STAT 8700 Homework 4

Brian Detweiler

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- 1. Consider data from a Normal population with unknown mean μ and variance σ^2 . A random sample of 100 observations is taken from this population, and the sample mean and variance were calculated to be 50 and 25 respectively.
- (a) If we choose to use a $N-Inv-\chi^2(40,0.64,1,16)$ prior distribution, write down the corresponding posterior distribution.

We are given the following:

$$n = 100$$
 $\overline{y} = 50$
 $s^2 = 25$
 $\mu_0 = 40$
 $\kappa_0 = 25$
 $\nu_0 = 1$
 $\sigma_0^2 = 16$

Now we can use these values to calculate the joint posterior distribution, $N-Inv-\chi^2(\mu_n,\sigma_n^2/\kappa_n;\nu_n,\sigma_n^2)$:

$$\mu_n = \frac{\kappa_0}{\kappa_0 + n} \mu_0 + \frac{n}{\kappa_0 + n} \overline{y}$$

$$= \frac{25}{25 + 100} 40 + \frac{100}{25 + 100} 50 = 48$$

$$\kappa_n = \kappa_0 + n$$

$$= 25 + 100 = 125$$

$$\nu_n = \nu_0 + n$$

$$= 1 + 100 = 101$$

$$\nu_n \sigma_n^2 = \nu_0 \sigma_0^2 + (n - 1)s^2 + \frac{\kappa_0 n}{\kappa_0 + n} (\overline{y}\mu_0)^2$$

$$= 1(16) + (100 - 1)25 + \frac{25(100)}{25 + 100} (50 - 40)^2 = 4491$$

$$\sigma_n^2 \approx 44.4653465347$$

And thus our joint posterior distributin is $N - Inv - \chi^2(48, 0.355722772278; 101, 44.4653465347)$.

(b) Either analytically or via simulation, construct 95% credible intervals for σ^2 and μ .

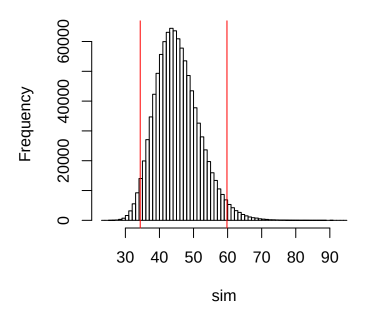
To simulate this, we first draw σ^2 from its marginal posterior distribution, $\sigma^2 | y \sim Inv - \chi^2(\nu_n, \sigma_n^2)$

```
library(geoR)
set.seed(124)

x <- seq(0, 100, by = 0.001)
nu_n <- 101
sigma_n_2 <- 44.4653465347

sim <- rinvchisq(n = 1000000, df = nu_n, scale = sigma_n_2)
hist(sim, breaks = 90, main = 'Distribution of variance')
lower <- sort(sim)[25000]
upper <- sort(sim)[975000]
abline(v=lower, col='red')
abline(v=upper, col='red')</pre>
```

Distribution of variance



A 95% credible interval for σ^2 is (34.3564188, 59.8085225).

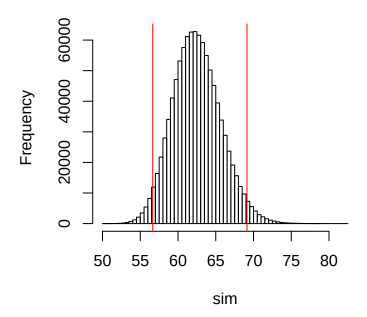
Then we sample from $N\left(\frac{\frac{\kappa_0}{\sigma^2}\mu_0 + \frac{n}{\sigma^2}\overline{y}}{\frac{\kappa_0}{\sigma^2} + \frac{n}{\sigma^2}}, \frac{1}{\frac{\kappa_0}{\sigma^2} + \frac{n}{\sigma^2}}\right)$ using the previous values for σ^2 .

```
sigma_2 <- sim
kappa_0 <- 25
mu_0 <- 40
```

```
n <- 100
y_bar <- 50
mu_n <- ((kappa_0 / sigma_2) * mu_0) + ((n / sigma_2) * y_bar) / ((kappa_0 / sigma_2) + (n / sigma_2))
sigma_2_kappa_n <- 1 / ((kappa_0 / sigma_2) + (n / sigma_2))
sim <- rnorm(n = 1000000, mu_n, sigma_2_kappa_n)
hist(sim, breaks = 90, main = 'Distribution of mean')

lower <- sort(sim)[25000]
upper <- sort(sim)[975000]
abline(v=lower, col='red')
abline(v=upper, col='red')</pre>
```

