# Lecture 17: <br> More Fun With Sparse Matrices 

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## Logistics

- Thanks for info on final project ideas.
- HW 2 due Monday!


## Life lessons from HW 2?

- Where an error occurs may not be where you observe it!
- Check against a slow, naive, obvious calculation.
- assert is your friend.
- Use version control (git, cvs, svn, ...).


## Reminder: Conjugate Gradients

What if we only know how to multiply by $A$ ?
About all you can do is keep multiplying!

$$
\mathcal{K}_{k}(A, b)=\operatorname{span}\left\{b, A b, A^{2} b, \ldots, A^{k-1} b\right\} .
$$

Gives surprisingly useful information!
If $A$ is symmetric and positive definite, $x=A^{-1} b$ minimizes

$$
\begin{aligned}
\phi(x) & =\frac{1}{2} x^{\top} A x-x^{\top} b \\
\nabla \phi(x) & =A x-b .
\end{aligned}
$$

Idea: Minimize $\phi(x)$ over $\mathcal{K}_{k}(A, b)$.
Basis for the method of conjugate gradients

## Convergence of CG

- KSPs are not stationary (no constant fixed-point iteration)
- Convergence is surprisingly subtle!
- CG convergence upper bound via condition number
- Large condition number iff form $\phi(x)$ has long narrow bowl
- Usually happens for Poisson and related problems
- Preconditioned problem $M^{-1} A x=M^{-1} b$ converges faster?
- Whence $M$ ?
- From a stationary method?
- From a simpler/coarser discretization?
- From approximate factorization?


## PCG

```
Compute \(r^{(0)}=b-A x\)
for \(i=1,2, \ldots\)
    solve \(M z^{(i-1)}=r^{(i-1)}\)
    \(\rho_{i-1}=\left(r^{(i-1)}\right)^{T} z^{(i-1)}\)
    if \(i==1\)
        \(p^{(1)}=z^{(0)}\)
    else
        \(\beta_{i-1}=\rho_{i-1} / \rho_{i-2}\)
        \(p^{(i)}=z^{(i-1)}+\beta_{i-1} p^{(i-1)}\)
    endif
    \(q^{(i)}=A p^{(i)}\)
    \(\alpha_{i}=\rho_{i-1} /\left(p^{(i)}\right)^{T} q^{(i)}\)
    \(x^{(i)}=x^{(i-1)}+\alpha_{i} p^{(i)}\)
    \(r^{(i)}=r^{(i-1)}-\alpha_{i} q^{(i)}\)
end
```


## Parallel work:

- Solve with $M$
- Product with $A$
- Dot products
- Axpys

Overlap comm/comp.

## PCG bottlenecks

Key: fast solve with $M$, product with $A$

- Some preconditioners parallelize better! (Jacobi vs Gauss-Seidel)
- Balance speed with performance.
- Speed for set up of $M$ ?
- Speed to apply $M$ after setup?
- Cheaper to do two multiplies/solves at once...
- Can't exploit in obvious way - lose stability
- Variants allow multiple products - Hoemmen's thesis
- Lots of fiddling possible with $M$; what about matvec with $A$ ?


## Thinking on (basic) CG convergence



Consider 2D Poisson with 5-point stencil on an $n \times n$ mesh.

- Information moves one grid cell per matvec.
- Cost per matvec is $O\left(n^{2}\right)$.
- At least $O\left(n^{3}\right)$ work to get information across mesh!


## CG convergence: a counting approach

- Time to converge $\geq$ time to propagate info across mesh
- For a 2D mesh: $O(n)$ matvecs, $O\left(n^{3}\right)=O\left(N^{3 / 2}\right)$ cost
- For a 3D mesh: $O(n)$ matvecs, $O\left(n^{4}\right)=O\left(N^{4 / 3}\right)$ cost
- "Long" meshes yield slow convergence
- 3D beats 2D because everything is closer!
- Advice: sparse direct for 2D, CG for 3D.
- Better advice: use a preconditioner!


## CG convergence: an eigenvalue approach

Define the condition number for $\kappa(L)$ s.p.d:

$$
\kappa(L)=\frac{\lambda_{\max }(L)}{\lambda_{\min }(L)}
$$

Describes how elongated the level surfaces of $\phi$ are.

- For Poisson, $\kappa(L)=O\left(h^{-2}\right)$
- CG steps to reduce error by $1 / 2=O(\sqrt{\kappa})=O\left(h^{-1}\right)$.

Similar back-of-the-envelope estimates for some other PDEs. But these are not always that useful... can be pessimistic if there are only a few extreme eigenvalues.

## CG convergence: a frequency-domain approach



Error $e_{k}$ after $k$ steps of CG gets smoother!

## Choosing preconditioners for 2D Poisson

- CG already handles high-frequency error
- Want something to deal with lower frequency!
- Jacobi useless
- Doesn't even change Krylov subspace!
- Better idea: block Jacobi?
- Q: How should things split up?
- A: Minimize blocks across domain.
- Compatible with minimizing communication!


