

# Synchronous and Asynchronous Parallel Simulated Annealing with Multiple Markov Chains

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**Abstract**—Simulated annealing is a general-purpose optimization technique capable of finding an optimal or near-optimal solution in various applications. However, the long execution time required for a good quality solution has been a major drawback in practice. Extensive studies have been carried out to develop parallel algorithms for simulated annealing. Most of them were not very successful, mainly because multiple processing elements (PEs) were required to follow a single Markov chain and, therefore, only a limited parallelism was exploited. In this paper, we propose new parallel simulated annealing algorithms which allow multiple Markov chains to be traced simultaneously by PEs which may communicate with each other. We have considered both synchronous and asynchronous implementations of the algorithms. Their performance has been analyzed in detail and also verified by extensive experimental results. It has been shown that for graph partitioning the proposed parallel simulated annealing schemes can find a solution of equivalent (or even better) quality up to an order of magnitude faster than the conventional parallel schemes. Among the proposed schemes, the one where PEs exchange information dynamically (not with a fixed period) performs best.

**Index Terms**—Asynchronous communication, graph partitioning, multiple Markov chains, parallel algorithm, simulated annealing, solution quality, speed-up.

## 1 INTRODUCTION

**S**IMULATED annealing (SA) is an iterative probabilistic algorithm which combines local search with Monte Carlo techniques [1]. Numerous researchers have demonstrated that SA can be very effective in many optimization problems such as TSP, placement and routing in VLSI design, logic minimization, code design and image processing, etc. One of the distinct features of the SA is its hill-climbing capability which allows escape from local optima. Also, it may start from any initial solution and converges to an optimal or near optimal solution [2]. In addition, SA is applicable to various classes of problems including discrete, nondifferentiable, or combinatorial problems, etc. However, the long execution time of SA has been the major drawback in practice.

There have been numerous efforts to make SA practical for problems of realistic size by shortening the execution time. These efforts can be classified into two major categories: algorithmic optimization (of sequential SA) and parallel processing.

Two types of approaches are prevalent in the category of optimizing SA: careful perturbation and cooling. In the approach of careful perturbation, the idea of range-limiting was introduced and successfully applied to function minimization and VLSI design [3], [4]. Since the execution time is proportional to the total number of trials, a heuristic

perturbation scheme may shorten the execution time by reducing the chance of rejection [3], [5]. In the other approach, most schemes utilize statistical quantities such as mean and variance of costs obtained during annealing. They usually modify the conventional annealing schedule to accelerate the execution of SA. However, it has been shown that modifying the annealing schedule does not improve both the solution quality and execution time [6], [7]. The execution time was reduced by a factor of 2 at best.

The advent of parallel computers has made it possible to achieve speed-up of orders of magnitude in various applications. Most parallel SA schemes previously proposed share the feature that multiple PEs follow a single search path (Markov chain) [8], [9], [10], [11], [12], [13] (to be referred to as SMC PSA). In SMC PSA, a perturbation at each PE and/or evaluation of the perturbation is executed in parallel, and then only one global solution is accepted in each iteration. While this approach preserves the sequential nature of SA, following a single search path might be an unnecessary restriction especially from the viewpoint of performance, i.e., the execution time and solution quality. That is, only a limited parallelism is exploited, which is believed to be the main reason why most of the previous PSAs have not been very successful.

Banerjee et al. implemented SMC PSA schemes for a standard cell placement problem on hypercube multiprocessors. The PSA schemes are based on a sequential version of SA (TimberWolf). The task (SA) is spatially partitioned and non-exact calculations by multiple PEs are allowed by assuming temporary locality. They concluded that such erroneous calculations can achieve a considerable speed-up without seriously affecting convergence [10]. However, this may be acceptable

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for particular applications only when a periodic synchronization is incorporated to compensate for the erroneous calculations. Consequently, either the synchronization overhead may increase the execution time or the erroneous calculations may deteriorate the solution quality. Also, a general purpose PSA scheme which takes advantage of the acceptance rate but follows a single Markov chain was proposed by Roussel-Ragot and Dreyfus. Besides the drawbacks due to the single Markov chain, their synchronization strategy using the acceptance rate does not appear to be very efficient [11].

The feasibility of solving combinatorial problems on a Boltzmann machine was investigated by Aarts and Korst [15]. However, it has been shown by computer simulation that the performance characteristics (convergence and speed) strongly depend on the type of problem. Also, a practical implementation on a massively parallel machine resulted in very small speed-up for placement problems [16]. These facts imply that the implementation of SA on a massively parallel machine is not yet suitable in practice due to its sequential and random natures.

In contrast to the abundance of SMC PSA schemes, there are very few schemes where PEs are allowed to follow multiple Markov chains (to be referred to as MMC PSA). Aarts and Korst described the idea of the MMC PSA, namely the *division algorithm*. However, it does not seem to have been fully developed. The idle and communication times which are inevitable in practice are not taken into account. In addition, PEs may exchange the information at non-quasi-equilibrium state in the proposed algorithm, which may not lead to an optimal solution [7].

In this paper, we propose a new MMC PSA which can overcome the potential drawbacks of the previous PSA schemes. One of the distinct features of our scheme is that PEs (following multiple Markov chains) communicate with each other dynamically, not with a fixed period. This yields a significant savings in communication time by allowing PEs to exchange only useful information. Also, it is shown that the information exchange can be implemented asynchronously in order to further improve performance of the proposed MMC PSA. The goals of the proposed PSA are

- 1) to achieve higher speed-up and efficiency,
- 2) to obtain a good solution quality (at least as good as that of the sequential SA), and
- 3) to realize problem independent performance.

This paper is organized as follows. In Section 2, the new MMC PSA and its asynchronous version are described, following our versions of SMC PSAs. In Section 3, The SMC and MMC PSAs are compared through theoretical analysis. The implementation results for graph partitioning are provided with discussions in Section 4, followed by conclusions in Section 5.

## 2 PARALLELIZATION OF SIMULATED ANNEALING

### 2.1 Simulated Annealing

SA, also called *statistical cooling*, has many attractive features including the convergence to a high quality solution, versatility, and ease of implementation. However, SA requires a careful setup of the basic strategies and parameters, e.g.,

- 1) how to define the configuration space and an appropriate cost function,
- 2) how to implement an efficient perturbation scheme and
- 3) how to design the cooling schedule including *initial temperature*, a rule for temperature decrement, *quasi-equilibrium conditions* for a temperature, and a *stop criterion*.

A typical structure of SA consists of two nested loops as shown in Fig. 1. It starts from an arbitrarily selected configuration  $s_0$  with an appropriate initial temperature ( $T_i$ ) and works to minimize a given *cost function*. At a fixed temperature, the inner loop repeatedly executes the following three step operation, to be referred to as *iteration*, until an inner loop break condition is satisfied. It randomly perturbs the current solution (or configuration), evaluates the corresponding cost, and accepts the new solution with the probability of  $\min(1, e^{-\frac{\Delta C}{T}})$  where  $\Delta C$  is the cost change due to the perturbation and  $T$  is the current temperature. The outer loop decreases temperature according to the rule,  $T \leftarrow \alpha T$ , where  $\alpha$ , the *cooling coefficient*, satisfies  $0 < \alpha < 1$ . It can be said that SA consists of a sequential chain of consecutive perturbation, evaluation and decision steps.

```

s ← s0;
T ← Ti;
While the outer loop break condition is not met do {
  While the inner loop break condition is not met do {
    Perturb current configuration s to s*;
    Evaluate cost function and find the difference;
    ΔC = C(s*) - C(s);
    Accept new configuration with probability min(1, e-ΔC/T);
  }
  T ← α T;
}

```

Fig. 1. Typical structure of simulated annealing.

Those parameters mentioned above, which control the execution of the nested loops are called *scheduling parameters*, i.e., *initial temperature* ( $T_i$ ), *cooling coefficient* ( $\alpha$ ), and *equilibrium conditions* (also known as *break conditions*) for the inner and outer loops. The execution time and solution quality are heavily dependent on the scheduling parameters. In the following, we will first describe our versions of SMC PSAs, mainly for comparison purposes, and then present the proposed MMC PSA. In evaluating performance of a PSA, we need to consider solution quality as well as execution speed. The execution speed may be quantified in terms of *speed-up* ( $S$ ) and *efficiency* ( $E$ ). The speed-up ( $S$ ) is defined as the ratio of the execution time (on one PE) by the sequential SA to that by the PSA (on  $N$  PEs) for an equivalent solution quality. In the ideal case,  $S$  would be equal to  $N$ . Efficiency ( $E$ ) is defined as the ratio of the actual speed-up to the ideal speed-up ( $N$ ).

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## 2.2 Single Markov Chain PSA

Let's define a couple of terms first. A *move* is defined to be a local disturbance of the solution (configuration), in which two PEs are involved (refer to Section 4.2 for examples). Note that a *perturbation* may result from more than one *move* by PEs, i.e., from a set of concurrent moves.

In SMC PSA, local moves at PEs which are involved in a perturbation form one global perturbation of configuration. The difference in the cost due to the configuration change is evaluated by all PEs cooperatively in parallel. One PE, the *scheduler*, takes care of the perturbation control, collection of local evaluation results, decision, and scheduling of the annealing.

In this section, three variations of SMC PSAs similar to the conventional PSA schemes are described. Those will be compared with the proposed MMC PSA. They are the *single move scheme*, the *multiple move scheme*, and the *hybrid move scheme*. Each of these variations has a different perturbation strategy.

### 2.2.1 Single Move Scheme

In the single move scheme, a perturbation results from a move. The scheduler PE randomly chooses one pair of PEs which are to interact with each other for perturbation. The primary PE of the two initiates the perturbation. For the perturbed configuration, the partial cost evaluated by each PE is collected at the scheduler. Thus, a perturbation is done by a pair of PEs while the evaluation is carried out by all PEs. Decision on the perturbation is made by the scheduler and is broadcast to all other PEs.

