Worksheet 5

Foundations of Bayesian Methodology

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Exercise 3 - Normal example in JAGS

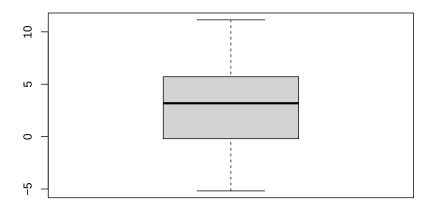
rjags interface to JAGS

The steps of list.factories and set.factories show and control over the status of factories in JAGS modules. The steps of jags.modules shows the names of the currently loaded modules and also loads or unloads JAGS modules.

```
# Load required libraries
library(rjags)
library(coda)
library(MASS)
library(FNN)
library(INLA)
library(runjags)
```

```
list.factories(type = "rng")
list.factories(type = "monitor")
list.factories(type = "sampler")
set.factory(name = "base::Slice", type = "sampler", state = FALSE)
list.factories(type = "sampler")
set.factory(name = "base::Slice", type = "sampler", state = TRUE)
list.factories(type = "sampler")
list.modules()
load.module("glm")
list.modules()
unload.module("glm")
list.modules()
```

The following lines of code provide us with a basic setup



```
summary(y)

## Min. 1st Qu. Median Mean 3rd Qu. Max.
## -5.1820  0.3307  3.1815  3.1884  5.6765 11.1440

sd(y)

## [1] 4.046692

# Define the parameters of the prior distributions
mu0 <- -3
sigma2_0 <- 4
a0 <- 1.6
b0 <- 0.4</pre>
```

Boxplot shows that the median of observed data is around 3, the 25% quantile and 75% quantile are around 0 and 5, respectively. In addition, the minimum and maximum are around -5 and 11, respectively.

This step gives observed values, sets up initial values and specifies model.

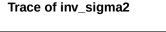
```
wb_inits <- list( mu=-0.2381084, inv_sigma2=0.3993192,
                  .RNG.name="base::Wichmann-Hill", .RNG.seed=314159)
modelString = " # open quote for modelString
model{
# likelihood
for (i in 1:N){
y[i] ~ dnorm( mu, inv_sigma2 )
# Priors
mu ~ dnorm( -3, 0.25 ) # prior for mu N(mu0, prec=1/sigma2_0)
inv_sigma2 ~ dgamma( 1.6, 0.4 ) # prior for precision G(a0, b0)
# transformations
# deterministic definition of variance
sigma2 <- 1/inv_sigma2</pre>
# deterministic definition of standard deviation
sigma <- sqrt(sigma2)</pre>
}
" # close quote for modelString
writeLines(modelString, con="TempModelexe3.txt") # write to a file
# model initiation
model.jags <- jags.model(</pre>
 file = "TempModelexe3.txt",
 data = wb_data,
 inits = wb_inits,
 n.chains = 1,
 n.adapt = 4000
)
## Compiling model graph
      Resolving undeclared variables
##
##
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 30
##
      Unobserved stochastic nodes: 2
##
      Total graph size: 41
##
## Initializing model
str(model.jags)
## List of 8
             :function ()
## $ ptr
## $ data
             :function ()
## $ model
              :function ()
## $ state
               :function (internal = FALSE)
## $ nchain :function ()
## $ iter :function ()
              :function ()
## $ sync
## $ recompile:function ()
## - attr(*, "class")= chr "jags"
```

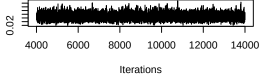
```
class(model.jags)
## [1] "jags"
attributes(model.jags)
## $names
                                           "state"
## [1] "ptr"
                   "data"
                               "model"
                                                       "nchain"
                                                                   "iter"
## [7] "sync"
                   "recompile"
##
## $class
## [1] "jags"
list.samplers(model.jags)
## $'bugs::ConjugateNormal'
## [1] "mu"
##
## $'bugs::ConjugateGamma'
## [1] "inv_sigma2"
update(model.jags, n.iter = 4000) # burn-in
# sampling
fit.jags.coda <- coda.samples(</pre>
 model = model.jags,
 variable.names = c("mu", "sigma2", "inv_sigma2"),
 n.iter = 10000,
 thin = 1
str(fit.jags.coda)
## List of 1
## $ : 'mcmc' num [1:10000, 1:3] 0.0802 0.0629 0.063 0.0611 0.0624 ...
## ..- attr(*, "dimnames")=List of 2
## ....$ : NULL
    .. ..$ : chr [1:3] "inv_sigma2" "mu" "sigma2"
## ..- attr(*, "mcpar")= num [1:3] 4001 14000 1
## - attr(*, "class")= chr "mcmc.list"
class(fit.jags.coda)
## [1] "mcmc.list"
attributes(fit.jags.coda)
## $class
## [1] "mcmc.list"
summary(fit.jags.coda)
```

Iterations = 4001:14000

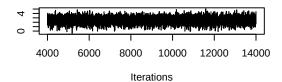
```
## Thinning interval = 1
## Number of chains = 1
## Sample size per chain = 10000
## 1. Empirical mean and standard deviation for each variable,
##
      plus standard error of the mean:
##
##
                  Mean
                            SD Naive SE Time-series SE
               0.06604 0.01696 0.0001696
## inv_sigma2
                                               0.0001839
## mu
               2.46929 0.70519 0.0070519
                                               0.0074453
              16.21924 4.52002 0.0452002
## sigma2
                                               0.0488709
##
## 2. Quantiles for each variable:
##
                 2.5%
##
                           25%
                                     50%
                                              75%
                                                   97.5%
## inv_sigma2 0.03686
                       0.05398
                                0.06455
                                          0.07649
                                                   0.103
## mu
              1.01783
                       2.01871
                                2.49325
                                          2.93789
                                                   3.821
              9.71207 13.07363 15.49151 18.52587 27.132
## sigma2
```

plot(fit.jags.coda)

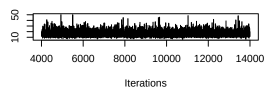




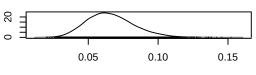
Trace of mu



Trace of sigma2

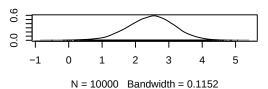


Density of inv_sigma2

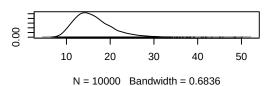


N = 10000 Bandwidth = 0.002822

Density of mu



Density of sigma2



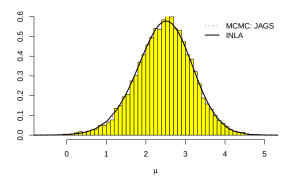
```
# store samples for each parameter from the chain into separate objects
m.fit.jags.coda <- as.matrix(fit.jags.coda)
mu.sim <- m.fit.jags.coda[,"mu"]
sigma2.sim <- m.fit.jags.coda[,"sigma2"]
inv_sigma2.sim <- m.fit.jags.coda[,"inv_sigma2"]

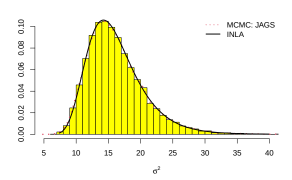
# plot for mean
rg <- range(inla.output$marginals.fixed$"(Intercept)"[,2])
truehist(mu.sim, prob=TRUE, col="yellow", xlab=expression(mu),ylim=rg)</pre>
```

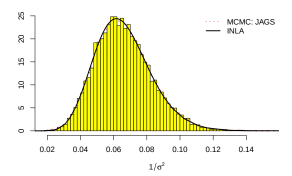
[1] -0.01205037

```
# plot for variance
m_var <-inla.tmarginal(function(x) 1/x, inla.output$marginals.hyperpar[[1]])</pre>
rg <- range(m_var[,2])</pre>
truehist(sigma2.sim, prob=TRUE, col="yellow", xlab=expression(sigma^2),ylim=rg, xlim=c(5,40))
lines(density(sigma2.sim), lty=3, lwd=3, col=2)
lines(m_var, lwd=2)
legend("topright",c("MCMC: JAGS","INLA"),lty=c(3,1),lwd=c(2,2),col=c(2,1),
       cex=1.0,bty="n")
# plot for precision
truehist(inv_sigma2.sim, prob=TRUE, col="yellow", xlab=expression(1/sigma^2))
lines(density(inv_sigma2.sim),lty=3,lwd=3, col=2)
lines(inla.output$marginals.hyperpar[[1]],lwd=2)
legend("topright",c("MCMC: JAGS","INLA"),lty=c(3,1),lwd=c(2,2),col=c(2,1),
       cex=1.0,bty="n")
KL.divergence(inla.rmarginal(length(inv sigma2.sim),
                              inla.output$marginals.hyperpar[[1]]),
              inv_sigma2.sim, k = 1)
```

