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A Supernodal Cholesky Factorization Algorithm for Shared-Memory Multiprocessors

E. G. Ng
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Engineering Physics and Mathematics Division

Mathematical Sciences Section

**A SUPERNODAL CHOLESKY FACTORIZATION ALGORITHM FOR
SHARED-MEMORY MULTIPROCESSORS**

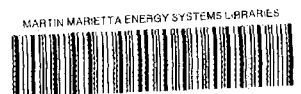
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A SUPERNODAL CHOLESKY FACTORIZATION ALGORITHM FOR SHARED-MEMORY MULTIPROCESSORS

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Abstract

This paper presents a new left-looking parallel sparse Cholesky factorization algorithm for shared-memory MIMD multiprocessors. The algorithm is particularly well-suited for vector supercomputers with multiple processors, such as the Cray Y-MP. The new algorithm uses supernodes in the Cholesky factor to improve performance by reducing indirect addressing and memory traffic. Earlier factorization algorithms have also used supernodes in this manner. The new algorithm, however, also uses supernodes to reduce the number of system synchronization calls, often by an order of magnitude or more in practice. Experimental results on a Sequent Balance 8000 and a Cray Y-MP demonstrate the effectiveness of the new algorithm. On eight processors of a Cray Y-MP, the new routine performs the factorization at rates exceeding one Gflop for several test problems from the Harwell Boeing test collection, none of which are exceedingly large by current standards.

1. Introduction

Large sparse symmetric positive definite systems arise frequently in many scientific and engineering applications. One way to solve such a system is to use Cholesky factorization. Let A be a symmetric positive definite matrix. The Cholesky factor of A , denoted by L , is a lower triangular matrix with positive diagonal such that $A = LL^T$. When A is sparse, fill occurs during the factorization; that is, some of the zero elements in A will become nonzero elements in L . In order to reduce time and storage requirements, only the nonzero positions of L are stored and operated on during *sparse* Cholesky factorization. Techniques for accomplishing this task and for reducing fill have been studied extensively (see [16] for details). In this paper we restrict our attention to the numerical factorization phase. We assume that the preprocessing steps, such as reordering to reduce fill and symbolic factorization to set up the compact data structure for L , have been performed. Details on the preprocessing can be found in [16].

In recent years, because of advances in computer architectures, there has been much interest in the solution of large sparse linear systems on high performance computers. In particular, there have been investigations into the solution of such problems on computers with multiple processors [18]. Basically, multiprocessor systems can be classified by how their memory is organized. In a shared-memory multiprocessor system, every processor has direct access to a globally shared memory. In this case, the processors can read from or write into the same memory location simultaneously. Of course, for data integrity, writing into the same memory location at any time by more than one processor must be synchronized. Examples of shared-memory multiprocessor systems include the Cray Y-MP, Encore Multimax, Sequent Balance, and Sequent Symmetry. Another way of organizing the memory in a multiprocessor system is to give each processor its own memory to which the owner alone has direct access. For one processor to access data in another processor's memory, the two processors must communicate with each other, for example, by message passing. Examples of distributed-memory multiprocessor systems include the NCUBE 3200 and 6400, and the Intel iPSC/2 and iPSC/i860. It should be noted that there are also hybrid multiprocessor systems in which both local and shared memory are available, such as the BBN Butterfly.

In this paper, we are concerned with the factorization of a sparse symmetric positive definite matrix A on a shared-memory multiprocessor system. This paper can be regarded as a sequel to [15], in which a parallel implementation of a sequential algorithm from [16] was described. We will show however that the number of synchronization operations (i.e., locking and unlocking operations) required by the parallel algorithm in [15] is relatively high; it is proportional to the number of nonzeros in the Cholesky factor L . The object of our paper is to describe a new version of the algorithm that reduces the amount of synchronization overhead by exploiting the *supernodal* structure found in the sparsity pattern of L . (The notion of supernodes will be introduced

in Section 3.) The role of supernodes in improving both left- and right-looking sparse Cholesky factorization algorithms is well documented [1,3,5,12,25,28]. The new parallel algorithm uses supernodes to reduce memory traffic and indirect indexing operations as previous algorithms have done, which is particularly important on vector supercomputers [1,3,5]. The primary contribution of the paper is the way supernodes are used to improve the parallel efficiency of a left-looking algorithm.

An outline of the paper is as follows. Section 2 reviews the sequential and parallel factorization algorithms discussed in [15]. Section 3 describes the notion of supernodes and their usefulness in a sequential sparse Cholesky factorization algorithm. A parallel supernodal Cholesky factorization algorithm will be presented in Section 3 as well. Section 4 provides experimental results on an IBM RS/6000, a Cray Y-MP, and a Sequent Balance 8000. Finally, Section 5 contains a few concluding remarks and discusses possible future work.

2. Background material

2.1. Notation and terminology

Assume that A is an $n \times n$ symmetric and positive definite matrix, and let L denote the Cholesky factor of A . We use $L_{*,j}$ and $L_{i,*}$ to represent respectively the j -th column and i -th row of L . The sparsity structures of column j and row i of L (excluding the diagonal entry) are denoted by $Struct(L_{*,j})$ and $Struct(L_{i,*})$, respectively. That is,

$$\begin{aligned} Struct(L_{*,j}) &:= \{s > j : l_{s,j} \neq 0\}, \\ Struct(L_{i,*}) &:= \{t < i : l_{i,t} \neq 0\}. \end{aligned}$$

Assume that $l_{k,j} \neq 0$ and suppose that $l_{k,j}$ is not the last nonzero in column j of L . The function $next(k,j)$ returns the row index of the *first* nonzero beneath $l_{k,j}$ in the column $L_{*,j}$ [15]. If $l_{k,j}$ is the last nonzero in $L_{*,j}$, then we define $next(k,j)$ to be $n + 1$.

The two computational tasks occurring at each step in the Cholesky factorization are scaling a vector and subtracting a multiple of a vector from another vector. These two tasks will be denoted by *cdiv* and *cmul*, respectively [14].

```

cdiv(j):
  lj,j ← (aj,j)1/2
  for i = j + 1 to n do
    li,j ← ai,j/lj,j
  end for

```

```

cmod( $j, k$ ),  $k < j$ :
  for  $i = j$  to  $n$  do
     $a_{i,j} \leftarrow a_{i,j} - l_{j,k} l_{i,k}$ 
  end for

```

Finally, if M is an $m \times n$ matrix, then $|M|$ denotes the number of nonzero elements in M .

2.2. Sequential sparse Cholesky factorization

We begin our discussion by first reviewing a sequential general sparse Cholesky factorization algorithm, details of which can be found in [16]. The algorithm is column-oriented and is a *left-looking* algorithm. That is, when column $L_{*,j}$ is to be computed, the algorithm modifies column $A_{*,j}$ with multiples of the previous columns of L , namely $L_{*,k}$, $1 \leq k \leq j - 1$. Of course, sparsity will be exploited when A is sparse. We will assume throughout that the nonzeros of A and L are stored by columns. The sequential factorization algorithm is given in Figure 2.1. This algorithm and its variations are widely used in many sparse matrix packages, such as SPARSPAK [7].

```

for  $j = 1$  to  $n$  do
  for  $k \in Struct(L_{j,*})$  do
    cmod( $j, k$ )
  end for
  cdiv( $j$ )
end for

```

Figure 2.1: A sequential sparse Cholesky factorization algorithm.

Since the algorithm in Figure 2.1 is column-oriented and the nonzeros of L are stored by columns, its implementation is quite straightforward except for the determination of the structure of row j of L (i.e., $Struct(L_{j,*})$). Instead of computing the structure of every row of L *prior* to the factorization, the factorization algorithm itself can efficiently generate these sets *during* the factorization, as shown in Figure 2.2. For each column $L_{*,j}$, we maintain a set \mathcal{S}_j of column indices, which will contain precisely the column indices belonging to $Struct(L_{j,*})$ when the column $L_{*,j}$ is computed.

