

- In theory, infinite number of iterations might be required to converge to exact solution
- In practice, iteration terminates when residual ||b Ax||, or some other measure of error, is as small as desired
- Iterative methods are especially useful when matrix *A* is sparse because, unlike direct methods, no fill is incurred



Jacobi Method

- Jacobi method requires nonzero diagonal entries, which can usually be accomplished by permuting rows and columns if not already true
- Jacobi method requires duplicate storage for *x*, since no component can be overwritten until all new values have been computed
- Components of new iterate do not depend on each other, so they can be computed simultaneously
- Jacobi method does not always converge, but it is guaranteed to converge under conditions that are often satisfied (e.g., if matrix is strictly diagonally dominant), though convergence rate may be very slow

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Gauss-Seidel Method

- Gauss-Seidel requires nonzero diagonal entries
- Gauss-Seidel does not require duplicate storage for *x*, since component values can be overwritten as they are computed
- But each component depends on previous ones, so they must be computed successively
- Gauss-Seidel does not always converge, but it is guaranteed to converge under conditions that are somewhat weaker than those for Jacobi method (e.g., if matrix is symmetric and positive definite)
- Gauss-Seidel converges about twice as fast as Jacobi, but may still be very slow

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Jacobi Method

Outline

 Beginning with initial guess x⁽⁰⁾, Jacobi method computes next iterate by solving for each component of x in terms of others

$$x_i^{(k+1)} = \left(b_i - \sum_{j \neq i} a_{ij} x_j^{(k)}\right) / a_{ii}, \quad i = 1, \dots, n$$

• If *D*, *L*, and *U* are diagonal, strict lower triangular, and strict upper triangular portions of *A*, then Jacobi method can be written

$$m{x}^{(k+1)} = m{D}^{-1} \left(m{b} - (m{L} + m{U}) m{x}^{(k)}
ight)$$

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Gauss-Seidel Method

- Faster convergence can be achieved by using each new component value as soon as it has been computed rather than waiting until next iteration
- This gives Gauss-Seidel method

$$x_i^{(k+1)} = \left(b_i - \sum_{j < i} a_{ij} x_j^{(k+1)} - \sum_{j > i} a_{ij} x_j^{(k)}\right) / a_{ii}$$

• Using same notation as for Jacobi, Gauss-Seidel method can be written

$$x^{(k+1)} = (D + L)^{-1} (b - Ux^{(k)})$$

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SOR Method

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- Successive over-relaxation (SOR) uses step to next Gauss-Seidel iterate as search direction with fixed search parameter ω
- SOR computes next iterate as

$$oldsymbol{x}^{(k+1)} = oldsymbol{x}^{(k)} + \omega \left(oldsymbol{x}^{(k+1)}_{GS} - oldsymbol{x}^{(k)}
ight)$$

where $x_{GS}^{(k+1)}$ is next iterate given by Gauss-Seidel

• Equivalently, next iterate is weighted average of current iterate and next Gauss-Seidel iterate

$$x^{(k+1)} = (1-\omega)x^{(k)} + \omega x_{GS}^{(k+1)}$$

• If A is symmetric, the SOR can be written as the application of a symmetric matrix; this is the Symmetric Succesive Over-Relaxation method

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SOR Method

- ω is fixed *relaxation* parameter chosen to accelerate convergence
- $\omega > 1$ gives *over-relaxation*, while $\omega < 1$ gives *under-relaxation* ($\omega = 1$ gives Gauss-Seidel method)
- SOR diverges unless 0 < ω < 2, but choosing optimal ω is difficult in general except for special classes of matrices
- With optimal value for ω, convergence rate of SOR method can be order of magnitude faster than that of Gauss-Seidel

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Conjugate Gradient Method

For special case of quadratic problem,

Negative gradient is residual vector

 $-\nabla \phi(\boldsymbol{x}) = \boldsymbol{b} - \boldsymbol{A}\boldsymbol{x} = \boldsymbol{r}$

Optimal line search parameter is given by

$$\alpha = \boldsymbol{r}_k^T \boldsymbol{s}_k / \boldsymbol{s}_k^T \boldsymbol{A} \boldsymbol{s}_k$$

- Successive search directions can easily be *A*-orthogonalized by three-term recurrence
- Using these properties, we obtain *conjugate gradient method* (*CG*) for linear systems

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Conjugate Gradient Method		

- Key features that make CG method effective
 - Short recurrence determines search directions that are *A*-orthogonal (conjugate)
 - Error is minimal over space spanned by search directions generated so far
- Minimum error property implies that method produces exact solution after at most *n* steps
- In practice, rounding error causes loss of orthogonality that spoils finite termination property, so method is used iteratively

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Preconditioning

- Convergence rate of CG can often be substantially accelerated by preconditioning
- Apply CG to $M^{-1}A$, where M is chosen so that $M^{-1}A$ is better conditioned than A, and systems of form Mz = y are easily solved

