# Bayesian Logistic Regression

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Clear environment, graphics and console

```
# Clear environment ####
rm(list = ls())

# Clear plots
graphics.off() # Clears plots, closes all graphics devices

# Clear console
cat("\014") # ctrl+L
```

Install required packages

### 1. Introduction

This is a Bayesian logistic regression related to red variants of the Portuguese "Vinho Verde" wine. There are 11 input variables (covariates) based on physiochemical tests. The output variable is quality and is a score between 0 and 10 which is based on sensory data.

# 2. Import data.

The dataset is described in the publication by Cortez, P., Cerdeira, A., Almeida, F., Matos, T., & Reis, J. (2009). Modeling wine preferences by data mining from physicochemical properties..

```
wine <- read.csv("winequality-red.csv")
head(wine)</pre>
```

```
##
     fixed.acidity volatile.acidity citric.acid residual.sugar chlorides
## 1
               7.4
                                0.70
                                             0.00
                                                              1.9
                                                                       0.076
## 2
               7.8
                                 0.88
                                             0.00
                                                              2.6
                                                                       0.098
## 3
               7.8
                                0.76
                                             0.04
                                                              2.3
                                                                       0.092
## 4
              11.2
                                0.28
                                                              1.9
                                                                       0.075
                                             0.56
## 5
               7.4
                                0.70
                                             0.00
                                                              1.9
                                                                       0.076
## 6
               7.4
                                0.66
                                                                       0.075
                                             0.00
                                                              1.8
     free.sulfur.dioxide total.sulfur.dioxide density
                                                           pH sulphates alcohol
## 1
                       11
                                             34 0.9978 3.51
                                                                    0.56
                                                                             9.4
## 2
                       25
                                                0.9968 3.20
                                                                    0.68
                                                                             9.8
## 3
                       15
                                             54 0.9970 3.26
                                                                    0.65
                                                                             9.8
```

```
## 4
                       17
                                              60 0.9980 3.16
                                                                    0.58
                                                                              9.8
## 5
                       11
                                             34 0.9978 3.51
                                                                    0.56
                                                                              9.4
## 6
                                                                    0.56
                       13
                                              40 0.9978 3.51
                                                                              9.4
##
     quality
## 1
           5
## 2
           5
## 3
           5
           6
## 4
## 5
           5
## 6
           5
```

### Check for missing values

There are no NA values in the dataset.

```
any(is.na(wine))
```

## [1] FALSE

# 3. Classify "good" wines

We want to implement a logistic regression, therefore we want a response variable which assumes values either 0 or 1. Suppose we consider "good" a wine with quality above 6.5 (included).

We add a column called good that is 1 if the quality is greater than 6.5 and 0 if quality is less than 6.5.

We also add a column of strings called drinkable, relating 1 and 0 to "good" and "bad" as appropriate.

```
wine %<>%
mutate(good = ifelse(quality > 6, 1, 0)) %>%
mutate(drinkable = ifelse(quality > 6, "good", "bad")) %>%
mutate(quality = factor(quality, levels=1:10))
```

Remove spaces from names to make references easier.

The structure of the dataset, str(wine), will confirm that there are no NA values, the feature names, the feature type (num, Factor, chr) and the first 10 values.

```
str(wine)
```

```
## 'data.frame': 1599 obs. of 14 variables:
## $ fixed_acidity : num 7.4 7.8 7.8 11.2 7.4 7.4 7.9 7.3 7.8 7.5 ...
## $ volatile_acidity : num 0.7 0.88 0.76 0.28 0.7 0.66 0.6 0.65 0.58 0.5 ...
## $ citric_acid : num 0 0 0.04 0.56 0 0 0.06 0 0.02 0.36 ...
## $ residual sugar : num 1.9 2.6 2.3 1.9 1.9 1.8 1.6 1.2 2 6.1 ...
```

```
$ chlorides
                             0.076 0.098 0.092 0.075 0.076 0.075 0.069 0.065 0.073 0.071 ...
                       : num
   ##
  $ total sulfur dioxide: num
                             34 67 54 60 34 40 59 21 18 102 ...
  $ density
##
                             0.998 0.997 0.997 0.998 0.998 ...
                       : num
##
   #q#
                       : num
                             3.51 3.2 3.26 3.16 3.51 3.51 3.3 3.39 3.36 3.35 ...
                       : num 0.56 0.68 0.65 0.58 0.56 0.56 0.46 0.47 0.57 0.8 ...
##
  $ sulphates
  $ alcohol
                             9.4 9.8 9.8 9.8 9.4 9.4 9.4 10 9.5 10.5 ...
                       : num
                       : Factor w/ 10 levels "1", "2", "3", "4", ...: 5 5 5 6 5 5 7 7 5 ...
## $ quality
                       : num
## $ good
                             0 0 0 0 0 0 0 1 1 0 ...
                             "bad" "bad" "bad" "bad" ...
   $ drinkable
                       : chr
```

# 4. Frequentist Logistic Regression

The logistic model is computed using the generalised linear model (glm) with good being a function of all other covariates except quality and drinkable.

```
fit <- glm(
  good ~ . - quality - drinkable,
  data = wine,
  family = binomial(link="logit")
)</pre>
```

