Translating Probability Density Functions: From R to BUGS and Back Again

by David S. LeBauer, Michael C. Dietze, Benjamin M. Bolker

Abstract The ability to implement statistical models in the BUGS language facilitates Bayesian inference by automating MCMC algorithms. Software packages that interpret the BUGS language include OpenBUGS, WinBUGS, and JAGS. R packages that link BUGS software to the R environment, including rjags and R2WinBUGS, are widely used in Bayesian analysis. Indeed, many packages in the Bayesian task view on CRAN (http://cran.r-project.org/web/views/Bayesian.html) depend on this integration. However, the R and BUGS languages use different representations of common probability density functions, creating a potential for errors to occur in the implementation or interpretation of analyses that use both languages. Here we review different parameterizations used by the R and BUGS languages, describe how to translate between the languages, and provide an R function, r2bugs.distributions, that transforms parameterizations from R to BUGS and back again.

| Distribution | Lang. | Parameterization | Use | Notes |
|-------------------|-----------|--|---|---|
| Normal | | , | | |
| | R | $\frac{1}{\sqrt{2\pi}\sigma}\exp\left(-\frac{(x-\mu)^2}{2\sigma^2}\right)$ | $dnorm(x, mean = \mu, sd = \sigma)$ | |
| | BUGS | $\sqrt{\frac{\tau}{2\pi}}\exp\left(-\left(x-\mu\right)^2\tau\right)$ | dnorm(mean = μ , precision = τ) | $	au = \left(\frac{1}{\sigma}\right)^2$ |
| log-Normal | | | | |
| | R | $\frac{1}{\sqrt{2\pi}\sigma x} \exp\left(-\frac{(\log(x)-\mu)^2}{(2\sigma^2)}\right)$ | $dlnorm(x, mean = \mu, sd = \sigma)$ | |
| | BUGS | $\frac{\sqrt{\tau}}{x} \exp\left(\frac{-\tau(\log(x)-\mu)^2}{2}\right)$ | dlnorm(mean = μ , precision = τ) | $	au = \left(rac{1}{\sigma} ight)^2$ |
| Binomial | | | | reverse parameter order |
| | R BUGS | $\binom{n}{x}p^x (1-p)^{n-x}$ same | dbinom(x, size = n, prob = p) dbin(prob = p, size = n) | |
| Negative Binomial | | | | reverse parameter order |
| | R | $\frac{\Gamma(x+n)}{\Gamma(n)x!}p^n(1-p)^x$ | dnbinom(x, size = n, prob = p) | size (n) is continuous |
| | BUGS | $\binom{x+r-1}{x}p^r(1-p)^x$ | dnegbin(prob = p, size = r) | size (r) is discrete |
| Weibull | | | | |
| | R | $\frac{a}{b}(\frac{x}{b})^{a-1}\exp\left(-\left(\frac{x}{b}\right)^a\right)$ | dweibull(x, shape = a, scale = b) | |
| | BUGS | $\nu\lambda x^{\nu-1}\exp\left(-\lambda x^{\nu}\right)$ | $dweib(shape = \nu, \; lambda = \lambda)$ | $\lambda = \left(\frac{1}{b}\right)^a$ |
| Gamma | | | | reverse parameter order |
| | R | $\frac{r^a}{\Gamma(a)} x^{a-1} \exp(-xr)$ $\frac{\lambda^r x^{r-1} \exp(-\lambda x)}{\Gamma(r)}$ | dgamma(x, shape = a, rate = r) | |
| | BUGS | $\frac{\lambda^r x^{r-1} \exp(-\lambda x)}{\Gamma(r)}$ | $dgamma(shape = r, lambda = \lambda)$ | |

Table 1: Summary of different parameterizations of common distributions used by R and BUGS. **Note:** For ease of reference, parameterizations follow the JAGS and R documentation; as a result, the table includes equivalent equations that appear different, either because JAGS and R use different names for the same parameter or because the equation has been rearranged. For example, the shape parameter of the *Gamma* distribution is *r* in the BUGS documentation and *a* in the R documentation. For the *Binomial*, *Negative Binomial*, and *Gamma* distributions, BUGS and R expect parameters in different order (the order of parameters matters since arguments are assigned based on position in BUGS and may be in R as well). R allows alternate parameterizations for the *Negative Binomial* and *Gamma* distributions, but these are not shown here. The variable *x* is implicit in all of the BUGS "Use" expressions. The *Beta*, *Poisson*, *Exponential*, and *Uniform* distributions have identical parameterizations in R and BUGS.

Probability density functions in R and BUGS

R and BUGS implement many of the same probability distribution functions, but they often parameterize the same distribution differently (Table 1). Although these probability distribution functions are clearly described in the documentation of their respective languages, we were unable to find a summary of these differences in one place. The motivation for this article is to document and clarify these differences. Our sources are the JAGS documentation (Plummer, 2010) and the documentation of individual R functions.

