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INTERACTING MULTIPLE –TRY ALGORITHMS WITH DIFFERENT PROPOSAL DISTRIBUTIONS

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Abstract_

We propose a new class of interacting Markov chain Monte Carlo (MCMC) algorithms designed for increasing the efficiency of a modified multiple-try Metropolis (MTM) algorithm. The extension with respect to the existing MCMC literature is twofold. The sampler proposed extends the basic MTM algorithm by allowing different proposal distributions in the multiple-try generation step. We exploit the structure of the MTM algorithm with different proposal distributions to naturally introduce an interacting MTM mechanism (IMTM) that expands the class of population Monte Carlo methods and builds connections with the rapidly expanding world of adaptive MCMC. We show the validity of the algorithm and discuss the choice of the selection weights and of the different proposals. We provide numerical studies which show that the new algorithm can perform better than the basic MTM algorithm and that the interaction mechanism allows the IMTM to efficiently explore the state space.

Keywords: Interacting Monte Carlo, Markov chain Monte Carlo, Multiple-try Metropolis,

Population Monte Carlo.

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1 Introduction

Markov chain Monte Carlo (MCMC) algorithms are now essential for the analysis of complex statistical models. In the MCMC universe, one of the most widely used class of algorithms is defined by the Metropolis-Hastings (MH) (Metropolis et al., 1953; Hastings, 1970) and its variants. An important generalization of the standard MH formulation is represented by the multiple-try Metropolis (MTM) (Liu et al., 2000). While in the MH formulation one accepts or rejects a single proposed move, the MTM is designed so that the next state of the chain is selected among multiple proposals. The multiple-proposal setup can be used effectively to explore the sample space of the target distribution and subsequent developments made use of this added flexibility. For instance, Craiu and Lemieux (2007) and Bédard et al. (2010) propose to use antithetic and quasi-Monte Carlo samples to generate the proposals and to improve the efficiency of the algorithm while Pandolfi et al. (2010b) and Pandolfi et al. (2010a) apply the multiple-proposal idea to a trans-dimensional setup and combine Reversible Jump MCMC with MTM.

This work further generalizes the MTM algorithm presented in Liu et al. (2000) in two directions. First, we show that the original MTM transition kernel can be modified to allow for different proposal distributions in the multiple-try generation step while preserving the ergodicity of the chain. The use of different proposal distributions gives more freedom in designing MTM algorithms for target distributions that require different proposals across the sample space. An important challenge remains the choice of the distributions used to generate the proposals and we propose to address it by expanding upon methods used within the population Monte Carlo class of algorithms.

The class of population Monte Carlo procedures (Cappé et al., 2004; Del Moral and Miclo, 2000; Del Moral, 2004; Jasra et al., 2007) has been designed to address the inefficiency of classical MCMC samplers in complex applications involving multimodal and high dimensional target distributions (Pritchard et al., 2000; Heard et al., 2006). Its formulation relies on a number of MCMC processes that are run in parallel while learning from one another about the geography of the target distribution.

A second contribution of the paper is finding reliable generic methods for constructing the proposal distributions for the MTM algorithm. We propose an interacting MCMC sampling design for the MTM that preserves the Markovian property. More specifically, in the proposed interacting MTM (IMTM) algorithm, we allow the distinct proposal distributions to use information produced by a population of auxiliary chains. We infer that the resulting performance of the MTM is tightly connected to the performance of the chains' population. In order to maximize the latter, we propose and compare via simulations a number of strategies that can be used to tune the auxiliary chains.

In the next section we discuss the IMTM algorithm, propose a number of alternative implementations and prove their ergodicity. In Section 3 we focus on some special cases of the IMTM algorithm and in Section 4 the performance of the methods proposed is demonstrated with simulations and real examples. We end the paper with a discussion of future directions for research.

2 Interacting Monte Carlo Chains for MTM

We begin by describing the MTM and its extension for using different proposal distributions.

2.1 Multiple-Try Metropolis With Different Proposal Distributions

Suppose that of interest is sampling from a distribution π that has support in $\mathcal{Y} \subset \mathbf{R}^d$ and is known up to a normalizing constant. Assuming that the current state of the chain is x, the update defined by the MTM algorithm of Liu et al. (2000) is described in Algorithm 1.

Note that while the MTM uses the same distribution to generate all the proposals, it is possible to extend this formulation to different proposal distributions without altering the ergodicity of the associated Markov chain.

Let $T_j(\cdot|x)$, with j = 1, ..., M, be a set of proposal distributions for which $T_j(y|x) > 0$ if and only if $T_j(x|y) > 0$. Define

$$w_j(x,y) = \pi(x)T_j(y|x)\lambda_j(x,y) \qquad j = 1,\dots,M$$

where $\lambda_j(x, y)$ is a nonnegative symmetric function in x and y that can be chosen by the user. The only requirement is that $\lambda_j(x, y) > 0$ whenever T(x, y) > 0. Then the MTM algorithm with different proposal distributions is given in Algorithm 2.

- 1. Draw M trial proposals y_1, \ldots, y_M from the proposal distribution $T(\cdot|x)$. Compute $w(y_j, x)$ for each $j \in \{1, \ldots, M\}$, where $w(y, x) = \pi(y)T(x|y)\lambda(y, x)$, and $\lambda(y, x)$ is a symmetric function of x, y.
- 2. Select y among the M proposals with probability proportional to $w(y_j, x), j = 1, ..., M$.
- 3. Draw x_1^*, \ldots, x_{M-1}^* variates from the distribution $T(\cdot|y)$ and let $x_M^* = x$.
- 4. Accept y with generalized acceptance probability

$$\rho = \min\left\{1, \frac{w(y_1, x) + \ldots + w(y_M, x)}{w(x_1^*, y) + \ldots + w(x_M^*, y)}\right\}$$

Algorithm 2. MTM with Different Proposal Distributions

- 1. Draw independently M proposals y_1, \ldots, y_M such that $y_j \sim T_j(\cdot|x)$. Compute $w_j(y_j, x)$ for $j = 1, \ldots, M$.
- 2. Select Y = y among the trial set $\{y_1, \ldots, y_M\}$ with probability proportional to $w_j(y_j, x), j = 1, \ldots, M$. Let J be the index of the selected proposal. Then draw $x_j^* \sim T_j(\cdot|y), j \neq J, j = 1, \ldots, M$ and let $x_J^* = x$.
- 3. Accept y with probability

$$\rho = \min\left\{1, \frac{w_1(y_1, x) + \dots + w_M(y_M, x)}{w_1(x_1^*, y) + \dots + w_M(x_M^*, y)}\right\}$$

and reject with probability $1 - \rho$.

It should be noted that Algorithm 2 is a special case of the interacting MTM presented in the next section and that the proof of ergodicity for the associated chain follows closely the proof given in Appendix A for the interacting MTM and is not given here.

2.2 General Construction

Undoubtedly, Algorithm 2 offers additional flexibility in organizing the MTM sampler. This section introduces generic methods for using a population of MCMC chains to define the proposal distributions.

Consider a population of N chains, $X^{(i)} = \{x_n^{(i)}\}_{n \in \mathbb{N}}$ and $i = 1, \ldots, N$. For full generality we assume that the *i*th chain has MTM transition kernel with M_i different proposals $\{T_j^{(i)}\}_{1 \leq j \leq M_i}$ (if we set $M_i = 1$ we imply that the chain has a MH transition kernel). The interacting mechanism allows each proposal distribution to possibly depend on the values of the chains at the previous step. Formally, if $\Xi_n = \{x_n^{(i)}\}_{i=1}^N$ is the vector of values taken at iteration $n \in \mathbb{N}$ by the population of chains, then we allow each proposal distribution used in updating the population at iteration n+1 to depend on Ξ_n . The mathematical formalization is used in the description of Algorithm 3. One expects that the chains in the population are spread throughout the sample space and thus the proposals generated are a good representation of the sample space \mathcal{Y} ultimately resulting in better mixing for the chain of interest.

The first step in Algorithm 3 suggests that each proposal distribution used in each parallel MTM chain is allowed to depend on the current states of all the chains in the population. However, this general formulation for the IMTM, though correct in theory, can be difficult to tune efficiently in a given practical problem. Before we move to discuss implementations that simplify and enhance the practical application of the IMTM algorithm, we prove below that the chain underlying Algorithm 3 is ergodic to π .

