Abstract—Loops are the largest source of parallelism in many scientific applications. Parallelization of irregular loop applications is a challenging problem to achieve scalable performance on large-scale multi-core clusters. Previous research proposed an effective Master-Worker model on clusters for distributed self-scheduling schemes that apply to parallel loops with independent iterations. However, this model has not been applied to large-scale clusters. In this paper, we present an extension of the distributed self-scheduling schemes implemented in a hierarchical Master-Worker model. Our experiments with different self-scheduling schemes demonstrate good scalability when scaling up to 8,192 processors.

Index Terms—Scalable, Master-Worker, Self-Scheduling, Hierarchical.

I. INTRODUCTION

Loops are the largest source of parallelism in many scientific applications. There are several loop scheduling schemes for loops with and without data dependencies on clusters. If the iterations of a loop have no dependencies, each iteration can be considered as a task and can be scheduled independently. Loops can be scheduled statically at compile-time. This type of scheduling has the advantage of minimizing the scheduling time overhead, but it may cause load imbalance when the loop style is not uniformly distributed. Dynamic scheduling adapts the assigned number of iterations whenever it is unknown in advance how large the loop tasks are. A self-scheduling algorithm is a dynamic algorithm for scheduling loop iterations. An important class of dynamic scheduling are the self-scheduling schemes [1], [2], [3], [4] and references therein. In UMA (Uniform Memory Access) parallel system, these schemes can be implemented using a critical section for the loop iterations and no need exists for dedicating a (master) processor to do the scheduling. This is why these schemes are called self-scheduling schemes. An affinity scheduling algorithm is proposed and studied to reduce communications overhead (nonlocal memory accesses) on shared-memory multiprocessors in [5], [6] and [7]. A feedback guided dynamic loop scheduling is introduced and studied in [8] and [9]. An adaptive weighted factoring which performs well for scheduling loops in parallel unstructured grid applications and N-Body simulations is proposed in [10]. Different self-scheduling algorithms have been proposed which dynamically assign chunks of variable sizes to processors. They differ from each other in the way they calculate the size of the chunk assigned to each processor. The self-scheduling algorithms were initially proposed for loops without dependencies for shared memory parallel systems and later extended to distributed computing systems [2]. There are also results focusing on loops with dependencies [11]. Recent research results have been reported on loop self-scheduling methods for multi-core, graphics processing units, grids, and cloud systems [12], [13], [14], [15], [16], [17], [18], [19], [20], [21], [22], [23], [24], [25], [26] and [27].

Many modern high performance computing platforms, such as clusters, grids and clouds, can be scaled to thousands of parallel processors, servers and workstations. Thus, scalability becomes an important issue which should be taken into consideration. The developers of high performance computing application programs may over-schedule resources which can cause load imbalance and low speedup. This is especially true for some nested loops when executed on large-scale clusters. Previous research, [17], [18] has developed some loop scheduling schemes to get good performance and load balancing for small-scale clusters with multi-core processors. A scalable two Masters model with small number of workers on a small size application is proposed in [13]. In this paper, we design a scalable hierarchical distributed Master-Worker model for self-scheduling schemes on large-scale clusters. We implement these schemes on a large-scale cluster of Texas Advanced Computing Center, University of Texas at Austin. Our experiments demonstrate the good scalability of the proposed schemes.

The rest of the paper is organized as follows. In Section 2, we review simple loop self-scheduling schemes. In Section 3, we review distributed self-scheduling schemes. In Section 4, we describe the proposed hierarchical distributed schemes. In Section 5 and 6, experiments and results are presented. In Section 7, conclusions are drawn.

II. LOOP SCHEDULING SCHEMES

Self-scheduling is an automatic loop scheduling method in which idle processors request new loop iterations to be assigned to them. We study these methods from the perspective...
of distributed systems. For this, we use the Master-Worker architecture model (Figure 1). Idle processors submit a request to the master for new loop iterations. The master node has three components: 1) Task Scheduler. It uses scheduling schemes to divide the whole work into small scheduled chunks; 2) Request Queue. If one or some workers are idle, they request more work from the master node. If the master node is busy serving another worker, the requesting workers are added into Request Queue and wait to be served; 3) Result Collector. When the workers finish their work they send a new request and send the computed results to the Result Collector.