Distribution of mean



A 95% credible interval for μ is (56.6518857, 69.1340067).

2. Two random variables are said to have a bivariate normal distribution with parameters, $\mu_U, \mu_V, \sigma_U^2, \sigma_V^2$, and ρ if they have the following density function:

$$f(u,v) = \frac{1}{2\pi\sigma_U \sigma_V \sqrt{1-\rho^2}} e^{-\frac{1}{2(1-\rho^2)} \left[\frac{(u-\mu_U)^2}{\sigma_U^2} + \frac{(v-\mu_V)^2}{\sigma_V^2} - \frac{2\rho(u-\mu_U)(v-\mu_V)}{\sigma_U \sigma_V} \right]}$$

where μ_U and σ_U^2 are the mean and variance of U, μ_V and σ_V^2 are the mean and variance of V, and ρ is the correlation between U and V.

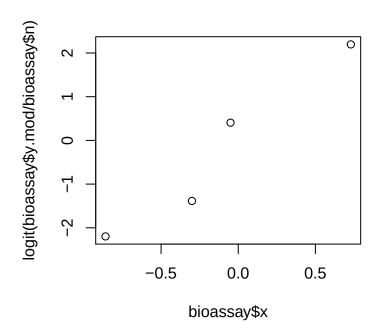
Replace the uniform prior on α and β in the analysis of the bioassay by a bivariate normal prior with $\alpha \sim Normal(0,4), \beta \sim Normal(10,100)$, and $corr(\alpha,\beta)=0.5$. Repeat all the computations and plots discussed in section 3.7 and in class.

The book does a quick maximum likelihood of α and β .

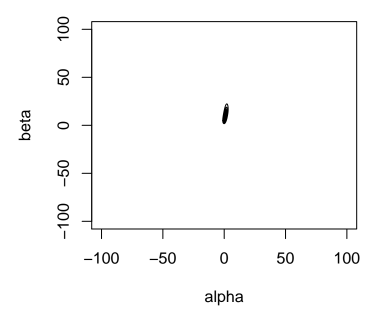
```
x \leftarrow c(-0.86, -0.3, -0.05, 0.73)
n \leftarrow c(5, 5, 5, 5)
y \leftarrow c(0, 1, 3, 5)
y.mod \leftarrow c(0.5, 1, 3, 4.5)
data <- cbind(y, n - y)
fit <- glm(data ~ x, family = binomial)</pre>
coef(fit)
## (Intercept)
     0.8465802 7.7488172
# This is the approximate covariance matrix:
summary(fit)$cov.unscaled
                 (Intercept)
## (Intercept) 1.038535 3.545987
                    3.545987 23.743865
## x
a <- coef(fit)[[1]]
b <- coef(fit)[[2]]</pre>
```

The maximum likelihood estimate for α is 0.8465802 and for β is 7.7488172.

