RECENT ADVANCES IN SPARSE DIRECT SOLVERS

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ABSTRACT

Direct methods for the solution of sparse systems of linear equations of the form $A x = b$ are used in a wide range of numerical simulation applications. Such methods are based on the decomposition of the matrix into a product of triangular factors (e.g., $A = L U$), followed by triangular solves. They are known for their numerical accuracy and robustness but are also characterized by a high memory consumption and a large amount of computations. Here we survey some research directions that are being investigated by the sparse direct solver community to alleviate these issues: memory-aware scheduling techniques, low-rank approximations, and distributed/shared memory hybrid programming.

INTRODUCTION

Context and objectives

Large sparse linear systems appear in numerous scientific applications. For example, the structural analysis of nuclear plants after the Fukushima events (March 2011) requires to solve large linear systems with typically tens or hundreds of millions of unknowns arising from numerical simulation at very large scale (an entire plant structure), and in very high resolution. Direct methods for the solution of sparse linear systems are known for their numerical robustness but usually have large computational requirements (as opposed to iterative methods). In particular, their large memory footprint often prevents their use, and achieving large scale parallelism (thousands of cores or more) is often difficult. Here we survey some recent techniques that aim at improving the scalability of direct methods. In the first section, we describe a memory-aware scheduling strategy that aims at improving the memory scalability of a variant of sparse Gaussian elimination called the multifrontal method. In the second section, we introduce different low-rank approximation techniques that aim at decreasing the space and time complexity of direct solvers by detecting data sparsity and introducing controlled approximations. Finally, in the last section, we show how to combine distributed-memory and shared-memory parallelism in order to exploit recent architectures.

Background

We briefly introduce some ingredients of sparse direct methods. We only provide simplistic descriptions of what is necessary to understand the following sections and we refer the reader to the bibliography for
more detail. We are to compute a factorization of a given matrix \( A \), \( A = LU \) if the matrix is unsymmetric, or \( A = LDL^T \) if the matrix is symmetric. Without loss of generality, in the rest of the paper we will assume that \( A \) is non-reducible. For a matrix \( A \) with an unsymmetric pattern (nonzero structure), we assume that the factorization takes place using the structure of \( A + A^T \), where the summation is structural. The fundamental structure on which we will rely in the following is called the elimination tree. A few equivalent definitions are possible (we recommend the survey by Liu (1990)); we use the following:

**Definition 1** Assume \( A = LU \) where \( A \) is a sparse, structurally symmetric, \( N \times N \) matrix. Then, the elimination tree of \( A \) is a tree of \( N \) nodes, with the \( i \)-th node corresponding to the \( i \)-th column of \( L \) and with the parent relations defined by:

\[
\text{parent}(j) = \min\{i : i > j \text{ and } \ell_{ij} \neq 0\}, \text{ for } j = 1, \ldots, N - 1
\]

In practice, nodes are amalgamated: nodes that represent columns and rows of the factors with similar structures are grouped together in a single node (often referred to as a supernode). The elimination tree is a workflow and dataflow graph for many direct methods. At this point, two main variants of sparse direct methods can be distinguished. In the multifrontal method (Duff & Reid (1983)), each node in the elimination tree is associated with a square dense matrix (referred to as a frontal matrix or front) with the following \( 2 \times 2 \) block structure:

\[
\begin{bmatrix}
F_{11} & F_{12} \\
F_{21} & F_{22}
\end{bmatrix}
\]

We refer to the variables associated with \( F_{11} \) and \( F_{22} \) as the fully-summed and non fully-summed variables, respectively. Factoring the matrix using the multifrontal method consists in a bottom-up traversal of the tree, following a topological order (a node is processed before its parent). Processing a node consists in:

- forming (or assembling) the frontal matrix by summing the rows and columns of \( A \) corresponding to the variables in the \((1, 1)\) block with temporary data that has been produced by the child nodes;

- eliminating the pivots in the \((1, 1)\) block \( F_{11} \); this is done through a partial factorization of the frontal matrix which produces the corresponding rows and columns of the factors stored in \( F_{11} \), \( F_{21} \) and \( F_{12} \).

