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A Supernodal Approach to Incomplete LU Factorization with Partial Pivoting

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We present a new supernode-based incomplete LU factorization method to construct a preconditioner for solving sparse linear systems with iterative methods. The new algorithm is primarily based on the ILUTP approach by Saad, and we incorporate a number of techniques to improve the robustness and performance of the traditional ILUTP method. These include new dropping strategies that accommodate the use of supernodal structures in the factored matrix and an area-based fill control heuristic for the secondary dropping strategy. We present numerical experiments to demonstrate that our new method is competitive with the other ILU approaches and is well suited for modern architectures with memory hierarchy.

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1. INTRODUCTION

As problem sizes increase with high-fidelity simulations demanding fine details on large, three-dimensional geometries, iterative methods based on preconditioned Krylov subspace techniques are attractive and cheaper alternatives to direct methods. A critical component of the iterative solution techniques is the construction of effective preconditioners. Physics-based preconditioners are quite effective for structured problems, such as those arising from discretized partial differential equations. On the other hand, methods based on incomplete LU decomposition are still regarded as

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generally robust "black-box" preconditioners for unstructured systems arising from a wide range of application areas. A variety of ILU techniques have been studied extensively in the past, including distinct dropping strategies, such as the level-of-fill structure-based approach (ILU(k)) [Saad 1996], the numerical threshold-based approach [Saad 1994], and, more recently the numerical inverse-based multilevel approach [Bollhöfer and Saad 2006; Bollhöfer et al. 2006]. The ILU(k) approach assigns a fill level to each nonzero entry, which characterizes the length of the shortest fill path leading to this fill-in [Hysom and Pothen 2001]. The entries with level of fill larger than k are dropped. Intuitively, this leads to good approximate factorization only if the fill-ins become smaller and smaller as the sequence of updates proceeds. The implementation of ILU(k) can involve a separate symbolic factorization stage to determine all the entries to be dropped. The threshold-based approach takes into account the numerical size of the entries in the factors, and drops those entries that are smaller than a numerical tolerance. A problem with threshold-based approaches is the difficulty of choosing an appropriate drop tolerance. This method tends to produce better approximations and is applicable for a wider range of problems, but its implementation is much more complex because the fill-in pattern must be determined dynamically. One of the most sophisticated threshold-based methods is ILUTP proposed by Saad [1994, 1996], which combines a dual dropping strategy with numerical pivoting ("T" stands for threshold, and "P" stands for pivoting). The dual dropping rule in $ILUTP(p, \tau)$ first removes the entries that are smaller than τ from the current factored row or column. Among the remaining entries, at most p largest entries are kept in order to control memory growth. Therefore, the dual strategy is somewhat in between the structural and numerical approaches.

Our method can be considered to be a variant of the ILUTP approach, and we have modified our high-performance direct solver SuperLU [Demmel et al. 1999a, 1999b] to perform incomplete factorization. A key component in SuperLU is the use of supernodes, which gives several performance advantages over a nonsupernodal (e.g., columnwise) algorithm. First, supernodes enable the use of higher-level BLAS kernels which improves data reuse in the numerical phase. Second, symbolic factorization traverses the supernodal-directed graph to determine the nonzero structures of L and U. Since the supernodal graph can be much smaller than the nodal (column) graph, the speed of this phase can be significantly improved. Last, the amount of indirect addressing is reduced while performing the scatter/gather operations for compressed matrix representation. Although the average size of the supernodes in an incomplete factor is expected to be smaller than in a complete factor because of dropping, we have attempted to retain supernodes as much as possible. We have adapted the dropping rules to incorporate the supernodal structures as they emerge during factorization. Therefore, our new algorithm has the combined benefits of retaining the numerical robustness of ILUTP as well as achieving fast construction and application of the ILU preconditioner. In addition, we have developed a number of new heuristics to enrich the existing dropping rules. We show that these new heuristics are helpful in improving the numerical robustness of the original ILUTP method.

A number of researchers have used blocking techniques to construct incomplete factorization preconditioners. But the extent to which the blocking has been applied is rather limited. For example, in device simulation, [Fan et al. 1996] used the four-by-four blocks which occur naturally when each grid point is associated with four variables after discretization of the coupled PDEs. Chow and Heroux [1998] used a predetermined block partitioning at a coarse level, and exploited fine-grain dense blocks to perform LU or ILU of the sparse diagonal blocks. Hénon et al. [2008] developed a general scheme for identifying supernodes in ILU(k), but it is not directly applicable to threshold-based dropping. Our algorithm is most similar to the method

proposed by Gupta and George [2008], and we extend it to the case of unsymmetric factorization with partial pivoting; see Section 3.2.

Our contributions can be summarized as follows. We adapt the classic dropping strategies of ILUTP in order to incorporate supernode structures and to accommodate dynamic supernodes due to partial pivoting. For the secondary dropping strategy, we propose an area-based fill control method, which is more flexible and numerically robust than the traditional column-based scheme. Furthermore, we incorporate several heuristics for adaptively modifying various threshold parameters as the factorization proceeds, which improves the robustness of the algorithm. Finally, the implementation of the algorithm has already been incorporated in the SuperLU Version 4.0 release, downloadable from the Internet.¹

The remainder of the article is organized as follows. In Section 2 we describe the test matrices and the performance metrics that will be used to evaluate the new algorithm. Section 3 describes in detail the new supernodal ILU algorithm together with various dropping strategies, and presents the numerical results. In Section 4 we give some remarks concerning implementation and software issues. Finally in Section 5 we compare our new code with the other codes that use different incomplete factorization algorithms.

2. NOTATIONS AND EXPERIMENTAL SETUP

We use A to denote the $n \times n$ coefficient matrix of the original linear system, and L and U to denote the incomplete triangular factors. The matrix F = L + U - I represents the filled matrix containing both L and U, and M = LU is the preconditioning matrix. D, D_r , D_c represent diagonal matrices, and P, P_r , P_c represent permutation matrices. #S denotes the cardinality of the set S. We use array section notation in Fortran and MATLAB (s:t) to refer to a range of integers $(s,s+1,\ldots,t)$. We use $\mathrm{nnz}(A)$ to denote the number of nonzeros in matrix A. The fill ratio refers to the ratio of the number of nonzeros in the filled matrix F over that in the original matrix A. Sometimes we need to refer to the fill ratio of a certain column f, that is, $\mathrm{nnz}(F(:,f))/\mathrm{nnz}(A(:,f))$. The fill ratio is a direct indicator of the memory requirement, because our code requires very little working storage. The number of operations is also related to the fill ratio, although it usually grows more linearly.

Our test machine is a Cray XT5 with 664 compute nodes operated at the National Energy Research Scientific Computing (NERSC) Center.² Each node contains two 2.4-GHz AMD Opteron quad-core processors, with 16 GB of memory. We use only one core of a node. Each core's theoretical peak floating-point performance is 9.6 Gflops. We use PathScale pathcc and pathf90 compilers with -03 -fPIC optimization flag.

We have used 232 matrices to evaluate our new ILU strategies. These include 227 matrices from the University of Florida Sparse Matrix Collection,³ and five matrices from the fusion device simulation [Jardin et al. 2007]. These are all unsymmetric matrices of medium to large size. In particular, the 227 matrices are all the real unsymmetric matrices in the UF collection which are of dimension 5000 to one million and have condition numbers below 10^{15} . The right-hand side vector is chosen such that the true solution vector is of all 1s. The iterative solver is restarted GMRES with our ILU as a right preconditioner (i.e., solving $P_rAM^{-1}y = P_rb$). The initial guess is a vector of all zeros. The stopping criterion is $||r_k|| = b - Ax_k||_2 \le \delta ||b||_2$, here we use $\delta = 10^{-8}$, which is in the order of the square root of IEEE double-precision machine epsilon. We set the dimension of the Krylov subspace to be 50 and the maximum iteration count to be 500.