After $L_{*,j}$ has been computed, j is inserted into \mathcal{S}_q , where q is the row index of the first nonzero beneath the diagonal in column j (i.e., $q = next(j, j)$). When the algorithm is ready to compute $L_{*,q}$, it will examine \mathcal{S}_q to find the columns of L needed to modify $A_{*,q}$. Among those columns it will find $L_{*,j}$, and thus it will perform **cmod**(q, j) as required. It is easy to see that the next column of A that $L_{*,j}$ will modify

```
for  $j = 1$  to  $n$  do
   $S_j \leftarrow \emptyset$ 
end for

for  $j = 1$  to  $n$  do
  for  $k \in S_j$  do
     $cmod(j, k)$ 
     $p \leftarrow next(j, k)$ 
    if  $p \leq n$  then
       $S_p \leftarrow S_p \cup \{k\}$ 
    end if
  end for
   $cdiv(j)$ 
   $q \leftarrow next(j, j)$ 
  if  $q \leq n$  then
     $S_q \leftarrow S_q \cup \{j\}$ 
  end if
end for
```

Figure 2.2: A sequential sparse Cholesky factorization algorithm, with the generation of row structure.

is given by $p = \text{next}(q, j)$. Hence, the algorithm puts j in \mathcal{S}_p for use when it later computes $L_{*,p}$. More informally, immediately after $L_{*,j}$ has been computed it begins “migrating” from one column of A to another as determined by the values of $\text{next}(*, j)$ (or equivalently the structure of $L_{*,j}$). The columns visited by $L_{*,j}$ are exactly those that must be modified by $L_{*,j}$. At any point during the factorization, $\mathcal{S}_i \cap \mathcal{S}_j = \emptyset$ for $i \neq j$. Consequently, the sets \mathcal{S}_j ($1 \leq j \leq n$) can be stored economically as linked lists using a single integer array of length n . This is the primary reason for generating the sets $\text{Struct}(L_{j,*})$ in this manner.

2.3. Sources of parallelism

As indicated in [15], there are two sources of potential parallelism in sparse Cholesky factorization. The first one is in performing *cmod* operations with the same “updating” column. Suppose $\text{Struct}(L_{*,j}) = \{i_1, i_2, \dots, i_\rho\}$, with $j < i_1 < i_2 < \dots < i_\rho$. When $L_{*,j}$ has been computed, columns i_1, i_2, \dots, i_ρ of A have to be modified by $L_{*,j}$. These *cmod*'s are independent: they can be performed simultaneously or in any order. Thus, if there are enough processors and if the nonzero entries of $L_{*,j}$ are available to these processors, the operations $\text{cmod}(i_1, j), \text{cmod}(i_2, j), \dots, \text{cmod}(i_\rho, j)$ can be performed concurrently. The independence of *cmod*'s using the same updating column but different target columns has nothing to do with the sparsity of L ; indeed, they are the primary source of parallelism in a *dense* column-based factorization.

Sparsity in L gives rise to large-grained parallelism that is not available in a dense factorization. Consider columns $L_{*,k}$ and $L_{*,j}$ where $j > k$. We shall say that $L_{*,j}$ *depends on* $L_{*,k}$ if $L_{*,j}$ cannot be completed until after $L_{*,k}$ has been completed. When neither $L_{*,j}$ depends on $L_{*,k}$ nor $L_{*,k}$ depends on $L_{*,j}$, the two columns are said to be *independent* of one another. The column dependencies are very simple when L is dense: since computation of $L_{*,j}$ requires modification of $A_{*,j}$ by a multiple of every column $L_{*,k}$ where $k < j$, $L_{*,j}$ depends on every such column $L_{*,k}$. To identify column dependencies in the sparse case, we introduce *elimination trees*.

Consider the Cholesky factor L . For each column $L_{*,j}$ having off-diagonal nonzero elements, we define $\text{parent}[j]$ to be the row index of the first off-diagonal nonzero in that column; that is, $\text{parent}[j] = \text{next}(j, j)$. For convenience, we define $\text{parent}[j]$ to be j when column $L_{*,j}$ has no off-diagonal nonzeros. The *elimination forest* of L is a graph \mathcal{T} with $\{1, 2, \dots, n\}$ as its node set, and an edge connecting i and j if and only if $j = \text{parent}[i]$ and $i \neq j$ [21,26]. It is also easy to show that \mathcal{T} is a tree if and only if the matrix A is *irreducible*. Without loss of generality, we will assume from now on that the given matrix A is irreducible, so that \mathcal{T} is indeed an *elimination tree*. We assume familiarity with the standard terminology associated with rooted trees: e.g., root, parent, child, ancestor, and descendant. We use the notation $\mathcal{T}[i]$ to denote the *subtree* rooted at node i ; that is, $\mathcal{T}[i]$ is a tree consisting of i and all of its descendants in \mathcal{T} .

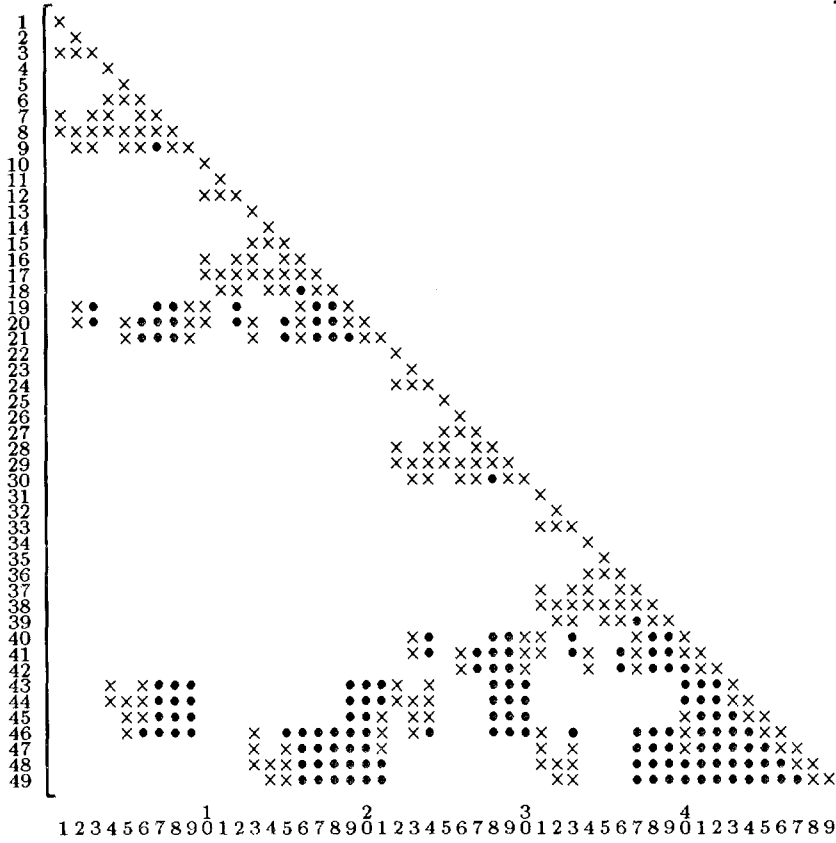


Figure 2.3: A matrix example defined on a 7×7 nine-point grid ordered by nested dissection. (each \times and \bullet refers to a nonzero in A and a fill entry in L , respectively.)

Consider the example in Figure 2.3, which contains the matrix and Cholesky factor associated with a 7×7 nine-point grid ordered by the nested dissection algorithm [13]. In the figure, each \times is a nonzero entry in the matrix A , and each \bullet is a fill entry in the Cholesky factor L . The reader may verify that the tree shown in Figure 2.4 is the elimination tree of the matrix L shown in Figure 2.3.

One of the many uses of elimination trees in sparse matrix computation is the analysis of column dependencies in sparse Cholesky factorization. (A survey of elimination trees and their applications in sparse matrix computations is contained in [22].) A key observation [21,26] is that $Struct(L_{j,*}) \subseteq T[j]$; that is, every $k \in Struct(L_{j,*})$ is a descendant of j in the elimination tree. Of course, column j of L cannot be completed until all columns in $Struct(L_{j,*})$ have been completed. Recursive application of this observation to the descendants of j demonstrates that column j of L cannot be completed until the columns associated with *all* descendants of j (i.e., *all* members of $T[j] - \{j\}$) have been completed. Moreover, $L_{*,j}$ does not depend on any other columns. Hence, columns i and j are independent if and only if $T[i]$ and $T[j]$ are disjoint subtrees. For