This scheme allows only one move between a PE pair per perturbation, allowing an elaborate annealing to be carried out. As a consequence, especially in the final stage of the annealing, this scheme could carefully proceed to a (near) optimal solution due to gradual change in the configuration. However, as shown in Fig. 2a, most PEs are idle when the selected PE pair is working on a perturbation. This degrades the parallelization efficiency.

### 2.2.2 Multiple Move Scheme

In the multiple move scheme, all PEs are paired up and they communicate with each other when performing moves. Primary PEs initiate a perturbation by selecting moves; there can be  $N/2$  moves per perturbation as shown in Fig. 2b. Then, each perturbed configuration is evaluated by the corresponding PEs and the local results are collected at the scheduler, which is in charge of the overall annealing scheduling. A decision is made and broadcast by the scheduler to update the intermediate configuration in the same way as in the single move scheme.

The multiple move scheme can change the cost more rapidly than the single move scheme since the configuration change per perturbation can be made larger. However, near the final stage of annealing, the solution (cost) being trapped in a local optimum may easily oscillate around a certain configuration and not converge toward a global optimum. In this case, it would be desirable to modify cost by a small amount. Also, since the decision is still made by one PE (scheduler) while the others are idle, a substantial degradation of efficiency may result in, especially when the number of PEs is large.

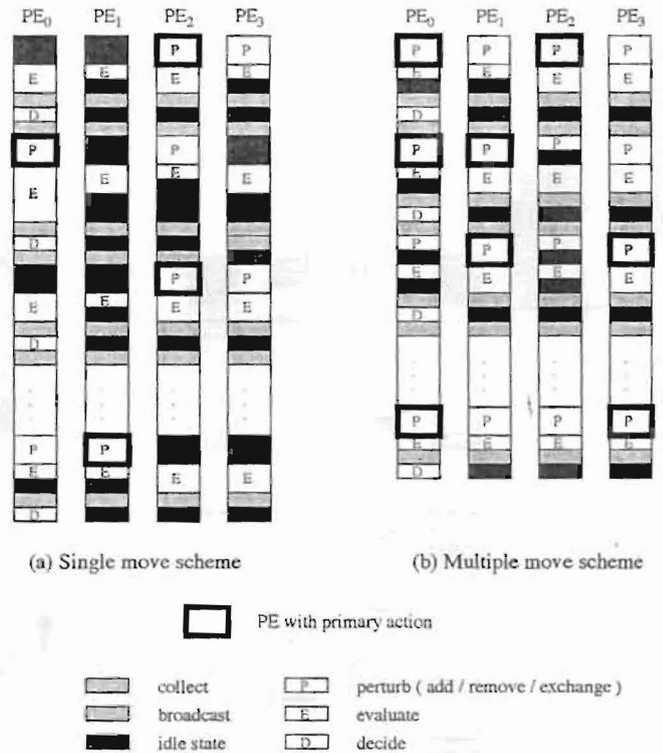


Fig. 2. SMC PSA schemes (a) single move scheme, (b) multiple move scheme.

### 2.2.3 Hybrid Move Scheme

The hybrid move scheme takes advantage of both the single move and the multiple move schemes. The basic idea appears to be equivalent to employing a range limiting function which, in general, restricts the range of moves according to the temperature [3].

In the hybrid move scheme, the *acceptance rate* (the number of perturbations accepted/the number of perturbations tried), is measured during annealing. It is used to adaptively determine the type of perturbation to be performed. The single move would be desirable in order to generate a small perturbation when the acceptance rate is low. At high acceptance rates, multiple moves to generate a large perturbation may lead to a fast reduction of the cost. Utilizing the acceptance rate does not add a heavy burden to the scheduler because the information at each PE can be delivered to the scheduler during the information exchange preceding a decision.

## 2.3 Multiple Markov Chain PSA

The approach of SMC PSA has at least two fundamental drawbacks. First, parallelism is limited since it is confined to a single search path. Second, a very high overhead is to be paid due to too frequent communication.

In order to improve performance of SMC PSA, one may relax the restriction that all PEs follow a single search path. One possibility is to have PEs trace multiple search paths at the same time, i.e., multiple Markov chain (MMC) PSA. Every PE has the entire search space, and the perturbation, evaluation and decision are performed by each PE individually. At the end of the algorithm, the best solution among the PEs is selected. The initial temperature, the

cooling coefficient and the equilibrium conditions can be independently decided and controlled by each PE. Since each PE randomly perturbs a configuration independently of others, it would be highly unlikely that any two PEs follows the same path. For better performance, PEs may be allowed to interact with each other as will be described later.

In order to guarantee a reasonable speed-up, the number of trials at a temperature is reduced by a certain factor, called *reduction factor* (refer to Section 3 for more details). Then, it is shown that we can still get an equivalent or even better solution.

Since PEs in the MMC PSA do not have to interact as frequently as in the SMC schemes, communication overhead will be significantly lower, which leads to a higher speed-up and efficiency. Also, the system is less likely to be trapped in a local optimum because multiple search paths are traced simultaneously.

Three schemes, namely, the *noninteracting scheme*, the *periodic exchange scheme*, and the *dynamic exchange scheme*, will be described below.

2.3.1 Noninteracting Scheme

Each PE independently perturbs the configuration, evaluates the cost, and decides on the perturbation. PEs do not interact during individual annealing processes until all PEs find their final solutions. Then the best of the solutions is saved and the others are discarded. One potential problem is that, as illustrated in Fig. 3a, all other PEs may have to wait for the PE with the longest search path (chain), resulting in idle time in every

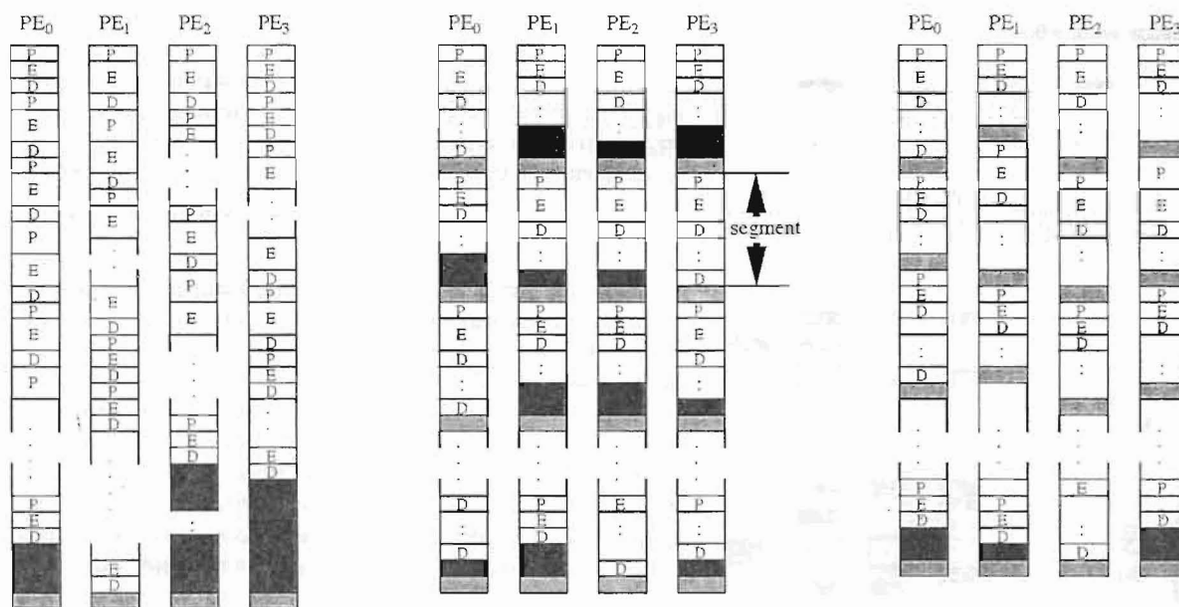
PE (except one), which may degrade the efficiency substantially.

In typical combinatorial problems, the computational load for perturbation and evaluation varies with configuration. Thus, as the annealing proceeds, the variance of the accumulated computational load among PEs becomes larger due to the random nature of SA, i.e., a larger variance in computation (or idle) time. Because PEs do not interact with each other until the end of SA, some of them may not perform useful computations. An efficient information exchange among PEs may be able to prevent PEs from performing unproductive computation or being idle so that the efficiency of parallelization can be improved.

2.3.2 Periodic Exchange Scheme

One possible way to improve the performance, referred to as the *periodic exchange scheme*, is illustrated in Fig. 3b. In this scheme, PEs exchange local information including the intermediate solutions and their costs with a fixed period. Then, each PE restarts from the best of the intermediate ones. Alternatively, to reduce the chance to be trapped, a random selection among the equilibrium solutions may be employed [11]. Also, depending on the intermediate cost distribution, the population update used in the genetic algorithm can be adopted [20]. We define a Markov chain between two successive information exchanges as a *segment*. Note that the multiple Markov chains are not completely independent, but they are within each segment.

Compared to the noninteracting scheme, communication overhead in this periodic exchange scheme would be



(a) Non-interacting MMC scheme

(b) Periodic exchange MMC scheme

(c) Asynchronous MMC scheme

- perturb / evaluate / decide
- idle state
- global state access

Fig. 3. MMC PSA schemes (a) noninteracting scheme, (b) periodic exchange scheme (synchronous), (c) asynchronous scheme.

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increased. However, each PE can utilize the information from other PEs such that the decrease in computations and idle times can be greater than the increase in communication overhead. For instance, a certain PE which is trapped in an inferior solution can recognize its state by comparing it with others and may accelerate the annealing procedure. That is, PEs may collectively converge to a better solution. Nevertheless, the period of the information exchange needs to be carefully selected depending on how the intermediate solutions affect the convergence of other PEs.

This scheme resembles the *division algorithm* with communication proposed in the literature [7]. However, note that this scheme does not require communication among all PEs at every temperature while the division algorithm does. Communication at every temperature would incur too much synchronization overhead.

