

CAS 741 (Development of Scientific Computing Software)

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Mixed-Precision Iterative Solver

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Linear Solver: the Good Old $\mathbf{Ax} = \mathbf{b}$ Problem

- Direct method: Gaussian elimination

$$\mathbf{Ax} = \begin{bmatrix} 1 & -1 & 3 \\ 1 & 1 & 0 \\ 3 & -2 & 1 \end{bmatrix} \begin{bmatrix} x_1 \\ x_2 \\ x_3 \end{bmatrix} = \begin{bmatrix} 11 \\ 3 \\ 3 \end{bmatrix} = \mathbf{b}$$

$$\mathbf{A}|\mathbf{b} = \left[\begin{array}{ccc|c} 1 & -1 & 3 & 11 \\ 1 & 1 & 0 & 3 \\ 3 & -2 & 1 & 3 \end{array} \right] \begin{array}{cc} \times 1 & \times 3 \\ \downarrow & \\ & \downarrow \end{array}$$

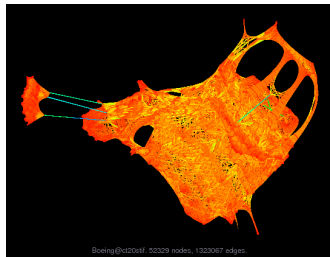
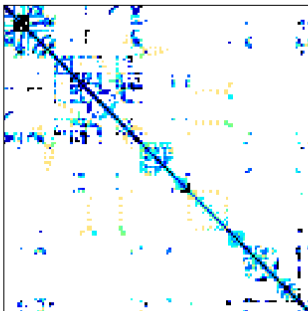
$$\mathbf{A}|\mathbf{b} \leftarrow \left[\begin{array}{ccc|c} 1 & -1 & 3 & 11 \\ 0 & 2 & -3 & -8 \\ 0 & 1 & -8 & -30 \end{array} \right]$$

Think Big, Think Sparse

- In structural simulations (e.g., finite element analysis), structures are modeled based on real-world physics, where forces and constraints are often localized.
- Each element or node in a structure is typically connected only to a few nearby elements or nodes.
- System of equations in structural simulations is assembled from local contributions.
- In contrast, in neural networks, dense weight matrices are used in fully connected layers.

Sparse Matrix Example: Boeing/ct20stif

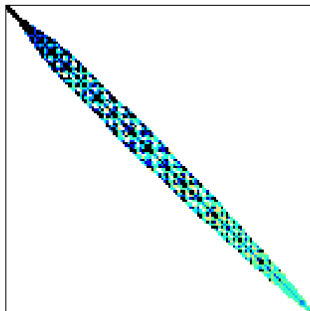
- $52\,329 \times 52\,329$
- 2 600 295 non-zeros
- ≈ 33 MB in file size



Boeing/ct20stif: CT20 Engine Block – Stiffness matrix

Sparse Matrix Example: Janna/Serena

- $1\,391\,349 \times 1\,391\,349$
- 64 131 971 non-zeros
- ≈ 847 MB in file size



Janna/Serena: gas reservoir simulation for CO₂ sequestration

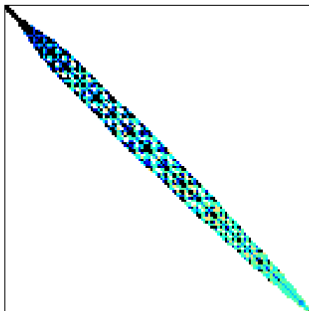
The Problem with Gaussian Elimination

- Store the whole matrix
- Memory bounded

$$\mathbf{A}|\mathbf{b} = \left[\begin{array}{ccc|c} 1 & -1 & 3 & 11 \\ 1 & 1 & 0 & 3 \\ 3 & -2 & 1 & 3 \end{array} \right] \begin{array}{cc} \times 1 & \times 3 \\ \downarrow & \\ & \downarrow \end{array}$$
$$\mathbf{A}|\mathbf{b} \leftarrow \left[\begin{array}{ccc|c} 1 & -1 & 3 & 11 \\ 0 & 2 & -3 & -8 \\ 0 & 1 & -8 & -30 \end{array} \right]$$