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- Typically, M is diagonal or triangular
- Types of preconditioners include
 - Diagonal or block-diagonal
 - SSOR
 - Incomplete factorization
 - Polynomial
 - Approximate inverse

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 $egin{aligned} eta_{k+1} = oldsymbol{r}_{k+1}^T oldsymbol{r}_{k+1} / oldsymbol{r}_k^T oldsymbol{r}_k \ oldsymbol{s}_{k+1} = oldsymbol{r}_{k+1} + eta_{k+1} oldsymbol{s}_k \end{aligned}$

 $\alpha_k = \boldsymbol{r}_k^T \boldsymbol{r}_k / \boldsymbol{s}_k^T \boldsymbol{A} \boldsymbol{s}_k$ $\boldsymbol{x}_{k+1} = \boldsymbol{x}_k + \alpha_k \boldsymbol{s}_k$ $\boldsymbol{r}_{k+1} = \boldsymbol{r}_k - \alpha_k \boldsymbol{A} \boldsymbol{s}_k$

 $m{x}_0 = {\sf initial guess}$ $m{r}_0 = m{b} - m{A}m{x}_0$

 $s_0 = r_0$ for k = 0, 1, 2, ...

Conjugate Gradient Method

quadratic function

end

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• If A is $n \times n$ symmetric positive definite matrix, then

attains minimum precisely when Ax = b

• For method of *steepest descent*, $s_k = -
abla \phi(x)$

Optimization methods have form

function $\phi(\boldsymbol{x}_k + \alpha \, \boldsymbol{s}_k)$ along \boldsymbol{s}_k

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Conjugate Gradient Method

 $\phi(x) = \frac{1}{2} \boldsymbol{x}^T \boldsymbol{A} \boldsymbol{x} - \boldsymbol{x}^T \boldsymbol{b}$

 $\boldsymbol{x}_{k+1} = \boldsymbol{x}_k + \alpha \, \boldsymbol{s}_k$

where α is search parameter chosen to minimize objective

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Conjugate Gradient Method

• Error is reduced at each iteration by factor of

$$(\sqrt{\kappa}-1)/(\sqrt{\kappa}+1)$$

on average, where

$$\kappa = \operatorname{cond}(\boldsymbol{A}) = \|\boldsymbol{A}\| \cdot \|\boldsymbol{A}^{-1}\| = \lambda_{\max}(\boldsymbol{A}) / \lambda_{\min}(\boldsymbol{A})$$

- Thus, convergence tends to be rapid if matrix is well-conditioned, but can be arbitrarily slow if matrix is ill-conditioned
- But convergence also depends on clustering of eigenvalues of *A*

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- CG is not directly applicable to nonsymmetric or indefinite systems
- CG cannot be generalized to nonsymmetric systems without sacrificing one of its two key properties (short recurrence and minimum error)
- Nevertheless, several generalizations have been developed for solving nonsymmetric systems, including GMRES, QMR, CGS, BiCG, and Bi-CGSTAB
- These tend to be less robust and require more storage than CG, but they can still be very useful for solving large nonsymmetric systems

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Parallel Implementation

- Iterative methods for linear systems are composed of basic operations such as
 - vector updates (saxpy)
 - inner products
 - matrix-vector multiplication
 - solution of triangular systems
- In parallel implementation, both data and operations are partitioned across multiple tasks
- In addition to communication required for these basic operations, necessary convergence test may require additional communication (e.g., sum or max reduction)

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- Sparse matrix A can be partitioned among tasks by rows, by columns, or by submatrices
- Partitioning by submatrices may give uneven distribution of nonzeros among tasks; indeed, some submatrices may contain no nonzeros at all
- Partitioning by rows or by columns tends to yield more uniform distribution because sparse matrices typically have about same number of nonzeros in each row or column



- Desired data locality can be achieved by partitioning graph of matrix, or partitioning underlying grid or mesh for finite difference or finite element problem
- For example, graph can be partitioned into p pieces by nested dissection, and vector components corresponding to nodes in each resulting piece assigned to same task, with neighboring pieces assigned to neighboring tasks
- Then matrix-vector product requires relatively little communication, and only between neighboring tasks

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- Chaco
- Jostle
- Meshpart
- Metis/ParMetis
- Mondriaan
- Party
- Scotch
- Zoltan



- Iterative methods typically require several vectors, including solution x, right-hand side b, residual r = b Ax, and possibly others
- Even when matrix *A* is sparse, these vectors are usually dense
- These dense vectors are typically uniformly partitioned among *p* tasks, with given task holding same set of component indices of each vector
- Thus, vector updates require no communication, whereas inner products of vectors require reductions across tasks, at cost we have already seen

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Row Partitioning of Sparse Matrix