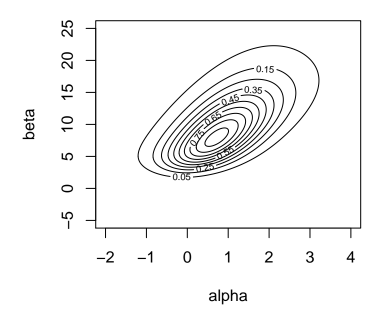
```
bivariate_normal <- function(u, v) {</pre>
  mu_U <- 0
  mu_V <- 10
  sigma_2_U <- 4
  sigma_2_V <- 100
  sigma_U <- sqrt(sigma_2_U)</pre>
  sigma_V <- sqrt(sigma_2_V)</pre>
  rho <- 0.5
  first <- 1 / (2 * pi * sigma_U * sigma_V * sqrt(1 - rho^2))
  second \leftarrow \exp(-(1 / (2 * (1 - rho^2))))
                 * ( ((u - mu_U)^2 / sigma_2_U)
                     + ((v - mu_V)^2 / sigma_2_V)
                     - (2 * rho * (u - mu_U) * (v - mu_V)) / (sqrt(sigma_2_U) * sqrt(sigma_2_V))))
  rval <- first * second</pre>
  return(rval)
bioassay <- data.frame(cbind(x, n, y, y.mod))</pre>
logit <- function(x) {</pre>
 log(x / (1-x))
inv_logit <- function(x){</pre>
exp(x) / (1 + exp(x))
plot(bioassay$x, logit(bioassay$y.mod / bioassay$n))
```



```
posterior <- function(a, b) {</pre>
  temp <-1
    x <- bioassay$x
    y <- bioassay$y
    n <- bioassay$n
    for (i in 1: length(x)) {
      temp <- (temp * (inv_logit(a + (b * x[i]))^y[i])
                   * ((1 - inv_logit(a + (b * x[i])))^(n[i] - y[i])))
  }
  bivariate_normal(a, b) * temp
}
posterior_contour <- function(alpha_min,</pre>
                                alpha_max,
                                grid_size_alpha,
                                beta_min,
                                beta_max,
                                grid_size_beta,
                                drawlabels = TRUE) {
  # Generate a list of alpha values
  alpha <- seq(alpha_min, alpha_max, length = grid_size_alpha)</pre>
  # Generate a list of beta values
  beta <- seq(beta_min, beta_max, length = grid_size_beta)</pre>
  # Evaluate the posterior density and all possible combinations of alpha and beta values.
  post.dens <- outer(alpha, beta, 'posterior')</pre>
```

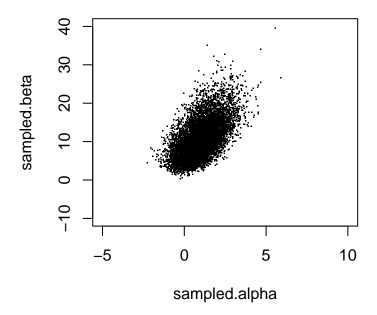


```
posterior_contour(-2, 4, 2001, -5, 25, 2001)
```



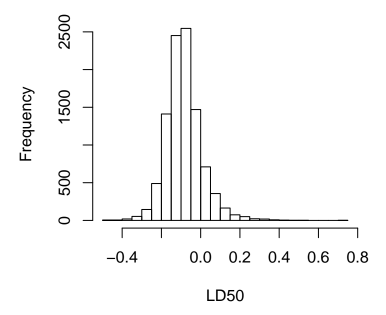
```
##Sampling from the joint posterior
posterior_sample <- function(alpha_min,</pre>
                               alpha_max,
                               grid_size_alpha,
                              beta_min,
                              beta_max,
                               grid_size_beta,
                              nsim) {
  # Generate a list of alpha values
    alpha <- seq(alpha_min, alpha_max, length = grid_size_alpha)</pre>
  # Generate a list of beta values
    beta <- seq(beta_min, beta_max,length = grid_size_beta)</pre>
  {\it \# Evaluate the posterior density and all possible combinations of alpha and beta values.}
    post.dens <- outer(alpha, beta, 'posterior')</pre>
  # Rescale the posterior so values are now relative to height of posterior mode.
    scaled.dens <- post.dens / max(post.dens)</pre>
  # Makes sure our discrete probability density sums to 1
    normalized.dens = scaled.dens / sum(scaled.dens)
    marg.alpha <- rowSums(normalized.dens)</pre>
    #Sample 'nsim' values of alpha and beta
```

```
# Define a vector to hold our alpha values
    sampled.alpha <- NULL</pre>
  #Define a vector to hold out beta values
    sampled.beta <- NULL</pre>
    alpha_width <- (alpha_max - alpha_min) / (grid_size_alpha - 1)</pre>
    beta_width <- (beta_max-beta_min) / (grid_size_beta - 1)</pre>
  for (i in 1:nsim) {
    #Generate a value of alpha from the marginal of alpha
        temp.alpha <- sample(alpha, 1, marg.alpha, replace = TRUE)</pre>
    # Translate the obtained alpha to the appropriate row of our grid
        j <- ((temp.alpha - alpha_min) / alpha_width) + 1</pre>
    # Calculate the condition distribution of beta, conditional on the obtained value of alpha
        conditional.beta <- normalized.dens[j, ] / sum(normalized.dens[j, ])</pre>
    # Obtain a beta from the conditional distribution of beta
        temp.beta <- sample(beta, 1, conditional.beta, replace = TRUE)</pre>
    # Add some noise to try to undiscretize the posterior
        temp.alpha <- temp.alpha + runif(1, -alpha_width / 2, alpha_width / 2)
        temp.beta <- temp.beta + runif(1, -beta_width / 2, beta_width / 2)</pre>
    # Add the generated alpha to the list of previously generated alpha
        sampled.alpha <- c(sampled.alpha, temp.alpha)</pre>
        sampled.beta<-c(sampled.beta, temp.beta)</pre>
  }
    data.frame(sampled.alpha, sampled.beta)
}
sims <- posterior_sample(-5, 10, 2001, -10, 40, 2001, 10000)
plot(sims, xlim = c(-5, 10), ylim = c(-10, 40), pch='.')
```