### 2.3.3 Dynamic Exchange Scheme

The statistical data obtained during execution may be utilized to adaptively control the SA process in each PE to further reduce the execution time. For example, the acceptance rate which is closely related to the annealing state can be utilized. The periodic exchanges may induce unnecessary and untimely information exchanges. Moreover, an intermediate solution derived at an insufficiently cooled state can hamper the convergence of other PEs.

We propose a new MMC PSA, the *dynamic exchange scheme*, which adaptively determines when information is to be exchanged. In the dynamic exchange scheme, whenever the following two conditions are satisfied, the information exchange among PEs is carried out. First, a certain period of time has to elapse, i.e., to allow each PE a sufficient independent annealing. Second, the acceptance rate is below a certain value, i.e., some PEs arrive at significantly better solutions. That is, PEs exchange information only when necessary rather than with a fixed period. In this way, PEs more efficiently guide each other to a higher quality solution. Also, communication time can be reduced substantially. In addition, to utilize the idle time, all PEs may continue (even after the termination criterion is met) their own annealing processes by exchanging their solutions until the PE, for which the termination criterion is met last, finds its final solution. In such a case, all PEs actually terminate their annealing simultaneously.

## 2.4 Asynchronous MMC PSA

The MMC PSAs described in the previous section are synchronous schemes in that all PEs should be ready before an information exchange takes place. While the synchronous MMC PSA is expected to find a global optimal or near optimal solution faster than the conventional SMC PSAs, there is still potential for considerable performance (especially speed) improvement. Each PE in the synchronous MMC PSA needs to wait for other PEs in each segment before it can communicate with them to get their current solutions (*local states*) for comparison as illustrated in Fig. 3b. Therefore, unless the segment length is the same for all paths or PEs (an extremely unlikely condition), efficiency of parallelization is degraded due to the idle time of each PE. As the variation in segment length among PEs increases, the efficiency decreases.

In order to further improve the performance of the MMC PSA, we propose asynchronous communication among PEs accessing the global state to reduce or eliminate the idle times. Each PE follows a separate search path, accesses the *global state* which consists of the current best solution and its cost whenever it finishes a segment, and updates the state if necessary. The global state is stored in a memory location that can be accessed by all PEs. Once a PE gets the global state, it proceeds to the next segment without any delay. For the asynchronous MMC PSA, the definition of *segment* needs to be generalized to a search path or Markov chain that a PE follows between two successive accesses (communications) to the global state. Note that this definition is valid for the synchronous MMC PSA also.

In the case of the asynchronous MMC PSA, each PE locks the global state before accessing it at the end of each segment, and unlocks it subsequently for other PEs. If the local state is better than the global state, it updates the global state by its local state. Otherwise, it updates its local state by the global state. Then, without any delay, it proceeds to the following segment. On the contrary, in the case of the synchronous MMC PSA, each PE accesses and updates the global state in the same manner, but needs to wait until all other PEs finish the updating in each segment. Then, the global state which is the best solution (and its cost) found in that segment is copied by all PEs before they initiate their next segments.

The asynchronous MMC PSA has the following features compared to the synchronous MMC PSA. First, no PE will be idle during annealing. That is, different lengths of segments do not cause idle states until a Markov chain ends since communication (global state access) is carried out asynchronously. Second, communication overhead itself is less in the asynchronous implementation than in the synchronous one. In the synchronous MMC PSA, local states are to be exchanged among PEs; during this time computations are not performed. In contrast, an isolated access to the global state is needed by each PE at the end of each segment in the asynchronous MMC PSA. Simultaneous accesses by PEs, which result in conflicts, are unlikely. Third, the probability of being trapped in a local optimum can be smaller depending on the cost function. This is mainly due to the fact that not all PEs start from the same state in each segment while they do in the synchronous MMC PSA.

## 3 PERFORMANCE ANALYSIS

In this section, we model the PSAs described in the previous section and compare their performance. The main purpose is to show analytically that the MMC PSA can perform better than the SMC PSA and that the asynchronous MMC PSA can be faster than the synchronous MMC PSA.

### 3.1 SMC PSA vs. MMC PSA

We consider the MMC PSA first. Let the time for one iteration (P/E/D) be denoted by  $t_{ii}$  ( $= t_p + t_e + t_d$ ) and  $I_k$  denote the number of trials at the  $k$ th temperature (outer) loop of  $PE_i$ . Then the total execution time of the non-interacting

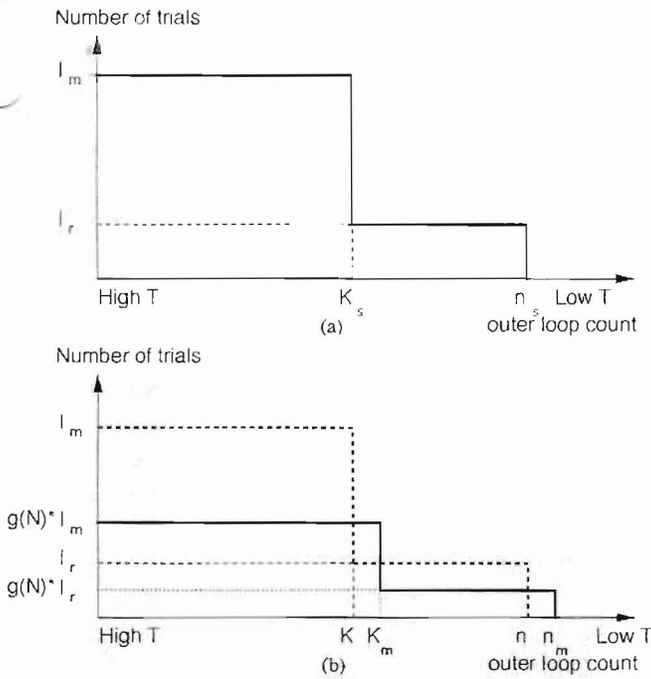


Fig. 4. Simplified annealing model (number of trials vs. outer loop count) for (a) SMC PSA and (b) MMC PSA.

MMC PSA with  $N$  PEs,  $t_{mmc}(N)$  can be written as follows:

$$t_{mmc}(N) = \max_{i \in \{1, \dots, N\}} \sum_{k=1}^{n_i} I_{ij} t_u + T_c, \quad (1)$$

where  $n_i$  is the final loop count of  $PE_i$  and  $T_c$  is the total communication time. For the interacting MMC PSAs, the first term becomes  $\sum_{j=1}^M \max_{i \in \{1, \dots, N\}} I_{ij} t_u$ , where  $M$  is the number of segments and  $I_{ij}$  is the total number of trials in the  $j$ th segment of  $PE_i$ .

It is reasonable to divide the temperature into higher and lower temperature zones by the *cut-off temperature*. The cut-off temperature ( $T_K$ ) is defined as a temperature at which the number of trials per temperature is noticeably reduced. Then, the annealing curve, the number of trials versus the outer loop count, can be approximated by a two-step curve as shown in Fig. 4. Note that the execution time is proportional to the area under the curve. The reduction factor,  $g(N)$ , which was introduced in Section 2.3, obviously affects the parallel execution time. A typical example of  $g(N)$  is  $\frac{1}{N}$ . For the MMC PSA, the number of trials at a temperature is usually limited by  $g(N) \cdot I_m$  from the initial temperature to the cut-off temperature and by  $g(N) \cdot I_r$  from the cut-off temperature to the final temperature, where  $I_m$  and  $I_r$  are the maximum numbers of trials allowed at high and low temperature regions in the sequential SA, respectively.

Let's denote the temperature (outer) loop count at the  $t$ -off temperature by  $K$  and the total outer loop count by  $n$  for the sequential SA. Then, the corresponding  $K_m$  and  $n_m$  for the MMC PSA can be represented as

$K_m = \max_{i \in \{1, \dots, N\}} K_i$  and  $n_m = \max_{i \in \{1, \dots, N\}} n_i$  where  $K_i$  and  $n_i$  are  $K$  and  $n$  of the  $PE_i$ , respectively. The outer loop counts at the final temperature ( $n_m$ ) and the cut-off temperature ( $K_m$ ) are random variables which vary with other parameters and the characteristics of cost distribution.

Let the times for perturbation, evaluation and decision be  $t_p$ ,  $t_e$ , and  $t_d$ , respectively, and the scheduling time per iteration be  $t_s$ . Then, the execution time of a sequential SA,  $t_{seq}$ , is

$$t_{seq} = (I_m K + I_r (n - K)) (t_p + t_e + t_d) + n t_s. \quad (2)$$

Similarly, referring to Fig. 4b, the execution time for the MMC PSA,  $t_{mmc}(N)$ , can be represented from (1) as follows:

$$t_{mmc}(N) = g(N) (I_m K_m + I_r (n_m - K_m)) (t_p + t_e + t_d) + M t_c \log_2 N. \quad (3)$$

In (3), it is assumed that the communication among  $N$  PEs can be completed in  $\log_2 N$  steps (e.g., in a hypercube), and  $t_c$  is the communication time between adjacent PEs. We note that  $M = 1$  for the noninteracting scheme, and  $M = \lceil n_m / L_p \rceil$  for the periodic exchange scheme, where  $L_p$  is the number of temperature loops between adjacent global state accesses (refer to Fig. 3).  $M$  for the dynamic exchange scheme is smaller than that for the periodic exchange scheme.

From (2) and (3), the speed-up for the MMC PSA,  $S_{mmc}$ , can be derived as

$$S_{mmc} = \frac{t_{seq}}{t_{mmc}(N)} = \frac{C_0 (t + t_d) + n t_s}{C_m (t + t_d) g(N) + M t_c \log_2 N + n_m t_s} \quad (4)$$

where  $t = t_p + t_e$ ,  $C_0 = I_m K + I_r (n - K)$ , and  $C_m = I_m K_m + I_r (n_m - K_m)$ .

