# Discussion of: Unbiased MCMC with couplings (Jacob et al)

#### **Darren Wilkinson**

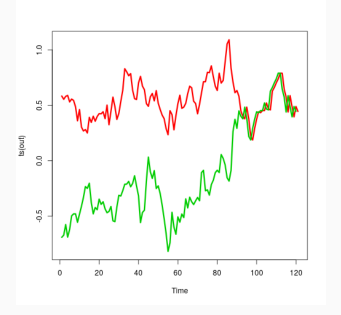
@darrenjw
darrenjw.github.io
Newcastle University and The Alan Turing Institute

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#### **Overview**

- Since in general we cannot initialise an MCMC chain with an exact sample from the target, we rely on asymptotic convergence to equilibrium, but diagnosing "burn in" and "convergence" is difficult
- The method outlined in the read paper does not solve the "exact sampling" problem, but instead provides a method for removing the bias from any estimates that are produced from the algorithm output, using a pair of coupled chains
- The approach is quite general, and can potentially be applied to many different kinds of MCMC algorithms — this paper concentrates on MH and Gibbs sampling (so I will, too), but there are a bunch of related papers exploring applications to other MCMC schemes

# Coupled Metropolis chains (with offset of 1)



#### Coupling versus convergence

• Time-averaged estimator:

$$H_{k:m}(X,Y) = \frac{1}{m-k+1} \sum_{t=k}^{m} h(X_t) + \sum_{t=k+1}^{\tau-1} \min\left(1, \frac{t-k}{m-k+1}\right) \{h(X_t) - h(Y_{t-1})\}$$

$$\equiv \mathsf{MCMC}_{k:m} + \mathsf{BC}_{k:m}$$

- Just the regular MCMC estimate with a correction term that terminates once the chains have coupled
- Coupling depends on the (arbitrary) proposed coupling mechanism as well as the mixing properties of the chain
- Possible to engineer early coupling (via initial conditions and MH kernel, for example), but this might be a dangerous strategy (see multi-modal example in paper)

4

## **Coupling Gibbs samplers**

- ullet The technique for coupling MH kernels works for multivariate as well as univariate samplers, though reflection maximal coupling works much better than a simple  $\gamma$ -maximal coupling in high dimensions
- However, it also makes sense to want to couple Gibbs samplers and other component-wise update algorithms
- For each component update of a Gibbs sampler, just couple the full-conditionals for the two chains
- Once all components have coalesced, the full state of the chain will be coalesced, and the coupling of the two chains will be faithful
- Little theory providing reassurance that this will happen in reasonable time, but seems to work empirically

## Coupled Gibbs sampler for an AR(1)

Consider an AR(1) process defined by

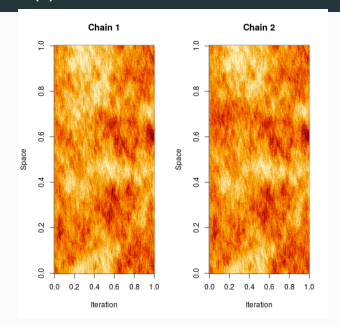
$$X_t = \alpha X_{t-1} + \epsilon_t, \qquad \epsilon_t \sim N(0, \sigma^2),$$

for  $t=1,2,\ldots,N$ , with periodic boundaries (so that  $X_0\equiv X_N$  and  $X_{N+1}\equiv X_1$ ), and stationary variance  $\sigma^2/(1-\alpha^2)$ . The full-conditional for each variable  $X_t$  is of the form

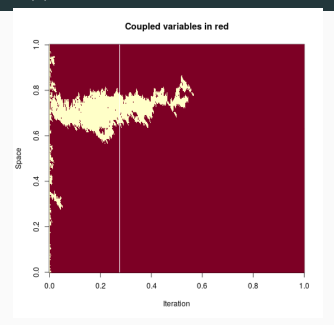
$$X_t | X_{t-1}, X_{t+1} \sim N\left(\frac{\alpha}{1+\alpha^2}(X_{t-1} + X_{t+1}), \frac{\sigma^2}{1+\alpha^2}\right).$$

We run two chains by cycling through each variable in turn and sampling from the coupled full conditionals. Here we use N=200 and  $\alpha=0.99$ , for n=500 iterations.