In order to give a representation of the IMTM transition density let us introduce the following notation. Let $T^{(i)}(y_{1:M_i}|x) = \prod_{k=1}^{M_i} T_k^{(i)}(y_k|\tilde{f}_n^{(i)}(x))$ and $T_{-j}^{(i)}(y_{1:M_i}|x) = \prod_{k\neq j}^{M_i} T_k^{(i)}(y_k|\tilde{f}_n^{(i)}(x))$ and define $dy_{1:M_i} = \prod_{k=1}^{M_i} dy_k$ and $dy_{-j} = \prod_{k\neq j}^{M_i} dy_k$.

The transition density associated to the population of chains is then

$$K(\Xi_n, \Xi_{n+1}) = \prod_{i=1}^{N} K_i(x_n^{(i)}, x_{n+1}^{(i)})$$
(1)

where

$$K_i(x,y) = \sum_{j=1}^{M_i} A_j^{(i)}(x,y) T_j^{(i)}(y|x) + \left(1 - \sum_{j=1}^{M_i} B_j^{(i)}(x)\right) \delta_x(y)$$
(2)

is the transition kernel associated to the i-th chain of algorithm with

$$A_{j}^{(i)}(x,y) = \int_{\mathcal{Y}^{2(M_{i}-1)}} \tilde{w}_{j}^{(i)}(y,x)\rho_{j}^{(i)}(x,y)T_{-j}^{(i)}(x_{1:M_{i}}^{*}|y)T_{-j}^{(i)}(y_{1:M_{i}}|x)dx_{-j}^{*}dy_{-j}dy$$

and

$$B_{j}^{(i)}(x) = \int_{\mathcal{Y}^{2(M_{i}-1)+1}} \rho_{j}^{(i)}(x,y) T_{-j}^{(i)}(x_{1:M_{i}}^{*}|y) T^{(i)}(y_{1:M_{i}}|x) dx_{-j}^{*} dy_{1:M_{i}}.$$

Algorithm 3. Interacting Multiple Try Algorithm (IMTM)

- For i = 1, ..., N
 - 1. Let $x = x_n^{(i)}$ and define the map $\tilde{f}_n^{(i)}(z) = (x_n^{(1:i-1)}, z, x_n^{(i+1:N)})^T$; for $j = 1, \ldots, M_i$ draw $y_j \sim T_j^{(i)}(\cdot | \tilde{f}_n^{(i)}(x))$ independently and compute

$$w_j^{(i)}(y_j, x) = \pi(y_j) T_j^{(i)}(x | \tilde{f}_n^{(i)}(y_j)) \lambda_j^{(i)}(y_j, x).$$

- 2. Select $J \in \{1, \ldots, M_i\}$ with probability proportional to $w_i^{(i)}(y_j, x), j = 1, \ldots, M_i$ and set $y = y_J$.
- 3. For $j = 1, ..., M_i$ and $j \neq J$ draw $x_j^* \sim T_j^{(i)}(\cdot | \tilde{f}_n^{(i)}(y))$, let $x_J^* = x_n^{(i)}$ and compute

$$w_j^{(i)}(x_j^*, y) = \pi(x_j^*) T_j^{(i)}(y | \tilde{f}_n^{(i)}(x_j^*)) \lambda_j^{(i)}(x_j^*, y).$$

4. Set $x_{n+1}^{(i)} = y$ with probability

$$\rho_i = \min\left\{1, \frac{w_1^{(i)}(y_1, x) + \ldots + w_{M_i}^{(i)}(y_{M_i}, x)}{w_1^{(i)}(x_1^*, y) + \ldots + w_{M_i}^{(i)}(x_{M_i}^*, y)}\right\}$$

and
$$x_{n+1}^{(i)} = x_n^{(i)}$$
 with probability $1 - \rho_i$.

In the above equations $\tilde{w}_{j}^{(i)}(y_{j}, x) = w_{j}^{(i)}(y_{j}, x) / (w_{j}^{(i)}(y, x) + \bar{w}_{-k}^{(i)}(y_{1:M_{i}}|x))$, with $j = 1, \ldots, M_{i}$ and $\bar{w}_{-j}^{(i)}(y_{1:M_{i}}|x) = \sum_{k \neq j}^{M_{i}} w_{k}^{(i)}(y_{k}, x)$, are the normalized weights used in the selection step of the IMTM algorithm and

$$\rho_j^{(i)}(x,y) = \min\left\{1, \frac{w_j^{(i)}(y,x) + \bar{w}_{-j}^{(i)}(y_{1:M_i}|x)}{w_j^{(i)}(x,y) + \bar{w}_{-j}^{(i)}(x_{1:M_i}^*|y)}\right\}$$

is the generalized MH ratio associated to a MTM algorithm.

The validity of the IMTM algorithm relies upon the detailed balance condition.

Theorem 1. The transition density $K_i(x_n^{(i)}, x_{n+1}^{(i)})$ associated to the *i*-th chain of the IMTM algorithm satisfies the conditional detailed balanced condition.

Proof See Appendix A.

Since each transition $K_i(x_n^{(i)}, x_{n+1}^{(i)})$, i = 1, ..., N has $\pi(x)$ as stationary distribution and satisfies the conditional detailed balance condition then the joint transition $K(\Xi_n, \Xi_{n+1}) = \prod_{i=1}^N K_i(x_n^{(i)}, x_{n+1}^{(i)})$ has $\pi(x)^N$ as a stationary distribution.

3 Practical Implementation

It should be noted that at each iteration of the IMTM the computational complexity of the algorithm is $\mathcal{O}(\sum_{i=1}^{N} M_i)$. This can become burdensome when the number of chains, N, and the number of proposals, M_i , are simultaneously large so one needs to decide between increasing the number of chains or the number of proposals. We distinguish two possible strategies in designing the interaction mechanism. The first strategy is to use a small number of chains, say $5 \leq N \leq 20$, and a number of proposals equal to the number of chains, i.e. $M_i = N$, for all $1 \leq i \leq N$. In this way all the chains can interact at each iteration of the algorithm and many search directions can be included among the proposals.

A second strategy is to use a higher number of chains, e.g. N = 100, in order to possibly have, at each iteration, a good approximation of the target or a much higher number of search directions for a good exploration of the sample space. This algorithm design strategy is common in Population Monte Carlo or Interacting MCMC methods. Clearly when a high number of chains is used within IMTM, it is necessary to set $M_i < N$, possibly $M_i = 1$ for each auxiliary chain.

In this section we discuss a few strategies to built the M_i proposals for each chain and in the simulation section we compare the two strategies outlined above.

3.1 Parsing the Population of Auxiliary Chains

Although one may run a large number of auxiliary chains, one may not want to use all the chains at each iteration of the IMTM. One approach that turned out to be successful in our applications consists in selecting a random subset of chains from the population in order to build the proposals. For ease of description, assume that $M_i = M < N$, for all chains, $1 \le i \le N$ N. Then, when updating the *i*-th chain of the population, we sample the random indices I_1, \ldots, I_{M-1} from the uniform distribution $\mathcal{U}\{1, \ldots, N\}$ and we let $I_M = i$. Then the *M* proposals used for chain *i* will be allowed to depend only on the current states of those chains with indices I_1, \ldots, I_M (which includes the index i). Using the notation introduced and letting $I_n^{(i)} = (I_1, \ldots, I_M)$ then the *M* proposals used for chain *i* at time *n* are sampled using $T_j^{(i)}(y|x_n^{(I_1)}, \ldots, x_n^{(I_M)})$, for all $j = 1, \ldots, M$. Our simulation experiments showed a good performance when we used a relatively simpler version in which the jth proposal depends only on the current state of chain I_j , i.e., it is sampled using $T_j^{(i)}(\cdot|x_n^{(I_j)})$, for all $j = 1, \ldots, M$. One can see that the interweaving of the chains is performed by allowing the proposals used in chain i to be sampled conditional not only on the current state of the chain, $x_n^{(i)}$, but also on the current states of those chains whose indices are sampled at random and stored in $I_n^{(i)}$.

Another important issue directly connected to the practical implementation of the IMTM is the choice of $\lambda_j^{(i)}(x, y)$. Previously suggested forms for the function $\lambda_i^{(i)}(x, y)$ (Liu et al., 2000) are:

a)
$$\lambda_j^{(i)}(x,y) = 2\{T_j^{(i)}(x|y) + T_j^{(i)}(y|x)\}^{-1}$$

b) $\lambda_j^{(i)}(x,y) = \{T_j^{(i)}(x|y)T_j^{(i)}(y|x)\}^{-\alpha}, \alpha > 0$

Little guidance is offered in the existent literature regarding the choice of λ and, to our knowledge, in most applications of the original MTM algorithm the default choice is $\lambda = 1$.