 $^{^{1} \}texttt{http://crd.lbl.gov/}^{\sim} \texttt{xiaoye/SuperLU/}$

²http://www.nersc.gov/nusers/systems/hopper/

 $^{^3}$ http://www.cise.ufl.edu/research/sparse/matrices

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We mainly use two performance metrics to assess the algorithms: memory requirement as reflected by the amount of fill in the preconditioners, and the total solution time of the ILU+GMRES iterative solver. To compare different solvers, or different ILU parameter configurations, we use the performance profiles similar to those proposed by Dolan and Moré [2002] to present the data. The idea of performance profiles is as follows. Given a set $\mathcal M$ of matrices and a set $\mathcal S$ of solvers, for each matrix $m \in \mathcal M$ and solver $s \in \mathcal S$, we use fr(m,s) and t(m,s) to denote the fill ratio and total time, respectively, needed to solve m by s. If s fails to solve m for any reason (e.g., out of memory, or exceeding maximum iteration limit), we set fr(m,s) and t(m,s) to be $+\infty$ (in practice, a very large number that is outside the range of our interest is sufficient). Then, for each solver s, we define the following two cumulative distribution functions as the profiles of fill ratio and time ratio, respectively:

Prob
$$f(s, x) = \frac{\#\{m \in \mathcal{M} : fr(m, s) \le x\}}{\#\mathcal{M}}, \quad x \in \mathbb{R},$$

and

$$Prob_{t}(s,x) = \frac{\#\left\{m \in \mathcal{M} : \frac{t(m,s)}{\min_{s \in \mathcal{S}}\{t(m,s)\}} \leq x\right\}}{\#\mathcal{M}}, \quad x \in \mathbb{R}.$$

Intuitively, $\operatorname{Prob}_f(s,x)$ shows the fraction of the problems that s could solve within the fill ratio x, and $\operatorname{Prob}_t(s,x)$ shows the fraction of the problems that s could solve within a factor of x of the best solution time among all the solvers. Therefore, the plots of different solvers in the x- Prob_f or x- Prob_t coordinate can be used to differentiate the strength of the solvers in the criteria of fill ratio or time ratio—the higher the curve, the more problems the corresponding solver could solve under the same fill or time limit.

We caution that, even though the performance profile is a powerful tool to present visually the overall performance comparison of different solvers, it cannot show the difference of the solvers for each individual matrix. Therefore, in addition to the performance profiles, we will show other specific results when there is a need.

3. INCOMPLETE FACTORIZATION WITH SUPERNODES

3.1 Left-Looking Supernodal ILUTP

Our base algorithm is the left-looking, partial pivoting, supernodal sparse LU factorization algorithm implemented in SuperLU [Demmel et al. 1999a, 1999b]. The factorization algorithm proceeds from left to right, using a supernode-panel updating kernel. A panel is simply a set of consecutive columns, which is used to enhance data reuse in the memory hierarchy; it enables the use of Level 3 BLAS. The panel size is an algorithmic blocking parameter. At each step of panel factorization, we obtain a panel in the U factor and a panel in the L factor.

Several preprocessing steps are used in SuperLU before the factorization kernel, including row/column equilibration and sparsity-preserving reordering of the columns. In the case of incomplete factorization, we found that it is often beneficial to include another preprocessing step to make the initial matrix more diagonally dominant (e.g., via a maximum weighted bipartite matching algorithm). For this, we use the HSL subroutine MC64 [Duff and Koster 1999; HSL 2007], which is based on the algorithm developed by Olschowka and Neumaier [1996]. The algorithm finds a permutation and the row/column scalings so that the scaled and permuted matrix has entries of the modulus 1 on the diagonal and off-diagonal entries of the modulus bounded by 1. We tested 232 matrices with or without MC64, and found that, using MC64, the ILU-preconditioned GMRES converged for 203 matrices with an average of 12 iterations,

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Algorithm 1. Left-looking, supernode-panel ILU algorithm
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(1) Preprocessing