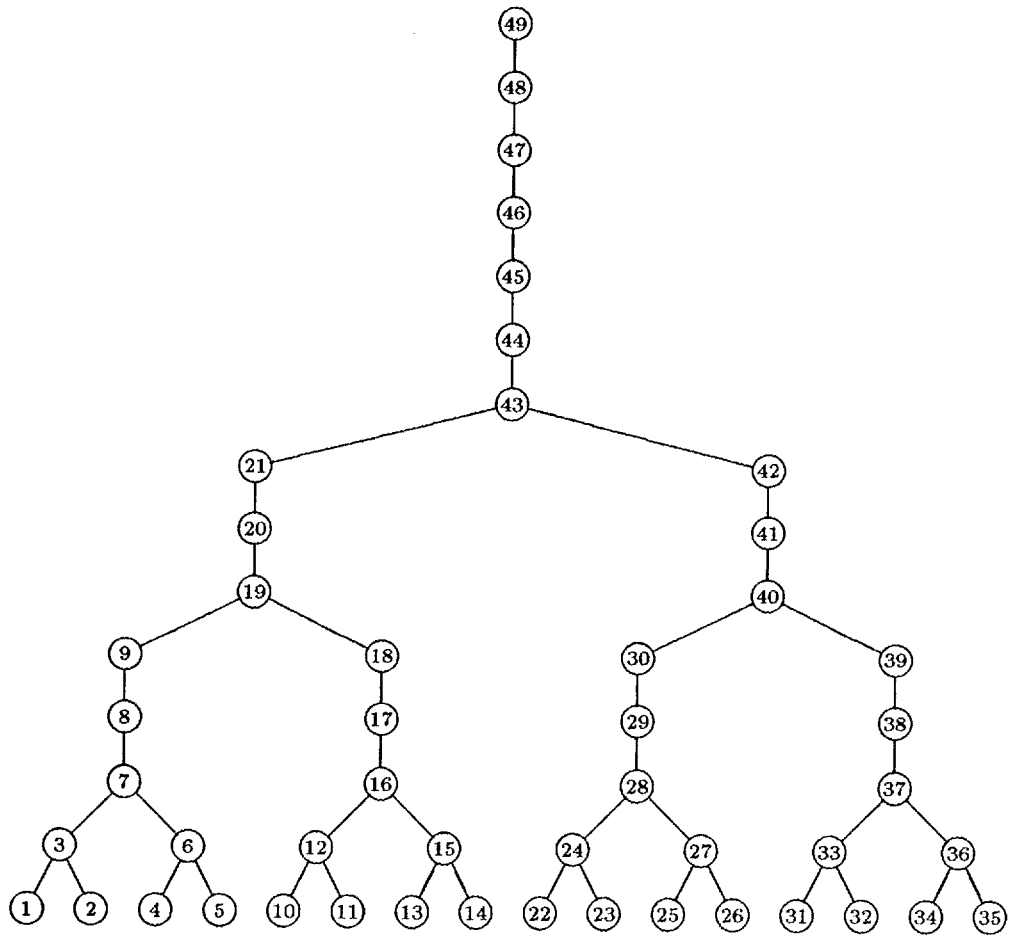


Figure 2.4: Elimination tree for the matrix shown in Figure 2.3.

example, column 41 in Figure 2.4 depends on columns 22 – 40, and depends on no other columns of the matrix. Columns 41 and 21 are independent because $T[41]$ and $T[21]$ are disjoint subtrees.

2.4. Parallel sparse Cholesky factorization

We now describe an algorithm for shared-memory multiprocessor systems that exploits these two sources of parallelism. (The algorithm was introduced in [15].) The task of computing column $L_{*,j}$ is referred to as a *column task* in the computation and is denoted by $Tcol(j)$. More precisely,

$$Tcol(j) := \{cmod(j, k) \mid k \in Struct(L_{*,j})\} \cup \{cdiv(j)\}.$$

The parallel algorithm maintains a pool of column tasks, and each processor will be responsible for performing a subset of these column tasks. The assignment of column tasks to processors is dynamic. When a processor is free, it will get a column task from the pool, perform the necessary *cmod* operations, and then carry out the required *cdiv* operation. When the processor has finished a column task, it will get another column task from the pool. Efficient implementation of this dynamic scheduling strategy requires that the pool of tasks be made available to all processors. This is particularly appropriate for shared-memory multiprocessor systems. This approach usually results in good load-balancing, as might be expected.

The parallel algorithm in [15] is presented in Figure 2.5. A few comments on the parallel algorithm are in order. First, note that it is quite similar to the algorithm in Figure 2.2. Second, we assume that the data reside in a globally-shared memory so that every processor can access the entire set of data. Third, since every processor will access the pool of tasks \mathcal{Q} , popping a column task from \mathcal{Q} is a critical section and must be performed in a synchronized manner.

Fourth, updating an index set \mathcal{S}_p requires another critical section since \mathcal{S}_p may be simultaneously updated by more than one processor. In Figure 2.5, we have used two primitives, *lock* and *unlock*, to synchronize this operation. The first primitive, *lock*, signals the beginning of a critical section and allows only one processor to proceed. If there is already a processor executing the critical section, a second processor attempting to enter the same section must wait until the first processor has exited the section. The second primitive, *unlock*, signals the end of a critical section, and its execution by one processor permits another processor to enter the critical section. The number of synchronization operations required to maintain the pool of tasks is $O(n)$. It is easy to see that the number of synchronization calls required to update each set \mathcal{S}_i is $O(|L_{i,*}|)$. Thus, the total number of synchronization calls required in the parallel algorithm is $O(|L|)$.

Global initialization:
 $Q \leftarrow \{Tcol(1), Tcol(2), \dots, Tcol(n)\}$
 for $j = 1$ **to** n **do**
 $S_j \leftarrow \emptyset$
 end for

Work performed by each processor:
 while $Q \neq \emptyset$ **do**
 pop $Tcol(j)$ from Q
 while column j requires further *cmod*'s **do**
 if $S_j = \emptyset$ **then**
 wait until $S_j \neq \emptyset$
 end if
 lock
 obtain k from S_j
 $q \leftarrow next(j, k)$
 if $q \leq n$ **then**
 $S_q \leftarrow S_q \cup \{k\}$
 end if
 unlock
 $cmod(j, k)$
 end while
 $cdiv(j)$
 $q \leftarrow next(j, j)$
 if $q \leq n$ **then**
 lock
 $S_q \leftarrow S_q \cup \{j\}$
 unlock
 end if
 end while

Figure 2.5: A parallel sparse Cholesky factorization algorithm for shared-memory multiprocessor machines.

3. Supernodal Cholesky factorization algorithms

Although the results reported in [15] indicated that the parallel algorithm in Figure 2.5 achieved good speed-up ratios, the algorithms in Figures 2.2 and 2.5 are far from optimal for at least two important reasons. First, both the sequential and parallel algorithms are poor at exploiting some of the hardware features available on advanced computer architectures, in particular, the pipelined arithmetic units on current vector supercomputers. Second, the number of synchronization operations connected with critical sections in the parallel algorithm is relatively high.

In this section, we discuss the notion of *supernodes* in the Cholesky factor of a sparse symmetric positive definite matrix, and show how these supernodes can be used to improve the algorithms in Figures 2.2 and 2.5. In particular, we show how both difficulties with the algorithm in [15] can be dealt with by taking advantage of the supernodal structure.

3.1. Notion of supernodes

In the Cholesky factor of a sparse symmetric positive definite matrix, columns with the “same” sparsity structure are often clustered together. Such a grouping of columns is referred to as a *supernode*¹. We define a supernode of a sparse Cholesky factor L to be a contiguous block of columns in L , $\{p, p + 1, \dots, p + q - 1\}$, such that

$$Struct(L_{*,p}) = Struct(L_{*,p+q-1}) \cup \{p + 1, \dots, p + q - 1\}.$$

It is quite easy to show that for $p \leq i \leq p + q - 2$, $Struct(L_{*,i}) = Struct(L_{*,p+q-1}) \cup \{i + 1, \dots, p + q - 1\}$. (For details consult [23,24]). Thus, the columns of the supernode $\{p, p + 1, \dots, p + q - 1\}$ have a dense diagonal block and have *identical* structure below row $p + q - 1$. Figure 3.1 shows a set of supernodes for the matrix of Figure 2.3. The partition of the columns of L into supernodes is often referred to as a *supernode partition*.

Apparently, the term “supernode” first appeared in [5], although the basic idea behind the term was used much earlier. For example, the notion of supernodes has played an important role in improving the efficiency of the minimum degree ordering algorithm [17] and the symbolic factorization process [27]. More recently, supernodes have been used to organize sparse numerical factorization algorithms around matrix-vector or matrix-matrix operations that reduce memory traffic, thereby making more efficient use of vector registers [3,5] or cache [1,25]. They play such a role in both the serial and the new parallel Cholesky factorization algorithms presented in this section.

Note that supernode partitions are not uniquely specified in our definition. Indeed, the choices of a supernode partition depend heavily on the *maximal* sets of contiguous

¹It is convenient to denote a column $L_{*,j}$ belonging to a supernode by its column index j . It should be clear by context when j is being used in this manner.

