

# Bayesian statistics with R

## 6. Bayesian analyses in R with the Jags software

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April 2022

## Bayes in practice

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## Software implementation (R friendly)

Oldies but goodies:

- WinBUGS, OpenBUGS: Where it all began.
- Jags: What we will use in this course.

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If you're not into coding:

- brms: Bayesian regression models with Stan.
- MCMCglmm: Generalised Linear Mixed Models.
- Check out the CRAN Task View: Bayesian Inference for more.

# Introduction to JAGS (Just Another Gibbs Sampler)

Martyn Plummer



## Real example

Impact of climatic conditions on white stork breeding success



mangl.at

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- We have collected data.
- We need to build a model - write down the likelihood.
- We need to specify priors for parameters.

## Read in the data

```
nbchicks <- c(151,105,73,107,113,87,77,108,118,122,112,120,122,89,69,71,
              53,41,53,31,35,14,18)

nbpairs <- c(173,164,103,113,122,112,98,121,132,136,133,137,145,117,90,80,
             67,54,58,39,42,23,23)

temp <- c(15.1,13.3,15.3,13.3,14.6,15.6,13.1,13.1,15.0,11.7,15.3,14.4,14.4,
          12.7,11.7,11.9,15.9,13.4,14.0,13.9,12.9,15.1,13.0)

rain <- c(67,52,88,61,32,36,72,43,92,32,86,28,57,55,66,26,28,96,48,90,86,
          78,87)

datax <- list(N = 23,
              nbchicks = nbchicks
```

## Write down the model

$$\text{nbchicks}_i \sim \text{Binomial}(\text{nbpairs}_i, p_i) \quad [\text{likelihood}]$$

$$\text{logit}(p_i) = a + b_{\text{temp}} \text{temp}_i + b_{\text{rain}} \text{rain}_i \quad [\text{linear model}]$$

$$a \sim \text{Normal}(0, 1000) \quad [\text{prior for } a]$$

$$b_{\text{temp}} \sim \text{Normal}(0, 1000) \quad [\text{prior for } b_{\text{temp}}]$$

$$b_{\text{rain}} \sim \text{Normal}(0, 1000) \quad [\text{prior for } b_{\text{rain}}]$$

## Build the model

```
{  
# Likelihood  
  for( i in 1 : N){  
    nbchicks[i] ~ dbin(p[i],nbpairs[i])  
    logit(p[i]) <- a + b.temp * temp[i] + b.rain * rain[i]  
  }  
  
# ...
```

## Specify priors

```
# Priors  
a ~ dnorm(0,0.001)  
b.temp ~ dnorm(0,0.001)  
b.rain ~ dnorm(0,0.001)  
}
```

**Warning:** Jags uses precision for Normal distributions (1 / variance)

## You need to write everything in a file

```
model <-  
paste("  
model  
{  
  for( i in 1 : N)  
  {  
    nbchicks[i] ~ dbin(p[i],nbpairs[i])  
    logit(p[i]) <- a + b.temp * temp[i] + b.rain * rain[i]  
  }  
a ~ dnorm(0,0.001)  
b.temp ~ dnorm(0,0.001)  
b.rain ~ dnorm(0,0.001)  
}  
")
```



## Alternatively, you may write a R function

```
logistic <- function() {  
  for( i in 1 : N)  
  {  
    nbchicks[i] ~ dbin(p[i],nbpairs[i])  
    logit(p[i]) <- a + b.temp * temp[i] + b.rain * rain[i]  
  }  
  
  # priors for regression parameters  
  a ~ dnorm(0,0.001)  
  b.temp ~ dnorm(0,0.001)  
  b.rain ~ dnorm(0,0.001)  
}
```

## Let us specify a few additional things

```
# list of lists of initial values (one for each MCMC chain)
```

```
init1 <- list(a = -0.5, b.temp = -0.5, b.rain = -0.5)
```

```
init2 <- list(a = 0.5, b.temp = 0.5, b.rain = 0.5)
```

```
inits <- list(init1,init2)
```

```
# specify parameters that need to be estimated
```

```
parameters <- c("a","b.temp","b.rain")
```

```
# specify nb iterations for burn-in and final inference
```

```
nb.burnin <- 1000
```

```
nb.iterations <- 2000 # beware: nb.iterations includes nb.burnin!
```

