}$$

Using bayes rule we find the following posterior:

$$\begin{aligned}h(\lambda|\mathbf{x}) &\propto h(\lambda)L(\mathbf{x}|\lambda) \propto \lambda^{\sum_{i=1}^n x_i + \alpha - 1} e^{-(n+\beta)\lambda} \\&\propto \text{gamma}(\sum_{i=1}^n x_i + \alpha, n + \beta)\end{aligned}$$

Poisson/Jeffreys prior

The density distribution for poisson is equal to

$$f(n|\lambda) = e^{-\lambda} \frac{\lambda^n}{n!}$$

The jeffreys prior $h(\theta)$ is a non informative prior distrubution for a parameter space and its proportionality is expressed as

$$\begin{aligned}h(\theta) &\propto \sqrt{\det I(\theta)} \\I(\theta) &= -E\left[\frac{\partial^2}{\partial \theta^2} \log f(x|\theta)\right] = \frac{1}{\theta}\end{aligned}$$

And the following jeffreys prior is thus

$$h(\theta) \propto \theta^{-\frac{1}{2}} I_{\theta>0}$$

The posterior is calculated as follows

$$h(\theta|x) \propto f(\mathbf{x}|\theta)h(\theta) \propto e^{-n\theta} \theta^{-\frac{1}{2} + \sum_{i=1}^n x_i}$$

which is in fact a gamma distribution

$$\theta|x \sim \text{Gamma}(\alpha = \frac{1}{2} + \sum_{i=1}^n x_i, \beta = n)$$