A bilingual translation function

To support the automation of model specification in JAGS with priors computed and stored in R (LeBauer et al., 2013), we developed a function to translate parameterizations of common probability distributions from R to BUGS (and back again, by specifying direction = 'bugs2r'). Parameter transformations, parameter order, and differences in function names are documented in Table 1 and implemented in the R function r2bugs.distributions.

```
r2bugs.distributions <- function(priors, direction = 'r2bugs') {</pre>
 priors$distn <- as.character(priors$distn)</pre>
 priors$parama <- as.numeric(priors$parama)</pre>
 priors$paramb <- as.numeric(priors$paramb)</pre>
 ## index dataframe according to distribution
        <- priors$distn %in% c('norm', 'lnorm')</pre>
                                                        # these have same transform
         <- grepl("weib", priors$distn)</pre>
                                                        # matches r and bugs version
 gamma <- priors$distn == 'gamma'</pre>
        <- grepl("chisq", priors$distn)
 chsq
                                                        # matches r and bugs version
         <- priors$distn %in% c('binom', 'bin')</pre>
                                                       # matches r and bugs version
 bin
        <- priors$distn %in% c('nbinom', 'negbin') # matches r and bugs version</pre>
 nbin
 ## Normal, log-Normal: Convert sd to precision
 exponent <- ifelse(direction == "r2bugs", -2, -0.5)
 priors$paramb[norm] <- priors$paramb[norm] ^ exponent</pre>
 ## Weibull
 if(direction == 'r2bugs'){
    ## Convert R parameter b to BUGS parameter lambda by l = (1/b)^a
    priors$paramb[weib] <- (1 / priors$paramb[weib]) ^ priors$parama[weib]</pre>
  } else if (direction == 'bugs2r') {
    ## Convert BUGS parameter lambda to BUGS parameter b by b = 1^{(-1/a)}
    priors$paramb[weib] <- priors$paramb[weib] ^ (- 1 / priors$parama[weib] )</pre>
 }
 ## Reverse parameter order for binomial and negative binomial
 priors[bin | nbin, c('parama', 'paramb')] <-</pre>
    priors[bin | nbin, c('paramb', 'parama')]
 ## Translate distribution names
 if(direction == "r2bugs"){
    priors$distn[weib] <- "weib"</pre>
    priors$distn[chsq] <- "chisqr"</pre>
    priors$distn[bin] <- "bin"
    priors$distn[nbin] <- "negbin"</pre>
 } else if(direction == "bugs2r"){
    priors$distn[weib] <- "weibull"</pre>
    priors$distn[chsq] <- "chisq"</pre>
    priors$distn[bin] <- "binom"</pre>
    priors$distn[nbin] <- "nbinom"</pre>
 return(priors)
```

A simple example

As an example, we take the R-parameterized prior distribution $X \sim \mathcal{N}(\mu = 10, \sigma = 2)$ and convert it to BUGS parameterization $X \sim \mathcal{N}(\mu = 10, \tau = 1/4)$. We specify a model in JAGS that allows us to sample directly from a prior distribution. The function works for each of the distributions in Table 1. This particular example is the JAGS implementation of 'rnorm(10000, 10, 2)' in R. It is presented as minimal demonstration; for a non-trivial application, see LeBauer et al. (2013).

```
r.distn <- data.frame(distn = "norm", parama = 10, paramb = 2)</pre>
bugs.distn <- r2bugs.distributions(r.distn)</pre>
sample.bugs.distn <- function(prior = data.frame(distn = "norm", parama = 0,</pre>
                                 paramb = 1), n = 10000) {
 require(rjags)
 model.string <- paste0(</pre>
    "model{Y ~ d", prior$distn,
    "(", prior$parama,
    ## chisqr has only one parameter
    ifelse(prior$distn == "chisqr", "", paste0(", ", prior$paramb)), ");",
    a <- x
 ## trick JAGS into running without data
 writeLines(model.string, con = "test.bug")
 j.model <- jags.model(file = "test.bug", data = list(x = 1))</pre>
 mcmc.object <- window(</pre>
    coda.samples(
      model = j.model, variable.names = c('Y'),
      n.iter = n * 4, thin = 2),
    start = n
 Y <- sample(as.matrix(mcmc.object)[,"Y"], n)
}
X <- sample.bugs.distn(bugs.distn)</pre>
```

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Bibliography

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David S. LeBauer Energy Biosciences Institute University of Illinois USA dlebauer@illinois.edu Benjamin M. Bolker Department of Mathematics & Statistics McMaster University Canada

Michael C. Dietze Department of Earth And Environment Boston University USA