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(1.1) (optional) Use MC64 to find a row permutation P_0 and row and column scaling
        factors D_r and D_c such that the entries on the diagonal of P_0D_rAD_c has the largest
        absolute values.
   (1.2) If Step (1.1) is not performed, do a simple row & column equilibration to obtain D_rAD_c
        in which the entry of largest absolute value in each row and column has value 1;
   (1.3) Compute a fill-reduction column permutation P_c;
(2) Factorization of P_0D_rAD_cP_s^T
   FOR each panel of columns DO
   (2.1) Symbolic factorization: determine which supernodes to the left will update the current
        panel and a topological order of updates;
   (2.2) Panel factorization:
         FOR each updating supernode DO
              Apply triangular solve to obtain the U part;
              Apply matrix-matrix multiplication to obtain the L part;
         END FOR
   (2.3) Inner factorization:
         FOR each column j in the panel \underline{DO}
              Update the current column j;
              Apply the dropping rule to the U part;
              Find pivot in this column;
              (optional) Modify the diagonal entry to handle zero-pivot breakdown;
              Determine supernode boundary:
              IF column j starts a new supernode THEN
                 Apply the dropping rule to the newly formed supernode L(:, s: j-1),
                    where s is the first column of this supernode;
              END IF
         END FOR
   END FOR
```

whereas, without MC64, only 170 matrices converged and the average iteration count was 11.

Our incomplete factorization algorithm retains most of the algorithmic ingredients from SuperLU, with the added dropping rules that are applied to the L and U factors on-the-fly [Li et al. 2009]. The description of the algorithm is given as Algorithm 1. The steps marked as **bold** correspond to the new steps introduced to perform ILU. Note that the factorization is performed on the matrix $P_r P_0 D_r A D_c P_c^T$, where D_r and D_c are diagonal scaling matrices, P_0 is the row permutation matrix returned from MC64, P_c is the column permutation matrix for sparsity preservation, and P_r is the row permutation matrix from partial pivoting. D_r , D_c , P_0 , and P_c are obtained before the factorization, and P_r is obtained during factorization.

Our primary dropping criteria are threshold-based and akin to the ILUTP variants [Saad 1994, 1996]. That is, while performing Gaussian elimination with partial pivoting, we set to zero the entries in L and U with the modulus smaller than a prescribed relative threshold τ , where $\tau \in [0, 1]$.

Since our compressed storage is column oriented for both L and U, the dropping rule is also column oriented. The upper triangular matrix U is stored in a standard compressed column format and we can easily remove the small entries while storing the newly computed column into the compressed storage, using the first criterion given in Figure 1.

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Threshold-based dropping criteria for $ILUT(\tau)$

- (1) Dropping entries in U: If $|u_{ij}| \le \tau ||A(:,j)||_{\infty}$, we set u_{ij} to zero.
- (2) Dropping entries in L: In a supernode L(:, s:t), if $RowSize(L(i, s:t)) \le \tau$, we set the entire ith row to zero, where s and t are the first and last columns of the supernode.

Fig. 1. The threshold-based dropping criteria, with the *L*-supernode.

The lower triangular matrix L is stored as a collection of supernodes. Recall that a supernode consists of a range (r:s) of columns in L with the triangular block on the diagonal being full, and the identical nonzero structure elsewhere among the columns. Our goal is to retain the supernodal structure of the complete factorization as far as possible. To this end, we either keep or drop an entire row of a supernode when it is formed. This is similar to what was first proposed in Gupta and George [2008] and Ng et al. [1999] for incomplete Cholesky (IC) factorizations. This supernodal dropping criterion is the second rule shown in Figure 1. Since partial pivoting is used, the magnitude of the entries in L is bounded by 1, and so the absolute quantity is the same as the relative quantity.

Note that we use the *RowSize()* function to determine whether the size of a row as a whole is considered to be small and can be dropped. There are several possibilities to define this metric. In our code, we provide three choices:

- $\begin{array}{ll} \text{(i)} & \textit{RowSize}(v) \stackrel{\text{def}}{=} \frac{\|v\|_1}{m}, \\ \text{(ii)} & \textit{RowSize}(v) \stackrel{\text{def}}{=} \frac{\|v\|_2}{\sqrt{m}}, \text{and} \end{array}$
- (iii) $RowSize(v) \stackrel{\text{def}}{=} ||v||_{\infty}$,

where m is the number of columns in the supernode (i.e., m = t - s + 1). The first two metrics can be interpreted as "generalized mean" of the vector entries, whereas the last one is a standard vector norm. The following relations hold for the three metrics above:

$$\frac{\|v\|_1}{m} \le \frac{\|v\|_2}{\sqrt{m}} \le \|v\|_{\infty}.$$

In case (i), a row can be dropped when the average magnitude of the entries is small, but some entries much larger than τ can also be dropped. This is the same criterion used by Gupta and George [2008] in their IC algorithm. Our experiments showed that this metric is much worse than (iii) for ILU, probably because many large entries are dropped. In case (iii), the use of ∞ -norm implies that, when row i is dropped, the magnitude of every entry in this row is smaller than τ , and hence these entries would also be dropped in a column-wise algorithm. Therefore, from a local viewpoint, this supernodal dropping rule retains more entries compared to a column-wise algorithm. In other words, in terms of the amount of dropping, (i) is the most aggressive and (iii) is the most conservative.

Throughout the rest of the article, the supernodal ILUTP variants are used. In Section 3.3, we will show the numerical results comparing our supernodal ILU to the column-wise ILU when setting the supernode size to be 1.

3.2 Adaptive, Area-Based Dropping Rules to Limit Memory Use

 $ILUTP(\tau)$ works well if there is sufficient memory, but it may still suffer from too much fill. We employ two new ideas to further control the amount of fill. One is dynamically adjusting parameter p in the dual dropping rule; another is dynamically adjusting τ .

Several methods have been proposed for the secondary dropping rule using parameter p. In Saad's [1994] original ILUT (p, τ) approach, p is the maximum number of nonzero entries (not the level-of-fill) allowed in each row of *F* (in a row-wise algorithm), and is fixed on input. Gupta and George [2008] suggested using $p(j) = \gamma \cdot \text{nnz}(A(:,j))$ for the jth column instead of a constant, where γ is an upper bound of the fill ratio given by the user. They also proposed a method of computing a secondary dropping tolerance by an interpolation formula rather than sorting the largest p entries, which is cheaper than the original ILUT (p,τ) . According to our experience, Gupta's heuristic depends largely on the distribution of the locations of the nonzero entries in F, and the fill ratio can be either very large or very small. The performance benefit realized by not sorting is also limited. In fact, sorting is not necessary here; a selection algorithm is more suitable, because we only need to find the pth largest entry. If the number of nonzero entries in F(:, j) is k, the pth largest entry can be found by quickselect in O(k)time on average, which is the same as interpolation. Even for a sorting-based selection algorithm, the complexity is merely $O(k \log k)$ on average using quicksort and is reasonably fast in practice. Our implementation uses quickselect.

We propose a new adaptive strategy for choosing p, which takes into account the overall fill ratio up to the current step. Given a user-desired upper bound of the overall fill ratio γ , we define a continuous, upper-bound function f(j) for each column $j, f: [1,n] \to [0,+\infty)$, which satisfies $f(n) \le \gamma$. Then at the jth column, if the current fill ratio