[1] -0.01645461







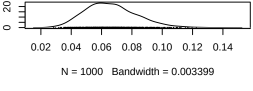
The empirical mean and standard deviation for each variable and corresponding quantiles are shown in the output. The traceplot as well as the plot of density indicate good mixing for three different parameters: precision, mean and variance. When compared to the result of INLA, whether plots or the results of Kullback-Leibler divergence proves the similarity between the output of INLA and JAGS.

```
wb_inits <- function() {</pre>
  list(mu = rnorm(1),
       inv_sigma2 = runif(1)
}
# model initialisation
model.jags <- jags.model(</pre>
  file = "TempModelexe3.txt",
  data = wb_data,
  inits = wb_inits,
  n.chains = 4,
  n.adapt = 4000
## Compiling model graph
      Resolving undeclared variables
      Allocating nodes
##
## Graph information:
##
      Observed stochastic nodes: 30
      Unobserved stochastic nodes: 2
##
##
      Total graph size: 41
##
## Initializing model
update(model.jags, n.iter = 4000) # burn-in
# sampling/monitoring
fit.jags.coda <- coda.samples(</pre>
  model = model.jags,
  variable.names = c("mu", "sigma2", "inv_sigma2"),
  n.iter = 10000,
  thin = 10
summary(fit.jags.coda)
##
## Iterations = 4010:14000
## Thinning interval = 10
## Number of chains = 4
## Sample size per chain = 1000
## 1. Empirical mean and standard deviation for each variable,
##
      plus standard error of the mean:
##
##
                             SD Naive SE Time-series SE
                  Mean
## inv_sigma2  0.06618  0.01693  0.0002677
                                               0.0002676
## mu
               2.45764 0.70427 0.0111355
                                                0.0117491
## sigma2
              16.16827 4.41901 0.0698708
                                                0.0698405
##
## 2. Quantiles for each variable:
##
                           25%
                                    50%
                 2.5%
## inv_sigma2 0.03755 0.0541 0.06481 0.07667 0.1028
```

```
## mu 1.01307 1.9981 2.47869 2.94081 3.8028
## sigma2 9.72874 13.0426 15.43075 18.48385 26.6341
```

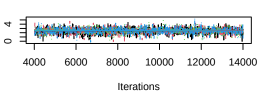
plot(fit.jags.coda)

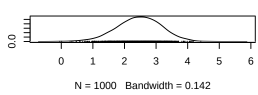
Trace of inv_sigma2 4000 6000 8000 10000 12000 14000 Iterations Trace of mu

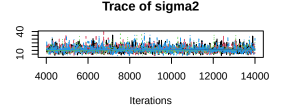


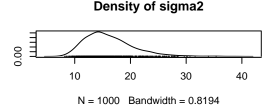
Density of mu

Density of inv_sigma2



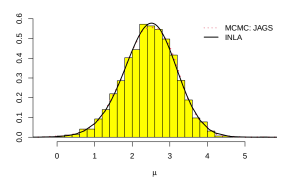


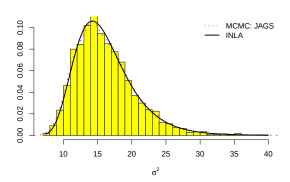


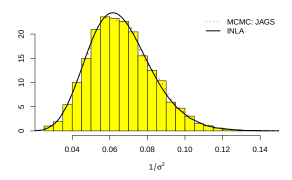


[1] 0.01068301

[1] -0.01211412







The empirical mean and standard deviation for each variable and corresponding quantiles are shown in the output. Traceplots and density plots of the four chain model show similar results when compared to the single chain model. Moreover we can find good mixing for the three parameters (mean, variance and precision); INLA and JAGS have similar results.

effectiveSize(fit.jags.coda)

```
## inv_sigma2 mu sigma2
## 4000.000 3639.148 4000.000
```

lapply(fit.jags.coda, effectiveSize)

```
## [[1]]
## inv_sigma2 mu sigma2
## 1000 1000 1000
##
## [[2]]
```

```
## 1000.0000 843.0896 1000.0000
##
## [[3]]
## inv_sigma2
                  mu
                          sigma2
## 1000.0000 796.0581 1000.0000
##
## [[4]]
## inv_sigma2
                           sigma2
                   mu
##
       1000
                  1000
                            1000
gelman.diag(fit.jags.coda,autoburnin=TRUE)
## Potential scale reduction factors:
##
##
            Point est. Upper C.I.
## inv_sigma2
                  1
## mu
                     1
                               1
## sigma2
                     1
##
## Multivariate psrf
##
## 1
gelman.plot(fit.jags.coda,autoburnin=TRUE)
geweke.diag(fit.jags.coda)
## [[1]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
## inv_sigma2
                         sigma2
                   mu
     1.7268
              0.1135 -1.4485
##
##
##
## [[2]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
                mu
## inv_sigma2
                         sigma2
             0.5908
##
      0.1833
                         0.4241
##
##
## [[3]]
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
## inv_sigma2
                  mu
                         sigma2
##
   0.30775
             -0.12090
                       -0.07339
##
##
## [[4]]
## Fraction in 1st window = 0.1
```

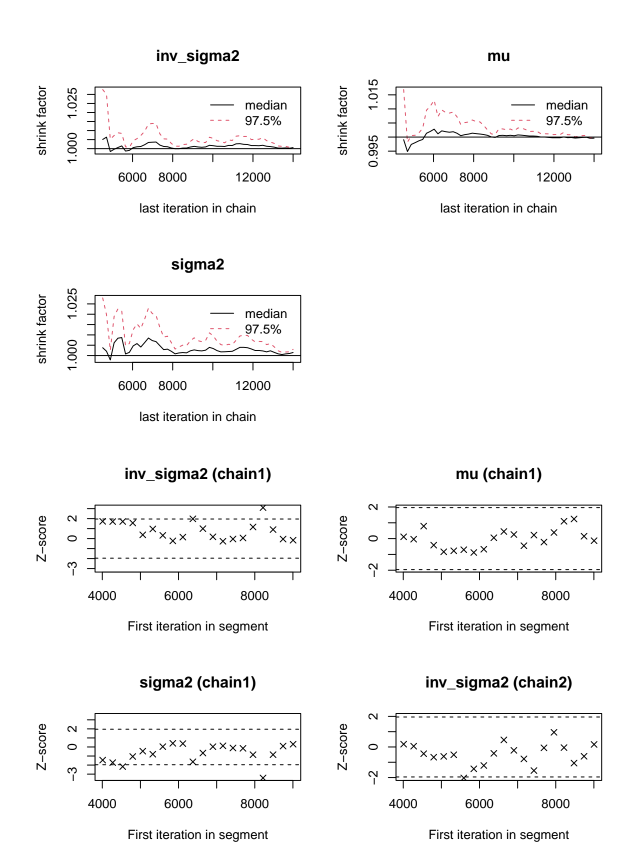
```
## Fraction in 2nd window = 0.5
##
## inv_sigma2
                              sigma2
                      \mathtt{mu}
## -0.058414
                0.280434
                            0.006464
geweke.plot(fit.jags.coda)
heidel.diag(fit.jags.coda)
## [[1]]
##
                                      p-value
##
              Stationarity start
##
              test
                            iteration
                                      0.213
## inv_sigma2 passed
                            1
## mu
              passed
                            1
                                      0.925
## sigma2
              passed
                            1
                                      0.207
##
              Halfwidth Mean
##
                                 Halfwidth
##
              test
## inv_sigma2 passed
                          0.0658 0.00102
                         2.4310 0.04354
## mu
              passed
                         16.1943 0.26482
## sigma2
              passed
##
## [[2]]
##
##
                                      p-value
              Stationarity start
##
              test
                            iteration
                                       0.618
## inv_sigma2 passed
                            1
## mu
              passed
                            1
                                       0.979
## sigma2
                            1
                                       0.213
              passed
##
##
              Halfwidth Mean
                                 Halfwidth
##
              test
                          0.0662 0.00104
## inv_sigma2 passed
              passed
                         2.4597 0.04807
                         16.1556 0.27077
## sigma2
              passed
##
## [[3]]
##
##
              Stationarity start
                                      p-value
##
              test
                            iteration
## inv_sigma2 passed
                                      0.389
## mu
                                      0.328
              passed
                            1
## sigma2
                                      0.536
              passed
                            1
##
##
              Halfwidth Mean
                                 Halfwidth
              test
                          0.0671 0.00104
## inv_sigma2 passed
                          2.4771 0.04948
## mu
              passed
## sigma2
              passed
                         15.9114 0.26671
##
## [[4]]
##
##
              Stationarity start
                                      p-value
                            iteration
##
              test
## inv_sigma2 passed
                            1
                                      0.815
## mu
                            1
                                      0.455
              passed
## sigma2
              passed
                            1
                                      0.527
```