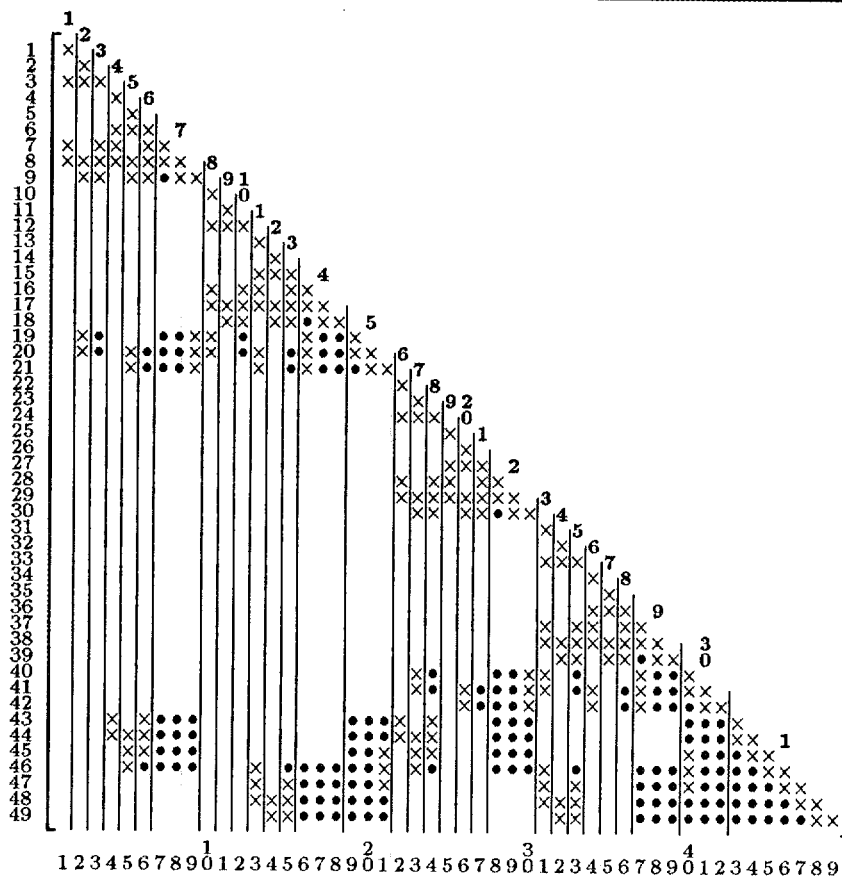


Figure 3.1: *Fundamental supernodes* in the matrix given in Figure 2.3. (Each \times and \bullet represents a nonzero in A and a fill in L , respectively. Numbers over diagonal entries label supernodes.)

columns that can be supernodes and from which one or more supernodes can be formed. We have used so-called *fundamental* supernodes in our algorithms. The set $\mathbf{K} = \{p, p + 1, \dots, p + q - 1\}$ is a fundamental supernode if \mathbf{K} is a maximal subset of contiguous columns that forms a supernode for which the following holds: for $i = 1, 2, \dots, q - 1$, the node $p + i - 1$ is the *sole child* of $p + i$ in the *elimination tree*. The notion of fundamental supernodes was introduced in [4] and was discussed extensively in [23]. The fundamental supernodes for our model problem are shown in Figure 3.1. Associated with any supernode partition is a *supernodal elimination tree*, which is obtained from the elimination tree essentially by collapsing the nodes (columns) in each supernode into a single node. The supernodal elimination tree for the partition in Figure 3.1 is shown in Figure 3.2, superimposed on the underlying elimination tree.

The primary reason for using the fundamental supernode partition in this application was pointed out in [23]: it is the coarsest supernode partition for which the supernode dependencies can be observed in the supernode elimination tree in a manner strictly analogous to the way the column dependencies are observed in the nodal elimination tree. Consequently, a fundamental supernode partition can be used more cleanly and naturally in a parallel factorization algorithm, where data dependencies are of great practical importance. Liu et al. [23] contains a full discussion of this point.

Given the matrix A , the supernode partition can be obtained by several means. When the ordering of the columns and rows of A is a minimum degree or nested dissection ordering, the partition can be obtained easily as a natural by-product of the reordering step. Otherwise, the supernode partition can be obtained directly from the structure of L after the symbolic factorization; it can also be obtained before the symbolic factorization using the algorithm given in [23].

3.2. Sequential supernodal Cholesky factorization

In this section we describe a left-looking sequential sparse Cholesky factorization algorithm that exploits the supernodal structure in L . The algorithm is not new; its variants have appeared in [5] and [25]. Let $\mathbf{K} = \{p, p + 1, \dots, p + q - 1\}$ be a supernode in L . Consider the computation of $L_{*,j}$ for some $j > p + q - 1$. Suppose column $A_{*,j}$ has to be modified by $L_{*,i}$ where $i \in \mathbf{K}$. It follows from the definition of supernodes that column $A_{*,j}$ will be modified by *all* columns of \mathbf{K} . In other words, a column $j > p + q - 1$ is either updated by *no* column of \mathbf{K} or *every* column of \mathbf{K} . This observation has some important ramifications for the performance of sparse Cholesky factorization. Loosely speaking, the columns in a supernode can now be treated as a single unit in the computation. Since the columns in a supernode have the same sparsity structure below the dense diagonal block, modification of a particular column $j > p + q - 1$ by these columns can be accumulated in a work vector using *dense* vector operations, and then applied to the target column using a single *sparse* vector operation that employs indirect addressing. Moreover, the use of *loop unrolling* in the

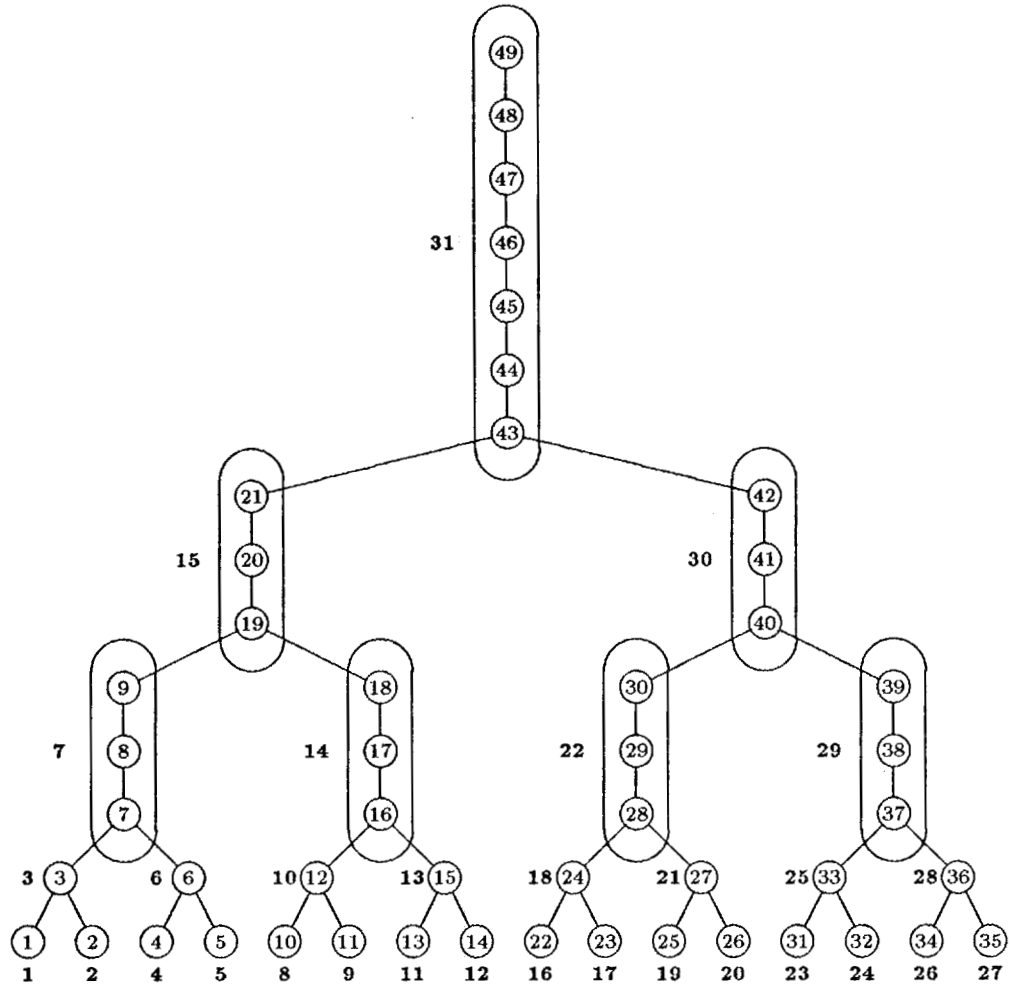


Figure 3.2: Supernodal elimination tree induced by the fundamental supernodes of the matrix shown in Figure 2.3. Ovals enclose supernodes that contain more than one node; nodes not enclosed by ovals are singleton supernodes. Bold-face numbers label supernodes.

accumulation, as described in [9], further reduces memory traffic. These issues have been addressed in detail in [1,3,5,25].

