The Cost of Iterative Computations

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• Iterative linear algebra computations
  • Eigenvalue problems, linear systems, etc.
  • Matrix $A$ typically large and sparse
• Ubiquitous in applications, but typically incapable of fully exploiting underlying hardware
• Requires the development of new methods, algorithms, and implementations (and hardware!) capable meeting energy and/or runtime constraints
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• Goal: determine the optimal computational approach for a given machine and a given instance of data $A$ and $b$
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• Goal: determine the optimal computational approach for a given machine and a given instance of data $A$ and $b$

What is the cost of an iterative computation?
What is a computation?

- problem
- instance (data, machine, etc.)
- implementation
- algorithm
- method
1. The Problem

Ultimately, the computation is performed in order to solve some *problem*

Running example: Solve $N \times N$ linear system $Ax = b$

Must also consider greater context:

• The origin of the problem dictates mathematical structure
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Consider the infinite dimensional problem

$$Gu = f$$

where $G: S \rightarrow S$ is a bounded invertible operator on Hilbert space $S$

The problem is approximated on a finite dimensional subspace $S_h \subset S$ by the finite dimensional operator $G_h$, giving

$$G_h u_h = f_h$$
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Choosing a basis for $S_h$ gives rise to the matrix problem

$$Ax = b$$
2. The Method

- Mathematical approach for transforming the data
- Many possible choices depending on the properties and structure of $A$
  - Dense $A \rightarrow$ LU (or Cholesky if SPD)
  - Large, sparse, and SPD $A \rightarrow$ Conjugate Gradient method
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Krylov subspace method
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Krylov subspace method constructs at step $i$ an approximation $A_i$ of $A$ with desired approximate solution

$$x_i = \rho_{i-1}(A_i)b \approx A^{-1}b = x$$

where $\rho_{i-1}(\lambda)$ is associated polynomial of degree at most $i - 1$. 
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where $\rho_{i-1}(\lambda)$ is associated polynomial of degree at most $i - 1$.

$A_i$ is obtained by restricting and projecting $A$ onto the $i$th Krylov subspace

$$\mathcal{K}_i(A, r_0) = \text{span}\{r_0, Ar_0, \ldots, A^{i-1}r_0\}$$

where $r_0 = b - Ax_0$,*

*for connection to infinite dimensional setting see [Málek & Strakoš, 2015]
Let $V_i = [v_1, \ldots, v_i]$ be a basis for the Krylov subspace $\mathcal{K}_i(A, r_0)$.

In step $i$, we have the operator (on a subspace of small dimension):

$$A_i = V_i V_i^* A V_i V_i^*$$
The conjugate gradient method

Let $V_i = [v_1, ..., v_i]$ be a basis for the Krylov subspace $\mathcal{K}_i(A, r_0)$

In step $i$, we have the operator (on a subspace of small dimension):

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$$T_i$$

In the conjugate gradient method, projected matrix is a tridiagonal Jacobi matrix $T_i$
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In each step $i$, solve the projected system

$$T_i y_i = \|r_0\| e_1$$

and update the approximate solution $x_i = x_0 + V_i y_i$
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and update the approximate solution $x_i = x_0 + V_i y_i$

(can be seen as a model reduction of the original system $Ax = b$)
The residuals in CG are of the form

\[ r_i = \rho^{CG}_i (A) r_0 \]

where \( \rho^{CG}_i (\lambda) \) is the CG polynomial which satisfies \( \rho^{CG}_0 (\lambda) = 1 \) and

\[
\rho^{CG}_i (\lambda) = \frac{\left( \lambda - \theta_1^{(i)} \right) \cdots \left( \lambda - \theta_i^{(i)} \right)}{(-1)^i \theta_1^{(i)} \cdots \theta_i^{(i)}}
\]

where \( \theta_1^{(i)}, \ldots, \theta_i^{(i)} \) are the eigenvalues of the Jacobi matrix \( T_i \)
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where \( \theta_1^{(i)}, \ldots, \theta_i^{(i)} \) are the eigenvalues of the Jacobi matrix \( T_i \)

CG polynomial is uniquely defined by the minimization problem

\[ \|x - x_i\|_A = \min_{\rho(0)=1, \deg(\rho) \leq i} \|\rho(A)(x - x_0)\|_A = \|\rho_i^{CG}(A)(x - x_0)\|_A \]

which is equivalent to

\[ \|x - x_i\|_A = \min_{z \in x_0 + \mathcal{K}_i(A, r_0)} \|x - z\|_A. \]

In each step \( i \), CG picks the approximate solution from the shifted Krylov subspace \( x_0 + \mathcal{K}_i(A, r_0) \) that **minimizes the A-norm (energy norm) of the error**.