At this step, the so-called Schur complement or contribution block is computed as \( F_{22} \leftarrow F_{22} - F_{21} \cdot F_{11}^{-1} \cdot F_{12} \) and stored in a temporary memory; it will be used to form the front associated with the parent node. Therefore, when a node is activated, it “consumes” the contribution blocks of its children.

In the multifrontal method, the active memory (at a given step in the factorization) consists of the front being processed and a set of contribution blocks that are temporarily stored and will be consumed at a later step.

Supernodal methods can also rely on the elimination tree, however a node in the tree is not associated with a dense matrix but simply with some columns and rows of the factors. There is no contribution block. Instead, the updates corresponding to a node are done by communicating pieces of factors to some of its ancestors or from some of its descendants (right-looking or left-looking approach, respectively). The multifrontal method usually exhibits good flop rates as it mainly consists of large BLAS 3 operations (for computing contribution blocks), and it has a simple communication pattern (from children nodes to their parent), but the abovementioned active memory can be quite large. Supernodal methods do not have this extra memory usage but they have more complicated communication patterns. We illustrate the difference between these two classes of methods in Figure 1.

Direct methods relying on the elimination tree lend themselves very naturally to parallelism since multiple processes can be employed to treat one, large enough, frontal matrix or supernode or to process concurrently frontal matrices or supernodes belonging to separate subtrees. These two sources of parallelism are commonly referred to as node and tree parallelism, respectively, and their correct exploitation is the key to achieving high performance on parallel supercomputers.
In this section we focus on the multifrontal method, although similar ideas could be applied to other variants. In the multifrontal method, the active memory (mentioned in the previous section) can be quite large and is often difficult to control in a parallel environment. We are interested in maximizing the memory efficiency \( e_{max}(p) = \frac{S_{seq}}{pS_{max}} \) where \( S_{seq} \) is the peak of active memory for a sequential execution, and \( S_{max} \) is the maximum peak of active memory for an execution on \( p \) processors. \( e_{max}(p) \) should be close to 1, which means that the peak of a given processor should be close to \( \frac{S_{seq}}{p} \). \( S_{max} \) depends on the node-to-process mapping followed during the factorization. Commonly-used mappings often rely on the proportional mapping; in this strategy, the processors assigned to a node of the elimination tree are distributed to its children proportionally to their weights (operation count or sequential peak of memory). For example, the tree shown in Figure 2(a) is mapped on 64 processors. Its child node “a” is mapped on \( \frac{70}{70+50+50} \cdot 64 = 26 \) processors. The three subtrees rooted at a, b, and c, are mapped on different processors and are processed in parallel. In this example, the maximum peak of active memory is at least \( \frac{70}{26} \) GB (which is reached in the ideal case where the efficiency in the subtree rooted at a is 1), which yields \( e_{max} \leq \frac{80}{64+1.6} = 0.40 \), which is far from 1.

We proposed a memory-aware mapping technique that aims at enforcing a user-given memory constraint \( M_0 \) which is the maximum allowed peak of active memory of a processor. The basic idea is to follow a proportional mapping and to reject the steps that lead to violate the memory constraint. Take Figure 2(a) and assume \( M_0 = 1.6 \) GB. As explained above, the memory peak for the subtree rooted at a is at least \( \frac{70}{26} \) GB, which is greater than \( M_0 \). Our memory-aware strategy is illustrated in Figure 2(b): instead of mapping the whole set of siblings using proportional mapping, we detect groups of siblings (here \{a\} and \{b, c\}) that can be mapped using proportional mapping without violating \( M_0 \). We enforce scheduling constraints that force these groups to start one after another (here the subtrees rooted at nodes b and c cannot start before the subtree rooted at a is completed). Here the efficiency becomes at least \( e_{max} = \frac{80}{64+1.6} = 0.8 \).
In Table 1, we provide results for 3 benchmark matrices. pancake2_3 is from a 3D problem in electromagnetism, HV15R is from computational fluid dynamics, and meca_raff6 is a thermo-mechanical coupling problem for EDF nuclear power plants. We compare the default scheduling strategy in MUMPS with our memory-aware scheduling. The results show that the active memory can be significantly reduced, at the price of a moderate increase in run time. The missing data for HV15R means that the factorization with the default scheduling ran out of memory (we show an estimated memory consumption).

