

Open Source Software Tools for Parallel Computation of Multiple MCMC Chains with WinBUGS

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BACKGROUND

- When used for a Bayesian analysis, Markov chain Monte Carlo (MCMC) simulations generate samples that approximate the joint posterior distribution of the sampled parameters.
- The sequential nature of MCMC simulation limits the benefits of parallel computation when applied to a single chain.
- Parallel computation of multiple chains, however, is an "embarrassingly parallel" problem that can substantially reduce computation time, and is relatively easy to implement using freely available software.

OBJECTIVE

- To develop open source software tools for parallel computation of multiple MCMC chains with WinBUGS [1] running in Microsoft Windows.

METHODS

- `bugsParallel` is a set of R [2] functions for distributed computing with WinBUGS 1.4.*.
 - It is a modified subset of the R2WinBUGS package [3].
 - Uses the R package `Rmpi` [4] to implement a network of Windows workstations.
 - Uses the `rlecuyer` package [5] for parallel random number generation.
 - This assures appropriately independent initial estimates when they are specified via a function that uses random number generation to generate the initial estimates.
 - The seeds used by each WinBUGS chain are generated in `bugsParallel` by the master node. The master generates a vector of random seeds – one per chain. The vector and chain identifier are passed to each slave that uses the appropriate element of that vector. The seed is passed to WinBUGS via the undocumented `set.seed` WinBUGS script command.
- Implementation involves installation of the following freely distributed software components:
 - MPICH2 for Windows [6], an implementation of the Message-Passing Interface (MPI) for parallel computation,
 - WinBUGS 1.4.3 [1],
 - R for Windows [2],
 - R packages `Rmpi` [4] and `rlecuyer` [5].
- `bugsParallel` is currently provided as an R script that the user accesses via a "source" command in a user written R script.
- Parallel computation with WinBUGS was previously reported by Gigris et al [7] using commercial computing grid software.

Illustrative Example

- The following R script and WinBUGS model were used to fit a sigmoid Emax model to a set of 125 simulated data points.
- The example was run on a ThinkMate Workstation with two Intel E5345 Quad-Core processors (2.33 GHz) with 16 GB RAM.
- Eight MCMC chains of 100,000 iterations each were computed.
 - The first 10,000 iterations were discarded.
 - The MCMC chains were thinned to retain 1 of each 100 iterations, i.e., 7200 MCMC samples remained for analysis.

Equation 1. Bayesian Model from Illustrative Example

$$E_i \sim N(\hat{E}_i, \sigma^2)$$

$$\hat{E}_i = \frac{E_{max} c_i^\gamma}{EC_{50}^\gamma + c_i^\gamma}$$

Where E_i is the "observed response."

Equation 2. Prior distributions:

$$E_{max} \sim U(0, 100)$$

$$\log(EC_{50}) \sim N(0, 10^6)$$

$$\gamma \sim U(0, 10)$$

$$\sigma \sim U(0, 1000)$$

Figure 1. R script from illustrative example

```
model.name = "FxExample1" #root name of modeling files
toolsDir = "C:/bugsTools"

library(Rmpi)
library(rlecuyer)
library(coda)
library(lattice)
source(paste(toolsDir,"/bglillespie.utilities.R",sep=""))
source(paste(toolsDir,"/bugsParallelAll.R",sep=""))
source(paste(toolsDir,"/bugs.tools.R",sep="")) # A few BUGS-specific utilities

#####
# Data management

xdata = read.csv("fxa.data.avg.csv")

#create WinBUGS data set
bugsdata = list(
  nobs = nrow(xdata),
  cobs = xdata$cavg,
  FXa = xdata$fxa.inh.avg)

#create initial estimates
bugsinit = function()
{
  Emax = runif(1,0,100),
  logEC50 = rnorm(1,log(100),0.4),
  gamma = 10*rbeta(1,0.25,5),
  sigma = exp(rnorm(1,log(5),0.2)))
}

# specify what variables to monitor
parameters = c("Emax","EC50","gamma","sigma","FxPred")

# specify the variables for which you want history and density plots
parameters.to.plot = c("deviance","Emax","EC50","gamma","sigma")

#####
# run WinBUGS

n.chains = 4
n.iter = 100000
n.burnin = 10000
n.thin = 100

mpi.spawn.Rslaves() # launched multiple R slave processes on available processors
mpi.set.rngstream() # initializes parallel random number generation

system.time(bugs.fit <- bugsParallel(data=bugsdata,init=bugsinit,
  parameters.to.covar=parameters,model.file=paste(getwd(),"\\" ,model.name,
  ".txt", sep=""),n.chains=n.chains,n.iter=n.iter,n.thin=n.thin,
  n.burnin=n.burnin,refresh=1,clearWD=TRUE,bugs.directory =
  "c:/Program Files/WinBUGS14/"))

# save scripts, data and results to a directory
save.model(bugs.fit,model.name)
```

Figure 1. R Script from Illustrative Example, Continued

```
#convert MCMC to formats suitable for post-processing
sims.array = apem(array(unlist(bugs.fit),dim=c(nrow(bugs.fit)[1]),
  ncol(bugs.fit[1,]),length(bugs.fit)),dimnames=c(dimnames(bugs.fit)[1]),
  list(NULL)),c(1,3,2))
posterior = array(as.vector(sims.array),dim=c(prod(dim(sims.array)[1:2]),
  dim(sims.array)[3]),dimnames=list(NULL,dimnames(sims.array)[1:3]))
```

#####
posterior distributions of parameters

```
# open graphics device
pdf(file = paste(model.name,"/",model.name,".plots.pdf",sep=""),width=6,
height=6)

# create history, density and Gelman-Rubin-Brooks plots, and a table of
summary stats
ptable = parameter.plot.table(sims.array[,,
  unlist(sapply(c(paste("##",parameters,to.plot,"$","sep="),
  paste("##",parameters,to.plot,"\\\",sep=")),grep,
  x.dimnames(sims.array)[1:3]))])

write.csv(signif(ptable,3),paste(model.name,"/",model.name,"",
  summary.csv",sep=""))

#####
#posterior distributions of parameters
```

```
pred = posterior[,grep(FxPred, dimnames(posterior)[2:3])]

x1 = xdata
x1$type = rep("observed",nrow(x1))
x1$cavg = rbind(x1,x1)
x2$fxa.ing.avg = as.vector(t(apply(pred,2,
  quantile,probs=c(0.95,0.5,0.05))))
x2$type = rep(c("##",median,"95%ile"),ea=nrow(x1))
x1= rbind(x1,x2)
x1 = [order(x1$type,x1$cavg),]
```

```
xplot(fxa.inh.avg~avg,x1,groups=type,
  panel=panel.superpose,pch=(NA,NA,NA,1),
  type=c("l","p"),lty=c(3,1,0),col=c("red","red","blue",
  "black"),lwd=3,plevel.list=cex1,par.strip.text=cex1,
  strip=function(...) strip.default(..., style = 1),
  xlab= list(label="time-averaged plasma drug concentration",cex=1.2),
  ylab= list(label="time-averaged Fxa inhibition (%)",cex=1.2))

dev.off() # close graphics device
```

```
mpi.close.Rslaves() # stop R slave processes
```