\( \Rightarrow \) CG (and other Krylov subspace methods) are highly nonlinear.
3. The Algorithm

- Many potential algorithms to choose from to implement a given method
- CG: Hestenes and Stiefel (1952) (HSCG)
  - Uses three 2-term recurrences for updating $x_i, r_i, p_i$

\[
\begin{align*}
  r_0 &= b - Ax_0, \quad p_0 = r_0 \\
  \text{for } i &= 1:n_{\text{max}} \\
  \alpha_{i-1} &= \frac{r_{i-1}^T r_{i-1}}{p_{i-1}^T Ap_{i-1}} \\
  x_i &= x_{i-1} + \alpha_{i-1} p_{i-1} \\
  r_i &= r_{i-1} - \alpha_{i-1} Ap_{i-1} \\
  \beta_i &= \frac{r_i^T r_i}{r_{i-1}^T r_{i-1}} \\
  p_i &= r_i + \beta_i p_{i-1}
\end{align*}
\]
\[ r_0 = b - Ax_0, \quad p_0 = r_0 \]

\[ s_0 = Ap_0, w_0 = Ar_0, z_0 = Aw_0, \]

\[ \alpha_0 = r_0^T r_0 / p_0^T s_0 \]

for \( i = 1 : \text{nmax} \)

\[ x_i = x_{i-1} + \alpha_{i-1} p_{i-1} \]

\[ r_i = r_{i-1} - \alpha_{i-1} s_{i-1} \]

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\[ \alpha_i = \frac{r_i^T r_i}{w_i^T r_i - (\beta_i / \alpha_{i-1}) r_i^T r_i} \]

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end
Pipelined CG [Ghysels and Vanroose, 2014]

\[ r_0 = b - Ax_0, \quad p_0 = r_0 \]
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for \( i = 1:\text{nmax} \)

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for \( i = 1 : n_{\text{max}} \)

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end
\[ r_0 = b - Ax_0, \quad p_0 = r_0 \]

for \( k = 0: \text{nmax}/s \)

Compute \( y_k \) and \( b_k \) such that \( A y_k = y_k b_k \) and

\[
\text{span}(y_k) = \mathcal{K}_{s+1}(A, p_{sk}) + \mathcal{K}_s(A, r_{sk})
\]

\[ G_k = y_k^T y_k \]

\[ x'_0 = 0, \quad r'_0 = e_{s+2}, \quad p'_0 = e_1 \]

for \( j = 1: s \)

\[
\alpha_{sk+j-1} = \frac{r'_{j-1}^T G_k r'_{j-1}}{p'_{j-1}^T G_k b_k p'_{j-1}} \]

\[ x'_j = x'_{j-1} + \alpha_{sk+j-1} p'_{j-1} \]

\[ r'_j = r'_{j-1} - \alpha_{sk+j-1} b_k p'_{j-1} \]

\[
\beta_{sk+j} = \frac{r'_{j}^T G_k r'_j}{r'_{j-1}^T G_k r'_{j-1}} \]

\[ p'_j = r'_j + \beta_{sk+j} p'_{j-1} \]

end

\[
[x_{s(k+1)}-x_{sk}, r_{s(k+1)}, p_{s(k+1)}] = y_k [x'_s, r'_s, p'_s] \]

end
\[ r_0 = b - Ax_0, p_0 = r_0 \]
for \( k = 0:n_{max}/s \)