The number of iterations a processor should be assigned is an important issue. Due to processors’ possible heterogeneity and communication overhead, assigning the wrong processor a large number of iterations at the wrong time, may cause load imbalancing. Also, assigning a small number of iterations may cause too much communication and scheduling overhead.

A. Notations:

The following are common notations used throughout the whole paper:

- $I$ is the total number of iterations or tasks of a parallel loop;
- $p$ is the number of workers (i.e. processors) in the parallel or distributed system which execute the computational tasks;
- $P_1, P_2, ..., P_p$ represent the $p$ workers in the system;
- A few consecutive iterations are called a chunk. $C_i$ is the chunk-size at the $i$-th scheduling step (where: $i = 1, 2, ...$);
- $N$ is the number of scheduling steps;
- $t_j$, $j = 1, ..., p$, is the execution time of $P_j$ to complete all its tasks assigned to it by the scheduling scheme;
- $T_p = \max_{j=1,...,p}(t_j)$, is the parallel execution time of the loop on all $p$ workers;

In a generic self-scheduling scheme, at the $i$-th scheduling step, the master computes the chunk-size $C_i$ and the remaining number of tasks $R_i$:

$$R_0 = I, \quad C_i = f(R_{i-1}, p), \quad R_i = R_{i-1} - C_i$$  \hspace{1cm} (1)

where $f(., .)$ is a function possibly of more inputs than just $R_{i-1}$ and $p$. Then the master assigns to a worker processor $C_i$ tasks. Imbalance depends on the execution time gap between $t_j$, for $j = 1, ..., p$. This gap may be large if the first chunk is too large or (more often) if the last chunk (called the critical chunk) is too small.

The different ways to compute $C_i$ has given rise to different scheduling schemes. Some widely used schemes are the following. These schemes are studied or extended in [1], [2], [3], [4], [5], [10] and references therein.

Trapezoid Self-Scheduling (TSS) $C_i = C_{i-1} - D$, with (chunk) decrement : $D = \left\lfloor \frac{(F-L)}{(N-1)} \right\rfloor$, where: the first and last chunk-sizes $(F,L)$ are user/compiler-input or $F = \left\lceil \frac{I}{2p} \right\rceil$, $L = 1$. The number of scheduling steps assigned: $N = \left\lceil \frac{2s+1}{(F+L)} \right\rceil$. Note that $C_N = F - (N-1)D$ and $C_N \geq 1$ due to integer divisions.

Factoring Self-Scheduling (FSS) $C_i = \left\lfloor \frac{R_{i-1}}{(\alpha p)} \right\rfloor$, where the parameter $\alpha$ is computed (by a probability distribution) or is suboptimally chosen $\alpha = 2$. The chunk-size is kept the same in each stage or round (in which all processors are assigned a chunk of the same size) before moving to the next stage. Thus $R_i = R_{i-1} - pC_i$ (where $R_0 = I$) after each stage.

Guided Self-Scheduling (GSS) $C_i = \left\lfloor \frac{R_{i-1}}{p} \right\rfloor$. In the last steps too many small chunks are assigned. It assigns large chunks initially, which implies reduced communication/scheduling overheads only in the beginning but small chunks later. A modified version $GSS(k)$ with minimum assigned chunk-size $k$ (chosen by the user) attempts to improve on the weaknesses of $GSS$.

III. DISTRIBUTED LOOP SCHEDULING SCHEMES FOR DISTRIBUTED SYSTEMS

Load balancing in distributed systems is a very important factor in achieving near optimal execution time. To obtain load balancing, loop scheduling schemes must take into account the processing speeds of the workers or processors forming the system. The processors’ speeds are not precise, since memory, cache structure and even the program type may affect the performance of processors. However, one could run experiments to obtain estimates of the throughputs and one could show that these schemes are quite effective in practice.

We next present the distributed loop scheduling schemes based on the Master-Worker architecture.

A. Terminology:

- $V_j = \text{Speed}(P_j)/\min_{1 \leq i \leq p}\{\text{Speed}(P_i)\}$, $j = 1, ..., p$, is the virtual power of $P_j$ (computed by the master), where $\text{Speed}(P_j)$ is the processing speed of $P_j$. That is a standardized computing power in the current cluster.
- $V = \sum_{j=1}^{p} V_j$ is the total virtual computing power of the cluster.
- $DC$ is the distributed chunk size for one worker request, in a single scheduling step of distributed self-scheduling scheme.