Here we propose to include in the construction of λ the information provided by the population of chains. Therefore we suggest to modify the above functions to

$$\begin{split} \mathbf{a}') \ \ \lambda_{j}^{(i)}(x,y) &= 2\nu_{j} \left\{ T_{j}^{(i)}(x|y) + T_{j}^{(i)}(y|x) \right\}^{-1} \\ \mathbf{b}') \ \ \lambda_{j}^{(i)}(x,y) &= \nu_{j} \left\{ T_{j}^{(i)}(x|y) T_{j}^{(i)}(y|x) \right\}^{-\alpha}, \ \alpha > 0, \end{split}$$

where the factor ν_j is

$$\nu_j = \frac{1}{N} \left[1 + \sum_{i=1}^N \mathbf{1}_{\{j\}} (J_{n-1}^{(i)}) \right], \quad j = 1, \dots, M,$$
(3)

and $J_{n-1}^{(i)}$ is the index of the proposal selected in the *i*th chain update at iteration n-1. It can be seen that the $\{\nu_j\}_{1\leq j\leq M}$ capture the behaviour of the auxiliary chains at the previous iteration. More precisely, ν_j will be relatively larger for those proposal distributions $T_j(\cdot|\cdot)$ whose samples have been selected as the potential next states for the chains in the population at iteration n-1. The modifications proposed for $\lambda(\cdot, \cdot)$ would increase the use of those proposal distributions favoured by the population of chains at previous iteration. Since ν_j depends only on samples generated at the previous step by the population of chains, the ergodicity of the IMTM chain is preserved. In the simulation section we compare the performance of IMTM coupled with either a') or b') when $\alpha = 1$.

3.2 Annealed IMTM

Our belief in IMTM's improved performance is underpinned by the assumption that the population of Monte Carlo chains is spread throughout the sample space. This can be partly achieved by initializing the chains using draws from a distribution overdispersed with respect to π (see also Jennison, 1993; Gelman and Rubin, 1992) and partly by modifying the stationary distribution for some of the chains in the population. Specifically, we consider the sequence of annealed distributions $\pi_t = \pi^t$ with $t \in \{\xi_1, \xi_2, \dots, \xi_N\}$, where $1 = \xi_1 > \xi_2 > \ldots > \xi_n$, for instance $\xi_t = 1/t$. When t, s are close temperatures, π_t is similar to π_s , but $\pi = \pi_1$ may be much harder to sample from than π_{ξ_N} , as has been long recognized in the simulated annealing and simulated tempering literature (see Marinari and Parisi, 1992; Geyer and Thompson, 1994; Neal, 1994). Therefore, it is likely that some of the chains designed to sample from π_1, \ldots, π_N have good mixing properties, making them good candidates for the population of MCMC samplers needed for the IMTM. Recent theoretical work by Atchadé et al. (2010) has build, in an adaptive setup, connections between the temperature ladder and the optimal scaling problem. Such an analysis goes beyond the scope of this paper; in the simulation section we compare three methods for constructing the temperature ladder $1 = \xi_1 > \xi_2 > \ldots > \xi_n$.

We consider the Monte Carlo population made of the N-1 chains having $\{\pi_2, \ldots, \pi_N\}$ as stationary distributions. However, the use of MTM for *each*

auxiliary chain may be redundant since for smaller ξ_i 's the distribution π_i is easy to sample from. For this reason, in our simulations we shall use the annealed IMTM (AIMTM) in which the first chain is ergodic to π is based on the IMTM transition kernel and each auxiliary chain is a MH chain (M = 1)with target π_i , $2 \leq i \leq N$. The AIMTM is described in Algorithm 4. In practice, we always use the current state of the chain ergodic to π ($\xi = 1$) among the states used for generating one of the proposals (e.g., in Algorithm 4 we automatically set $I_1 = 1$ and let I_2, \ldots, I_M be sampled at random).

An additional gain could be obtained if the auxiliary chains' transition kernels are modified using adaptive MCMC strategies (see also Chauveau and Vandekerkhove, 2002, for another example of adaption for interacting chains). However, letting the auxiliary chains adapt indefinitely results in complex theoretical justifications for the IMTM which go beyond the scope of this paper and will be presented elsewhere. Our recommendation is to use finite adaptation for the auxiliary chains prior to the start of the IMTM. One could take advantage of multi-processor computing units and use parallel programming to increase the computational efficiency of this approach.

The adaptation of $\lambda_j^{(i)}$, through the weights ν_j defined in (3), should be used cautiously in this case. The aim of the annealing procedure is to allow the higher temperatures chains to explore widely the sample space and to improve the mixing of the MTM chain. Using ν_j the context of annealed IMTM could arbitrarily penalize some of the higher temperature proposals and reduce the effectiveness of the annealing strategy. For this reason we do not consider using adaptive λ 's for AIMTM.

Finally, we would like to note that it is possible to obtain a Monte Carlo approximation of a quantity of interest by using the output produced by *all the chains* in the population. For example let

$$\mathcal{I} = \int_{\mathcal{Y}} h(x) \pi(x) dx$$

be the quantity of interest where h is a test function. It is possible to approximate this quantity as follows

$$\mathcal{I}_{NT} = \frac{1}{T} \sum_{n=1}^{T} \frac{1}{\bar{\zeta}} \sum_{j=1}^{N} h(x_n^{(j)}) \zeta_j(x_n^{(j)})$$

where $x_n^{(i)}$ is the output of the *i*-th chain ergodic to target π^{ξ_i} at time *n*, for all $n = 1, \ldots, T$ and all $i = 1, \ldots, N$, $\zeta_j(x) = \pi(x)/\pi^{\xi_j}(x)$ are the importance weights and $\bar{\zeta} = \sum_{j=1}^N \zeta_j(x_n^{(j)})$.

- For i = 1
 - 1. Let $x = x_n^{(i)}$ and sample I_1, \ldots, I_M from $\mathcal{U}\{1, \ldots, N\}$.
 - 2. For j = 1, ..., M draw $y_j \sim T_j^{(i)}(\cdot | x_n^{(I_j)})$ independently and (a) If $I_j \neq 1$ set

$$w_j^{(i)}(y_j, x) = \pi(y_j) T_j^{(i)}(x | x_n^{(I_j)}) \lambda_j^{(i)}(y_j, x)$$

(b) If $I_j = 1$ set

$$w_j^{(i)}(y_j, x) = \pi(y_j) T_j^{(i)}(x|y_j) \lambda_j^{(i)}(y_j, x).$$

- 3. Select $J \in \{1, \ldots, M\}$ with probability proportional to $w_j^{(i)}(y_j, x), j = 1, \ldots, M$ and set $y = y_J$.
- 4. Let $x_J^* = x_n^{(i)}$ and for $j = 1, ..., M, j \neq J$,
 - (a) If $I_j \neq 1$ draw $x_j^* \sim T_j^{(i)}(\cdot | x_n^{(I_j)})$, (b) If $I_j = 1$ draw $x_j^* \sim T_j^{(i)}(\cdot | x_n)$

(b) If
$$I_j = 1$$
 draw $x_j^* \sim T_j^{(0)}(\cdot | y)$

- 5. Compute $w_i^{(i)}(x_j^*, y)$ using the same rule as in 2.
- 6. Set $x_{n+1}^{(i)} = y$ with probability ρ_i , where ρ_i is the generalized MH ratio of the IMT algorithm and $x_{n+1}^{(i)} = x_n^{(i)}$ with probability $1 - \rho_i$.
- For i = 2,..., N we perform the usual MH update using proposal distribution T⁽ⁱ⁾ for chain i, that is:
 - 1. Let $x = x_n^{(i)}$ and update the proposal function $T^{(i)}(\cdot|x)$.
 - 2. Draw $y \sim T^{(i)}(\cdot|x)$ and compute

$$\rho_i = \min\left\{1, \frac{\pi(y)^{\xi_i} T^{(i)}(x|y)}{\pi(x)^{\xi_i} T^{(i)}(y|x)}\right\}.$$

3. Set $x_{n+1}^{(i)} = y$ with probability ρ_i and $x_{n+1}^{(i)} = x_n^{(i)}$ with probability $1 - \rho_i$.

4 Simulation Results

4.1 Beta Mixture Model

Mixture models have been widely applied in many fields to capture heterogeneity in the data and Bayesian inference for such models represents a challenging statistical issue. More specifically, in the Bayesian analysis of mixture models, as the posterior distribution of a k-components mixture is invariant with respect to permutation of the labels of the parameters and exhibits k! modes. Sampling from the posterior is therefore a challenging problem which rarely can be solved successfully by the conventional singlechain MCMC methods. More efficient sampling algorithms are thus needed. As emphasized by Jasra et al. (2007) in the context of Bayesian mixture models, population Monte Carlo methods allow to sample efficiently from the posterior distribution.

