# The s-Step Conjugate Gradient Method in Finite Precision 

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## SIAM CSE '19

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## Conjugate Gradient on the World's Fastest Computer

## Summit - IBM Power System AC922

| Site: | Oak Ridge National Laboratory |
| :--- | :--- |
| Manufacturer: | IBM |
| Cores: | $2,282,544$ |
| Memory: | $2,801,664$ GB |
| Processor: | IBM POWER9 22C 3.07 GHz |
| Interconnect: | Dual-rail Mellanox EDR Infiniband |
| Performance |  |
| Theoretical peak: | 187,659 TFlops/s |
| LINPACK benchmark: | 122,300 Tflops/s |
| HPCG benchmark: | 2,926 Tflops/s |
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LINPACK benchmark (dense $A x=b$, direct) $65 \%$ efficiency

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| Site: | Oak Ridge National Laboratory | LINPACK benchmark (dense $A x=b$, direct) $65 \%$ efficiency <br> - HPCG benchmark (sparse $A x=b$, iterative) $1.5 \%$ efficiency |
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## The Conjugate Gradient (CG) Method

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\begin{aligned}
& r_{0}=b-A x_{0}, \quad p_{0}=r_{0} \\
& \text { for } i=1 \text { nmax } \\
& \qquad \begin{array}{l}
\alpha_{i-1}=\frac{r_{i-1}^{T} r_{i-1}}{p_{i-1}^{T} A p_{i-1}} \\
x_{i}=x_{i-1}+\alpha_{i-1} p_{i-1} \\
r_{i}=r_{i-1}-\alpha_{i-1} A p_{i-1} \\
\\
\qquad \beta_{i}=\frac{r_{i}^{T} r_{i}}{r_{i-1}^{T} r_{i-1}} \\
\text { end }
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## The Conjugate Gradient (HSCG) Method

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\text { end } \\
p_{i}=r_{i}+\beta_{i} p_{i-1}
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\end{aligned}
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Iteration Loop

Sparse Matrix $\times$ Vector

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\end{array} \\
& \text { end }
\end{aligned}
$$


$\Rightarrow$ Communication bottleneck!

## s-step Krylov subspace methods

- Idea: Compute blocks of $s$ iterations at once
- Compute updates in a different basis
- Communicate every $s$ iterations instead of every iteration
- Reduces number of synchronizations per iteration by a factor of $s$
- An idea rediscovered many times...
- First related work: s-dimensional steepest descent, least squares
- Khabaza ('63), Forsythe ('68), Marchuk and Kuznecov ('68)
- Flurry of work on s-step Krylov methods in '80s/early '90s: see, e.g., Van Rosendale (1983); Chronopoulos and Gear (1989)
- Resurgence of interest in recent years due to growing problem sizes; growing relative cost of communication


## s-step CG

$r_{0}=b-A x_{0}, p_{0}=r_{0}$
for $k=0: n \max / s$
Compute $\mathcal{Y}_{k}$ and $\mathcal{B}_{k}$ such that $A \underline{\mathcal{Y}}_{k}=\mathcal{Y}_{k} \mathcal{B}_{k}$ and

$$
\operatorname{span}\left(\mathcal{Y}_{k}\right)=\mathcal{K}_{s+1}\left(A, p_{s k}\right)+\mathcal{K}_{s}\left(A, r_{s k}\right)
$$

$$
\mathcal{G}_{k}=\mathcal{Y}_{k}^{T} \mathcal{Y}_{k}
$$

$$
x_{0}^{\prime}=0, r_{0}^{\prime}=e_{s+2}, p_{0}^{\prime}=e_{1}
$$

$$
\text { for } j=1: s
$$

$$
\begin{aligned}
& \alpha_{s k+j-1}=\frac{r_{j-1}^{\prime T} \mathcal{G}_{k} r_{j-1}^{\prime}}{p_{j-1}^{\prime T} \mathcal{G}_{k} \mathcal{B}_{k} p_{j-1}^{\prime}} \\
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\end{aligned}
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end
$\left[x_{s(k+1)}-x_{s k}, r_{s(k+1)}, p_{s(k+1)}\right]=\mathcal{Y}_{k}\left[x_{s}^{\prime}, r_{s}^{\prime}, p_{s}^{\prime}\right]$
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Outer Loop

Compute basis
O(s) SPMVs

$$
\mathcal{G}_{k}=y_{k}^{T} \mathcal{Y}_{k}
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## The effects of finite precision

Well-known that roundoff error has two effects:

1. Delay of convergence

- No longer have exact Krylov subspace
- Can lose numerical rank deficiency
- Residuals no longer orthogonal Minimization of $\left\|x-x_{i}\right\|_{A}$ no longer exact

2. Loss of attainable accuracy

- Rounding errors cause true residual $b-A x_{i}$ and updated residual $r_{i}$ deviate!