Compute \( Y_k \) and \( B_k \) such that \( AY_k = Y_k B_k \) and
\[
\text{span}(Y_k) = \mathcal{K}_{s+1}(A,p_{sk}) + \mathcal{K}_s(A,r_{sk})
\]
\[ G_k = Y_k^T Y_k \]
\[ x'_0 = 0, r'_0 = e_{s+2}, p'_0 = e_1 \]
for \( j = 1:s \)

\[ \alpha_{sk+j-1} = \frac{r'_{j-1}^T G_k r'_{j-1}}{p'_{j-1}^T G_k B_k p'_{j-1}} \]
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\[ [x_{s(k+1)} - x_{sk}, r_{s(k+1)}, p_{s(k+1)}] = Y_k [x'_s, r'_s, p'_s] \]
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for \( k = 0 : \text{nmax}/s \)

Compute \( Y_k \) and \( B_k \) such that \( AY_k = Y_kB_k \) and \( \text{span}(Y_k) = \mathcal{K}_{s+1}(A, p_{sk}) + \mathcal{K}_s(A, r_{sk}) \)

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for \( j = 1 : s \)

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[\[ x_{s(k+1)} - x_{sk}, r_{s(k+1)}, p_{s(k+1)} \] = \[ Y_k [x'_s, r'_s, p'_s] \]

end

end
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Compute \( y_k \) and \( B_k \) such that \( A y_k = y_k B_k \) and

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\[ G_k = y_k^T y_k \]

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\[ [x_{s(k+1)}-x_{sk}, r_{s(k+1)}, p_{s(k+1)}] = \text{span}[y_k, x'_s, r'_s, p'_s] \]

end
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for \( k = 0 : \text{nmax}/s \)

Compute \( y_k \) and \( B_k \) such that \( A y_k = y_k B_k \) and

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\( x'_0 = 0, r'_0 = e_{s+2}, p'_0 = e_1 \)

for \( j = 1 : s \)

\[
\alpha_{sk+j-1} = \frac{r_{j-1}' T g_k r_{j-1}'}{p_{j-1}' T g_k B_k p_{j-1}'}
\]

\[
x'_j = x'_{j-1} + \alpha_{sk+j-1} p'_{j-1}
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r'_j = r'_{j-1} - \alpha_{sk+j-1} B_k p'_{j-1}
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\beta_{sk+j} = \frac{r'_j T g_k r'_j}{r'_{j-1} T g_k r'_{j-1}}
\]

\[
p'_j = r'_{j} + \beta_{sk+j} p'_{j-1}
\]

end

\[
[x_{sk(k+1)}, r_{sk(k+1)}, p_{sk(k+1)}] = y_k [x'_s, r'_s, p'_s]
\]

end

**Graphical Representation**

- **Outer Loop**
  - Compute basis \( O(s) \) SPMVs
  - \( O(s^2) \) Inner Products (one synchronization)

- **Inner Loop**
  - Local Vector Updates (no comm.)

- **End Inner Loop**

- **Inner Outer Loop**

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  - Programming language, parallelization, machine precision, etc.

- Instance: the particular data $A, b$, stopping criteria, particular machine
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How is cost defined?

Work on extending the Turing machine model to numerical algorithms motivated by apparent lack of formal notion of cost ...

"There is not even a formal definition of algorithm in the subject ... Thus we view numerical analysis as an eclectic subject with weak foundations."

[Blum, Cucker, Shub, Smale, 1998]

"Phrases like 'cost', 'gap', etc., used within mathematical formalism, are ambiguous. Rigorous proofs require rigorous definitions."

[Iserles, 2000]
1. Cost is well-defined in terms of energy and/or runtime
   • Measurable and meaningful even in total absence of a formal Turing machine model

2. The computational transformation of the data, consisting of a problem, method, algorithm, implementation, and instance is the key concept.
   • Focus on the algorithm misses the bigger picture
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Important! : for iterative computations:

\[
\text{cost} = (\text{cost per iteration}) \times (\text{number of iterations})
\]
The cost per iteration

- Cost per iteration depends on particular algorithm, implementation, and data/machine
- Single iteration broken down into computational kernels (SpMV, inner products, etc.)
- The cost of these kernels can be modeled in terms of computation (flops) and communication (data movement)
The cost per iteration

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- Single iteration broken down into computational kernels (SpMV, inner products, etc.)