We consider here a Bayesian mixture of normals that was previously used by Jasra et al. (2005) and Jasra et al. (2007) for comparing the performance of different population Monte Carlo methods. Let $y_1, \ldots y_n$ be ni.i.d. samples with density

$$\sum_{h=1}^{K} \tau_h f(y|\mu_h, \eta_h^{-1})$$
(4)

where K is the number of mixture components and $f(y_i|\mu_h, \eta_h^{-1})$ is the density of a normal distribution with location parameter μ_h and precision parameter η_h . The weights $\tau_h \ge 0, h = 1, \ldots, K$ of the mixture are such that $\sum_{h=1}^{K} \tau_h = 1$. We assume the following priors (see also Jasra et al., 2005; Richardson and Green, 1997).

$$\mu_j \sim \mathcal{N}(\xi, \kappa^{-1}), \quad \eta_j \sim \mathcal{G}a(\alpha, \beta), \quad \tau_{1:k-1} \sim \mathcal{D}ir(\delta)$$
 (5)

where $\mathcal{N}(\xi, \kappa^{-1})$, $\mathcal{G}a(\alpha, \beta)$ and $\mathcal{D}ir(\delta)$ are, respectively, the normal distribution with location ξ and precision κ , the gamma distribution with shape parameter α and scale parameter β and the symmetric Dirichlet distribution, with parameter δ .

We will use the problem of sampling from the posterior distribution defined by the model above as a benchmark for comparing the IMTM methods proposed in this paper with other population Monte Carlo algorithms based on MH kernels. We assume we have available a dataset of 100 (simulated) samples from an equally weighted, (i.e. $\tau_j = 1/k$ for $j = 1, \ldots, k$) normal mixture with k = 4 components with true means, $(\mu_1, \mu_2, \mu_3, \mu_4)^T =$ $(-3, 0, 3, 6)^T$, and equal standard deviations $\eta_k^{-1/2} = 0.55$, $1 \le k \le 4$. The algorithms being compared below are the following:

- **MH** A population of Monte Carlo algorithms in which all the N parallel chains have random walk MH (RWMH) kernels in which the *j*-th Gaussian proposal distribution has covariance $\sigma_j^2 \mathbf{I}$ where $\sigma_j = 0.01 + 0.59 * j/N$ for all $1 \le j \le N$ such that the acceptance rates obtained for the population of chains are between 10-60%.
- **MH1** A population of Monte Carlo algorithms in which each of the N parallel chains run a RWMH algorithm whose proposal distribution is a mixture of 4 normal densities. The standard deviations of the proposals are divided equally between 0.01 and 0.3.
- **MH2** A population of Monte Carlo algorithms in which each of the N transition kernels is a mixture of four RWMH kernels with same standard deviations as those defined for MH2.

The above algorithms do not allow interaction between the parallel chains which is arguably less flexible than the IMTM setup. Therefore we include in our comparison the above three algorithms to which we apply the cross-over interaction introduced by Liang and Wong (2001). The different chains of the population have the same target thus the acceptance-probability of the cross-over move is one.

- MH.c.o The MH algorithm described above with cross-over moves.
- MH1.c.o The MH1 algorithm described above with cross-over moves.
- MH2.c.o The MH2 algorithm described above with cross-over moves.

The six algorithms described above are compared with the following IMTM samplers:

- **IMTM-TA** An IMTM algorithm with N chains defined as in Section 3.1 and using $\lambda_j^{(i)}(x,y) = 2\{T_j^{(i)}(x|y) + T_j^{(i)}(y|x)\}^{-1}$ weights. The *j*-th proposal uses $T_j^{(i)}(y|x) = N(x, \sigma_j^2 \mathbf{I})$ where $\sigma_j = 0.01 + 0.59 * j/M$ for all $1 \le j \le M, 1 \le i \le N$.
- **IMTM-TA-a** The same algorithm as IMTM-TA but with adaptive weights $\lambda_j^{(i)}(x,y) = 2\nu_j \{T_j^{(i)}(x|y) + T_j^{(i)}(y|x)\}^{-1}$ where ν_j is defined as in (3)
- **IMTM-IS** An IMTM algorithm identical to IMTM-TA but using $\lambda_j^{(i)}(x, y) = \{T_j^{(i)}(x|y)T_j^{(i)}(y|x)\}^{-1}$ weights.

IMTM-IS-a The same algorithm as IMTM-IS but with adaptive weights $\lambda_j^{(i)}(x,y) = \nu_j \{T_j^{(i)}(x|y)T_j^{(i)}(y|x)\}^{-1}$ where ν_j is defined as in (3).

The comparison is made with respect to the estimation of the marginal means μ_1, \ldots, μ_4 . We consider T = 100,000 iterations and J = 100 chains for the MH, MH1, MH2, MH.c.o, MH1.c.o and MH2.c.o algorithms. For all the IMTM algorithms we used T = 10,000 iterations, produced from running N = 100 chains each with M = 10 proposals.

We observed from all the simulation experiments that IMTM-TA and IMTM-IS have similar performances, so we present the graphical results only for IMTM-TA. A typical output of the IMTM-TA algorithm is given in the top panel of Figure 1 which shows, for one of the chains in the population, the traces for each of the four coordinates sampled, μ_1 , μ_2 , μ_3 and μ_4 . We notice that the chain is able to switch frequently between the different modes of the posterior distribution and this compares favourably with the MH, MH1 and MH2 algorithms, with and without cross-over. The MH chains rarely switch between modes as can be seen also in Figure 1 of Jasra et al. (2007).

In order to give an alternative representation of the raw output of the population of chains we follow Frühwirth-Schnatter (2006) and present in Figure 2 the samples produced by each algorithm. The bottom panel in Figure 2 has been produced by projecting the samples on all the planes (μ_i, μ_j) with $i \neq j$ (in total we have K(K-1) = 12 such planes) and then superimposing all the plots into a single one. As discussed in Frühwirth-Schnatter (2006) the number of simulation clusters in this graphical representation, for a K-components mixture, is K(K-1) = 12, that is equal to 12 in our example. In the same panel we show (red line) the trajectory of one of the chains.

In Figure 2 we show samples produced by the other algorithms considered in the comparison. The six populations of chains, MH, MH1, MH2, with and without cross-over, are able to visit different modes of the posterior. Note that the samples from the population of MH chains are usually not evenly distributed across the different posterior modes. Moreover the single chains of the population of the MH algorithms usually visit only one of the clusters and are not able to visit the other clusters. In each panel the red line illustrates the path of a single chain. One can easily notice the difficulty of the MH, MH1 or MH2 chains without cross-over to explore the posterior surface. The red lines shown in the right-side panels crystallize the effect of the cross-over moves on the mixing property of the population of interacting chains. Each chain is now able to visit many modes and this results in improved efficiency for the class of MH algorithms considered here. However,

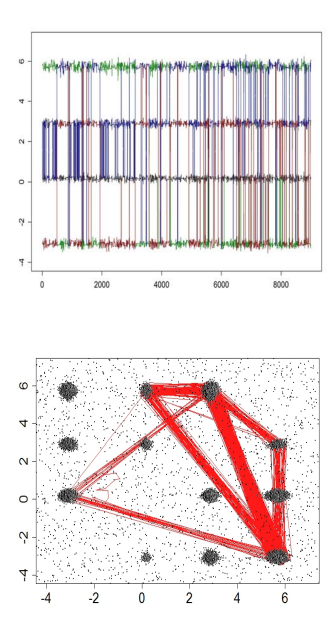


Figure 1: Top panel: Trace plots generated using 9,000 samples obtained for μ_1, \ldots, μ_4 (the first 1,000 samples have been discarded) from one of the IMTM-TA1 chains. Bottom panel: The dots represent the projection of the values sampled by the IMTM-TA population of chains on the (μ_i, μ_j) planes, with $i \neq j$. The trajectory of one of the chains is projected in red on the plane (μ_1, μ_2) .

