Solution of Sparse Positive Definite Systems on a Shared-Memory Multiprocessor

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ON A SHARED-MEMORY MULTIPROCESSOR

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ABSTRACT

Algorithms and software for performing sparse Cholesky decomposition and using the Cholesky factors in the solution of sparse symmetric positive definite systems on serial computers have reached a high state of development. In this paper we present algorithms for performing these two phases on a shared-memory multiprocessor computer, along with some numerical experiments demonstrating their performance on a Sequent Balance 8000 system.

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1. Introduction

This article deals with the problem of solving a large sparse positive definite system of equations $Ax = b$ on a shared memory multiprocessor system. In [3], a parallel algorithm was developed for solving dense positive definite systems in such an environment, so this article can be regarded as a sequel to that work, in which the sparsity of the problem is addressed and exploited.

The solution of large sparse positive definite systems typically involves four distinct steps [6]:

1. **Ordering**
   Find a good ordering $P$ for $A$. That is, find a permutation matrix $P$ so that $PAP^T$ has a sparse Cholesky factor $L$ (i.e., $PAP^T = LL^T$).

2. **Symbolic factorization**
   Determine the structure of the Cholesky factor of $PAP^T$, and set up a data structure that exploits the sparsity of $L$.

3. **Numerical factorization**
   Place the elements of $PAP^T$ into the data structure, and then compute $L$.

4. **Triangular solution**
   Using the computed $L$, solve the triangular systems $Ly = Pb$, $L^T z = y$, and then set $x = P^T z$.

The problems of implementing an ordering algorithm and a symbolic factorization algorithm on a multiprocessor machine are major projects and will be considered elsewhere. In this paper we develop and test parallel algorithms for steps 3 and 4.

The computing regime we adopt employs the notion of a pool of tasks whose parallel execution is controlled by a self-scheduling discipline [7]. In our context, the tasks are those computations associated with columns of the coefficient matrix, and thus have a well-defined order associated with them.

In some parallel algorithms, specific tasks are mapped onto specific processors in advance of initiating the computation. In this situation, effective (static) load balancing among the processors requires that the distribution of work be reasonably uniform. Self-scheduling can be regarded as a mechanism for implementing dynamic load balancing; $p$ processes are initiated to perform $T$ tasks $(p \leq T)$. When a given process completes a task, it checks to see if any unassigned tasks remain, and if so it is assigned the next one. Thus, if a process happens to have drawn a relatively small task, it will become free to perform another one sooner than a process occupied by a larger task. In this way, processors tend to be kept busy even if the tasks vary in their computational requirements.
This self-scheduled pool-of-tasks approach is flexible in that it is not very strongly dependent on the number of processors available. However, its effective use depends upon certain computational and hardware characteristics.

Since the pool of tasks must be made available to each processor, there must be either a significant amount of shared global memory, or very high communication bandwidth among processors. This is particularly important if the tasks involve a substantial amount of data and/or if the computation involved with each task is small compared to the time required for the processor to acquire the task.

This latter issue of granularity is complicated in the context of using the self-scheduling pool-of-tasks approach. On one hand, unless the overhead associated with the initiation of a process is small, one would prefer to have the computation associated with each task quite substantial, so as to amortize the initiation overhead over a large amount of computation. On the other hand, large tasks imply that there will be fewer of them, and the problem of allocating them to processors in a way that will keep them all gainfully employed over time will be more difficult.

In light of the remarks above, we will assume that $T \gg p$, and that the balance between hardware speed and task size is such that the time associated with assigning a task to a processor is low compared to the time it takes to perform that task. Computers for which such assumptions are reasonable include the Cray X-MP, Elxsi, Encore, Flex, and the Sequent, each of which has a moderate number of processors (4-30) and considerable memory, all attached to a very fast bus.

An outline of the paper is as follows. In Section 2 we briefly review the ideas in [3] and some basic results about sparse Cholesky factorization. We then describe a parallel version of the algorithm. In Section 3 we consider the parallel solution of sparse triangular systems. In Section 4 we present numerical experiments which demonstrate the performance characteristics of the algorithms. These were performed on a Sequent multiprocessor, which has a shared-memory architecture. Finally, Section 5 contains our concluding remarks.