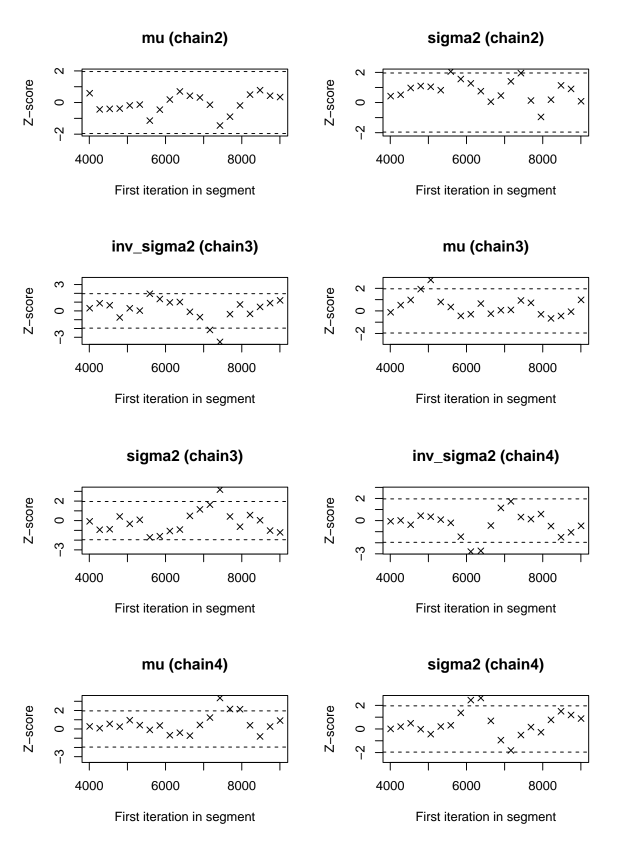
##

```
## Halfwidth Mean Halfwidth
## test
## inv_sigma2 passed 0.0656 0.00109
## mu passed 2.4627 0.04278
## sigma2 passed 16.4118 0.29194
```

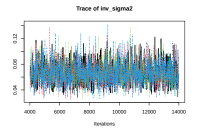
raftery.diag(fit.jags.coda)

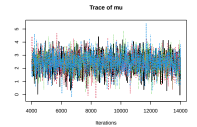
```
## [[1]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
## You need a sample size of at least 3746 with these values of q, r and s
##
## [[2]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
## You need a sample size of at least 3746 with these values of q, r and s
##
## [[3]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
## You need a sample size of at least 3746 with these values of q, r and s
## [[4]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
## You need a sample size of at least 3746 with these values of q, r and s
```

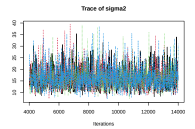




```
coda:::traceplot(fit.jags.coda)
# "DIC" penalised expected deviance computation
dic1<-dic.samples(model=model.jags, n.iter=1000, type="popt")</pre>
```







ESS: The Effective Sample Size of the three parameters are shown in the table. The ESS must be large enough to get stable inferences for quantities of interest. In the case of 10000 iterations, the global ESS of around 4000 is large enough.

	Global	1st Chain	2nd Chain	3rd Chain	4th Chain
Precision	3863.425	981.847	1000.000	881.578	1000
Mean	4099.911	1199.804	900.106	1000.000	1000
Variance	3931.376	931.376	1000.000	1000.000	1000

Table 1: Effective Sample Size

Gelman-Rubin-Brooks: The estimated shrink factors and corresponding upper CI of three parameters of Gelman and Rubin's convergence diagnostic are within the acceptable range. In addition, Gelman-Rubin-Brooks plot shows the evolution of shrink factor as the number of iterations increases. The trends of the median of shrink factors are mostly stable among three parameters in the range of 6000 to 14000 iterations.

Geweke: The results of Geweke's convergence diagnostic for 4 chaines are all in the range of -2 to 2. (However, the Geweke-Brooks plot indicates that the 4th chain has more samples who hold a Z-score that is not in the range of -2 to 2, which proves that the 4th chain has not reached equilibrium.) Newest pdf version doesnt support observations in brackets

Heidelberger & Welch: Heidelberger and Welch's convergence diagnostic shows a similar result as Geweke-Brooks plot. (The variance in the 4th chain fails the stationary test indicating non convergence therefore a longer MCMC run is needed). All parameters (except the beforementioned one) pass the halfwidth test which means that the length of the sample is long enough to estimate the mean with sufficient accuracy. **Newest pdf version doesnt support observations in brackets**

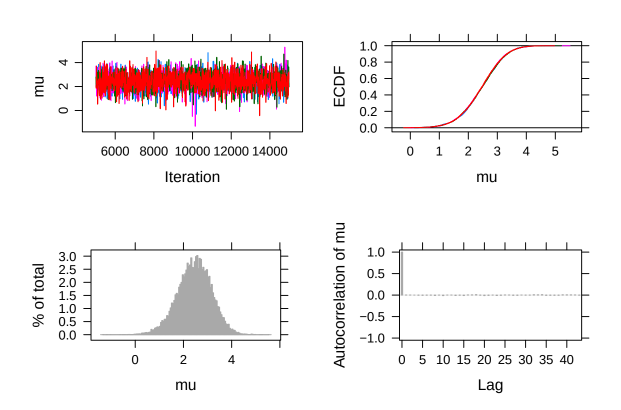
Raftery & Lewis Raftery and Lewis's diagnostic gives the minimum sample size based on zero auto-correlation. All the four chains need a minimum sample size of 3746.

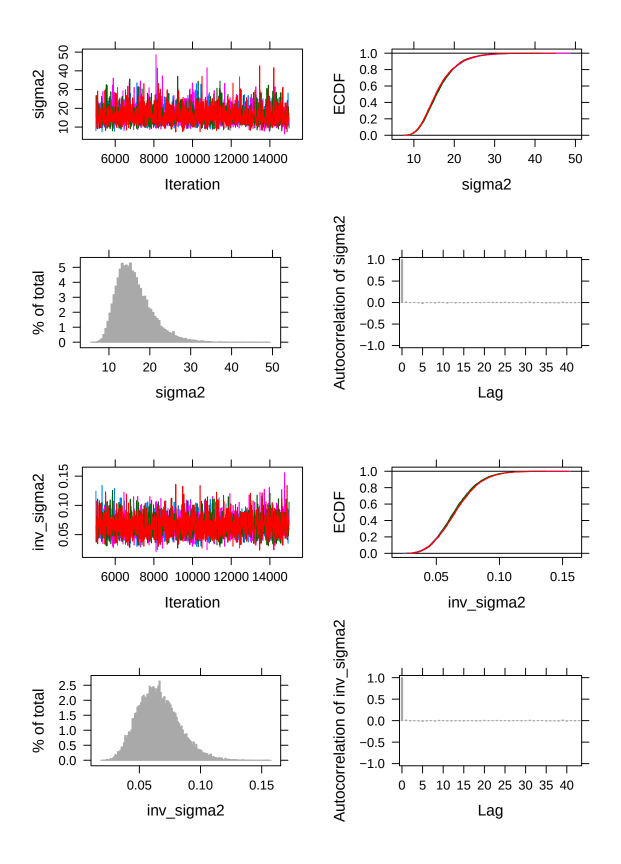
Trace plot Again the traceplots show that each of the four chains explores the distribution pretty good.

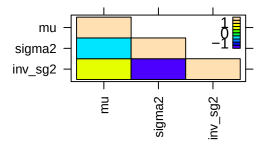
runjags interface with a link to a file

```
inits=wb_inits,
                      n.chains=4,
                      burnin=4000,
                      sample=5000,
                      adapt=1000,
                      thin=2)
## Compiling rjags model...
## Calling the simulation using the rjags method...
## Note: the model did not require adaptation
## Burning in the model for 4000 iterations...
## Running the model for 10000 iterations...
## Simulation complete
## Calculating summary statistics...
## Calculating the Gelman-Rubin statistic for 3 variables....
## Finished running the simulation
plot(fit.runjags)
## Generating plots...
print(fit.runjags)
##
## JAGS model summary statistics from 20000 samples (thin = 2; chains = 4; adapt+burnin = 5000):
##
              Lower95
                        Median Upper95
                                                        SD Mode
                                                                     MCerr MC%ofSD
                                             Mean
                                                                 0.0049834
## mu
                 1.055
                       2.4814 3.8224
                                           2.4547 0.70858 --
                                                                               0.7
                8.8395
                        15.528 25.331 16.289
                                                  4.4801
                                                             --
                                                                  0.031392
                                                                               0.7
## sigma2
## inv_sigma2 0.034094 0.064399 0.098675 0.065721 0.016858
                                                           -- 0.00011489
                                                                               0.7
##
                                 psrf
##
             SSeff
                       AC.20
## mu
             20217 -0.012233 0.99994
             20367 0.0036286 1.0003
## sigma2
## inv_sigma2 21530 0.0029035 1.0003
## Total time taken: 3.2 seconds
# CODA
fit.runjags.coda<-as.mcmc.list(fit.runjags)</pre>
summary(fit.runjags.coda)
## Iterations = 5001:14999
## Thinning interval = 2
## Number of chains = 4
## Sample size per chain = 5000
##
## 1. Empirical mean and standard deviation for each variable,
##
     plus standard error of the mean:
##
##
                           SD Naive SE Time-series SE
                  Mean
              2.45469 0.70858 0.0050104 0.0049845
## mu
            16.28939 4.48009 0.0316790
                                              0.0315059
## sigma2
## inv_sigma2 0.06572 0.01686 0.0001192
                                              0.0001152
```

##
2. Quantiles for each variable:
##
2.5% 25% 50% 75% 97.5%
mu 0.99944 1.99540 2.4814 2.94218 3.7834
sigma2 9.73985 13.10978 15.5283 18.66358 27.0930
inv_sigma2 0.03691 0.05358 0.0644 0.07628 0.1027







Using plot function except ordinary plots that appeared before, autocorrelation plots and a cross-correlation plot are produced.