## Significant coefficients

The significant coefficients are labelled in the last column of the summary() function. Most significant coefficients are:

```
• sulphates (\Pr(\alpha > 6.924 = 4.39\text{e-}12)),
• alcohol(\Pr(\alpha > 5.724 = 1.04\text{e-}08)),
• volatile_acidity (\Pr(\alpha > |-3.291| = 9.99\text{e-}4)), and
• total_sulfur_dioxide (\Pr(\alpha > |-3.378| = 7.31\text{e-}4))
```

```
fit %>% summary()
```

```
##
## Call:
  glm(formula = good ~ . - quality - drinkable, family = binomial(link = "logit"),
##
       data = wine)
##
## Deviance Residuals:
##
      Min
                1Q
                     Median
                                   3Q
                                          Max
## -2.9878 -0.4351 -0.2207 -0.1222
                                        2.9869
##
## Coefficients:
##
                         Estimate Std. Error z value Pr(>|z|)
## (Intercept)
                        2.428e+02 1.081e+02
                                               2.247 0.024660 *
## fixed_acidity
                        2.750e-01 1.253e-01
                                              2.195 0.028183 *
## volatile_acidity
                       -2.581e+00 7.843e-01 -3.291 0.000999 ***
## citric_acid
                        5.678e-01 8.385e-01
                                              0.677 0.498313
## residual_sugar
                        2.395e-01 7.373e-02
                                              3.248 0.001163 **
```

```
## chlorides
                       -8.816e+00 3.365e+00 -2.620 0.008788 **
## free_sulfur_dioxide 1.082e-02 1.223e-02 0.884 0.376469
## total sulfur dioxide -1.653e-02 4.894e-03 -3.378 0.000731 ***
## density
                      -2.578e+02 1.104e+02 -2.335 0.019536 *
## pH
                       2.242e-01 9.984e-01 0.225 0.822327
                       3.750e+00 5.416e-01 6.924 4.39e-12 ***
## sulphates
## alcohol
                       7.533e-01 1.316e-01 5.724 1.04e-08 ***
## ---
## Signif. codes: 0 '***' 0.001 '**' 0.05 '.' 0.1 ' ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
##
      Null deviance: 1269.92 on 1598 degrees of freedom
## Residual deviance: 870.86 on 1587 degrees of freedom
## AIC: 894.86
##
## Number of Fisher Scoring iterations: 6
```

# 5. Impact on outcome by varying total\_sulfur\_dioxide

By fixing all coefficients and varying one, we can see the impact of one particular coefficient has on the probabilistic outcome of the logistic regression.

As a comparison I have fixed all significant coefficients in turn (total\_sulfur\_dioxide, volatile\_acidity, sulphates and alcohol)

First, we save coefficients to their own variable.

```
b0 <- fit$coefficients[1]  # Intercept = 242.76251933
b1 <- fit$coefficients[2]  # fixed_acidity = 0.27495289
b2 <- fit$coefficients[3]  # volatile_acidity = -2.58100211
b3 <- fit$coefficients[4]  # citric_acid = 0.56779433
b4 <- fit$coefficients[5]  # residual_sugar = 0.23946420
b5 <- fit$coefficients[6]  # chlorides = -8.81636544
b6 <- fit$coefficients[7]  # free_sulfur_dioxide = 0.01082060
b7 <- fit$coefficients[8]  # total_sulfur_dioxide = -0.01653061
b8 <- fit$coefficients[9]  # density = -257.79757874
b9 <- fit$coefficients[10]  # pH = 0.22418522
b10 <- fit$coefficients[11]  # sulphates = 3.74987886
b11 <- fit$coefficients[12]  # alcohol = 0.75333905</pre>
```

Then save each individual mean values

```
fixed_acidity_mean <- mean(wine$fixed_acidity)
volatile_acidity_mean <- mean(wine$volatile_acidity)
citric_acid_mean <- mean(wine$citric_acid)
residual_sugar_mean <- mean(wine$residual_sugar)
chlorides_mean <- mean(wine$chlorides)
free_sulfur_dioxide_mean <- mean(wine$free_sulfur_dioxide)
total_sulfur_dioxide_mean <- mean(wine$total_sulfur_dioxide)
density_mean <- mean(wine$density)
pH_mean <- mean(wine$pH)
sulphates_mean <- mean(wine$sulphates)
alcohol_mean <- mean(wine$alcohol)</pre>
```

Compute the range of each significant coefficient, total\_sulfur\_dioxide, sulphates, alcohol, and volatile\_acidity

```
total_sulfur_dioxide_range <- seq(from=min(wine$total_sulfur_dioxide), to=max(wine$total_sulfur_dioxide) sulphates_range <- seq(from=min(wine$sulphates), to=max(wine$sulphates), by=0.002) alcohol_range <- seq(from=min(wine$alcohol), to=max(wine$alcohol), by=0.05) volatile_acidity_range <- seq(from=min(wine$volatile_acidity), to=max(wine$volatile_acidity), by=0.002)
```

