## Parallel Numerical Algorithms

Chapter 3 – Dense Linear Systems Section 3.2 – LU Factorization

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#### Outline

- LU Factorization
  - Motivation
  - Gaussian Elimination
- Parallel Algorithms for LU
  - Fine-Grain Algorithm
  - Agglomeration Schemes
  - Mapping Schemes
  - Scalability
- Partial Pivoting

#### LU Factorization

System of linear algebraic equations has form

$$Ax = b$$

where  ${\bf A}$  is given  $n \times n$  matrix,  ${\bf b}$  is given n-vector, and  ${\bf x}$  is unknown solution n-vector to be computed

 Direct method for solving general linear system is by computing LU factorization

$$A = LU$$

where L is unit lower triangular and U is upper triangular

#### LU Factorization

• System Ax = b then becomes

$$LUx = b$$

Solve lower triangular system

$$Ly = b$$

by forward-substitution to obtain vector y

Finally, solve upper triangular system

$$Ux = y$$

by back-substitution to obtain solution x to original system

## Gaussian Elimination Algorithm

LU factorization can be computed by Gaussian elimination as follows, where  $\boldsymbol{U}$  overwrites  $\boldsymbol{A}$ 

```
\begin{array}{ll} \text{for } k=1 \text{ to } n-1 & \{ \text{ loop over columns } \} \\ \text{for } i=k+1 \text{ to } n & \{ \text{ compute multipliers } \\ \ell_{ik}=a_{ik}/a_{kk} & \text{ for current column } \} \\ \text{end} & \\ \text{for } j=k+1 \text{ to } n & \{ \text{ apply transformation to } \\ a_{ij}=a_{ij}-\ell_{ik}a_{kj} & \text{ remaining submatrix } \} \\ \text{end} & \\ \text{end} & \\ \text{end} & \\ \text{end} & \\ \end{array}
```

#### Gaussian Elimination Algorithm

- In general, row interchanges (pivoting) may be required to ensure existence of LU factorization and numerical stability of Gaussian elimination algorithm, but for simplicity we temporarily ignore this issue
- Gaussian elimination requires about  $n^3/3$  paired additions and multiplications, so model serial time as

$$T_1 = \gamma \, n^3 / 3$$

where  $\gamma$  is time required for multiply-add operation

ullet About  $n^2/2$  divisions also required, but we ignore this lower-order term

# Loop Orderings for Gaussian Elimination

 Gaussian elimination has general form of triple-nested loop in which entries of L and U overwrite those of A

 Indices i, j, and k of for loops can be taken in any order, for total of 3! = 6 different ways of arranging loops

## Loop Orderings for Gaussian Elimination

- Different loop orders have different memory access patterns, which may cause their performance to vary widely
- Right-looking orderings (loop over k is outermost) perform updates to the trailing matrix (update all  $a_{ij}$  for  $i, j \geq k$ ) eagerly
- Left-looking orderings (loop over k is innermost) update the trailing matrix lazily (updates to  $a_{ij}$  done only when all entries  $a_{i'j'}$  with  $\min(i',j') < \min(i,j)$  have been updated)
- Right-looking ordering achieve better read-locality (the same divisor and outer-product vectors are reused)
- Left-looking ordering achieve better write-locality (entries
  of A may be changed in memory only once)

#### Gaussian Elimination Algorithm

Right-looking form of Gaussian elimination

```
\begin{array}{l} \text{for } k=1 \text{ to } n-1 \\ \text{ for } i=k+1 \text{ to } n \\ \ell_{ik}=a_{ik}/a_{kk} \\ \text{ end} \\ \text{ for } j=k+1 \text{ to } n \\ \text{ for } i=k+1 \text{ to } n \\ a_{ij}=a_{ij}-\ell_{ik}\,a_{kj} \\ \text{ end} \\ \text{ end} \\ \text{ end} \end{array}
```

• Multipliers  $\ell_{ik}$  computed outside inner loop for greater efficiency

#### Parallel Algorithm

#### **Partition**

• For i, j = 1, ..., n, fine-grain task (i, j) stores  $a_{ij}$  and computes and stores