For small  $t_s$  and  $t_c$ , if there is no significant difference between  $K$  and  $K_m$ , and  $n$  and  $n_m$ , the speed-up would be nearly  $1/g(N) = N$  assuming  $t \gg t_s, t_c$ . However, as the unit communication time ( $t_c$ ), the number of PEs ( $N$ ) or the total number of segments (communication phases) ( $M$ ), increases, the speed-up is degraded. Even if the two MMC PSA schemes with communication complete at the same outer loop count ( $n_m$ ), the amount of communication required by the dynamic exchange scheme is less than that by the periodic exchange scheme.

Now, consider the SMC PSA. Suppose that the perturbation and evaluation steps are parallelized, but not the decision step. The execution time for the SMC PSA can be written as (refer to Figs. 2 and 4)

$$t_{smc}(N) = (I_m K_s + I_r (n_s - K_s)) \left( \frac{t_p + t_e}{N} + t_d N + t_s + f_c t_c \log_2 N \right) + n_s t_s. \quad (5)$$

In (5),  $t'_c$  is the unit communication time of the SMC PSA which corresponds to the  $t_c$  of the MMC PSA,  $f_c$  is the number of communications per iteration, and  $t_0$  is the parallelization overhead for data management (e.g., the scheduler PE prepares data for parallel perturbation and evaluation, and the information to be broadcast to all PEs; refer to Section 2.2).  $t_c$

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and  $t'_c$  depend on data size, communication algorithm and computer system, and  $t_o$  depends on the perturbation and evaluation strategies.

Then, the speed-up of SMC PSA is derived as follows:

$$S_{smc} = \frac{t_{seq}}{t_{smc}(N)} = \frac{C_0(t + t_d) + nt_s}{C_s \left( \frac{t}{N} + t_o N + t_d + f_c t'_c \log_2 N \right) + n_s t_s} \quad (6)$$

where  $C_s = I_m K_s + I_r (n_s - K_s)$ .

To compare the two strategies (SMC and MMC),

$$\frac{S_{mmc}}{S_{smc}} = \frac{t_{smc}}{t_{mmc}(N)} = \frac{C_s \left( \frac{t}{N} + t_o N + t_d + f_c t'_c \log_2 N \right) + n_s t_s}{C_m (t + t_d) g(N) + M t_c \log_2 N + n_m t_s} \quad (7)$$

Even though the size of data exchanged at a time in the SMC PSA may be smaller than that in the MMC PSA ( $t'_c \leq t_c$ ), the total communication time of the SMC PSA is much larger than that of the MMC PSA due to the fact that PEs in the SMC PSA communicate much more frequently, i.e.,  $C_s f_c t'_c \log_2 N \gg M t_c \log_2 N$  since  $C_s f_c \gg M$ . Under the assumptions that  $t_s, t_o \ll t, C_s \approx C_m$  and  $g(N) = 1/N, \frac{S_{mmc}}{S_{smc}} > 1$ , i.e., the MMC PSA is faster than the SMC PSA. The assumptions are valid in most SA problems since

- 1)  $t_p$  and  $t_e$  are usually much larger than the scheduling time,  $t_s$ ,
- 2) the total number of trials in the MMC PSA can be made similar to that in the SMC PSA and
- 3) the reduction factor  $g(N) = 1/N$  is applicable because the sufficient cooling can lead PEs to achieve a quasi-equilibrium in the proposed interacting MMC PSA.

Even if a less drastic reduction factor like  $g(N) = \frac{1}{\sqrt{N}}$  is employed (then more trials are made, therefore, a better solution can be obtained),  $S_{mmc} > S_{smc}$  is usually satisfied under reasonable conditions such as  $t : t_o : t'_c = 10 : 1 : 1$ . Only in case that  $t \gg f_c t'_c \log_2 N, S_{smc}$  can approach  $S_{mmc}$ .

It would be interesting to see how problem-dependent the performance (speed-up) of each strategy is. For this purpose, one may employ the derivative of speed-up with respect to  $t$  where  $t = t_p + t_e$ , i.e.,  $\frac{\partial S_{mmc}}{\partial t}$  and  $\frac{\partial S_{smc}}{\partial t}$ . Note that a different problem would have a different value of  $t$ .

From (4) and (6), we can derive the following derivatives when  $t_s, t_o, t_d \ll t$ , and  $g(N) = 1/N$ ,

$$\frac{\partial S_{mmc}}{\partial t} = \frac{\frac{C_s}{C_m} (M t_c \log_2 N) + n_m t_s}{\left( \frac{t}{N} + \frac{M t_c \log_2 N + n_m t_s}{C_m} \right)^2} \quad (8)$$

and

$$\frac{\partial S_{smc}}{\partial t} = \frac{\frac{C_s}{C_s} (t_o N + f_c t'_c \log_2 N + (1 - \frac{1}{N}) t_d)}{\left( \frac{t}{N} + \frac{n_s t_s}{C_s} \right)} \quad (9)$$

respectively.

If  $C_s = C_m$  and  $t_c, t'_c \ll t$ , the second term of the denominator in (8) is much smaller than the first term due to  $C_m t \gg N n_m t_s$ . Also, the second term of the denominator in (9) is

much smaller than the first term due to  $C_s t \gg N n_s t_s$ . Comparing the numerators of (8) and (9),  $C_m > \frac{M}{f_c}$  and  $C_m t_o > \frac{n_m}{N} t_s$  are satisfied in most applications. Therefore, the following inequality holds in most cases:

$$\frac{\partial S_{smc}}{\partial t} > \frac{\partial S_{mmc}}{\partial t} \quad (10)$$

That is, the performance (speed-up) of MMC PSA is more consistent than that of SMC PSA as the optimization problem varies. This is mainly due to the fact that the MMC PSA parallelizes the whole annealing steps while the SMC PSA does only a part which can vary significantly with problem.

### 3.2 Synchronous MMC PSA vs. Asynchronous MMC PSA

The length of a segment can be formulated as a random variable which has a certain distribution characterized by a probability density function (pdf). In the following analysis, it will be assumed that the random variable for segment length has the same distribution for all PEs and at all temperatures, given a problem and a scheduling scheme. This assumption is mainly for simplicity of analysis. Later, whenever necessary and possible, relaxing the assumption or the effect of deviation from the assumption will be discussed.

Let random variable  $l_{ik}$  denote segment length for the  $k$ th segment of  $PE_i$ . The unit of the segment length is time and is always non-negative. The mean and standard deviation of  $l_{ik}$  are  $\mu_{ik}$  and  $\sigma_{ik}$ , respectively. Under the above assumption  $\mu_{ik} = \mu$  and  $\sigma_{ik} = \sigma$  for all  $i, k$ .

Let the total execution times of the asynchronous and synchronous MMC PSAs be  $t_{asyn}$  and  $t_{syn}$ , respectively. Suppose that  $N$  PEs are employed and each PE follows  $M$  segments. In the case of the synchronous MMC PSA, all PEs are synchronized at the end of each segment. Therefore,  $t_{syn}$  can be written as follows:

$$t_{syn} = \sum_{k=1}^M \max_{i \in \{1, \dots, N\}} l_{ik} \quad (11)$$

In the case of the asynchronous MMC PSA, each PE proceeds without synchronization until the end of the entire simulated annealing when local solutions are compared. Hence,  $t_{asyn}$  can be represented as follows:

$$t_{asyn} = \max_{i \in \{1, \dots, N\}} \left( \sum_{k=1}^M l_{ik} \right) \quad (12)$$

From (11) and (12), one can see that  $t_{syn}$  can never be smaller than  $t_{asyn}$ . They become equal when the outcome of  $l_{ik}$  doesn't vary with  $i$ , which is practically impossible. For a more quantitative comparison, let  $l_{max}$  denote the maximum possible outcome (or upper bound) of  $l_{ik}$ . It is reasonable to assume that  $l_{max}$  depends on  $\mu$  and  $\sigma$ . Noting that  $l_{max}$  is larger for a larger  $\sigma$ , given a class of underlying distribution, one may write

$$l_{max} = \mu(1 + h(\bar{\sigma})), \quad (13)$$

where  $h(\cdot)$  is a monotonically increasing positive function

indicating how much  $l_{max}$  is deviated from  $\mu$ , and  $\bar{\sigma}$  is the standard deviation normalized by the mean, i.e.,  $\frac{\sigma}{\mu}$ .

One may try to analytically derive  $l_{max}$  by assuming a specific pdf for  $l_{ik}$ . However, it may not be worthwhile due to the following reasons. First, it is almost impossible to derive an easily appreciable closed form formula describing maximum outcome for samples of the sum of random variables,  $t_{asyn}$ , though it is relatively easy for samples of a random variable. Second, it may not be feasible to build a probabilistic model, e.g., pdf, that precisely fits a given practical problem. Third, in most cases, a performance measure derived with the first and second order moments, i.e., mean and standard deviation, is sufficiently accurate and meaningful.

From (11),

$$t_{syn} = M l_{max} = \mu(1 + h(\bar{\sigma}))M. \quad (14)$$

The standard deviation of  $l_i = \sum_{k=1}^M l_{ik}$ , which is the total segment length of  $PE_i$  in the asynchronous MMC PSA, is  $\sigma\sqrt{M(M-1)\rho + M}$  where  $\rho$  is the correlation coefficient between segments  $l_{ij}$  and  $l_{ii}$  for  $j \neq i$ , and  $0 \leq \rho \leq 1$ . After normalization by its mean which is  $\mu M$ , it becomes  $\bar{\sigma}\sqrt{\frac{(M-1)\rho + 1}{M}}$ . With this result, (12) can be expressed as

$$t_{asyn} = \mu M \left( 1 + h\left(\bar{\sigma}\sqrt{\frac{(M-1)\rho + 1}{M}}\right) \right). \quad (15)$$

Now, denote the ratio of  $t_{syn}$  to  $t_{asyn}$  by  $r$ , which indicates the relative improvement of the execution time (it will be simply referred to as the *improvement factor*) of asynchronous MMC PSA over synchronous MMC PSA. It needs to be emphasized that  $r$  is not the speed-up ( $S_{mmc}$ ) of asynchronous MMC PSA over sequential SA. From (14) and (15),

$$r = \frac{t_{syn}}{t_{asyn}} = \frac{1 + h(\bar{\sigma})}{1 + h\left(\bar{\sigma}\sqrt{\frac{(M-1)\rho + 1}{M}}\right)} \quad (16)$$

where  $M \geq 1$ .