Autocorrelation plots gives an overview of autocorrelation in different thinning interval. For all three parameters, when the lag is larger than 1, the autocorrelation is close to 0.

Cross-correlation plot shows the correlation between parameters. The correlation between variance and mean is close to 0, while variance and precision have a strong negative correlation which is not surprising given that the precision is the inverse of the variance.

R2jags wrapper to rjags interface to JAGS several chains

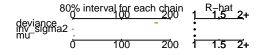
```
wb data <- list( N=30,
                  y=c(3.048,2.980,2.029,7.249,-0.259,3.061,4.059,6.370,7.902,1.926,
                      9.094,10.489,-0.384,-3.096,2.315,5.830,-1.542,-1.544,5.714,
                      -5.182,3.828,-4.038,2.169,5.087,-0.201,4.880,3.302,3.859,
                      11.144,5.564)
)
#define parameters
params<-c("mu", "sigma2", "inv_sigma2")</pre>
# define inits
inits1 <- list(mu=rnorm(1), inv_sigma2=runif(1),</pre>
                .RNG.name="base::Super-Duper", .RNG.seed=1)
inits2 <- list(mu=rnorm(1), inv sigma2=runif(1),</pre>
                .RNG.name="base::Wichmann-Hill", .RNG.seed=2)
wb_inits <- list(inits1,inits2)</pre>
fit.R2jags<-jags(data=wb_data,
                  inits=wb inits,
                  parameters.to.save=params,
                  model.file="05normal_exmple_JAGS.txt",
```

```
n.chains=2,
n.iter=50000,
n.burnin=4000,
n.thin=5,
DIC = TRUE,
    jags.seed = 321,
    refresh =100,
    digits = 4,
     jags.module = c("glm","dic"))
## module glm loaded
```

```
## Compiling model graph
## Resolving undeclared variables
## Allocating nodes
## Graph information:
## Observed stochastic nodes: 30
## Unobserved stochastic nodes: 2
## Total graph size: 41
##
## Initializing model
```

```
# Standard plots of the monitored variables
plot(fit.R2jags)
```

Bugs model at "05normal_exmple_JAGS.txt", fit using jags, 2 chains, each with 50000 iterations (first 4000 discarded)

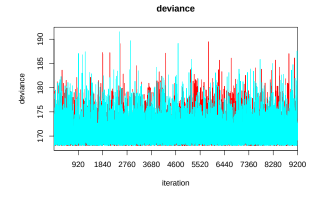


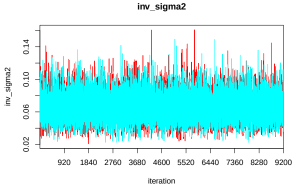
medians and 80% intervals

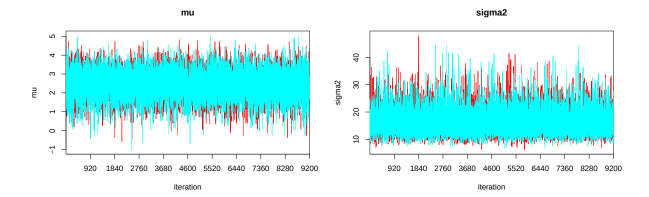
```
# Display summary statistics
print(fit.R2jags)
```

Inference for Bugs model at "O5normal_exmple_JAGS.txt", fit using jags,

```
## 2 chains, each with 50000 iterations (first 4000 discarded), n.thin = 5
##
   n.sims = 18400 iterations saved
              mu.vect sd.vect
                                                                 97.5% Rhat n.eff
##
                                 2.5%
                                           25%
                                                   50%
                                                           75%
## inv_sigma2
                0.066
                       0.017
                                0.037
                                         0.054
                                                 0.064
                                                         0.076
                                                                 0.103 1.001 14000
                        0.710
                                1.015
                                        1.997
                                                 2.474
                                                         2.938
## mu
                2.458
                                                                 3.805 1.001 18000
## sigma2
               16.268
                        4.455
                                9.751 13.144 15.508 18.597 27.078 1.001 14000
                        2.653 168.077 168.906 170.107 172.040 177.876 1.001 18000
              170.864
## deviance
##
## For each parameter, n.eff is a crude measure of effective sample size,
## and Rhat is the potential scale reduction factor (at convergence, Rhat=1).
##
## DIC info (using the rule, pD = var(deviance)/2)
## pD = 3.5 and DIC = 174.4
## DIC is an estimate of expected predictive error (lower deviance is better).
# traceplot
traceplot(fit.R2jags)
# CODA
fit.R2jags.coda<-as.mcmc(fit.R2jags)</pre>
summary(fit.R2jags.coda)
##
## Iterations = 4001:49996
## Thinning interval = 5
## Number of chains = 2
## Sample size per chain = 9200
##
## 1. Empirical mean and standard deviation for each variable,
##
      plus standard error of the mean:
##
                             SD Naive SE Time-series SE
##
                   Mean
              170.86446 2.65305 0.0195586
## deviance
                                                0.0193999
## inv sigma2
                0.06575 0.01677 0.0001236
                                                0.0001222
                2.45814 0.70964 0.0052316
                                                0.0051393
## m11
               16.26773 4.45496 0.0328424
                                               0.0323722
## sigma2
##
## 2. Quantiles for each variable:
##
##
                   2.5%
                                        50%
                                                   75%
                                                          97.5%
                              25%
## deviance
              168.07654 168.90602 170.10684 172.04030 177.8755
## inv_sigma2
                0.03693
                          0.05377
                                    0.06448
                                               0.07608
                                                         0.1026
                          1.99698
                                    2.47410
                                               2.93849
## mu
                1.01469
                                                         3.8053
## sigma2
                9.75132 13.14450 15.50786 18.59687 27.0781
```







The function R2jags::jags computes deviance when DIC is true, where DIC is an estimate of expected predictive error and lower deviance is better. Besides, using plot function, the quantiles for each variable are visualized.

Exercise 4 - Logistic regression in JAGS

```
# original covariate values
x <- c(0.0028, 0.0028, 0.0056, 0.0112, 0.0225, 0.0450)

# the centered covariate values (centered dose) from the Mice data from Collett
x_centered <- x - mean(x)

# number of mice deaths
y <- c(26, 9, 21, 9, 6, 1)

# total number of mice
n <- c(28, 12, 40, 40, 40, 40)

d.mice <- data.frame(
    x, y, n, x_centered, y/n, n-y
)
colnames(d.mice) <- c("$x$", "$y$", "$n$", "centered $x$", "$p$", "$alive$")
knitr::kable(d.mice, align="c", caption="Mice data from Collett (2003)")</pre>
```

Table 2: Mice data from Collett (2003)

\overline{x}	y	n	centered x	p	alive
0.0028	26	28	-0.0121833	0.9285714	2
0.0028	9	12	-0.0121833	0.7500000	3
0.0056	21	40	-0.0093833	0.5250000	19
0.0112	9	40	-0.0037833	0.2250000	31
0.0225	6	40	0.0075167	0.1500000	34
0.0450	1	40	0.0300167	0.0250000	39

Logistic model:

$$logit(p_i) = ln\left(\frac{p_i}{1 - p_i}\right) = \alpha + \beta x_i$$

$$p_i = \frac{\exp(\alpha + \beta x_i)}{1 + \exp(\alpha + \beta x_i)}$$

fit.classic <- $glm(cbind(y, (n-y)) \sim x_centered, data = d.mice, family = binomial)$

Classic Approach

summary(fit.classic)

```
##
## Call:
## glm(formula = cbind(y, (n - y)) ~ x_centered, family = binomial,
      data = d.mice)
##
## Deviance Residuals:
##
                          3
           0.4474 -0.9319 -2.2893 0.7546
## 3.0784
##
## Coefficients:
               Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.9800 0.2399 -4.085 4.41e-05 ***
                        26.3630 -5.564 2.63e-08 ***
## x_centered -146.6927
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 92.287 on 5 degrees of freedom
## Residual deviance: 18.136 on 4 degrees of freedom
## AIC: 40.805
## Number of Fisher Scoring iterations: 5
knitr::kable(coef(summary(fit.classic)), align="c", caption="Summary results for classic approach")
```

Table 3: Summary results for classic approach

	Estimate	Std. Error	z value	$\Pr(> z)$
(Intercept) x centered	-0.9800475 -146.6927209	$0.2399331 \\ 26.3629619$	-4.084669 -5.564349	4.41e-05 $0.00e+00$

	-0.01218	-0.00938	-0.00378	0.00752	0.03002
0	5	19	31	34	39
1	35	21	9	6	1

```
fit.glm <- glm(y_binary ~ x_centered, data = d.mice1, family = binomial)
summary(fit.glm)</pre>
```

```
##
## Call:
## glm(formula = y_binary ~ x_centered, family = binomial, data = d.mice1)
##
## Deviance Residuals:
      Min
            1Q
                    Median
                                  3Q
                                         Max
## -1.5337 -1.0030 -0.0957
                            0.8589
                                      3.2826
##
## Coefficients:
##
              Estimate Std. Error z value Pr(>|z|)
## (Intercept) -0.9796
                          0.2399 -4.084 4.43e-05 ***
                           26.3629 -5.564 2.63e-08 ***
## x_centered -146.6927
## ---
## Signif. codes: 0 '*** 0.001 '** 0.01 '* 0.05 '.' 0.1 ' 1
## (Dispersion parameter for binomial family taken to be 1)
##
      Null deviance: 261.37 on 199 degrees of freedom
## Residual deviance: 187.22 on 198 degrees of freedom
## AIC: 191.22
##
## Number of Fisher Scoring iterations: 6
```