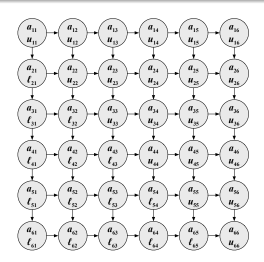
$$\left\{ \begin{array}{ll} u_{ij}, & \text{if } i \leq j \\ \ell_{ij}, & \text{if } i > j \end{array} \right.$$

yielding 2-D array of  $n^2$  fine-grain tasks

#### Communicate

- Broadcast entries of A vertically to tasks below
- Broadcast entries of L horizontally to tasks to right

#### Fine-Grain Tasks and Communication



## Fine-Grain Parallel Algorithm

```
for k = 1 to \min(i, j) - 1
    recv broadcast of a_{kj} from task (k, j)
                                                                { vert bcast }
    recv broadcast of \ell_{ik} from task (i, k)
                                                                { horiz bcast }
                                                                { update entry }
    a_{ij} = a_{ij} - \ell_{ik} a_{ki}
end
if i < j then
    broadcast a_{ij} to tasks (k, j), k = i + 1, \ldots, n
                                                                { vert bcast }
else
    recv broadcast of a_{ij} from task (j,j)
                                                                { vert bcast }
                                                                { multiplier }
    \ell_{ij} = a_{ij}/a_{jj}
    broadcast \ell_{ij} to tasks (i, k), k = j + 1, \ldots, n
                                                                { horiz bcast }
end
```

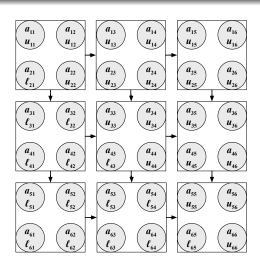
#### Agglomeration

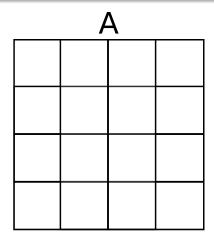
#### Agglomerate

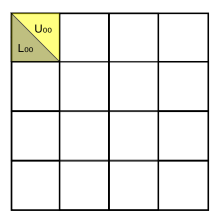
With  $n \times n$  array of fine-grain tasks, natural strategies are

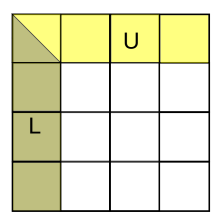
- 2-D: combine  $k \times k$  subarray of fine-grain tasks to form each coarse-grain task, yielding  $(n/k)^2$  coarse-grain tasks
- 1-D column: combine *n* fine-grain tasks in each column into coarse-grain task, yielding *n* coarse-grain tasks
- 1-D row: combine n fine-grain tasks in each row into coarse-grain task, yielding n coarse-grain tasks

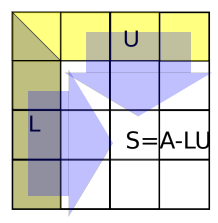
#### 2-D Agglomeration







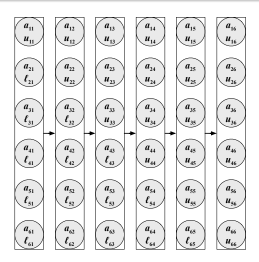




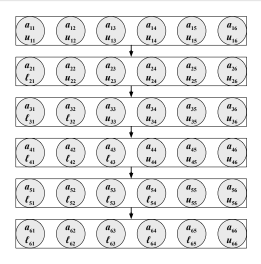
## Coarse-Grain 2-D Parallel Algorithm

```
for k=1 to n-1
    broadcast \{a_{kj}: j \in \textit{mycols}, j \geq k\} in processor column
    if k \in mycols then
        for i \in myrows, i > k
            \ell_{ik} = a_{ik}/a_{kk}
                                                     { multipliers }
        end
    end
    broadcast \{\ell_{ik}: i \in \textit{myrows}, i > k\} in processor row
    for j \in mycols, j > k
        for i \in myrows, i > k,
             a_{ij} = a_{ij} - \ell_{ik} \, a_{ki}
                                                       { update }
        end
    end
end
```

#### 1-D Column Agglomeration



#### 1-D Row Agglomeration

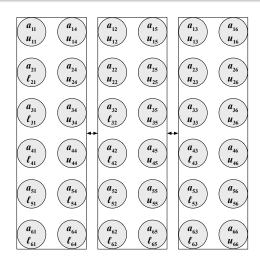


#### Mapping

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- 2-D: assign  $(n/k)^2/p$  coarse-grain tasks to each of p processors treating target network as 2-D mesh, using
  - blocked mapping (aggregating into larger blocks)
  - cyclic mapping of blocks, yielding block-cyclic layout
- 1-D: assign n/p coarse-grain tasks to each of p processors treating target network as 1-D mesh, using
  - blocked mapping (aggregating into panels)
  - cyclic mapping of rows/cols, yielding row-cyclic or column-cyclic layout