It is clear from (16) that  $r \geq 1$  since  $0 \leq \rho \leq 1$  and  $h(\cdot)$  is monotonically increasing, i.e., the asynchronous MMC PSA is faster than the synchronous MMC PSA unless  $\rho = 1$  for which  $r = 1$ . Though (16) is based on the assumptions mentioned above, several meaningful observations can be drawn from the equation.

### 3.2.1 Global State Access Frequency

First, consider the effect of  $M$  on  $r$ . Though  $M$  is the total number of global state accesses (or segments), it may be referred to as *global state access frequency* to indicate how often a PE accesses the global state, which should be linearly proportional to  $M$ . The global state access frequency,  $M$ , is one of the major parameters which differentiates the synchronous and asynchronous schemes. As  $M$  increases, PEs access the global state more frequently, PEs will be idle for a longer period of time in both PSAs. However, the idle time in the asynchronous scheme, which occurs only in the final segment, is relatively smaller than that in the synchronous scheme. This is due to the property that the nor-

malized standard deviation ( $\bar{\sigma}$ ) of a sum of random variables cannot be larger than the sum of the normalized standard deviations of individual random variables, as derived earlier. This difference becomes larger as  $M$  increases as can be seen in (16).

From (16), it is easy to see that  $r$  increases first and tends to saturate as the global state access frequency increases. This benefit of using the asynchronous MMC PSA is greater when the normalized standard deviation is larger. As the correlation between segments decreases,  $r$  becomes more advantageous to use the asynchronous scheme. The improvement factor  $r$  is maximized, given  $\bar{\sigma}$  and  $M$  when  $\rho = 0$  (segments are uncorrelated).

### 3.2.2 Number of Markov Chains

In the above derivation and discussion, the number of Markov chains is assumed to have no effect on  $l_{max}$  which influences the improvement factor. Theoretically, the maximum value of a random variable is fixed given an underlying distribution. When sampling the random variable in practice, the maximum of the samples or outcomes may not be the same as the theoretical one. The sample maximum is expected to approach the theoretical one as the number of samples increases. This practical aspect was not taken into account when  $l_{max}$  was formulated (refer to (13)). To make the improvement factor formula more realistic, (13) may be modified as follows:

$$l_{max} = \mu(1 + h(\bar{\sigma}))f(N) \quad (17)$$

where  $f(N)$  is a monotonically increasing function of  $N > 1$ ,  $0 < f(2) < 1$ , and  $f(N)$  approaches one as  $N$  increases.

$f(N)$  can be considered as a calibrating function which indicates the *actual* effect of  $h(\bar{\sigma})$  or  $\bar{\sigma}$  on  $l_{max}$  as a function of  $N$ . As the number of PEs,  $N$ , increases, the improvement factor  $r$  (of the asynchronous MMC PSA over the synchronous MMC PSA) increases, but eventually saturates.

### 3.2.3 Communication Overhead

Another advantage of employing the asynchronous MMC PSA is that communication time itself can be saved (though it is not included in the equations of this section). In the synchronous MMC PSA implemented on a message passing multiprocessor, the local states are to be collected at a certain PE which will then derive the global state from them and broadcast it back to all other PEs. This data sharing would require communication time proportional to  $D \log_2 N$  on a hypercube, for example, where  $D$  is the size of data (global state) to be exchanged and  $N$  the number of PEs [18]. If the synchronous MMC PSA is implemented on a shared memory multiprocessor, the communication time would be  $O(DN)$  assuming that there is only one copy of the global state on the shared memory.

In the asynchronous MMC PSA, each PE just needs to access and update, if necessary, the global state independent of other PEs. Therefore, the communication time would be  $O(D)$  on a shared memory system assuming that there is no conflict among PEs in the global state access. Of course, in the worst case when all PEs try to access the global state at the same time, the complexity would be  $O(DN)$ . However, this worst case is very unlikely to occur. On a message

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passing multiprocessor, each PE may send its local state to a controller at the end of each segment. Then, the controller returns the global state (after updating it by the local state if necessary). The communication time complexity is the same as on a shared memory system. Therefore, the saving in communication time by the asynchronous scheme can be substantial when  $N$  is large.

### 3.2.4 Solution Quality

Any faster scheme which degrades solution quality significantly would not be acceptable unless speed is more important than solution quality. All PEs in the synchronous MMC PSA initiate each segment from the same global state (solution). However, a different PE in the asynchronous MMC PSA may see a different global state depending on the order in which PEs access the global state.

Suppose that a PE accesses the global state at the end of each segment. If a local solution (state) of the PE is better than the global solution, the PE updates the global solution by its solution. If not, the PE discards its solution and initiates next segment from the global solution (state). For example, suppose the PE with the best solution happens to update the global state *first* in a certain segment. Then, all PEs would start from the same global state at least in that segment. Unless this happens, some PEs would have initial solutions for a segment which are different from others. As in the genetic algorithm [20], this will reduce the probability that the global state is trapped in a local optimum. Therefore, the solution quality is not expected to be worse than that by the synchronous MMC PSA.

### 3.2.5 Problem Dependency

The cost function may vary with the problem. If the cost function, more specifically density or distribution of local optima, changes, it is likely for the pdf of  $l_k$  to change resulting in a different  $\sigma$ . For a larger  $\bar{\sigma}$ , a higher  $r$  would be obtained as can be seen from (16).

## 4 APPLICATION TO GRAPH PARTITIONING PROBLEM

In order to experimentally examine the performance of the proposed MMC PSAs (synchronous and asynchronous), we have applied them to graph partitioning, which is NP-complete. Many engineering problems can be formulated as a graph partitioning problem [21], [22], [23], [24].

### 4.1 Problem Formulation

An *undirected* graph  $G = (V, E)$  consists of a finite set of nodes (vertices)  $V$  and a finite set of edges (links)  $E$ , where an element of  $E$  is an *unordered* pair of nodes  $u$  and  $v$  in  $V$ , i.e., an edge in  $E$  is represented as  $e_{u,v} = (u, v) \in E$ , where  $u, v \in V$ . A *subgraph*  $(W, F)$  can be defined such that  $W \subseteq V$  and  $F \subseteq E$ . If nonempty sets  $V_1$  and  $V_2$  are subsets of  $V$  such that  $V_1 \cup V_2 = V$  and  $V_1 \cap V_2 = \emptyset$ ,  $\{V_1, V_2\}$  is said to be a *partition*. *Cut* is a set of edges the removal of which results in a partition. The weight of a node  $u$  is denoted by  $\omega_v(u)$  and the weight of an edge  $(u, v)$  by  $\omega_e((u, v))$ .

### 4.1.1 Graph Partitioning Problem

Given an undirected graph  $G = (V, E)$  with node and edge weights,  $(\omega_v(\cdot), \omega_e(\cdot))$ , where  $V$  is a set of nodes and  $E$  is a set of edges, find a partition of  $V$  into  $R$  disjoint sets  $\{V_1, V_2, \dots, V_R\}$  such that the cost function below is minimized.

The cost function, which is to be minimized, is chosen as

$$C(\{V_1, \dots, V_R\}) = aM(\{V_1, \dots, V_R\})^x + bL(\{V_1, \dots, V_R\})^y \quad (18)$$

where

$$M(\{V_1, \dots, V_R\}) = \max\{\|V_1\|, \dots, \|V_R\|\},$$

and

$$L(\{V_1, \dots, V_R\}) = \sum_{(u,v) \in \delta(V_i, V_j), 1 \leq i, j \leq R, i \neq j} \omega_e((u, v)),$$

where  $\|V_i\| = \sum_{v_n \in V_i} \omega_v(v_n)$  and  $\delta(V_i, V_j)$  is the cut of the sets  $V_i$  and  $V_j$ , which is defined as

$$\delta(V_i, V_j) = \{(u, v) \in E \mid u \in V_i \wedge v \in V_j\}.$$

Suppose that  $a = b = x = y = 1$  with  $\omega_v(\cdot) = 1$  and  $\omega_e(\cdot) = 1$  for all nodes and edges in the graph. Then,  $C(\cdot) = M + L$ , where  $M$  is the largest number of nodes in a subgraph and  $L$  is the total number of edges between subgraphs. In this case, minimizing  $C(\cdot)$  requires balancing the node distribution among subgraphs and at the same time reducing the number of edges lying between subgraphs. By choosing an appropriate set of values for the parameters  $(a, b, x, y)$ , one may formulate a cost function suitable for a given application. Also, it needs to be pointed out that our approach of MMC PSA is not limited to graph partitioning only, but applicable to any optimization problems.

### 4.2 Perturbation Algorithm

In SA, the next candidate configuration (partition) is determined by a perturbation scheme. In the graph partitioning problem, we define three ways of perturbation, i.e., *add*, *remove*, and *exchange*, (each of which is referred to as a *move*), where an *add* in a subgraph corresponds to a *remove* in the paired subgraph. Note that a perturbation may result from multiple moves when there are more than two partitions (subgraphs). One of the moves is selected at random according to the node distribution. The add, remove and exchange are uniformly generated except in the extreme cases (e.g., the difference in the number of nodes exceeds a certain constant or the selected subgraphs have only non-removable nodes). Fig. 5 shows the perturbation examples where (a) is the initial configuration and the resulting configurations after a move—(b) add, (c) remove and (d) exchange—in view of the subgraph A (the *primary* subgraph).