We refer the reader to (Rouet (2012)) for a more detailed description and more results.

**LOW-RANK APPROXIMATIONS**

Matrices coming from elliptic Partial Differential Equations (PDEs) have been shown to have a low-rank property: well defined off-diagonal blocks of their Schur complements can be approximated by low-rank products. Given a suitable ordering of the matrix, such approximations can be computed using truncated SVD or rank-revealing QR factorizations, given a low-rank truncation parameter $\varepsilon$. The resulting representation offers a substantial reduction of the memory requirements and many of the basic dense algebra operations can be performed with less operations. Even when $\varepsilon$ is set to maintain full accuracy (e.g., $\varepsilon = 10^{-14}$ for double-precision arithmetic) dramatic savings can be obtained both in memory consumption and in complexity of a sparse factorization; alternatively, with bigger threshold values, approximated factorizations can be computed for example to precondition iterative solvers at a cost which is only a fraction of the cost of a standard, full-rank factorization. Here we present two low-rank representations that can be embedded within sparse direct solvers: Block Low-Rank techniques and Hierarchically Semi-Separable representations.

**Block Low-Rank techniques**

A flexible, efficient technique can be used to represent fronts with low-rank subblocks based on a storage format called Block Low-Rank (BLR). Unlike other formats such as $H$-matrices and HSS matrices (see next section), the BLR approach is based on a flat, non-hierarchical blocking of the matrix which is
defined by conveniently clustering the associated unknowns. A BLR representation of a front $F$ is shown in equation from Figure 3(a) where we assume that $p$ subblocks have been defined over the fully-summed variables part of $F$, and $q$ subblocks over the non fully-summed variables part. Subblocks $\tilde{B}_{i,j}$ of size $m_{i,j} \times n_{i,j}$ and numerical rank $k_{i,j}(\varepsilon)$ are approximated by a low-rank product $U_{i,j}V_{i,j}^T$ at accuracy $\varepsilon$. Note that diagonal subblocks $B_{i,i}$ are never approximated because they are assumed to be full-rank.

\[
\tilde{F} = \begin{bmatrix}
B_{1,1} & \cdots & \tilde{B}_{1,p} & \cdots & \tilde{B}_{1,p+q} \\
\vdots & \ddots & \vdots & \ddots & \vdots \\
\tilde{B}_{p,1} & \cdots & B_{p,p} & \cdots & \tilde{B}_{p,p+q} \\
\vdots & \cdots & \vdots & \ddots & \vdots \\
\tilde{B}_{p+q,1} & \cdots & B_{p+q,p} & \cdots & \tilde{B}_{p+q,p+q}
\end{bmatrix}
\]

(a) BLR matrix definition.

(b) Structure of a BLR matrix. The lighter a block is, the smaller is its rank.

Figure 3: BLR matrix definition and structure example.

In order to achieve a satisfactory reduction in both complexity and memory footprint, subblocks have to be chosen to be as low-rank as possible (e.g., with exponentially decaying singular values) which can be achieved by clustering the unknowns in such a way that an admissibility condition (see Bebendorf (2008)) is satisfied. This condition states that a subblock $\tilde{B}_{i,j}$, interconnecting variables of $i$ with variables of $j$, will have a low rank if variables of $i$ and variables of $j$ are far away in the domain, intuitively, because the associated variables are likely to have a weak interaction. In practice, the fully-summed variables of each front are groups by partitioning (with a tool such as METIS) the related adjacency graph whereas the non fully-summed variables are grouped using a partitioning induced by those of the fully-summed variables of ancestor nodes (see Amestoy et al. (2012)). Although compression rates may not be as good as those achieved with hierarchical formats such as those described in the next section\(^1\), BLR offers a good flexibility thanks to its simple, flat structure. In a parallel environment, this allows for an easier distribution and handling of the frontal matrices. Also, numerical pivoting can be more easily done within a BLR matrix without perturbing much the structure. Lastly, converting a matrix from the standard representation to BLR and vice versa, is much cheaper with respect to the case of hierarchical matrices. This allows to switch back and forth from one format to the other whenever needed at a reasonable cost; this is, for example, done to simplify the assembly operations that are extremely complicated to perform in any low-rank format. All these points make BLR easy to adapt to any multifrontal solver without a complete rethinking of the code.