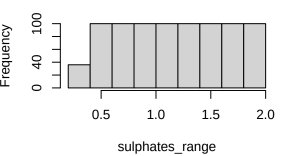
Plot of the range histogram of each of the four most significant covariates.

```
par(mfrow=c(2,2))
hist(total_sulfur_dioxide_range)
hist(sulphates_range)
hist(alcohol_range)
hist(volatile_acidity_range)
```

### Histogram of total\_sulfur\_dioxide\_rang

# O 50 100 200 300 total sulfur dioxide range

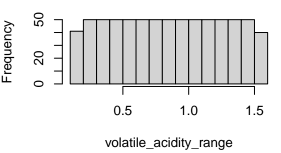
## Histogram of sulphates\_range



# Histogram of alcohol\_range



# Histogram of volatile\_acidity\_range



Calculate probabilities for each significant coefficient

```
total_sulfur_dioxide_GOOD <- b0 + b1*fixed_acidity_mean + b2*volatile_acidity_mean + b3*citric_acid_mean + b4*residual_sugar_mean + b5*chlorides_mean + b6*free_sulfur_dioxide_mean + b7*total_sulfur_dioxide_range + b8*density_mean + b9*pH_mean + b10*sulphates_mean + b11*alcohol_mean sulphates_GOOD <- b0 + b1*fixed_acidity_mean + b2*volatile_acidity_mean + b3*citric_acid_mean + b4*residual_sugar_mean + b5*chlorides_mean + b6*free_sulfur_dioxide_mean + b7*total_sulfur_dioxide_mean + b8*density_mean + b9*pH_mean + b10*sulphates_range + b11*alcohol_mean
```

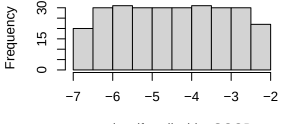
```
alcohol_GOOD <- b0 + b1*fixed_acidity_mean + b2*volatile_acidity_mean +</pre>
  b3*citric_acid_mean + b4*residual_sugar_mean + b5*chlorides_mean + b6*free_sulfur_dioxide_mean +
  b7*total_sulfur_dioxide_mean + b8*density_mean + b9*pH_mean + b10*sulphates_mean + b11*alcohol_range
volatile_acidity_GOOD <- b0 + b1*fixed_acidity_mean + b2*volatile_acidity_range +
  b3*citric_acid_mean + b4*residual_sugar_mean + b5*chlorides_mean + b6*free_sulfur_dioxide_mean +
  b7*total_sulfur_dioxide_mean + b8*density_mean + b9*pH_mean + b10*sulphates_mean + b11*alcohol_mean
```

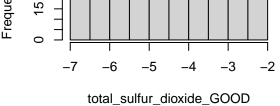
Histograms of what????

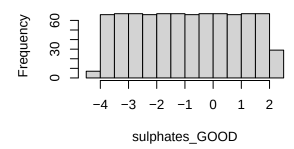
```
par(mfrow=c(2,2))
hist(total_sulfur_dioxide_GOOD)
hist(sulphates_GOOD)
hist(alcohol GOOD)
hist(volatile_acidity_GOOD)
```

# Histogram of total\_sulfur\_dioxide\_GOC

# Histogram of sulphates\_GOOD

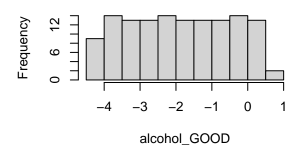


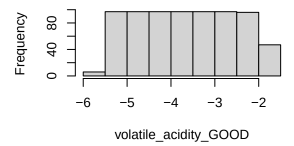




# Histogram of alcohol\_GOOD

# Histogram of volatile\_acidity\_GOOD





Calculate logistic probabilities for each

```
total_sulfur_dioxide_probs <- exp(total_sulfur_dioxide_GOOD)/(1 + exp(total_sulfur_dioxide_GOOD))
sulphates_probs <- exp(sulphates_GOOD) / (1 + exp(sulphates_GOOD))</pre>
alcohol_probs <- exp(alcohol_GOOD) / (1 + exp(alcohol_GOOD))</pre>
volatile_acidity_probs <- exp(volatile_acidity_GOOD) / (1 + exp(volatile_acidity_GOOD))
```

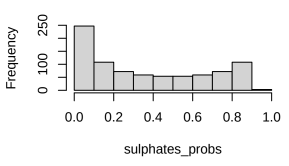
Plot logistic probabilities

par(mfrow=c(2,2))
hist(total\_sulfur\_dioxide\_probs)
hist(sulphates\_probs)
hist(alcohol\_probs)
hist(volatile\_acidity\_probs)

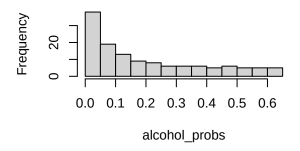
# Histogram of total\_sulfur\_dioxide\_prol

# 0.00 0.04 0.08 total\_sulfur\_dioxide\_probs

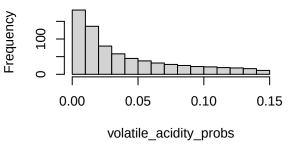
# **Histogram of sulphates\_probs**



# Histogram of alcohol\_probs

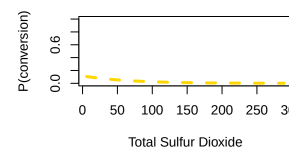


# Histogram of volatile\_acidity\_probs

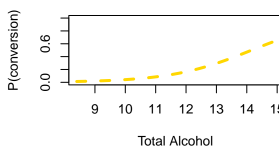


### Plot the results.

# **Probability of conversion**



# Probability of conversion



The plots of each of the four covariates are the probabilities across the range of # 6. Bayesian Logistic Regression

The following is a Bayesian logistic regression analysis.