$$\frac{\operatorname{nnz}(F(:,1:j))}{\operatorname{nnz}(A(:,1:j))} \tag{1}$$

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exceeds f(j), we choose the largest possible value p such that, when we keep the largest p entries, the current fill ratio is bounded by f(j). This criterion can be adapted to our supernodal algorithm as follows. For a supernode with k columns and j being the index of the last column of the current supernode, p may be computed as

$$p = \max \left\{ \frac{f(j) \cdot \operatorname{nnz}(A(:, 1:j)) - \operatorname{nnz}(F(:, 1:j-k))}{k}, \quad k \right\}. \tag{2}$$

In other words, if we keep the largest p rows of this supernode, the current fill ratio is guaranteed not to exceed f(j). The second k term in $\max\{\cdots\}$ is to ensure that we do not drop any row in the diagonal block of the supernode. We call this scheme an area-based $ILUT(p, \tau)$, with adaptive p.

This is more flexible than the column-based method in that it allows a larger amount of fill for certain columns as long as the cumulative fill ratio in the previous columns is small. At the end of the factorization, the total fill ratio is still bounded by γ because of the condition $f(n) \leq \gamma$.

The above description of the area-based strategy is generic, and may be used in any implementation. We now introduce a specific f(j) that is suitable for the SuperLU implementation. Since L and U are stored in different data structures, and dropping in L is invoked after a complete supernode is formed, it is sensible to use different f(j) for L and U. For a column-based method, at the jth column, the simplest way is to split γ proportionally with j:(n-j) ratio for U(:,j) and L(:,j). For our area-based approach, we may choose two functions, $f_L(j)$ for L and $f_U(j)$ for U, as long as $f_L(n) + f_U(n) \leq \gamma$. A simple way is to assign $f_L(n)$ and $f_U(n)$ to be the areas of L(:,j) and U(:,j) relative to F(:,1:j), as follows:

$$f_U(j) = \frac{j}{2n} \gamma, \quad f_L(j) = \left(1 - \frac{j}{2n}\right) \gamma.$$
 (3)

Then we split the fill quota proportionally with the $f_U(j)$: $f_L(j)$ ratio.

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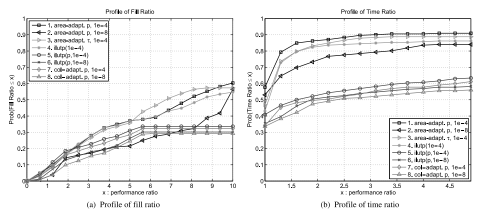


Fig. 2. Performance profiles after incorporating the secondary dropping rules; $\gamma = 10$. The running time is the total GMRES solution time with the various ILU preconditioners.

A problem is that dropping in U could be very constraining for small j. Then we can simply use $f_U(j) = \gamma/2$. With this, even though it could happen that $f_L(j) + f_U(j) \ge \gamma$ in the middle of the factorization, $f_L(n) + f_U(n) \le \gamma$ holds in the end, and the total memory is still bounded. Since we do not apply the dropping rules toward the end in order to reduce the number of zero pivots (see Section 3.4), we need to reserve some quota for them by reducing $f_L(j) + f_U(j)$. In addition, we need to allow more fill-ins in L than in U, because the dense diagonal blocks are stored in L, and some small entries in L are located in the rows with large norm and hence are not dropped. As a result, we propose to use the following heuristics:

$$f_U(j) = \frac{\gamma}{2} \times 90\%, \quad f_L(j) = \left(1 - \frac{j}{2n}\right)\gamma$$
 (4)

In conjunction with the dynamic, area-based strategy for choosing p, we devised an adaptive scheme for choosing τ as well. Specifically, let $\tau(1) = \tau_0$ be the user-input threshold. At column j, if the fill ratio given by Equation (1) is larger than f(j), we increase τ at the next step: $\tau(j+1) = \min\{1, 2\tau(j)\}$, forcing more entries to be dropped. Otherwise, we decrease τ as $\tau(j+1) = \max\{\tau_0, \tau(j)/2\}$, retaining more entries. Note that we always maintain $\tau(j) \in [\tau_0, 1]$. An advantage of this adaptive τ is that, even though the initial τ chosen by the user is too small, the code can self-adjust properly as the factorization proceeds.

We now present the results of the tests comparing various parameter settings of ILU. When secondary dropping is used, we set $\gamma = 10$. Our ILU configurations include the following:

- (1) area-based adaptive p, $\tau = 10^{-4}$;
- (2) area-based adaptive p, $\tau = 10^{-8}$;
- (3) area-based adaptive τ , $\tau_0 = 10^{-4}$, no secondary dropping;
- (4) ILUTP(τ), $\tau = 10^{-4}$;
- (5) ILUTP(p, τ), $p = \gamma \cdot \text{nnz}(A)/n, \tau = 10^{-4}$;
- (6) ILUTP $(p, \tau), p = \gamma \cdot \text{nnz}(A)/n, \tau = 10^{-8};$
- (7) column-based adaptive p (i.e., $p(j) = \gamma \cdot \text{nnz}(A(:, j))$), $\tau = 10^{-4}$;
- (8) column-based adaptive p (i.e., $p(j) = \gamma \cdot \text{nnz}(A(:, j))$), $\tau = 10^{-8}$.

Figure 2 shows the performance profiles of the fill ratio and the time ratio for the 232 test matrices. We see that a small τ such as 10^{-8} is generally not good, because

it generates too much fill and takes long to compute the solution. It is therefore not efficient to rely only on the secondary dropping rule. The threshold-based dropping criterion in Figure 1 must play a significant role. If we do not use the secondary dropping, the fraction of the problems solved within a fill-in constraint is still smaller than after using our area-based secondary dropping; see the curve corresponding to "ilutp(1e-4)" in Figure 2(a). Both the nonadaptive and the adaptive column-based secondary dropping heuristics are much worse than the area-based secondary dropping. This is clearly seen from the bottom four curves in both Figures 2(a) and 2(b).

A key conclusion is that our new area-based scheme is much more robust than the column-based scheme; it is also better than ILUTP(τ) when the fill ratio does not exceed the user-desired γ (10 in these cases). ILUTP(τ) becomes better only when the fill ratio is unbounded (i.e., allowing it to exceed γ). This is consistent with the intuition that an ILU preconditioner tends to be more robust with more fill-ins.

It is possible that the actual fill ratio is larger than the preset parameter γ although this occurs occasionally. There are two reasons for this to occur. First, our dropping rules do not drop entries in the dense diagonal blocks. Therefore, when there are some large supernodes in L, these blocks would contribute to a large memory growth even if τ is large. Second, we never drop any entries in several trailing columns (see Section 3.4), and usually there are a lot of fill-ins toward the end of the factorization. If not exceeding the memory limit is important to the user, we recommend that a slightly smaller γ be used.

Figure 2(b) shows the runtime comparison of the solvers. In this plot, the matrices with fill ratios larger than 10 are considered as failures. Thus the comparison is made under the same memory constraint, and none of the solvers are allowed to consume significantly more memory than the others. The top three solvers are much better than the others. The area-based adaptive-p or adaptive- τ schemes have quite similar performance, with the former having a slight edge over the latter.

Taking into account both memory and time, we see that the secondary dropping helps achieve a good tradeoff, with controlled fill-in and the solver not being much slower. Either the scheme corresponding to the line with "square" (area-based adaptive p, $\tau = 10^{-4}$) or the scheme corresponding to the line with "right triangle" (adaptive τ , $\tau_0 = 10^{-4}$) can be used as a default setting in the code.

3.3 Comparison of the Supernodal and the Column-wise algorithms

We have performed experiments to compare the supernodal ILU and the columnwise ILU (by setting the maximum supernode size to 1). We use both metrics (ii) $RowSize(v) \stackrel{\text{def}}{=} \|v\|_2/\sqrt{m}$ and (iii) $RowSize(v) \stackrel{\text{def}}{=} \|v\|_{\infty}$ to measure the row size in the dropping rule (2) of Figure 1.