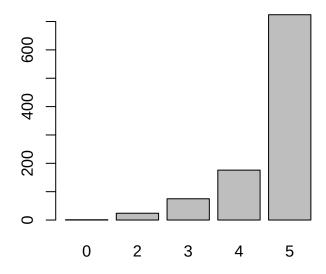


Posterior Distribution of LD50 # LD50 is the dose at which the death rate is 50%, only makes sense fo
LD50 <- (-sims\$sampled.alpha[sims\$sampled.beta > 0] / sims\$sampled.beta[sims\$sampled.beta > 0])
hist(LD50, br = 35)

Histogram of LD50



```
# Posterior Predictive distribution for a dose level x = 0.25, n = 5
# y~Bin(5, theta)
# logit(theta) = alpha + beta*x
theta <- inv_logit(sims$sampled.alpha + (sims$sampled.beta * 0.25))
y <- rbinom(1000, 5, theta)
barplot(table(y))</pre>
```

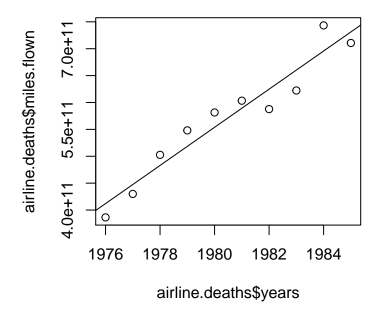


3. Consider the airline fatalities data discussed in the previous exercise. Let us suppose that we now assume that the number of fatal accidents in year t follows a Poisson distribution with mean $\alpha + \beta t$.

```
years <- c(1976:1985)
fatal.accidents <- c(24, 25, 31, 31, 22, 21, 26, 20, 16, 22)
passenger.deaths <- c(734, 516, 754, 877, 814, 362, 764, 809, 223, 1066)
death.rate <- c(0.19, 0.12, 0.15, 0.16, 0.14, 0.06, 0.13, 0.13, 0.03, 0.15)
airline.deaths <- as.data.frame(cbind(years, fatal.accidents, passenger.deaths, death.rate))
miles.flown <- (airline.deaths$passenger.deaths / airline.deaths$death.rate) * 1000000000
airline.deaths$miles.flown <- miles.flown</pre>
```

```
years fatal.accidents passenger.deaths death.rate miles.flown
## 1
       1976
                          24
                                           734
                                                     0.19 386315789474
                          25
## 2
       1977
                                           516
                                                     0.12 430000000000
## 3
       1978
                          31
                                           754
                                                     0.15 502666666667
## 4
       1979
                          31
                                           877
                                                     0.16 548125000000
## 5
       1980
                          22
                                                     0.14 581428571429
                                           814
## 6
       1981
                          21
                                           362
                                                     0.06 603333333333
## 7
       1982
                          26
                                           764
                                                     0.13 587692307692
## 8
       1983
                          20
                                           809
                                                     0.13 622307692308
## 9
       1984
                          16
                                           223
                                                     0.03 743333333333
## 10 1985
                          22
                                          1066
                                                     0.15 710666666667
```

```
plot(airline.deaths$miles.flown ~ airline.deaths$years)
abline(lm(airline.deaths$miles.flown ~ airline.deaths$years))
```



