The Reverse Cuthill-McKee Algorithm in Distributed-Memory

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Abstract—Ordering vertices of a graph is key to minimize fill-in and data structure size in sparse direct solvers, maximize locality in iterative solvers, and improve performance in graph algorithms. Except for naturally parallelizable ordering methods such as nested dissection, many important ordering methods have not been efficiently mapped to distributed-memory architectures. In this paper, we present the first-ever distributed-memory implementation of the reverse Cuthill-McKee (RCM) algorithm for reducing the profile of a sparse matrix. Our parallelization uses a two-dimensional sparse matrix decomposition. We achieve high performance by decomposing the problem into a small number of primitives and utilizing optimized implementations of these primitives. Our implementation shows strong scaling up to 1024 cores for smaller matrices and up to 4096 cores for larger matrices.

I. INTRODUCTION

Reordering a sparse matrix to reduce its bandwidth or profile can speed up many sparse matrix computations [1], [2]. For example, a matrix with a small profile is useful in direct methods for solving sparse linear systems since it allows a simple data structure to be used. It is also useful in iterative methods because the nonzero elements will be clustered close to the diagonal, thereby enhancing data locality. Given a symmetric matrix A, a bandwidth-reduction ordering aims to find a permutation P so that the bandwidth of PAP^{T} is small. Since obtaining a reordering to minimize bandwidth is an NPcomplete problem [3], various heuristics are used in practice such as Cuthill-McKee, Reverse Cuthill-McKee (RCM), and Sloan's algorithms [4], [5], [6]. This paper solely focuses on the RCM algorithm [5] because, with careful algorithm design, it is amenable to massive distributed-memory parallelism – the primary topic of interest of this paper.

The need for distributed memory RCM algorithms is driven by the move to extreme-scale computing. In a large scale scientific application, the matrix has often been distributed already by the time one arrives at the numerical phases of sparse matrix computations. Hence, it would be a waste of time to gather a distributed matrix onto a single processor to compute an ordering using sequential or multithreaded algorithms. Furthermore, by clustering the nonzeros closer to the diagonal, RCM ordering not only increases cache performance of iterative solvers, but it can often restrict the communication

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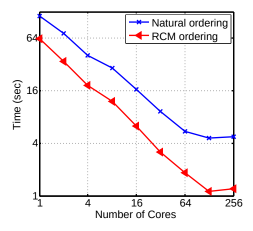


Fig. 1: The time to solve thermal2 using the conjugate gradient method with block Jacobi preconditioner in PETSc on NERSC/Edison. thermal2 has 1.2M rows and columns and 4.9M nonzeros. The bandwidths of the original and RCM-permuted matrix are 1,226,000 and 795, respectively.

to resemble more of a nearest-neighbor pattern. Figure 1 illustrates the performance effects of RCM ordering on a preconditioned conjugate gradient solver of the popular PETSc package [7]. Notice that the performance benefit of RCM ordering actually increases as the number of cores increases, possible due to reduced communication costs. Our goal in this paper is therefore to design and develop a scalable distributed-memory parallel implementation of the RCM algorithm [5].

Similar to many sparse matrix computations, RCM ordering has been shown to be a difficult problem to parallelize [8]. The RCM algorithm relies on the repeated application of breadth-first search (BFS) and its computational load is highly dynamic, especially if the graph has high diameter. The problem exacerbates on high concurrency, where load imbalance and communication overhead degrade the performance of the parallel algorithm. RCM is harder to parallelize than BFS primarily for two reasons. First, RCM imposes additional restrictions on how the vertices are numbered in the graph traversal, limiting available parallelism. Second, RCM is often used for matrices with medium-to-high diameter while most of the existing work on parallel BFS has focused on low-diameter graphs such as synthetic graphs used by the Graph500 benchmark. A higher diameter increases the crit-

ical path of level-synchronous BFS algorithms, also limiting available parallelism.

In this paper, we aim to overcome these challenges by using the graph-matrix duality and replacing unstructured graph operations by structured matrix/vector operations. We make the following contributions:

- We present a scalable distributed-memory algorithm for RCM ordering. Our algorithm relies on a handful of bulksynchronous parallel primitives that are optimized for both shared-memory and massively parallel distributed memory systems.
- The quality (bandwidth and envelope) of ordering from our distributed-memory implementation is comparable to the state-of-the-art and remains insensitive to the degree of concurrency.
- We provide a hybrid OpenMP-MPI implementation of the RCM ordering that attains up to 38× speedup on matrices from various applications on 1024 cores of a Cray XC30 supercomputer. We provide detailed performance evaluation on up to 4096 cores, which sheds light on the performance bottlenecks and opportunities for future research.