As shown in Figure 4, the $O(N^2)$ complexity of a standard, full rank solution of a 3D problem (of $N$ unknowns) from an 11-point stencil is reduced to $O(N^{4/3})$ when using the BLR format. In Table 2, we report results on two matrices from real-life application: Geoazur128 is a 3D seismic wave propagation study and TH RAFF7 is a 3D thermal test-case in structural engineering for EDF nuclear power plants. The results show that the number of operations and the memory peak can be substantially reduced.

Detailed information about these techniques can be found in Amestoy et al. (2012).

Hierarchically Semi-Separable matrices

Hierarchically semi-separable (HSS) matrices (Vandebril et al. (2005)) are another type of low-rank representation that can be embedded within a sparse solver. The resulting HSS-sparse factorization can be used as a direct solver or preconditioner depending on the application’s accuracy requirement and the characteristics of the PDEs. If the randomized sampling compression technique is employed in compression, we

\(^1\)This is the object of ongoing research.
algorithms to solve the linear systems (Chandrasekaran et al. (2006)). We can apply the above HSS low-rank

Figure 4: Operation count of the BLR multifrontal factorization of a matrix coming from the Laplacian operator discretized with a 3D 11-point stencil, revealing a \(O(N^{4/3})\) complexity for \(\varepsilon = 10^{-14}\).

can show that for the 3D model problems, the HSS-sparse factorization costs \(O(N)\) flops for discretized matrices from certain PDEs and \(O(N^{4/3})\) for broader classes of PDEs (including non-selfadjoint and indefinite discretized PDEs; cf., Xia (2012)). This complexity is much lower than the \(O(N^2)\) cost of the traditional, exact sparse factorization method.

Informally, the HSS representation partitions the off-diagonal blocks of a dense matrix in a hierarchical fashion; these off-diagonal blocks are approximated by compact forms, such as truncated SVD. A key property of HSS is that the orthogonal bases are desired to be nested following the hierarchical partitioning. This leads to asymptotically fast construction and factorization algorithms. Figure 5 illustrates a block \(8 \times 8\) HSS representation of \(A\), for which the hierarchical structure and the generators \(U_i, V_i, R_i\), and \(B_i\) are succinctly depicted by the HSS tree on the right side. As a special example, its leading block \(4 \times 4\) part looks like the following, where \(t_7\) is the index set associated with node 7 of the HSS tree:

\[
A|_{t_7 \times t_7} \approx \begin{pmatrix}
D_1 & U_1 B_1 V_1^H \\
U_2 B_2 V_1^H & D_2 \\
U_4 R_4 & U_5 R_5 \\
U_4 B_4 V_2^H & W_2^H V_2^H
\end{pmatrix}
\begin{pmatrix}
U_1 R_1 \\
U_2 R_2 \\
B_6 (W_1^H V_1^H) \\
U_5 B_5 V_5^H
\end{pmatrix}\begin{pmatrix}
D_3 \\
D_4 \\
B_3 (W_4^H V_4^H) \\
U_4 B_4 V_5^H
\end{pmatrix}
\]

Figure 5: Pictorial illustrations of a block \(8 \times 8\) HSS form and the corresponding HSS tree \(T\).