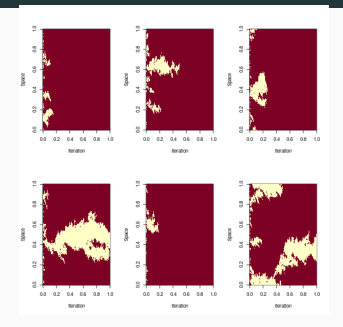
# Coupled AR(1) chains



# Coupled AR(1) variables



# Coupling behaviour for pairs of AR(1) chains



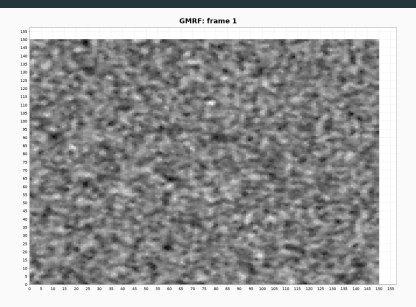
#### Gibbs sampler for a GMRF

 For a more challenging/interesting example, consider a Gaussian Markov random field (on a torus) with full-conditionals

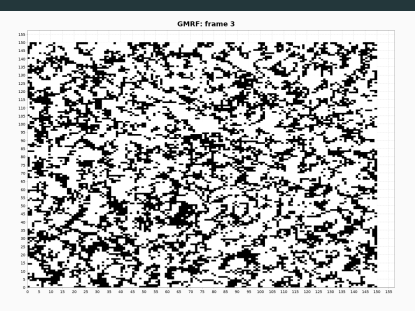
$$X_{i,j}|\cdot \sim N\left(\frac{\alpha}{4}(X_{i-1,j} + X_{i+1,j} + X_{i,j-1} + X_{i,j+1}), \sigma^2\right).$$

• Exact simulation is possible, but here we run two chains by cycling through each pixel in turn and sample from the coupled full conditionals. Here we use a  $150 \times 150$  grid (22.5k variables) with  $\alpha = 0.99$ , and  $\sigma^2 = 1$ , and run chains until they couple in order to study the coupling time distribution (for 1k replicates).

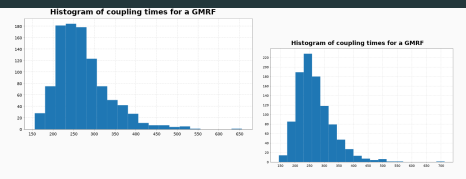
## Gibbs sampler for a GMRF



## Coupling of pixels for a GMRF



# Coupling time distribution



- Histograms of coupling time distributions (for 1k replicates), based on  $\gamma$ -maximal coupling (left) and reflection maximal coupling (right)
- Very similar profiles, and execution times, with the reflection coupling being slightly faster in terms of CPU time

## Coupling time distribution

- Using the standard maximal coupling, the mean coupling time is around 272, and median around 260. The maximum observed coupling time is 656.
- Using the reflection maximal coupling, the mean coupling time is around 268 and the median is around 257. The maximum observed coupling time is 712.
- This is consistent with other experiments I've done. For univariate kernels, the reflection coupling is no worse than the standard  $(\gamma)$  coupling, and sometimes slightly better
- The reflection coupling is really designed for high dimensional Gaussian kernels, where it encourages positive correlation between generated values
- In the univariate case, it's not immediately obvious that the reflection coupling should be better than the standard coupling

#### Parallel chains

- One of the main motivations for obtaining unbiased estimation of posterior expectations is to fix one of the issues with parallel chains MCMC
- We would like to run many chains independently on different processors and then pool results in some way
- In this case any bias in the chains will not "average out" across multiple processors, since there will be non-negligible bias in every chain
- Unbiased estimators can be safely averaged to get new unbiased estimators with reduced variance
- However, the debiasing doesn't actually eliminate burn-in it
  just corrects for it, so you still have repeated burn-in, limiting
  the scalability of the parallel chains approach

#### Using coupled chains in parallel

- ullet The paper recommends choosing k as a high quantile of the coupling time distribution, and m a multiple of k
- This is very consistent with the approach that would be taken for a (biased, uncoupled) parallel chains approach, with k playing the role of "burn-in"
- It might be "dangerous" to choose k this way if early coupling has been (artificially) engineered
- In any case, any non-zero k limits speed-up of parallel coupled chains relative to "one long run", via Amdahl's law for parallel chains MCMC (Wilkinson, 2005):

$$\mathsf{SpeedUp}(N) = \frac{b+n}{b+\frac{n}{N}} \xrightarrow{N} \frac{b+n}{b},$$

for burn-in b and monitoring run n on N processors (eg. asymptotic limit of 10 for n=9b).

#### Conclusion

- The method proposed in the paper is often straightforward to implement, and seems to work well on a range of problems
- It is potentially useful in the context of parallelisation of MCMC algorithms, since averaging unbiased estimators is relatively "safe", but it is not clear that it fundamentally solves the "parallel MCMC" problem, since some kind of "burn-in" must still be repeated for every pair of chains
- Many open questions regarding the co-development of MCMC and coupling algorithms for overall efficiency
- Overall, an interesting and discussable contribution to the literature, so it gives me great pleasure to propose the vote of thanks!

My materials: https://github.com/darrenjw/unbiased-mcmc