Recent work on parallel sparse Cholesky factorization includes the following. In [1], Duff considers the implementation of the multifrontal approach for solving sparse symmetric systems on a shared-memory multiprocessor, and in [4], the authors of this paper have proposed a parallel sparse Cholesky factorization algorithm for a distributed-memory multiprocessor.
2. Parallel Sparse Cholesky Factorization

2.1. Parallel Dense Column-Cholesky

In [3], the column-Cholesky formulation is recommended for parallel Cholesky factorization of dense symmetric positive definite linear systems on a shared-memory multiprocessor. Following [3], we let $T_{col}(j)$ be the task that computes the $j$-th column of the Cholesky factor. Each such task consists of the following two types of subtasks:

1. $cmd(j, k)$ : modification of column $j$ by column $k$ ($k < j$);
2. $cdiv(j)$ : division of column $j$ by a scalar.

We first review the version of dense column-Cholesky in [3]. The self-scheduling of the tasks $T_{col}(j)$ is implemented by maintaining a vector of flags $ready$. They can be viewed as a vector of semaphores to ensure synchronization among these column tasks. The value $ready[j]$ indicates whether column $j$ is ready to be used for modification of subsequent columns. The following gives an algorithmic description of the implementation.

\[
\begin{align*}
\text{for } j & := 1 \text{ to } n \\
\quad & \text{ready}[j] := 0; \\
\text{for } j & := 1 \text{ to } n \\
\quad & \text{schedule } T_{col}(j); \\
\end{align*}
\]

The task $T_{col}(j)$ can then be implemented as follows.

\[
\begin{align*}
\text{for } k & := 1 \text{ to } j - 1 \\
\text{begin} \\
\quad & \text{wait until } ready[k] = 1; \\
\quad & \text{do } cmd(j, k); \\
\text{end} \\
\quad & \text{do } cdiv(j); \\
\quad & \text{ready}[j] := 1; \\
\end{align*}
\]

It should be clear that the columns of the matrix become $ready$ in order of the column subscripts, so that the tasks are completed in the sequence:

$$ T_{col}(1), T_{col}(2), \cdots, T_{col}(n) .$$

We now introduce a different version of parallel dense factorization. It is the basis for the parallel sparse column-Cholesky algorithm to be described in Section 2.2, but is easier to understand. Its purpose here is to facilitate a better
appreciation of the underlying idea for the sparse case.

The new formulation maintains a set of $n$ non-overlapping linked lists, one for each column of the matrix. Since they are non-overlapping, an $n$-vector $link$ will be enough to implement them. In the following discussion, $link^m[j]$ denotes the $m$-th element in the linked list for column $j$; for example, $link^3[j] = link[link[link[j]]]$. We assume that the lists are null-terminated, so that the $j$-th list is given by:

$$link[j], link^2[j], \ldots, link^r[j],$$

where for some $r$, $link^{r+1}[j] = 0$. These linked lists are often used in the sequential algorithms for sparse Cholesky factorization [10]. Furthermore, we define $next(j,k)$ to be the row subscript of the next nonzero in column $k$ of $L$ immediately beneath $L_{jk}$. That is, $next(j,k) = j + 1$. (In the dense case, $next(j,k)$ is independent of $k$, but $next(j,k)$ will depend on both $j$ and $k$ in the sparse case.) The new algorithm is shown below.

```plaintext
for j := 1 to n
  begin
    link[j] := 0;
    nmod[j] := j - 1;
  end
for j := 1 to n
  schedule Tcol(j);
```

Each task $Tcol(j)$ is as follows.
while $nmod[j] > 0$
begin
    wait until $link[j] > 0$;
    $k := link[j]$;  /* remove first column $k$ from $j$-th list */
    $link[j] := link[k]$;

    do $cmod(j, k)$;

    $nmod[j] := nmod[j] - 1$;
    $nextnz := next(j, k)$;
    if $nextnz \leq n$ then
        begin  /* add $k$ to $next(j, k)$-th list */
            $link[k] := link[nextnz]$;
            $link[nextnz] := k$;
        end
    end

    do $cdiv(j)$;

    $nextnz := next(j, j)$;
    if $nextnz \leq n$ then
        begin  /* add $j$ to $next(j, j)$-th list */
            $link[j] := link[nextnz]$;
            $link[nextnz] := j$;
        end
end

In this version, $nmod[j]$ is used to keep track of the number of column modifications that remain to be performed on column $j$; and it is initialized to $j-1$. For column $j$, the linked list:

$$link[j], link^2[j], \ldots$$

gives the columns that are currently ready to modify column $j$. As each (say, column $k$) is used for the column modification, it is removed from the linked list for $j$ and passed onto that for $next(j, k)$ (namely, column $j+1$ in the dense case). Moreover, after the $cdiv(j)$ operation, column $j$ is now ready to modify subsequent columns, and it is placed in the linked list for $next(j, j)$ (which is again equal to $j+1$).