In Figure 3.3, we present a supernodal Cholesky factorization algorithm, which is quite similar to the one in Figure 2.2. In Figure 2.2, \mathcal{S}_j identifies the columns of L needed to modify $A_{*,j}$ when $L_{*,j}$ is computed. Incorporating supernodes into the algorithm, we exploit the fact that columns in the same supernode update the same set of columns outside the supernode. Thus, \mathcal{S}_j will identify the *supernodes* needed to modify $A_{*,j}$ when $L_{*,j}$ is to be computed. In Figure 3.3, we have adopted the following notation. Supernodes are denoted by bold capital letters, and in order to keep the notation simple \mathbf{K} is to be interpreted in one of two different senses, depending on the context in which it appears. In one context, \mathbf{K} is interpreted as the set of columns composing the supernode, i.e., $\mathbf{K} = \{p, p+1, \dots, p+q-1\}$. In other lines of the algorithm, the supernodes are treated as an ordered set of loop indices $\mathbf{1}, \mathbf{2}, \dots, \mathbf{K}, \dots, \mathbf{N}$, where $\mathbf{K} < \mathbf{J}$ if and only if $p < p'$, where p and p' are the first columns of \mathbf{K} and \mathbf{J} , respectively. This dual-purpose notation is illustrated in Figure 3.1, where the supernode labels are written over the diagonal entries, yet we can still write $\mathbf{30} = \{40, 41, 42\}$, for example. We denote both the last supernode and the number of supernodes by \mathbf{N} .

Suppose $\mathbf{K} = \{p, p+1, \dots, p+q-1\}$. Whenever $j > p+q-1$ and $l_{j,p+q-1} \neq 0$, the task $cmod(j, \mathbf{K})$ consists of the operations $cmod(j, k)$ where $k = p, p+1, \dots, p+q-1$. If, however, $j \in \mathbf{K}$, then $cmod(j, \mathbf{K})$ consists of the operations $cmod(j, k)$, for $k = p, p+1, \dots, j-1$. Suppose $L_{*,\ell}$ is the last column in a supernode \mathbf{K} and let $l_{j,\ell} \neq 0$. Then $next(j, \mathbf{K})$ is defined to be $next(j, \ell)$. Similarly, we define $next(\mathbf{K}, \mathbf{K})$ to be $next(\ell, \ell)$.

To reiterate the advantage of exploiting the supernodal structure of L , we note that the operation $cmod(j, \mathbf{K})$ for $j \notin \mathbf{K}$ can be accumulated in work storage by a sequence of dense vector operations (**saxpy** using the BLAS terminology [19]), after which the accumulated column modifications can be applied to the target column $L_{*,j}$ using a single column operation that requires indirect addressing. Execution of the operation $cmod(j, \mathbf{J})$ for $j \in \mathbf{J}$ is even easier, requiring no work storage or indirect addressing. In both cases, loop unrolling can be employed to reduce memory traffic, thereby improving the utilization of pipelined arithmetic units, especially on vector supercomputers. These capabilities are not available in the “nodal” Cholesky factorization algorithm in Figure 2.2.

3.3. Parallel supernodal Cholesky factorization

As far as we know, the first attempt to parallelize a supernodal Cholesky factorization algorithm was described in [28]. Using the notation in Figure 3.3, the basic idea in [28] is to partition the work in $cmod(j, \mathbf{K})$ and $cmod(j, \mathbf{J})$ evenly among the available processors. This approach is similar to that employed in the LAPACK project [2],

```
for  $j = 1$  to  $N$  do
   $S_j \leftarrow \emptyset$ 
end for

for  $J = 1$  to  $N$  do
  for  $j \in J$  (in order) do
    for  $K \in S_j$  do
       $cmod(j, K)$ 
       $q \leftarrow next(j, K)$ 
      if  $q \leq n$  then
         $S_q \leftarrow S_q \cup \{K\}$ 
      end if
    end for
     $cmod(j, J)$ 
     $cdiv(j)$ 
  end for
   $q \leftarrow next(J, J)$ 
  if  $q \leq n$  then
     $S_q \leftarrow S_q \cup \{J\}$ 
  end if
end for
```

Figure 3.3: A sequential supernodal Cholesky factorization algorithm.

where, in the interest of software portability and reliability, use of multiple processors occurs *strictly within* each call to some computationally intensive variant of a matrix-matrix multiply (BLAS3) or matrix-vector multiply (BLAS2) kernel subroutine. Hence each call to the kernel involves a fork-and-join operation. For large dense matrices, where the vectors are quite long and each call to the kernel routine typically involves a substantial amount of work, this approach is quite effective [8]. For sparse matrices, however, short vectors and a limited amount of work within a typical call to the kernel routine make it quite difficult to implement this approach in an effective manner. The performance of the code in [28] apparently suffers from these defects, and the stripmining technique used to distribute the tasks $cmod(j, \mathbf{K})$ and $cmod(j, \mathbf{J})$ among the processors greatly shortens the vector lengths, which is quite detrimental on the target machine, a Cray Y-MP multiprocessor.

In this section, we describe a different way to parallelize the supernodal Cholesky factorization algorithm in Figure 3.3. Our parallel version, shown in Figure 3.4, exploits far more of the potential parallelism than the technique used in [28].

A few comments on the algorithm in Figure 3.4 are in order. First, note that a supernode \mathbf{J} is inserted into \mathcal{S}_q , where $q = next(\mathbf{J}, \mathbf{J})$, only after the last column of \mathbf{J} has been completed. Thus, when a processor working on column q obtains \mathbf{J} from \mathcal{S}_q , *all* columns of the supernode \mathbf{J} are available for updating column q of A . This is potentially inefficient, as the columns of \mathbf{J} are not made available to update other columns of A as soon as they have been completed. An alternative to this approach is to insert \mathbf{J} into \mathcal{S}_q as soon as the *first* column of the supernode has been completed. Of course, when a processor working on column q obtains \mathbf{J} from \mathcal{S}_q , not all columns of \mathbf{J} are necessarily available. Some flags must be maintained so that the processor can determine which columns of \mathbf{J} have been completed. This approach attempts to make every column of L available to update other columns as soon as it has been completed. We have implemented both approaches. Preliminary tests indicate that the difference in performance for these two approaches is extremely small. We have chosen to use the approach shown in Figure 3.4 because it is much simpler to describe and it simplifies our implementation, especially the incorporation of loop unrolling into the code.

Another remark concerns the $cmod(j, \mathbf{K})$ operations. As in the sequential case, since all the columns in \mathbf{K} share the same sparsity structure below the diagonal entry of the last member of \mathbf{K} , these operations can be accumulated in work storage using dense vector operations. Extra care is required however when performing the $cmod(j, \mathbf{J})$ operation. Since some of the columns $j' < j$ belonging to \mathbf{J} may not be completed, a flag has to be associated with each column to record the column's current status. The flag for a particular column is set immediately after the column has been completed. Since the flag for a particular column is set only by the processor that computes the column, synchronization is not needed.

The number of `lock` and `unlock` synchronization operations required in the parallel supernodal Cholesky factorization algorithm is often much smaller than that required

Global initialization:
 $Q \leftarrow \{Tcol(1), Tcol(2), \dots, Tcol(n)\}$
 for $j = 1$ **to** N **do**
 $S_j \leftarrow \emptyset$
 end for

Work performed by each processor:
 while $Q \neq \emptyset$ **do**
 pop $Tcol(j)$ from Q
 let J be the supernode containing column j
 while column j requires further *cmod*'s **do**
 if $S_j = \emptyset$ **then**
 wait until $S_j \neq \emptyset$
 end if
 lock
 obtain K from S_j
 $q \leftarrow next(j, K)$
 if $q \leq n$ **then**
 $S_q \leftarrow S_q \cup \{K\}$
 end if
 unlock
 cmod(j, K)
 end while
 cmod(j, J)
 cdiv(j)
 if j is the last column of supernode J **then**
 $q \leftarrow next(J, J)$
 if $q \leq n$ **then**
 lock
 $S_q \leftarrow S_q \cup \{J\}$
 unlock
 end if
 end if
 end while

Figure 3.4: A parallel supernodal Cholesky factorization algorithm for shared-memory multiprocessor machines.

in the nodal version, as the following discussion shows. One of the key features in Figures 3.3 and 3.4 is that each set \mathcal{S}_j contains supernodes, rather than columns as it did in Figures 2.2 and 2.5. It is easy to see in Figure 3.3 that each supernode \mathbf{J} has to be inserted into $O(|L_{*,\ell}|)$ sets, where $L_{*,\ell}$ is the last column of \mathbf{J} . Consequently, the total number of synchronizations required in Figure 3.4 depends on the number of *compressed subscripts* [27], which is basically the number of nonzero entries in the last columns of all supernodes. To illustrate the reduction in the amount of synchronization, consider a model $m \times m$ grid problem using either a 5-point or a 9-point operator. Suppose the grid points are labelled using the nested dissection algorithm [13]. It is easy to show that the number of nonzeros in L is $O(m^2 \log m)$ [13] and the number of compressed subscripts is $O(m^2)$ [27]. Thus, the amount of synchronization in the parallel supernodal Cholesky factorization is reduced by a factor of $\log m$. Experimental results in the next section show the reductions in a collection of test problems.