- The cost of these kernels can be modeled in terms of computation (flops) and communication (data movement)

- Simplified cost model: If a processor performs $F$ flops, and sends/receives $S$ messages containing a total of $W$ words, we then model its worst-case cost as

$$\text{cost} = \gamma F + \beta W + \alpha S$$

where $\gamma$ is cost of a flop on local data, $\beta$ is cost per word of data moved (inverse bandwidth), $\alpha$ is cost per message (latency)

  - Machine structure: CPUs, GPUs, half-precision tensor cores, accelerators, etc.

- Rates of improvement: $\gamma \gg \beta \gg \alpha$
  - Communication more expensive than computation, trend will continue
Describing CG Convergence

Infinite precision: CG convergence rate depends strongly on the distribution of eigenvalues

\[
\frac{\|x - x_i\|_A}{\|x - x_0\|_A} \leq \min_{\rho(0)=1} \max_{1 \leq j \leq N} |\rho(\lambda_j)|
\]

Let \(d\) be the number of distinct eigenvalues of \(A\). For \(i = 1, \ldots, d - 1\), there exist \(i + 1\) distinct eigenvalues of \(A\), \(\hat{\lambda}_1, \ldots, \hat{\lambda}_{i+1}\), such that

\[
\min_{\rho(0)=1} \max_{1 \leq j \leq N} |\rho(\hat{\lambda}_j)| = \left( \sum_{k=1}^{i+1} \prod_{j=1}^{i+1} \frac{|\hat{\lambda}_j|}{|\hat{\lambda}_j - \hat{\lambda}_k|} \right)^{-1}
\]

[Greenbaum, 1979]
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\[
\frac{\|x - x_i\|_A}{\|x - x_0\|_A} \leq \min_{\rho(0)=1} \max_{1 \leq j \leq N, \deg(\rho) \leq i} |\rho(\lambda_j)|
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\[
\begin{align*}
\min_{\rho(0)=1} \max_{1 \leq j \leq N, \deg(\rho) \leq i} |\rho(\hat{\lambda}_j)| &= \left( \sum_{k=1}^{i+1} \prod_{j=1 \atop j \neq k}^{i+1} \frac{|\hat{\lambda}_j|}{|\hat{\lambda}_j - \hat{\lambda}_k|} \right)^{-1} \\
&= \left( \frac{\sqrt{\kappa(A)} - 1}{\sqrt{\kappa(A)} + 1} \right)^i \\
&= 2 \left( \frac{\sqrt{\kappa(A)} - 1}{\sqrt{\kappa(A)} + 1} \right)^i \\
\end{align*}
\]

\[\kappa(A) = \frac{\lambda_N}{\lambda_1}\]

Frequent (over)simplification: estimate by replacing set of eigenvalues of \(A\) by continuous interval \([\lambda_1, \lambda_N]\) and use scaled and shifted Chebyshev polynomials:

\[
\frac{\|x - x_i\|_A}{\|x - x_0\|_A} \leq 2 \left( \frac{\sqrt{\kappa(A)} - 1}{\sqrt{\kappa(A)} + 1} \right)^i
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\]

Chebyshev polynomial-based bound holds for any distribution of eigenvalues between \(\lambda_1\) and \(\lambda_N\) and any distribution of the components of the initial residuals in the individual invariant subspaces!
Describing CG Convergence

Infinite precision: CG convergence rate depends strongly on the distribution of eigenvalues

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Let \( d \) be the number of distinct eigenvalues of \( A \). For \( i = 1, \ldots, d - 1 \), there exist \( i + 1 \) distinct eigenvalues of \( A, \hat{\lambda}_1, \ldots, \hat{\lambda}_{i+1} \), such that

\[
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\]

Chebyshev polynomial-based bound holds for \textit{any} distribution of eigenvalues between \( \lambda_1 \) and \( \lambda_N \) and \textit{any} distribution of the components of the initial residuals in the individual invariant subspaces!