The Problem with Gaussian Elimination

- Sparsity preservation



Janna/Serena: gas resevoir simulation for CO₂ sequestration

Iterative Refinement

- A method to enhance the accuracy of a solution to $\mathbf{Ax} = \mathbf{b}$

The algorithm goes:

1. Solve $\mathbf{Ax}_0 = \mathbf{b}$ *approximately* to get an initial solution \mathbf{x}_0 .
2. Compute the residual $\mathbf{r} = \mathbf{b} - \mathbf{Ax}_0$, which measures how far \mathbf{x}_0 is from being an exact solution.
3. Solve $\mathbf{Ae} = \mathbf{r}$ for the correction vector \mathbf{e} .
4. Update (“refine”) the solution as $\mathbf{x}_1 = \mathbf{x}_0 + \mathbf{e}$.
5. Repeat steps 2–4 until the residual \mathbf{r} is small enough.

Iterative Refinement in Mixed-Precision

1. Solve $\mathbf{Ax}_0 = \mathbf{b}$ *approximately* in **low** precision.
2. Compute the residual $\mathbf{r} = \mathbf{b} - \mathbf{Ax}_0$ in **high** precision.
3. Solve $\mathbf{Ae} = \mathbf{r}$ for the correction vector \mathbf{e} in **low** precision.
4. Update the solution as $\mathbf{x}_1 = \mathbf{x}_0 + \mathbf{e}$ in **medium** precision.
5. Repeat steps 2–4 until the residual \mathbf{r} is small enough.

Key points:

- Trade computational complexity for space complexity
- Choose algorithms that can preserve sparsity: sparse LU factorization, General Minimal Residual Method (GMRES)
- Inner solves could also be iterative and in mixed-precision
- Possible matrix-free implementation

Goals

- GS1 Given some matrix \mathbf{A} and column vector \mathbf{b} , the solver should iteratively find \mathbf{x} satisfying the equation $\mathbf{Ax} = \mathbf{b}$ until the norm of the residual $\mathbf{r} = \mathbf{Ax} - \mathbf{b}$ is smaller than some tolerance ϵ , or the maximum number of iterations n_{iter} is exhausted, whichever comes first.
- GS2 Given some combinations of floating point precision configuration, the solver should perform internal steps such as matrix factorizations, triangular solves, and residual computation in these configured precisions.

Goals (Non-functional)

- GS3 The solver should offer a quantifiable performance or resource utilization advantage over other competing sparse linear solvers.
- GS4 The library should offer a set of streamlined public application programming interfaces (APIs), such that when integrated into other software as a dependent library, the interfaces are self-contained, readable and easy to consume.

Stretch Goals

- GS5 With the existing solver implementation being the baseline, the refactored solver should produce more accurate results, lowering the norm of the residual by at least 1 order of magnitude.
- GS6 The solver should optimize existing algorithms such that given the same set of inputs, it produces results with the same accuracy in notably less time.

Inputs

Variable	Description
A	$n \times n$ matrix
b	n -vector
ϵ	a solution is found if the norm of the residual is less than ϵ
n_{iter}	the maximum number of iterations to perform
u_f	factorization precision
u_w	working precision
u_r	precision in which the residuals are computed

Outputs

Variable	Description
\mathbf{x}	a numerical solution to the linear system
err	the norm of the residual

Assumptions and Constraints

- A1 Matrix \mathbf{A} is quasi-definite. (LC1)
- A2 The precisions follow the order $u_f \leq u_w \leq u_r$, with u_r being the highest precision.
- C1 Use preconditioned GMRES for solving the error correction vector.

Questions