Bayesian Approach

```
library(rjags)
library(coda)
library(ggplot2)
```

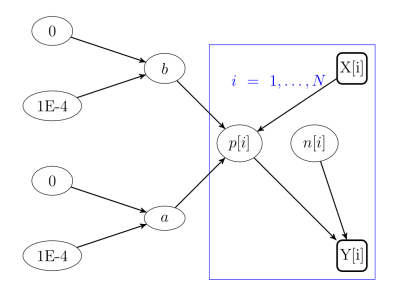
```
modelString <- "model{
  for (i in 1:length(y)) {
    y[i] ~ dbin(p[i],n[i])
    p[i] <- ilogit(alpha + beta * x[i])
}

alpha ~ dnorm(0, 1.0E-04)
beta ~ dnorm(0, 1.0E-04)
}"

writeLines(modelString, con="LogitModel.txt")</pre>
```

```
## Alternatively
modelString <- model{
  for (i in 1:length(y)) {
    y[i] ~ dbern(p[i])
    p[i] <- ilogit(alpha + beta * x[i])
  }

alpha ~ dnorm(0, 1.0E-04)
  beta ~ dnorm(0, 1.0E-04)
}"</pre>
```



```
## Set seed for reproducible results
set.seed(44566)

## Generate initial values based on estimates in classical logistic regression
inits.alpha <- coef(summary(fit.classic))[1, 1] +
   coef(summary(fit.classic))[1, 2] * rnorm(4)

inits.beta <- coef(summary(fit.classic))[2, 1] +
   coef(summary(fit.classic))[2, 2] * rnorm(4)

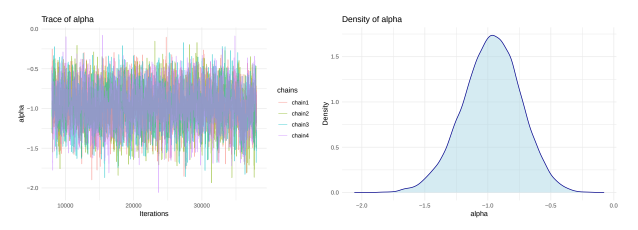
## Generate data list for JAGS
dat.jags <- list(y=y, x=x_centered, n=n)</pre>
## Set initial values and reader good for reproducible results
```

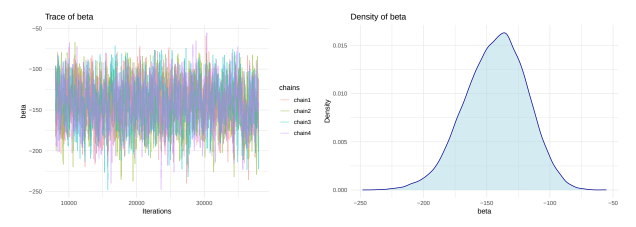
```
## Set initial values and random seed for reproducible results
inits.jags <- list(list(alpha=inits.alpha[1], beta=inits.beta[1],</pre>
                         .RNG.name="base::Wichmann-Hill", .RNG.seed=314159),
                   list(alpha=inits.alpha[2], beta=inits.beta[2],
                         .RNG.name="base::Marsaglia-Multicarry", .RNG.seed=159314),
                   list(alpha=inits.alpha[3], beta=inits.beta[3],
                         .RNG.name="base::Super-Duper", .RNG.seed=413159),
                   list(alpha=inits.alpha[4], beta=inits.beta[4],
                         .RNG.name="base::Mersenne-Twister", .RNG.seed=143915))
## Compile JAGS model
model.jags <- jags.model(</pre>
  file = "LogitModel.txt",
  data = dat.jags,
  inits = inits.jags,
  n.chains = 4,
  n.adapt = 4000
```

```
## Compiling model graph
## Resolving undeclared variables
## Allocating nodes
```

```
## Graph information:
##
      Observed stochastic nodes: 6
##
      Unobserved stochastic nodes: 2
##
      Total graph size: 37
## Initializing model
## Burn-in
update(model.jags, n.iter = 4000)
## Sampling
fit.bayesian <- coda.samples(</pre>
  model = model.jags,
  variable.names = c("alpha", "beta"),
  n.iter = 30000,
  thin = 3
)
```

```
m.fit.bayesian <-as.matrix(fit.bayesian)</pre>
d.chains <- data.frame(</pre>
  iterations = rep(seq(8003, 38000, by=3), times=4),
  alpha = m.fit.bayesian[, "alpha"],
  beta = m.fit.bayesian[, "beta"],
  chains = rep(c("chain1", "chain2", "chain3", "chain4"), each=10000)
ggplot(d.chains, aes(x=iterations, y=alpha, color=chains)) + geom_line(alpha=0.5) +
  labs(title="Trace of alpha", x="Iterations") + theme_minimal()
ggplot(d.chains, aes(x=alpha, y=..density..)) +
  geom_density(color="darkblue", fill="lightblue", alpha=0.5) +
  labs(title="Density of alpha", y="Density") + theme_minimal()
{\tt ggplot(d.chains, aes(x=iterations, y=beta, color=chains)) + geom\_line(alpha=0.5) +}
  labs(title="Trace of beta", x="Iterations") + theme_minimal()
ggplot(d.chains, aes(x=beta, y=..density..)) +
  geom_density(color="darkblue", fill="lightblue", alpha=0.5) +
  labs(title="Density of beta", y="Density") + theme_minimal()
```





```
d.summary <- t(rbind(
   colMeans(m.fit.bayesian),
   apply(m.fit.bayesian, 2, function(x) sd(x)),
   apply(m.fit.bayesian, 2, function(x) quantile(x, probs=c(0.025, 0.5, 0.975)))
))

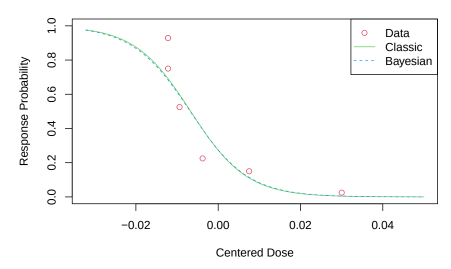
colnames(d.summary) <- c("Mean", "SD", "2.5%", "Median", "97.5%")
knitr::kable(d.summary, align="c", caption="Summary results for Bayesian approach")</pre>
```

Table 5: Summary results for Bayesian approach

	Mean	SD	2.5%	Median	97.5%
alpha	-0.971492	0.2318099	-1.448174	-0.9648464	-0.5344214
beta	-142.984631	24.4544352	-193.589700	-141.7772812	-98.2082827

```
## Inverse logit function
ilogit <- function(alpha, beta, x) {</pre>
  tmp <- exp(alpha + beta * x)</pre>
  pi <- tmp / (1 + tmp)
  return(pi)
## Extract estimates from classic and Bayesian models
alpha.classic <- coef(summary(fit.classic))[1, 1]</pre>
beta.classic <- coef(summary(fit.classic))[2, 1]</pre>
alpha.bayesian <- d.summary[1, 1]</pre>
beta.bayesian <- d.summary[2, 1]</pre>
x.grid <- seq(min(x centered)-0.02, max(x centered)+0.02, length.out=100)
y.pred.classic <- ilogit(alpha=alpha.classic, beta=beta.classic, x=x.grid)
y.pred.bayesian <- ilogit(alpha=alpha.bayesian, beta=beta.bayesian, x=x.grid)
plot(y.pred.classic ~ x.grid, col=3, type="l", ylim=c(0, 1), xlab="Centered Dose",
     ylab="Response Probability", main="Logistic curves with aggregate data")
lines(y.pred.bayesian ~ x.grid, col=4, lty=2)
points(x=x_centered, y=y/n, col=2)
legend("topright", legend=c("Data", "Classic", "Bayesian"),
       col=2:4, lty=c(NA, 1, 2), pch=c(1, NA, NA))
```

Logistic curves with aggregate data



Exercise 5 - CODA for logistic regression in JAGS



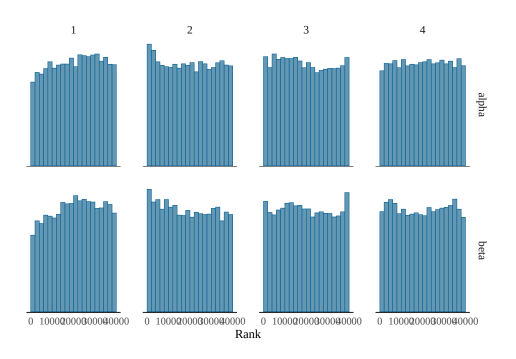


Figure 1: Rank Plot.