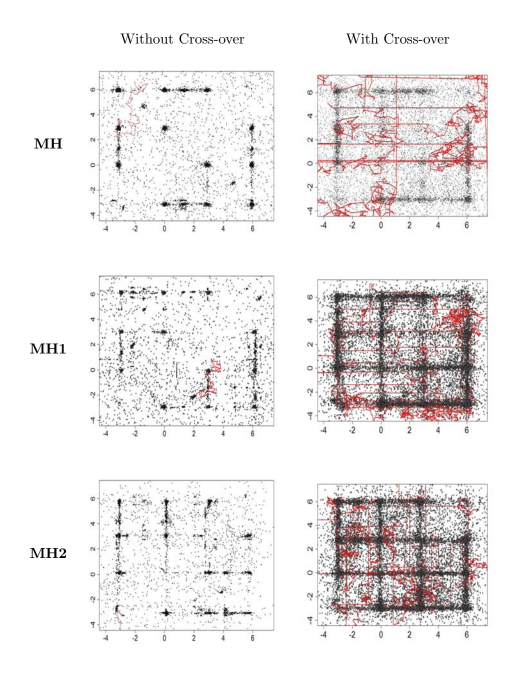


Figure 2: The dots represent the projection of the values sampled by the population of MH chains considered in the simulation on the (μ_i, μ_j) planes, with $i \neq j$. The plots illustrate the samples obtained without cross-over (left column) and with cross-over (right column) for the MH (top row), MH1 (middle row) and MH2 (bottom row). The trajectory of one of the chains is shown in red.

one can glance the superiority of the IMTM-TA algorithm by looking at one of the paths shown in the bottom panel of Figure 1 where it can be seen that the chain visits often many modes of the posterior distribution.

The efficiency improvement is also noticeable from the autocorrelation functions (ACF) shown in Figure 3. For each method included in the comparison, the curves shown are obtained by averaging the ACF estimates over the N chains of the population and over 10 replicates. The MH with cross-over are more efficient then the parallel MH algorithms but still less efficient than the IMTM algorithms.

The results in Table 1 show that the IMTM algorithms are generally able to produce more efficient estimates than the MH class of algorithms considered in the comparison. Given the plots described above, it is not surprising that by adding the cross-over moves brings the efficiency of the MH, MH1 and MH2 closer to that of the IMTM samplers, especially when the number of parallel chains is large (N=100). However when we reduce the number of chains (e.g. N = 20) the performance of the MH algorithms (with and without cross-over) is clearly inferior to that of the IMTM algorithms. Interestingly, the estimation results of the IMTM-TA and IMTM-IS perform similarly whether we choose to adapt the weights λ_i or not.

4.1.1 Comparison in the Presence of Annealing

The performance of the MH, MH1 and MH2 populations can be improved by combining them with an annealing procedure. Our interest, here, lies in comparing AIMTM with the algorithms MH, MH1, MH2 which are modified to incorporate an annealing-based strategy. We consider once again two variants of the AIMTM defined by the choice of weights λ_j . Specifically, we consider AIMTM-TA and AIMTM-IS as Algorithm 4 with, respectively, the same λ 's as IMTM-TA and IMTM-IS.

We also consider the uniform, logarithmic and power tempering schemes that were also suggested by Jasra et al. (2007), and are defined as:

$$\xi_{i} = \xi_{i-1} - \frac{1}{N}$$

$$\xi_{i} = \log(\xi_{i-1} + 1) / \log(Q), \quad Q > 0$$

$$\xi_{i} = (\xi_{i-1} - Q)^{\psi}, \quad \psi > 0, \ Q \in (0, 1)$$

for i = 2, ..., N with $\xi_1 = 1$. The three tempering schemes will be denoted with M1, M2 and M3 in what follows. For the MH algorithms we will build chain *i* ergodic to π^{ξ_i} and construct different scales for the chains of

	N=100				N=20					
	1	2	3	4	MSE	1	2	3	4	MSE
MH	0.81	0.42	2.08	1.06	18.83	0.39	0.69	0.67	2.28	26.76
	(4.22)	(4.37)	(4.39)	(4.10)		(5.35)	(5.16)	(6.02)	(3.15)	
MH1	0.72	0.21	0.62	0.91	5.42	0.10	0.17	0.66	0.78	7.35
	(2.12)	(2.09)	(2.14)	(2.19)		(2.47)	(1.89)	(2.49)	(2.91)	
MH2	0.99	1.89	1.47	1.01	3.30	0.11	2.80	0.42	0.37	5.09
	(1.57)	(1.73)	(1.87)	(1.89)		(1.99)	(1.71)	(1.98)	(1.85)	
MH c.o.	1.87	1.09	1.91	1.66	7.89	1.74	1.11	1.01	1.75	11.02
	(2.52)			(2.92)						
MH1 c.o.	0.65	0.21	1.59	1.46	2.77	0.51	0.22	1.83	1.12	3.51
	(1.86)	(1.35)	(1.24)	(1.35)		(1.48)	(1.91)	(1.27)	(1.91)	
MH2 c.o.	1.11	1.69	1.27	1.26	2.17	0.59	1.68	0.97	1.14	2.26
	(1.33)	(1.34)	(1.76)	(1.29)		(1.43)	(1.16)	(1.36)	(1.58)	
IMTM-IS	1.40	1.52	1.37	1.42	1.05	1.36	1.39	1.61	1.69	1.42
	(1.01)	(0.98)	(1.22)	(0.87)		(0.98)	(1.20)	(1.12)	(1.42)	
IMTM-IS-a	1.37	1.44	1.58	1.54	0.49	1.31	1.71	1.35	1.72	1.18
	(0.83)	(0.56)	(0.71)	(0.64)		(0.81)	(0.97)	(1.23)	(1.24)	
IMTM-TA	1.31	1.46	1.53	1.61	0.52	1.29	1.21	1.70	1.32	0.89
	(0.38)	(1.06)	(0.48)	(0.73)		(1.34)	(1.05)	(0.31)	(0.59)	
IMTM-TA-a	1.56	1.39	1.60	1.37	0.47	1.63	1.75	1.61	1.44	0.85
	(0.48)	(0.91)	(0.76)	(0.42)		(0.76)	(0.86)	(1.02)	(0.97)	

Table 1: Estimates of μ_1, \ldots, μ_4 and its standard deviations (in parenthesis). For the parallel MHs, without and with cross-over (c.o.), we considered alternatively N = 20 and N = 100 chains and T = 100,000 iterations. For the IMTMs, without and with adaptation (IMTM-TA-a and IMTM-IS-a), we consider N = 100 chains T = 10,000 iterations and J = 10 different proposals (selected randomly within the other chains of the population). Values results from average over 10 runs of the different algorithms. For expository purposes, we report, for each algorithm and the different population sizes and the Mean Square Error (MSE), averaged over the parameters, with respect to the theoretical value that is 1.5.

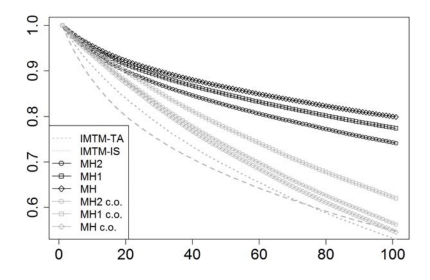


Figure 3: Autocorrelation functions for the methods considered. The curves are obtained by averaging over the population of chains used for each algorithm and over the 10 replicated runs of each algorithm.

the population as in Jasra et al. (2007). For the *i*-th chain the proposal variance $\sigma_i = \sigma_1/(1 + \gamma_i)$ with $\sigma_1 = 0.5$.

For the logarithmic scheme M2 we consider Q = 2.25 and for the power scheme M3 we set Q = 0.001 and $\psi = 3/2$ as suggested in Jasra et al. (2007).

We report the estimates for each mean μ_i in in Table 2. The AE column shows the maximum bias (over the four means). One can see easily that, on average, the AIMT yields the smallest errors within each tempering scheme. Note that the results are not directly comparable with the ones in Table 1 because in the experiments without tempering all the chains of the population have the same target and have been used in order to estimate the parameters of the mixture. In the experiments with the different tempering schemes we consider, for each algorithm, the output of the chain with $\xi_i = 1$, which has the target π . The results in Table 2 show that the AIMTM algorithms outperform the population of MH, MH1 and MH2 chains for all the three different tempering schemes. The logarithm and power decay schemes seem to give the best result when combined with the AIMTM.