II. PRELIMINARIES

A. Serial Algorithms

Let n be the number of columns in a symmetric matrix \mathbf{A} . Let $f_i(\mathbf{A})$ be the the column subscript of the first nonzero element in column i of \mathbf{A} : $f_i(\mathbf{A}) = \min\{j \mid a_{ij} \neq 0\}$. The i-th bandwidth $\beta_i(\mathbf{A})$ is defined as $\beta_i(\mathbf{A}) = i - f_i(\mathbf{A})$. The overall bandwidth of matrix \mathbf{A} is denoted $\beta(\mathbf{A})$ and is defined as $\beta(\mathbf{A}) = \max\{\beta_i(\mathbf{A}) \mid 1 \leq i \leq n\}$. Using these notations, the envelope $Env(\mathbf{A})$ is defined as:

$$Env(\mathbf{A}) = \{\{i, j\} \mid 0 < j - i < \beta_i(\mathbf{A}), 1 < i, j < n\}.$$

The quantity $|Env(\mathbf{A})|$ is called the profile or envelope size of \mathbf{A} .

Reverse Cuthill-McKee ordering, or RCM, introduced by George [9], is a variant of the Cuthill-McKee ordering [4], which aims at reducing the *bandwidth* of a sparse symmetric matrix \mathbf{A} . This is of particular importance when the matrix is to be stored using a profile based format. Finding a reordering of the rows/columns of \mathbf{A} corresponds to the process of labeling vertices of the graph $\mathcal{G}(\mathbf{A})$ associated with \mathbf{A} .

RCM repeatedly labels vertices adjacent to the current vertex v_i until all have been labeled, as depicted in Algorithm 1. The algorithm essentially processes vertices levels by levels and labels them in reverse order. The resulting reordered matrix often has a smaller profile [9].

For the graph $\mathcal{G}(\mathbf{A})=(V,E)$ associated with \mathbf{A},V is the set of vertices and E is the set of edges in $\mathcal{G}(\mathbf{A})$. The number of vertices in $\mathcal{G}(\mathbf{A})$ is denoted n=|V| and m is the number of edges m=|E|. The *eccentricity* [10] of a vertex v is defined as $\ell(v)=\max\{d(v,w)\mid w\in V\}$, where d(v,w) is the graph distance between v and w.

Algorithm 1: Reverse Cuthill-McKee algorithm

```
1 v_1 \leftarrow r

2 for i=1 to n do

3 Find all the unnumbered neighbors of the vertex v_i

4 Label the vertices found in increasing order of degree

5 The Reverse Cuthill-McKee ordering is given by w_1, w_2, \ldots, w_n where w_i = v_{n-i+1} for i=1, \ldots, n.
```

The rooted level structure [11] of a vertex $v \in V$ is the partioning $\mathcal{L}(v)$ of V satisfying

$$\mathcal{L}(v) = \{L_0(v), L_1(v), \dots, L_{\ell(v)}(v)\},\$$

where

$$L_0(v) = \{v\}, \quad L_1(v) = Adj(L_0(v)),$$

$$L_i(v) = Adj(L_{i-1}(v)) \setminus L_{i-2}(v), \quad i = 2, 3, \dots, \ell(v).$$

The *length* of $\mathcal{L}(v)$ corresponds to the eccentricity $\ell(v)$ while the *width* $\nu(v)$ of $\mathcal{L}(v)$ is defined by

$$\nu(v) = \max\{|L_i(v)| \mid 0 \le i \le \ell(v)\}.$$

The first vertex being labeled strongly impacts the bandwidth of the permuted matrix. Experience shows that it is better to start with a node having a large eccentricity. A peripheral vertex is a vertex with maximum eccentricity. Finding such vertex is prohibitively expensive, and a common heuristic is to use a pseudo-peripheral vertex instead [12], [5]. A pseudo-peripheral vertex is a vertex displaying a high eccentricity, as close to the graph diameter as possible. Gibbs et al. [13] introduced an algorithm to find such a vertex, which is later refined by George and Liu [12]. The process, given in Algorithm 2, starts with an arbitrary vertex in V and computes its rooted level structure. Then, a vertex in the last level is picked and the corresponding level structure is computed. This process is repeated until the number of levels in the rooted level structure converges. Computing the level structure corresponds to a BFS of the graph $\mathcal{G}(\mathbf{A})$.

Algorithm 2: An algorithm to find a pseudo-peripheral vertex r

```
1 r \leftarrow \{ \text{arbitrary vertex in } V \}
2 \mathcal{L}(r) \leftarrow \{ L_0(r), L_1(r), \dots, L_{\ell(r)}(r) \}
3 \text{nlvl} \leftarrow \ell(r) - 1
4 while \ell(r) > \text{nlvl} do
5 \text{nlvl} \leftarrow \ell(r)
6 (Shrink last level): Choose a vertex v in L_{\ell(r)}(r) of minimum degree.
7 \mathcal{L}(v) \leftarrow \{ L_0(v), L_1(v), \dots, L_{\ell(v)}(v) \}
8 \text{l} r \leftarrow v
```

TABLE I: Matrix-algebraic primitives needed for the RCM algorithm.