The figure illustrates the histograms of the ranked posterior drawn, ranked over all chains for α and β . Whereas traditional trace plots visualize how the chains mix over the course of sampling, rank histograms

visualize how the values from the chains mix together in terms of ranking. An ideal plot would show the rankings mixing or overlapping in a uniform distribution. See Vehtari et al. (2019) for details.

It can be seen in the figure that the rank histograms nearly follow a normal distribution.

Convergence diagnostics

Heidelberger & Welch (Convergence to stationarity)

heidel.diag(fit.bayesian)

```
## [[1]]
##
##
         Stationarity start
                                 p-value
##
         test
                      iteration
## alpha passed
                      1
                                 0.418
## beta passed
                                 0.331
                      1
##
##
         Halfwidth Mean
                           Halfwidth
##
         test
                     -0.96 0.0202
## alpha passed
## beta passed
                   -141.40 2.6148
##
## [[2]]
##
##
         Stationarity start
                                 p-value
##
         test
                      iteration
                                 0.239
## alpha passed
                      1
                      1
                                 0.353
## beta passed
##
##
         Halfwidth Mean
                            Halfwidth
##
         test
## alpha passed
                     -0.979 0.0224
## beta passed
                   -144.290 2.9099
##
## [[3]]
##
##
         Stationarity start
                                 p-value
##
         test
                      iteration
                                 0.908
## alpha passed
                      1
## beta passed
                                 0.931
##
##
         Halfwidth Mean
                            Halfwidth
##
         test
                     -0.977 0.0211
## alpha passed
## beta passed
                   -143.131 2.9174
##
## [[4]]
##
                                 p-value
##
         Stationarity start
##
         test
                      iteration
                                 0.674
## alpha passed
                      1
## beta passed
                                 0.772
##
##
         Halfwidth Mean
                           Halfwidth
```

##

test

```
## alpha passed -0.97 0.0214
## beta passed -143.12 2.7860
```

The output from heidel.diag shows the summary results for the four generated chains. The diagnostic on one hand checks if the length of the sample is long enough and on the other hand checks if the means are estimated from a converged chain.

The stationarity test uses the Cramer-von-Mises statistic to test the null hypothesis that the sampled values come from a stationary distribution. The half-width test calculates a 95% confidence interval for the mean, using the portion of the chain which passed the stationarity test.

We see that both tests are passed for the four chains and all p-values are larger than $0.05 = \alpha$. We conclude that the sampled values come from a stationary distribution and the length of the sample is considered long enough to estimate the mean with sufficient accuracy.

Raftery & Lewis (Convergence to ergodic average)

raftery.diag(fit.bayesian)

##

```
## [[1]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
##
          Burn-in Total Lower bound
                                        Dependence
##
          (M)
                    (N)
                           (Nmin)
                                        factor (I)
                                         15.1
##
    alpha 54
                   56421
                           3746
    beta 120
                   132465 3746
                                         35.4
##
##
##
## [[2]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
##
  Probability (s) = 0.95
##
##
          Burn-in Total Lower bound Dependence
##
                                        factor (I)
          (M)
                    (N)
                           (Nmin)
    alpha 90
                   99945 3746
                                        26.7
##
    beta 108
                   124650 3746
                                        33.3
##
##
## [[3]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
  Probability (s) = 0.95
##
##
          Burn-in Total Lower bound Dependence
##
          (M)
                    (N)
                           (Nmin)
                                        factor (I)
                   61911
##
    alpha 63
                           3746
                                        16.5
##
    beta 120
                   149892 3746
                                        40.0
##
##
## [[4]]
```

```
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
##
          Burn-in Total Lower bound
                                        Dependence
##
          (M)
                                        factor (I)
                    (N)
                           (Nmin)
                                        19.4
                          3746
##
    alpha 60
                   72732
    beta 105
                   126930 3746
                                        33.9
##
```

The output from raftery.diag shows the summary results for the four generated chains. raftery.diag is a run length control diagnostic based on a criterion of accuracy of estimation of the quantile q. The dependence factor I ($I = \frac{M+N}{N_{\min}}$) indicates to which extent autocorrelation inflates the required sample size. I > 5 indicates strong autocorrelation which may be due to a poor choice of starting value, high posterior correlations or "stickiness" of the MCMC algorithm. We see that the dependence factors for the four chains are all larger than 5 (depending on the random sample) and hence strong autocorrelation exists.

Geweke (Convergence to stationarity)

geweke.diag(fit.bayesian)

```
## [[1]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
##
   alpha
            beta
## 0.9601 0.5386
##
##
## [[2]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
## alpha beta
## 1.538 1.370
##
##
## [[3]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
##
   alpha
            beta
## 0.8589 1.4979
##
## [[4]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
## alpha
            beta
## 0.6512 0.6046
```

The output from <code>geweke.diag</code> shows the summary results for the four generated chains. <code>geweke.diag</code> is a convergence diagnostic for Markov chains based on a test for equality of the means of the first and last part of a Markov chain (by default the first 10% and the last 50%). If the samples are drawn from the stationary distribution of the chain, the two means are equal and Geweke's statistic has an asymptotically standard normal distribution. The test statistic is a standard Z-score.

The idea behind the Geweke's diagnostic is that in a long enough chain whose trace plots suggest convergence to the target distribution, we assume the second half of the chain has converged to the target distribution and we test if the first 10% can be treated as burn-in. So we mimic the simple two-sample test of means: if the mean of the first 10% is not significantly different from the last 50%, then we conclude the target distribution converged somewhere in the first 10% of the chain. So we'll use this 10% as burn-in.

We see that for the four chains the absolute values of Z-scores for variables alpha and beta are smaller than 2, which indicates that the equilibrium may have been reached.

geweke.plot(fit.bayesian)

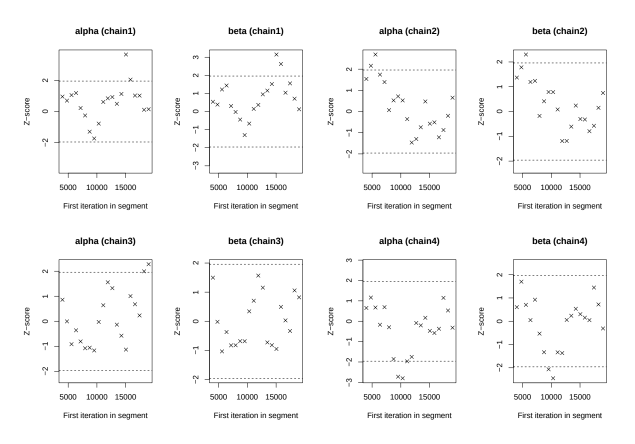


Figure 2: Geweke Plot.

If geweke.diag indicates that the first and last part of a sample from a Markov chain are not drawn from the same distribution, it may be useful to discard the first few iterations to see if the rest of the chain has "converged". geweke.plot shows what happens to Geweke's Z-score when successively larger numbers of iterations are discarded from the beginning of the chain. To preserve the asymptotic conditions required for Geweke's diagnostic, the plot never discards more than half the chain. For beta (chain1), alpha (chain2), and alpha (chain3), there exist several segments out of the 95% confidence bands (|Z| > 2). The plot shows that we mainly obtain z-score values below an absolute value of 2. So we can assume that the chain has converged.

Gelman and Rubin's convergence diagnostic

```
gelman.diag(fit.bayesian, autoburnin=TRUE)
```

```
## Potential scale reduction factors:
##
## Point est. Upper C.I.
## alpha 1 1.01
## beta 1 1.01
##
## Multivariate psrf
##
## 1
```

The idea of Gelman and Rubin's convergence diagnostic is to run multiple chains from widely dispersed starting values and perform an Analysis of Variance to see if the between-chain variability (B) is large in relation to the average variability within (W+B) the pooled chain.

$$\sqrt{\hat{R}} \approx \sqrt{\frac{W+B}{W}}$$

 \hat{R} is so-called "potential scale reduction factor" and it is calculated for each variable. We see that the "Potential scale reduction factors" for both alpha and beta are close to 1, which indicates that the between-chain variability (B) is close to 0. In other words, the separate four chains have mixed quite well.

```
gelman.plot(fit.bayesian, autoburnin=TRUE)
```

The gelman.plot shows the evolution of Gelman and Rubin's shrink factor as the number of iterations increases. We see that the shrink factors for both variables quickly decrease to 1 after 10000 iterations and the variability of shrink factors becomes more stable as the number of iterations keeps increasing.