3.4. Scheduling column tasks

Our discussion thus far has ignored an important issue: the scheduling of the column tasks $Tcol(j)$ on the available processors. While we have found this issue to be less important on shared-memory machines than it is on distributed-memory machines, it nonetheless deserves some attention, and is likely to be of more consequence as shared-memory machines with substantially more processors become available in the future. Again, the column-dependency information contained in the elimination tree is indispensable in dealing with this problem.

We use the following simple technique to schedule the column tasks. Before the factorization begins, all columns are placed into the column task pool *in the order in which they will be selected from the pool*. Thus, the pool of tasks can be viewed as a static queue. A scheduling, then, is essentially the order in which the column tasks are placed in the static queue.

The goal of the heuristic we use to order the column tasks in the queue is to exploit as much as possible the high-level parallelism available for sets of independent columns. Recall that $Tcol(i)$ and $Tcol(j)$ are independent column tasks if i and j belong to two disjoint subtrees in the elimination tree. Consider the example in Figure 2.4. Columns associated with the leaves of the elimination tree (1, 2, 4, 5, 10, 11, 13, 14, 22, 23, 25, 26, 31, 32, 34 and 35) are independent, since they belong to disjoint subtrees. If there are enough processors available, the corresponding column tasks can be performed in parallel *with no delays due to data-dependencies*. Similarly, columns 9, 18, 30 and 39 are independent. Thus, $Tcol(9)$, $Tcol(18)$, $Tcol(30)$ and $Tcol(39)$ can be carried out concurrently if the column tasks associated with the subtrees $\mathcal{T}[8]$, $\mathcal{T}[17]$, $\mathcal{T}[29]$ and $\mathcal{T}[38]$ have already been completed. The elimination tree therefore provides a natural way to schedule the column tasks. The order in which the column tasks are placed in the work pool \mathcal{Q} is generated by a *breadth-first, bottom-up* traversal of a post-ordered

version of the elimination tree. The same strategy was used in [15].

4. Numerical experiments

4.1. Test problems

problem	brief description
BCSSTK13	Stiffness matrix – fluid flow generalized eigenvalues
BCSSTK14	Stiffness matrix – roof of Omni Coliseum, Atlanta
BCSSTK15	Stiffness matrix – module of an offshore platform
BCSSTK16	Stiffness matrix – Corp. of Engineers dam
BCSSTK17	Stiffness matrix – elevated pressure vessel
BCSSTK18	Stiffness matrix – R.E.Ginna nuclear power station
BCSSTK23	Stiffness matrix – portion of a 3D globally triangular bldg
BCSSTK24	Stiffness matrix – winter sports arena
BCSSTK25	Stiffness matrix – 76 story skyscraper
BCSSTK29	Stiffness matrix – buckling model of the 767 rear bulkhead
BCSSTK30	Stiffness matrix – off-shore generator platform (MSC NASTRAN)
BCSSTK31	Stiffness matrix – automobile component (MSC NASTRAN)
BCSSTK32	Stiffness matrix – automobile chassis (MSC NASTRAN)
BCSSTK33	Stiffness matrix – pin boss (auto steering component), solid elements
NASA1824	Structure from NASA Langley, 1824 degrees of freedom
NASA2910	Structure from NASA Langley, 2910 degrees of freedom
NASA4704	Structure from NASA Langley, 4704 degrees of freedom
NASASRB	Structure from NASA Langley, shuttle rocket booster

Table 4.1: List of test problems.

Most of the test problems used in our numerical experiments were taken from the Harwell-Boeing Test Collection [11]. A brief description of the problems is given in Table 4.1. In the experiments, each matrix was initially ordered using an implementation of the minimum degree algorithm due to Liu [20], followed by a postordering of the elimination tree [22]. The reason for postordering the elimination tree is that the algorithms in [23] were used to compute the fundamental supernodes and the symbolic factorization, and they require such a postordering. Some statistics, such as the size of each matrix, nonzero counts for both A and L , number of subscripts required to represent the supernodal structure of L (denoted by $\mu(L)$), the number of fundamental supernodes in L , and the number of floating-point operations² are provided in Table 4.2.

²A single floating-point operation is either a floating-point addition or a floating-point multiplication, and is denoted by “flop”.

problem	n	$ A $	$ L $	$\mu(L)$	N	flops
BCSSTK13	2,003	83,883	271,671	28,621	599	58,550,598
BCSSTK14	1,806	63,454	112,267	17,508	503	9,793,431
BCSSTK15	3,948	117,816	651,222	61,614	1,295	165,035,094
BCSSTK16	4,884	290,378	741,178	50,365	691	149,100,948
BCSSTK17	10,974	428,650	1,005,859	94,225	2,595	144,269,031
BCSSTK18	11,948	149,090	662,725	116,807	7,438	140,907,823
BCSSTK23	3,134	45,178	420,311	49,018	1,522	119,155,247
BCSSTK24	3,562	159,910	278,922	22,331	414	32,429,194
BCSSTK25	15,439	252,241	1,416,568	205,513	7,288	283,732,315
BCSSTK29	13,992	619,488	1,694,796	174,770	3,231	393,045,158
BCSSTK30	28,924	2,043,492	3,843,435	229,670	3,689	928,323,809
BCSSTK31	35,588	1,181,416	5,308,247	330,896	8,304	2,550,954,465
BCSSTK32	44,609	2,014,701	5,246,353	374,507	6,927	1,108,686,016
BCSSTK33	8,738	591,904	2,546,802	124,532	1,201	1,203,491,786
NASA1824	1,824	39,208	73,699	12,587	527	5,160,949
NASA2910	2,910	174,296	204,403	25,170	599	21,068,943
NASA4704	4,704	104,756	281,472	35,339	1,245	35,003,786
NASASRB	54,870	2,677,324	11,904,998	592,254	8,027	4,672,895,526

Table 4.2: Characteristics of test problems.

Legend:

n : number of equations,

$|A|$: number of nonzeros in A ,

$|L|$: number of nonzeros in L , including the diagonal,

$\mu(L)$: number of row subscripts required to represent the supernodal structure of L ,

N : number of fundamental supernodes in L ,

flops: number of floating-point operations required to compute L .

Throughout this section, we use `colfct` to refer to the column-based approach to Cholesky factorization used in Figures 2.2 and 2.5. Likewise, we use `supfct` to refer to supernode-based approach used in Figures 3.3 and 3.4. For each approach there are two distinct but similar routines: a serial routine and a parallel routine. Thus, there are four routines, each implementing one of the algorithms found in Figures 2.2, 2.5, 3.3, and 3.4. It is worth noting that the serial `colfct` routine is a version of SPARSPAK's `gsfct` routine that has been slightly modified for fair comparison with the other routines, which were written from scratch.

4.2. Numerical results on an IBM RS/6000

The primary purpose of Table 4.3 is to show the impact on performance of various levels of loop unrolling in the `supfct` routines. The kernel subroutine called by both the serial and parallel `supfct` routines is capable of unrolling the outer loop of a column-oriented matrix-vector multiply, as in [9]. (The loop unrolling performs multiple `saxpy`'s in a single loop.) Such loop unrolling cannot be introduced into the `colfct` routines because they are not cognizant of the supernode structure on which the technique depends. Loop unrolling levels $\ell = 1, 2, 4,$ and 8 have been tried on several machines.

problem	colfct	supfct with loop unrolling			
		$\ell=1$	$\ell=2$	$\ell=4$	$\ell=8$
BCSSTK13	9.86	7.39	5.98	5.45	5.23
BCSSTK14	1.74	1.33	1.07	1.02	1.00
BCSSTK18	24.02	18.59	15.17	13.93	13.60
BCSSTK23	19.91	14.88	11.94	10.74	10.52
BCSSTK24	5.59	4.10	3.27	3.02	2.95
NASA1824	.94	.74	.62	.59	.58
NASA2910	3.68	2.83	2.32	2.13	2.13
NASA4704	6.09	4.53	3.71	3.42	3.37

Table 4.3: Factorization times in seconds for various levels of loop-unrolling on an IBM RS/6000 (model 320).

Table 4.3 records the results of these tests on an IBM RS/6000 workstation (model 320). Our double precision Fortran code was compiled using the IBM Fortran compiler `xlf` with optimization turned on (i.e., `xlf -O`). The results for some of the smaller problems in our test set are reported in Table 4.3.

The first thing to note is that `supfct` with no loop unrolling ($\ell=1$) is significantly faster than `colfct`. We believe the improvement is due to better use of the cache by `supfct`, which is due, in turn, to the reduction in indirect addressing and increased locality of the data references obtained via supernodes and careful attention to certain implementation details. On other machines we have tried, `supfct` with no loop unrolling generally runs no faster than `colfct`, and quite often runs slightly slower.

