MultiBUGS: A parallel implementation of the BUGS modelling framework for faster Bayesian inference

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Background

BUGS is general-purpose Bayesian modelling software that implements Markov chain Monte Carlo (MCMC).

Started in 1989: ClassicBUGS, then WinBUGS, then OpenBUGS

Ideas from BUGS widely adopted

- JAGS (Plummer, 2017)
- NIMBLE (Valpine et al., 2017)
- Related ideas are used in Stan (Carpenter et al., 2017)

Latest version is MultiBUGS:

- Available to download from https://www.multibugs.org
- This talk is based on Goudie et al. (?2019)

Background & motivation

Impossible or extremely time-consuming to use OpenBUGS with a huge amount of data.

- OpenBUGS uses only a single CPU/core/thread
- Increases in single-thread performances slowing
- Number of cores available increasing

Aim: to make the speed-ups of multi-core computation available to applied statisticians using BUGS for general models, without requiring any knowledge of parallel programming

Note: not aiming to improve mixing properties of the Markov chain, simply to run it faster
Parallelisation in MultiBUGS

MultiBUGS implements two levels of parallelisation.

Simple approach – run each of multiple, independent MCMC chains on a separate CPU or core (Bradford and Thomas, 1996)

- Useful for assessing convergence e.g. the Brooks-Gelman-Rubin diagnostic
- Burn-in time isn’t shortened

More complicated approach – use multiple CPUs/cores for a single MCMC chain

- Aim to shorten the per-iteration computation time by identifying tasks that can be calculated in parallel
- MultiBUGS parallelises the following tasks:
  1. “Likelihood” computation
  2. Sampling of conditionally-independent components

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Selected other parallelisation approaches

1. Speculatively consider sequence of MCMC steps, evaluate each on a separate core.

   e.g. Brockwell (2006)

2. Modify Metropolis-Hastings algorithm by proposing a sequence of candidate points in parallel.

   e.g. Calderhead (2014).

3. Run parts of the model on separate cores and then combine

   e.g. Scott et al. (2016), Goudie et al. (2019)


Trivial illustrative example (“seeds”)

A random-effects logistic regression without outcome $r_i$ and covariates $X_{1i}$ and $X_{2i}$ (21 observations)

$$r_i \sim \text{Bin}(p_i, n_i)$$

$$\logit(p_i) = \alpha_0 + \alpha_1 X_{1i} + \alpha_2 X_{2i} + \alpha_{12} X_{1i} X_{2i} + \beta_i$$

$$\beta_i \sim \text{N}(\mu_\beta, \sigma^2_\beta)$$

$$\alpha_0, \alpha_1, \alpha_2, \alpha_{12} \sim \text{N}(\mu_\alpha, \sigma^2_\alpha)$$

$$\sigma_\beta \sim \text{Unif}(\sigma_{\text{min}}, \sigma_{\text{max}})$$
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Generic algorithm used by BUGS

At each MCMC iteration, BUGS does the following:

```
for v in S do
    Do something involving \( p(v | V_{-v}) \)
end for
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\textbf{for} \ v \ \textbf{in} \ S \ \textbf{do}

Do something involving \( p(v \mid V_{-v}) \)

\textbf{end for}

The conditional distribution \( p(v \mid V_{-v}) \) of a node \( v \in S \), given the other nodes \( V_{-v} \), is

\[
p(v \mid V_{-v}) \propto p(v \mid pa(v)) \times \prod_{u \in ch(v)} p(u \mid pa(u))
\]

\[
= p(v \mid pa(v)) \times L(v)
\]

\[
= \text{“prior” term} \times \text{“likelihood” term}
\]
Generic algorithm used by BUGS

At each MCMC iteration, BUGS does the following:

\begin{verbatim}
for v in S do
    Evaluate the “prior” \( p(v \mid \text{pa}(v)) \)
    for u ∈ ch(v) do
        Evaluate “likelihood” component \( p(u \mid \text{pa}(u)) \)
    end for
    etc ...
end for
\end{verbatim}

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Type 1 – Splitting likelihood computation

When a parameter has many children, the likelihood is the product of many terms.

\[ L(v) = \prod_{u \in \text{ch}(v)} p(u | \text{pa}(u)) \]

But, with a partition of the children \( \text{ch}(v) = \{ \text{ch}^{(1)}(v), \ldots, \text{ch}^{(C)}(v) \} \),

\[ L(v) = \left[ \prod_{u \in \text{ch}^{(1)}(v)} p(u | \text{pa}(u)) \right] \times \left[ \prod_{u \in \text{ch}^{(2)}(v)} p(u | \text{pa}(u)) \right] \times \ldots \times \left[ \prod_{u \in \text{ch}^{(C)}(v)} p(u | \text{pa}(u)) \right] \]
Type 2 – Parallelising sampling of parameters

When a model includes a large number of parameters then computation may be slow in aggregate, even if sampling of each individual parameter is fast.

But parameters that do not directly depend on each other can be updated simultaneously.

More precisely, parameters in a mutually conditionally-independent set $W \subseteq S$ can be updated simultaneously. That is, $W$ satisfying

$$\text{all } w_1, w_2 \in W \ (w_1 \neq w_2) \text{ satisfy } w_1 \perp w_2 \mid V \setminus W$$

If not all parameters can be collated into a single $W$, form a series of $W$s and sample in turn.
Identifying sets $W$

Define the **topological depth** of a node $v \in V$ recursively, starting from the nodes with no parents.

$$d(v) = \begin{cases} 
0 & \text{if } \text{pa}(v) = \emptyset \\
1 + \max_{u \in \text{pa}(v)} d(u) & \text{otherwise}
\end{cases}$$
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Depth sets $D^h = \{ v \in S : d(v) = h \}$
Identifying sets $W$

Define the topological depth of a node $v \in V$ recursively, starting from the nodes with no parents.

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Depth sets $D^h = \{v \in S : d(v) = h\}$

Parameters in a set $W$ within a single depth set $D^h$ are mutually conditionally-independent, given the other nodes $V \setminus W$, if the parameters in $W$ have no child node in common.
Which type of parallelism to exploit for each parameter in the model?

MultiBUGS aims to

- Parallelise the evaluation of the “likelihood” of ‘fixed effect’-like parameters (Type 1)
- Parallelise sampling of ‘random effect’-like parameters (Type 2)
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Let $\bar{ch} = \text{mean}_v |ch(v)|$ be the mean number of children across parameters

**Heuristic algorithm:**
Consider each depth set $D^h$ in turn, starting with the ‘deepest’ set