Function	Arguments	Returns	Serial Complexity	Communication	
IND	x: a sparse vector x : a sparse vector x : local indices of nonzero entries of x		O(nnz(x))	None	
SELECT	x: a sparse vector y : a dense vector $expr$: logical expr. on y assume $size(x) = size(y)$	$z \leftarrow$ an empty sparse vector for $i \in \text{IND}(x)$ if $(expr(y[i]))$ then $z[i] \leftarrow x[i]$	O(nnz(x))	None	
SET	x: a sparse vector y: a dense vector	for $i \in IND(x)$ $y[i] \leftarrow x[i]$	O(nnz(x))	None	
SPMSPV	A: a sparse matrix x: a sparse vector SR: a semiring	returns $\mathbf{A} \cdot x$	$\sum_{k \in \text{IND}(x)} nnz(\mathbf{A}(:,k))$	AllGather & AlltoAll on subcommunicator [14]	
REDUCE	x: a sparse vector y: a dense vector a reduction operation	$\begin{aligned} & \text{mv} = \text{maximum value in } y \\ & \text{for } i \in \text{IND}(x) \\ & \text{mv} \leftarrow \min\{\text{mv}, \ y[i]\} \end{aligned}$	O(nnz(x))	AllReduce	
SORTPERM	x: a sparse vector y: a dense vector	$T \leftarrow$ an empty array of tuples for $i \in \text{IND}(x)$ $T[i] \leftarrow (x[i], y[i], i)$ sort T and return the permutation	$O(nnz(x) \log nnz(x))$	AllToAll	

B. Previous Work on Parallel RCM

There is a strong connection between BFS, finding a pseudo-peripheral vertex and computing the RCM ordering. However, RCM is often used for matrices with higher diameter than graphs for which parallel BFS is often optimized [14], [15]. In addition, computing the actual RCM ordering is even harder to parallelize because the vertices within each level of the traversal tree has to be ordered by degree.

Computing sparse matrix orderings in parallel have received intermittent attention over the last few decades. While RCM is often used to accelerate iterative solvers, one of its first reported supercomputer-scale implementations was in the context of direct solvers by Ashcraft et al. [16] on a CRAY X-MP. It is hard to compare results from 30 years ago, where both the architectures and the sizes of matrices were significantly different. Karantasis et al. [8] recently studied the shared-memory parallelization of various reordering algorithms including RCM.

III. ALGORITHMS BASED ON MATRIX ALGEBRA

In this section, we present a distributed memory algorithm for computing the RCM ordering that takes advantage of the equivalence between graph algorithms and sparse matrix algebra to exploit latest hardware platforms. The nnz() function computes the number of nonzeros in its input, e.g., $nnz(\mathbf{x})$ returns the number of nonzeros in \mathbf{x} . We utilize the Matlab colon notation: $\mathbf{A}(:,i)$ denotes the ith column, and $\mathbf{A}(i,:)$ denotes the ith row.

A. Primitives for the RCM algorithm

With an aim to design a scalable parallel algorithm, we redesign the sequential algorithms for RCM ordering (Algorithm 1) and finding a pseudo-peripheral vertex (Algorithm 2) in terms of matrix-algebraic operations. For this purpose, we use the primitives summarized in Table I. A sparse vector is

used to represent a subset of vertices. Consider that a subset of n_1 unique vertices V_1 is represented by a sparse vector x. Then, for each vertex v_i in V_1 , there is a nonzero entry in the i-th location of x (i.e., $x[i] \neq 0$), and the number of nonzeros nnz(x) in x is equal to n_1 . The nonzero entries of the sparse vector can store arbitrary values depending on the context of the algorithm. By contrast, a dense vector y stores information about all vertices in the graph, hence the length of y is always n. In sparse matrix computations, we often are interested in where the nonzero elements are in the sparse matrices. Hence, if \mathbf{A} is an $n \times n$ sparse symmetric matrix, then we use a graph G = (V, E) to describe the sparsity structure of \mathbf{A} : $V = \{v_1, v_2, \cdots, v_n\}$, where v_i corresponds to row/column i of \mathbf{A} , for $1 \leq i \leq n$, and $\{v_i, v_i\} \in E$ when $\mathbf{A}_{i,j} \neq 0$.

Let x be a sparse vector, y be a dense vector, expr be a logical unary operation and A be a sparse matrix. IND(x) returns the indices of nonzero entries in x. SELECT(x, y, expr)returns every nonzero entry x[i] of x where expr(y[i]) is true. SET(y, x) replaces values of y by corresponding nonzero entries of x (other entries of y remain unchanged). RE-DUCE(x, y, op) considers only the nonzero indices I of the sparse vector x and performs a reduction on the values of the dense vector y[I] using the operation op. In Table I, we provide an example where the reduction operation is to find the minimum value. SORTPERM(x, y) creates a tuple (x[i], y[i], i) for each nonzero index i in x. The list of tuples are lexicographically sorted and the function returns the sorted permutation (the indices passed to the sorting routine are used to obtain the permutation). Finally, SPMSPV(\mathbf{A}, x, SR) performs a sparse matrix-sparse vector multiplication over the semiring SR. For the purposes of this work, a semiring is defined over (potentially separate) sets of 'scalars', and has its two operations 'multiplication' and 'addition' redefined. We refer to a semiring by listing its scaling operations, such as the