The improvements due to loop unrolling shown in Figure 4.3 are fairly typical of

what we have observed on other machines, too. While the benefits of loop unrolling levels higher than $\ell=4$ are minimal on the IBM RS/6000, the point of diminishing returns is usually higher on other machines. Experience has shown $\ell=8$ to be a good overall choice for the machines we have worked with. In all subsequent experiments, sequential and parallel `supfct` use loop unrolling to level $\ell=8$.

4.3. Numerical results on a Sequent Balance 8000

Next, we compare the performance of parallel `colfct` with that of parallel `supfct` on a Sequent Balance 8000, a shared-memory multiprocessor with 12 processors and 16 Mbytes of memory. The parallel routines used Sequent Fortran compiler directives to access the parallel capabilities of the machine and to perform the necessary synchronization operations. The Sequent's Fortran preprocessor transformed these compiler directives into appropriate Fortran code, which, in turn, issued the required system subroutine calls. The Fortran source code was compiled using the Fortran compiler `fortran` with the optimization and preprocessing options turned on (i.e., `fortran -04 -mp`).

Table 4.4 contains factorization times and speed-up ratios (enclosed in parentheses) for runs on some of the smaller problems in our test set. Since we are interested primarily in comparing the ability of `colfct` and `supfct` to exploit multiple processors, each speed-up ratio is formed by dividing the time required for a parallel run into the time required for a serial run of the same method. Note that the serial time is quite distinct from the time required by the parallel algorithm on a single processor, which is always greater.

problem	method	serial	parallel				
			p=1	p=2	p=4	p=7	p=10
BCSSTK13	<code>colfct</code>	1147.2	1299.4 (0.88)	652.2 (1.8)	333.7 (3.4)	195.6 (5.9)	140.7 (8.2)
	<code>supfct</code>	874.4	878.3 (1.00)	440.3 (2.0)	225.9 (3.9)	132.0 (6.6)	93.7 (9.3)
BCSSTK14	<code>colfct</code>	195.2	225.6 (0.87)	114.2 (1.7)	58.7 (3.3)	34.8 (5.6)	25.5 (7.7)
	<code>supfct</code>	155.4	157.5 (0.99)	79.6 (2.0)	40.4 (3.8)	23.4 (6.6)	16.8 (9.3)
BCSSTK18	<code>colfct</code>	2790.9	3152.3 (0.89)	1589.7 (1.8)	810.0 (3.4)	477.3 (5.8)	346.3 (8.1)
	<code>supfct</code>	2144.4	2179.3 (0.98)	1097.9 (2.0)	556.7 (3.9)	326.2 (6.6)	234.8 (9.1)
BCSSTK23	<code>colfct</code>	2328.8	2627.9 (0.89)	1322.5 (1.8)	670.2 (3.5)	387.0 (6.0)	275.1 (8.5)
	<code>supfct</code>	1755.1	1776.8 (0.99)	893.1 (2.0)	447.5 (3.9)	259.1 (6.8)	182.6 (9.6)
BCSSTK24	<code>colfct</code>	640.3	733.8 (0.87)	371.6 (1.7)	188.8 (3.4)	110.7 (5.8)	80.2 (8.0)
	<code>supfct</code>	493.9	500.4 (0.99)	250.6 (2.0)	126.8 (3.9)	73.1 (6.8)	52.1 (9.5)
NASA1824	<code>colfct</code>	104.9	122.4 (0.86)	62.0 (1.7)	32.0 (3.3)	19.4 (5.4)	14.7 (7.1)
	<code>supfct</code>	84.8	86.8 (0.98)	43.9 (1.9)	22.4 (3.8)	13.2 (6.4)	9.6 (8.8)
NASA2910	<code>colfct</code>	417.0	482.7 (0.86)	242.5 (1.7)	124.1 (3.4)	73.2 (5.7)	53.3 (7.8)
	<code>supfct</code>	330.0	337.0 (0.98)	167.6 (2.0)	85.5 (3.9)	49.5 (6.7)	35.2 (9.4)
NASA4704	<code>colfct</code>	691.9	791.8 (0.87)	401.3 (1.7)	204.2 (3.4)	119.8 (5.8)	86.6 (8.0)
	<code>supfct</code>	539.4	542.5 (0.99)	274.6 (2.0)	138.7 (3.9)	80.2 (6.7)	57.0 (9.5)

Table 4.4: Factorization times in seconds (and speed-ups) on a Sequent Balance 8000.

Comparing the factorization times for the two methods in the last column ($p=10$) clearly indicates the superiority of `supfct` over `colfct`. Indeed the differences in their performance on these problems are large and remarkably consistent, ranging from a

low of 47.5%³ to a high of 57.8%. Two observations largely account for the superior performance of `supfct`. First, the loop unrolling discussed in the previous section is quite valuable on the Sequent also. The effects of loop unrolling are apparent in the serial runs, and they are quite consistent among the problems, with improvements in performance ranging from a low of 23.7% to a high of 32.7%. The benefits of loop unrolling are largely preserved in the parallel implementation of `supfct`.

Second, `supfct`'s speed-up ratios are consistently better than `colfct`'s; for `colfct` they range from 7.1 to 8.5, and for `supfct` they range from 8.8 to 9.6. The "speed-up" ratios for parallel runs on a single processor ($p=1$) suggest that one of the primary reasons for `colfct`'s inferior speed-up ratios is the high synchronization overhead incurred by the method. Since there is no contention for access to the critical sections of the code when the parallel codes are run on a single processor, it is likely that the relative difference in synchronization overhead costs is significantly greater on 10 processors. The speed-ups for `colfct` on 10 processors are nonetheless quite respectable (7.1-8.5).

4.4. Numerical results on a Cray Y-MP

Finally, we compare parallel `colfct` and parallel `supfct` on a Cray Y-MP, a powerful vector supercomputer with 8 processors and 128 Mwords of memory. The code run on this machine was the same code run on the Sequent, with a few minor changes required to take care of machine-dependent constructs for exploiting parallelism. Again, the loop unrolling level used by `supfct` was $\ell=8$, and Fortran compiler directives were used to exercise the machine's parallel capabilities and to perform the necessary synchronization operations. The code was compiled using the Fortran compiler `cf77` with optimization (the default) and preprocessing options on. (i.e., `cf77 -Zu`).

The top half of Table 4.5 reports factorization times and speed-up ratios (enclosed in parentheses) for both methods applied to some small problems in our test set. The bottom half of the table records performance data for `supfct` on the remaining problems in our test set.

Not surprisingly, `supfct` performs much better than `colfct` on this machine. Loop unrolling is more effective on the Cray Y-MP than it is on the Sequent. Comparing the serial runs for the two methods, we find differences in performance ranging from a low of 53% to a high of 132%, due to loop unrolling and reduced indirect indexing in `supfct`. Similar results have been reported previously in [5]. We also find that `supfct` parallelizes much better than `colfct`. For example, on eight processors ($p=8$) the speed-up ratios for `supfct` range from a low of 6.0 to a high of 6.9, which is quite good, especially on such small problems. The speed-up ratios for `colfct`, however, are very poor, ranging from a low of 2.0 to a high of 3.7. As was the case on the Sequent, the "speed-ups" obtained on a single processor indicate that the high synchronization costs

³The base for each percentage is the smaller of the two times. This applies to percentages presented later in this section as well.