With this HSS representation, we can use the ULV factorization and the accompanying solution algorithms to solve the linear systems (Chandrasekaran et al. (2006)). We can apply the above HSS low-rank
approximation algorithms to the dense submatrices in the sparse multifrontal and supernodal factorization algorithms. We report on some results using a multifrontal code specialized in the solution of the discretized Helmholtz equation on regular grids; matrices are complex, unsymmetric, highly-indefinite, and in single-precision arithmetic. We compare runs with the same code in a classical multifrontal mode, and in the HSS-sparse mode (the tolerance is $\varepsilon = 10^{-4}$). We solve for one right-hand side and apply iterative refinement in the solution. Using HSS techniques speeds up the solution process, and gains increase with the size of the problem, which is expected since, for this kind of equations, the complexity is $O(N^{4/3})$ instead of $O(N^2)$ for a full-rank approach. However the flop rates are smaller which is due to the fact that HSS kernels operate on small blocks, while regular kernels only perform large BLAS3 operations. The size of factors is significantly reduced but the gains on the maximum peak of memory are not as large because this particular code does not apply HSS techniques on contribution blocks (this is work in progress). The communication volume is also reduced when HSS kernels are used. Finally, using iterative refinement, the HSS-enabled code is almost as stable as the pure multifrontal code (the scaled residual is close to machine precision).

**Table 3: Solution of the discretized Helmholtz equation on a $k \times k \times k$ grid.**

<table>
<thead>
<tr>
<th></th>
<th>Processors</th>
<th>100</th>
<th>200</th>
<th>300</th>
<th>400</th>
</tr>
</thead>
<tbody>
<tr>
<td>MF</td>
<td>Total time (s)</td>
<td>89.0</td>
<td>1530.2</td>
<td>4218.2</td>
<td>6376.4</td>
</tr>
<tr>
<td></td>
<td>Gflops/s</td>
<td>600.6</td>
<td>2275.7</td>
<td>9505.6</td>
<td>35477.3</td>
</tr>
<tr>
<td></td>
<td>Factors size (GB)</td>
<td>16.6</td>
<td>280.0</td>
<td>1450.1</td>
<td>4636.1</td>
</tr>
<tr>
<td></td>
<td>Maximum peak per proc. (GB)</td>
<td>0.5</td>
<td>1.9</td>
<td>2.5</td>
<td>2.0</td>
</tr>
<tr>
<td></td>
<td>Communication volume (GB)</td>
<td>83.1</td>
<td>2724.7</td>
<td>26867.8</td>
<td>165299.3</td>
</tr>
<tr>
<td>HSS</td>
<td>Total time (s)</td>
<td>122.7</td>
<td>1069.5</td>
<td>2265.3</td>
<td>3859.3</td>
</tr>
<tr>
<td></td>
<td>Gflops/s</td>
<td>207.8</td>
<td>720.4</td>
<td>2576.6</td>
<td>6494.8</td>
</tr>
<tr>
<td></td>
<td>Factors size (GB)</td>
<td>10.7</td>
<td>112.9</td>
<td>434.3</td>
<td>845.3</td>
</tr>
<tr>
<td></td>
<td>Maximum peak per proc. (GB)</td>
<td>0.5</td>
<td>1.7</td>
<td>2.1</td>
<td>0.4</td>
</tr>
<tr>
<td></td>
<td>Communication volume (GB)</td>
<td>93.6</td>
<td>2241.2</td>
<td>18621.1</td>
<td>143300.0</td>
</tr>
<tr>
<td></td>
<td>$\max_{i,j} \frac{</td>
<td>Ax_i - bj</td>
<td>}{</td>
<td></td>
<td>A</td>
</tr>
</tbody>
</table>

**INTRODUCING SHARED-MEMORY PARALLELISM IN DISTRIBUTED-MEMORY SOLVERS**

Recent architectures often consist of increasingly complex shared-memory nodes. Using *distributed-memory programming* (task parallelism, e.g., with MPI) within such nodes is not the best way to exploit them since they tend to have smaller and smaller amounts of per-core memory and non-uniform memory accesses (NUMA). It is thus necessary to combine distributed-memory programming for inter-node computations with *shared-memory programming* (thread parallelism, e.g., with OpenMP) for intra-nodes computations. Here we describe recent work carried out in MUMPS and SuperLU_DIST on this topic.

