The Landscape of Sparse Ax=b Solvers

 $\begin{array}{ll} \underline{\text{Direct}} & \underline{\text{Iterative}} \\ A = LU & y' = Ay \end{array}$

Nonsymmetric

Symmetric positive definite

Pivoting GMRES, BiCGSTAB, ...

Cholesky Conjugate gradient

More General

More Robust

More Robust



Less Storage

$$x_0 = 0$$
 approx solution $r_0 = b$ residual = b - Ax

$$r_0 = b$$
 residual = b - Ax

$$d_0 = r_0$$
 search direction

for
$$k = 1, 2, 3, \dots$$

$$X_k = X_{k-1} + ...$$

$$r_k = \dots$$

new approx solution

new residual

$$d_k = \dots$$

new search direction

$$x_0 = 0$$
 approx solution

$$r_0 = b$$
 residual = b - Ax

$$d_0 = r_0$$
 search direction

for
$$k = 1, 2, 3, \dots$$

$$\alpha_k = \dots$$

$$x_k = x_{k-1} + \alpha_k d_{k-1}$$

$$r_k = \dots$$

step length

new approx solution

new residual

$$d_k = \dots$$

new search direction

Conjugate gradient iteration

$$\begin{array}{lll} x_0 = 0, & r_0 = b, & d_0 = r_0 \\ \hline \textbf{for} & k = 1, 2, 3, \dots \\ & \alpha_k = \left(r^T_{k-1}r_{k-1}\right) / \left(d^T_{k-1}Ad_{k-1}\right) & \text{step length} \\ & x_k = x_{k-1} + \alpha_k \, d_{k-1} & \text{approx solution} \\ & r_k = r_{k-1} - \alpha_k Ad_{k-1} & \text{residual} \\ & \beta_k = \left(r^T_k \, r_k\right) / \left(r^T_{k-1}r_{k-1}\right) & \text{improvement} \\ & d_k = r_k + \beta_k \, d_{k-1} & \text{search direction} \end{array}$$

- One matrix-vector multiplication per iteration
- Two vector dot products per iteration
- Four n-vectors of working storage

Conjugate gradient: Orthogonal sequences

- Krylov subspace: $K_t(A, b) = \text{span}(b, Ab, A^2b, \dots, A^{t-1}b)$
- Conjugate gradient algorithm:

for
$$t = 1, 2, 3, ...$$

find $x_t \in K_t(A, b)$
such that $r_t = (b - Ax_t) \perp K_t(A, b)$

- Notice $r_t \in K_{t+1}(A, b)$, so $r_t \perp r_k$ for all k < t
- Similarly, the "directions" are A-orthogonal:

$$(x_t - x_{t-1})^T \cdot A \cdot (x_k - x_{k-1}) = 0$$

The magic: Short recurrences. . .

A is symmetric => can get next residual and direction from the previous one, without saving them all.

Conjugate gradient: Convergence

- In exact arithmetic, CG converges in n steps (completely unrealistic!!)
- Accuracy after t steps of CG is related to:
 - consider polynomials of degree t that are equal to 1 at 0.
 - how small can such a polynomial be at all the eigenvalues of A?
- Thus, eigenvalues close together are good.
- Condition number: $\kappa(A) = \|A\|_2 \|A^{-1}\|_2 = \lambda_{\max}(A) / \lambda_{\min}(A)$
- Residual is reduced by a constant factor by $O(\varkappa^{1/2}(A))$ iterations of CG.

Preconditioners

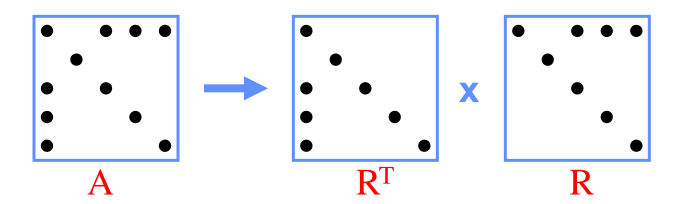
- Suppose you had a matrix B such that:
 - 1. condition number $\kappa(B^{-1}A)$ is small
 - 2. By = z is easy to solve
- Then you could solve (B⁻¹A)x = B⁻¹b instead of Ax = b
 - Actually $(B^{-1/2}AB^{-1/2})$ $(B^{1/2}x) = B^{-1/2}b$, but never mind...
- B = A is great for (1), not for (2)
- B = I is great for (2), not for (1)
- Domain-specific approximations sometimes work
- B = diagonal of A sometimes works
- Better: blend in some direct-methods ideas. . .