## 1-D Column Agglomeration with Cyclic Mapping



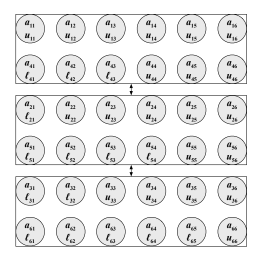
## 1-D Column Agglomeration

- Matrix rows need not be broadcast vertically, since any given column is contained entirely in only one process
- But there is no parallelism in computing multipliers or updating any given column
- Horizontal broadcasts still required to communicate multipliers for updating

# Coarse-Grain 1-D Column Parallel Algorithm

```
for k=1 to n-1
    if k \in mycols then
        for i = k + 1 to n
            \ell_{ik} = a_{ik}/a_{kk}
                                               { multipliers }
        end
    end
    broadcast \{\ell_{ik}: k < i \leq n\}
                                               { broadcast }
    for j \in mycols, j > k
        for i = k + 1 to n
                                               { update }
            a_{ij} = a_{ij} - \ell_{ik} a_{kj}
        end
    end
end
```

## 1-D Row Agglomeration with Cyclic Mapping

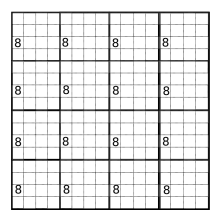


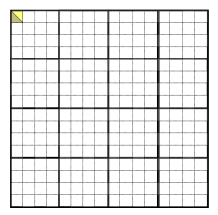
## 1-D Row Agglomeration

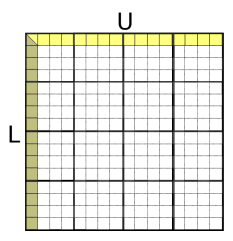
- Multipliers need not be broadcast horizontally, since any given matrix row is contained entirely in only one process
- But there is no parallelism in updating any given row
- Vertical broadcasts still required to communicate each row of matrix to processors below it for updating

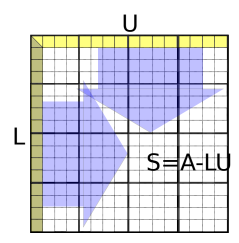
## Coarse-Grain 1-D Row Parallel Algorithm

```
for k=1 to n-1
    broadcast \{a_{kj}: k \leq j \leq n\}
                                                { broadcast }
    for i \in myrows, i > k,
        \ell_{ik} = a_{ik}/a_{kk}
                                                { multipliers }
    end
    for i = k + 1 to n
        for i \in myrows, i > k,
             a_{ij} = a_{ij} - \ell_{ik} a_{kj}
                                                { update }
        end
    end
end
```









#### Performance Enhancements

- Each processor becomes idle as soon as its last row and column are completed
- With block mapping, in which each processor holds contiguous block of rows and columns, some processors become idle long before overall computation is complete
- Block mapping also yields unbalanced load, as computing multipliers and updates requires successively less work with increasing row and column numbers
- Cyclic or reflection mapping improves both concurrency and load balance

#### Performance Enhancements

Performance can also be enhanced by overlapping communication and computation

- At step k, each processor completes updating its portion of remaining unreduced submatrix before moving on to step k+1
- Broadcast of each segment of row k+1, and computation and broadcast of each segment of multipliers for step k+1, could be initiated as soon as relevant segments of row k+1 and column k+1 have been updated by their owners, before completing remainder of their updating for step k
- This look-ahead strategy enables other processors to start working on next step earlier than they otherwise could