Gelman-Rubin diagnostic using stable variance estimators

```
stable.GR(fit.bayesian)
```

```
## $psrf
## [1] 1.000276 1.000462
##
## $mpsrf
## [1] 1.000591
##
## $means
##
          alpha
##
     -0.9716766 -143.0065650
## $n.eff
## [1] 3121.102
##
## $blather
## [1] FALSE
```

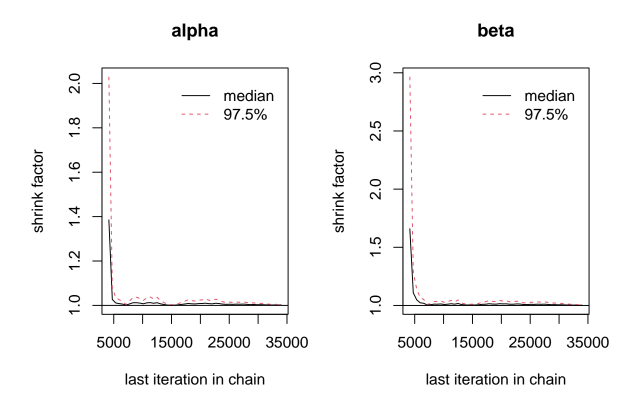


Figure 3: Gelman plot.

stable.GR extends Gelman-Rubin diagnostic using stable variance estimators. We see that the univariate potential scale reduction factors for both alpha and beta are calculated and they are close to 1. Multivariate psrf is also calculated by taking into account the interdependence of the chain's components. The PSRFs decrease to 1 as the chain length increases. When the PSRF becomes sufficiently close to 1, the sample collected by the Markov chain has converged to the target distribution. Means of variables (alpha and beta) and the effective sample size are also reported in the output.

Conclusions:

- Heidel: We conclude that the sampled values come from a stationary distribution and the length of the sample is considered long enough to estimate the mean with sufficient accuracy.
- Raftery & Lewis: The dependence factors for the four chains are larger than 5 and hence autocorrelation exists.
- Geweke: We assume that the equilibrium has been reached. We achieve convergence towards the target distribution.
- Gelman & Rubin: The between-chain variability is close to 0. The separate four chains have mixed quite well.
- $\bullet\,$ Gelman & Rubin for stable variance: The chains have converged to the target distribution.

All in all, it can be concluded that no issues arose during the diagnostics.

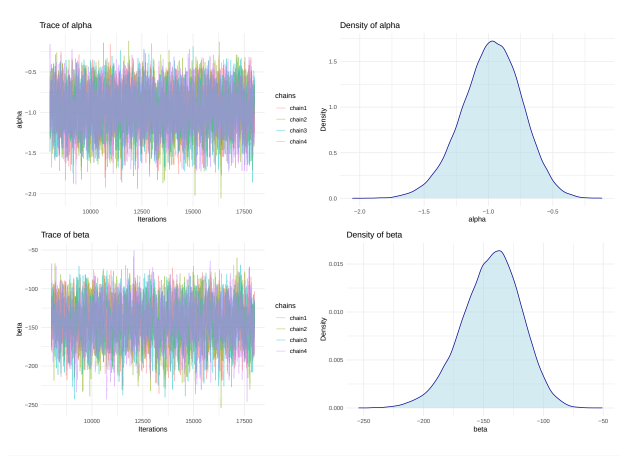
Re-run the MCMC simulation

In the following steps the MCMC simulation is adapted to the findings from sub task 5.2. We keep the same model as before but we change n.iter from 10'000 to 30'000 and n.thin from 1 to 3, to reduce

the autocorrelation. The number of burn-in iterations (n.burnin= 4000) is kept.

```
## Compile JAGS model
model2.jags <- jags.model(</pre>
  file = "LogitModel.txt",
  data = dat.jags,
 inits = inits.jags,
 n.chains = 4,
  n.adapt = 4000
)
## Compiling model graph
##
      Resolving undeclared variables
##
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 6
      Unobserved stochastic nodes: 2
##
##
      Total graph size: 37
##
## Initializing model
## Burn-in (increase burn-in iterations)
update(model2.jags, n.iter = 6000)
\mbox{\tt\#\#} 
 Initialize starting points (let JAGS initialize) and set seed
inits.jags <- list(list(.RNG.name="base::Wichmann-Hill", .RNG.seed=314159),</pre>
                   list(.RNG.name="base::Marsaglia-Multicarry", .RNG.seed=159314),
                   list(.RNG.name="base::Super-Duper", .RNG.seed=413159),
                   list(.RNG.name="base::Mersenne-Twister", .RNG.seed=143915))
## Sampling
fit2.jags.coda <- coda.samples(</pre>
  model = model2.jags,
 variable.names = c("alpha", "beta"),
 n.iter = 50000,
  thin = 5
)
summary(fit2.jags.coda)
##
## Iterations = 6005:56000
## Thinning interval = 5
## Number of chains = 4
## Sample size per chain = 10000
##
## 1. Empirical mean and standard deviation for each variable,
##
      plus standard error of the mean:
##
##
                        SD Naive SE Time-series SE
              Mean
          -0.9742 0.2327 0.001164
                                           0.00443
## alpha
## beta -143.4471 24.7007 0.123504
                                            0.57337
## 2. Quantiles for each variable:
##
                                  50%
                                            75%
##
             2.5%
                       25%
                                                   97.5%
                            -0.9673
                                       -0.8141 -0.5387
## alpha -1.454 -1.125
## beta -195.488 -159.414 -142.0939 -126.3137 -98.4751
```

```
m.fit2.jags.coda <-as.matrix(fit2.jags.coda)</pre>
d.chains <- data.frame(</pre>
  iterations = rep(8001:18000, times=4),
  alpha = m.fit2.jags.coda[, "alpha"],
  beta = m.fit2.jags.coda[, "beta"],
  chains = rep(c("chain1", "chain2", "chain3", "chain4"), each=10000),
  alphaRanks = rank(m.fit2.jags.coda[, "alpha"]),
  betaRanks = rank(m.fit2.jags.coda[, "beta"])
)
ggplot(d.chains, aes(x=iterations, y=alpha, color=chains)) + geom_line(alpha=0.5) +
  labs(title="Trace of alpha", x="Iterations") + theme_minimal()
ggplot(d.chains, aes(x=alpha, y=..density..)) +
  geom_density(color="darkblue", fill="lightblue", alpha=0.5) +
  labs(title="Density of alpha", y="Density") + theme_minimal()
ggplot(d.chains, aes(x=iterations, y=beta, color=chains)) + geom_line(alpha=0.5) +
  labs(title="Trace of beta", x="Iterations") + theme_minimal()
ggplot(d.chains, aes(x=beta, y=..density..)) +
  geom_density(color="darkblue", fill="lightblue", alpha=0.5) +
  labs(title="Density of beta", y="Density") + theme_minimal()
```



Compared to the rank plots obtained with the original model, the rank plots for β in the new model have a higher resamblance with a normal distribution. The rank plots of α are comparable.

mcmc_rank_hist(fit2.jags.coda)

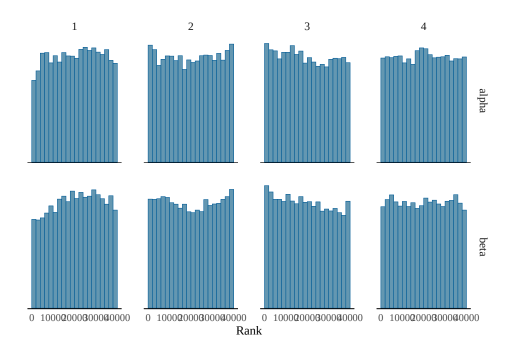


Figure 4: Rank Plot.

Heidelberger & Welch (Convergence to stationarity) heidel.diag(fit2.jags.coda)

```
## [[1]]
##
##
                            p-value
       Stationarity start
##
       test iteration
## alpha passed
                   1
                            0.238
## beta passed
                            0.145
##
##
       Halfwidth Mean
                       Halfwidth
##
       test
## alpha passed
                 -0.965 0.0155
## beta passed -142.054 2.0840
##
## [[2]]
##
##
        Stationarity start
                            p-value
##
        test
                   iteration
## alpha passed
                            0.745
                   1
## beta passed
                   1
                            0.579
##
##
                         Halfwidth
       Halfwidth Mean
##
       test
## alpha passed
                 -0.973 0.0184
## beta passed -143.460 2.3698
##
## [[3]]
##
##
       Stationarity start
                            p-value
##
       test iteration
## alpha passed
                   1
                            0.675
## beta passed
                  1
                            0.449
```

```
##
##
         Halfwidth Mean
                            Halfwidth
##
         test
## alpha passed
                     -0.984 0.0187
## beta passed
                   -144.788 2.2626
##
## [[4]]
##
                                p-value
##
         Stationarity start
##
         test
                      iteration
## alpha passed
                      1
                                0.577
## beta passed
                                0.816
##
##
         Halfwidth Mean
                            Halfwidth
##
         test
## alpha passed
                     -0.974 0.0168
## beta passed
                   -143.486 2.2647
```

The p-values of the Heidelberg diagnostics have slightly changed. But the interpretation stays the same above.

```
## Raftery & Lewis (Convergence to ergodic average)
raftery.diag(fit2.jags.coda)
```

```
## [[1]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
##
          Burn-in Total Lower bound Dependence
          (M)
##
                   (N)
                                       factor (I)
                          (Nmin)
##
   alpha 75
                   80250 3746
                                       21.4
   beta 120
                   129150 3746
                                       34.5
##
##
## [[2]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
##
          Burn-in Total Lower bound Dependence
##
                   (N)
                          (Nmin)
                                       factor (I)
          (M)
                                       23.9
##
   alpha 75
                   89400 3746
   beta 135
                   145920 3746
                                       39.0
##
##
##
## [[3]]
##
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
##
          Burn-in Total Lower bound Dependence
                                       factor (I)
##
          (M)
                   (N)
                          (Nmin)
## alpha 75
                   76845 3746
                                       20.5
## beta 135
                   140820 3746
                                       37.6
```

```
##
##
## [[4]]
## Quantile (q) = 0.025
## Accuracy (r) = +/- 0.005
## Probability (s) = 0.95
##
##
          Burn-in Total Lower bound Dependence
##
          (M)
                    (N)
                                        factor (I)
                           (Nmin)
##
    alpha 90
                    105900 3746
                                        28.3
    beta 150
                   147495 3746
                                        39.4
```

The values for the dependence factor I have increased, so we obtain an even stronger autocorrelation than before.

```
## Geweke (Convergence to stationarity)
geweke.diag(fit2.jags.coda)
```

```
## [[1]]
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
   alpha
            beta
## 0.6664 1.1324
##
##
## [[2]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
## alpha
            beta
## 1.3241 0.4805
##
##
## [[3]]
##
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
##
     alpha
              beta
## 0.01028 0.29911
##
##
## [[4]]
## Fraction in 1st window = 0.1
## Fraction in 2nd window = 0.5
##
##
     alpha
              beta
## -0.1841 -0.2226
```

The obtained absolute values for α and β are all below 2, which indicates convergence to the target distribution. The plot shows that some z-score points of the different chains lie above or below 2.0, but only a few which is acceptable.