Compared to the previous algorithm (using the $ready$ vector), this version has several drawbacks. Two $n$-vectors are required in the current version. More importantly, some form of critical section has to be set up during the update of the $link$ vector. This will avoid simultaneous update of the $link$ vector by different Tom tasks. Such a mechanism is not necessary in the $ready$-version. However, the
changes are important in order to take advantage of the parallelism derived from the sparsity of $L$.

2.2. Parallel Sparse Column-Cholesky

As we have pointed out in the previous subsection, general sparse matrix packages (on serial machines) often use a variant of the link-version described in Section 2.1 for sparse Cholesky factorization; examples are the Yale sparse matrix package [2] and SPARSPAK [5]. One important difference from the dense case is that next $(j, k)$ is no longer always equal to $j + 1$. Recall that next $(j, k)$ is the row subscript of the next nonzero in column $k$ of $L$ immediately beneath $L_{jk}$. Hence, next $(j, k)$ will depend on both $j$ and $k$ in the sparse case. More precisely, next $(j, k)$ depends on the structure of $L$. Thus column $k$ (after a cmod) or $j$ (after a cddiv) of $L$ will not be passed to the $(j + 1)$-st list in general, but instead will be passed to the list determined by the structure of the Cholesky factor. For simplicity, if $L_{jk}$ is the last nonzero in column $k$, we put next $(j, k) = n + 1$.

The link-version for the sparse factorization is similar to that for the dense factorization in Section 2.1. One difference is that nmod $[j]$ should be initialized to be $\eta(L_{j*})$, where $\eta(L_{j*})$ is the number of offdiagonal nonzeros in the $j$-th row of $L$. This information can be obtained easily from the structure of $L$ (which is computed in the symbolic factorization phase). Also the function next can be evaluated easily if the nonzeros of a column are stored in ascending order of the row subscripts.

Another implementational difference from the dense case is in the execution of cmod $(j, k)$. In the sparse case, the columns $L_j$ and $L_k$ are stored in a compact form. To perform cmod $(j, k)$, we need to unpack the compact form of the column $L_j$, so that modifications from other columns can be done efficiently. This implies that the processor executing the task $Tcol(j)$ requires a local working array of size $n$ to facilitate the column update operations on column $j$. However the data structure used for storing $L$ in the parallel sparse Cholesky algorithm is the same as that in the sequential sparse Cholesky algorithm, since it is stored in a global memory and is accessible by all processors.

In terms of behavior, the dense and the sparse cases differ in the order the cddiv's are performed. In the dense case, the cddiv's are performed in a sequential order. However, in the sparse case, only a few cmod's have to be applied to column $j$ and cddiv $(j)$ can be performed once all the necessary cmod's have been applied. Thus not only can the cmod's be carried out in parallel in the sparse case, some of the cddiv's may also be completed simultaneously.

In the discussion above, we have not addressed the problem of scheduling the tasks. We have assumed that the column tasks are scheduled according to the ordering of the columns. Recall from Section 1 that the columns of $A$ are ordered at the beginning so as to reduce fill in the Cholesky factor. (In our experiments,
we have used a variant of the minimum degree algorithm [8].) Thus we implicitly assume that such an ordering is appropriate for parallel computation on a shared-memory multiprocessor. In general, because the Cholesky factor is sparse, the completion time of parallel sparse Cholesky factorization will depend on how the column tasks are scheduled. It is possible to reorder the columns of the permuted matrix \( PAP^T \) so that the Cholesky factor has the same amount of fill, but when the new ordering is used to schedule the column tasks, the completion time on a shared-memory multiprocessor will be reduced. See [9] for details.

3. Parallel Sparse Triangular Solutions

In this section we consider the parallel solutions of the triangular systems

\[Lv = u \quad \text{and} \quad L^Tw = v,\]

where \( L \) and \( u \) are either given or computed elsewhere. It is assumed that the elements of \( L \) are stored column by column. We shall first describe a parallel algorithm for the backward solve and then present two parallel algorithms for the forward solve.