Preconditioned conjugate gradient iteration

$$\begin{aligned} x_0 &= 0, \quad r_0 = b, \quad d_0 = B^{\text{-1}} r_0, \quad y_0 = B^{\text{-1}} r_0 \\ \underline{\text{for}} \quad k &= 1, 2, 3, \dots \\ \alpha_k &= \left(y^T_{k\text{-1}} r_{k\text{-1}}\right) / \left(d^T_{k\text{-1}} A d_{k\text{-1}}\right) \quad \text{step length} \\ x_k &= x_{k\text{-1}} + \alpha_k \; d_{k\text{-1}} & \text{approx solution} \\ r_k &= r_{k\text{-1}} - \alpha_k \, A d_{k\text{-1}} & \text{residual} \\ y_k &= B^{\text{-1}} r_k & \text{preconditioning solve} \\ \beta_k &= \left(y^T_k r_k\right) / \left(y^T_{k\text{-1}} r_{k\text{-1}}\right) & \text{improvement} \\ d_k &= y_k + \beta_k \; d_{k\text{-1}} & \text{search direction} \end{aligned}$$

- One matrix-vector multiplication per iteration
- One solve with preconditioner per iteration

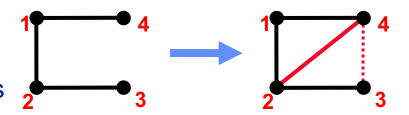
Incomplete Cholesky factorization (IC, ILU)



- Compute factors of A by Gaussian elimination, but ignore fill
- Preconditioner B = $R^TR \approx A$, not formed explicitly
- Compute B⁻¹z by triangular solves (in time nnz(A))
- Total storage is O(nnz(A)), static data structure
- Either symmetric (IC) or nonsymmetric (ILU)

Incomplete Cholesky and ILU: Variants

- Allow one or more "levels of fill"
 - unpredictable storage requirements



- Allow fill whose magnitude exceeds a "drop tolerance"
 - may get better approximate factors than levels of fill
 - unpredictable storage requirements
 - choice of tolerance is ad hoc
- Partial pivoting (for nonsymmetric A)
- "Modified ILU" (MIC): Add dropped fill to diagonal of U or R
 - A and R^TR have same row sums
 - good in some PDE contexts

Incomplete Cholesky and ILU: Issues

Choice of parameters

- good: smooth transition from iterative to direct methods
- bad: very ad hoc, problem-dependent
- tradeoff: time per iteration (more fill => more time)
 vs # of iterations (more fill => fewer iters)

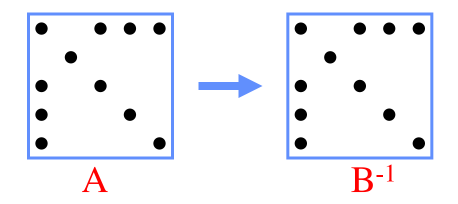
Effectiveness

- condition number usually improves (only) by constant factor (except MIC for some problems from PDEs)
- still, often good when tuned for a particular class of problems

Parallelism

- Triangular solves are not very parallel
- Reordering for parallel triangular solve by graph coloring

Sparse approximate inverses



- Compute $B^{-1} \approx A$ explicitly
- Minimize $\|\mathbf{A}\mathbf{B}^{-1} \mathbf{I}\|_{F}$ (in parallel, by columns)
- Variants: factored form of B⁻¹, more fill, . .
- Good: very parallel, seldom breaks down
- Bad: effectiveness varies widely

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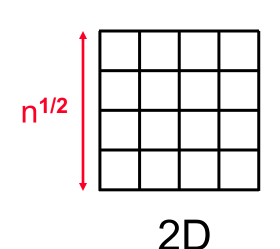
More Robust

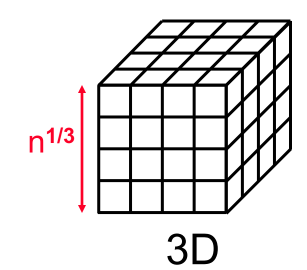


Less Storage

Complexity of linear solvers

Time to solve model problem (Poisson's equation) on regular mesh





Dense Cholesky:	O(n ³)	O(n ³)
Sparse Cholesky:	O(n ^{1.5})	O(n ²)
CG, exact arithmetic:	O(n²)	O(n²)
CG, no precond:	O(n ^{1.5})	O(n ^{1.33})
CG, modified IC:	O(n ^{1.25})	O(n ^{1.17})
Low-stretch trees:	O~(n)	O~(n)
Multigrid:	O(n)	O(n)

Hierarchy of matrix classes (all real)

- General nonsymmetric
- Diagonalizable
- Normal
- Symmetric indefinite
- Symmetric positive (semi)definite = Factor width n
 - Factor width 2 < k < n
- Diagonally dominant SPSD = Factor width 2
- Generalized Laplacian = Symm diag dominant M-matrix
- Graph Laplacian

Other Krylov subspace methods

- Nonsymmetric linear systems:
 - GMRES:

```
for i = 1, 2, 3, ...

find x_i \in K_i(A, b) such that r_i = (Ax_i - b) \perp K_i(A, b)

But, no short recurrence => save old vectors => lots more space

(Usually "restarted" every k iterations to use less space.)
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- BiCGStab, QMR, etc.:
 Two spaces K_i (A, b) and K_i (A^T, b) w/ mutually orthogonal bases
 Short recurrences => O(n) space, but less robust
- Convergence and preconditioning more delicate than CG
- Active area of current research
- Eigenvalues: Lanczos (symmetric), Arnoldi (nonsymmetric)