**Improvement of shared-memory parallelism in distributed-memory multifrontal methods**

In order to exploit shared-memory parallelism at the thread level, the simplest approach consists in making many cores collaborate on the same node of the elimination tree, using multithreaded BLAS and OpenMP directives to parallelize the work outside the BLAS calls (fork-join model). This approach is generally efficient, especially on large 3D problems. In order to go further, it is interesting to also exploit tree parallelism in shared-memory environments. In distributed-memory parallelism, a commonly-used technique consists in defining a separating layer in the tree $\mathcal{L}_{ps}$ such that both tree and node parallelism are applied above $\mathcal{L}_{ps}$ and only tree parallelism is applied below it. We adapted this idea to shared-memory en-
environments with the A vår L 0 algorithm (L’Excellent and Sid-Lakhdar (2012)), that computes a layer $L_{th}$ such that only node parallelism is applied above $L_{th}$ but tree parallelism is now applied below it (see Figure 6).

Figure 6: Layers $L_{ps}$ and $L_{th}$ to exploit tree parallelism in distributed and shared memory environments.

In multi-core environments, NUMA (Non-Uniform Memory Access) architectures allow for a hierarchical organization of memory banks. On such architectures, it is critical for performance to take into account memory affinity and memory allocation policies. By default, the localalloc policy, which consists in mapping the memory pages on the local memory of the processor that first touches them, is applied. However, several other policies exist. Among them, the interleave policy consists in allocating the memory pages on all memory banks in a round-robin fashion, such that the allocated memory is spread over all the physical memory. Applying the localalloc policy on data structures below $L_{th}$, in order to achieve a better data locality and cache exploitation, and the interleave policy on the ones used above $L_{th}$, in order to improve the bandwidth, has a dramatic effect on performance, as summarized in Table 4 when only 1 MPI process is used. This table also shows that the use of OpenMP directives to parallelize the work outside BLAS calls provides some gains, compared to the sole use of multithreaded BLAS.

On matrices such as Serena (Table 4), the sole effect of the A vår L 0 algorithm is not so large (1081.42 seconds down to 893.64 seconds). However, the effect of memory interleaving without $L_{th}$ is even smaller (1081.42 seconds down to 1006.66 seconds). The combined use of A vår L 0 and memory interleaving brings a huge gain: 1081.42 seconds down to 530.63 seconds (increasing the speed-up from 7.3 to 14.7 on 24 cores). Hence, on large matrices, the main benefit of A vår L 0 is to make the interleaving become very efficient by applying it only above $L_{th}$. This also shows that, in the implementation, it is critical to separate the work arrays for local threads below $L_{th}$ and for the more global approach in the upper part of the tree, allowing the application of different memory policies below and above $L_{th}$ (localalloc and interleave, respectively).

Table 4: Factorization times in seconds and effects of the interleave memory allocation policy with node parallelism and with A vår L 0 on an AMD 24 cores (Istanbul) system using MUMPS with a single MPI process.

<table>
<thead>
<tr>
<th>Matrix</th>
<th>Serial reference</th>
<th>Node parallelism only</th>
<th>Node parallelism only</th>
<th>ALG L 0</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Threaded BLAS</td>
<td>Threaded BLAS + OpenMP</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>Interleaving off on</td>
<td>Interleaving off on</td>
<td></td>
</tr>
<tr>
<td>AUDI</td>
<td>1535.8</td>
<td>269.8 260.8</td>
<td>231.8 225.5</td>
<td>158.0 110.0</td>
</tr>
<tr>
<td>conv3D64</td>
<td>3001.4</td>
<td>518.5 563.1</td>
<td>497.5 496.9</td>
<td>439.0 303.6</td>
</tr>
<tr>
<td>Serena</td>
<td>7845.4</td>
<td>1147.6 1058.0</td>
<td>1081.4 1006.7</td>
<td>893.7 530.6</td>
</tr>
</tbody>
</table>

These results show important aspects that should be considered inside a shared-memory node. Tuning the specific shared-memory factorization kernels inside each computing node is another aspect that should be considered to further improve performance. Optimizing the number of threads inside each MPI process can also have a significant impact on performance.