To perform a Bayesian analysis via Metropolis-Hastings algorithm we write a function defining the target distribution of the Beta coefficients. We work in terms of log posterior to avoid numerical problems with computer computation.

### Likelihood.

The likelihood is a product of Bernoulli likelihoods for each combination of covariate and response variable for the i-th term.

$$L(\beta; y, x) = \prod_{i=1}^{n} \left( \frac{\exp(x_i \beta)}{1 + \exp(x_i \beta)} \right)^{y_i} \left( 1 - \frac{\exp(x_i \beta)}{1 + \exp(x_i \beta)} \right)^{1 - y_i}$$

## Log-likelihood

$$\log L(\beta; y, x) = \sum_{i=1}^{n} y_i(x_i \beta) - \log[1 + \exp(x_i \beta)]$$

### Prior

Independent normal prior distributions for each  $\beta_j$ , j = 0, 1, ..., k, such that  $\beta_j \sim N(0, 100)$ 

### Log posterior distribution

```
lpost.LR <- function(beta, x, y) {</pre>
  # beta: vector of coefficients
  # x: covariates (regressors)
  # y: response variable
  # eta is the matrix product of the covariates and the coefficients
  eta <- as.numeric(x %*% beta)</pre>
  # logp: y=1, in terms of logistic function
  logp <- eta - log(1 + exp(eta)) # in log scale</pre>
  # logq: y=0, expressed in terms of logistic function
  logq \leftarrow log(1 - exp(logp))
  # sum of both contributions: when y=1 + when y=0
  logl \leftarrow sum(logp[y==1]) + sum(logq[y==0])
  # log prior: normal indep prior dists $N(0, 100)$
  lprior <- sum(dnorm(beta, 0, 10, log = T))</pre>
  # return log posterior (likelihood + prior)
  return(log1 + lprior)
}
```

The MLE for each coefficient can be used to initialise the simulation matrix

```
fit <- glm(
    #good ~ volatile_acidity + total_sulfur_dioxide + sulphates + alcohol, # test model
    good ~ . - quality - drinkable, # full model
    data = wine,
    family = binomial(link="logit")
)</pre>
```

Set seed for reproducible results

```
set.seed(1234)
```

### Number of simulations

```
S <- 10<sup>4</sup>
```

Set X and y variables

### Initialisations for the coefficients

The first initialisation contains the MLE estimates for each coefficient. The remaining three initialisations are randomly selected values between 500 and 1000 from the uniform distribution runif(48, 500, 1000).

```
# init <- matrix(data=c(runif(20, min = 100, max = 200)), nrow=4, ncol=5, byrow = T) # for testing init <- matrix(data=c(runif(48, min = 500, max = 1000)), nrow=4, ncol=12, byrow = T)
```

### Run a Metropolis-Hastings algorithm

```
# First initialisation
beta_mat_init1 <- matrix(NA, nrow = S, ncol = ncol(X))</pre>
k <- ncol(beta mat init1)</pre>
#beta mat init1[1,] <- init[1,]
beta_mat_init1[1,] <- as.numeric(coefficients(fit)) # initialise with MLE of each coefficient
acc <- 0
for (iter in 2:S) {
  # simulate all (k) values using previous value of beta as mean and set sd
 beta_star <- rnorm(k, beta_mat_init1[iter-1,], 5)</pre>
  # compute target distribution for proposed value
  newpost = lpost.LR(beta_star, X, y)
  # compute target distribution for old value
  oldpost = lpost.LR(beta_mat_init1[iter-1,], X, y) # symmetric dist => no ratio computed
  # acceptance step, in log scale
  if (runif(1,0,1) > exp(newpost - oldpost)) {
    # chain doesn't move
    beta_mat_init1[iter,] = beta_mat_init1[iter-1,]
  } else {
    # add to chain and add 1 to counter
    beta mat init1[iter,] = beta star
    acc=acc + 1
 }
  #if (iter%%1000 == 0) {print(c(iter, acc/iter))}
print(c(iter, acc/iter))
```

### ## [1] 1.000e+04 3.046e-01

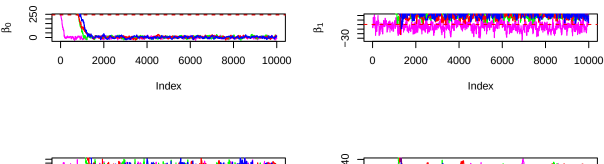
```
# Second initialisation
beta_mat_init2 <- matrix(NA, nrow = S, ncol = ncol(X))
k <- ncol(beta_mat_init2)
beta_mat_init2[1,] <- init[2,]
acc <- 0
for(iter in 2:S){
   beta_star <- rnorm(k, beta_mat_init2[iter-1,], 5)
   newpost = lpost.LR(beta_star, X, y)
   oldpost = lpost.LR(beta_mat_init2[iter-1,], X, y)
   if (runif(1,0,1) > exp(newpost - oldpost)) {
      beta_mat_init2[iter,] = beta_mat_init2[iter-1,]
   } else {
```