The perturbation scheme, which is referred to as the *heuristic rearrangement algorithm*, is incorporated into the inner loop to efficiently perturb the configuration. In the heuristic rearrangement algorithm, the nodes in the primary subgraph (which is randomly chosen among all subgraphs) are sorted periodically (not in every iteration) in the ascending order according to internal degree (where the internal degree,  $deg(v)$ , is defined as the sum of weights of

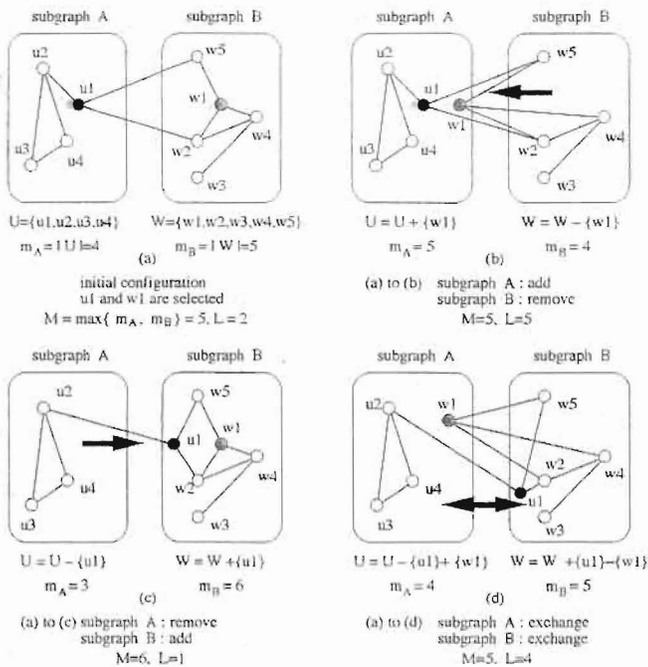


Fig. 5. Perturbation examples (a) initial configuration, (b) add, (c) remove, (d) exchange.

internal edges incident on the node  $v_i$ , where an internal edge is an edge that does not lie between subgraphs). The top node in the sorted list is involved in the move immediately following the sorting. Selection of nodes to be moved or exchanged is basically random, but is designed such that no node is completely excluded.

This heuristic rearrangement algorithm generates the next configuration more effectively than a simple random perturbation since it can reduce the unnecessary perturbations to a great extent by utilizing the connectivity information of subgraph nodes.

4.3 Communication Algorithms

Information exchange among PEs is necessary for both types of PSAs described earlier. In the SMC PSA, communication is indispensable when subgraphs are perturbed, when the locally evaluated costs at different PEs are collected at a certain PE, and when the decision is broadcast. In the MMC PSA, local information needs to be exchanged to update the global state (the current configuration and its cost) when each PE wants to check other PEs status during annealing and when all PEs finish annealing. Thus, three different types of communications are needed:

- 1) broadcast,
- 2) collect, and
- 3) exchange.

These communications can be efficiently implemented using the *pseudobinary tree* embedded in the hypercube [18].

4 Results and Discussions

The proposed PSA algorithms have been applied to the IEEE 30-node and 118-node standard networks, labeled as G30 and G118, respectively, a randomly generated 200 node weighted graph (G200), where the range of node weight (10, 20) and the

range of edge weight (1, 5) and 1,200 node graph (G1200). Their performance has been evaluated under the same conditions (parameters listed in Section 4.4.1). All performance figures, cost and execution time, are the averaged (not best) ones.

The PSA schemes have been compared based on the minimum cost found and the parallel execution time. In order to evaluate the convergence property of the proposed schemes, the best results obtained by the sequential algorithm of Irving and Sterling, [21] are referred to, which was reported to find significantly better solutions than an iterative improvement algorithm. The performance results of the proposed schemes for the cost function in (18) ((a, b, x, y) are usually set to (1, 1, 2, 3) as in [21]) are provided for G30 (average degree: 2.73), G118 (average degree: 3.05), G200 (average degree: 2.77), and G1200 (average degree: 2.98).

For further comparison, another SMC PSA scheme proposed by Roussel-Ragot and Dreyfus, [11], to be referred to as Modified R-R (MRR), has been modified for better adaptation to the graph partitioning problem and implemented. It utilizes the acceptance rate in determining next solution. In the MRR, PEs asynchronously continue annealing until at least one PE finds an acceptable solution. Then, one of the accepted solutions is randomly selected when the acceptance rate is high ( $> 1/N$ ) while the best solution is chosen when the acceptance rate is low ( $< 1/N$ ) [11]. Note that PEs have to communicate with each other frequently especially when the temperature is high.

4.4.1 Test Environment and Parameters

A graph is initially partitioned at random. The initial temperature ( $T_i$ ) for each PE is not predetermined identically, but is independently derived in the beginning with randomly selected sample configurations. In addition, the standard deviation of the intermediate solutions at each PE is calculated at every temperature to determine the next temperature [19]. In the following, the annealing parameters for the graph partitioning problem, which are empirically determined, are summarized:

- maximum number of temperature decrement : 250
- maximum number of iterations at a temperature ( $I_m$ ):  $k \cdot 100 \cdot (R - 1) \cdot g(N)$   
( $k = 1$  for G30,  $k = 2$  for G118,  $k = 3$  for G200,  $k = 10$  for G1200 where  $g(N)$  is usually set to  $1/N$ .)
- minimum number of iterations at a temperature ( $I_r$ ) :  $\max(0.3 I_m, 5)$
- static temperature decrement factor  $\alpha$  (cooling rate): 0.95
- threshold acceptance rate for hybrid move : 0.5

The synchronous MMC PSA schemes have been compared with the SMC PSA schemes on the Intel iPSC/2, a message-passing system. The synchronous and asynchronous MMC PSAs have been tested on a *shared-memory* multiprocessor system, BBN GP1000. The typical implementation results are tabulated in Tables 1 through 6 where cost and speed-up are the averaged ones for multiple executions in each case.

4.4.2 SMC PSA

Among the SMC PSA schemes, the hybrid move scheme has found the best configuration in all cases tested, as

PSA/SSA	Seq
SMC	3.6
PSA	1.1
MMC	3.8
PSA	1.1
SSA	1.1

(—) In the single

N	cost
1	341
2	442
4	396
8	370
16	485

N	cost
1	481
2	523
4	661
8	843
16	861

N	cost
1	1.27
2	1.33
4	2.02
8	3.24
16	4.57

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TABLE 1  
PERFORMANCE OF PSA SCHEMES FOR IEEE  
30-NODE NETWORK (G30), (x, y) = (2, 3)

PSA/SSA scheme	(a, b) = (1, 3)						(a, b) = (10, 1)					
	N=2, R=2			N=4, R=4			N=4, R=4			N=8, R=8		
	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S
SMC	600	17948	0.5	1657	10731	1.4	1539	8251	1.5	3010	6668	3.6
PSA	—	—	—	2268	7625	2.0	1810	7880	1.6	3113	5854	4.1
MMMC	633	18774	0.5	1657	9942	1.6	1539	8240	1.5	3104	7743	3.1
SSA	553	5362	1.7	1636	4359	3.5	1322	3900	3.2	2675	3459	6.9
Irving	649	n/a	—	1617	n/a	—	1322	n/a	—	n/a	n/a	—

(—) In the SMC, the multiple move and hybrid move schemes are the same as the single move scheme for N = R = 2.

TABLE 2  
PERFORMANCE OF PSA SCHEMES  
FOR G30, G118, AND G1200

N	SMC PSA						MMC PSA					
	Hybrid Move			Modified R-R			Noninteracting			Dynamic Exchange		
	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S
1	340	15910	1	—	—	—	340	15910	1	—	—	—
2	442	12240	1.3	340	26520	0.6	320	6311	1.6	320	4482	1.7
4	396	6696	2.4	571	29703	0.5	365	3963	3.2	330	3153	3.3
8	370	2864	5.6	768	14451	1.1	338	2482	6.4	320	1974	6.5
16	487	2796	5.7	391	12757	1.2	320	2098	7.6	320	1302	12.2

(a) G30, R=2, (a,b)=(1,1), (x,y)=(2,3)

N	SMC PSA						MMC PSA					
	Hybrid Move			Modified R-R			Noninteracting			Dynamic Exchange		
	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S
1	4812	33159	1	—	—	—	4812	33159	1	—	—	—
2	5230	19804	1.7	6885	32460	1.0	4417	17047	1.9	3824	15736	2.1
4	6616	10738	3.0	8058	21717	1.5	4712	9584	3.5	4210	8880	3.7
8	8436	7712	4.3	14191	29417	1.2	6857	4901	6.8	4210	4510	7.3
16	8614	6109	5.4	9659	23169	1.4	5231	2883	11.5	4481	2429	13.7

(b) G118, R=2, (a,b)=(1,1), (x,y)=(2,3)

N	SMC PSA						MMC PSA					
	Hybrid Move			Modified R-R			Noninteracting			Dynamic Exchange		
	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S
1	1.27e6	3219	1	—	—	—	1.27e6	3219	1	—	—	—
2	1.33e6	2118	1.5	1.73e6	2660	1.2	1.13e6	1462	2.2	1.13e6	1461	2.2
4	2.02e6	999	3.2	3.28e6	1778	1.8	1.50e6	727	4.4	1.37e6	704	4.6
8	3.24e6	787	4.1	3.62e6	1594	2.1	1.83e6	382	8.4	1.11e6	338	9.5
16	4.57e6	733	4.4	4.96e6	1668	1.9	2.67e6	239	13.4	1.83e6	213	15.1

(c) G1200, R=2, (a,b)=(1, 1), (x,y)=(2,3)

shown in Tables 1 and 3. In terms of the execution time, the multiple move scheme takes the least amount of time, and the single move scheme takes the longest since it changes the graph configuration gradually. While the multiple move scheme can rapidly perform the initial stages of annealing, it is susceptible to local minima and oscillations in the final stages. The single move is better at the final stage of the annealing. This is the reason why the hybrid move

TABLE 3  
PERFORMANCE OF PSA SCHEMES FOR IEEE  
118-NODE NETWORK (G118), (x, y) = (2, 3)

PSA/SSA scheme	N=2, R=2			N=4, R=4			N=8, R=8			N=16, R=16		
	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S	cost	T[ms]	S
SMC	5230	19804	1.2	7526	33471	1.3	27625	31350	3.1	175841	38663	4.6
PSA	—	—	—	9369	30901	1.4	30416	23138	4.2	314657	23803	7.4
MMMC	—	—	—	6856	32837	1.3	24965	26265	3.7	103967	29917	5.9
SSA	6885	32460	0.8	12017	40579	1.0	64529	35597	2.7	175785	43277	4.1
Static Noninter	4417	17047	1.5	4819	12849	3.3	14353	17669	5.5	85380	15967	11.1
Periodic Exch	4283	16084	1.5	4188	10946	3.8	14012	14166	6.9	90975	12015	14.8
Dynamic Exch	3824	15736	1.6	3964	10640	3.9	11489	13648	7.1	74232	11184	15.8
Seq. Heuristic SA (N=1)	5336	24725	1.0	4265	41924	1.0	14353	97180	1.0	85409	177233	1.0

scheme performs best among the SMC PSA schemes.