## Run Jags

```
# load R2jags
library(R2jags)
# run Jags
storks <- jags(data = datax,
               inits = inits,
               parameters.to.save = parameters,
               #model.file = "code/logistic.txt",
               model.file = logistic, # if a function was written
               n.chains = 2,
               n.iter = nb.iterations,
               n.burnin = nb.burnin)

storks
```

## Inspect parameter estimates

```
#> Compiling model graph
#>   Resolving undeclared variables
#>   Allocating nodes
#> Graph information:
#>   Observed stochastic nodes: 23
#>   Unobserved stochastic nodes: 3
#>   Total graph size: 181
#>
#> Initializing model
#> Inference for Bugs model at "code/logistic.txt", fit using jags,
#> 2 chains, each with 2000 iterations (first 1000 discarded)
#> n.sims = 2000 iterations saved
```

#>	mu.vect	sd.vect	2.5%	25%	50%	75%	97.5%	Rhat	n.eff
#> a	1.545	0.081	1.441	1.514	1.549	1.585	1.648	1.194	110
#> b.rain	-0.162	0.063	-0.274	-0.203	-0.163	-0.123	-0.041	1.031	560
#> b.temp	0.026	0.059	-0.096	-0.009	0.027	0.063	0.144	1.003	1600
#> deviance	206.200	30.419	201.803	202.664	203.739	205.378	210.718	1.082	2000

## Your turn: Practical 5

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## Assess convergence

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## Reminder – MCMC Algorithm

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- For the MCMC algorithm, the posterior distribution is only needed to be known up to proportionality.
- Once the stationary distribution is reached we can regard the realisations of the chain as a (dependent) sample from the posterior distribution (and obtain Monte Carlo estimates).
- We consider some important implementation issues.

## MCMC – Proposal Distribution

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- To implement a MCMC algorithm, we often need to specify a proposal distribution from which we generate candidate value then accept/reject.
- This typically involves
  - specifying a given distribution family (e.g. normal, uniform), and then,
  - setting the parameters of the given distribution.
- Although the exact distribution specified is essentially arbitrary – it will have a significant effect on the performance of the MCMC algorithm.

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- In order to balance the size of the proposed moves with the chance of accepting them the proposal variance is often tuned to obtain a mean acceptance probability of 20 – 40%.



## Why is the proposal distribution so important?

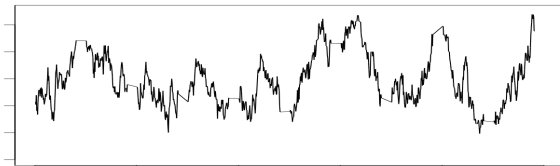
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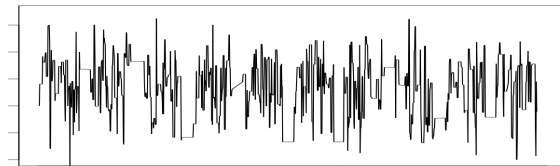
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- Automatic in Jags – ouf!
- The movement around the parameter space is often referred to as **mixing**.

## Good/Bad Traces

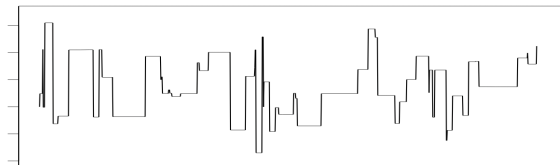
*Small  
moves -  
bad*



*good*



*Large  
moves -  
bad*



0

500

1000

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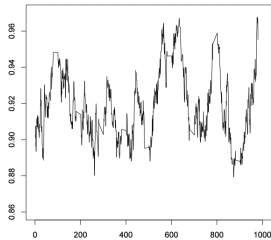
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- Autocorrelation function (ACF) plots are a convenient way of displaying the strength of autocorrelation in the given sample values.