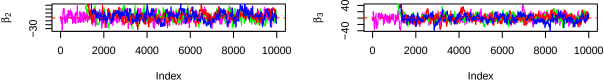
```
beta_mat_init2[iter,] = beta_star
    acc=acc + 1
 }
  #if (iter%%1000==0) {print(c(iter,acc/iter))}
print(c(iter, acc/iter))
## [1] 1.000e+04 2.045e-01
# Third initialisation
beta_mat_init3 <- matrix(NA,nrow=S,ncol=ncol(X))</pre>
k <- ncol(beta_mat_init3)</pre>
beta_mat_init3[1,] <- init[3,]</pre>
acc <- 0
for(iter in 2:S){
  beta_star <- rnorm(k,beta_mat_init3[iter-1,], 5)</pre>
 newpost=lpost.LR(beta_star,X,y)
  oldpost=lpost.LR(beta_mat_init3[iter-1,],X,y)
  if(runif(1,0,1)>exp(newpost-oldpost)){
    beta_mat_init3[iter,]=beta_mat_init3[iter-1,]
  } else{
    beta_mat_init3[iter,]=beta_star
    acc=acc+1
  #if(iter%%1000==0){print(c(iter,acc/iter))}
print(c(iter, acc/iter))
## [1] 1.000e+04 2.057e-01
# Forth initialisation
beta_mat_init4 <- matrix(NA,nrow=S,ncol=ncol(X))</pre>
k <- ncol(beta mat init4)</pre>
beta_mat_init4[1,] <- init[4,]</pre>
acc <- 0
for(iter in 2:S){
  beta_star <- rnorm(k,beta_mat_init4[iter-1,], 5)</pre>
  newpost=lpost.LR(beta_star,X,y)
  oldpost=lpost.LR(beta_mat_init4[iter-1,],X,y)
  if(runif(1,0,1)>exp(newpost-oldpost)){
    beta_mat_init4[iter,]=beta_mat_init4[iter-1,]
  } else{
    beta_mat_init4[iter,]=beta_star
    acc=acc+1
  #if(iter%%1000==0){print(c(iter,acc/iter))}
print(c(iter, acc/iter))
```

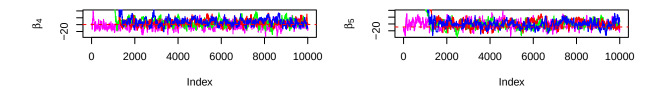
## [1] 1.00e+04 2.14e-01

Plot the chains for each coefficient (same plot)

```
par(mfrow=c(3,2))
plot(beta_mat_init1[,1], type="1", col="magenta", ylab=expression(beta[0]))
lines(beta_mat_init2[,1], type="l", col="green")
lines(beta_mat_init3[,1], type="l", col="red")
lines(beta_mat_init4[,1], type="1", col="blue")
abline(h=fit$coefficients[1],col="red",lty=2)
plot(beta_mat_init1[,2], type="l", col="magenta", ylab=expression(beta[1]))
lines(beta_mat_init2[,2], type="1", col="green")
lines(beta_mat_init3[,2], type="1", col="red")
lines(beta_mat_init4[,2], type="1", col="blue")
abline(h=fit$coefficients[2],col="red",lty=2)
plot(beta_mat_init1[,3], type="1", col="magenta", ylab=expression(beta[2]))
lines(beta_mat_init2[,3], type="1", col="green")
lines(beta_mat_init3[,3], type="l", col="red")
lines(beta_mat_init4[,3], type="1", col="blue")
abline(h=fit$coefficients[3],col="red",lty=2)
plot(beta_mat_init1[,4], type="1", col="magenta", ylab=expression(beta[3]))
lines(beta_mat_init2[,4], type="1", col="green")
lines(beta_mat_init3[,4], type="1", col="red")
lines(beta_mat_init4[,4], type="l", col="blue")
abline(h=fit$coefficients[4],col="red",lty=2)
plot(beta_mat_init1[,5], type="l", col="magenta", ylab=expression(beta[4]))
lines(beta_mat_init2[,5], type="1", col="green")
lines(beta_mat_init3[,5], type="1", col="red")
lines(beta_mat_init4[,5], type="1", col="blue")
abline(h=fit$coefficients[5],col="red",lty=2)
plot(beta_mat_init1[,6], type="l", col="magenta", ylab=expression(beta[5]))
lines(beta_mat_init2[,6], type="1", col="green")
lines(beta_mat_init3[,6], type="1", col="red")
lines(beta_mat_init4[,6], type="1", col="blue")
abline(h=fit$coefficients[6],col="red",lty=2)
```



