Argonne Training Program on Extreme-Scale Computing

ATPESC 2023

Krylov Solvers and Algebraic Multigrid with *hypre*

Sarah Osborn, Ulrike Meier Yang
Lawrence Livermore National Laboratory

July 30 – August 11, 2023
Outline

• What are Krylov Solvers?
• Why are they used?
• Why multigrid methods?
• Algebraic multigrid software
• Hypre software library – interfaces
  – Why different interfaces?
• How does multigrid work?
• Unstructured vs structured multigrid solvers
Iterative Solvers

• Solve linear system $Ax = b$, where $A$ is a large sparse matrix of size $n$

• Direct solvers (e.g., Gaussian elimination) too expensive

• Iterative solvers

• Richardson iteration:
  
  \[ x^{n+1} = x^n + (b - Ax^n) \]
  \[ e^{n+1} = (I - A)e^n \]

• Introduce a preconditioner $B$:
  
  \[ x^{n+1} = x^n + B(b - Ax^n) \]
  \[ e^{n+1} = (I - BA)e^n \]

• Jacobi: $B = D^{-1}$; Richardson: $B = \lambda I$
Generalized Minimal Residual (GMRES)

• $x^{n+1} = x^n + B(b - Ax^n)$
• $\Rightarrow x^{n+1} = \sum_{i=0}^{n} \alpha_i (BA)^i Bb$
• $x^{n+1} \in K^n = \text{span}\{Bb, (BA)Bb, (BA)^2 Bb, \ldots, (BA)^n Bb\}$

Krylov space

• Now optimize by defining $x^{n+1}$ through
  $$\min_{x^{n+1} \in K^n} \|B(Ax^{n+1} - b)\|$$

• Construct a new basis for $K^n$ through orthonormalization
  $$\{q_0 = \frac{Bb}{\|Bb\|}, q_1, \ldots, q_n\}$$

• Solve the minimization in the new basis
• $q_i$ also called search directions
Some comments on GMRES

• GMRES consists of fairly simple operations:
  – Inner products and norms (global reductions)
  – Vector updates (embarrassingly parallel)
  – Matvecs (nearest neighbor updates)
  – Application of preconditioner (can be very complicated)

• Often used restarted as GMRES(k), i.e., after k iterations throw out $q_i$ and start again using latest approximation

• Many variants to reduce and/or overlap communication (pipelined GMRES, etc)
Other Krylov solvers

• Conjugate Gradient (CG)
  – For symmetric positive definite matrices
  – Possesses like GMRES an orthogonality property
  – Uses a three-term concurrence
  – Requires only two inner products and a norm per iteration

• BiCGSTAB (Biconjugate Gradient Stabilized)
  – Like CG uses a three-term recurrence relation
  – No orthogonality property, can break down
  – Requires several inner products and a norm at each iteration (and two matvecs)
  – More erratic convergence than GMRES, but needs generally less memory
Hands-on Exercises: Krylov methods (First Set of Runs)

- Go to https://xsdk-project.github.io/MathPackagesTraining2023/lessons/krylov_amg_hypre/

- Poisson equation: \(-\Delta \varphi = \text{RHS}\)

  with Dirichlet boundary conditions \(\varphi = 0\)

- Grid: cube

- Finite difference discretization:
  - Central differences for diffusion term
  - 7-point stencil
Multigrid linear solvers are optimal ($O(N)$ operations), and hence have good scaling potential

- Weak scaling – want constant solution time as problem size grows in proportion to the number of processors
Multigrid (MG) uses a sequence of coarse grids to accelerate the fine grid solution

- Error on the fine grid
- Error approximated on a smaller coarse grid

Multigrid V-cycle

- smoothing (relaxation)
- prolongation (interpolation)
- restriction

Algebraic multigrid (AMG) only uses matrix coefficients
No actual grids!
AMG Building Blocks

Setup Phase:
- Select coarse “grids”
- Define interpolation: $P^m$, $m = 1, 2, ...$
- Define restriction: $R^m$, $m = 1, 2, ..., \text{ often } R^m = (P^m)^T$
- Define coarse-grid operators: $A^{(m+1)} = R^m A^m P^m$

Galerkin product

Solve Phase:

Relax $A^m u^m = f^m$

\begin{align*}
&\text{Compute } r^m = f^m - A^m u^m \\
&\text{Restrict } r^{m+1} = R^m r^m
\end{align*}

Correct $u^m \leftarrow u^m + e^m$

Interpolate $e^m = P^m e^{m+1}$

Solve $A^{(m+1)} e^{m+1} = r^{m+1}$
Multigrid software

- ML, MueLu included in TRILINOS
- GAMG in PETSc
- The hypre library provides various algebraic multigrid solvers, including multigrid solvers for special problems e.g., Maxwell equations, ...
- ...