Algorithm 3: Reverse Cuthill-McKee algorithm in terms of matrix-algebraic operations. **Inputs:** a sparse adjacency matrix A, a dense vector \mathcal{D} storing the degrees of all vertices, and a pseudo-peripheral vertex r. **Output:** the RCM ordering \mathcal{R} .

```
1 \mathcal{R} \leftarrow -1
                           // Dense vector storing the ordering of all vertices; initialized to -1
2 \mathcal{L}_{cur} \leftarrow \{r\}, \mathcal{L}_{next} \leftarrow \phi
                                                                                  // Current and next BFS levels (frontier)
\mathbf{3} \ \mathcal{R}[r] \leftarrow 0
                                                                                                                 // label of r is set to 0
4 nv \leftarrow 1
                                                                                            // Number of vertices labeled so far
5 while \mathcal{L}_{cur} \neq \phi do
        \mathcal{L}_{cur} \leftarrow \text{Set}(\mathcal{L}_{cur}, \mathcal{R})
        \mathcal{L}_{next} \leftarrow \text{SPMSPV}(\mathbf{A}, \mathcal{L}_{cur}, \text{SR=(select2nd, min))}
                                                                                               // Visit neighbors of the frontier
        \mathcal{L}_{next} \leftarrow \text{Select}(\mathcal{L}_{next}, \mathcal{R} = -1)
                                                                                                               // Keep unvisited vertices
8
                                                          // Lexicographically sorted permutation of vertices in
        \mathcal{R}_{next} \leftarrow \text{SORTPERM}(\mathcal{L}_{next}, \mathcal{D})
         the next frontier based on the (parent_order, degree) pair
        \mathcal{R}_{next} \leftarrow \mathcal{R}_{next} + nv
                                                                                                                              // Global ordering
10
        nv \leftarrow nv + nnz(R_{next})
11
        \mathcal{R} \leftarrow \text{Set}(\mathcal{R}, \mathcal{R}_{next})
                                                                               // Set orders of the newly visited vertices
12
        \mathcal{L}_{cur} \leftarrow \mathcal{L}_{next}
13
14 return \mathcal{R} in the reverse order
```

Algorithm 4: Algorithm to find a pseudo-peripheral vertex r in terms of matrix-algebraic operations. **Inputs:** a sparse adjacency matrix \mathbf{A} and a dense vector \mathcal{D} storing the degrees of all vertices. **Output:** a pseudo-peripheral vertex r.

```
1 r \leftarrow \{\text{arbitrary vertex in } V\}
 2 \ell \leftarrow 0, \text{nlvl} \leftarrow \ell - 1
                                                                  // Number of levels in current and previous BFS trees
 3 while \ell > \text{nlvl do}
         \mathcal{L} \leftarrow -1
                                         // Dense vector storing the BFS level that each vertex belongs to
 4
         \mathcal{L}_{cur} \leftarrow \{r\}, \mathcal{L}_{next} \leftarrow \phi
                                                                                          // Current and next BFS levels (frontier)
5
         nlvl \leftarrow \ell, \mathcal{L}[r] \leftarrow 0
                                                                                                                           // Level of r is set to 0
6
7
              \mathcal{L}_{cur} \leftarrow \text{Set}(\mathcal{L}_{cur}, \mathcal{L})
8
              \mathcal{L}_{next} \leftarrow \text{SPMSPV}(\mathbf{A}, \mathcal{L}_{cur}, \text{SR=(select2nd, min))}
                                                                                                                                         // Visit neighbors
9
              \mathcal{L}_{next} \leftarrow Select(\mathcal{L}_{next}, \mathcal{L} = -1)
                                                                                                                         // Keep unvisited vertices
10
              if \mathcal{L}_{next} \neq \phi then
11
                  \mathcal{L} \leftarrow \text{Set}(\mathcal{L}, \mathcal{L}_{next})
                                                                                              // Set levels of newly visited vertices
12
                \mathcal{L}_{cur} \leftarrow \mathcal{L}_{next}
13
              \ell \leftarrow \ell + 1
14
         while \mathcal{L}_{next} \neq \phi
15
         r \leftarrow \text{Reduce}(\mathcal{L}_{cur}, \mathcal{D})
                                                                                        // Find the vertex with the minimum degree
16
```

(multiply, add) semiring. The usual semiring multiply for BFS is select2nd, which returns the second value it is passed. The BFS semiring is defined over two sets: the matrix elements are from the set of binary numbers, whereas the vector elements are from the set of integers.

B. The RCM algorithm using matrix-algebraic primitives

Algorithm 3 describes the RCM algorithm using the operations from Table I. Here, we assume that the graph is connected. The case for more than connected components can be handled by repeatedly invoking Algorithm 3 for each connected component. Algorithm 3 takes a sparse adjacency matrix $\bf A$ and a pseudo-peripheral vertex r as inputs, and returns the RCM ordering as a dense vector \mathcal{R} .

The *i*-th element $\mathcal{R}[i]$ of \mathcal{R} is initialized to -1 and it retains

the initial value until the *i*-th vertex is visited and labeled. Let \mathcal{L}_{cur} and \mathcal{L}_{next} be the set of vertices in the current and the next BFS level, respectively. \mathcal{L}_{cur} is called the BFS frontier (the set of current active vertices). The algorithm starts by labeling the pseudo-peripheral vertex r, inserting it into \mathcal{L}_{cur} .