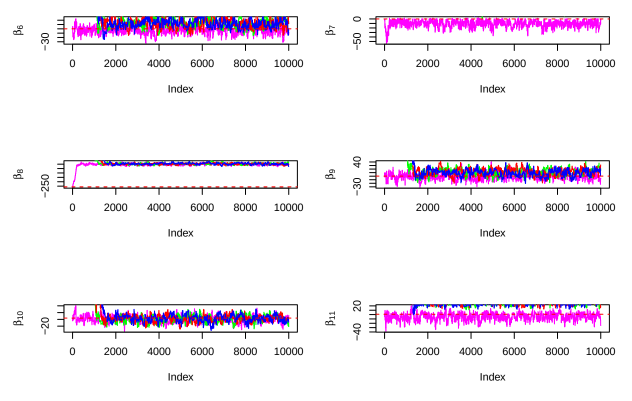




```
par(mfrow=c(3,2))
plot(beta_mat_init1[,7], type="l", col="magenta", ylab=expression(beta[6]))
lines(beta_mat_init2[,7], type="1", col="green")
lines(beta_mat_init3[,7], type="1", col="red")
lines(beta mat init4[,7], type="1", col="blue")
abline(h=fit$coefficients[7],col="red",lty=2)
plot(beta_mat_init1[,8], type="l", col="magenta", ylab=expression(beta[7]))
lines(beta_mat_init2[,8], type="1", col="green")
lines(beta_mat_init3[,8], type="1", col="red")
lines(beta_mat_init4[,8], type="1", col="blue")
abline(h=fit$coefficients[8],col="red",lty=2)
plot(beta_mat_init1[,9], type="l", col="magenta", ylab=expression(beta[8]))
lines(beta_mat_init2[,9], type="1", col="green")
lines(beta_mat_init3[,9], type="1", col="red")
lines(beta_mat_init4[,9], type="1", col="blue")
abline(h=fit$coefficients[9],col="red",lty=2)
plot(beta_mat_init1[,10], type="l", col="magenta", ylab=expression(beta[9]))
lines(beta_mat_init2[,10], type="l", col="green")
lines(beta mat init3[,10], type="l", col="red")
lines(beta_mat_init4[,10], type="1", col="blue")
abline(h=fit$coefficients[10],col="red",lty=2)
```

```
plot(beta_mat_init1[,11], type="l", col="magenta", ylab=expression(beta[10]))
lines(beta_mat_init2[,11], type="l", col="green")
lines(beta_mat_init3[,11], type="l", col="red")
lines(beta_mat_init4[,11], type="l", col="blue")
abline(h=fit$coefficients[11],col="red",lty=2)

plot(beta_mat_init1[,12], type="l", col="magenta", ylab=expression(beta[11]))
lines(beta_mat_init2[,12], type="l", col="green")
lines(beta_mat_init3[,12], type="l", col="red")
lines(beta_mat_init4[,12], type="l", col="blue")
abline(h=fit$coefficients[12],col="red",lty=2)
```



## COMMENT

## 7. Posterior Predictive Distribution

Approximate the posterior predictive distribution of an unobserved variable characterised by the following values for each covariate:

```
fixed_acidity <- 7.5
volatile_acidity <- 0.6
citric_acid <- 0.0
residual_sugar <- 1.7
chlorides <- 0.085</pre>
```

```
free_sulfur_dioxide <- 5</pre>
total_sulfur_dioxide <- 45
density <- 0.9965
pH < -3.4
sulphates <- 0.63
alcohol <- 12
S <- 20000
beta_mat2 <- matrix(NA, nrow = S, ncol = ncol(X))</pre>
beta_mat2[1,] <- as.numeric(coefficients(fit))</pre>
y_new <- c(1)</pre>
\# x_new \leftarrow c(1, 5, 0.63, 12, 0.6) \# for testing
x_new <- c(1, fixed_acidity, volatile_acidity, citric_acid, residual_sugar,</pre>
           chlorides, free_sulfur_dioxide, total_sulfur_dioxide, density,
           pH, sulphates, alcohol)
new model
library(mvtnorm)
# prediction
Omega_prop <- solve(t(X) %*% X)</pre>
k <- ncol(beta_mat2)</pre>
acc <- 0
for(iter in 2:S)
  # 1. Propose a new set of values
  beta_star <- rmvnorm(1, beta_mat2[iter-1,], 1.5 * Omega_prop)</pre>
  # 2. Compute the posterior density on the proposed value and on the old value
  newpost=lpost.LR(t(beta_star), X, y)
  oldpost=lpost.LR(matrix(beta_mat2[iter-1,], ncol=1), X, y)
  # 3. Acceptance step
  if (runif(1, 0, 1) > exp(newpost - oldpost)) {
    beta_mat2[iter,] = beta_mat2[iter-1,]
  } else {
    beta_mat2[iter,] = beta_star
    acc = acc + 1
  # 4. Print the stage of the chain
  if (iter%%1000 == 0){ print(c(iter, acc/iter)) }
  # 5. Prediction
  p_new <- exp(sum(beta_mat2[iter,] * x_new) ) / (1 + exp(sum(beta_mat2[iter,] * x_new) ))</pre>
 y_new[iter] <- rbinom(1,1,prob=p_new)</pre>
## [1] 1000.000
                    0.197
## [1] 2000.0000
                   0.1945
```