			M1		
	1	2	3	4	AE
MH	1.81	0.73	1.02	1.79	0.97
MH1	0.64	1.62	0.91	1.59	0.86
MH2	0.81	1.75	1.12	1.99	0.69
AIMTM1-IS	1.83	1.43	1.98	1.37	0.48
AIMTM1-TA	0.89	1.15	1.92	1.81	0.61
			M2		
	1	2	3	4	AE
MH	0.84	0.72	1.41	0.93	0.78
MH1	1.67	1.57	1.06	1.84	0.54
MH2	1.71	1.32	1.52	1.01	0.49
AIMTM1-IS	1.44	1.91	1.37	1.26	0.41
AIMTM1-TA	1.86	1.19	1.51	1.49	0.36
			M3		
	1	2	3	4	AE
MH	0.82	1.25	0.83	0.97	0.68
MH1	1.79	1.42	1.33	0.98	0.52
MH2	0.99	1.27	1.63	1.69	0.51
AIMTM1-IS	1.19	1.97	1.16	1.12	0.47
AIMTM1-TA	1.37	1.04	1.86	1.77	0.46

Table 2: Estimates of μ_1, \ldots, μ_4 . For the MHs we considered N = 100 chains and T = 100,000 iterations. For the AIMTM1 we consider N = 100 chains T = 10,000 iterations and J = 10 proposals. For expository purposes, we report the maximum absolute bias (AE) for each algorithm and tempering scheme.

The gain in efficiency with respect to the populations of MH-type algorithms is evident also from the ACF functions presented in Figure 4). The ACF have been obtained by averaging over 10 independent runs of the algorithms considered in the comparison.

4.1.2 Multivariate Normal Mixture

We compare, for high-dimensional target distribution, the population Monte Carlo algorithms (MHs with cross-over and IMTM-TA) described in the previous section. The target considered for the comparison is the following multivariate mixture of two normals with a sparse variance-covariance structure

$$\frac{1}{3}\mathcal{N}_{20}(\mu_1, \Sigma_1) + \frac{2}{3}\mathcal{N}_{20}(\mu_2, \Sigma_2)$$
(6)

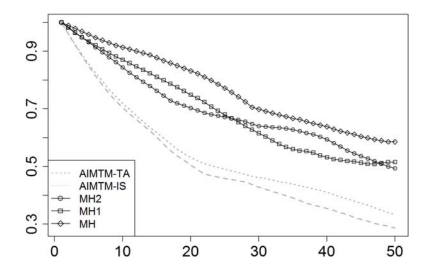


Figure 4: Autocorrelation functions obtained by averaging over 10 independent runs of each algorithm for the population of MH and AMTM chains.

with $\boldsymbol{\mu}_1 = (3, \ldots, 3)'$, $\boldsymbol{\mu}_2 = (10, \ldots, 10)'$ and Σ_j , with j = 1, 2, generated independently from a Wishart distribution $\Sigma_j \sim \mathcal{W}_{20}(\nu, Id_{20})$ where ν is the degrees of freedom parameter of the Wishart. In the experiments we set $\nu = 21$.

In the comparison we considered N = 20 chains and T = 100,000 iterations for the parallel MHs, with cross-over and N = 20 chains, T = 10,000iterations and M = 10 for the IMTM-TA algorithm. The proposal distributions for the *i*th chain, $T_j^{(i)}(\mathbf{y}|\mathbf{x}_n^{(i)})$, is Gaussian with variance-covariance matrix $\Lambda_i = (0.1 + 5i)\mathbf{I}_{20}$ for all $j = 1, \ldots, M$. For the proposals of chains in the MH1 and MH2 populations we consider Gaussian random walk with scales in the same range of the IMTM proposal scales.

The autocorrelation function of the chains (average over 20 dimensions of each chain, the different chains of the population and the 10 different runs of the population Monte Carlo algorithms) is given in Fig. 5. The values of the ACF for the IMTM (see Fig. 5) are less than those for the MH with cross-over. We conclude that in this example the population of MTM chains outperforms, in terms of estimation efficiency, the populations of MHs with

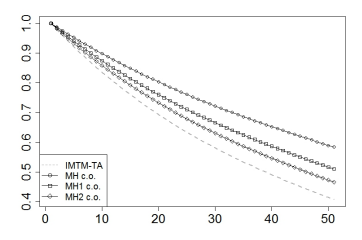


Figure 5: Autocorrelation function (ACF) for MH with cross-over and IMTM classes of algorithms. The ACF result from an average over the 20 components of the multivariate chain, the different chains of the population and over 10 replicates.

cross-over.

4.2 Stochastic Volatility

The estimation of the stochastic volatility (SV) model due to Taylor (1994) still represents a challenging issue in both off-line (Celeux et al. (2006)) and sequential (Casarin and Marin (2009)) inference contexts. The first challenging issue in inference for SV models is related to the nonlinear structure of the model which makes parameter estimation difficult. Another main difficulty is due to the high dimension of the sampling space which hinders the use of the data-augmentation and prevents a reliable joint estimation of the parameters and the latent variables. As highlighted in Casarin et al. (2009) using multiple chains with a chain interaction mechanism could lead to a substantial improvement in the MCMC method for this kind of model.

We consider the SV model given in Celeux et al. (2006)

$$\begin{array}{rcl} y_t | h_t & \sim & \mathcal{N}\left(0, e^{h_t}\right) \\ h_t | h_{t-1}, \boldsymbol{\theta} & \sim & \mathcal{N}\left(\alpha + \phi h_{t-1}, \sigma^2\right) \\ h_0 | \boldsymbol{\theta} & \sim & \mathcal{N}\left(0, \sigma^2/(1 - \phi^2)\right) \end{array}$$

with t = 1, ..., T and $\boldsymbol{\theta} = (\alpha, \phi, \sigma^2)$. For the parameters we assume the noninformative prior (see Celeux et al., 2006)

$$\pi(\boldsymbol{\theta}) \propto 1/(\sigma\beta)\mathbb{I}_{(-1,1)}(\phi)$$

where $\beta^2 = \exp(\alpha)$. In order to simulate from the posterior we consider the full conditional distributions and apply a Gibbs algorithm. If we define $\mathbf{y} = (y_1, \ldots, y_T)$ and $\mathbf{h} = (h_0, \ldots, h_T)$ then the full conditionals for β and ϕ are the inverse gamma distributions

$$\beta^{2}|\mathbf{h}, \mathbf{y} \sim \mathcal{IG}\left((T-1)/2, \sum_{t=1}^{T} y_{t}^{2} \exp(-h_{t})/2\right)$$
$$\sigma^{2}|\phi, \mathbf{h}, \mathbf{y} \sim \mathcal{IG}\left((T-1)/2, \sum_{t=2}^{T} (h_{t} - \phi h_{t-1})^{2}/2 + h_{1}^{2}(1 - \phi^{2})\right)$$

and ϕ and the latent variables have non-standard full conditionals

$$\pi(\phi|\sigma^{2}, \mathbf{h}, \mathbf{y}) \propto (1 - \phi^{2})^{1/2} \exp\left(-\frac{\phi^{2}}{2\sigma^{2}} \sum_{t=2}^{T-1} h_{t}^{2} - \frac{\phi}{\sigma^{2}} \sum_{t=2}^{T} h_{t} h_{t-1}\right) \mathbb{I}_{(-1,+1)}(\phi)$$

$$\pi(h_{t}|\alpha, \phi, \sigma^{2}, \mathbf{h}, \mathbf{y}) \propto \exp\left(-\frac{1}{2\sigma^{2}} \left((h_{t} - \alpha - \phi h_{t-1})^{2} - (h_{t+1} - \alpha - \phi h_{t})^{2}\right) - \frac{1}{2} \left(h_{t} + y_{t}^{2} \exp(-h_{t})\right)\right).$$

In order to sample from the posterior we use the IMTM-IS within Gibbs algorithm. More specifically, in the IMTM step for ϕ , we follow Celeux et al. (2006), and use as proposal, a truncated normal distribution on (-1, 1) with mean and variance

$$\sum_{t=2}^{T} h_t h_{t-1} \Big/ \sum_{t=2}^{T-1} h_t^2 \quad \text{and} \quad \sigma^2 \Big/ \sum_{t=1}^{T-1} y_t^2$$