3.1. Parallel Backward Solve

Let \( Tw(j) \) denote the task that computes \( w_j \). A parallel algorithm for computing \( w \) is given below and it makes use of the vector \( \text{ready} \) described in the previous section.

\[
\text{for } j := 1 \text{ to } n \\
\quad \text{ready}[j] := 0; \\
\text{for } j := n \text{ down to } 1 \\
\quad \text{schedule } Tw(j); \\
\]

Here each task \( Tw(j) \) is given as follows.

\[
\text{for each offdiagonal nonzero } L_{kj} \text{ in the } j\text{-th column, do} \\
\quad \text{begin} \\
\quad \quad \text{wait until } ready[k] = 1; \\
\quad \quad \text{do } \text{wmod}(j, k); \\
\quad \text{end} \\
\quad \text{do } \text{wdiv}(j); \\
\quad \text{ready}[j] := 1; \\
\]

The functions \( \text{wmod}(j, k) \) and \( \text{wdiv}(j) \) are similar to \( \text{cmod} \) and \( \text{cdiv} \) in the Cholesky factorization respectively. More precisely, \( \text{wmod}(j, k) \) modifies \( v_j \) by \( w_k \) using \( L_{kj} \), and \( \text{wdiv}(j) \) divides \( v_j \) by \( L_{jj} \) to yield the result \( w_j \). Note that once a component \( w_j \) is computed, it can be used to modify any \( v_i, i < j \). The flag
ready[j] is used to signal when w_j is available. The parallel algorithm described above is efficient in terms of accessing the elements of \( L \), since \( L \) is stored column by column.

When \( L \) is dense, the \( w_{mod} \)'s can be performed in parallel, but the \( w_{div} \)'s are completed sequentially. However, when \( L \) is sparse, \( w_{mod}(j,k) \) is performed only when \( L_{kj} \) is nonzero. As a result, not only can the \( w_{mod} \)'s be carried out in parallel, some of the \( w_{div} \)'s may also be completed simultaneously. Thus the algorithm described above exploits the parallelism inherent in the backward solve and that provided by the sparsity of \( L \).

3.2. Parallel Forward Solve

Now we consider the problem of solving the lower triangular system \( Lv = u \) in parallel. We first recall that the elements of \( L \) are assumed to be stored column by column. Let \( Tv(j) \) denote the task that computes \( v_j \). In the following discussion, \( w_{mod}(j,k) \) modifies \( u_j \) by \( v_k \) using \( L_{jk} \), and \( w_{div}(j) \) divides \( u_j \) by \( L_{jj} \) to yield the result \( v_j \).

We shall consider the dense case first. We note that \( v_j \) can be computed only when \( u_j \) has been modified by the product of \( v_i \) and \( L_{ji} \), \( 1 \leq i \leq j - 1 \). This requires accessing the elements of \( L \) by rows. Note that our ultimate goal is to develop a parallel algorithm for sparse forward solve. Since we assume that, when \( L \) is sparse, the nonzeros are stored column by column in a compact data structure, it is awkward to access the elements by rows. This difficulty can be alleviated by using the ideas in the link-version of the parallel algorithm for Cholesky factorization given in Section 2. That is, the nonzeros in row \( j \) of \( L \) that will be used to modify \( u_j \) are put in a linked list and the linked lists are updated after these nonzeros are used. However, it should be noted that the amount of computing is small in forward solve compared to the overhead (for synchronization purposes and updating the linked lists) involved. Indeed, preliminary experiments on a Sequent multiprocessor have shown that the performance of this approach is poor.

We now describe a second approach for parallel forward solve. It makes use of only \( n_{mod} \) and it is column-oriented. The algorithm is shown below. (Recall that \( \eta(L_{j*}) \) denotes the number of offdiagonal nonzeros in row \( j \) of \( L \).)

```plaintext
for j := 1 to n
    nmod[j] := \( \eta(L_{j*}) \);
for j := 1 to n
    schedule Tv(j);
```

Here each task \( Tv(j) \) is defined as follows.
wait until nmod[j] = 0;
do vdiv(j);
for each offdiagonal nonzero L_{kj} in column j, do
begin
    do vmod(k,j);
    nmod[k] := nmod[k] - 1;
end

In this version, once v_{kj} is computed, it can be used, together with column j of L, to modify the right hand side vector. Thus accessing the elements of L is very efficient.