(a) If we let y_t represent the number of fatal accidents in year t, write down $p(y_t|\alpha,\beta)$ the likelihood for year t in terms of the parameters α , and β .

$$t \sim Poisson(\alpha + \beta t)$$
$$p(y_t | \alpha, \beta) = \frac{(\alpha + \beta t)^{y_t}}{y_t!} e^{-(\alpha + \beta t)}$$

(b) If we assume uniform priors on α and β , write the posterior density for (α, β) .

With a uniform prior, we have $p(\alpha, \beta) \propto 1$. So the posterior is proportional to the likelihood.

We scale the years and let t=1 represent 1975, let t=2 represent 1976, and so on.

$$\begin{split} p(\alpha,\beta|y) &\propto p(\alpha,\beta)p(y|\alpha,\beta) \\ &\propto p(\alpha,\beta) \prod_{i=1}^{10} p(y_i|\alpha,\beta) \\ &\propto \prod_{i=1}^{10} p(y_i|\alpha,\beta) \\ &= \prod_{i=1}^{10} \frac{(\alpha+\beta t)^{y_t}}{y_t!} e^{-(\alpha+\beta t)} \end{split}$$

(c) Following the same idea as the boassay example (and the previous question) create a grid of possible α and β values on which to evaluate the joint posterior and plot the contours. Start with large ranges for α and β and refine based on the countour plot. Include all your iterations in your answer, not just you final grid and contour plot.

First, we create our posterior probability density,

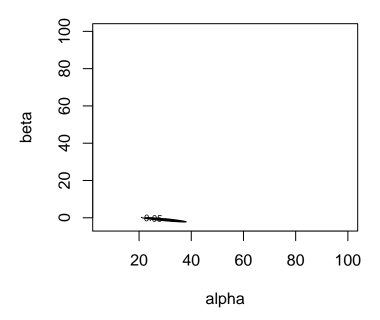
```
posterior <- function(a, b) {
   rval <- 1
   y <- airline.deaths$fatal.accidents
     for (t in 1: length(y)) {
        rval <- (rval * dpois(y[t], (a + b*t)))
   }

   return(rval)
}</pre>
```

```
posterior_contour <- function(alpha_min,</pre>
                                alpha_max,
                                grid_size_alpha,
                                beta_min,
                                beta_max,
                                grid_size_beta,
                                drawlabels = TRUE) {
  # Generate a list of alpha values
  alpha <- seq(alpha_min, alpha_max, length = grid_size_alpha)</pre>
  # Generate a list of beta values
  beta <- seq(beta_min, beta_max, length = grid_size_beta)</pre>
  # Evaluate the posterior density and all possible combinations of alpha and beta values.
  post.dens <- outer(alpha, beta, 'posterior')</pre>
  # Set NANs to O
  post.dens[is.nan(post.dens)] <- 0</pre>
  post.dens[is.na(post.dens)] <- 0</pre>
  # Rescale the posterior so values are now relative to height of posterior mode.
  scaled.dens <- post.dens / max(post.dens)</pre>
  # Draw contour plot.
  contour(alpha,
          beta,
          scaled.dens,
          levels = c(0.05, 0.15, 0.25, 0.35, 0.45, 0.55, 0.65, 0.75, 0.85, 0.95),
          xlab="alpha",
          ylab="beta",
          drawlabels = drawlabels)
```