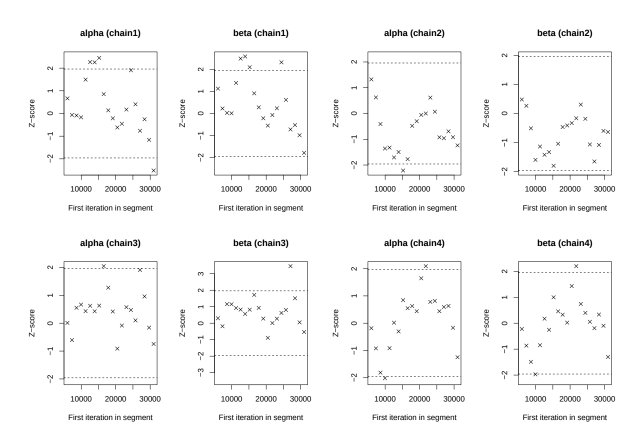


Figure 5: Geweke Plot.

```
## Gelman and Rubin's convergence diagnostic
gelman.diag(fit2.jags.coda, autoburnin=TRUE)
```

```
## Potential scale reduction factors:
##
## Point est. Upper C.I.
## alpha 1 1
## beta 1 1
##
## Multivariate psrf
##
## 1
```

The Gelman and Rubin diagnostic output is equivalent to the one from the original model.

```
gelman.plot(fit2.jags.coda, autoburnin=TRUE)
```

```
stable.GR(fit2.jags.coda)

## $psrf
## [1] 1.000152 1.000240
##
```

\$mpsrf

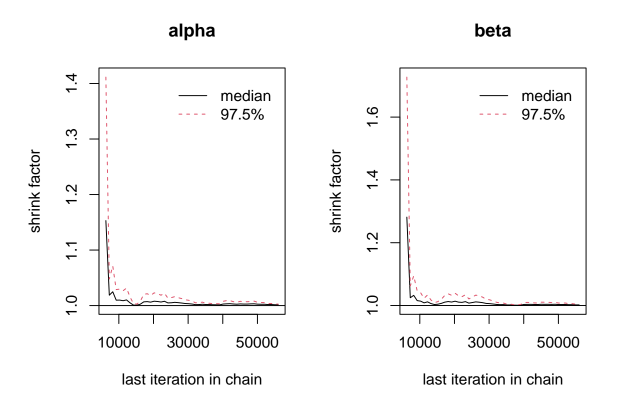


Figure 6: Gelman plot.

```
## [1] 1.00034
##
## $means
## alpha beta
## -0.9744474 -143.4753578
##
## $n.eff
## [1] 5129.801
##
## $blather
## [1] FALSE
```

The univariate potential scale reduction factors for both alpha and beta are close to 1. Compared to the original model the effective sample size has been increased.

Exercise 6 - EES

Run the code from the previous exercise with mice data with only one chain monitoring beta under the following two conditions:

1. After an adaptation phase of 1000 and a burn-in of 4000 draw a sample of 1000 observations in one chain with thinning set to 1.

```
## Initialize starting points (let JAGS initialize) and set seed
inits3.jags <- list(list(.RNG.name="base::Wichmann-Hill", .RNG.seed=314159))</pre>
```

```
## Compile JAGS model
model3.jags <- jags.model(</pre>
 file = "LogitModel.txt",
 data = dat.jags,
 n.chains = 1,
 inits = inits3.jags,
  n.adapt = 1000
## Compiling model graph
##
      Resolving undeclared variables
##
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 6
      Unobserved stochastic nodes: 2
##
##
      Total graph size: 37
## Initializing model
## Burn-in (increase burn-in iterations)
update(model3.jags, n.iter = 4000)
## Posterior Sampling
fit3.jags.coda <- coda.samples(</pre>
 model = model3.jags,
 variable.names = c("beta"), #monitoring only beta
 n.iter = 1000,
  thin = 1
)
```

2. After an adaptation phase of 1000 and a burn-in of 4000 draw a sample of 10000 observations in one chain with thinning set to 10.

```
## Initialize starting points (let JAGS initialize) and set seed
inits4.jags <- list(list(.RNG.name="base::Wichmann-Hill", .RNG.seed=314159))</pre>
## Compile JAGS model
model4.jags <- jags.model(</pre>
 file = "LogitModel.txt",
 data = dat.jags,
 n.chains = 1,
 inits = inits4.jags,
 n.adapt = 1000
)
## Compiling model graph
      Resolving undeclared variables
      Allocating nodes
## Graph information:
##
      Observed stochastic nodes: 6
##
      Unobserved stochastic nodes: 2
##
      Total graph size: 37
##
```

Initializing model

```
## Burn-in (increase burn-in iterations)
update(model4.jags, n.iter = 4000)

## Posterior Sampling
fit4.jags.coda <- coda.samples(
    model = model4.jags,
    variable.names = c("beta"), #monitoring only beta
    n.iter = 10000,
    thin = 10
)</pre>
```

(a) For which of the above conditions the ESS estimates will be larger and why?

```
# Function to test if a vector is monoton decreasing,
# a boolean value is returned
monotone <- function(vec){</pre>
 a <- TRUE
 if(length(vec) == 1){
   return(a)
 for(i in 2:length(vec)){
    if(vec[i] > vec[i-1]){
      a <- FALSE
      break;
    }
 }
 return(a)
}
# Function to find the lag to stop the ESS
# calculation compare:
#
#
       Geyer (1992),
#
      "Practical Markov Chain Monte Carlo".
#
       Statistical Science, 7: 473-511
\# Gamma_i = qamma(2*i) + qamma(2*i + 1)
# m is the greatest integer at which Gamma_i > 0
# and Gamma_i is monotone for i = 1, \ldots, m.
# Thereby gamma(i) is the sample autocorrelation
# at lag i.
# Parameter:
# vec - sample vector (mcmc object)
# Output
# m <- greatest integer where both criteria are
      fulfilled
geyer <- function(vec){</pre>
 g <- c()
 res <- 1
  for(i in 1:(length(vec)/2 - 1)){
    g \leftarrow c(g,
           autocorr(vec, lags = i) + autocorr(vec, lags = i + 1)
    if(monotone(g) == FALSE || g[i] < 0){</pre>
```

```
break
    }
  }
  if(i==1){
    res <- 1
  else{
    res <- i-1
  return(res)
}
# Function to calculate the effective sample
# size for one MCMC chain.
#
# Parameter:
# mcmc - mcmc object
# M - number of sampled values
# Output:
# effective sample size
ess <- function(mcmc, M){
  m <- geyer(mcmc)</pre>
  y \leftarrow M / (1 + 2 * sum(autocorr(mcmc, lag = 1:(2*m +1))))
  return(y)
}
```

$$ESS = N_{eff} = \frac{M}{1 + 2\sum_{k>1}^{\infty} ACF(k)}$$

where $\text{ACF}(k) = \text{corr}(\theta_t^*, \theta_{t+k}^*)$

For practical computation, the infinite sum in the definition of ESS may be stopped earlier. Here the stopping is defined by the criteria of Geyer 1992.

We have in the second model a thinning parameter of 10 and thus expect the autocorrelation in the MCMC-sample to have less correlation as we pick only the 10th entry each time and put it into our MCMC-sample. Thus the sum will be smaller for the less correlated sample (fit4.jags.coda). M is in both models the same with 1000. Therefore we expect the ESS to be larger in the second model!

(b) To check your answer: Apply both the 05ess.R code and the function effectiveSize from the coda package. Compare the ESS estimates with those obtained with the n.eff function from package stableGR (Vats and Knudson, 2021). Please report your findings.

Table 6: Effective sample size with different thinning and functions.

method	thinning	n_eff
ESS-Script	1	11.96032
ESS-Script	10	115.43639
stableGR	1	10.92997
stableGR	10	92.17642

As expected, the effective sample size is with both methods higher in the model with thinning = 10 compared to thinning = 1.