Obviously, some form of critical section must be set up when performing the \textit{vmod}'s and decrementing \textit{nmod}. This is particularly important in the sparse case since the sparsity of L may cause a component of u to be modified simultaneously by several components of v. Thus, in general, there must be some form of synchronization lock for each component of u. The second algorithm therefore requires the same amount of storage as the link version (assuming a synchronization lock and an integer location both require the same amount of space), but the overhead is smaller (since no linked lists have to be maintained). However, the second algorithm suffers from the drawback that it requires n synchronization locks (n is the order of L). There will be a difficulty if the multiprocessor system provides only a small number of such locks.

4. Numerical Results

The numerical experiments were performed on a Sequent Balance 8000, which is a multiprocessor system with shared memory. On our system, there are 8 processors attached to a high speed bus, and these processors share 8M bytes of global memory. The operating system running on the Sequent multiprocessor is Dynix, which is a variant of the Unix operating system. Special library functions are provided for creating multiple cooperating processes and for synchronizing these processes.

The parallel algorithms described in this paper were implemented in FORTRAN and the programs were compiled using the Sequent FORTRAN compiler. The programs have been tested on p processors, where 1 \leq p \leq 7 (the eighth processor was reserved for operating system functions). The test problems used in the experiments were matrices defined on a sequence of L-shaped finite element meshes, and these matrices have been reordered using a variant of the minimum degree algorithm [8]. For comparison purposes, we have run SPARSPAK, which is a software package for solving large sparse symmetric positive definite systems on serial machines [5], on a single processor in order to obtain the "sequential" times. The results are provided in Table 4.1.
Table 4.1: Performance of the sequential algorithms.
(Times are in seconds.)

| \( n \) | \(| A |\) | Sequential Algorithms |
|----------|--------|-----------------------|
|          |        | Factorization | Forward Solve | Backward Solve |
| 265      | 1753   | 1.350       | .150         | .150          |
| 406      | 2716   | 2.950       | .267         | .267          |
| 577      | 3889   | 4.917       | .400         | .417          |
| 778      | 5272   | 8.350       | .600         | .600          |
| 1009     | 6865   | 11.700      | .817         | .817          |
| 1270     | 8668   | 16.433      | 1.083        | 1.083         |
| 1561     | 10681  | 23.733      | 1.433        | 1.400         |
| 1882     | 12904  | 32.950      | 1.800        | 1.783         |
| 2233     | 15337  | 42.450      | 2.217        | 2.167         |
| 2614     | 17980  | 55.567      | 2.733        | 2.650         |
| 3025     | 20833  | 67.817      | 3.233        | 3.117         |
| 3466     | 23896  | 92.583      | 3.917        | 3.800         |

Table 4.1: Performance of the sequential algorithms.
(Times are in seconds.)

The results of the multiple-processor experiments are presented in Tables 4.2-4.7. In order to gain some insights into the performance of the parallel algorithms, we have computed the speedup ratios \( \sigma \), which are defined to be

\[
\sigma = \frac{\text{time required by the sequential algorithm}}{\text{time required by the parallel algorithm on } p \text{ processors}}.
\]

Thus a parallel algorithm on \( p \) processors has a good performance if the speedup ratios are close to \( p \).

Table 4.2 contains the execution times (in seconds) and speedup ratios of the parallel sparse numeric factorization algorithm on \( p \) processors, for \( 2 \leq p \leq 7 \). It is worthwhile to point out that the speedup ratios on a fixed number of processors improve as the problem size increases. When \( p = 2 \), the speedup ratios approach 1.8 for our set of test problems. Thus the efficiency (which is defined to be \( \sigma / p \)) approaches 90%. On 7 processors, the efficiency approaches 77% and we expect it to improve as the problem size increases. Table 4.3 contains the performance statistics of running the parallel sparse numeric factorization algorithm on only 1 processor. This provides an indication of the amount of synchronization overhead in the parallel algorithm. As one can see, the overhead is fairly small. From these results we conclude that our parallel sparse numeric factorization algorithm is capable of producing good efficiencies for large enough problems.
Table 4.2: Performance of the parallel sparse numeric factorization algorithm on $p$ processors. (Times are in seconds).