problem	method	serial	parallel				
			p=1	p=2	p=4	p=6	p=8
BCSSTK13	colfct	.929	1.347 (.69)	.685 (1.4)	.379 (2.5)	.314 (3.0)	.301 (3.1)
	supfct	.439	.493 (.89)	.249 (1.8)	.128 (3.4)	.089 (4.9)	.069 (6.4)
BCSSTK14	colfct	.238	.391 (.61)	.203 (1.2)	.126 (1.9)	.121 (2.0)	.121 (2.0)
	supfct	.156	.185 (.84)	.093 (1.7)	.048 (3.2)	.033 (4.7)	.026 (6.0)
BCSSTK15	colfct	2.485	3.471 (.72)	1.770 (1.4)	.962 (2.6)	.768 (3.2)	.728 (3.4)
	supfct	1.071	1.172 (.91)	.594 (1.8)	.299 (3.6)	.204 (5.2)	.157 (6.8)
BCSSTK16	colfct	2.444	3.549 (.69)	1.814 (1.3)	.995 (2.5)	.834 (2.9)	.812 (3.0)
	supfct	1.067	1.178 (.91)	.590 (1.8)	.299 (3.6)	.202 (5.3)	.154 (6.9)
BCSSTK17	colfct	2.712	4.121 (.66)	2.142 (1.3)	1.223 (2.2)	1.104 (2.5)	1.094 (2.5)
	supfct	1.373	1.540 (.89)	.773 (1.8)	.392 (3.5)	.267 (5.1)	.206 (6.7)
BCSSTK18	colfct	2.288	3.284 (.70)	1.675 (1.4)	.928 (2.5)	.767 (3.0)	.729 (3.1)
	supfct	1.314	1.481 (.89)	.741 (1.8)	.382 (3.4)	.265 (5.0)	.213 (6.2)
BCSSTK23	colfct	1.755	2.408 (.73)	1.219 (1.4)	.652 (2.7)	.514 (3.4)	.473 (3.7)
	supfct	.798	.879 (.91)	.441 (1.8)	.224 (3.6)	.153 (5.2)	.125 (6.4)
BCSSTK24	colfct	.674	1.071 (.63)	.551 (1.2)	.329 (2.0)	.302 (2.2)	.301 (2.2)
	supfct	.338	.381 (.89)	.190 (1.8)	.096 (3.5)	.065 (5.2)	.050 (6.8)
BCSSTK25	supfct	2.580	2.872 (.90)	1.433 (1.8)	.731 (3.5)	.504 (5.1)	.394 (6.5)
BCSSTK29	supfct	2.939	3.195 (.92)	1.602 (1.8)	.810 (3.6)	.547 (5.4)	.421 (7.0)
BCSSTK30	supfct	5.816	6.301 (.92)	3.154 (1.8)	1.590 (3.7)	1.073 (5.4)	.815 (7.1)
BCSSTK31	supfct	12.607	13.299 (.95)	6.653 (1.9)	3.330 (3.8)	2.249 (5.6)	1.706 (7.4)
BCSSTK32	supfct	7.773	8.491 (.92)	4.249 (1.8)	2.134 (3.6)	1.442 (5.4)	1.100 (7.1)
BCSSTK33	supfct	5.831	6.146 (.95)	3.074 (1.9)	1.539 (3.8)	1.040 (5.6)	.788 (7.4)
NASA1824	supfct	.114	.137 (.83)	.069 (1.7)	.036 (3.2)	.025 (4.6)	.019 (6.0)
NASA2910	supfct	.287	.331 (.87)	.166 (1.7)	.084 (3.4)	.058 (4.9)	.045 (6.4)
NASA4704	supfct	.429	.483 (.89)	.243 (1.8)	.124 (3.5)	.085 (5.0)	.065 (6.6)
NASASRB	supfct	23.563	24.850 (.95)	12.404 (1.9)	6.202 (3.8)	4.179 (5.6)	3.164 (7.4)

Table 4.5: Factorization times in seconds (and speed-ups) on a Cray Y-MP.

incurred by colfct seriously degrade its parallel performance. Indeed, on this machine the overhead appears to be considerably higher than it was on the Sequent. This high overhead, combined with the fast floating-point computational rates on this machine, probably account for most of the degradation in parallel performance of colfct.

The performance of supfct on the large problems in the bottom half of the table is consistently good. Ignoring the three smallest problems in this portion of the table (NASA1824, NASA2910, and NASA4704), speed-up ratios range from a low of 6.5 to a high of 7.4. On six out of seven of these problems the speed-up ratio is 7.0 or greater, with the 6.5 speed-up ratio reserved for the problem requiring the least work, namely BCSSTK25.

Table 4.6 compares the performance of supfct with the parallel supernodal factorization algorithm used in [28], which we will designate as supfct-SVY. The performance figures are expressed in Mflops⁴, as is commonly done for vector supercomputers such as the Cray Y-MP. We report the performance of both codes on those problems in our test set for which results for supfct-SVY were available to us. The performance data for supfct-SVY were obtained from an unpublished manuscript [29].

Consider the results obtained on 8 processors ($p=8$). On six of the twelve prob-

⁴Mflops (megaflops) are millions of floating-point operations per second. Gflops (gigaflops) are billions of floating-point operations per second.

problem	method	serial	parallel	
			p=4	p=8
BCSSTK15	supfct	154.1	551.9	1051.2
	supfct-SVY	197.8	301.1	320.8
BCSSTK16	supfct	139.7	498.6	968.2
	supfct-SVY	190.8	287.5	297.4
BCSSTK23	supfct	149.3	531.9	953.2
	supfct-SVY	191.6	293.3	315.1
BCSSTK24	supfct	95.9	337.8	648.5
	supfct-SVY	139.4	168.2	168.7
BCSSTK30	supfct	159.6	583.8	1139.0
	supfct-SVY	212.2	350.0	375.0
BCSSTK31	supfct	202.3	766.0	1495.3
	supfct-SVY	251.4	566.3	689.2
BCSSTK32	supfct	142.6	519.5	1007.9
	supfct-SVY	193.5	291.4	307.0
BCSSTK33	supfct	206.4	782.0	1527.3
	supfct-SVY	258.4	593.1	717.2
NASA1824	supfct	45.3	143.3	271.5
	supfct-SVY	64.3	69.8	69.8
NASA2920	supfct	73.4	250.8	468.1
	supfct-SVY	97.5	121.6	121.4
NASA4704	supfct	81.6	282.3	538.4
	supfct-SVY	117.0	143.5	143.5
NASASRB	supfct	198.3	753.4	1476.9
	supfct-SVY	250.6	531.6	625.2

Table 4.6: Factorization computational rates (Mflops) on a Cray Y-MP.

lems, `supfct` performs the factorization at over a Gflop, with highs of 1.48 Gflops on NASASRB, 1.50 Gflops on BCSSTK31, and 1.53 Gflops on BCSSTK33. For two others problems, the computational rate is nearly a Gflop: .97 Gflop on BCSSTK16 and .95 Gflop on BCSSTK23. Thus, for 8 out of 12 of the problems, `supfct` computed the factorization at nearly a Gflop or more. Table 4.2 indicates that the remaining four problems (NASA1824, NASA2910, NASA4704, and BCSSTK24) are quite small. Moreover, in Table 4.5 we see that serial `supfct` requires less than half a second to factor any of these matrices.

The performance of `supfct-SVY` is much poorer due to the problems with this approach mentioned earlier in Section 3.3. The code runs at less than a Gflop on every problem, despite having significantly higher serial efficiency due to assembly language programming of the compute-intensive kernel routines and other machine-specific optimizations. Our parallel implementation of `supfct` is a Fortran 77 code, with *no* machine-specific optimizations.

5. Concluding remarks

We have implemented a new parallel sparse Cholesky factorization algorithm for shared-memory multiprocessors. This new left-looking algorithm uses techniques from [5] and [15]: it uses supernodes to reduce indirect addressing and memory traffic [5], and it decomposes the computation into column tasks $T_{col}(j)$ and schedules these tasks dynamically on the available processors [15]. Incorporation of supernodes into the algorithm in [15] reduces the synchronization overhead required to manage the row structure sets S_q from $O(|L|)$ to $O(\mu(L))$, where $\mu(L)$ is the number of row subscripts required to represent the supernodal structure of L . In practice, $\mu(L)$ is often much smaller than $|L|$; consequently, contention for the critical sections is likely to be much higher in `colfct` than in `supfct`. Since the algorithms use a single lock variable for the critical sections, the sections are executed serially. Thus, the serial component of `supfct` is in practice much smaller than that of `colfct`. Our tests indicate that this is the single most important factor contributing to the new algorithm's superior parallel performance.

Right-looking sparse Cholesky algorithms (i.e., multifrontal algorithms) for shared-memory multiprocessors have appeared in [6,10,29]. These algorithms exploit supernodes in much the same way that the new parallel left-looking algorithm does. Parallelizing the multifrontal algorithm however is considerably more complicated than parallelizing the simpler left-looking algorithms. For instance, parallel multifrontal Cholesky for shared-memory machines can no longer use a simple and efficient stack to manage the "update" matrices required by the method. Methods for dealing with this fragmented component of work storage are necessarily more complicated and storage inefficient [10,29]. On the other hand, `supfct` requires only a modest amount of work storage, which can be determined *before* the numerical factorization begins. Break-

ing multifrontal Cholesky factorization into tasks and scheduling these tasks on the available processors is also more complicated than it is for the left-looking algorithms. The additional complexity has led to parameterized implementations, where the performance of the code is quite sensitive to parameter selection (see [10] and especially [29]).

There are however significant advantages enjoyed by the multifrontal method; e.g., it is a superior out-of-core method and is better able to improve performance by loading and reusing data in cache. It appears to us that the multifrontal method very likely will always be the method of choice for out-of-core sparse Cholesky factorization. However, we think that a block-to-block left-looking algorithm may be quite competitive with the multifrontal method at exploiting cache to improve performance. Such an algorithm would be built around a $cmod(\mathbf{J}, \mathbf{K})$ operation that updates the appropriate subset of columns from supernode \mathbf{J} with all the columns of the updating supernode \mathbf{K} . With the rising importance of cache memory on recent supercomputers and workstations, exploring this approach on current serial and parallel machines is a promising area for future work.

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