The convergence of the MRR (SMC) PSA seems to be poor since the random selection rule (which is designed to preserve the property of the sequential SA) at high temperatures may not be suitable for the graph partitioning problem. The solution quality is usually worse than those by our SMC PSA schemes as can be seen in Tables 1, 2, and 3. Also, the speed-up is noticeably lower in the MRR PSA than in our SMC PSA schemes especially for a large N. In the MRR PSA, a larger amount of information (perturbation, evaluation and decision results) needs to be exchanged at every step among PEs to generate N different states.

It is observed that the SMC PSA sometimes performs even worse than the sequential version especially for a small size problem (Table 1). For a small size of graph, the computational load in the perturbation and evaluation steps (which are to be parallelized) is relatively small compared to the total execution time, and the relative communication overhead is larger. That is, the overhead can offset the speed gain from the limited exploitation of parallelism in computation. For the MRR PSA, in which even evaluation is not parallelized, the performance degradation is more apparent.

The single Markov chain approach is shown not to be suitable for efficient parallelization of the SA algorithm particularly for the graph partitioning problem, since the perturbation and evaluation of the configuration (for which the workload is shared by PEs) do not occupy a large portion of the execution time. In addition, the communication overhead involved in each perturbation and decision can be overwhelming.

4.4.3 Synchronous MMC PSA

Among the synchronous MMC PSAs, the proposed dynamic exchange scheme, which utilizes the acceptance rate in determining when to communicate, usually performs best in terms of both the solution quality and the execution time, as shown in Tables 1, 2, and 3. The periodic exchange scheme often completes its execution earlier than the non-interacting MMC scheme. However, as the problem size grows and the number of subgraphs (subnetworks) increases, it is likely to be trapped at a local minimum (Table 1 for N = R = 8 and Table 3 for N = R = 16). It can be observed that the MMC PSA performs much better in terms of the solution quality and/or the execution time than the SMC PSA in most cases. This superiority of the MMC PSA becomes more visible as the number of PEs employed, N, increases. The speed-ups reported in the tables

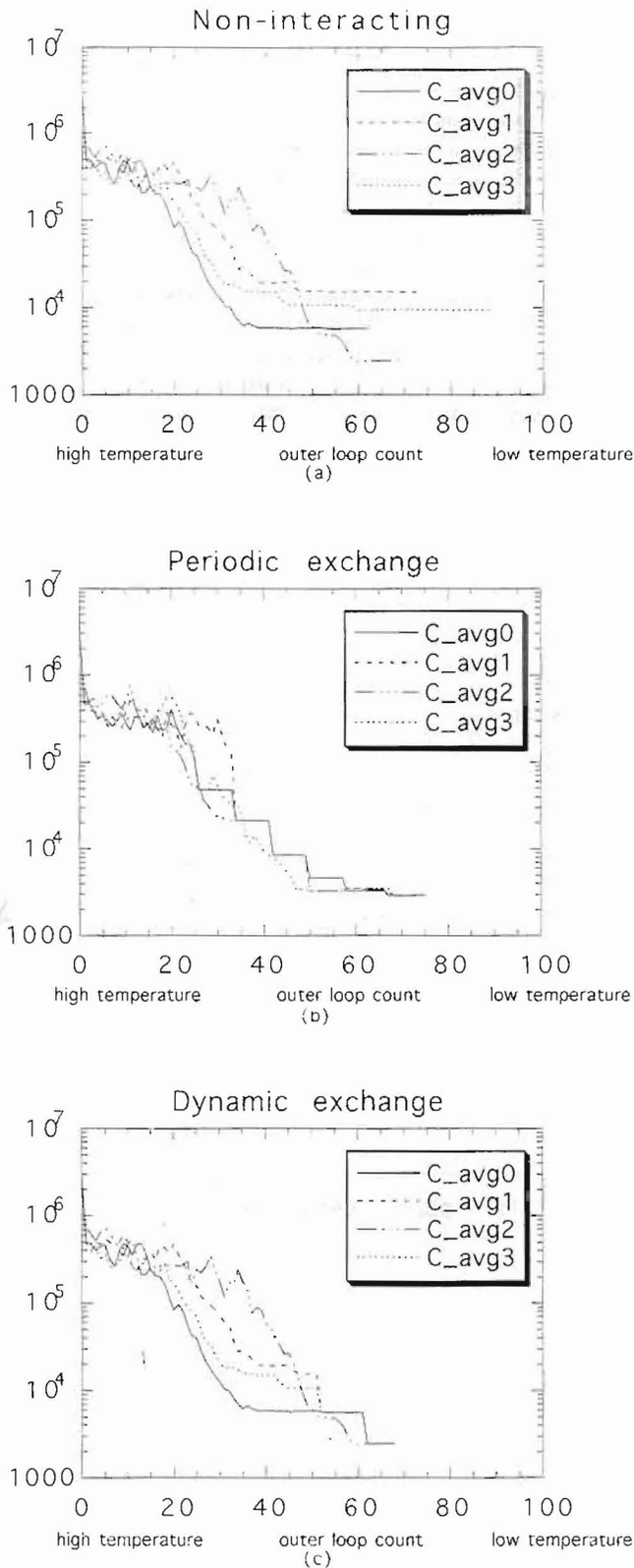


Fig. 6. Annealing curves of MMC PSAs ( $N = 4$  for G118) (a) noninteracting, (b) periodic exchange, (c) dynamic exchange.

are based on different qualities of solutions due to the random nature of SA. However, note that the solution quality by the MMC PSAs is better than that by the SMC PSAs in almost all cases (all cases for the dynamic exchange scheme). Therefore,

it is expected that higher speed-ups would be achieved if we somehow force the same quality of solution in both of MMC and SMC PSAs.

These observations may be explained as follows. First, since multiple search paths (chains) are followed in the MMC PSA, the probability of searching the local space where a global (or near) optimal solution is located would be higher for the MMC PSAs. Once most PEs (at least one PE) get on the right track, the annealing toward the near global optimum solution could be accelerated especially in the interacting MMC PSAs, i.e., a shorter execution time. Second, the SMC PSA requires more frequent communications (broadcast, collect, and exchange) among PEs since each PE needs to get the updated configuration and cost for every perturbation. This less communication overhead of the MMC PSA is another major factor which contributes to their better performance. Since the communication time is proportional to the number of PEs,  $N$ , the SMC PSA performs significantly worse than the MMC PSA, especially for a larger  $N$ . Third, in the MMC PSA, the P/E/D steps are fully (equivalently speaking) parallelized while in the SMC PSA the perturbation and decision (with updating the new graph partition in sequence) may not be executed fully in parallel.

The typical annealing curves (average cost vs. temperature) for the MMC PSAs are shown in Fig. 6. Each curve in the non-interacting scheme represents a normal sequential annealing curve, but only one of them leads to the final (best) solution. Note that the PE which completes annealing last has not found the best solution. The periodic exchange MMC PSA may reduce the redundant or unnecessary computation. However, the periodic exchange does not always result in positive effects as can be seen in Tables 1 and 3 and the annealing curve in Fig. 6b. Overly frequent information exchanges can lead to oscillation, hampering careful annealing processes and eventually deteriorating the convergence. In other words, only the sufficiently annealed solutions are worthwhile to be exchanged. This is why the dynamic exchange scheme improves the solution quality and also the execution time in most cases (Tables 1, 2 and 3).

Fig. 7 shows how the solution quality varies depending on the reduction factor,  $g(N)$ , for the G30 and G118. As the number of PEs (or Markov chains),  $N$ , increases, the solution quality (average cost) is improved to a certain point and then is degraded in general. A more drastic reduction factor like  $1/N$  usually has the turning point at small  $N$  (compared to  $1/\sqrt{N}$  or  $1/\log_2(N+1)$ ). That is, there can be a trade-off between the solution quality and execution time (remember that the speed-up,  $S$ , is almost inversely proportional to  $g(N)$ ).

The problem dependencies of the SMC and MMC PSAs are compared in Fig. 8. In these experiments, the system parameters (except  $t = t_p + t_e$ ) like communication time, obtained from the iPSC/2, were used. Fig. 8a shows that for a certain value of  $t$ , it would be possible for the speed-up of the SMC PSA to decrease after some point as  $N$  increases. Also, the derivatives of  $S_{mmc}$  and  $S_{smc}$  with respect to  $t$  (refer to (8) and (9)) are plotted in Fig. 8b. It is obvious that  $S_{mmc}$  is much less problem-dependent than  $S_{smc}$ .

Fig. 7. Eff exchange results).