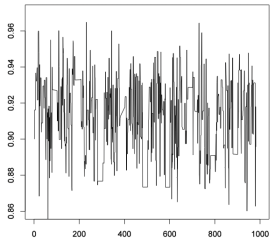
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- Autocorrelation function (ACF) plots are a convenient way of displaying the strength of autocorrelation in the given sample values.
- ACF plots provide the autocorrelation between successively sampled values separated by  $k$  iterations, referred to as lag, (i.e.  $\text{cor}(\theta_t, \theta_{t+k})$ ) for increasing values of  $k$ .

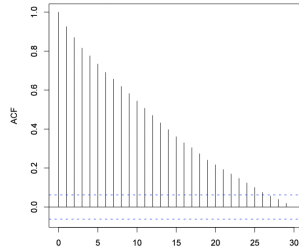
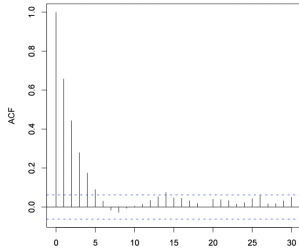
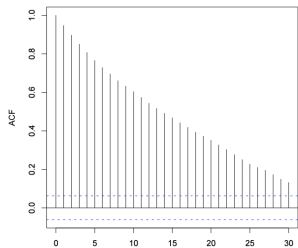
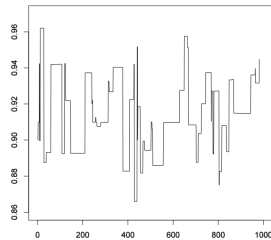
*Small moves*



*OK*



*Big moves*

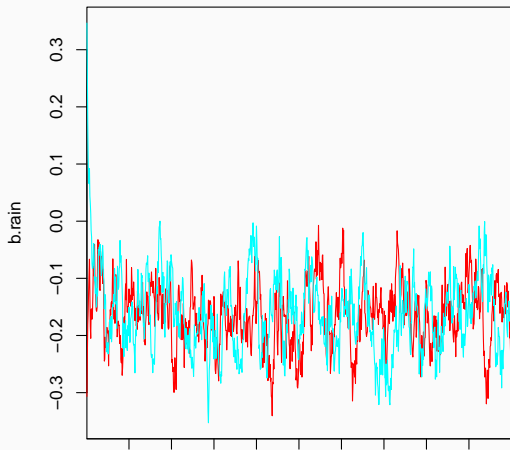




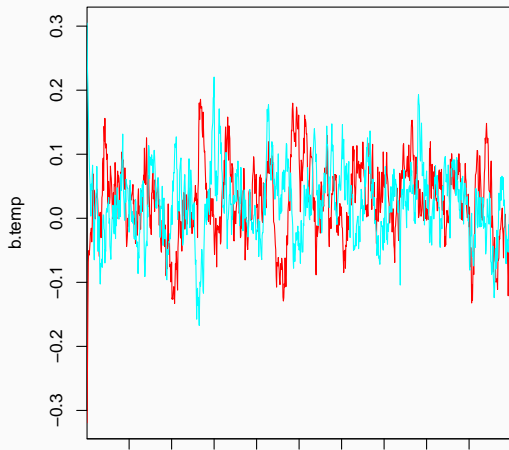
## Traceplots for the storks

```
traceplot(storks, mfrow = c(1, 2), varname = c('b.rain', 'b.temp'), ask = FALSE)
```