## Restrictive Additive Schwartz (RAS)



## Restrictive Additive Schwartz (RAS)



- Get ghost cell data
- Solve everything local (including neighbor data)
- Update local values for next step
- Default strategy in PETSc


## Multilevel Ideas

- RAS propogates information by one processor per step
- For scalability, still need to get around this!
- Basic idea: use multiple grids
- Fine grid gives lots of work, kills high-freq error
- Coarse grid cheaply gets info across mesh, kills low freq

More on this another time.

## CG performance

Two ways to get better performance from CG:

1. Better preconditioner

- Improves asymptotic complexity?
- ... but application dependent

2. Tuned implementation

- Improves constant in big-O
- ... but application independent?

Benchmark idea (?): no preconditioner, just tune.

## Tuning PCG

```
Compute \(r^{(0)}=b-A x\)
for \(i=1,2, \ldots\)
    solve \(M z^{(i-1)}=r^{(i-1)}\)
    \(\rho_{i-1}=\left(r^{(i-1)}\right)^{T} Z^{(i-1)}\)
    if \(i==1\)
        \(p^{(1)}=z^{(0)}\)
    else
        \(\beta_{i-1}=\rho_{i-1} / \rho_{i-2}\)
        \(p^{(i)}=z^{(i-1)}+\beta_{i-1} p^{(i-1)}\)
    endif
    \(q^{(i)}=A p^{(i)}\)
    \(\alpha_{i}=\rho_{i-1} /\left(p^{(i)}\right)^{T} q^{(i)}\)
    \(x^{(i)}=x^{(i-1)}+\alpha_{i} p^{(i)}\)
    \(r^{(i)}=r^{(i-1)}-\alpha_{i} q^{(i)}\)
end
```

- Most work in $A, M$
- Vector ops synchronize
- Overlap comm, comp?


## Tuning PCG

Compute $r^{(0)}=b-A x$

$$
p_{-1}=0 ; \beta_{-1}=0 ; \alpha_{-1}=0
$$

$$
s=L^{-1} r^{(0)}
$$

$$
\rho_{0}=s^{T} s
$$

$$
\text { for } i=0,1,2, \ldots
$$

Check convergence ( $\left\|r_{i+1}\right\|$ )
$\beta_{i}=\rho_{i+1} / \rho_{i}$
end
Split $z=M^{-1} r$ into $s, w_{i}$
Overlap

$$
w_{i}=L^{-T} s
$$

- $p_{i}^{T} q_{i}$ with $x$ update

$$
p_{i}=w_{i}+\beta_{i-1} p_{i-1}
$$

$$
q_{i}=A p_{i}
$$

- $s^{T} s$ with $w_{i}$ eval

$$
\gamma=p_{i}^{\top} q_{i}
$$

- Computing $p_{i}, q_{i}, \gamma$

$$
x_{i}=x_{i-1}+\alpha_{i-1} p_{i-1}
$$

- Pipeline $r_{i+1}, s$ ?

$$
\alpha_{i}=\rho_{i} / \gamma_{i}
$$

- Pipeline $p_{i}, w_{i}$ ?

$$
r_{i+1}=r_{i}-\alpha q_{i}
$$

$$
s=L^{-1} r_{i+1}
$$

$$
\rho_{i+1}=s^{T} s
$$

Parallel Numerical LA,
Demmel, Heath, van der Vorst

## Tuning PCG

Can also tune

- Preconditioner solve (hooray!)
- Matrix multiply
- Represented implicitly (regular grids)
- Or explicitly (e.g. compressed sparse column)

Or further rearrange algorithm (Hoemmen, Demmel).

## Tuning sparse matvec

- Sparse matrix blocking and reordering (Im, Vuduc, Yelick)
- Packages: Sparsity (Im), OSKI (Vuduc)
- Available as PETSc extension
- Optimizing stencil operations (Datta)


## Reminder: Compressed sparse row storage



for $i=1: n$

$$
y[i]=0 ;
$$

$$
\text { for } j j=\operatorname{ptr}[i] \text { to ptr[i+1]-1 }
$$

$$
y[i]+=A[j j] * x[\operatorname{col}[j]] ;
$$

end
end
Problem: y[i] += A[jj]*x[col[j]];

## Memory traffic in CSR multiply

Memory access patterns:

- Elements of $y$ accessed sequentially
- Elements of $A$ accessed sequentially
- Access to $x$ are all over!

Can help by switching to block CSR.
Switching to single precision, short indices can help memory traffic, too!

## Parallelizing matvec



- Each processor gets a piece
- Many partitioning strategies
- Idea: re-order so one of these strategies is "good"


## Reordering for matvec

SpMV performance goals:

- Balance load?
- Balance storage?
- Minimize communication?
- Good cache re-use?

Also reorder for

- Stability of Gauss elimination,
- Fill reduction in Gaussian elimination,
- Improved performance of preconditioners...


