Bayesian data analysis demo 3.6

Aki Vehtari, Markus Paasiniemi

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Binomial regression and grid sampling for Bioassay data (BDA3 p. 74-)

ggplot2, and gridExtra are used for plotting, tidyr for manipulating data frames

```
library(ggplot2)
theme_set(theme_minimal())
library(gridExtra)
library(tidyr)
library(dplyr)
library(purrr)
```

The Dataset

Call:

```
df1 <- data.frame(</pre>
 x = c(-0.86, -0.30, -0.05, 0.73),
 n = c(5, 5, 5, 5),
 y = c(0, 1, 3, 5)
```

${\bf Trying\ out\ } logistic\ regression$

We employ Maximum Likelihood Estimation in order to find the α and β .

glm(formula = theta ~ x, family = binomial, data = df1)

```
theta \leftarrow df1$y / df1$n
theta
## [1] 0.0 0.2 0.6 1.0
df1$theta <- theta
df1
##
         x n y theta
## 1 -0.86 5 0 0.0
## 2 -0.30 5 1
                 0.2
## 3 -0.05 5 3
                 0.6
## 4 0.73 5 5
                 1.0
log_fits <- glm( theta ~ x, family = binomial, data=df1 )</pre>
## Warning in eval(family$initialize): non-integer #successes in a binomial glm!
summary(log_fits)
```

```
## Deviance Residuals:
##
                     2
          1
                                3
## -0.07708
              0.03637 -0.02625
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
##
                  0.8466
                              2.2787
                                       0.372
                                                 0.710
## (Intercept)
                  7.7488
                             10.8957
                                       0.711
                                                 0.477
## x
##
  (Dispersion parameter for binomial family taken to be 1)
##
##
##
       Null deviance: 3.158282 on 3 degrees of freedom
## Residual deviance: 0.010948 on 2 degrees of freedom
## AIC: 5.3992
##
## Number of Fisher Scoring iterations: 7
As we can see that the maximum likelihood estimate of (\hat{\alpha}, \hat{\beta}) is (0.8, 7.7).
Plot data
ggplot(df1, aes(x=x, y=y)) +
    geom_point(size=2, color='red') +
    scale_x_continuous(breaks = df1$x, minor_breaks=NULL, limits = c(-1.5, 1.5)) +
    scale_y_continuous(breaks = 0:5, minor_breaks=NULL) +
    labs(title = 'Bioassay', x = 'Dose (log g/ml)', y = 'Number of deaths') +
    theme(panel.grid.major = element_blank())
     Bioassay
   5
   4
Number of deaths
   1
   0
                        -0.86
                                       -0.30 -0.05
                                                                    0.73
                                          Dose (log g/ml)
```

Compute the posterior density in grid

- usually should be computed in logarithms!
- with alternative prior, check that range and spacing of A and B are sensible

```
A = seq(-4, 8, length.out = 50)
B = seq(-10, 40, length.out = 50)
```

Make vectors that contain all pairwise combinations of A and B

```
cA <- rep(A, each = length(B))
cB <- rep(B, length(A))</pre>
```

Make a helper function to calculate the log likelihood given a dataframe with x, y, and n and evaluation points a and b. For the likelihood see BDA3 p. 75 log1p(x) computes log(x+1) in numerically more stable way.

```
logl <- function(df, a, b)
df['y']*(a + b*df['x']) - df['n']*log1p(exp(a + b*df['x']))</pre>
```

Calculate likelihoods: apply logl function for each observation ie. each row of data frame of x, n and y

```
p <- apply(df1, 1, logl, cA, cB) %>%
  # sum the log likelihoods of observations
# and exponentiate to get the joint likelihood
rowSums() %>% exp()
```

Sample from the grid (with replacement)

Add random jitter, see BDA3 p. 76

```
samp_A <- samp_A + runif(nsamp, (A[1] - A[2])/2, (A[2] - A[1])/2)
samp_B <- samp_B + runif(nsamp, (B[1] - B[2])/2, (B[2] - B[1])/2)</pre>
```

Create data frame

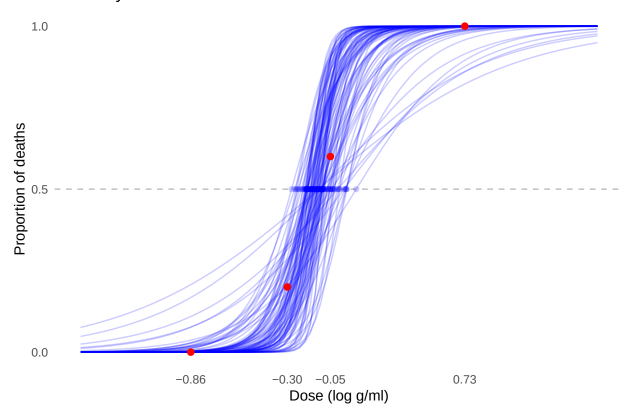
```
samps <- data_frame(ind = 1:nsamp, alpha = samp_A, beta = samp_B) %>%
mutate(ld50 = - alpha/beta)
```

```
## Warning: `data_frame()` was deprecated in tibble 1.1.0.
## Please use `tibble()` instead.
```

Plot draws of logistic curves

```
ppost + geom_hline(yintercept = 0.5, linetype = 'dashed', color = 'gray') +
   geom_point(data=samps[1:100,], aes(x=ld50, y=0.5), color='blue', alpha=0.2)
```

Bioassay



Create a plot of the posterior density

```
# limits for the plots
xl <- c(-2, 8)
yl <- c(-2, 40)
pos <- ggplot(data = data.frame(cA ,cB, p), aes(cA, cB)) +
    geom_raster(aes(fill = p, alpha = p), interpolate = T) +
    geom_contour(aes(z = p), colour = 'black', size = 0.2) +
    coord_cartesian(xlim = xl, ylim = yl) +
    labs(title = 'Posterior density evaluated in grid', x = 'alpha', y = 'beta') +
    scale_fill_gradient(low = 'yellow', high = 'red', guide = F) +
    scale_alpha(range = c(0, 1), guide = F)</pre>
```

Plot of the samples

```
sam <- ggplot(data = samps) +
  geom_point(aes(alpha, beta), color = 'blue') +
  coord_cartesian(xlim = xl, ylim = yl) +
  labs(title = 'Posterior draws', x = 'alpha', y = 'beta')</pre>
```

Combine the plots

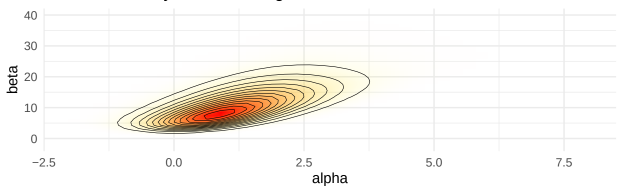
```
grid.arrange(pos, sam, nrow=2)
```

Warning: It is deprecated to specify `guide = FALSE` to remove a guide. Please

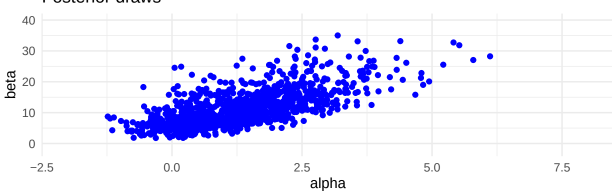
```
## use `guide = "none"` instead.
```

Warning: It is deprecated to specify `guide = FALSE` to remove a guide. Please
use `guide = "none"` instead.

Posterior density evaluated in grid



Posterior draws



Plot of the histogram of LD50 $\,$