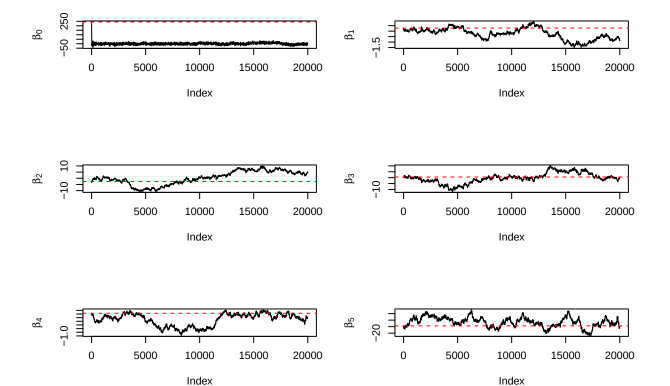
## [1] 3000.0000000

0.1936667

```
## [1] 4000.000
                   0.201
## [1] 5000.0000
                    0.2026
## [1] 6000.0000
                    0.2035
## [1] 7000.0000000
                       0.2041429
## [1] 8000.00000
                     0.20125
## [1] 9000.0000000
                       0.2042222
## [1] 1.000e+04 2.048e-01
## [1] 1.100000e+04 2.048182e-01
## [1] 1.200000e+04 2.051667e-01
## [1] 1.300000e+04 2.047692e-01
## [1] 1.40e+04 2.04e-01
## [1] 1.500e+04 2.044e-01
## [1] 1.6000e+04 2.0525e-01
## [1] 1.700000e+04 2.058235e-01
## [1] 1.800000e+04 2.055556e-01
## [1] 1.900000e+04 2.061579e-01
## [1] 2.0000e+04 2.0725e-01
```

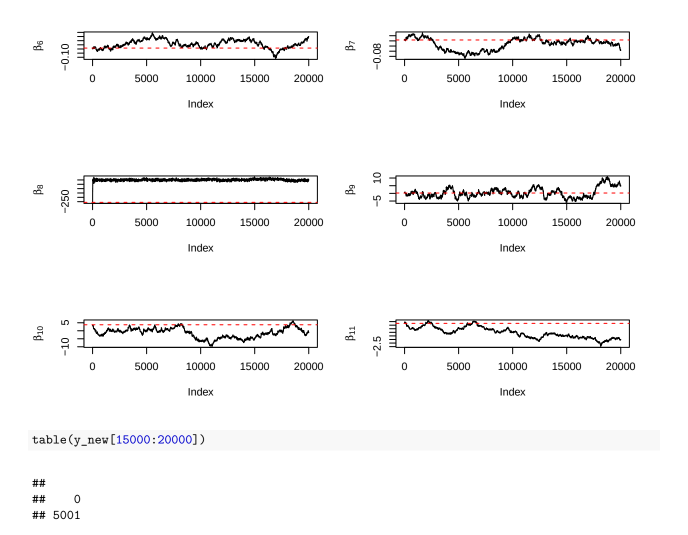
### Plots

```
par(mfrow=c(3,2))
plot(beta_mat2[,1],type="l", ylab=expression(beta[0]))
abline(h=fit$coefficients[1],col="red",lty=2)
plot(beta_mat2[,2],type="l", ylab=expression(beta[1]))
abline(h=fit$coefficients[2],col="red",lty=2)
plot(beta_mat2[,3],type="l", ylab=expression(beta[2]))
abline(h=fit$coefficients[3],col="red",lty=2)
plot(beta_mat2[,4],type="l", ylab=expression(beta[3]))
abline(h=fit$coefficients[4],col="red",lty=2)
plot(beta_mat2[,5],type="l", ylab=expression(beta[4]))
abline(h=fit$coefficients[5],col="red",lty=2)
plot(beta_mat2[,6],type="l", ylab=expression(beta[5]))
abline(h=fit$coefficients[6],col="red",lty=2)
```



### Plots

```
par(mfrow=c(3,2))
plot(beta_mat2[,7],type="l", ylab=expression(beta[6]))
abline(h=fit$coefficients[7],col="red",lty=2)
plot(beta_mat2[,8],type="l", ylab=expression(beta[7]))
abline(h=fit$coefficients[8],col="red",lty=2)
plot(beta_mat2[,9],type="l", ylab=expression(beta[8]))
abline(h=fit$coefficients[9],col="red",lty=2)
plot(beta_mat2[,10],type="l", ylab=expression(beta[9]))
abline(h=fit$coefficients[10],col="red",lty=2)
plot(beta_mat2[,11],type="l", ylab=expression(beta[10]))
abline(h=fit$coefficients[11],col="red",lty=2)
plot(beta_mat2[,12],type="l", ylab=expression(beta[11]))
abline(h=fit$coefficients[12],col="red",lty=2)
```



# 8. metrop() analysis of Q6

return(log1 - sum(beta^2) / 8)