\( \Rightarrow \) \textit{Linearization of highly nonlinear phenomena}
Example: diagonal matrix with $N = 25$, 
\[ \lambda_1 = 0.1, \lambda_N = 100 \]
\[ \lambda_j = \lambda_1 + \frac{j - 1}{N - 1} (\lambda_N - \lambda_1) \gamma^{N-j} \]
RHS: $b_i = 1/5$

\[ \gamma = 0.1 : \]
\[ \gamma = 0.9 : \]
"Soon after the introduction of $\kappa(A)$ for error analysis, Hestenes and Stiefel showed that this quantity also played a role in complexity analysis. More precisely, they showed that the number of iterations of the conjugate gradient method (assuming infinite precision) needed to ensure that the current approximation to the solution of a linear system attained a given accuracy is proportional to $\sqrt{\kappa(A)}$. "

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- The Chebyshev polynomial-based bound does not appear anywhere in the original 1952 paper of Hestenes and Stiefel (1952)
- First appearance in the literature in [Daniel, 1967], although he did not identify it with the CG "convergence rate":
  
  "Assuming only that the spectrum of the matrix $A$ lies inside the interval $[\lambda_1, \lambda_N]$, we can do no better than [the $\kappa(A)$-based bound]."
The effects of rounding errors

Well-known that roundoff error has two effects:

1. Delay of convergence
   - No longer have Krylov subspace
   - Can lose numerical rank deficiency
   - Residuals no longer orthogonal
     - Minimization of $\|x - x_i\|_A$

2. Loss of attainable accuracy
   - Rounding errors cause true residual $b - Ax_i$ and updated residual $r_i$ deviate!

$A$: bcsstk03 from SuiteSparse,
b: equal components in the eigenbasis of $A$, $\|b\| = 1$
$N = 112, \kappa(A) \approx 6.8e6$
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Algorithms designed for HPC

• Computational complexity is a poor measure of runtime/energy cost
  • Cost depends heavily on communication complexity

• Communication-avoiding algorithms
  • Prove lower bounds on amount of data moved, number of messages
  • Find algorithms that meet lower bounds
  • Many successes in direct numerical linear algebra
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• Extension to iterative numerical linear algebra is tricky
  • Idea: Modify algorithms to reduce synchronization cost over fixed number of steps
  • Long history of work on synchronization-reducing algorithms mathematically equivalent to HSCG (e.g., pipelined CG, s-step CG)
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*But how many steps are required to converge to prescribed accuracy?*
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  **But how many steps are required to converge to prescribed accuracy?**

  **Can we even still converge to prescribed accuracy?**
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  But how many steps are required to converge to prescribed accuracy?
  Can we even still converge to prescribed accuracy?

• Makes little sense to claim these algorithms to be "high-performance algorithms" or "exascale algorithms" without answering these questions
  • Must understand behavior in finite precision and potential amplification of rounding errors
Maximum attainable accuracy

- Accuracy $\|x - \hat{x}_i\|$ generally not computable, but $x - \hat{x}_i = A^{-1}(b - A\hat{x}_i)$.
- Size of the true residual, $\|b - A\hat{x}_i\|$, used as computable measure of accuracy.
- Rounding errors cause the true residual, $b - A\hat{x}_i$, and the updated residual, $\hat{r}_i$, to deviate.
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- Rounding errors cause the true residual, $b - A\hat{x}_i$, and the updated residual, $\hat{r}_i$, to deviate

- Writing $b - A\hat{x}_i = \hat{r}_i + b - A\hat{x}_i - \hat{r}_i$,

$$||b - A\hat{x}_i|| \leq ||\hat{r}_i|| + ||b - A\hat{x}_i - \hat{r}_i||$$

- As $||\hat{r}_i|| \to 0$, $||b - A\hat{x}_i||$ depends on $||b - A\hat{x}_i - \hat{r}_i||$
Accuracy $\|x - \hat{x}_i\|$ generally not computable, but $x - \hat{x}_i = A^{-1}(b - A\hat{x}_i)$

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As $\|\hat{r}_i\| \rightarrow 0$, $\|b - A\hat{x}_i\|$ depends on $\|b - A\hat{x}_i - \hat{r}_i\|$