A: bcsstk03 from SuiteSparse, $b$ : equal components in the eigenbasis of $A,\|b\|=1$

$$
N=112, \kappa(A) \approx 7 \mathrm{e} 6
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N=112, \kappa(A) \approx 7 \mathrm{e} 6
$$ residual $r_{i}$ deviate!

Much work on these results for CG; See Meurant and Strakoš (2006) for a thorough summary of early developments in finite precision analysis of Lanczos and CG

## s-step CG



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s-step CG with monomial basis ( $\left.\mathcal{Y}=\left[p_{i}, A p_{i}, \ldots, A^{s} p_{i}, r_{i}, A r_{i}, \ldots A^{s-1} r_{i}\right]\right)$


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Convergence delay and attainable accuracy worse with increasing s!

## s-step CG

Even assuming perfect parallel scalability with $s$ (which is usually not the case due to extra SpMVs and inner products), already at $s=4$ we are worse than HSCG in terms of number of synchronizations!


## Maximum attainable accuracy

- Accuracy $\left\|x-\hat{x}_{i}\right\|$ generally not computable, but $x-\hat{x}_{i}=A^{-1}\left(b-A \hat{x}_{i}\right)$
- Size of the true residual, $\left\|b-A \hat{x}_{i}\right\|$, used as computable measure of accuracy


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- Writing $b-A \widehat{x}_{i}=\hat{r}_{i}+b-A \hat{x}_{i}-\hat{r}_{i}$,

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- As $\left\|\hat{r}_{i}\right\| \rightarrow 0,\left\|b-A \hat{x}_{i}\right\|$ depends on $\left\|b-A \hat{x}_{i}-\hat{r}_{i}\right\|$
- Many results on bounding attainable accuracy, e.g.: Greenbaum (1989, 1994, 1997), Sleijpen, van der Vorst and Fokkema (1994), Sleijpen, van der Vorst and Modersitzki (2001), Björck, Elfving and Strakoš (1998) and Gutknecht and Strakoš (2000).


## Maximum attainable accuracy of HSCG

- In finite precision HSCG, iterates are updated by

$$
\hat{x}_{i}=\hat{x}_{i-1}+\hat{\alpha}_{i-1} \hat{p}_{i-1}-\boldsymbol{\delta} \boldsymbol{x}_{\boldsymbol{i}} \quad \text { and } \quad \hat{r}_{i}=\hat{r}_{i-1}-\hat{\alpha}_{i-1} A \hat{p}_{i-1}-\boldsymbol{\delta} \boldsymbol{r}_{\boldsymbol{i}}
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$$

- Let $f_{i} \equiv b-A \hat{x}_{i}-\hat{r}_{i}$


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$\left\|f_{i}\right\| \leq O(\varepsilon) \sum_{m=0}^{i} N_{A}\|A\|\left\|\hat{x}_{m}\right\|+\left\|\hat{r}_{m}\right\| \quad$ van der Vorst and $\mathrm{Ye}, 2000$
$\left\|f_{i}\right\| \leq O(\varepsilon)\|A\|\left(\|x\|+\max _{m=0, \ldots, i}\left\|\hat{x}_{m}\right\|\right) \quad$ Greenbaum, 1997
$\left\|f_{i}\right\| \leq O(\varepsilon) N_{A}\||A|\|\left\|A^{-1}\right\| \sum_{m=0}^{i}\left\|\hat{r}_{m}\right\| \quad$ Sleijpen and van der Vorst, 1995

## Sources of local roundoff error in s-step CG

Computing the $s$-step Krylov subspace basis:

$$
A \underline{\mathcal{Y}}_{k}=\hat{\mathcal{Y}}_{k} \mathcal{B}_{k}+\Delta \mathcal{Y}_{k}
$$

Updating coordinate vectors in the inner loop:

$$
\begin{aligned}
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Recovering CG vectors for use in next outer loop:

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For s-step CG: $i \equiv s k+j$

$$
\left\|f_{s k+j}\right\| \leq\left\|f_{0}\right\|+\varepsilon \Gamma_{k} \sum_{m=1}^{s k+j}(1+N)\|A\|\left\|\hat{x}_{m}\right\|+\left\|\hat{r}_{m}\right\|
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Conditioning of computed "s-step basis" plays a huge role in determining numerical behavior!