**b.rain**

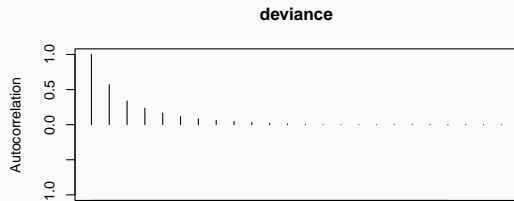
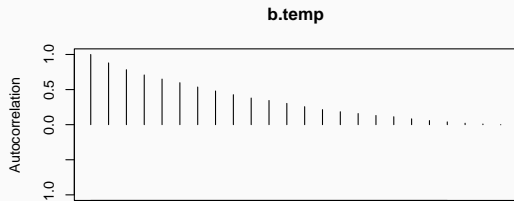
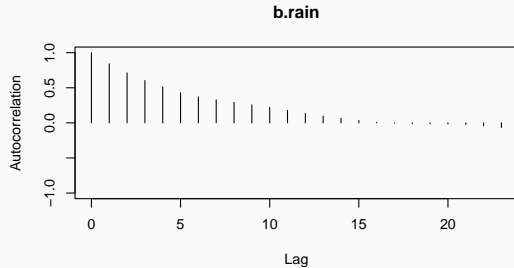
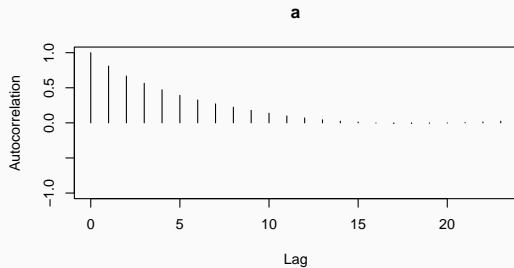


**b.temp**



# Autocorrelation for the storks

```
autocorr.plot(as.mcmc(storks), ask = FALSE)
```



## How do good chains behave?

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- Once there, explore efficiently: The post-convergence sample size required for suitable numerical summaries.

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- In practice, we must discard observations from the start of the chain and just use observations from the chain once it has converged.
- The initial observations that we discard are referred to as the **burn-in**.
- The simplest method to determine the length of the burn-in period is to look at trace plots.

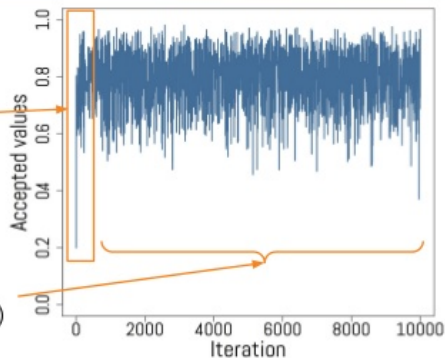


## Burn-in (if simulations cheap, be conservative)

Discard initial guesses that are still far from optimum: the

**BURN-IN**

These numbers should be a good  
**sample of the Posterior**  $P(\phi \mid \text{data})$



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- The effective sample size ( $n_{\text{eff}}$ ) measures chain length while taking into account the autocorrelation of the chain.
  - $n_{\text{eff}}$  is less than the number of MCMC iterations.
  - Check the  $n_{\text{eff}}$  of every parameter of interest.
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  - Check the  $n_{\text{eff}}$  of any interesting parameter combinations.
- We need  $n_{\text{eff}} \geq 100$  independent steps.

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## Potential scale reduction factor

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- Measures the ratio of the total variability combining multiple chains (between-chain plus within-chain) to the within-chain variability. Asks the question is there a chain effect? Very much alike the  $F$  test in an ANOVA.



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- Values near 1 indicates likely convergence, a value of  $\leq 1.1$  is considered acceptable.
- Necessary condition, not sufficient; In other words, these diagnostics cannot tell you that you have converged for sure, only that you have not.

## n.eff and $\hat{R}$ for the storks

storks

```
#> Inference for Bugs model at "code/logistic.txt", fit using jags,  
#> 2 chains, each with 2000 iterations (first 1000 discarded)  
#> n.sims = 2000 iterations saved  
#>          mu.vect sd.vect   2.5%   25%   50%   75%   97.5% Rhat  
#> a          1.545   0.081   1.441   1.514   1.549   1.585   1.648 1.194  
#> b.rain     -0.162   0.063  -0.274  -0.203  -0.163  -0.123  -0.041 1.031  
#> b.temp      0.026   0.059  -0.096  -0.009   0.027   0.063   0.144 1.003  
#> deviance 206.200  30.419 201.803 202.664 203.739 205.378 210.718 1.082  
#>  
#> For each parameter, n.eff is a crude measure of effective sample size,  
#> and Rhat is the potential scale reduction factor (at convergence, Rhat=  
#>  
#> DIC info (using the rule,  $pD = \text{var}(\text{deviance})/2$ )
```

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- Run multiple chains from arbitrary starting places (initial values).
- Assume convergence when all chains reach same regime.
- Discard initial burn-in phase.
- Check autocorrelation, effective sample size and  $\hat{R}$ .