Table I compares various performance metrics of the supernodal (S-ILU) versus column-wise (C-ILU) ILU. Each number in the table is the average of the corresponding metric taken over the number of matrices where both versions succeed. As can be seen, using either metric in the dropping rule, S-ILU is faster than C-ILU, although the speed advantage is not great compared to that in a complete factorization. This is mainly due to two reasons: (1) because of the row dropping criterion for supernode, S-ILU has more fill-ins and floating-point operations, and (2) the average supernode size is small after dropping (less than three columns on average). More fill-in in the supernodal versions translates into many more floating-point operations, especially in the conservative case when the ∞ -norm is used, for which S-ILU performs over 3.5 times more operations than C-ILU. The larger the supernodes, the more fill-ins and operations are incurred in S-ILU. From our experiments, we found that we have to set a smaller cutoff for the maximum supernode size (maxsuper) than in complete

Table I. Comparison of Supernodal ILU (S-ILU, with maxsuper=20) and Column-wise ILU (C-ILU, with maxsuper=1) Using "Mean" Statistics About Various Metrics; Two RowSize() Functions Used; Times are in Seconds. $\tau=10^{-4}, \gamma=10$.

	Factor construction					Total time		
	Fill-ratio	s-Node size (columns)	Flops (10 ⁹)	Factor time	Iters	One triangular solution time	Iter time	
	(ii) $RowSize(v) \stackrel{\text{def}}{=} \frac{\ v\ _2}{\sqrt{m}}$; 138 problems for which both versions succeeded							
S-ILU	4.2	2.80	7.60	39.69	21.6	0.0092	2.93	42.68
C-ILU	3.7	1	2.65	65.15	20.0	0.0079	2.55	67.75
	(iii) $RowSize(v) \stackrel{\text{def}}{=} v _{\infty}$; 134 problems for which both versions succeeded							
S-ILU	4.2	2.72	9.45	54.44	20.5	0.0109	3.40	57.90
C-ILII	3.6	1	2.58	74 10	19.8	0.0090	2.88	77 04

factorization (i.e., when a supernode size exceeds maxsuper columns, we will break this supernode into two supernodes). Otherwise the extra operations would offset the BLAS-3 benefit. For example, in S-ILU, maxsuper = 20 strikes a good balance in overall speed, whereas, in the complete LU, we often use maxsuper = 100.

In these tests, we excluded one matrix *cage13*, because the factorization time was about 1000x slower than the average, and the inclusion of this matrix would severely skew the statistics of the mean metric.

3.4 Handling Breakdown Due to Zero Pivots

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In the case of LU factorization with partial pivoting, zero pivots may occur due to numerical cancellations when the matrix is nearly singular. However, for an incomplete LU factorization, zero pivots may occur more often because of dropping, which has nothing to do with numerical cancellation.

To illustrate this, let us consider the following two 2×2 matrices:

$$A_1 = \begin{bmatrix} a & b \\ c & 0 \end{bmatrix}, \qquad A_2 = \begin{bmatrix} c & 0 \\ a & b \end{bmatrix}, \qquad (bc \neq 0)$$

Assume that the column permutation is the identity. Thus, if $|c| < \tau |a|$, the (2, 1) entry will be dropped, and the (2, 2) entry will become zero, causing ILU to break down. Assuming that a, b, and c are drawn independently from the uniform distribution in [-1, 1], we have

$$Prob\{u_{22}=0\}=\frac{\tau}{2}>0.$$

In general, let us assume that the nonzero entries of a nonsingular matrix satisfy a uniform distribution in [-1,1]. Then for a given sparsity pattern, if there exist a row permutation P and a column j such that (PA)(j:n,j)=0 and (PA)(1:j-1,1:j-1) is nonsingular, the probability of encountering a zero pivot in the jth column would be positive. We can show that $(\tau/2)^{\operatorname{nnz}(A)-n}$ is a lower bound of the probability. Let B=PA and suppose that j is the minimum column index which satisfies B(j:n,j)=0. The probability that all the pivots of the first j-1 columns are diagonal entries of B and all the off-diagonal entries are dropped (with this condition, definitely there will be a zero pivot in the jth column) is

$$\begin{split} \prod_{i=1}^{j-1} \prod_{k>i} \text{Prob} \big\{ |B(k,i)| \, < \, \tau \, |B(i,i)| \big\} \; &= \; \prod_{i=1}^{j-1} \left(\frac{\tau}{2} \right)^{\text{nnz}(B(i+1:n,i))} \\ &\geq \; \left(\frac{\tau}{2} \right)^{\text{nnz}(A(:,1:j-1)) - (j-1)} \, \geq \, \left(\frac{\tau}{2} \right)^{\text{nnz}(A) - n} \, . \end{split}$$

Algorithm 2. Classical column-wise MILU for column *j*

- (1) Obtain the current filled column F(:, j);
- (2) Pivot: Choose pivot row *i*, such that $i = \arg \max_{i>j} |f_{ij}|$; Swap rows *i* and *j*;
- (3) Apply a dropping rule to F(:, j);
- (4) Compute the sum of dropped entries in F(:, j): $s = \sum_{\text{dropped}} f_{ij}$; Set $f_{jj} := f_{jj} + s$;
- (5) Separate *U* and *L*: U(1:j,j) := F(1:j,j); L(j:n,j) := F(j:n,j)/F(j,j);

The last inequality comes from the fact that there is at least one nonzero entry in each column of a nonsingular matrix.

On the other hand, if j is the first column index that encounters zero pivot, the matrix must satisfy the condition that (PA)(j:n,j)=0 and (PA)(1:j-1,1:j-1) is nonsingular.

Usually, many zero pivots occur in the last columns, because it is more probable at the end than in the beginning that all the nonzero entries of a column are permuted to the upper triangular part. To mitigate this, we stop dropping when the column index is larger than $\max\{n-2N_s, n\times 95\%\}$, where N_s is the maximum size of a supernode. That is, the factorization is almost completed. According to our experiments, this helps reduce a large fraction of the zero pivots for some matrices.

We have devised a simple adaptive mechanism to handle the situation when a zero pivot indeed occurs. At column j, when we encounter $u_{jj}=0$, we set it to $\hat{\tau} \|A(:,j)\|_{\infty}$ to ensure the factorization can continue and U is nonsingular after the factorization. This is equivalent to adding a small perturbation $\hat{\tau}$ to l_{ij} at the current step. If $\hat{\tau}=\tau$, the perturbation we add to u_{jj} will not exceed the upper bound of the error propagated by dropping entries. In our code, we choose $\hat{\tau}(j)=10^{-2(1-j/n)}$, which is an increasing function with the column index, rather than a constant. This prevents the diagonal entries of U from being too small; otherwise, it could result in a very ill-conditioned preconditioner.

Adding a small perturbation on the zero diagonal is a simple remedy to enable the factorization to complete. This is an acceptable solution when not many zero pivots occur; otherwise, the preconditioner can be quite ill-conditioned even though the factorization completes, making this ineffective. Some other methods were proposed to handle the breakdown, such as the delayed pivoting [Gupta and George 2008] and the multilevel method [Bollhöfer and Saad 2006]. We plan to investigate them in the future. But our comparisons show that our current ILU scheme is very competitive with a multilevel ILU scheme as in ILUPACK [Bollhöfer et al. 2006], see Section 5.

3.5 Modified ILU (MILU)

The MILU techniques were introduced to reduce the effect of dropping by compensating for the discarded entries [Saad 1996]. For systems arising from the discretization of second-order elliptic PDEs, it is generally admitted that the modified incomplete Cholesky (MIC) factorization is more efficient than the unmodified one [Dupont et al. 1968; Greenbaum 1997; Gustaffson 1978; Notay 1994; van der Vorst 1990]. However for general systems, very little is known either theoretically or empirically whether MILU helps. Our empirical experience shows that sometimes it can be better than the unmodified ILU.

The basic idea is to add the dropped entries in a row or column to the diagonal of U. The commonly used strategy has an appealing property that it preserves the rowsum relation $P_rAe = \tilde{L}\tilde{U}e$ for a row-wise algorithm or column-sum relation $e^TP_rA = e^T\tilde{L}\tilde{U}$ for a column-wise algorithm, where \tilde{L} and \tilde{U} are incomplete factors. Algorithm 2 gives

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Algorithm 3. SMILU-1: Supernodal MILU for column *j*

(1) Obtain the current filled column F(:, j);