<table>
<thead>
<tr>
<th>$n$</th>
<th>$p = 2$</th>
<th>$p = 3$</th>
<th>$p = 4$</th>
<th>$p = 5$</th>
<th>$p = 6$</th>
<th>$p = 7$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>time</td>
<td>$\sigma$</td>
<td>time</td>
<td>$\sigma$</td>
<td>time</td>
<td>$\sigma$</td>
</tr>
<tr>
<td>265</td>
<td>.883</td>
<td>1.53</td>
<td>.667</td>
<td>2.02</td>
<td>.567</td>
<td>2.38</td>
</tr>
<tr>
<td>406</td>
<td>1.850</td>
<td>1.59</td>
<td>1.317</td>
<td>2.24</td>
<td>1.067</td>
<td>2.76</td>
</tr>
<tr>
<td>577</td>
<td>3.017</td>
<td>1.63</td>
<td>2.150</td>
<td>2.29</td>
<td>1.700</td>
<td>2.89</td>
</tr>
<tr>
<td>778</td>
<td>5.017</td>
<td>1.66</td>
<td>3.517</td>
<td>2.37</td>
<td>2.767</td>
<td>3.02</td>
</tr>
<tr>
<td>1009</td>
<td>7.000</td>
<td>1.67</td>
<td>4.917</td>
<td>2.38</td>
<td>3.867</td>
<td>3.03</td>
</tr>
</tbody>
</table>

Table 4.3: Performance of the parallel sparse numeric factorization algorithm on 1 processor. (Times are in seconds).

<table>
<thead>
<tr>
<th>$n$</th>
<th>$p = 1$</th>
<th>$\sigma$</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>time</td>
<td>$\sigma$</td>
</tr>
<tr>
<td>265</td>
<td>1.633</td>
<td>.83</td>
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<td>406</td>
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</tr>
<tr>
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<td>9.583</td>
<td>.87</td>
</tr>
<tr>
<td>1009</td>
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<td>1270</td>
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</tr>
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<td>1561</td>
<td>26.683</td>
<td>.89</td>
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<td>1882</td>
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</tr>
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<td>3466</td>
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<td>.92</td>
</tr>
</tbody>
</table>

The results for the parallel sparse forward solve algorithm (column-oriented version) are given in Tables 4.4-4.5. Note that the performance of the column-oriented algorithm is fairly good despite the fact that updating a component of the right hand side vector must be done in a synchronized manner. The efficiency is about 45%.
Table 4.4: Performance of the parallel sparse forward solve algorithm (column-oriented version) on \( p \) processors. (Times are in seconds).

<table>
<thead>
<tr>
<th>( n )</th>
<th>( p = 2 )</th>
<th>( p = 3 )</th>
<th>( p = 4 )</th>
<th>( p = 5 )</th>
<th>( p = 6 )</th>
<th>( p = 7 )</th>
</tr>
</thead>
<tbody>
<tr>
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<td>1.21</td>
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Table 4.5: Performance of the parallel sparse forward solve algorithm (column-oriented version) on 1 processor. (Times are in seconds).

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<th>( p = 1 )</th>
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Tables 4.6 and 4.7 contain the results of the parallel sparse backward solve algorithm. The performance is between those of the factorization and column-oriented forward solve algorithms. We can achieve an efficiency of about 60%. As we can see from Tables 4.3, 4.5 and 4.7, the synchronization overhead in the parallel forward and backward solves is higher than that in parallel numeric factorization. This is because the forward and backward solve algorithms require relatively little computing.
To summarize our results, we have plotted the speedup ratios versus the problem sizes in Figures 4.1-4.3.
Figure 4.1: Speedup graph for parallel sparse numeric factorization.
Figure 4.2: Speedup graph for parallel sparse forward solve. (Column-oriented version.)
5. Concluding Remarks

In this paper, we have developed algorithms for sparse Cholesky factorization and triangular solutions that are suitable for multiprocessor systems with shared memory. We have presented numerical experiments performed on a Sequent multiprocessor system to demonstrate the efficiencies of our algorithms. The results indicate that inherent parallelism in the problem and parallelism provided by sparsity can be exploited on shared-memory multiprocessor systems. Good efficiency is achieved in the sparse factorization because the synchronization overhead is small compared to the amount of computing required. However, the efficiencies in the triangular solutions are relatively poor since the amount of computing is small and is approximately the same as the synchronization overhead.

Even though the programs were written for a particular multiprocessor system with shared memory, they can be used with minor changes on any multiprocessor system with shared memory that provides synchronization primitives. Parallel algorithms for ordering and symbolic factorization are under
investigation and the results will be described elsewhere.

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