We refer the reader to L’Excellent and Sid-Lakhdar (2012) for further reference.
Multithreaded Schur-complement update in SuperLU_DIST

The distributed-memory sparse direct solver SuperLU_DIST (Li & Demmel, 2003) was initially designed to use only MPI to exploit task parallelism. In SuperLU_DIST, the computational cost of numerical factorization is typically dominated by the trailing submatrix update, where each process updates several independent blocks of the trailing submatrix at each step. Recently, we incorporated light-weight OpenMP threads in each MPI process to update disjoint sets of the independent blocks in parallel. There are several options as to how to assign the independent blocks to the threads. We use a combination of 1D/2D assignment of blocks to threads. When the number of columns of the trailing submatrix is greater than the number of threads, a process can assign its local supernodal columns to the threads in a 1D block fashion; i.e., the \( t \)-th thread updates \((t - 1) \cdot h\)-th to \((t \cdot h - 1)\)-th columns, where \( h = \frac{n_c}{n_t} \), \( n_t \) is the number of threads, and \( n_c \) is the number supernodal columns assigned to this process (see Figure 7(a)). Since these columns are contiguous in memory, each thread can access the columns with a relatively small stride. However, with this layout, the number of usable threads is constrained by the number of columns. Therefore, when the number of columns is smaller than the number of threads, we divide the columns row-wise as well, then assign the blocks to threads in a 2D cyclic fashion; namely the \((i, j)\)-th block is assigned to \((b_r \cdot t_c + b_c)\)-th thread, where the threads are organized into a \( t_r \times t_c \) grid (i.e., \( n_t = t_r \cdot t_c \)), \( b_r = \text{mod}(i, t_r) \), and \( b_c = \text{mod}(j, t_c) \) (see Figure 7(b)). Since the blocks assigned to a thread are not contiguous in memory, accessing these blocks incurs some overhead. The advantage is that it offers more parallelism than the 1D layout.

This hybrid programming paradigm obtained significant reduction in memory usage while achieving the same level of parallel efficiency as the pure MPI paradigm. As a result, in comparison to the pure MPI paradigm which failed due to the per-core memory constraint, this hybrid paradigm could use more cores on each node and reduce the factorization time on the same number of nodes.

![Figure 7: Mapping of threads to supernodal blocks.](image)

We tested this OpenMP-enabled SuperLU_DIST solver on the Cray-XE6 machine at NERSC (hopper). Hopper consists of 6,384 NUMA nodes. Each node has two twelve-core AMD MagnyCours 2.1-GHz processors, giving 24 cores per node. Each MagnyCours has two dies of six-core each. The memory access times within a die, outside a die, and outside a MagnyCours are different. The total memory per node is 32GB and per core is 1.3GB. We used 16 nodes and varied the number of OpenMP threads associated with each MPI task. Figure 8 shows the running times for a matrix from simulations of accelerator cavities, using different MPI/OpenMP configurations. We see that the best time with the fixed node count of 16 was obtained by the hybrid paradigm. The missing data points mean that the code ran out of memory in that MPI/thread combination. This clearly shows that the hybrid paradigm was able to better utilize the resources available on the compute nodes. The figure also shows the total high watermark of the memory used by the solver, which includes the size of the \( L, U \) factors, the solver’s auxiliary memory and the system memory (e.g., MPI buffers). When the same number of cores is used, threading helps reduce memory usage significantly. The reduction is in the MPI system buffers. OpenMP itself does not generate much memory overhead; this can
be seen in the following MPI/thread configurations: $32 \times 1$, $32 \times 2$ and $32 \times 4$.


![Figure 8: Time and memory using 16 nodes with various combinations of MPI tasks and OpenMP threads.](image)

**REFERENCES**