Maximum attainable accuracy of HSCG

- In finite precision HSCG, iterates are updated by

\[ \hat{x}_i = \hat{x}_{i-1} + \hat{a}_{i-1} \hat{p}_{i-1} - \delta x_i \]  

and

\[ \hat{r}_i = \hat{r}_{i-1} - \hat{a}_{i-1} A \hat{p}_{i-1} - \delta r_i \]
In finite precision HSCG, iterates are updated by

\[ \hat{x}_i = \hat{x}_{i-1} + \hat{\alpha}_{i-1}\hat{p}_{i-1} - \delta x_i \quad \text{and} \quad \hat{r}_i = \hat{r}_{i-1} - \hat{\alpha}_{i-1}A\hat{p}_{i-1} - \delta r_i \]

Let \( f_i \equiv b - A\hat{x}_i - \hat{r}_i \)

\[
\begin{align*}
    f_i &= b - A(\hat{x}_{i-1} + \hat{\alpha}_{i-1}\hat{p}_{i-1} - \delta x_i) - (\hat{r}_{i-1} - \hat{\alpha}_{i-1}A\hat{p}_{i-1} - \delta r_i) \\
    &= f_{i-1} + A\delta x_i + \delta r_i \\
    &= f_0 + \sum_{m=1}^{i}(A\delta x_m + \delta r_m)
\end{align*}
\]
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= f_{i-1} + A\delta x_i + \delta r_i
= f_0 + \sum_{m=1}^i (A\delta x_m + \delta r_m)
\]

\[
\|f_i\| \leq \mathcal{O}(\varepsilon) \sum_{m=0}^i N_A \|A\| \|\hat{x}_m\| + \|\hat{r}_m\| \quad \text{van der Vorst and Ye, 2000}
\]

\[
\|f_i\| \leq \mathcal{O}(\varepsilon) \|A\| (\|x\| + \max_{m=0,\ldots,i} \|\hat{x}_m\|)
\quad \text{Greenbaum, 1997}
\]

\[
\|f_i\| \leq \mathcal{O}(\varepsilon) N_A \|A\| \|A^{-1}\| \sum_{m=0}^i \|\hat{r}_m\| \quad \text{Sleijpen and van der Vorst, 1995}
\]
Computed explicitly: $q_i \equiv A p_i$

Pipelined CG uses 3 auxiliary recurrences:

\[ s_i \equiv A p_i, \quad w_i = A r_i, \quad z_i \equiv A q_i \]

\[
\hat{p}_i = \hat{r}_i + \beta_i \hat{p}_{i-1} + \delta_i^p \\
\hat{s}_i = \hat{w}_i + \beta_i \hat{s}_{i-1} + \delta_i^s \\
\hat{z}_i = \hat{q}_i + \beta_i \hat{z}_{i-1} + \delta_i^z \\
\hat{x}_i = \hat{x}_{i-1} + \alpha_i \hat{p}_{i-1} + \delta_i^x \\
\hat{r}_i = \hat{r}_{i-1} - \alpha_i \hat{s}_{i-1} + \delta_i^r \\
\hat{w}_i = \hat{w}_{i-1} - \alpha_i \hat{z}_{i-1} + \delta_i^w
\]
Attainable Accuracy of Pipelined CG

\[ f_i = f_0 - \sum_{j=0}^{i} \hat{a}_j g_j - \sum_{j=0}^{i} (A\delta_j^x + \delta_j^r) \]

\( f_i \equiv b - A\hat{x}_i - \hat{r}_i \)
Attainable Accuracy of Pipelined CG

\[ f_i = f_0 - \sum_{j=0}^{i} \hat{\alpha}_j g_j - \sum_{j=0}^{i} (A\delta_j^x + \delta_j^y) \]

\[ g_j = \left( \prod_{k=1}^{j} \beta_k \right) g_0 + \sum_{k=1}^{j} \left( \prod_{\ell=k+1}^{j} \beta_{\ell} \right) (A\delta_k^p - \delta_k^s) + h_k \]

\[ h_k = h_0 + \sum_{\ell=0}^{k-1} ((A\delta_{\ell}^r - \delta_{\ell}^w) - \hat{\alpha}_\ell c_\ell) \]

\[ c_\ell = \left( \prod_{m=1}^{\ell} \beta_m \right) c_0 + \sum_{m=1}^{\ell} \left( \prod_{n=m+1}^{\ell} \beta_n \right) (A\delta_m^q - \delta_m^z) \]