- All of these provide different flavors of multigrid and provide excellent performance for suitable problems
- Focus here on hypre
The *hype* software library provides structured and unstructured multigrid solvers

- Used in many applications

- Displays excellent weak scaling and parallelization properties on BG/Q type architectures

![Elasticity/Plasticity](image1)

![Electro-magnetics](image2)

![Magneto-hydrodynamics](image3)

![Facial surgery](image4)

![Quantum Chromodynamics](image5)

![Subsurface simulations](image6)

![Graph](image7)
(Conceptual) linear system interfaces are necessary to provide “best” solvers and data layouts.
Why multiple interfaces? The key points

• Provides natural “views” of the linear system

• Eases some of the coding burden for users by eliminating the need to map to rows/columns

• Provides for more efficient (scalable) linear solvers

• Provides for more effective data storage schemes and more efficient computational kernels
hypre supports these system interfaces

• Structured-Grid (Struct)
  – logically rectangular grids

• Semi-Structured-Grid (SStruct)
  – grids that are mostly structured
  – Examples: block-structured grids,
    structured adaptive mesh refinement grids,
    overset grids
  – Finite elements

• Linear-Algebraic (IJ)
  – general sparse linear systems
BoomerAMG is an algebraic multigrid method for unstructured grids

- **Interface**: SStruct, IJ
- **Matrix Class**: ParCSR

- Originally developed as a general matrix method (i.e., assumes given only $A$, $x$, and $b$)
- Various coarsening, interpolation and relaxation schemes
- Automatically coarsens “grids”
- Can solve systems of PDEs if additional information is provided
- Can also be used through PETSc and Trilinos
- Can now also be used on GPUs (CUDA, HIP)
Complexity issues

- Coarse-grid selection in AMG can produce unwanted side effects.
- Operator (RAP) “stencil growth” reduces efficiency.
- For BoomerAMG, we will also consider complexities:
  - Operator complexity: $C_{op} = (\sum_{i=0}^{L} \text{nnz}(A_i))/\text{nnz}(A_0)$
  - Affects flops and memory
  - Generally, would like $C_{op} < 2$, close to 1
- Can control complexities in various ways:
  - Varying strength threshold
  - More aggressive coarsening
  - Operator sparsification (interpolation truncation, non-Galerkin approach)
- Needs to be done carefully to avoid excessive convergence deterioration.
ParCSRMatrix data structure

- Based on compressed sparse row (CSR) data structure
- Consists of two CSR matrices:
  - One containing local coefficients connecting to local column indices
  - The other (Offd) containing coefficients with column indices pointing to off processor rows
- Also contains a mapping between local and global column indices for Offd
- Requires much indirect addressing, integer computations, and computations of relationships between processes etc.,
SMG and PFMG are semicoarsening multigrid methods for structured grids

- Interface: Struct
- Matrix Class: Struct
- SMG uses plane smoothing in 3D, where each plane “solve” is affected by one 2D V-cycle
- SMG is very robust
- PFMG uses simple pointwise smoothing, and is less robust
- Note that stencil growth is limited for SMG and PFMG (to at most 27 points per stencil in 3D)
- Constant-coefficient versions
- Can be used on GPUs (CUDA, HIP, RAJA, Kokkos)
PFMG is an algebraic multigrid method for structured grids

- Matrix defined in terms of grids and stencils
- Uses semicoarsening
- Simple 2-point interpolation → limits stencil growth to at most 9pt (2D), 27pt (3D)
- Optional non-Galerkin approach (Ashby, Falgout), uses geometric knowledge, preserves stencil size
- Pointwise smoothing
- Highly efficient for suitable problems
Structured-Grid System Interface (Struct)

• Appropriate for scalar applications on structured grids with a fixed stencil pattern

• Grids are described via a global \(d\)-dimensional index space (singles in 1D, tuples in 2D, and triples in 3D)

• A box is a collection of cell-centered indices, described by its “lower” and “upper” corners

• The grid is a collection of boxes

• Matrix coefficients are defined via stencils

\[
\begin{bmatrix}
    \text{s}_4 \\
    \text{s}_1 \\
    \text{s}_0 \\
    \text{s}_2 \\
    \text{s}_3
\end{bmatrix}
= \begin{bmatrix}
    -1 & 4 & -1 \\
    -1 & & \end{bmatrix}
\]
StructMatrix data structure

- Stencil
  \[
  \begin{bmatrix}
  S4 \\
  S1 \\
  S0 \\
  S2 \\
  S3 \\
  \end{bmatrix}
  =
  \begin{bmatrix}
  -1 & 4 & -1 \\
  \end{bmatrix}
  \]

- Grid boxes: [(-3,1), (-1,2)] [(0,1), (2,4)]

- Data Space: grid boxes + ghost layers:
  [(-4,0), (0,3)] , [(-1,0), (3,5)]

- Data stored

- Operations applied to stencil entries per box (corresponds to matrix (off) diagonals from a matrix point of view)
Algebraic multigrid as preconditioner

- Generally algebraic multigrid methods are used as preconditioners to Krylov methods, such as conjugate gradient (CG) or GMRES.
- This often leads to additional performance improvements.