The while loop (lines 5-13 of Algorithm 3) explores the vertices level-by-level until the frontier \mathcal{L}_{cur} becomes empty. Vertices in \mathcal{L}_{cur} have been already labeled in the previous iteration or during initialization. Hence each iteration of the while loop traverses the unvisited neighbors \mathcal{L}_{next} of \mathcal{L}_{cur} and labels vertices in \mathcal{L}_{next} .

At the beginning of the while loop of Algorithm 3, the values of the sparse vector \mathcal{L}_{cur} are set to the labels of vertices. Next, we discover vertices \mathcal{L}_{next} that are adjacent to the current frontier \mathcal{L}_{cur} by multiplying **A** by \mathcal{L}_{cur} over

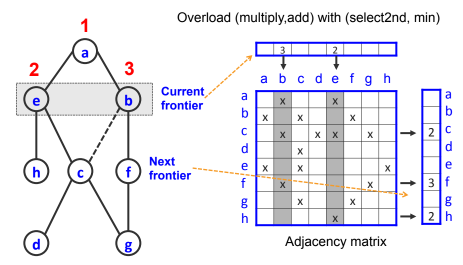


Fig. 2: An example of using specialized sparse matrix-sparse vector multiplication in exploring the next-level vertices in the RCM algorithm. On the left, we show a BFS tree rooted at vertex a. The current frontier is the set of two vertices $\{e, b\}$. Labels of the already explored vertices are shown in red numbers. The adjacency matrix representing the graph is shown on the right. The input sparse vector corresponding to the current frontier is show at the top of the matrix. The sparse vector has two nonzero entries corresponding to e and b. The values of the nonzero entries are the labels of the vertices. The SpMSpV algorithm over a (select2nd, min) semiring then selects the columns of the matrix corresponding to the nonzero indices of the vector shown in gray columns and retains the minimum product in each row where there exists at least one nonzero entry in the selected columns. The indices of the output vector represent the vertices in the next level while the values in the vector denote the labels of parent vertices. The (select2nd, min) semiring ensures that a child v always attaches itself to a parent with the minimum label among all of v is visited neighbors. Therefore, for vertex v, vertex v0 is selected as a parent instead of vertex v1 because the former has a smaller label than the latter.

(select2nd, min) semiring (line 7). Here the overloaded multiplication operation selec2nd passes the labels of parents to the children and the overloaded addition operation min ensures that each vertex in \mathcal{L}_{next} becomes a child of a vertex in \mathcal{L}_{cur} with the minimum label. This step is explained in Figure 2 with an example. Notice that the specific operator overloading by a (select2nd, min) semiring makes the vertex exploration deterministic, and parallelizing the specialized SpMSpV becomes more challenging than traditional BFS. Thankfully, the linear algebraic formulation with its operator overloading feature hides the RCM-specific complexity and is able to achieve high performance despite the deterministic nature of the computation.

After the vertices in the next level are discovered via SpMSpV, previously labeled vertices are removed from \mathcal{L}_{next} (line 8). The next step is to label the vertices in \mathcal{L}_{next} . For this purpose, we sort vertices in \mathcal{L}_{next} based on the labels of the parents of \mathcal{L}_{next} and the degrees of vertices \mathcal{L}_{next} . The sorted permutation of vertices in \mathcal{L}_{next} is used to set their labels. Finally, we update the labels of the newly visited vertices (line 12) and proceed to the next iteration of the while loop. When the current frontier becomes empty, Algorithm 3 returns with the RCM ordering by reversing the labels of the vertices.

C. Finding the pseudo-peripheral vertex

Algorithm 4 finds a pseudo-peripheral vertex using matrixalgebraic operations from Table I. Similar to Algorithm 2, Algorithm 4 starts with an arbitrary vertex r. Initially, the number of levels in the current and previous BFSs are set to 0 and -1, respectively, so that the while loop in line 4 iterates at least twice before termination. Each iteration of the while loop in line 4 runs a full BFS starting with r. As before, \mathcal{L}_{cur} and \mathcal{L}_{next} represent the subsets of vertices in the current and next levels of the BFS. \mathcal{L} stores the BFS level to which each vertex belongs (-1 denotes an unvisited vertex). The do-while loop (lines 8-16) performs the BFS traversal similar to Algorithm 3. The overloaded addition operation of the SPMSPV at line 12 is set to min only for convenience. It can be replaced by any equivalent operation because the order of vertices within a level of BFS does not matter in the discovery of a pseudoperipheral vertex. At the end of the BFS, we find a vertex with the minimum degree from the last non-empty level and make it the root for the next BFS (line 17). The while loop at line 4 terminates when the number of levels in the latest BFS is not greater than the previous BFS. At this point, Algorithm 4 returns the pseudo-peripheral vertex r.

IV. DISTRIBUTED MEMORY ALGORITHM

In order to parallelize the RCM algorithm, we simply need to parallelize the matrix-algebraic functions described in Table I.