- Suppose that each task is assigned n/p rows, yielding p tasks, where for simplicity we assume that p divides n
- In dense matrix-vector multiplication, since each task owns only n/p components of vector operand, communication is required to obtain remaining components
- If matrix is sparse, however, few components may actually be needed, and these should preferably be stored in neighboring tasks
- Assignment of rows to tasks by contiguous blocks or cyclically would not, in general, result in desired proximity of vector components

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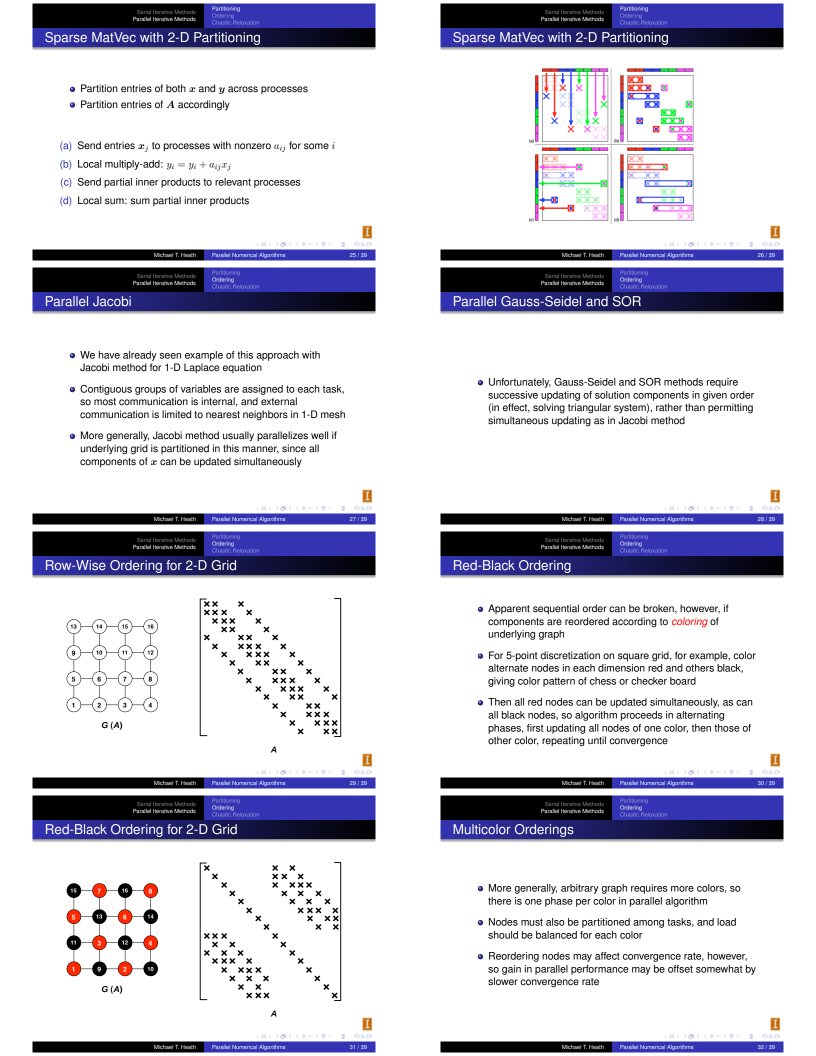
Graph Partitioning Methods

- Combinatorial refinement (e.g., Kernighan-Lin)
- Level structure
- Coordinate bisection
- Inertial
- Spectral
- Geometric
- Multilevel

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Two-Dimensional Partitioning			

$\begin{bmatrix} 1 & 6 & 0 & 0 & 0 & 0 & 0 & 0 \\ 5 & 1 & 9 & 0 & 5 & 0 & 0 & 0 \\ 8 & 0 & 1 & 7 & 0 & 0 & 0 & 0 \\ 0 & 0 & 2 & 1 & 0 & 0 & 0 & 7 \\ 0 & 0 & 0 & 0 & 1 & 8 & 0 & 0 \\ 0 & 4 & 0 & 0 & 3 & 1 & 3 & 0 \\ \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \\ x_4 \\ x_5 \\ x_6 \end{bmatrix}$	×××× ×××××××××××××××××××××××××××××××××

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 More generally, multicolor ordering of graph of matrix enhances parallel performance of sparse triangular solution by identifying sets of solution components that can be computed simultaneously (rather than in usual sequential order for triangular solution)

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Asynchronous or Chaotic Relaxation

Parallel Iterative Methods

- Using updated values for solution components in Gauss-Seidel and SOR methods improves convergence rate, but limits parallelism and requires synchronization
- Alternatively, in computing next iterate, each processor could use most recent value it has for each solution component, rather than waiting for latest value on any processor
- This approach, sometimes called *anynchronous* or *chaotic* relaxation, can be effective, but stochastic behavior complicates analysis of convergence and convergence rate

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