Figure 2. WinBUGS Model for Sigmoid Emax Model Example

```
model{

  for(i in 1:nobs){
    FXa[i] ~ dnorm(FxHat[i],tau) #likelihood
    FxPred[i] ~ dnorm(FxHat[i],tau) #simulation to generate posterior
    #predictive distributions
    FxHat[i] <- Emax * pow(cobs[i],gamma) / (pow(EC50,gamma) +
    pow(cobs[i],gamma))
  }

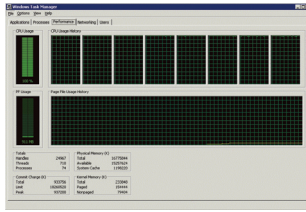
  Emax ~ dunif(0,100)
  logEC50 ~ dnorm(0,0.000001)
  log(EC50) < logEC50
  gamma ~ dunif(0,10)
  sigma ~ dunif(0,10)
  tau <- 1/(sigma*sigma)

}
```

RESULTS

- Elapsed time for computation of 8 MCMC chains of 100,000 iterations
 - 8 processors: 243 seconds
 - 1 processor: 1806 seconds

Figure 3. Screen Shot of Windows Task Manager



Screenshot of Windows Task Manager showing 100% CPU usage of 8 processors during execution of example problem.

Figure 4. MCMC History Plot

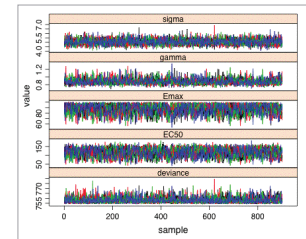


Figure 5. Marginal Posterior Distributions of Model Parameters

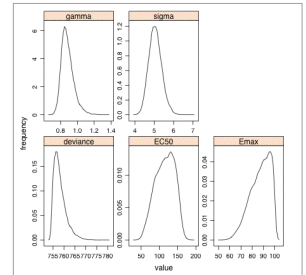
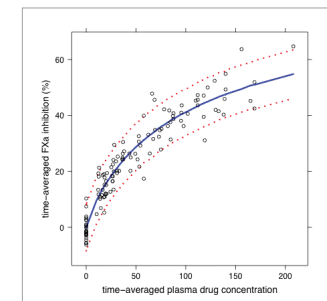


Figure 6. Data and Model Predictions (Posterior Median and 90% Credible Intervals)



DISCUSSION

- `bugsParallel` provides a practical open source option for parallel computation of multiple MCMC chains using WinBUGS in a MS Windows environment.
- `bugsParallel` is freely available from Metrum Institute (<http://metruminstitute.org/>).
- The remaining software components required for implementation are also freely available at the sites listed under "References."
- Integration with R permits data management, MCMC and analysis of MCMC results within a single software environment.
- Works across multiple machines or multiple processors within a single machine.
- Limitations of current version of `bugsParallel`
 - Limited to a grid of Windows computers/processors.
 - Lack of substantial "bullet-proofing" makes it poorly suited for a large-scale grid.
 - E.g., it does not implement "bullet-proofing" such as identifying and aborting "stuck" processes and cleaning up after crashed processes.

REFERENCES

- D.J Lunn, A.Thomas, N.Best, D.Spiegelhalter. (2000) WinBUGS -- a Bayesian modelling framework: concepts, structure, and extensibility. Statistics and Computing, 10:325–337. (<http://www.mrc-bsu.cam.ac.uk/bugs/winbugs/contents.shtml>)
- R Development Core Team. (2007) R: A Language and Environment for Statistical Computing. (<http://www.r-project.org/>)
- <http://cran.wustl.edu/src/contrib/Descriptions/R2WinBUGS.html> and <http://www.stat.columbia.edu/~gelman/bugsR/>
- <http://cran.wustl.edu/src/contrib/Descriptions/Rmpi.html> and <http://www.stats.wo.ca/faculty/yu/Rmpi>
- <http://cran.wustl.edu/src/contrib/Descriptions/rlecuyer.html>
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- IG Gigris, T.Schaible, P.Nandy, F.De Ridder, J.Mathers, S.Mohanty. (2007) Parallel Bayesian methodology for population analysis. PAGE Meeting, June 13-15, 2007, Copenhagen, Denmark (http://www.page-meeting.org/pdf_assets/6232-PARALLEL%20Gigris_POSTER_final.pdf).