Master:
- (1) Compute \( V_j \) for each worker
  (a) Receive \( \text{Speed}(P_j) \);
  (b) Compute all \( V_j \);
  (c) Send all \( V_j \);
- (2) Assign work and get the results
  (a) While there are unassigned tasks, if a request arrives, put it in the Request Queue.
  (b) Pick a request from the queue and get its virtual power \( V_j \). If there are computed results in this request, Result Collector receives them first. Then Task Scheduler compute the next chunk size \( DC \) to assign. The followings are the DTSS, DFSS and DGSS algorithms to compute the next chunk \( DC \):

**DTSS:**

\( \text{Current} \) is chunk size in the current step of TSS.

Initialization: \( F = \left\lceil \frac{I}{2P} \right\rceil, L = 1, N = \left\lceil \frac{2J}{(F+L)} \right\rceil, \) \( D = \left\lfloor \frac{(F-L)}{(N-1)} \right\rfloor, \text{Current} = F \)

**Algorithm 1 Calculate DC**

\[
DC = 0;
for \ k = 1 \rightarrow V_j \ do
\begin{align*}
DC &= DC + \text{Current}; \\
\text{Current} &= \text{Current} - D;
\end{align*}
end for
return DC;

**DFSS:**

\( DC_{sum} \) is the assigned work in the current stage.

Initialization: \( R = I, \alpha = 2.0, DC_{sum} = 0 \)

**Algorithm 2 Calculate DC**

\[
DC = \left\lceil \frac{R}{(\alpha V)} \right\rceil * V_j; \\
DC_{sum} = DC_{sum} + DC; \\
\text{if} \ (\text{Master has assigned all the work in the current stage}) \hspace{1cm} \text{then} \\
\begin{align*}
&\{ \text{Goto next stage and update the remaining work.} \} \hspace{1cm} \text{end if} \\
&\text{R} = R - DC_{sum}; \\
&DC_{sum} = 0;
\end{align*}
end if
return DC;

**DGSS:**

Initialization: \( R = I \)

**Algorithm 3 Calculate DC**

\[
DC = \left\lceil \frac{R}{(A)} \right\rceil * V_j; \\
R = R - DC; \\
\text{return} \ DC;
\]

**Worker**:

- (1) Send \( \text{Speed}(P_j) \);
- (2) Send a request;
- (3) Wait for a reply;
  IF (There is unassigned work)
  \{ \text{Compute the new work; } \\
  \text{Return the results and send another request; } \\
  \text{Go back to (2); } \\
  \}
ELSE
  Terminate;

**IV. Hierarchical Distributed Schemes**

When considering a scheduling scheme using the Master-Worker model for concurrent computing, several issues must be considered: the scalability, the communication and synchronization overhead, and the load balancing.

All the policies, where a single node (the master) is in charge with the work distribution and collecting the results, may cause degradation in performance as the problem size increases. This means that for a large size problem (and for a large number of processors) the master could become a bottleneck. There are two major kinds of overhead in simple Master-Worker architectures. The first one is: if workers send back the computed results, it may take a long time to gather the computed results. The communication overhead is expensive in a distributed memory system such as a cluster, where long communication latency can be encountered. Another kind of overhead occurs when many workers send work requests at the same time and only one worker can be served from the request queue and the others have to wait. This is time consuming, especially in the case of a single request queue, when the task scheduler is slow or the scheduling schemes are complicated.

It is known that distributed policies usually do not perform as well as the simple Master-Worker policies (i.e. using a single master), for small problem sizes and small number of workers. This is because the algorithm and the implementation of distributed schemes usually add a non-trivial overhead.

We consider a logical hierarchical architecture as a good model for scalable systems and we propose a new hierarchical approach for addressing the bottleneck problems in the Master-Worker schemes.

Instead of making one master process responsible for all the workload distribution, several master processes are introduced. Thus, the hierarchical structure contains a lower level, consisting of worker processes, and several superior levels, of master processes. On top, the hierarchy has an overall supermaster. The workers’ role is to perform the computations following a Master-Worker self-scheduling method for the problem that is to be solved. This scheme is called a Hierarchical Distributed Scheme.

Figure 2 shows this design for two levels of master processes, one supermaster and two master nodes. The task scheduler resides in the supermaster and it uses distributed scheduling schemes (DTSS/DFSS/DGSS) to compute small scheduled chunks for each master node and send to master nodes’ Task Pools. When the Task Pool of a master node is empty, it asks for more work (from the supermaster) in order
to fill the Task Pool until there is no more work. The master node accepts a worker request, places it into the request queue and gets a scheduled chunk from the Task Pool and serves the top request from Request Queue. Also, the master node is in charge of gathering the computed results from workers. There are multiple Request Queues and Result Collectors distributed in different master nodes, which can share the responsibilities.