One of the most difficult issues is related to the choice of the proposal distribution for h_t . In this paper we follow a standard approach based

on the second-order Taylor approximation of the term $\exp\{h_t\}$, in the full conditional of h_t , around the mean μ_t of the distribution of $h_t|h_{t-1}, \phi, \sigma^2$. The approach has been introduced by Shephard and Pitt (1997) and has been adapted to the context of iterated importance sampling by Celeux et al. (2006). The proposal distribution for h_1 is a normal with mean

$$\frac{\phi h_2 \sigma^{-2} + 0.5 \exp(-\phi h_2) y_1^2 (1+\phi h_2) \beta^{-2} - 0.5}{\sigma^{-2} + 0.5 \exp(-\phi h_2) y_1^2 \beta^{-2}}$$

and variance $(\sigma^{-2} + 0.5 \exp(-\phi h_2)y_1^2\beta^{-2})^{-1}$. The proposal for h_t with $t = 2, \ldots, T-1$ is a normal with mean

$$\frac{(1+\phi^2)\mu_t\sigma^{-2}+0.5\exp(-\mu_t)y_t^2(1+\mu_t)\beta^{-2}-0.5}{(1+\phi^2)\sigma^{-2}+0.5\exp(-\mu_t)y_t^2\beta^{-2}}$$

and variance $((1 + \phi^2)\sigma^{-2} + 0.5 \exp(-\mu_t)y_t^2\beta^{-2})^{-1}$. The proposal for h_T is a normal with mean

$$\frac{\phi h_{T-1} \sigma^{-2} + 0.5 \exp(-\phi h_{T-1}) y_T^2 (1 + \phi h_{T-1}) \beta^{-2} - 0.5}{\sigma^{-2} + 0.5 \exp(-\phi h_{T-1}) y_T^2 \beta^{-2}}$$

and variance $(\sigma^{-2} + 0.5 \exp(-\phi h_{T-1})y_T^2 \beta^{-2})^{-1}$.

The single-move Gibbs sampler updates sequentially the latent variables and this makes the classic hybrid Metropolis within Gibbs algorithm inefficient. A possible remedy (see Shephard and Pitt , 1997) consists in simulating jointly some blocks of the latent variables (blocking). Our IMTM algorithms can be extended to consider blocking procedure. In our experiments we did not find efficiency improvements in applying blocking techniques to our simulated datasets. Moreover, the main goal of our simulation study is to demonstrate the IMTM algorithm's ability to break down the dependence in the single-move sampler and thus to improve the efficiency of the Monte Carlo sample.

In the simulation experiments we consider the two parameter settings $(\alpha, \phi, \sigma^2) = (0, 0.99, 0.01)$ and $(\alpha, \phi, \sigma^2) = (0, 0.9, 0.1)$ which correspond, in a financial stock market context, to daily and weekly frequency data respectively. Note that as reported in Casarin and Marin (2009) inference in the daily example is more difficult. We compare the the IMTM within Gibbs algorithms with a population of MH within Gibbs in terms of Mean Square Error (MSE) for the parameters and of cumulative RMSE for the latent variables. We carry out the comparison through the MSE and the SD by running the algorithms on 20 independent simulated datasets of 200 observations. In the comparison we take into account the computational cost and for the population of MH we consider N = 20 chains and T = 50,000iterations and for the IMTM-IS within Gibbs we use N = 20 interacting chains, each with M = 5 proposals, and T = 10,000 iterations.

The results for the parameter estimation when applying IMTM-IS are presented in Table 3 and show an effective improvement in the estimates, both for weekly and daily data, when compared to the results of a MH algorithm with an equivalent computational load.

		Daily Data				Weekly Data			
θ	Value	MSE			Value	MSE			
		IMTM-IS	MH			IMTM-IS	MH		
α	0	0.03018	0.07392	α	0	0.00202	0.00597		
		(0.00583)	(0.00201)			(0.00179)	(0.00139)		
ϕ	0.99	0.19853	0.29871	ϕ	0.9	0.01512	0.08183		
		(0.02038)	(0.04423)			(0.03920)	(0.04011)		
σ^2	2 0.01	0.00204	0.01373	σ^2	0.1	0.00892	0.07405		
		(0.00241)	(0.00191)			(0.00201)	(0.00293)		

Table 3: Mean square error (MSE) and standard deviation (in parenthesis) for the parameter estimation with IMTM-IS and MH within Gibbs algorithms. Left panel: daily datasets. Right panel: weekly dataset.

We show here that an approach based on the use of multiple-try interacting chains can also break-down the dependence structure in the output of the sampler thus improving the efficiency of the posterior simulation for the latent variables.

More specifically Figure 6 exhibit the estimated ACFs for the MH (black lines) and MTM (gray lines) class of algorithms, for the 200 components associated to the latent process $\{h_t\}_{t=1,...,T}$ with T = 200. The ACF for each latent variable h_t results from the average over the chains of the MH and MTM populations and over 10 independent run of each algorithm on the same set of simulated data. In Figure 6, the top panel shows the results for the daily dataset and the bottom panel for the weekly dataset. In both daily and weekly setups the IMTM-IS overperforms in terms of estimation efficiency the population of parallel MH.

Note that these results is similar to the results obtained for SV models in Celeux et al. (2006), Casarin and Marin (2009) and Casarin et al. (2009) for population Monte Carlo algorithms. We can conclude that the IMTM shares similar properties of other population Monte Carlo algorithms with the advantage that the convergence of the algorithm relies upon the detail balance condition and no further theoretical results are needed. We compare our IMTM with MH and left for future research a comparison with importance sampling based methods such as the Popopulation Monte Carlo methods or Sequential Monte Carlo methods described in Jasra et al. (2007).

Figure 7 show the HPD region at the 90% (gray areas) and the mean (black lines) of the cumulative RMSE of each algorithm for the weekly (top) and daily data (bottom panel). The statistics have been estimated from 20 independent experiments. The average RMSE shows that, in both parameter settings considered here, the IMTM (dashed black line) is more efficient than the standard MH algorithm (solid black line).

4.3 Loss of Heterozigosity Application

We consider here the problem of the genetic instability of esophageal cancers. During a neoplastic progression the cancer cells undergo a number of genetic changes and possibly lose entire chromosome sections. The loss of a chromosome section containing one allele by abnormal cells is called *Loss* of *Heterozygosity* (LOH). The LOH can be detected using laboratory assays on patients with two different alleles for a particular gene. Chromosome regions containing genes which regulate cell behavior, are hypothesized to have a high rates of LOH. Consequently the loss of these chromosome sections disables important cellular controls.

Chromosome regions with high rates of LOH are hypothesized to contain *Tumor Suppressor Genes* (TSGs), whose deactivation contributes to the development of esophageal cancer. Moreover the neoplastic progression is thought to produce a high level of background LOH in all chromosome regions.

In order to discriminate between "background" and TSGs LOH, the Seattle Barrett's Esophagus research project (Barrett et al. (1996)) has collected LOH rates from esophageal cancers for 40 regions, each on a distinct chromosome arm. The labeling of the two groups is unknown so Desai (2000) suggest to consider a mixture model for the frequency of LOH in both the "background" and TSG groups.

We consider the hierarchical Beta-Binomial mixture model proposed in

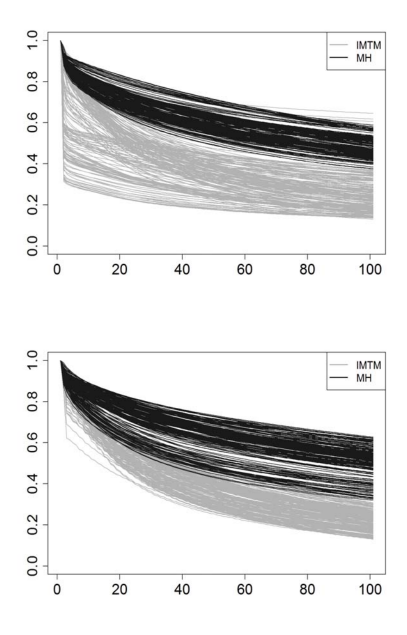


Figure 6: ACF of the population of MH and MTM chains for the 200 components associated to the latent process $\{h_t\}_{t=1,\dots,T}$, for daily (top) and weekly (bottom) datasets. The ACFs are averaged over the different chains of the population.