Classic porous media diffusion problem:

\[-\nabla \cdot \kappa \nabla u = f\]

with \(\kappa\) having jumps of 2-3 orders of magnitude.

Weak scaling: 32x32x32 grid points per core, BG/Q.
Hands-on Exercises: Algebraic multigrid (Second Set of Runs)

• Go to https://xsdk-project.github.io/MathPackagesTraining2023/lessons/krylov_amg_hypre/

• Poisson equation: \(-\Delta \varphi = \text{RHS}\)

  with Dirichlet boundary conditions \(\varphi = 0\)

• Grid: cube

• Finite difference discretization:
  – Central differences for diffusion term
  – 7-point stencil
Porting to GPUs required inclusion of new programming models and different strategies for structured/unstructured interfaces

• Strategy for structured interface and solvers
  – Include new programming models (CUDA, HIP, RAJA, Kokkos, OMP, and SYCL) in hypre_BoxLoops (macros that operate on data in loops).

• Strategy for unstructured interface and solvers CSR-based matrix data structures
  – Modularize into smaller chunks/kernels to be ported to CUDA for Nvidia GPUs initially
  – Convert CUDA kernels to HIP for AMD GPUs and SYCL for Intel GPUs
  – Develop new algorithms for portions not suitable for GPUs (interpolation operators, smoothers)
    → different defaults for CPU and GPU use
  – Various special solvers (e.g., Maxwell solver AMS, ADS, AME, pAIR, MGR) built on BoomerAMG benefit from this strategy
Structured multigrid methods perform significantly better than unstructured ones on CPUs and - even more - on GPUs

ThetaGPU
GPU: 1 Nvidia A100
CPU: 16 MPI tasks

Used optimal settings for AMG, which are different for CPU and GPU!

Speedups at n=200

<table>
<thead>
<tr>
<th>Speedup</th>
<th>GPU/CPU</th>
<th>PFMG/AMG</th>
</tr>
</thead>
<tbody>
<tr>
<td>CPU Speedup</td>
<td>13.2</td>
<td>1.7</td>
</tr>
<tr>
<td>GPU Speedup</td>
<td>28.5</td>
<td>3.8</td>
</tr>
</tbody>
</table>
Most gains of PFMG over AMG in setup phase
Successfully used hypre on Frontier (AMD GPUs) for solving complex multiphysics simulations

Single-phase flow (Poisson-like problem)

- Weak scaling with BoomerAMG/GMRES(50)
- Time complexity \( \sim O(\log(N)) \); Iteration counts \( \sim O(1) \).
Frontier (AMD GPUs) results - Solved system with 80B DOFs using less than 25% of the machine

Mechanics (linear elasticity problem)

- Weak scaling with BoomerAMG/GMRES(40)
Hands-on Exercises: Comparing GPU to CPU Performance

Algebraic Multigrid methods (Third Set of Runs)

• Go to https://xsdk-project.github.io/MathPackagesTraining2023/lessons/krylov_amg_hypre/

• Poisson equation: \(-\Delta \varphi = \text{RHS}\)

  with Dirichlet boundary conditions \(\varphi = 0\)

• Grid: cube

• Finite difference discretization:
  – Central differences for diffusion term
  – 7-point stencil
Thank you!
This work was performed under the auspices of the U.S. Department of Energy by Lawrence Livermore National Laboratory under contract DE-AC52-07NA27344. Lawrence Livermore National Security, LLC.

This work was supported by the U.S. Department of Energy Office of Science, Office of Advanced Scientific Computing Research (ASCR), Scientific Discovery through Advanced Computing (SciDAC) program, and by the Exascale Computing Project, a collaborative effort of the U.S. Department of Energy Office of Science and the National Nuclear Security Administration.

Disclaimer
This document was prepared as an account of work sponsored by an agency of the United States government. Neither the United States government nor Lawrence Livermore National Security, LLC, nor any of their employees makes any warranty, expressed or implied, or assumes any legal liability or responsibility for the accuracy, completeness, or usefulness of any information, apparatus, product, or process disclosed, or represents that its use would not infringe privately owned rights. Reference herein to any specific commercial product, process, or service by trade name, trademark, manufacturer, or otherwise does not necessarily constitute or imply its endorsement, recommendation, or favoring by the United States government or Lawrence Livermore National Security, LLC. The views and opinions of authors expressed herein do not necessarily state or reflect those of the United States government or Lawrence Livermore National Security, LLC, and shall not be used for advertising or product endorsement purposes.