A. Data distribution and storage

We use the Combinatorial BLAS (CombBLAS) framework [17], which distributes its sparse matrices on a 2D $p_r \times p_c$ processor grid. Processor P(i,j) stores the submatrix \mathbf{A}_{ij} of dimensions $(m/p_r) \times (n/p_c)$ in its local memory. Vectors are also distributed on the same 2D processor grid. CombBLAS supports different formats to store its local submatrices. In this work, we use the CSC format as we found it to be the fastest for the SpMSpV operation with very sparse vectors, which is often the case for matrices where RCM is commonly used. CombBLAS uses a vector of {index, value} pairs for storing sparse vectors. To balance load across processors, we randomly permute the input matrix \mathbf{A} before running the RCM algorithm.

B. Analysis of the distributed algorithm

We measure communication by the number of *words* moved (W) and the number of *messages* sent (S). The cost of communicating a length m message is $\alpha + \beta m$ where α is the latency and β is the inverse bandwidth, both defined relative to the cost of a single arithmetic operation. Hence, an algorithm that performs F arithmetic operations, sends S messages, and moves W words takes $T = F + \alpha S + \beta W$ time.

Since the amount of work is variable across BFS iterations, we analyze the aggregate cost over the whole BFS. For ease of analysis, matrix and vector nonzeros are assumed to be i.i.d. distributed. We also assume a square processor grid $p_c = p_r = \sqrt{p}$. Number of complete breadth-first searches is denoted by |iters|. The value of |iters| is exactly one in Algorithm 3, but it can be more than one in Algorithm 4.

We leverage the 2D SpMSpV algorithms implemented in CombBLAS. The complexity of parallel SpMSpV algorithm has been analyzed in this context recently [18], hence we just state the result here:

$$T_{\text{SPMSPV}} = O\Big(\frac{m}{p} + \beta \big(\frac{m}{p} + \frac{n}{\sqrt{p}}\big) + \left|\text{iters}\right| \alpha \sqrt{p}\Big).$$

As described before, the SORTPERM function returns the sorted permutation of the vertices in the next frontier \mathcal{L}_{next} based on the lexicographic order of (parent_label, degree, vertex_id), where parent_label is the label of the parent of a vertex in \mathcal{L}_{next} . We observe that parents of \mathcal{L}_{next} are a subset of the current frontier \mathcal{L}_{cur} and vertices in \mathcal{L}_{cur} are incrementally labeled in the previous iteration. Hence, the number of unique parent labels is less than or equal to $nnz(\mathcal{L}_{cur})$. This observation motivates us to design a distributed bucket sort algorithm where the i-th processor is responsible for sorting vertices whose parent's labels fall in the range

$$\left[\frac{nnz(\mathcal{L}_{cur})}{p}i, \frac{nnz(\mathcal{L}_{cur})}{p}(i+1)\right).$$

Hence, processors exchange tuples using AllToAll and the responsible processors sort tuples locally. After sorting, the vertex indices associated with the tuples work as the inverse permutation. Processors conduct another round of AllToAll (only the indices) to obtain the sorted permutation.

Name		Dimensions	BW (pre-RCM)
Description	Spy Plot	Nonzeros	BW (post-RCM)
•			Pseudo-diameter
nd24k 3D mesh problem		72K×72K 29M	68,114 10,294 14
Ldoor structural prob.		952K×952K 42.49M	686,979 9,259 178
Serena gas reservoir simulation	No. of the last of	1.39M×1.39M 64.1M	81,578 81,218 58
audikw_1 structural prob		943K×943K 78M	925,946 35,170 82
dielFilterV3real higher-order finite element		1.1M×1.1M 89.3M	1,036,475 23,813 84
Flan_1565 3D model of a steel flange		1.6M×1.6M 114M	20,702 20,600 199
Li7Nmax6 nuclear configuration interaction calculations		664K×664K 212M	663,498 490,000 7
Nm7 nuclear configuration interaction calculations		4M×4M 437M	4,073,382 3,692,599 5
nlpkkt240 Sym. indefinite KKT matrix		78M×78M 760M	14,169,841 361,755 243

Fig. 3: Structural information on the sparse matrices used in our experiments. All matrices, except two, are from the University of Florida sparse matrix collection [21]. Li7Nmax6 and Nm7 [22] are from nuclear configuration interaction calculations.

The total number of nonzeros sorted is exactly the sum of the frontier sizes over all iterations, which is O(n). Hence the per-process cost of SORTPERM is upper bounded by

$$T_{\text{SORTPERM}} = O\left(\frac{n\log n}{p} + \beta \frac{n}{p} + |\text{iters}| \alpha p\right),$$

using personalized all-to-all [19]. We found our specialized bucket sort to be faster than state-of-the-art general sorting libraries, such as HykSort [20].

V. RESULTS

A. Experimental Platform

We evaluated the performance of parallel RCM algorithm on Edison, a Cray XC30 supercomputer at NERSC. In Edison, nodes are interconnected with the Cray Aries network using a Dragonfly topology. Each compute node is equipped with 64 GB RAM and two 12-core 2.4 GHz Intel Ivy Bridge

TABLE II: The bandwidth and runtime of the shared-memory RCM implementation in SpMP. SpMP did not finish in 30 minutes for Nm7. On Nm7 and nlpkkt240 matrices, our distributed-memory implementation ran out of memory on a single node of Edison.