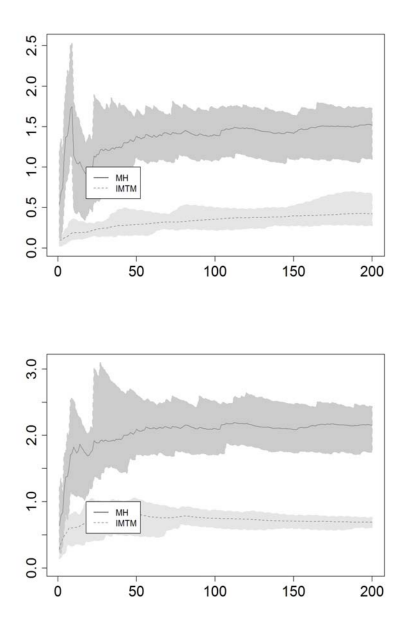


Figure 7: Cumulative RMSE for the IMTM (dashed line) and MH (solid line) and the 90% HPD regions for the IMTM (light gray) and MH (dark gray) estimated on 20 independent experiments for both the daily (top) and weekly (bottom) datasets.

Warnes (2001)

$$f(x, n|\eta, \pi_1, \pi_2, \gamma) = \eta \binom{n}{x} \pi_1^x (1 - \pi_1)^{n-x} +$$
(7)
(1 - η) $\binom{n}{x} \frac{\Gamma(1/\omega_2)}{\Gamma(\pi_2/\omega_2)\Gamma((1 - \pi_2)/\omega_2)} \frac{\Gamma(x + \pi_2/\omega_2)\Gamma(n - x + (1 - \pi_2)/\omega_2)}{\Gamma(n + 1/\omega_2)}$

with x number of LOH sections, n the number of examined sections, $\omega_2 = \exp\{\gamma\}/(2(1 + \exp\{\gamma\}))$. Let $\mathbf{x} = (x_1, \ldots, x_m)$ and $\mathbf{n} = (n_1, \ldots, n_m)$ be a set of observations from $f(x, n | \eta, \pi_1, \pi_2, \gamma)$ and let us assume the following priors

$$\eta \sim \mathcal{U}[0,1], \quad \pi_1 \sim \mathcal{U}[0,1], \quad \pi_2 \sim \mathcal{U}[0,1] \quad \text{and} \quad \gamma \sim \mathcal{U}[-30,30]$$
(8)

with \mathcal{U} the uniform distribution on [a, b]. Then the posterior distribution is

$$\pi(\eta, \pi_1, \pi_2, \gamma | \mathbf{x}, \mathbf{n}) \propto \prod_{j=1}^m f(x_j, n_j | \eta, \pi_1, \pi_2, \gamma)$$
(9)

The parametric space is of dimension four: $(\eta, \pi_1, \pi_2, \gamma) \in [0, 1]^3 \times [-30, 30]$ and the posterior distribution has two well-separated modes making it difficult to sample using generic methods.

We apply the IMTM-IS algorithm M = 4 proposal functions selected between a population of N = 100 chains. The values of the population of chains (dots) at the last iteration on the subspace (π_1, π_2) is given in Figure 8. The IMTM-IS is able to visit both regions of the parameter space confirming the analysis of Craiu et al. (2009) and Warnes (2001).

5 Conclusions

In this paper we propose a new class of interacting multiple-try Metropolis algorithms that extends the existing literature in two directions. First , the multiple try transition has been extended to allow the use of different proposal distribution and second, we propose a new interacting Monte Carlo algorithm for increasing the efficiency of MTM. We give a proof of validity of the algorithm and show on real and simulated examples the effective improvement in the mixing property and exploration ability of the resulting interacting chains. We note here that the use of antithetic and stratified sampling discussed by Craiu and Lemieux (2007) can be extended naturally to the IMTM sampler. Future work will focus on building stronger ties between IMTM and the emerging area of adaptive MCMC.

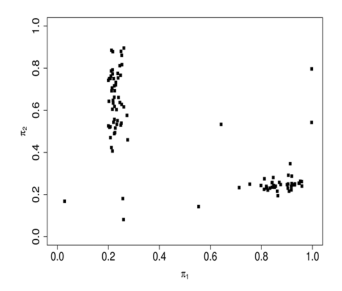


Figure 8: Values of the population of chains (dots) at the last iteration on the subspace (π_1, π_2) . The interaction is given by M = 4 proposal functions randomly selected between the population of N = 100 chains.

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Appendix A

Proof

Without loss of generality, we can set $M_i = N$, $\forall i$ and $x_n^{(i)} = x$. Fixed the *i*-th chain, the conditional detailed balance is proved. This ensures the ergodicity of the chain.

Following the notations in Algorithm 3, let us define the following quan-

tities

$$\bar{w}^{(i)}(y_{1:N}|x) = \sum_{j=1}^{N} w_j^{(i)}(y_j, x), \quad \bar{w}_{-k}^{(i)}(y_{1:N}|x) = \sum_{j \neq k}^{N} w_j^{(i)}(y_j, x)$$

and

$$S_N(J) = \frac{1}{\bar{w}^{(i)}(y_{1:N}|x)} \sum_{j=1}^N \delta_j(J) w_j^{(i)}(y_j,x)$$

with $J \in \mathcal{J} = \{1, \ldots, N\}$ the empirical measure generated by different pro-

posals and by the normalized selection weights. Let $T^{(i)}(dy_{1:N} | x) = \bigotimes_{j=1}^{N} T_j^{(i)}(dy_j | \tilde{f}_n^{(i)}(x))$ the joint proposal for the multiple try and define $T_{-k}^{(i)}(dy_{1:N} | x) = \bigotimes_{j \neq k}^{N} T_j^{(i)}(dy_j | \tilde{f}_n^{(i)}(x))$. Let A(x, y) be the actual transition probability for moving from x to y in the IMTM (Algorithm 3). Suppose that $x \neq y$, then the transition is a results two steps. The first step is a selection step which can be written as $y = y_J$ and $x_J^* = x$ with the random index J sampled from the empirical measure $S_N(J)$. The second step is a accept/reject step based on the generalized MH ratio which involves the generation of the auxiliary values x_j^* for $j \neq J$. Then

$$\begin{split} &\pi(x)A(x,y) = \\ &= \pi(x)\int_{\mathcal{Y}^{N}} T^{(i)}(dy_{1:N} \mid x) \int_{\mathcal{J}} S_{N}(dJ) \int_{\mathcal{Y}^{N-1} \times \mathcal{Y}^{2}} T^{(i)}_{-J}(dx_{1:N}^{*} \mid y) \times \\ &\quad \times \delta_{x}(dx_{J}^{*})\delta_{y_{J}}(dy) \min\left\{1, \frac{\bar{w}^{(i)}(y_{1:N} \mid x)}{\bar{w}^{(i)}(x_{1:N}^{*} \mid y)}\right\} \\ &= \pi(x)\sum_{j=1}^{N} \int_{\mathcal{Y}^{N-1}} T^{(i)}_{-j}(dy_{1:N} \mid x) T^{(i)}_{j}(y \mid \tilde{f}^{(i)}_{n}(x)) \int_{\mathcal{Y}^{N-1}} T^{(i)}_{-j}(dx_{1:N}^{*} \mid y) \times \\ &\quad \times \frac{w_{j}^{(i)}(y, x)}{w_{j}^{(i)}(y, x) + \bar{w}^{(i)}_{-j}(y_{1:N} \mid x)} \min\left\{1, \frac{w_{j}^{(i)}(y, x) + \bar{w}^{(i)}_{-j}(y_{1:N} \mid x)}{w_{j}^{(i)}(x, y) + \bar{w}^{(i)}_{-j}(x_{1:N}^{*} \mid y)}\right\} \\ &= \sum_{j=1}^{N} \frac{w_{j}^{(i)}(x, y)w_{j}^{(i)}(y, x)}{\lambda_{j}^{(i)}(y, x)} \int_{\mathcal{Y}^{2(N-1)}} T^{(i)}_{-j}(dy_{1:N} \mid x) \times \\ &\quad \times T^{(i)}_{-j}(dx_{1:N}^{*} \mid y) \min\left\{\frac{1}{w_{j}^{(i)}(y, x) + \bar{w}^{(i)}_{-j}(y_{1:N} \mid x)}, \frac{1}{w_{j}^{(i)}(x, y) + \bar{w}^{(i)}_{-j}(x_{1:N}^{*} \mid y)}\right\} \end{split}$$

which is symmetric in x and y.

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