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- Increase burn-in, sample more.



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- Reparameterize:
  - Standardize covariates.
  - Non-centering:  $\alpha \sim N(0, \sigma)$  becomes  $\alpha = z\sigma$  with  $z \sim N(0, 1)$ .

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- Something wrong with your model?
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  - Use simulations.
- Change your sampler. Upgrade to Nimble or Stan.

**MCMC makes you queens and kings  
of the stats world**

---

## Get all values sampled from posteriors

```
res <- as.mcmc(storks) # convert outputs in a list
res <- rbind(res[[1]],res[[2]]) # put two MCMC lists on top of each other
head(res)
#>           a      b.rain      b.temp  deviance
#> [1,] 0.07080353 -0.30662224 -0.31932056 1225.8420
#> [2,] 0.41577870 -0.23901959 -0.23508941  773.3837
#> [3,] 0.68847213 -0.18098311 -0.14169900  509.4181
#> [4,] 0.80746015 -0.15732435 -0.06811845  418.3528
#> [5,] 0.92786687 -0.12241767 -0.05058431  351.4461
#> [6,] 1.03149945 -0.09174747 -0.05743429  307.0004
tail(res)
#>           a      b.rain      b.temp  deviance
#> [1995,] 1.544116 -0.1747998 0.007950735 201.9061
#> [1996,] 1.532711 -0.1635601 0.022560051 201.7395
```

## Compute a posteriori $\Pr(\text{rain} < 0)$

```
# probability that the effect of rainfall is negative  
mean(res[, 'b.rain'] < 0)  
#> [1] 0.9945
```

## Compute a posteriori $\Pr(\text{temp} < 0)$

```
# probability that the effect of temperature is negative  
mean(res[, 'b.temp'] < 0)  
#> [1] 0.315
```



## Get credible interval for the rain effect

```
quantile(res[, 'b.rain'], c(0.025, 0.975))
```

```
#>           2.5%           97.5%
```

```
#> -0.27437161 -0.04117215
```

## Get credible interval for the temperature effect

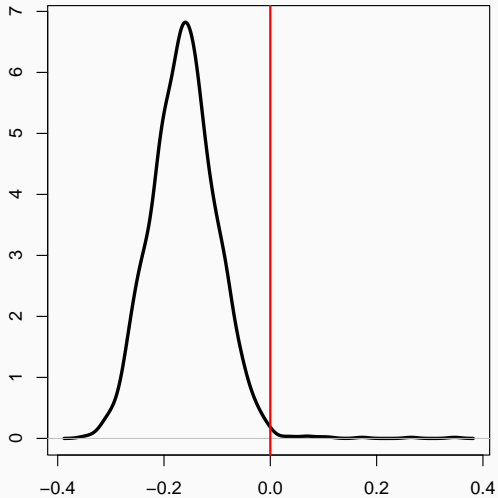
```
quantile(res[, 'b.temp'], c(0.025, 0.975))
```

```
#>           2.5%           97.5%
```

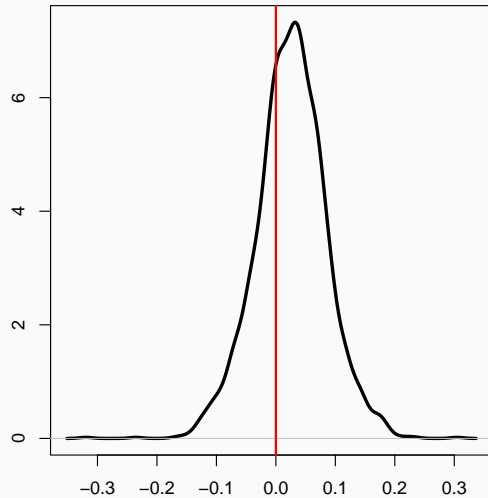
```
#> -0.09607297  0.14354530
```

# Graphical summaries

Rainfall



Temperature



## **Your turn: Practical 6**

---