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**Attainable Accuracy of Pipelined CG**

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Local rounding errors all potentially amplified!
\[ A : \text{bcsstk03 from SuiteSparse}, \]
\[ b : \text{equal components in the eigenbasis of } A, \|b\| = 1 \]
\[ N = 112, \kappa(A) \approx 6.8e6 \]
Attainable accuracy of s-step CG

\[ f_i \equiv b - A\hat{x}_i - \hat{r}_i \]

For CG:

\[ \|f_i\| \leq \|f_0\| + \varepsilon \sum_{m=1}^{i} (1 + N)\|A\|\|\hat{x}_m\| + \|\hat{r}_m\| \]

[see C., 2015]
For CG:

\[ f_i \equiv b - A\hat{x}_i - \hat{r}_i \]

For CG:

\[ \|f_i\| \leq \|f_0\| + \varepsilon \sum_{m=1}^{i} (1 + N)\|A\|\|\hat{x}_m\| + \|\hat{r}_m\| \]

For s-step CG: \( i \equiv sk + j \)

\[ \|f_{sk+j}\| \leq \|f_{sk}\| + \varepsilon \Gamma_k \sum_{\ell=1}^{j} (1 + N)\|A\|\|\hat{x}_{sk+\ell}\| + \|\hat{r}_{sk+\ell}\| \]

where \( \Gamma_k = c \cdot \|\hat{y}_k\| \|\hat{y}_k\| \), \( c \) is a low-degree polynomial in \( s \)

Local rounding errors amplified; amplification is "local" within s-steps

[see C., 2015]
Attainable accuracy of s-step CG

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Local rounding errors amplified; amplification is "local" within s-steps

\[ \|f_{sk+j}\| \leq \|f_0\| + \varepsilon \bar{\Gamma}_k \sum_{m=1}^{sk+j} (1 + N)\|A\|\|\hat{x}_m\| + \|\hat{r}_m\| \]

where \( \bar{\Gamma}_k = \max_{m \leq k} \Gamma_m \)

Accuracy of s-step CG in prec. \( \varepsilon \leftrightarrow \) Accuracy of CG in prec. \( \varepsilon \bar{\Gamma}_k \)

[see C., 2015]
s-step CG with monomial basis ($\mathbf{Y} = [p_i, Ap_i, ..., A^s p_i, r_i, Ar_i, ... A^{s-1} r_i]$)

$A$: bcsstk03 from SuiteSparse,
$b$: equal components in the eigenbasis of $A$, $\|b\| = 1$
$N = 112, \kappa(A) \approx 6.8e6$
Even assuming cost per iteration decreases by factor of $s$, already at $s = 4$ we are worse than HSCG in terms of number of synchronizations!

\[ A: \text{bcsstk03 from SuiteSparse}, \]
\[ b: \text{equal components in the eigenbasis of } A, \|b\| = 1 \]
\[ N = 112, \kappa(A) \approx 6.8e6 \]
A different problem...

\( A: \text{nos4} \) from SuiteSparse,
\( b: \) equal components in the eigenbasis of \( A \) and \( \|b\| = 1 \)
\( N = 100, \kappa(A) \approx 2e3 \)

If application only requires
\[
\frac{\|x - \hat{x}_i\|_A}{\|x - x_0\|_A} \approx 10^{-6},
\]
any of these algorithms will work!
Towards understanding convergence delay

• CG method = matrix formulation of Gauss-Christoffel quadrature (see [Liesen & Strakoš, 2013])

• Coefficients $\alpha$ and $\beta$ (related to entries of $T_i$) determine distribution functions $\omega^{(i)}(\lambda)$ which approximate distribution function $\omega(\lambda)$ determined by inputs $A, b, x_0$ in terms of the $i$th Gauss-Christoffel quadrature

• $A$-norm of CG error for $f(\lambda) = \lambda^{-1}$ given as scaled quadrature error

$$
\int \lambda^{-1} d\omega(\lambda) = \sum_{\ell=1}^{i} \omega^{(i)}_{\ell} \left\{ \theta^{(i)}_{\ell} \right\}^{-1} + \frac{\|x - x_i\|^2_A}{\|r_0\|^2} = \left( \frac{\|x - x_0\|^2_A}{\|r_0\|^2} \right)
$$
Towards understanding convergence delay