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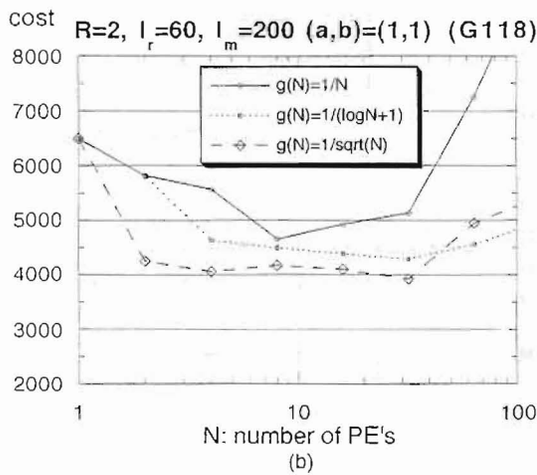
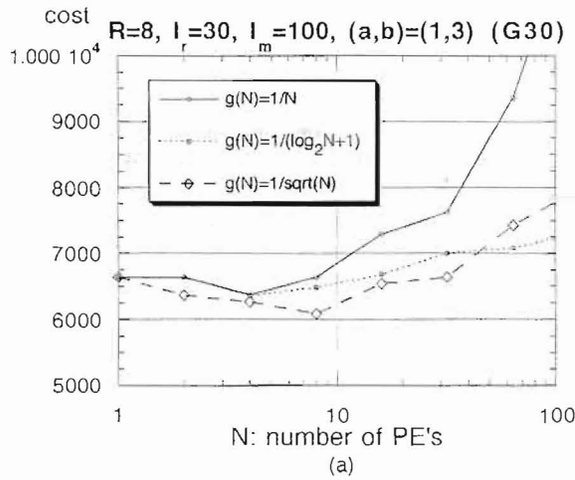


Fig. 7. Effect of reduction factor  $g(N)$  for MMC PSA (dynamic exchange) for (a) G30, (b) G118 ( $N > 32$ : sequential simulation results).

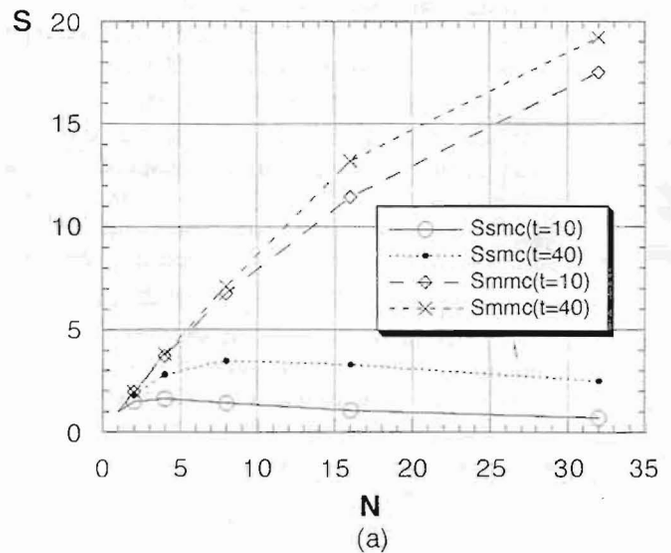
4.4.4 Asynchronous MMC PSA

The improvement factor,  $r$ , reported in Tables 4, 5, and 6 should be considered to be conservative since the solutions by the asynchronous MMC PSA are mostly better than those by its counterpart, the synchronous MMC PSA.

It is clear that the asynchronous MMC PSA outperforms the synchronous MMC PSA in speed as analyzed in Section 3.2. For the cases considered in this study, performance of MMC PSAs has been improved up to 43% by the asynchronous scheme in execution speed (refer to Table 4b). As can be seen in Table 4,  $r$  becomes larger as the global state access frequency,  $M$ , increases. Also, a larger problem (e.g., a larger graph, a larger number of partitions, a more complicated cost function) tends to result in a larger improvement factor since the segment length varies more relative to its mean when the problem (graph) size increases.

From Table 5, it can be observed that  $r$  increases as the number of Markov chains or PEs increases. This is because the maximum value (outcome) of segment length would be larger in general when there is a larger number of segments. This improvement in execution speed is also partially due to the

Speed-up (S) for  $t=10, t=40$



$dS/dt$  vs.  $t (=t_p + t_e)$  for G118

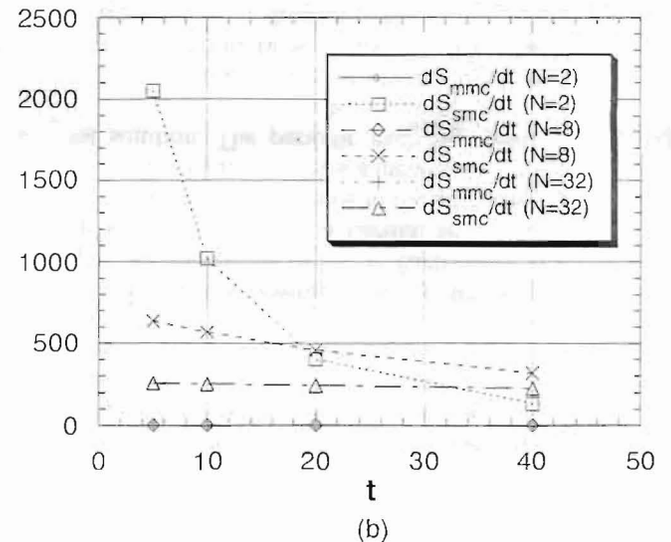


Fig. 8. (a) Speed-up ( $S$ ) and (b)  $\partial S/\partial t$  for SMC and MMC PSA. Note that the three curves for MMC in (b) are overlapped at the bottom (SMC: single move scheme, MMC: dynamic exchange scheme).

communication overhead which grows with  $N$  faster in the synchronous MMC PSA than in the asynchronous MMC PSA (refer to Tables 6b and 6d). For reference, the speed-up,  $S$ , of the asynchronous MMC PSA over the sequential SA is included in Table 5. It is seen that, in most cases, a high parallelization efficiency is obtained.

The percentage idle and communication (global state access) times (i.e., normalized by the total execution time) with  $N$  or  $M$  varied are provided in Table 6. For a fair comparison, the (absolute) idle and communication times are normalized by the total execution time which changes with  $N$  and  $M$ . First, it is clear that the percentage idle time outweighs the percentage communication time in both



TABLE 6  
 PERCENTAGE COMMUNICATION AND IDLE TIMES OF MMC PSAs:  
 $(g(N) = \frac{1}{N}, (t_{i_s}, t_{i_a}), (t_{c_s}, t_{c_a}))$ : PERCENTAGES OF IDLE  
 AND COMMUNICATION TIMES

M	Synchronous		Asynchronous		t <sub>i</sub> ratio	t <sub>c</sub> ratio
	t <sub>i<sub>s</sub></sub> (%)	t <sub>c<sub>s</sub></sub> (%)	t <sub>i<sub>a</sub></sub> (%)	t <sub>c<sub>a</sub></sub> (%)		
6	9.7	0.37	7.2	0.04	1.3	9.0
12	8.0	0.68	4.7	0.13	1.7	5.2
50	14.2	2.62	6.3	0.54	2.3	4.9
125	23.3	5.94	6.1	1.38	3.8	4.3

(a) G30 (N = 8, R = 4)

N	Synchronous		Asynchronous		t <sub>i</sub> ratio	t <sub>c</sub> ratio
	t <sub>i<sub>s</sub></sub> (%)	t <sub>c<sub>s</sub></sub> (%)	t <sub>i<sub>a</sub></sub> (%)	t <sub>c<sub>a</sub></sub> (%)		
2	3.7	0.24	1.7	0.11	2.1	2.1
4	7.6	0.74	3.3	0.25	2.3	3.0
8	14.2	2.62	6.3	0.54	2.3	4.9
16	32.0	7.95	8.2	1.09	3.9	7.3

(b) G30 (M = 50, R = 4)

M	Synchronous		Asynchronous		t <sub>i</sub> ratio	t <sub>c</sub> ratio
	t <sub>i<sub>s</sub></sub> (%)	t <sub>c<sub>s</sub></sub> (%)	t <sub>i<sub>a</sub></sub> (%)	t <sub>c<sub>a</sub></sub> (%)		
6	10.9	0.37	4.9	0.004	2.2	9.7
12	12.5	0.69	4.6	0.07	2.7	9.6
50	23.3	2.54	4.9	0.26	4.7	9.6
125	35.4	5.63	5.8	0.65	6.1	8.6

(c) G118 (N = 16, R = 16)

N	Synchronous		Asynchronous		t <sub>i</sub> ratio	t <sub>c</sub> ratio
	t <sub>i<sub>s</sub></sub> (%)	t <sub>c<sub>s</sub></sub> (%)	t <sub>i<sub>a</sub></sub> (%)	t <sub>c<sub>a</sub></sub> (%)		
2	7.9	0.04	2.7	0.02	2.9	2.4
4	13.5	0.15	3.4	0.04	3.9	3.6
8	18.5	0.58	3.5	0.10	5.2	5.7
16	23.3	2.54	4.9	0.26	4.7	9.6

(d) G118 (M = 50, R = 16)

asynchronous implementations have been considered. Their performance has been analyzed and experimentally compared to other PSAs for graph partitioning. The following conclusions may be drawn based on our extensive experimental results:

- The MMC PSAs can find a solution of better or equivalent quality faster than the SMC PSAs.
- The performance of the MMC PSAs is much less problem-dependent than that of the SMC PSAs.
- Among the MMC PSAs, the dynamic exchange scheme outperforms others in speed and solution quality.
- It is possible to reduce the execution time of the MMC PSA significantly by employing asynchronous information exchange.
- The quality of solution by the asynchronous MMC PSA is as good as (or even better than) that by the synchro-

nous version.

- The speed-up of the asynchronous MMC PSA over the synchronous version becomes larger for a greater number of PEs (or Markov chains) involved or a higher global state access frequency.

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