```
(2) Apply dropping rule 1) of Figure 1 to F(1:j,j), and set U(1:j,j):=F(1:j,j);
(3) Compute the sum of the dropped entries in U(:,j):s=\sum_{\text{dropped}}u_{ij};
(4) Pivot: Choose pivot row i, such that i=\arg\max_{i\geq j}|\mathbf{f_{ij}}+\mathbf{s}|;
\operatorname{Swap\ rows\ } i \text{ and } j, \text{ and set } u_{jj}:=f_{ij}+s;
(5) \underline{\operatorname{IF}}\ j \text{ starts\ } a \text{ new\ supernode\ } \underline{\operatorname{THEN}}
\operatorname{Let}\ (r:t) \text{ be the newly formed supernode; } (t\equiv j-1)
\operatorname{Apply\ dropping\ rule\ } 2) \text{ of\ Figure\ } 1 \text{ to\ the\ supernode\ } L(:,r:t);
\operatorname{For\ each\ column\ } k \text{ in\ the\ supernode\ } (r\leq k\leq t):
\operatorname{compute\ the\ sum\ of\ the\ dropped\ entries:\ } S_k=\sum_{i \text{ dropped\ }} l_{ik};
\operatorname{set\ } u_{kk}:=u_{kk}+S_k\cdot u_{kk};
\operatorname{END\ IF;}
```

the procedure to perform a column-wise MILU with partial pivoting. Note that for the upper triangular part $f_{ij} = u_{ij}$, whereas for the lower triangular part $f_{ij} = l_{ij}u_{,ij}$ because of division by the diagonal entry $f_{,ij}$.

We have modified the classical column-wise MILU procedure to accommodate our supernodal dropping criteria. Recall that in Algorithm 1, we apply the dropping rule (2) in Figure 1 only after a new supernode in L is formed. The consequence of this "delayed" dropping is that, at the time a column is processed for pivoting, the computed sum s may contain fewer dropped entries and the diagonal is not compensated enough. Therefore, the column-sum relation is not preserved.

Our supernodal version of the MILU strategy works as follows. For each column of U, we accumulate in s the sum of the dropped entries. Later on, s is not only added to the diagonal but is also used during pivot selection in the lower triangular part. The procedure is outlined in Algorithm 3 and we call it SMILU-1. This algorithm ensures that the pivot has the largest magnitude after the entries are dropped from the upper triangular part. However, we cannot guarantee that the pivot still has a relatively large absolute value after the entries in the lower triangular are dropped in a future step when the supernode is formed. The pivot could become small or even zero after we apply the dropping rule to L (i.e., after applying (5) in Algorithm 3.) This may cause the factor U to be ill-conditioned, resulting in an unstable preconditioner. A slightly modified algorithm, which we call SMILU-2, provides a remedy. Here, we take s to be the absolute value of the dropped sum: $s = |\sum_{\text{dropped}} u_{ij}|$, and the diagonal u_{ij} is set to be $u_{jj} := f_{ij} + \text{sign}(f_{ij})s$. This ensures that the magnitude of the pivot is nondecreasing after diagonal compensation, thereby avoiding small pivots. An alternative method is to accumulate the one-norm of the dropped vector: $\hat{s} = \sum_{\text{dropped}} |u_{ij}|$. Thus, the pivots would have larger absolute values compared to what would be in SMILU-2, and we expect the condition number of U to be smaller; we call this SMILU-3.

Another twist with MILU is to incorporate diagonal perturbations. In the earlier investigations of MIC (e.g., Notay [1994] and the references therein), people found that the unperturbed MIC may lead to unpredictable behavior of the largest eigenvalues of the preconditioned matrix. Different perturbations were introduced which could upper-bound the eigenvalues for special problems, such as Stieltjes matrices. One of the best is Notay's [1994] DRIC, a dynamic, relaxed IC. We adopted this scheme in our MILU variants above. Specifically, the diagonal u_{ij} is modified as follows:

- (1) Choose $\alpha = n^{-\frac{1}{d}}$ (for discretized PDEs, d can be the dimension of the space).
- (2) Set $\alpha_j = s/u_{jj}$, s is computed as in Step (3) of Algorithm 3.

Table II. Comparison of Various (M)ILUTP(τ , p)+GMRES Algorithms and SuperLU; Column "Zero pivots"
Indicates Number of Zero Pivots Encountered During ILU Factorization; Default Secondary Dropping
Strategy Is Used

		Converge	Slow	Diverge	Memory	Zero pivots	Avg. iterations
$\tau = 10^{-4}$	$ILUTP(\tau, p)$	140	42	48	2	2438	23
	SMILU-1	128	69	33	2	936	30
	SMILU-2	133	66	31	2	255	30
	SMILU-3	132	69	29	2	242	40
$\tau = 10^{-6}$	$ILUTP(\tau, p)$	128	53	49	2	2462	29
	SMILU-1	120	71	39	2	382	28
	SMILU-2	128	63	39	2	317	39
	SMILU-3	126	69	35	2	299	32
$(\tau = 0.0)$	SuperLU	222	0	0	10	0	1

- $\begin{array}{ll} \text{(3) Choose } \omega_j = \mathrm{sign}(\alpha_j) \, \min \left\{ \left| \frac{2(1-\alpha)}{\alpha_j} \right|, 1 \right\} & \text{(i.e., } -1 \leq \omega_j \leq 1). \\ \text{(4) Set } u_{jj} := f_{ij} + s \cdot \omega_j \text{ in Step (4) of Algorithm 3.} \end{array}$

Intuitively speaking, this ensures that we do not compensate too much on the diagonal if the sum of the dropped entries is large. Note that Notay provided theoretical arguments that the above choice of ω_i provides upper spectral bound for certain class of matrices, but no theoretical justifications exist for general unsymmetric matrices. We observed only empirically that it improves convergence in some cases.

In Table II, we compare the performance of various ILUTP+GMRES algorithms and direct solver SuperLU. We classify the GMRES convergence history in three categories: converge, slow (meaning the preset maximum iteration count 500 is exceeded although the residual norm is still decreasing, i.e, $\delta < \|r\|_2 / \|b\|_2 < 1$), and diverge (meaning $||r||_2 \ge ||b||_2$). The column labeled *memory* means the code runs out of memory. The (unmodified) ILUTP(τ) usually works very well; however, when it fails, it is often due to too many zero pivots. Looking at the slow and diverge columns, we see that for a number of matrices the MILU algorithms could succeed if they are allowed to continue with more iterations, whereas ILU would fail completely.

The performance profiles of the various MILU algorithms show that there is little difference to choose between the variants, with SMILU-2 being slightly better than the other two MILU schemes, and the MILU schemes are slightly worse than ILUTP. Therefore, we do not recommend it as the default in the software, but make it available as an option.

3.6 Threshold Partial Pivoting

For some matrices with band structure or close to diagonally dominant, sometimes we can trade partial pivoting for a sparser factorization. Therefore, we provide a relaxed pivoting strategy that gives preference to the diagonal entries. We use a threshold parameter $\eta \in [0, 1]$ to facilitate this. If $|f_{jj}| \ge \eta \max_{i \ge j} \{|f_{ij}|\}$, the diagonal entry f_{jj} is used as the pivot. Thus, $\eta = 1.0$ corresponds to partial pivoting, and $\eta = 0.0$ corresponds to diagonal pivoting. Usually, η cannot be too small if the numerical property of the matrix is unknown because the magnitude of the entries in L can grow as much as η^{-1} . In general, even though the pivot growth can be a bit larger than 1, the dropped entries are still relatively small. Tables III and IV present the numerical results when varying the pivot tolerance η , without and with secondary dropping, respectively.

When secondary dropping is not used (Table III), (threshold) pivoting is more reliable than no pivoting at all, because general matrices are not close to being diagonally dominant.

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Table III. Effect of Threshold Parameter η	with I	LUTP((τ)
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η		1.0	0.1	0.01	0.001	0
$\tau = 10^{-4}$	Number of successes	199	203	201	201	191
	Average fill ratio	17.3	17.0	15.3	15.6	10.0
	Maximum fill ratio	371	312	98	100	92
	Average iterations	7.8	11.7	11.5	15.4	14.7
$\tau = 10^{-6}$	Number of successes	209	209	208	205	203
	Average fill ratio	34.5	34.7	33.0	31.5	28.6
	Maximum fill ratio	1200	1200	1200	1200	1200
	Average iterations	4.1	3.7	3.8	3.3	6.6

Table IV. Effect of the Threshold Parameter η with ILUTP (τ, p) , Using Area-Based Secondary Dropping, $\gamma = 10$

n		1.0	0.1	0.01	0.001	0
$\tau = 10^{-4}$	Number of successes	132	140	138	136	142
	Average fill ratio	4.3	4.3	4.2	4.2	4.1
	Maximum fill ratio	9.6	9.6	9.6	9.6	9.6
	Average iterations	18	23	22	26	28
$\tau = 10^{-6}$	Number of successes	125	127	129	129	129
	Average fill ratio	5.3	5.4	5.4	5.4	5.4
	Maximum fill ratio	9.7	9.7	9.7	9.7	9.7
	Average iterations	31	30	30	33	29

When secondary dropping is used (Table IV), the situation is not very conclusive, and it is difficult to choose a good η . This is mainly because the influence of drop tolerance becomes insignificant in the presence of secondary dropping. Comparing to Table III, it is clear that the numbers of problems successfully solved are much smaller, and the iteration counts are much larger. The fill ratios are bounded by γ and are much smaller than those in Table III.