Implementation of the metrop() function was through the MCMC Package Example, Charles J. Geyer

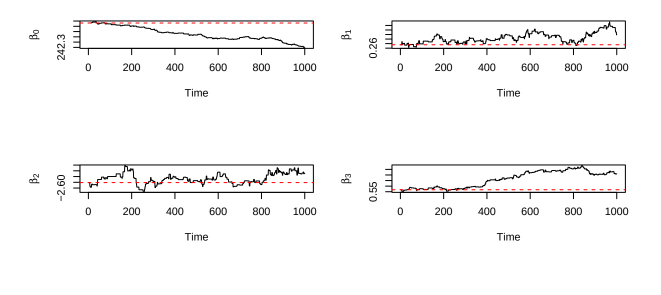
```
library(mcmc)
# out <- glm(wine$quality ~ wine$volatile_acidity + wine$total_sulfur_dioxide + wine$sulphates + wine$a
# summary(out)

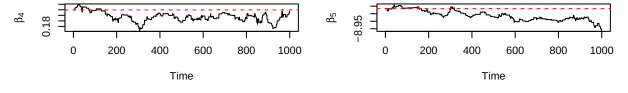
# full mode!
out <- glm(good ~ . - quality - drinkable, data=wine, family=binomial, x=TRUE)

lupost_factory <- function(x, y) function(beta) {
   eta <- as.numeric(x %*% beta)
   logp <- ifelse(eta < 0, eta - log1p(exp(eta)), - log1p(exp(- eta)))
   logq <- ifelse(eta < 0, - log1p(exp(eta)), - eta - log1p(exp(- eta)))
   log1 <- sum(logp[y == 1]) + sum(logq[y == 0])</pre>
```

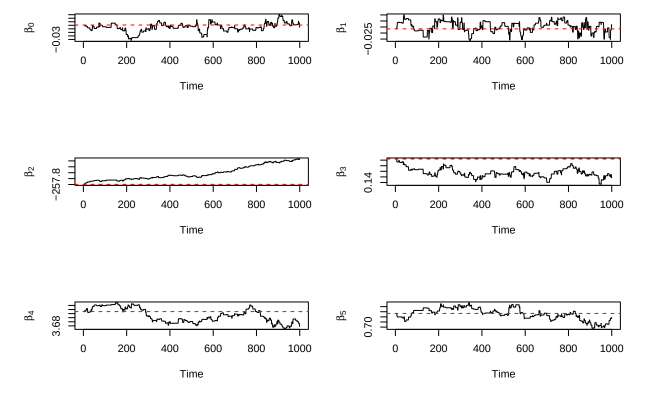
```
lupost <- lupost_factory(out$x, out$y)</pre>
set.seed(1234)
beta.init <- as.numeric(coefficients(out))</pre>
out <- metrop(lupost, beta.init, 1e3)</pre>
names(out)
   [1] "accept"
                        "batch"
                                        "initial"
                                                        "final"
                                                                         "accept.batch"
## [6] "initial.seed" "final.seed"
                                        "time"
                                                         "lud"
                                                                         "nbatch"
## [11] "blen"
                        "nspac"
                                        "scale"
                                                         "debug"
Look at the acceptance rate.
out$accept
## [1] 0
This is very low, so we can adjust the scale parameter to find an acceptance rate around 20%.
out <- metrop(out, scale = 0.0075)
out$accept
## [1] 0.213
par(mfrow=c(3,2))
plot(ts(out$batch)[,1], ylab=expression(beta[0]))
abline(h=fit$coefficients[1],col="red",lty=2)
plot(ts(out$batch)[,2], ylab=expression(beta[1]))
abline(h=fit$coefficients[2],col="red",lty=2)
plot(ts(out$batch)[,3], ylab=expression(beta[2]))
abline(h=fit$coefficients[3],col="red",lty=2)
```

plot(ts(out\$batch)[,4], ylab=expression(beta[3]))
abline(h=fit\$coefficients[4],col="red",lty=2)
plot(ts(out\$batch)[,5], ylab=expression(beta[4]))
abline(h=fit\$coefficients[5],col="red",lty=2)
plot(ts(out\$batch)[,6], ylab=expression(beta[5]))
abline(h=fit\$coefficients[6],col="red",lty=2)





```
par(mfrow=c(3,2))
plot(ts(out$batch)[,7], ylab=expression(beta[0]))
abline(h=fit$coefficients[7],col="red",lty=2)
plot(ts(out$batch)[,8], ylab=expression(beta[1]))
abline(h=fit$coefficients[8],col="red",lty=2)
plot(ts(out$batch)[,9], ylab=expression(beta[2]))
abline(h=fit$coefficients[9],col="red",lty=2)
plot(ts(out$batch)[,10], ylab=expression(beta[3]))
abline(h=fit$coefficients[10],col="red",lty=2)
plot(ts(out$batch)[,11], ylab=expression(beta[4]))
abline(h=fit$coefficients[11],col="red",lty=2)
plot(ts(out$batch)[,12], ylab=expression(beta[5]))
abline(h=fit$coefficients[12],col="red",lty=2)
```



## VISUAL COMPARISON