Graphs	SpMP				Distributed RCM		
	$_{ m BW}$	Runtime (sec)			Runtime (sec)		
		1t	6t	24t	1t	6t	24t
nd24k	10,608	0.26	0.06	0.03	1.45	0.38	0.12
ldoor	9,099	3.25	0.52	0.28	4.63	1.52	0.74
Serena	85,229	1.64	0.49	0.66	7.75	2.26	1.08
audikw_1	34,202	1.31	0.34	0.16	7.31	1.99	0.81
dielFilterV3real	25,436	1.99	0.73	0.46	8.63	2.37	0.95
Flan_1565	20,849	1.86	0.44	0.17	12.11	3.88	1.35
Li7Nmax6	443,991	4.62	1.48	0.87	20.28	4.91	2.85
Nm7	-	-	-	_	-	_	-
nlpkkt240	346,556	57.21	25.17	9.92	-	-	-

processors, each with 30 MB L3 cache. We used Cray's MPI implementation, which is based on MPICH2. We used OpenMP for intra-node multithreading and compiled the code with gcc 5.2.0 with -02 -fopenmp flags. In our experiments, we only used square process grids because rectangular grids are not supported in CombBLAS [17]. When p cores were allocated for an experiment, we created a $\sqrt{p/t} \times \sqrt{p/t}$ process grid, where t was the number of threads per process. In our hybrid OpenMP-MPI implementation, all MPI processes performed local computation followed by synchronized communication rounds. Local computation in every matrixalgebraic kernel was fully multithreaded using OpenMP. Only one thread in every process made MPI calls in the communication rounds.

B. Matrix Suite

The sparse matrix test suite used in our experiments are shown in Figure 3. These matrices came from a set of real applications, where either a sparse system Ax = b is solved or an eigenvalue problem $Ax = \lambda x$ is solved. The matrices were chosen to represent a variety of different structures and nonzero densities. Since RCM is only well defined on symmetric matrices, all matrices are symmetric.

In the last column of Figure 3, the original (pre-RCM) as well as final (post-RCM) bandwidth of the matrix are shown. In the majority of the cases, RCM effectively reduces the bandwidth. Serena and Flan_1565 seem to be the only two matrices where RCM was ineffective in that regard.

C. Shared-memory performance

Our implementation is fully multithreaded to take advantage of the shared-memory parallelism available within a node of modern supercomputers. Here, we compare the quality and runtime of our algorithm with the RCM implementation in SpMP (Sparse Matrix Pre-processing) by Park et al. [23], which is based on optimization from [24] and on the algorithm presented in [8]. The results from SpMP is shown in Table II. For four out of eight matrices where SpMP was able to compute RCM, the RCM ordering from our distributed-memory algorithm (shown in Figure 3) yields smaller bandwidths than SpMP. SpMP is faster than our implementation

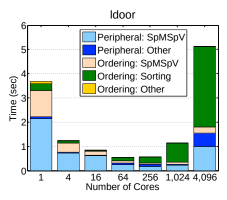


Fig. 6: Breakdown of total runtime when computing RCM ordering for ldoor using one thread per MPI process (flat MPI) on Edison.

in shared-memory due to our distributed-memory parallelization overheads. However, SpMP sometimes loses efficiency across NUMA domains. For example, SpMP slows down for Serena on 24 cores compared to 6 cores.

One very important thing to note is that in order to compute an ordering using a serial or multithreaded implementation of the RCM algorithm such as SpMP, the matrix structure has to be gathered on a single node. Indeed, in many real-life applications, the matrix is already distributed and this mandatory communication step has a non-negligible cost. For example, it takes over 9 seconds to gather the nlpkkt240 matrix from being distributed over 1024 cores into a single node/core. This time is approximately $3\times$ longer than computing RCM using our algorithm on the same number of cores. One of the key benefit of our approach is that it does not require this step. Adding those times to the time required to compute the ordering itself using SpMP makes our approach highly competitive and often faster.

D. Distributed-memory performance

We ran the distributed-memory RCM algorithm on up to 4096 cores of Edison. All performance results shown in this section used six threads per MPI process as it performed the best. Figure 4 shows the strong scaling of the distributed-memory RCM algorithm for nine graphs from Table 3. To better understand the performance, we break down the runtime into five parts at each concurrency where the height of a bar denotes the total runtime of identifying a pseudo-peripheral vertex and computing the RCM ordering.

Our distributed algorithm scales up to 1024 cores for all of the nine graphs as shown in Figure 4. The RCM algorithm attains the best speedup of $38\times$ for Li7Nmax6 and $27\times$ on nd24k on 1024 cores. By contrast, it achieves $10\times$ and $13\times$ speedups for ldoor and Flan_1565. The sharp drop in parallel efficiency on these graphs is due to their relatively high diameters. The level-synchronous nature of our BFS incurs high latency costs and decreases the amount of work per processor on high-diameter graphs. Figure 4 shows that SPMSPV is usually the most expensive operation on