Based on our experience, it is better not to use secondary dropping when memory is not at a premium. As a comparison, for complete factorization with $\eta = 0.1$, the average fill ratio is 90.7 and the maximum fill ratio is 1577.2.

4. COMMENTS ON THE SOFTWARE

In this section, we describe a few implementation difficulties encountered while developing our incomplete factorization, and summarize the input parameters introduced to the ILU routine.

4.1 Difficulty with Symmetric Pruning

Symmetric pruning is a technique to find a smaller graph (symmetric reduction) in place of $G(L^T)$ that maintains the path-preserving property. Using symmetric reduction can speed up the depth-first search traversals (i.e., the symbolic factorization) which are interleaved with the numerical factorization steps. Specifically, at step j, the symmetric reduction of the current factor L(:, 1:j) is obtained by removing all the nonzero entries l_{rs} for which $l_{ts}u_{st} \neq 0$ for some $t < \min(r, j)$ [Demmel et al. 1999a]. That is, in L, the nonzero entries below the first matching nonzero pair in column and row s < j of the factor F(1:j,1:j) can be removed. Consider the following 4×4

matrix A, the filled matrix F (using the given elimination order), and the symmetric reduction R:

$$A = \begin{bmatrix} \bullet & \bullet & \bullet & \bullet \\ \bullet & \bullet & \bullet \\ \bullet & \bullet & \bullet \end{bmatrix}, \quad F = \begin{bmatrix} \bullet & \bullet & \bullet & \bullet \\ \bullet & \bullet & \circ & \circ \\ \bullet & \circ & \circ & \circ \end{bmatrix}, \quad R = \begin{bmatrix} \bullet & \bullet & \bullet & \bullet \\ \bullet & \bullet & \circ & \circ \\ \bullet & \bullet & \circ & \circ \\ \bullet & \circ & \circ & \circ \end{bmatrix}.$$

In F and R, a symbol \circ indicates a fill-in entry. In R, a symbol \ominus indicates a removed entry from symmetric pruning, that is, l_{41} is removed due to the matching nonzero pair l_{21} and u_{12} . If G(F) is used in the depth-first traversal, the entry l_{43} is obtained by the following path 4 :

$$3 \xrightarrow{A} 1 \xrightarrow{F} 4$$
.

When using the reduced graph G(R), the above path is replaced by the following one, and the reachability is maintained:

$$3 \xrightarrow{A} 1 \xrightarrow{R} 2 \xrightarrow{R} 4$$
.

However, in an incomplete factorization, if the magnitude of l_{42} is smaller than the threshold, it would be dropped both in F and in R. Then the edge $2 \stackrel{R}{\longrightarrow} 4$ does not exist anymore. The entry l_{43} would be missing if R were used for the depth-first search, and similarly for l_{44} . The erroneous R is shown below, where \otimes indicates a dropped entry in ILU.

$$R_{ilu} = \begin{bmatrix} \bullet & \bullet & \bullet & \bullet \\ \bullet & \bullet & \circ & \circ \\ & \bullet & & \bullet \\ \ominus & \otimes & & \end{bmatrix}.$$

We thought about several ways to mitigate this problem, such as delayed pruning or protecting pruned entries from dropping. But their implementations would incur nontrivial costs in runtime and memory. We did some tests to evaluate the benefit of pruning. For complete factorization, even if the pruned graph is very small, that is, of size R less than 5% that of F, the total speedup is usually no more than 20%. For incomplete factorization, since the fill ratio is often much smaller (i.e., F is already quite small), we expect the benefit of pruning would be less. Therefore, we decided not to use a reduced graph.

4.2 Zero Pivots and Relaxed Supernodes

In SuperLU's complete factorization, we use relaxed supernodes to increase the average size of supernodes (or block size). We group several columns at the bottom of the column elimination tree into an artificial supernode [Demmel et al. 1999a]. The column elimination tree is the elimination tree (etree) of $|A|^T |A|$, which shows the columns' dependencies for any row permutation (partial pivoting). That is, the relaxed

⁴We use the convention that an edge is directed from a column to a row of the matrix.

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Options	Default			
Drop tolerance (τ)	10^{-4}			
RowSize()	∞-norm			
Secondary dropping	Area-based adaptive-p			
Fill-ratio bound (γ)	10			
SMILU	No			
MC64	ON			
Equilibration	ON			
Column permutation	COLAMD [Davis et al. 2004]			
Diagonal threshold (n)	0.1			

Table V. Default Values of Parameters of ILU Routine xGSITRF

supernodes at the bottom of the etree will not be modified by any other columns outside these supernodes. Given a postordered etree, this means that the nonzero row structure of a column L(:,j) must be disjoint from that of a later supernode (r:s)>j. Otherwise, there exists a numerical assignment such that a common row i can be selected as a pivot at step j, making supernode (r:s) dependent on column j. Therefore, selecting any pivot column j has no impact on supernode (r:s).

On the other hand, in an incomplete factorization, if zero pivot occurs in column j due to dropping, we cannot choose a random row below the diagonal as pivot, because it could overlap with a row in the future relaxed supernodes, which in essence changes the etree structure and dependency. Therefore, we must choose a pivot row which does not appear in any later relaxed supernode.

4.3 Tunable Parameters in the ILU Routine

The new ILU routine is named xGSITRF. It takes a structure options as the first argument, which contains a set of parameters to control how the ILU decomposition will be performed. The default values of these parameters are listed in Table V, which are set by calling the routine ilu_set_default_options().

Based on our experience, we provide the following guidelines regarding how to adjust the parameters if the defaults do not work.

- Equilibration is necessary, and MC64 is usually helpful.
- If there are many zero pivots and the preconditioner is too ill conditioned, you could try the modified variants SMILU-2 or SMILU-3.
- If the fill ratio is still small, you may try a smaller τ .
- If you run out of memory, you may try a smaller γ and a smaller η .

5. COMPARISON WITH THE OTHER PRECONDITIONERS

A number of preconditioning packages using incomplete factorization algorithms have been developed for unsymmetric matrices. In this section, we compare the performance of our new ILU preconditioner with SPARSKIT, a basic toolkit for sparse matrix computations,⁵ and ILUPACK [Bollhöfer et al. 2006].

These preconditioners are used in the standard GMRES iterative solver.

The algorithm in SPARSKIT is the original ILUTP algorithm proposed by Saad, and is the closest to our algorithm. The primary differences are (1) SPARSKIT performs factorization row by row and does not use supernodes, whereas ours is a column-wise

⁵http://www-users.cm.umn.edu/~saad/software/SPARSKIT/sparskit.html

algorithm and exploits supernodes; (2) the secondary dropping in SPARSKIT is row-based and p is a fixed constant, whereas ours is area based with adaptive p.

ILUPACK uses a very different approach to ours and SPARSKIT; it is an inverse-based method, and uses a relatively new multilevel approach to handle small pivots. The inverse-based approach attempts to control the size of the inverse of the preconditioner so that the condition number of the preconditioner is under control. This objective is achieved indirectly: at step k of the factorization, the algorithm monitors the norm of the kth row of L^{-1} . If that exceeds the prescribed bound ν , implying no suitable pivot can be chosen at this step, then row k and column k are moved to the end, and the factorization continues to the next row/column. After all the suitable pivots are chosen, the current level is considered to be complete, and the factorization starts a new level, which is comprised of all the delayed rows and columns from the previous level.

In our comparisons, we use the latest versions of SPARSKIT (version 2.0) and ILU-PACK (version 2.3). We use pathf90 -03 -fPIC to compile SPARSKIT which is in Fortran, and pathcc -03 -fPIC to compile SuperLU and ILUPACK which are in C, and linked with the AMD Core Math Library (ACML) for the BLAS routines. In our experiments, we tried to keep the similar parameter settings for all three codes:

- SuperLU 4.0: $\tau = 10^{-4}$, area-based secondary dropping with $\gamma = 5$ or 10, diagonal threshold $\eta = 0.1$;
- SPARSKIT: $\tau = 10^{-4}$, secondary dropping with $p = \gamma \cdot nnz(A)/n(A)$, where $\gamma = 5$ or 10, diagonal threshold $\eta = 0.1$;
- ILUPACK: $\tau = 10^{-4}$, $\nu = 5$, $\gamma = 5$ or 10 (corresponding to param. elbow in the code).