#### **Execution Time for 1-D Agglomeration**

- With 1-D column agglomeration, each processor factorizes panels of b columns, then broadcasts them to perform the trailing matrix update
- While work-efficient  $W_p = \Theta(n^3)$ , the concurrency in computational cost is constrained by panel factorization

$$F_p(n,b) = \Theta((n/b)nb^2 + n^3/p)$$

so we need b < n/p to maintain  $F_p(n,b) = \Theta(n^3/p)$ 

The overall execution time is given by

$$T_p(n,b) = \Theta\Big((n/b)T_p^{\text{bcast}}(nb) + \gamma F_p(n,b)\Big)$$

• It is generally minimized by picking  $b = \Theta(n/p)$ 

$$T_p(n, b) = \Theta(\alpha p \log p + \beta n^2 + \gamma n^3/p)$$

#### **Execution Time for 2-D Agglomeration**

- With 2-D agglomeration and block-cyclic mapping, a processor factorizes a  $b \times b$  diagonal block, broadcasts it to a column and row of processors, which update the panels and broadcast them to perform the trailing matrix updates
- The computational cost is constrained by lack of concurrency in the diagonal

$$F_p(n,b) = O(n^3/p + nb^2 + n^2b/\sqrt{p})$$

The overall execution time is given by

$$T_p(n,b) = \Theta\left((n/b)\left(T_{\sqrt{p}}^{\text{beast}}(b^2) + T_{\sqrt{p}}^{\text{beast}}(nb/\sqrt{p})\right) + \gamma F_p(n,b)\right)$$

• It is generally minimized by picking  $b = n/\sqrt{p}$ 

$$T_p(n) = T_p(n, n/\sqrt{p}) = \Theta(\alpha\sqrt{p}\log p + \beta n^2/\sqrt{p} + \gamma n^3/p)$$

#### Scalability for 2-D Agglomeration

- Cannon's algorithm for matrix multiplication (2-D agglomeration), could achieve strong scaling speed-up  $p_s = O((\gamma/\alpha)n^2)$  and unconditional weak scaling
- The SUMMA algorithm, which was based on broadcasts, achieved slightly inferior scaling due to a  $\Theta(\log(p))$  term on the latency cost
- The execution time of 2-D agglomeration for LU is the same as of SUMMA, so the efficiency and scaling characteristics are the same
- On the other hand, it is not possible to achieve strong scaling to  $O((\gamma/\alpha)n^3/\log(n))$  processors as the depth of the usual LU algorithm is D=n, meaning the maximum speed-up is  $p_s=\Theta(\max_p S_p)=O(Q_1/D)=O(n^2)$

## Partial Pivoting

- Row ordering of A is irrelevant in system of linear equations
- Partial pivoting takes rows in order of largest entry in magnitude of leading column of remaining unreduced matrix
- This choice ensures that multipliers do not exceed 1 in magnitude, which reduces amplification of rounding errors
- In general, partial pivoting is required to ensure existence and numerical stability of LU factorization

#### Partial Pivoting

Partial pivoting yields factorization of form

$$PA = LU$$

where P is permutation matrix

ullet If PA=LU, then system Ax=b becomes

$$PAx = LUx = Pb$$

which can be solved by forward-substitution in lower triangular system  $\boldsymbol{L}\boldsymbol{y}=\boldsymbol{P}\boldsymbol{b}$ , followed by back-substitution in upper triangular system  $\boldsymbol{U}\boldsymbol{x}=\boldsymbol{y}$ 

#### Parallel Partial Pivoting

- Partial pivoting complicates parallel implementation of Gaussian elimination and significantly affects potential performance
- With 2-D algorithm, pivot search is parallel but requires communication within processor column  $(S = \Omega(n \log(p)))$  and inhibits overlap
- With 1-D column algorithm, pivot search requires no communication but is purely serial
- Once pivot is found, index of pivot row must be communicated to other processors, and rows must be explicitly or implicitly interchanged in each process

#### Alternatives to Partial Pivoting

- Because of negative effects of partial pivoting on parallel performance, various alternatives have been proposed that limit pivot search
  - tournament pivoting (perform tree of partial pivoting on different subsets of matrix rows, selecting b at a time)
  - threshold pivoting (use local rows as pivots if the diagonal entries are within threshold of column norm)
  - pairwise pivoting (eliminate n(n-1)/2 entries by as many 2-by-2 transformations  $L_iP_i$ , where  $L_i$  is unit-lower triangular and  $P_i$  is a permutation matrix, applied to appropriate row pairs)
- Stability generally slightly worse in theory and for particularly hard test-cases
- Better stability without worrying about pivoting may be achieved via QR factorization

## Communication vs. Memory Tradeoff

- If explicit replication of storage is allowed, then lower communication volume is possible
- As with matrix multiplication, algorithms that leverage all available memory to reduce communication cost to the maximum extent possible
- If sufficient memory is avaiable, then these algorithms can achieve provably optimal communication

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