The hierarchical distributed scheduling scheme is described as follows:

Supermaster:
1. Compute $V_j$ for each Worker
   - (a) Receive Workers’ Speed($P_j$) from Masters;
   - (b) Compute all $V_j$;
2. Assign work to Masters
   - (a) While there are unassigned tasks, if a Master request arrives, put it in the queue;
   - (b) Pick a request from the queue and get the Workers virtual power $V_j$ under the requesting Master. Using distributed self-scheduling schemes (i.e. DTSS, DFSS and DGSS) to compute small scheduled chunks for each Worker under the requesting Master. Then Master may store the chunks into its Task Pool.

Master:
1. Compute $V_j$ for each Worker
   - (a) Receive Speed($P_j$) from its Workers;
   - (b) Send these Speed($P_j$) to Super Master;
2. Request work to Super Master to fill Task Pool;
3. Assign work to Workers;
   - (a) If there are unassigned tasks, if a Worker request arrives, put it into the Request Queue. Pick a request from the Request Queue, Result Collector receives computing results first. Then get a chunk from Task Pool and send this chunk to requesting worker;
   - (b) If there are not unassigned tasks, request more work to Supermaster;
   - (c) If there is no work left, go back to (2);

Worker :
1. Send Speed($P_j$) to its Master;
2. Send a request to its Master.

(3) Wait for a reply;
IF (There is unassigned work)
{
  Compute the new work;
  Return the results and send another request;
  Go back to (2);
}
ELSE
  Terminate;

V. IMPLEMENTATION AND EXPERIMENTS

A. Applications

- Mandelbrot Set [28]

The Mandelbrot Set is a doubly nested loop without dependencies. The computation of one column of the Mandelbrot matrix is considered the smallest schedulable unit. The Mandelbrot Set loop is an irregular loop in terms of unpredictable iteration task sizes. Thus this kind of loop causes load imbalance in the parallel computation. The following loops are used for computing the Mandelbrot Set.

MSetLSM(MSet,nx,ny,xmin,xmax,ymin,ymax,maxiter)
BEGIN
  FOR iy = 0 TO ny-1 DO
    cy = ymin+iy*(ymax - ymin)/(ny - 1)
    FOR ix = 0 TO nx-1 DO
      cx = xmin+ix*(xmax - xmin)/(nx - 1)
      MSet[ix][iy]=MSetLevel(cx,cy,maxiter)
    END FOR
  END FOR
END

MSetLevel(cx,cy,maxiter)
BEGIN
  x = y = x2 = y2 = 0.0, iter = 0
  WHILE(iter<maxiter)AND(x2+y2<2.0)DO
    temp = x2 - y2 + cx
    y = 2*x*y + cy
    x = temp
    x2 = x*x
    y2 = y*y
    iter = iter + 1
  END WHILE
RETURN(iter)
END

- Adjoint Convolution

This application involves computation of decreasing task sizes. Thus, it can cause load imbalance in the parallel computation. The $i_{th}$ iteration’s time is $O(N^2 - i)$.

BEGIN
  FOR I = 1 TO N * N DO
    FOR J = I TO N * N DO
      A(I) = A(I) + X * B(J) * C(J - I)
    END FOR
  END FOR
provides a 1GB/sec point to point bandwidth.

by InfiniBand technology in a full-CLOS topology which
memory limit is 32 GB per node. The nodes are interconnected
Opteron Quad-Core 64-bit processors and 16 cores total. The
nodes’ Operating System is Linux and the nodes are managed
Computing Center) in University of Texas at Austin. The
Ranger cluster system is located at TACC (Texas Advanced

B. Platform

We use the Ranger cluster system as our platform. The
Ranger cluster system is located at TACC (Texas Advanced Computing Center) in University of Texas at Austin. The
nodes’ Operating System is Linux and the nodes are managed by Rocks 4.1 cluster toolkit. Each node has four AMD
Opteron Quad-Core 64-bit processors and 16 cores total. The
memory limit is 32 GB per node. The nodes are interconnected by InfiniBand technology in a full-CLOS topology which
deprovides a 1GB/sec point to point bandwidth.