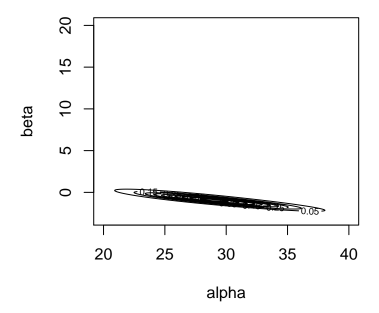
Then plot the contours

posterior_contour(6, 100, 2001, -3, 100, 2001)



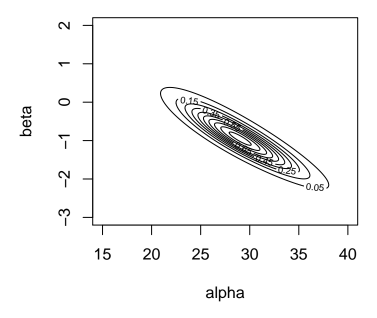
We can zoom in further,

posterior_contour(20, 40, 2001, -3, 20, 2001)



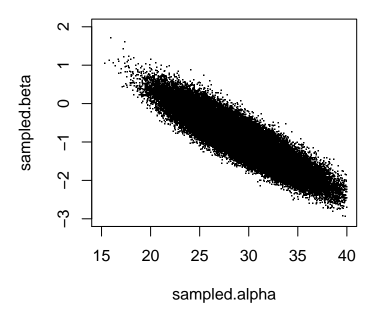
And now, we know just about where the ranges are, so we'll zoom in on those,

posterior_contour(15, 40, 2001, -3, 2, 2001)



(d) Simulate 100,000 values of α and β from the joint posterior and plot the histogram of the posterior density of the expected number of fatal accidents in 1986, $\alpha + 1986\beta$.

```
##Sampling from the joint posterior
posterior_sample <- function(alpha_min,</pre>
                               alpha max,
                               grid_size_alpha,
                              beta_min,
                              beta_max,
                               grid_size_beta,
                              nsim) {
  # Generate a list of alpha values
  alpha <- seq(alpha_min, alpha_max, length = grid_size_alpha)</pre>
  # Generate a list of beta values
    beta <- seq(beta_min, beta_max,length = grid_size_beta)</pre>
  # Evaluate the posterior density and all possible combinations of alpha and beta values.
    post.dens <- outer(alpha, beta, 'posterior')</pre>
  # Set NANs to O
  post.dens[is.nan(post.dens)] <- 0</pre>
  # Rescale the posterior so values are now relative to height of posterior mode.
  scaled.dens <- post.dens / max(post.dens)</pre>
  # Makes sure our discrete probability density sums to 1
    normalized.dens = scaled.dens / sum(scaled.dens)
    marg.alpha<-rowSums(normalized.dens)</pre>
    #Sample 'nsim' values of alpha and beta
  # Define a vector to hold our alpha values
    sampled.alpha <- NULL
  #Define a vector to hold out beta values
    sampled.beta <- NULL</pre>
    alpha_width <- (alpha_max - alpha_min) / (grid_size_alpha - 1)</pre>
    beta_width <- (beta_max-beta_min) / (grid_size_beta - 1)</pre>
  for (i in 1:nsim) {
    #Generate a value of alpha from the marginal of alpha
        temp.alpha <- sample(alpha, 1, marg.alpha, replace = TRUE)</pre>
    # Translate the obtained alpha to the appropriate row of our grid
        j <- ((temp.alpha - alpha_min) / alpha_width) + 1</pre>
```



- (e) Use your simulated values of α and β to simulate the number of fatal accidents in 1986. Use your simulations to construct a 95% predictive (credible) interval.
- (f) Return to your simulated values of β , calculate (well, estimate) $P(\beta < 0)$, that is, the probability that the number of fatal accidents per year is decreasing.

```
prob.decreasing <- length(sims$sampled.beta[sims$sampled.beta < 0]) / length(sims$sampled.beta)</pre>
```

There is a 95.829% chance the number of fatal accidents is decreasing.