## Reminder: Sparsity and partitioning



Want to partition sparse graphs so that

- Subgraphs are same size (load balance)
- Cut size is minimal (minimize communication)

Matrices that are "almost" diagonal are good?

## Reordering for bandedness



Natural order


RCM reordering

Reverse Cuthill-McKee

- Select "peripheral" vertex $v$
- Order according to breadth first search from $v$
- Reverse ordering


## From iterative to direct

- RCM ordering is great for SpMV
- But isn't narrow banding good for solvers, too?
- LU takes $O\left(n b^{2}\right)$ where $b$ is bandwidth.
- Great if there's an ordering where $b$ is small!


## Skylines and profiles

- Profile solvers generalize band solvers
- Use skyline storage; if storing lower triangle, for each row $i$ :
- Start and end of storage for nonzeros in row.
- Contiguous nonzero list up to main diagonal.
- In each column, first nonzero defines a profile.
- All fill-in confined to profile.
- RCM is again a good ordering.


## Beyond bandedness

- Bandedness only takes us so far
- Minimum bandwidth for 2D model problem? 3D?
- Skyline only gets us so much farther
- But more general solvers have similar structure
- Ordering (minimize fill)
- Symbolic factorization (where will fill be?)
- Numerical factorization (pivoting?)
- ... and triangular solves


## Reminder: Matrices to graphs

- $A_{i j} \neq 0$ means there is an edge between $i$ and $j$
- Ignore self-loops and weights for the moment
- Symmetric matrices correspond to undirected graphs


## Troublesome Trees



One step of Gaussian elimination completely fills this matrix!

## Terrific Trees



Full Gaussian elimination generates no fill in this matrix!

## Graphic Elimination





Eliminate a variable, connect all neighbors.

## Graphic Elimination

Consider first steps of GE

$$
\begin{aligned}
\mathrm{A}(2: \text { end, } 1)= & A(2: \text { end }, 1) / A(1,1) ; \\
A(2: \text { end, } 2: \text { end })= & A(2: \text { end }, 2: \text { end })-\ldots \\
& \text { A }(2: \text { end, } 1) \star A(1,2: \text { end }) ;
\end{aligned}
$$

Nonzero in the outer product at $(i, j)$ if $A(i, 1)$ and $A(j, 1)$ both nonzero - that is, if $i$ and $j$ are both connected to 1 .

General: Eliminate variable, connect remaining neighbors.

## Terrific Trees Redux



Order leaves to root $\Longrightarrow$ on eliminating $i$, parent of $i$ is only remaining neighbor.

## Nested Dissection



- Idea: Think of block tree structures.
- Eliminate block trees from bottom up.
- Can recursively partition at leaves.
- Rough cost estimate: how much just to factor dense Schur complements associated with separators?
- Notice graph partitioning appears again!
- And again we want small separators!


## Nested Dissection

Model problem: Laplacian with 5 point stencil (for 2D)

- ND gives optimal complexity in exact arithmetic (George 73, Hoffman/Martin/Rose)
- 2D: $O(N \log N)$ memory, $O\left(N^{3 / 2}\right)$ flops
- 3D: $O\left(N^{4 / 3}\right)$ memory, $O\left(N^{2}\right)$ flops


## Minimum Degree

- Locally greedy strategy
- Want to minimize upper bound on fill-in
- Fill $\leq$ (degree in remaining graph $)^{2}$
- At each step
- Eliminate vertex with smallest degree
- Update degrees of neighbors
- Problem: Expensive to implement!
- But better varients via quotient graphs
- Variants often used in practice


## Elimination Tree

- Variables (columns) are nodes in trees
- $j$ a descendant of $k$ if eliminating $j$ updates $k$
- Can eliminate disjoint subtrees in parallel!


## Cache locality

Basic idea: exploit "supernodal" (dense) structures in factor

- e.g. arising from elimination of separator Schur complements in ND
- Other alternatives exist (multifrontal solvers)


## Pivoting

Pivoting is a tremendous pain, particularly in distributed memory!

- Cholesky - no need to pivot!
- Threshold pivoting - pivot when things look dangerous
- Static pivoting - try to decide up front

What if things go wrong with threshold/static pivoting?
Common theme: Clean up sloppy solves with good residuals

## Direct to iterative

Can improve solution by iterative refinement:

$$
\begin{aligned}
P A Q & \approx L U \\
x_{0} & \approx Q U^{-1} L^{-1} P b \\
r_{0} & =b-A x_{0} \\
x_{1} & \approx x_{0}+Q U^{-1} L^{-1} P r_{0}
\end{aligned}
$$

Looks like approximate Newton on $F(x)=A x-b=0$. This is just a stationary iterative method!
Nonstationary methods work, too.

## Variations on a theme

If we're willing to sacrifice some on factorization,

- Single precision + refinement on double precision residual?
- Sloppy factorizations (marginal stability) + refinement?
- Modify $m$ small pivots as they're encountered (low rank updates), fix with $m$ steps of a Krylov solver?