## Choosing a Polynomial Basis

- Recall: in each outer loop of s-step CG, we compute bases for some Krylov subspaces, e.g., $\mathcal{K}_{s+1}\left(A, p_{i}\right)=\operatorname{span}\left\{p_{i}, A p_{i}, \ldots, A^{s} p_{i}\right\}$


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- Simple loop unrolling gives monomial basis, e.g., $\mathcal{Y}_{k}=\left[p_{m}, A p_{m}, \ldots, A^{s} p_{m}\right]$
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- Improve basis condition number to improve numerical behavior: Use different polynomials to compute a basis for the same subspace.
- Two choices based on spectral information that usually lead to wellconditioned bases:
- Newton polynomials
- Chebyshev polynomials


## "Backwards-like" analysis of Greenbaum

- Anne Greenbaum (1989): finite precision CG with matrix $A$ behaves like exact CG run on a larger matrix $\tilde{A}$ whose eigenvalues lie in tight clusters around the eigenvalues of $A$


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- Based on work of Chris Paige for finite precision Lanczos (1976, 1980):
- Complete rounding error analysis
- Computed eigenvalues lie between extreme eigenvalues of A to within a small multiple of machine precision
- At least one small interval containing an eigenvalue of $A$ is found by the Nth iteration
- The algorithm behaves as if it used full reorthogonalization until a close eigenvalue approximation is found
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- Can we make similar statements for s-step variants?


## Roundoff Error in Lanczos vs. s-step Lanczos

Finite precision Lanczos process: ( $A$ is $N \times N$ with at most $n$ nonzeros per row)

$$
\begin{gathered}
A \widehat{V}_{m}=\hat{V}_{m} \hat{T}_{m}+\hat{\beta}_{m+1} \hat{v}_{m+1} e_{m}^{T}+\delta \widehat{V}_{m} \\
\hat{V}_{m}=\left[\hat{v}_{1}, \ldots, \hat{v}_{m}\right], \quad \delta \hat{V}_{m}=\left[\delta \hat{v}_{1}, \ldots, \delta \hat{v}_{m}\right], \quad \hat{T}_{m}=\left[\begin{array}{ccccc}
\hat{\alpha}_{1} & \hat{\beta}_{2} & & \\
\hat{\beta}_{2} & \ddots & \ddots & \\
& \ddots & \ddots & \hat{\beta}_{m} \\
& & \hat{\beta}_{m} & \hat{\alpha}_{m}
\end{array}\right]
\end{gathered}
$$

for $i \in\{1, \ldots, m\}$,

$$
\begin{aligned}
\left\|\delta \hat{v}_{i}\right\|_{2} & \leq \varepsilon_{1} \sigma \\
\hat{\beta}_{i+1}\left|\hat{v}_{i}^{T} \hat{v}_{i+1}\right| & \leq 2 \varepsilon_{0} \sigma \\
\left|\hat{v}_{i+1}^{T} \hat{v}_{i+1}-1\right| & \leq \varepsilon_{0} / 2 \\
\left|\hat{\beta}_{i+1}^{2}+\hat{\alpha}_{i}^{2}+\hat{\beta}_{i}^{2}-\left\|A \hat{v}_{i}\right\|_{2}^{2}\right| & \leq 4 i\left(3 \varepsilon_{0}+\varepsilon_{1}\right) \sigma^{2}
\end{aligned}
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Lanczos [Paige, 1976]

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\begin{aligned}
& \varepsilon_{0}=O(\varepsilon N) \\
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s-step Lanczos [C., Demmel, 2015]:

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\begin{aligned}
& \varepsilon_{0}=O\left(\varepsilon N \Gamma^{2}\right) \\
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$$
\Gamma=c \cdot \max _{\ell}\left\|\hat{\mathcal{Y}}_{\ell}^{+}\right\|\left\|\hat{\mathcal{Y}}_{\ell}\right\|
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## Convergence of Ritz Values in s-step Lanczos

- All results of Paige [1980], e.g., loss of orthogonality $\rightarrow$ eigenvalue convergence, hold for s-step Lanczos as long as

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If $\Gamma \approx 1$ :
s-step Lanczos behaves the same numerically as classical Lanczos


Problem: Diagonal matrix with $n=100$ with evenly spaced eigenvalues between $\lambda_{\text {min }}=0.1$ and $\lambda_{\max }=100$; random starting vector

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## Top plots:

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| :--- | :--- |
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Bottom Plots:

+ True eigenvalues
- Computed Ritz values

Bounds on range of computed Ritz values

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$$
s=12
$$

## Top plots:

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| :---: |
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$$
\Gamma \leq 2 \times 10^{6}
$$




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## A different problem...

$A$ : nos4 from UFSMC,
$b$ : equal components in the eigenbasis
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Need adaptive, problem-dependent approach based on understanding of finite precision behavior!