VI. RESULTS

In this section, we compare the performance of the various
schemes, non-hierarchical (single master) and hierarchical (2
masters, 4 masters, 8 masters, 16 masters) and with a number
of workers (processors) from 256 to 8,192. The Mandelbrot
Set computation domain is [-2.0, 2.0] × [-2.0, 2.0] and its size
is 200K × 200K. The Adjoint Convolution has a size of 800
× 800 and the arrays are generated randomly.

In order to avoid too many small chunks at the end of
scheduling which may introduce unnecessary synchronization
overhead, we add a threshold to terminate if the chunk size
drops below it. In our experiment, the threshold equals 5,
which means the master can not assign a chunk with size
less than 5, except possibly the last chunk.

We test the HDTSS, HDFSS and HDGSS schemes dis-
cussed in section IV. All workers are treated (by the schemes)
as having the same computing power. The execution time is
measured in seconds.

The performance presented in Figure 3 and Figure 4 is orga-
nized from left to right in doubling numbers of workers using
HDTSS, HDFSS, HDGSS schemes for Mandelbrot Set and
Adjoint Convolution. It can be observed that the hierarchical
distributed scheme with more master nodes can achieve better
performance improvement. The 2-Masters’ model scales well
upto 512 workers, however past this point the execution time
does not decrease as the number of workers increases. The 16-
Masters shows the best scalability because when the number
of workers doubles, the execution time is halved. The load
balancing issue can be solved by the original self-scheduling
schemes (TSS, FSS and GSS), which have been demonstrated
to be effective scheduling schemes in both shared memory
systems and distributed memory systems. In our experiments,
the performance of HDFSS and HDGSS are a little better
than HDTSS because HDFSS and HDGSS may generate more
small chunks at the end to balance the workload across the
computation. These two schemes introduce more synchroniza-
tion problems (i.e. more chunks and more work requests).
However, hierarchical distributed schemes have distributed
queues and these synchronization points take really little time,
which can lead to good load balancing.

In Figure 5 and Figure 6, we show the non-overlapped
Fig. 5. The non-overlapped communication and synchronization overhead $T_{\text{overhead}}'$ of Mandelbrot Set

Fig. 6. The non-overlapped communication and synchronization overhead $T_{\text{overhead}}'$ of Adjoint Convolution

Fig. 7. The speedup of Mandelbrot Set using hierarchical distributed schemes

Fig. 8. The speedup of Adjoint Convolution using hierarchical distributed schemes
communication and synchronization overhead \(T^\prime_{\text{overhead}}\) with increasing number of workers, \(T^\prime_{\text{overhead}} = T_{\text{total}} - T_{\text{computation}}\). In our experiments, there are some overlapping between computation and communication for efficient computing. The computation time can be measured exactly but the total communication overhead is difficult to capture. So we use \(T^\prime_{\text{overhead}}\) to represent the sum of non-overlapped communication and synchronization overhead. In our results, when more masters are used, \(T^\prime_{\text{overhead}}\) are smaller. The 16-masters’ model has the best performance for our results, because it has more result collectors and distributed task queues residing on master nodes. This helps to reduce the synchronization overhead and especially the communication overhead, which may be the most slowest part for large problems in distributed memory systems.

Figure 7 and Figure 8 shows the speedup of the three hierarchical distributed schemes for Mandelbrot Set and Adjoint Convolution. The x-axis represents \(\log_2(p)\). The speedup is computed by \(S_p = \frac{T_1}{T_p}\). \(T_1\) is the execution time for the non hierarchical distributed scheme with 256 workers, \(T_p\) is the execution time with \(p\) workers. It can be observed that as the number of workers increases, the 16-masters’ hierarchical distributed scheme scales well up to 8,192 workers. The non hierarchical distributed scheme’s scalability is the worst.

VII. CONCLUSION AND FUTURE WORK

In this paper, we studied and implemented (in MPI) hierarchical distributed loop scheduling schemes. We showed that non hierarchical loop scheduling algorithms with Master-Worker model does not scale well when there are hundreds of workers in the system. We proposed and implemented a hierarchical distributed model for self-scheduling schemes on large-scale clusters that maintains the load balancing properties of some well known loop scheduling schemes, and also shows better scalability on large-scale clusters. There are past results on thread based and work-stealing based implementation of loop parallelization \[29][30]. In the future, we plan to study these approaches in our method.

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