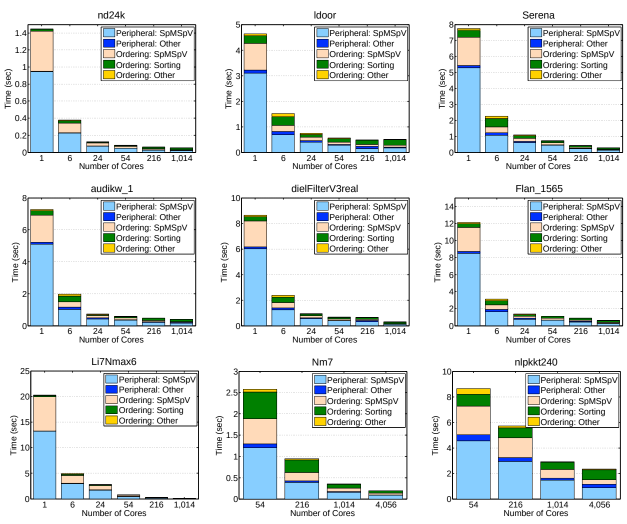


Fig. 4: Runtime breakdown of distributed-memory RCM on different graphs on Edison. Six threads per MPI process are used when the number of cores is greater than or equal to six. Nm7 and nlpkkt240 ran out of memory on a single node of Edison.

lower concurrency. However, SORTPERM starts to dominate on high concurrency because it performs an AllToAll among all processes, which has higher latency. The size of the matrix also contributes to the scalability of the distributed-memory RCM algorithm. For example, the largest two matrices in our test set (Nm7 and nlpkkt240) continue to scale on more than 4K cores whereas smaller problems do not scale beyond 1K cores. This demonstrates that our algorithm can scale on even higher core count if larger matrices are given as inputs.

Since the distributed-memory SPMSPV is the most expensive step of our RCM algorithm, we investigate its performance more closely. Figure 5 shows the breakdown of computation and communication time of the distributed SPMSPV primitive. We observe that on lower concurrency, SPMSPV is dominated by its computation, as expected. The communication time of SPMSPV starts to dominate the computation time on higher concurrency, and the crossover point where the communication becomes more expensive than computation depends on the properties of the matrices. Graphs with higher

diameters have higher communication overhead than graphs with low diameters. For example, ldoor has one of the highest pseudo-diameters among problems we considered. Hence, the RCM algorithm on ldoor becomes communication bound on 1K cores. By contrast, other lower diameter graphs with similar sizes continue to compute bound and scale beyond 1K cores on Edison.

We have also experimented with a flat MPI approach. This non-threaded RCM implementation had higher communication overhead and ran slower than the multithreaded implementation, especially on higher concurrencies. For example, the flat MPI implementation took $5\times$ longer to compute RCM on the ldoor matrix using 4096 cores of Edison as shown in Figure 6.

VI. CONCLUSION AND FUTURE WORK

We have introduced the first distributed-memory implementation of the RCM algorithm. In particular, we have described how the RCM algorithm can be reformulated using the sparse

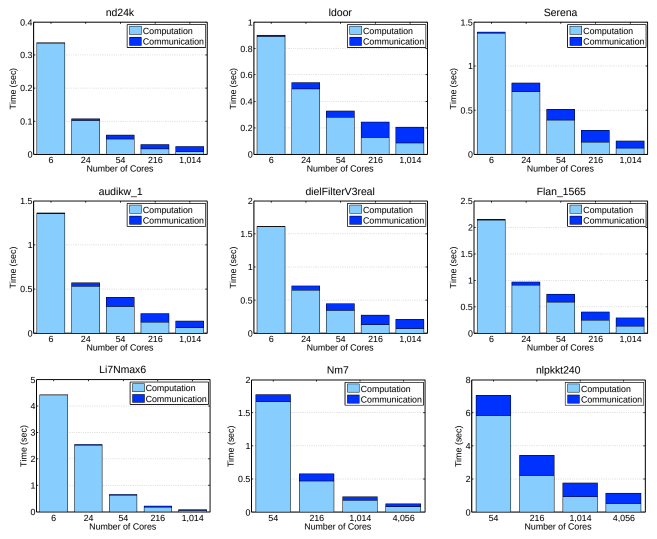


Fig. 5: Computation and communication time in all SpMSpV calls on different graphs on Edison. Six threads per MPI process are used.

matrix / graph duality, so that the parallel implementation can be accomplished using a small set of parallel primitives that have been highly optimized. Our experiments have shown that the distributed-memory implementation of the RCM algorithm presented in this work scales well up to 1024 processors for smaller matrices, and even to 4096 cores for the largest problems we have evaluated.

We have shown that this new approach is overall faster than the traditional approach that gathers the matrix structure on a single node, followed by the application of a serial or a multithreaded implementation of the RCM algorithm, and then redistributing the permuted matrix. More importantly, our approach removes the memory bottleneck that may be caused by having to store the entire graph with a single node.

A scalable implementation of the RCM algorithm is of prime importance for many iterative solvers that benefit from reordering the matrix. We have assessed this on a sample problem using the conjugate gradient method with block Jacobi preconditioner available in PETSc.

Immediate future work involves finding alternatives to sorting (i.e. global sorting at the end, or not sorting at all and sacrifice some quality). Longer term work would investigate alternative BFS formulations that are not level-synchronous, as the existing approach has trouble scaling to high-diameter graphs.

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