- CG method = matrix formulation of Gauss-Christoffel quadrature (see [Liesen & Strakoš, 2013])

- Coefficients $\alpha$ and $\beta$ (related to entries of $T_i$) determine distribution functions $\omega^{(i)}(\lambda)$ which approximate distribution function $\omega(\lambda)$ determined by inputs $A, B, x_0$ in terms of the $i$th Gauss-Christoffel quadrature

- A-norm of CG error for $f(\lambda) = \lambda^{-1}$ given as scaled quadrature error

  $$
  \int \lambda^{-1} d\omega(\lambda) = \sum_{\ell=1}^{i} \omega^{(i)}_{\ell} \{\theta^{(i)}_{\ell}\}^{-1} + \frac{\|x - x_i\|_A^2}{\|r_0\|^2} = \left(\frac{\|x - x_0\|_A^2}{\|r_0\|^2}\right)
  $$

- For particular CG implementation, can the computed $\hat{\omega}^{(i)}(\lambda)$ be associated with some distribution function $\hat{\omega}(\lambda)$ related to the distribution function $\omega(\lambda)$, i.e.,

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  $$

  where $F_i$ is small relative to error term?
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- For classical CG, yes; proved by Greenbaum [1989]

- For pipelined CG and s-step CG, THOROUGH ANALYSIS NEEDED!
Designing preconditioners

- Approach: design preconditioner $M$ such that preconditioned linear system $M^{-1}Ax = M^{-1}b$ converges in few iterations
- Frequent assertion:

  \[ \text{\# of clusters of eigenvalues} = \text{\# iterations for Krylov subspace method to converge} \]
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Example: diagonal matrix with $N = 25$, $\lambda_1 = 0.1$, $\lambda_N = 100$, $\gamma = 0.65$

\[ \lambda_j = \lambda_1 + \frac{j - 1}{N - 1} (\lambda_N - \lambda_1) \gamma^{N-j} \]

RHS: $b_i = 1/5$
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**Argument also fails for GMRES** (see [Greenbaum & Strakoš, 1994], [Greenbaum, Pták, Strakoš, 1996], [Arioli, Pták, Strakoš, 1998])

Example [Liesen & Tichý, 2004]:

- **exact GMRES** on
  - $A = \text{gallery('prolate', 40)}$
  - RHS: $b_i = 1/\sqrt{N}$

Note: the matrix $A$ is normal!
Other considerations:

- If $A$ is ill conditioned, $M$ is likely to be ill conditioned

\[ \hat{z}_i = (M^{-1}A + \Delta) \hat{r}_i \quad \|\Delta\|_2 \leq O(\varepsilon)N^{3/2}(\kappa(A) + \kappa(M))\|M^{-1}A\|_F \]

- Even if preconditioned system is well conditioned, application can still introduce significant roundoff error
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  • Fewer iterations, but potentially much more expensive

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"The impetus towards 'general' methods and 'general' software, which can cater to many different problems in a broad category, might be inimical to progress. Once we classify mathematical problems by their structural features, broad categories are much too coarse."
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- Only hope: algorithms that are able to adapt to the data

"Good computation requires the algorithm to respond to the data it is producing and change the allocation of computing resources (step size, size of the grid, number of iterations,..., even the discretization method itself) accordingly."

[Baxter and Iserles, 2003]
"Numerical analysis lies at the meeting point of pure mathematics, computer science, and application areas. It often attracts some degree of hostility from all three."

[Baxter and Iserles, *On the foundations of computational mathematics*, Handbook of numerical analysis, 11 (2003), pp.3-34]
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- All perspectives are valuable and can lead to interesting insights; none alone gives complete description
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• All perspectives are valuable and can lead to interesting insights; none alone gives complete description
• For bigger picture, must considered all aspects of a computation together → holistic approach needed
• Confluence of data science/informatics and computational science
  • Motivating changes in hardware, new algorithms, new approaches
  • Not every linear system comes from a PDE!
References


Thank you!
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