We use the default sparsity reordering options, which are different among the three codes due to different numerical pivoting strategies. Our ILU performs partial pivoting with row swappings and uses a column reordering method such as Column Approximate Minimum Degree [Davis et al. 2004], for which the underlying graph model is the adjacency graph of $|A|^T|A|$. This attempts to minimize an upper bound on the fill *regardless of row interchanges*. ILUPACK does not perform pivoting and uses a symmetric reordering method such as Approximate Minimum Degree [Amestoy et al. 1996], for which the underlying graph model is the adjacency graph of $|A|^T + |A|$.

SPARSKIT does not have sparsity reordering phase; we added a preprocessing step for this purpose. Since SPARSKIT performs elimination row-wise with column interchanges for partial pivoting, we need to compute a sparsity ordering based on the graph of $|A||A|^T$, and apply this permutation to the rows of A before the factorization. In our experiment, this permutation is computed by applying COLAMD to A^T . But we found that the quality of the ILU preconditioner with reordering is worse than that from the default row ordering. Therefore, in the following, we present the SPARSKIT results from the default row ordering.

Figure 3(a) shows the fill profiles of the three preconditioners with $\gamma=5$ in the secondary dropping. For smaller allowable fill ratio, ILUPACK could solve a few more problems than our ILU. However, when the fill ratio is close to the prescribed limit γ , our code can solve more problems. Both our ILU and ILUPACK are better than SPARSKIT. For a larger γ , the curves of profile are changed; see Figure 3(b). The left plots are different than just cutting off at $\gamma=5$ in the right figure.

Figure 4 compares the runtime of the GMRES solver using the three preconditioners. This shows that our area-based adaptive $ILUTP(\tau, p)$ is generally superior to the other two preconditioners. Recall that p is a function of γ and is computed adaptively at each step using Equation (2). Only at the leftmost part of Figure 4(b), when allowing the similar amount of time, ILUPACK and SPARSKIT are slightly better.

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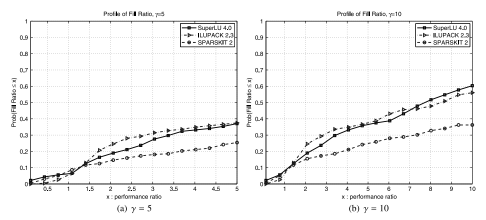


Fig. 3. Comparison of fill ratio between our supernodal ILUTP, ILUPACK and SPARSKIT.

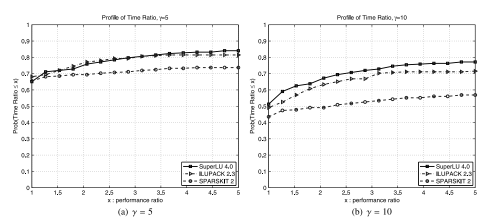


Fig. 4. Comparison of runtime of the GMRES solver using the three preconditioners.

Figure 5 compares the number of iterations of the GMRES solver using the three preconditioners. This is very much correlated to the runtime profiles shown in Figure 4. For $\gamma=5$, our ILUTP preconditioner and ILUPACK can solve almost the same number of problems within a prescribed number of iterations, and can solve more than SPARSKIT does.

Looking only at the performance profiles in Figures 3–5, our code and ILUPACK seem to be comparable: ours is slightly faster and ILUPACK maintains slightly lower fill. On the other hand, none of the preconditioners can succeed with all the problems, even though they are designed as general-purpose preconditioners. This is different from direct methods, which almost always succeed as long as there is enough memory. Therefore, in addition to performance profiles, we need to examine preconditioners from different perspectives. Among the 232 problems, our code succeeds with 140 problems, and ILUPACK succeeds with 130 problems. There is a common set of 100 problems on which both codes succeed. This shows that, even though the two preconditioners have similar success rates they succeed with different sets of problems, and so the two methods can be considered complementary to one another, and both have practical values in existence.

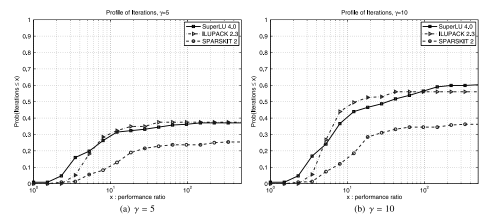


Fig. 5. Comparison of iteration count of the GMRES solver using the three preconditioners.

6. CONCLUSIONS

We adapted the classic dropping strategies of ILUTP in order to incorporate supernode structures and to accommodate dynamic supernodes due to partial pivoting. For the secondary dropping strategy, we proposed an area-based fill control mechanism which is more flexible and numerically more stable than the traditional column-based scheme. Furthermore, we incorporated several heuristics for adaptively modifying various threshold parameters as the factorization proceeds, which improves the robustness of the algorithm. The numerical experiments show that our new supernodal ILU algorithm is competitive with the inverse-based multilevel ILU method implemented in ILUPACK. The new ILU routine is already released to public in SuperLU Version 4.0, which can be downloaded from the Internet.

In the future, we plan to investigate different methods for handling zero pivots to enhance stability of the factorization, to add more adaptivity, and to study the preconditioning effect with the other iterative solvers.

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REFERENCES

AMESTOY, P. R., DAVIS, T. A., AND DUFF, I. S. 1996. An approximate minimum degree ordering algorithm. SIAM J. Matrix Anal. Appl. 17, 4, 886–905. Tech rep.

BOLLHÖFER, M. AND SAAD, Y. 2006. Multilevel preconditioners constructed from inverse-based ILUs. SIAM J. Sci. Comput. 27, 5, 1627–1650.

BOLLHÖFER, M., SAAD, Y., AND SCHENK, O. 2006. ILUPACK-preconditioning software package. Technical University of Braunschweig, Braunschweig, Germany. http://ilupack.tu-bs.de.

CHOW, E. AND HEROUX, M. A. 1998. An object-oriented framework for block preconditioning. ACM Trans. Math. Softw. 24, 159–183.

DAVIS, T. A., GILBERT, J. R., LARIMORE, S., AND NG, E. 2004. A column approximate minimum degree ordering algorithm. *ACM Trans. Math. Softw. 30*, 3, 353–376.

DEMMEL, J. W., EISENSTAT, S. C., GILBERT, J. R., LI, X. S., AND LIU, J. W. H. 1999a. A supernodal approach to sparse partial pivoting. SIAM J. Matrix Anal. Appl. 20, 3, 720–755.

DEMMEL, J. W., GILBERT, J. R., AND LI, X. S. 1999b. SuperLU users' guide. Tech. rep. LBNL-44289. Lawrence Berkeley National Laboratory, Berkeley, CA. http://crd.lbl.gov/~xiaoye/SuperLU/.

 $^{^6}$ http://crd.lbl.gov/ $^\sim$ xiaoye/SuperLU/

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- DOLAN, E. D. AND MORÉ, J. J. 2002. Benchmarking optimization software with performance profiles. *Math. Program. 91*, 2, 201–203.
- DUFF, I. S. AND KOSTER, J. 1999. The design and use of algorithms for permuting large entries to the diagonal of sparse matrices. SIAM J. Matrix Anal. Appl. 20, 4, 889–901.
- DUPONT, T., KENDALL, R. P., AND H. H. RACHFORD, J. 1968. An approximate factorization procedure for solving self-adjoint elliptic difference equations. SIAM J. Numer. Analy. 5, 559–573.
- FAN, Q., FORSYTH, P. A., MCMACKEN, J. R. E., AND TANG, W.-P. 1996. Performance issues for iterative solvers in device simulation. SIAM J. Sci. Comput. 17, 1, 100–117.
- GREENBAUM, A. 1997. Iterative Methods for Solving Linear Systems. SIAM, Philadelphia, PA.
- GUPTA, A. AND GEORGE, T. 2008. Adaptive techniques for improving the performance of incomplete factorization preconditioning. Tech. rep. RC 24598(W0807-036). IBM Research, Yorktown Heights, NY.
- GUSTAFFSON, I. 1978. A class of first order factorization methods. BIT 18, 142-156.
- HÉNON, P., RAMET, P., AND ROMAN, J. 2008. On finding approximate supernodes for an efficient ILU(k) factorization. *Parall. Comput.* 34, 345–362.
- HSL. 2007. HSL. http://www.cse.scitech.ac.uk/nag/hsl/hsl.shtml.
- HYSOM, D. AND POTHEN, A. 2001. A scalable parallel algorithm for incomplete factor preconditioning. SIAM J. Sci. Comput. 22, 6, 2194–2215.
- JARDIN, S. C., BRESLAU, J., AND FERRARO, N. 2007. A high-order implicit finite element method for integrating the two-fluid magnetohydrodynamic equations in two dimensions. J. Comput. Phys. 226, 2146-2174.
- LI, X., SHAO, M., YAMAZAKI, I., AND NG, E. 2009. Factorization-based sparse solvers and preconditioners. In *Proceeding the Conference*, Journal of Physics: Conference Series 180 (2009) 012015, Institute of Physics Publishing, San Diego, CA.
- NG, E. G., PEYTON, B. W., AND RAGHAVAN, P. 1999. A blocked incomplete Cholesky preconditioner for hierarchical-memory computers. In *Proceedings of Iterative Methods in Scientific Computations IV*. D. R. Kincard and A. C. Elster, Eds. IMACS, 211–222.
- NOTAY, Y. 1994. DRIC: A dynamic version of the RIC method. Numer. Lin. Alg. Appl. 1, 6, 511-532.
- OLSCHOWKA, M. AND NEUMAIER, A. 1996. A new pivoting strategy for Gaussian elimination. *Lin. Alg. Appl. 240*, 131–151.
- SAAD, Y. 1994. ILUT: A dual threshold incomplete LU factorization. Numer. Lin. Alg. Appl. 1, 4, 387-402.
- SAAD, Y. 1996. Iterative Methods for Sparse Linear Systems. PWS Publishing Company, Boston, MA.
- VAN DER VORST, H. 1990. The convergence behaviour of preconditioned CG and CGS in the presence of rounding errors. In *Preconditioned Conjugate Gradient Methods*, O. Axelsson and L. Kolotilina, Eds., Lecture Notes in Mathematics, vol. 14570. Springer Verlag, Berlin, Germany.

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