## Adaptive s-step CG

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- We can approximate an upper bound on this quantity by

$$
\frac{\left\|f_{m+s}-f_{m}\right\|}{\|A\|\|x\|} \lesssim \varepsilon\left(1+\kappa(A) \Gamma_{k} \frac{\max _{j \in\{0, \ldots, s\}}\left\|\hat{r}_{m+j}\right\|}{\|A\|\|x\|}\right) \quad f_{i} \equiv b-A \hat{x}_{i}-\hat{r}_{i}
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- If our application requires relative accuracy $\varepsilon^{*}$, we must have

$$
\Gamma_{k} \equiv c \cdot\left\|\hat{\mathcal{Y}}_{k}^{+}\right\|\left\|\left|\hat{y}_{k}\right|\right\| \lesssim \frac{\varepsilon^{*}}{\varepsilon \max _{j \in\{0, \ldots, s\}}\left\|\hat{r}_{m+j}\right\|}
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\frac{\left\|f_{m+s}-f_{m}\right\|}{\|A\|\|x\|} \lesssim \varepsilon\left(1+\kappa(A) \Gamma_{k} \frac{\max _{j \in\{0, \ldots, s\}}\left\|\hat{r}_{m+j}\right\|}{\|A\|\|x\|}\right) \quad f_{i} \equiv b-A \hat{x}_{i}-\hat{r}_{i}
$$

- If our application requires relative accuracy $\varepsilon^{*}$, we must have

$$
\Gamma_{k} \equiv c \cdot\left\|\hat{y}_{k}^{+}\right\|\left\|\left|\hat{y}_{k}\right|\right\| \lesssim \frac{\varepsilon^{*}}{\varepsilon \max _{j \in\{0, \ldots, s\}}\left(\left\|\hat{r}_{m+j} \mid\right\|\right)}
$$

- $\left\|\hat{r}_{i}\right\|$ large $\rightarrow \Gamma_{k}$ must be small; $\left\|\hat{r}_{i}\right\|$ small $\rightarrow \Gamma_{k}$ can grow


## Adaptive s-step CG

- Consider the growth of the relative residual gap caused by errors in outer loop $k$, which begins with global iteration number $m$
- We can approximate an upper bound on this quantity by

$$
\frac{\left\|f_{m+s}-f_{m}\right\|}{\|A\|\|x\|} \lesssim \varepsilon\left(1+\kappa(A) \Gamma_{k} \frac{\max _{j \in\{0, \ldots, s\}}\left\|\hat{r}_{m+j}\right\|}{\|A\|\|x\|}\right) \quad f_{i} \equiv b-A \hat{x}_{i}-\hat{r}_{i}
$$

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$$

- $\left\|\hat{r}_{i}\right\|$ large $\rightarrow \Gamma_{k}$ must be small; $\left\|\hat{r}_{i}\right\|$ small $\rightarrow \Gamma_{k}$ can grow
$\Rightarrow$ adaptive s-step approach [C., 2018]
- $s$ starts off small, increases at rate depending on $\left\|\hat{r}_{i}\right\|$ and $\varepsilon^{*}$


## Improving Adaptive s-step CG

- Method of Meurant and Tichý (2018) for cheap approximation of extremal Ritz values
- Uses Cholesky factors of Lanczos tridiagonal $T_{i}, T_{i}=L_{i} L_{i}^{T}$
- Use $\alpha$ and $\beta$ computed during each iteration to incrementally update estimates of $\left\|L_{i}\right\|_{2}^{2}=\lambda_{\max }\left(T_{i}\right) \approx \lambda_{\max }(A),\left\|L_{i}^{-1}\right\|_{2}^{-2}=\lambda_{\min }\left(T_{i}\right) \approx$ $\lambda_{\text {min }}(A)$
- Essentially no extra work, no extra communication


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- Essentially no extra work, no extra communication
- Can be used in two ways in adaptive algorithm

1. Incrementally refine estimate of $\kappa(A)$ (used in determining which $s$ to use)
2. Incrementally refine parameters used to construct Newton or Chebyshev polynomials
$A=494$ bus from SuiteSparse $b_{\mathrm{i}}=1 / \sqrt{N}$

$-s$-step CG

- adptv. $s$-step CG - N
- adptv. $s$-step CG - C
_ classical CG

Number of global synchronizations

| Fixed s-step | Improved adaptive s-step <br> w/Newton | Improved adaptive s-step <br> w/Chebyshev | classical CG |
| :---: | :---: | :---: | :---: |
| - | 59 | 53 | 414 |

## Summary

- In order to truly claim that a modified variant of a Krylov subspace method is suitable for HPC/more efficient than the classical approach, we must understand its behavior in finite precision
- In s-step variants of Krylov subspace methods, local roundoff errors are amplified by a factor related to the conditioning of the computed "s-step bases"
- Bounds on maximum attainable accuracy
- Working towards understanding convergence delay
- Understanding finite precision behavior can allow us to develop adaptive approaches that are both accurate and efficient


## Thank you!

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