```plaintext
if a parameter has more than $2 \times \bar{ch}$ children then
    Parallelise evaluation of this parameter’s “likelihood” (Type 1)
else
    Sample this parameter in parallel, if possible (Type 2)
end if
```
Computation schedule for illustrative example, with 4 cores

There are 26 stochastic parameters.
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Topological depths:
- $d(\beta_1) = \cdots = d(\beta_{21}) = 2$
- $d(\alpha_0) = \cdots = d(\alpha_{12}) = d(\sigma_\beta) = 1$
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1. Parameters $\beta_1, \ldots, \beta_{21}$
   - Likelihood evaluation not parallelised these parameter have only 1 child and $\bar{\text{ch}} \approx 4.8$.
   - But, $\beta_1, \ldots, \beta_{21}$ are mutually conditionally-independent and so sampling can be parallelised
   - $21 \mod 4 \neq 0$ so we will have idle cores

<table>
<thead>
<tr>
<th>Row</th>
<th>Core 1</th>
<th>Core 2</th>
<th>Core 3</th>
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   - \( 21 \mod 4 \neq 0 \) so we will have idle cores

2. Parameters \( \alpha_0, \alpha_1, \alpha_2, \alpha_{12} \) and \( \sigma_\beta \)
   - All of these parameters have 21 children \( \bar{ch} \approx 4.8 \), so likelihood computation is parallelised

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Implementation notes

Each core keeps

- The complete DAG, the computation schedule, and associated sampling algorithms
- A copy of the current state of the MCMC
- Two pseudo-random number generation (PRNG) streams
  1. “Core-specific” stream, initialised with a different seed for each core
     - Used when we wish to sample independently across cores
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For Metropolis-Hastings: the prior, the sampling of new value, and Metropolis test (redundantly) replicated on every core, using “common” PRNG stream
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**Sampling parallelisation:** Just delete the nodes that are sampled elsewhere from the list of nodes to be updated. Sample using “core-specific” PRNG stream, and propagate new values across cores using `Allgather`.
Hierarchical regression example

Based on an analysis linked database of methadone prescriptions given to opioid dependent patients in Scotland (Gao et al., 2016)

425,112 observations, with the following structure:

- Geographic regions \( (i = 1, \ldots, 8) \)
  - Containing patients (20410 in total)
    - Each of whom may have multiple prescriptions

For some measurements patient-level identifiers are available:

\[
y_{ijk} = \sum_{m=1}^{4} \beta_m \times x_{mij} + u_i + v_i \times r_{ijk} + w_{ij} + \varepsilon_{ijk}
\]

For other measurements no patient-level identifier is available:

\[
z_{il} = \lambda + u_i + v_i \times s_{il} + \eta_{il}
\]

Timings

In OpenBUGS, running chains 2 for 15,000 iterations takes about 32 hours.

In MultiBUGS:

- Sampling of pairs of random-effect means and variances parallelised;
- Sampling of person-level random effects $w_{ij}$ parallelised, except for the component corresponding to the person with the most observations (176 observations)
- The likelihood computation of all the other parameters in the model is parallelised
When is MultiBUGS quicker than OpenBUGS?

Independent-chain parallelisation is almost always be advantageous whenever sufficient cores are available, since no communication across cores is needed.

Within-chain parallelisation will be most useful for models involving parameters with a large number of likelihood terms and/or a large number of conditionally independent parameters.

- e.g. many standard regression-type models involving both fixed and random effects

For models without these features, the overheads of within-chain parallelisation may outweigh the gains on some computing hardware.

Note the mixing properties are the same as in OpenBUGS (but the exact samples will differ due to different PRNG stream use)
Outlook

MultiBUGS 1.0 is finished – https://www.multibugs.org

MultiBUGS 2.0¹ uses a more efficient system for communicating partial models to cores

Released version requires Windows, but we did port MultiBUGS 1.0 to Ubuntu

A compendium of ‘big models’ would be useful

Martyn Plummer is adopting a similar idea in JAGS (using OpenMP)

¹https://github.com/MultiBUGS/MultiBUGS